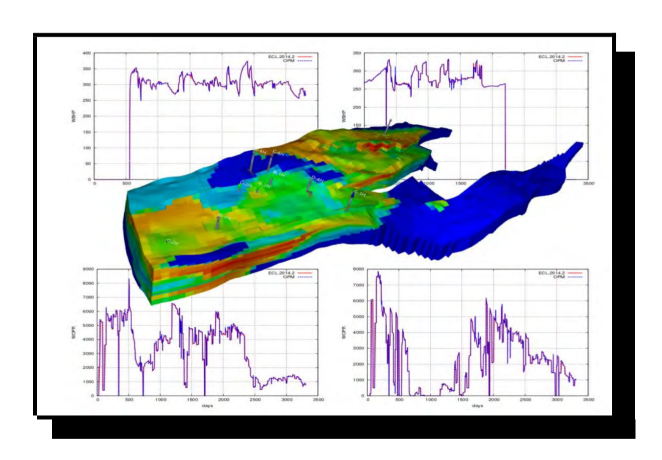
OPEN POROUS MEDIA

Flow Documentation Manual



OPM FLOW VERSION: 2018-10 MANUAL REVISION: Rev-1

OPEN POROUS MEDIA

FLOW DOCUMENTATION MANUAL (2018-10)

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1 INTRODUCTION

OPM Flow is a fully-implicit, black-oil simulator capable of running industry-standard simulation models. The simulator is implemented using automatic differentiation to enable rapid development of new fluid models.

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Model Formulation:

- · Back-oil with dissolved gas and vaporized oil.
- Rock-dependent capillary and relative-permeability curves.
- · End-point scaling and hysteresis.
- Oil vaporization controls (VAPPARS).

EOR Options:

- The Polymer Model in OPM Flow is based on a black-oil polymer formulation, which is developed by extending the black-oil model with a polymer component. The effects of the polymer mixing are simulated based on the Todd-Longstaff mixing model, and adsorption, dead pore space, and permeability reduction effects are also considered. A logarithmic shear thinning/thickening model has also been incorporated since the 2015-10 Release (see Flow-polymer). Note that Polymer model has now been incorporated into the main OPM Flow simulator and is no longer a separte simulator.
- The Solvent Model in OPM Flow extra component is again based on extending the black-oil oil formulation with a fourth component the simulator by adding a solvent component to the gas phase (see *Flow-solvent*). Note that Solvent model has now been incorporated into the main OPM Flow simulator and is no longer a separate simulator.

Description of Geology:

- Rectilinear and fully-unstructured grid.
- · Corner-point grids from Eclipse input, including fault and region multipliers, minpv/pinch, etc.

Well and Group Controls:

- Bottom-hole pressure and surface/reservoir rate.
- Group controls.
- Shut/stop/open individual completions.
- History-matching wells.

Input and Output:

- General reader/parser for Eclipse input decks.
- XML-based or simple text-format input of additional parameters.
- Flexible output of summary and restart files in Eclipse format.
- · logging to terminal and print file.

Simulation Technology:

- Fully-implicit in time.
- Two-point flux approximation in space with upstream-mobility weighting.
- Flexible assembly through the use of automatic differentiation.
- Block-structured linear solver with ILU0 pre-conditioner.
- Adaptive step-size controls.

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2 INSTALLING AND RUNNING FLOW

2.1 Installing Flow

OPM Flow can be installed and used in a variety of ways. Perhaps the simplest way is to use the operating system's package manager to install the OPM modules as binary packages, including OPM Flow and other OPM programs. That way all prerequisite libraries will automatically be installed on the system. This is supported on Ubuntu Linux version 16.04 and Red Hat Enterprise (or CentOS) version 6 or 7.

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OPM Flow can also be used via a Docker container. This allows OPM Flow to run on any system that supports Docker, including Microsoft Windows, without installing prerequisite libraries or affecting any other software on your system.

Finally, OPM Flow can be installed by compiling from source on Linux or macOS systems.

2.1.1 UBUNTU LINUX 16.04 (64-BIT VERSION ONLY) AND HIGHER

The easiest way to install the Ubuntu packages is to first add the OPM personal package archive (ppa). In order to do that we need to install the apt-add-repository command.

```
sudo apt-get update
sudo apt-get install software-properties-common
```

Then we add the repository, and run update again:

```
sudo apt-add-repository ppa:opm/ppa
sudo apt-get update
```

At this point, all the OPM modules should be available to install. To see a list of (for example) the opm-simulators packages:

```
apt-cache search opm-simulators
```

Then, to install the opm-simulators programs (including Flow) and their dependencies,

```
sudo apt-get install mpi-default-bin
sudo apt-get install libopm-simulators-bin
```

Notes

- The mpi install above is required for the mpi libraries which the OPM executables are linked to, even those that are not intended to be run with mpirun. The reason the OPM packages do not depend on that package is that by Debian policy the user should be allowed to choose which MPI implementation to use (above we picked the default set by Ubuntu).
- 2) If an old versions of prerequisite libraries have already been installed (for example from installing a previous release of OPM) it may be necessary to upgrade them via the following commands:

```
sudo apt-get update
sudo apt-get upgrade
```

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3) Some users have experienced trouble with OPM Flow simply aborting with no error message. In some cases that can be related to locale issues. Check by running the "locale" command. OPM Flow requires the "C" or an English locale to run correctly. One way to fix this can be to put the following in your ".bash_profile" (note the initial period or full stop in the filename), and open a new terminal (command) window:

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```
LANG="en_US.UTF-8"
export LANG
LC_ALL="en_US.UTF-8"
export LC_ALL
```

2.1.2 RED HAT ENTERPRISE OR CENTOS (VERSION 6 OR 7)

First add the OPM package repository:

```
sudo yum-config-manager --add-repo \
http://www.opm-project.org/package/opm.repo
```

The OPM software is split in several packages. To list all available OPM packages one can use:

```
sudo yum search opm-
```

For example you can install the opm-simulators binary package to get access to the OPM Flow reservoir simulator:

```
sudo yum install opm-simulators-bin
```

Development packages are available using the -devel suffix. For example, to install the opm-upscaling development package use:

```
sudo yum install opm-upscaling-devel
```

2.1.3 Using a Docker container

See the tutorial on the OPM website Running Flow in Docker.

2.1.4 Installing from source

See instructions on the OPM website Building from Source.

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2.2 RUNNING FLOW

If OPM Flow is installed on your system typing the following command:

flow [OPTIONS] [CASENAME]

from your terminal will start the simulator and run the case specified by CASENAME.DATA.

Alternatively, you can type the path to the binary and the .DATA file, as per the example below:

path_to_binary/flow path_to_data/CASENAME

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By default output files are generated in the same folder as the .DATA file.

The user can supply various options to the simulator. For instance for the 2018-04 release:

flow output_dir=foo CASENAME

or for the 2018-10 release the command:

flow --ecl-output-dir=foo CASENAME

will send the output files to the foo directory.

Note that there must be no spaces around the equals sign. A list of command line options is given in the following sections for various releases of OPM Flow.

If OPM Flow is installed with parallel capabilities.

mpirun -np 4 flow CASENAME

will start Flow on four nodes etc.

Note

On Red Hat based distributions, including centOS distributions, one needs to install the openmpi version of the simulator, opm-simulators-openmpi-bin, via:

sudo yum install opm-simulators-openmpi-bin

and then set the MPI version to use. The way this is done is by first making the module command available, by running the following command:

. /usr/share/Modules/init/bash

Then query for the installed modules with:

module avail

And lastly, you tell the system it to use openmpi with:

module add mpi/openmpi-x86_64

On Ubuntu based distributions, there is no need for this as openmpi is installed and mpirun just works.

Additional tutorials for running OPM Flow is available on OPM website in the *Tutorials* section.

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2.2.1 COMMAND LINE OPTIONS FOR OPM FLOW 2018-04

OPM Flow accepts command line options to control various aspects of the simulator's run time behavior, as described in the table below. To give a command line option use "key=value" syntax, with no spaces around the equals sign. It is also possible to put multiple options together in a parameter file. To do so, put one option on each line of the file with ".param" as the extension and pass that filename as a command line parameter to OPM Flow.

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This section describes the command line options up to the 2018-04 release of OPM Flow, post this release the command line options were changed to be the same as eWoms/ebos¹ command line parameters, as described in section 2.2.2 Command Line Options for OPM Flow 2018-10. It is anticipated that this section will be removed from the manual once the 2018-10 and later versions are firmly established.

OPM Flow 2018-04 Command Line Options			
No.	Variable Name	Description	Default
I	output_ecl	A Boolean value set to true or false that turns on (true) or off (false) output to the .*RST and .*SMRY files.	true
2	output	A character string that defines the output to *.PRT and *.DEBUG files:	all
		0) none: No output to the files.	
		log or false: Output logging information only.	
		2) all or true: Output everything.	
		For example to just output logging information use: output=log or output=false	
3	output_dir	Set the directory to which output files are written.	deck location
4	restart_double_si	A Boolean value set to true or false that switches on (true) or off (false) double precision in restart files and stores all restart data in SI units rather than using the unit family (METRIC, FIELD etc.) used in the input deck.	false
		The option improves the quality of the restart.	
5	async_output	A Boolean value set to true or false that sets the output mode to be asynchronously (true), letting the simulator continue computing the next time step while writing results to restart and summary files.	true
6	newton_use_gmres	A Boolean value set to true or false that when set to true OPM Flow will use GMRes instead of BiCGStab as linear solver within the Newton iterations.	false
7	linear_solver_reduction	A real positive double precision value that sets the tolerance for the linear solver. The linear solver convergences when the residual is reduced sufficiently.	0.01

eWorms is a C++ software framework concerned with fully-implicit numerical models for flow and transport in porous media. It it is developed as an integral part of the Open Porous Media (OPM) initiative.

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	OPM Flow 2018-04 Command Line Options				
No.	Variable Name	Description	Default		
8	linear_solver_maxiter	A positive integer value that defines the maximum number of linear iterations.	150		
9	linear_solver_verbosity	A positive integer value that defines the output from linear solver: 1) 0: no extra output 2) 1: output per solution iteration 3) >1: output per iteration	0		
10	linear_solver_ignoreconvergencefailure	A Boolean value set to true or false that if set to true convergences failures in the linear solver are ignored. This option should be used with care, as the results may be unreliable.	false		
11	linear_solver_use_amg	A Boolean value set to true or false that if set to true OPM Flow will use amg as the pre-conditioner instead of ilu.	false		
12	ilu_relaxation	A real positive double precision value that sets the relaxation parameter for the ILU pre-conditioner.	0.9		
13	ilu_fillin_level	A positive integer value that sets the fill in level for the ILU pre-conditioner.	0		
14	dp_max_rel	A real positive double precision value that sets the maximum allowed relative change of pressure per iteration.	0.3		
15	ds_max	A real positive double precision value that sets the maximum allowed change in saturation per iteration.	0.2		
16	dr_max_rel	A real positive double precision value that sets the maximum allowed relative change in dissolved gas and vaporized oil per iteration	le+09		
17	dbhp_max_rel	A real positive double precision value that sets the maximum allowed relative change in BHP per iteration.	I		
18	dwell_fraction_max	A real positive double precision value that sets the maximum allowed change in well's volume per iteration.	le+07		
19	tolerance_mb	A real positive double precision value that sets the maximum mass balance error.	le-05		
20	tolerance_cnv	A real positive double precision value that specifies the maximum non-linear tolerance error.	0.01		
21	tolerance_wells	A real positive double precision value that defines the maximum non-linear error for the well equations.	0.0001		
22	max_strict_iter	A positive integer value that sets the maximum number of non-linear iterations. After this maximum value has been exceeded only the mass balance error is checked.	8		

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	OPM Flow 2018-04 Command Line Options					
No.	Variable Name	Description	Default			
23	solve_welleq_initially	A Boolean value set to true or false that switches on (true) or off (false) the solving of the well equations as a pre-processing step.	true			
		Note that the well equations are always added to the full system and solved until converged.				
24	use_update_stabilization	A Boolean value set to true or false that switches on (true) or off (false) the stabilized Newton option.	true			
		This option may improve convergence for some cases.				
25	relax_max	A real positive double precision value that is used to tune the stabilized Newton option.	0.5			
26	relax_type	A character string that sets relaxation type of the stabilized Newton option.	dampen			
27	max_iter	A positive integer that sets the maximum number of non-linear iterations.	10			
28	min_iter	A positive integer that sets the minimum number of non-linear iterations.	I			
29	output_terminal	A Boolean value set to true or false that turns on (true) or off (false) output to terminal.	true			
30	use_TUNING	A Boolean value set to true or false that that instructs OPM Flow to read the time stepping parameters from the <i>TUNING - Numerical Tuning Control</i> keyword in the input deck, if set to true.	false			
		Note that only the first record of the TUNING keyword is processed.				
31	timestep.adaptive	A Boolean value set to true or false that that turns on (true) or off (false) adaptive time stepping. If set to false the report time steps provided by the input deck are used.	true			
32	solver.restartfactor	A real positive double precision value that sets the time step chop factor of the time step after convergence failure.	0.33			
		For example, if the current non-convergent time step is 30 days and <i>solver.restartfactor</i> is set to the default value, then the time step will be repeated using 0.33 * 30 days as the time step, that is 9.9 days.				
33	timestep.control.maxgrowth	A real positive double precision value that sets the maximum allowed value a time step can be increased by, subject to the maximum allowable time step size set by the timestep.max_timestep_in_days parameter.	3			
		For example, if the current time step has converged at 10 days and solver.restartfactor is set to the default value, then the next time step will be 3.0 * 10 days, that is at 30 days.				
34	timestep.max_timestep_in_days	A real positive double precision value that sets the maximum allowed time step size in days.	365			

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	OPM Flow 2018-04 Command Line Options				
No.	Variable Name	Description	Default		
35	solver.restart	A positive integer that defines the number of allowed consecutive restarts (or time step chops) before the simulation is terminated.	10		
36	solver.verbose	A Boolean value set to true or false that switches on (true) or off (false) solver specific output.	true		
37	timestep.verbose	A Boolean value set to true or false that switches on (true) or off (false) time step specific output.	true		
38	timestep_in_days	A real double precision value that sets the size of initial time step in days. The default value of -I sets the initial time step to	-1		
		be solver.restartfactor * the length of the first report step.			
39	full_timestep_initially	Try to use the report steps as time steps.	false		
40	timestep.timestep_in_days_after_event	A real double precision value that sets maximum allowed time step after an event; for example, when a well is open or closed etc.	-1		
		The default value of -I means that events to do effect the time stepping.			
41	timestep.control	A character string that defines the time stepping control algorithm and is set to one of the following:	pid		
		pid: PID controller based adaptive time step control as suggested by Turek and Kuzmin ² .			
		 pid+iteration: Use PID and linear iteration numbers to guide the time step. 			
		 pid+newtoniteration: Use PID and non- linear iterations numbers to guide the time step. 			
		Hardcoded: Use time steps supplied by user. Via timestep.control.filename			
42	timestep.control.tol	A real double precision value that sets the tolerance for PID (only used with the pid and pid+options defined by the <i>timestep.control</i> option).	0.1		
43	timestep.control.targetiteration	A positive integer that sets the target number of linear / non-linear iterations. This option can only be only used by pid+iterations and pid+newtoniteration defined by the <i>timestep.control</i> option.	8		

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Algebraic Flux Correction III. Incompressible Flow Problems. Uni Dortmund, Turek and Kuzmin, January, 2006 (DOI: 10.1007/3-540-27206-2 8)

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No.	Variable Name	Description	Default
44	timestep.control.filename	A character string that specifies a file name where time steps are specified. For instance generated by the ecl_summary application in libecl as per the following UNIX command line:	
		<pre>path_to_libecl_applications/ ecl_summary DECK TIME > filename</pre>	
		Where:	
		DECK is the name of the data deck you want to get the time steps from,	
		TIME tells the application to return the timing for the run, and	
		"filename" is the name of the file the times are piped to.	

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Table 2.1: OPM Flow 2018-04 Command Line Options

As mentioned previously, it is also possible to put multiple options together in a parameter file, by using a filename ending with the extension ".param" and then passing that filename as a command line parameter to OPM Flow.

Note

As the run parameters will effect the results and run times of a simulation it may be worth considering having a parameter file for each run of the form CASENAME.param. This can be then be used to audit any runs when evaluating the results and run times.

The following example is taken for the Norne project and the comments in the example (preceded with "//") explain the parameter setting used in the file.

Example

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```
// Set Min Newtonian Solver iterations to 1 and Max to 15
min_iter=1
max_iter=15
// Set Residual Threshold for Time Step Chop (Restart Solver)
//
max_residual_allowed=1e5
//
// USE BiCG Solver
newton_use_gmres=false
// Set Linear Solver Parameters
linear_solver_reduction=0.01
linear_solver_maxiter=50
linear_solver_restart=40
// ------
// TIME STEPPING PARAMETERS
//
// Set Time Stepping Scheme Option to Adaptive and Control Scheme
timestep.adaptive=true
timestep.control=pid+iteration
// PID Control Tolerance (default = 1e-3)
//
timestep.control.tol=4e-5
// Set Target iteration that is the Sum of all Linear Iterations Over All
// Newton Iterations per Time Step
timestep.control.targetiteration=8
// Set the Minimum Allowed Value a Time Step Can be Decreased
// After the Solver Failes to Converge
//
solver.restartfactor=0.3
// Set the Maximum Allowed Value a Time Step Can be Increased by,
// Subject to the Maximum Allowable Time Step Size
timestep.control.maxgrowth=1.6
// Max Number of Restarts (Time Step Chops) Before the Simulation is Terminated
solver.restart=10
//
// Verbosity of Solver and Adaptive Time Stepping
solver.verbose=true
timestep.verbose=true
```

2.2.2 COMMAND LINE OPTIONS FOR OPM FLOW 2018-10

OPM Flow release 2018-10 and beyond have switched to the eWoms/ebos³ command line interface. Previously the simulator used only OPM Flow specific command line parameters which were internally translated to the equivalent eWoms/ebos command line parameters. The pre 2018-10 set of command line parameters are now depreciated but are still documented in section 2.2.1 Command Line Options for OPM Flow 2018-04 for backward compatibility with previous releases of the simulator. For the 2018-10 release a combination of OPM Flow and eWoms/ebos command line parameters are available as tabulated in Table 2.2 on this and subsequent pages. It is anticipated that future releases of OPM Flow will fully migrate to the eWoms/ebos command line parameter set.

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OPM Flow accepts command line options to control various aspects of the simulator's run time behavior, as described in the table below. To give a command line option use "key=value" syntax, with no spaces around the equals sign. It is also possible to put multiple options together in a parameter file. To do so, put one option on each line of the file with ".param" as the extension and pass that filename as a command line parameter to OPM Flow using:

flow --parameter-file=CASENAME.PARAM

to instruct OPM Flow to read the parameter file.

	OPM Flow 2018-10 Command Line Options			
No.	Variable Name	Description	Default	
	General eWor	ms/ebos Command Line Parameters		
I	-h orhelp	A character string that causes OPM Flow to print a help message that gives a brief description of the available command line parameters.	N/A	
2	dbph-max-rel	A real positive value that sets maximum relative change of the bottom-hole pressure in a single iteration	1.0	
3	dp-max-rel	A real positive double precision value that sets the maximum allowed relative change of pressure per iteration.	0.3	
4	ds-max	A real positive double precision value that sets the maximum allowed change in saturation per iteration.	0.2	
5	dwell-fraction-max	A real positive double precision value that sets the maximum allowed change in well's volume fraction per iteration.	0.2	
6	ecl-deck-file-name	A character string that defines the name of the OPM Flow input file which contains the simulator's ECLIPSE formatted input deck to be simulated.	N/A	
7	ecl-output-double-precision	A Boolean value set to true or false that switches on (true) or off (false) double precision in restart files. Useful for 'perfect' restarts.	false	
8	ecl-output-interval	An integer value defining the number of report steps that ought to be skipped between two writes of restart files results.	I	

³ eWorms is a C++ software framework concerned with fully-implicit numerical models for flow and transport in porous media. It it is developed as an integral part of the Open Porous Media (OPM) initiative.

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	OPM Flow	2018-10 Command Line Options	
No.	Variable Name	Description	Default
9	enable-adaptive-time-stepping	A Boolean value set to true or false that that turns on (true) or off (false) adaptive time stepping. If set to false the report time steps provided by the input deck are used.	true
10	enable-async-ecl-output	A Boolean value set to true or false that sets the output mode to be asynchronously (true), letting the simulator continue computing the next time step while writing results to restart and summary files.	true
11	enable-async-vtk-output	A Boolean value set to true or false that sets the output mode to be asynchronously (true), letting the simulator continue computing the next time step while writing results to the VTK output files.	true
12	enable-dry-run	A Boolean value set to true or false that specifies if the simulation should actually run, (true) or just check the input deck (false).	true
		This option is equivalent to activating the NOSIM keyword in the RUNSPEC section of the input deck (see section 5.2.35 NOSIM – Activates the No Simulation Mode for Data File Checking).	
13	enable-ecl-output	A Boolean value set to true or false to write the binary output which is compatible with the ECLIPSE commercial simulator (restart and summary files).	true
14	enable-opm-rst-file	A Boolean value set to true or false to write OPM specific data sets to the commercial simulators restart file to enable restart of an OPM Flow run (true), or not to write the data (false).	true
15	enable-storage-cache	A Boolean value set to true or false that turns on (true) or off (false) storing previous storage terms and avoid re-calculating them.	true
16	enable-terminal-output	A Boolean value set to true or false that turns on (true) or off (false) high-level information about the simulation's progress to the terminal	true
17	enable-tuning	A Boolean value set to true or false that that instructs OPM Flow to read the time stepping parameters from the <i>TUNING - Numerical Tuning Control</i> keyword in the input deck, if set to true. Note that only the first record of the TUNING	false
		keyword is processed.	
18	enable-vtk-output	A Boolean value set to true or false that turns on (true) or off (false) a global switch for writing VTK files.	false
19	enable-write-all-solutions	A Boolean value set to true or false that turns on (true) or off (false) the writing of all solutions to disk instead of only the ones for the report steps.	false

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NI.	OPM Flow 2018-10 Command Line Options				
No.	Variable Name	Description	Default		
	OPM Flow Spe	cific Command Line Parameters			
20	flow-linear-solver-verbosity	A positive integer value that defines the output from linear solver:	0		
		I) 0: no extra output			
		2) I: output per solution iteration			
		3) >1: output per iteration			
21	flow-newton-max-iterations	A positive integer that defines the maximum number of Newton iterations per time step used by the simulator.	20		
22	flow-newton-min-iterations	A real positive value that sets the minimum number of Newton iterations per time step used by the simulator.	I		
	General eWom	s/ebos Command Line Parameters			
23	force-disable-fluid-in-place-output	A Boolean value set to true or false that instructs OPM Flow to not to print the Fluid In-Place report after each report time step (true) or not (false).	false		
24	full-time-step-initially	A Boolean value set to true or false that instructs OPM Flow to always attempt to finish a report step using a single time step (true) or not (false).	false		
25	ilu-fillin-level	A positive integer value that sets the fill in level for the ILU pre-conditioner.	0		
26	ilu-redblack	A Boolean value set to true or false that instructs OPM Flow to use red-black partitioning for the ILU pre-conditioner. (true) or not (false).	false		
27	ilu-relaxation	A real positive double precision value that sets the relaxation factor of the linear solver's ILU preconditioner	0.9		
28	ilu-reorder-spheres	A Boolean value set to true or false specifies OPM Flow to reorder the entries of the matrix in the red-black ILU pre-conditioner in spheres starting at an edge (true) or not (false)	false		
		If false the original ordering is preserved in each color. Otherwise try to ensure D4 ordering (in a 2D structured grid, the diagonal elements are consecutive).			
29	initial-time-step-in-days	A real double precision value that sets the size of initial time step in days.	1.0		
30	linear-solver-ignore-convergence- failure	A Boolean value set to true or false that if set to true convergences failures in the linear solver are ignored.	false		
		This option should be used with care, as the results may be unreliable.			
31	linear-solver-max-iter	A positive integer value that defines the maximum number of linear iterations.	200		
32	linear-solver-reduction	A real positive double precision value that sets the tolerance for the linear solver. The linear solver convergences when the residual is reduced sufficiently.	0.01		

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	OPM Flow 2018-10 Command Line Options			
No.	Variable Name	Description	Default	
33	linear-solver-require-full-sparsity- pattern	A Boolean value set to true or false that if set to true the simulator will produce the full sparsity pattern for the linear solver.	false	
34	linear-solver-restart	A positive integer value that sets the number of iterations after which GMRES is restarted.	40	
35	matrix-add-well-contributions	A Boolean value set to true or false that if set to true explicitly specifies the influences of wells between cells in the Jacobian and pre-conditioner matrices.	false	
36	max-inner-iter-ms-wells	A positive integer value that defines the maximum number of inner iterations for multi-segment wells.	10	
37	max-pressure-change-ms-wells	A real positive value that defines the maximum relative pressure change for a single iteration of the multisegment well model.	200000	
38	max-residual-allowed	A real positive value that sets the absolute maximum tolerance for residuals without cutting the time step size.	1.0 x 10 ⁷	
39	max-single-precision-days	A real positive value that set the maximum time step size where single precision floating point arithmetic can be used solving for the linear systems of equations.	20.0	
40	max-strict-iter	A positive integer that sets the maximum number of Newton iterations before relaxed tolerances are used for the CNV convergence criterion.	8	
41	max-welleq-iter	A positive integer that defines the maximum number of iterations to determine the solution to the well equations.	15	
42	milu-variant	A defined character string that specifies which variant of the modified-ILU preconditioner ought to be used. Possible variants are:	ILU	
		I) ILU (default, plain ILU),		
		MILU_I (lump diagonal with dropped row entries),		
		 MILU_2 (lump diagonal with the sum of the absolute values of the dropped row entries), 		
		 MILU_3 (if diagonal is positive add sum of dropped row entrires. Otherwise substract them), 		
		 MILU_4 (if diagonal is positive add sum of dropped row entrires. Otherwise do nothing. 		
		The default is ILU		
43	newton-max-relax	A real positive value that sets the maximum relaxation factor of a Newton iteration used by the simulator.	0.5	
44	newton-relaxation-type	A character string that defines the type of relaxation used in Newton's method., default is dampen.	dampen	
45	output-dir	A character string that defines the directory to which OPM Flow to write the ECLIPSE compatible output files (restart and summary files).	N/A	

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		ow 2018-10 Command Line Options	
No.	Variable Name	Description	Default
46	output-interval	A positive integer that specifies the number of report steps between two consecutive writes of restart data.	I
47	output-mode	A character string that defines the output to *.PRT and *.DEBUG files:	all
		I) none: No output to the files.	
		2) log or false: Output logging information only.	
		3) all or true: Output everything.	
		For example to just output logging information use:output=log oroutput=false	
48	parameter-file	A character string that defines the name of a parameter file which contains the simulator's set of run-time parameters, as listed in this table.	N/A
49	preconditioner-add-well- contributions	A Boolean value set to true or false that switches on (true) or off (false) the influences of wells between cells for the pre-conditioner matrix only.	false
50	pri-var-oscilation-threshold	A real positive vale that defines the threshold value for the primary variable switching conditions after its meaning has switched to hinder oscilations.	I x 10-5
51	print-parameters	A positive integer value that request that the <u>run</u> time parameters be printed at the start of the run:	2
		1) 0: No output to the files.	
		2) I: Output *.DBG file	
		3) 2: Output to *.DBG and *.PRT files (default)	
52	print-properties	A positive integer value that request that the <u>compile</u> time parameters be printed at the start of the run:	2
		1) 0: No output to the files.	
		2) I: Output *.DBG file	
		3) 2: Output to *.DBG and *.PRT files (default)	
53	solve-welleq-initially	A Boolean value set to true or false that switches on (true) or off (false) the solving of the well equations as a pre-processing step.	true
		Note that the well equations are always added to the full system and solved until converged.	
54	solver-growth-factor	A real positive value that specifies the allowed value a time step can be increased by, subject to the maximum allowable time step size set by the -flow-solver-max-time-step-in-days parameter.	2.0
		For example, if the current time step has converged at 10 days and <i>flow-solver-growth-factor</i> is set to the default value, then the next time step will be 2.0 * 10 days, that is at 20 days.	
55	solver-max-growth	A real positive value that specifies the maximum allowed value a time step can be increased by after a report time step. subject to the maximum allowable time step size set by the —flow-solver-max-time-step-in-days parameter.	3.0

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	OPM Flow 2018-10 Command Line Options			
No.	Variable Name	Description	Default	
56	solver-max-restarts	A positive integer that defines the number of allowed consecutive restarts (or time step chops) before the simulation is terminated.	10	
57	solver-max-time-step-in-days	A real positive double precision value that specifies the maximum allowed time step size in days.	365	
58	solver-restart-factor	A real positive double precision value that sets the time step chop factor of the time step after a convergence failure.	0.33	
		For example, if the current non-convergent time step is 30 days and <i>-flow-solver-restart-factor</i> is set to the default value, then the time step will be repeated using 0.33 * 30 days as the time step, that is 9.9 days.		
59	solver-verbosity	A positive integer that specifies the "chattiness" of the non-linear solver.	I	
60	threads-per-process	A positive integer value that stipulates the maximum number of threads to be instantiated per process ('-I' means 'automatic').	I	
61	time-step-after-event-in-days	A real double precision value that sets maximum allowed time step after an event; for example, when a well is open or closed etc.	-1	
		The default value of -I means that events to do effect the time stepping.		
62	time-step-control	A character string that defines the time stepping control algorithm and is set to one of the following:	pid	
		 pid: PID controller based adaptive time step control as suggested by Turek and Kuzmin⁴. 		
		pid+iteration: Use PID and linear iteration numbers to guide the time step.		
		 pid+newtoniteration: Use PID and non-linear iterations numbers to guide the time step. 		
		4) Hardcoded: Use time steps supplied by user. Via timestep.control.filename		
63	time-step-control-decay-rate	A real positive value that specifies the decay rate a time step can be decreased by after the number of target iterations has been exceeded	0.75	

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Algebraic Flux Correction III. Incompressible Flow Problems. Uni Dortmund, Turek and Kuzmin, January, 2006 (DOI: 10.1007/3-540-27206-2 8)

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	OPM Flow 2018-10 Command Line Options			
No.	Variable Name	Description	Default	
64	time-step-control-file-name	A character string that specifies a filename where time steps are specified. The default is the character string timesteps	timesteps	
		For instance generated by the ecl_summary application in libecl as per the following UNIX command line:		
		<pre>path_to_libecl_applications/ ecl_summary DECK TIME > filename Where:</pre>		
		DECK is the name of the data deck you want to get the time steps from,		
		TIME tells the application to return the timing for the run, and		
		"filename" is the name of the file the times are piped to.		
65	time-step-control-growth-rate	A real positive value that specifies the growth rate of the time step size of the number of target iterations is undercut.	1.25	
66	time-step-control-target-iterations	A positive integer that defines the number of linear iterations which the time step control scheme should aim for (if applicable).	30	
67	time-step-control-target-newton- iterations	A positive integer that specifies the number of of Newtonian iterations which the time step control scheme should aim for (if applicable).	8	
68	time-step-control-tolerance	A real double precision value that sets the tolerance for PID (only used with the pid and pid+ options defined by the <i>-flow-timestep.control</i> option).	0.1	
69	time-step-verbosity	A positive integer that specifies the "chattiness" during the time integration.	I	
70	tolerance-cnv	A real positive double precision value that specifies the maximum non-linear tolerance error. This is the local convergence tolerance (maximum of local saturation errors).	0.01	
71	tolerance-cnv-relaxed	A real positive value that defines the relaxed local convergence tolerance that applies for iterations after the iterations with the strict tolerance.	1.0 x 10°	
72	tolerance-mb	A real positive double precision value that sets the maximum mass balance error, that is the tolerated mass balance error relative to total mass present.	1.0 x 10 ⁵	
73	tolerance-pressure-ms-wells	A real positive double precision value that specifies the tolerance for the pressure equations for multisegment wells.	1000	
74	tolerance-well-control	A real positive double precision value that sets the maximum tolerance for the well control equations.	1.0 x 10 ⁻⁷	
75	tolerance-wells	A real positive double precision value that defines the maximum non-linear error for the well equations.	0.0001	

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	OPM Flow		
No.	Variable Name	Description	Default
76	-update-equations-scaling	A Boolean value set to true or false that switches on (true) or off (false) the updating of the scaling factors for mass balance equations during the simulation.	false
77	use-amg	A Boolean value set to true or false that if set to true OPM Flow will use AMG as the linear solver's preconditioner	false
78	use-cpr A Boolean value set to true or false that when set to true OPM Flow will use CPR as the linear solver's preconditioner		false
79	use-gmres	A Boolean value set to true or false that when set to true OPM Flow will use GMRes instead of BiCGStab as linear solver within the Newton iterations.	false
80	use-inner-iterations-ms-wells	A Boolean value set to true or false that when set to true the simulator will use nested iterations for multisegment wells.	true
81	use-multisegment-well	A Boolean value set to true or false that when set to true the simulator will use the well model for multi-segment wells instead of the one for single-segment wells.	false
82	use-update-stabilization	A Boolean value set to true or false that switches on (true) or off (false) the stabilized Newton option.	true
		This option may improve convergence for some cases.	
	VTK Graphic	cs Command Line Parameters ⁵	
83	vtk-write-average-molar-masses	A Boolean value set to true or false that turns on (true) or off (false) the output of the average phase mass in the VTK output files.	false
84	vtk-write-densities	A Boolean value set to true or false that turns on (true) or off (false) the output of the phase densities to the VTK output files	true
85	vtk-write-dof-index	A Boolean value set to true or false that turns on (true) or off (false) the output of the index of the degrees of freedom to the VTK output files.	false
86	vtk-write-extrusion-factor	A Boolean value set to true or false that turns on (true) or off (false) the output of the of the extrusion factor of the degrees of freedom to the VTK output files.	false
87	vtk-write-filter-velocities	A Boolean value set to true or false that turns on (true) or off (false) the output of the filter velocities of the phases in the VTK output files.	false
88	vtk-write-fugacities	A Boolean value set to true or false that turns on (true) or off (false) the output of the component fugacities to the VTK output files.	false

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Virtualization ToolKit ("VTK") files used to visualize the 3D results from the model using ParaView, an open-source, multiplatform general purpose data analysis and visualization application (https://www.paraview.org/overview/). For ECLIPSE compatible output files OPMS's ResInsight 3D visualization package can be used which is specifically tailored for OPM Flow and the commercial simulator.

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	OPM Flow 2018-10 Command Line Options			
No.	Variable Name	Description	Default	
89	vtk-write-fugacity-coeffs	A Boolean value set to true or false that turns on (true) or off (false) the output of the component fugacity coefficients to the VTK output files.	false	
90	vtk-write-gas-dissolution-factor	A Boolean value set to true or false that turns on (true) or off (false) the output of the gas dissolution factor (Rs) of the observed oil to the VTK output files.	false	
91	vtk-write-gas-formation-volume- factor	A Boolean value set to true or false that turns on (true) or off (false) the output of the gas formation volume factor (Bg) to the VTK output files.	false	
92	vtk-write-gas-saturation-pressure	A Boolean value set to true or false that turns on (true) or off (false) the output of the saturation pressure of gas (Pg,sat) to the VTK output files'	false	
93	vtk-write-intrinsic-permeabilities	A Boolean value set to true or false that turns on (true) or off (false) the output of the intrinsic permeability to the VTK output files.	false	
94	vtk-write-mass-fractions	A Boolean value set to true or false that turns on (true) or off (false) the output of the mass fractions to the VTK output files.	false	
95	vtk-write-mobilities	A Boolean value set to true or false that turns on (true) or off (false) the output of the phase mobilities to the VTK output files.	false	
96	vtk-write-molarities	A Boolean value set to true or false that turns on (true) or off (false) the output of the component molarities to the VTK output files.	false	
97	vtk-write-mole-fractions	A Boolean value set to true or false that turns on (true) or off (false) the output of the mole fractions to the VTK output files.	true	
98	vtk-write-oil-formation-volume- factor	A Boolean value set to true or false that turns on (true) or off (false) the output of the oil formation volume factor (Bo) to the VTK output files.	false	
99	vtk-write-oil-saturation-pressure	A Boolean value set to true or false that turns on (true) or off (false) the output of the saturation pressure of oil (Po,sat) to the VTK output files.	false	
100	vtk-write-oil-vaporization-factor	IA Boolean value set to true or false that turns on (true) or off (false) the output of the oil vaporization factor (Rv) of the observed gas to the VTK output files.	false	
101	vtk-write-porosity	A Boolean value set to true or false that turns on (true) or off (false) the output of the porosity array to the VTK output files.	true	
102	vtk-write-potential-gradients	A Boolean value set to true or false that turns on (true) or off (false) the output of the phase pressure potential gradients to the VTK output files.	false	
103	vtk-write-pressures	A Boolean value set to true or false that turns on (true) or off (false) the output of the phase pressures to the VTK output files.	true	

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	OPM Flow 2018-10 Command Line Options			
No.	Variable Name	Description	Default	
104	vtk-write-primary-vars	A Boolean value set to true or false that turns on (true) or off (false) the output of the primary variables to the VTK output files.	false	
105	vtk-write-primary-vars-meaning	A Boolean value set to true or false that turns on (true) or off (false) the output of the how the primary variables should be interpreted to the VTK output files.	false	
106	vtk-write-process-rank A Boolean value set to true or false that turns on (true) or off (false) the output of the MPI process rank to the VTK output files.		false	
107	vtk-write-relative-permeabilities	·		
108	vtk-write-saturated-gas-oil- vaporization-factor	A Boolean value set to true or false that turns on (true) or off (false) the output of the oil vaporization factor (Rv,sat) of oil saturated gas to the VTK output files.	false	
109	vtk-write-saturated-oil-gas- dissolution-factor	A Boolean value set to true or false that turns on (true) or off (false) the output of the gas dissolution factor (Rs,sat) of gas saturated oil to the VTK output files.	false	
110	vtk-write-saturation-ratios	A Boolean value set to true or false that turns on (true) or off (false) the output of the ratio of the actually and maximum dissolved component of the mixture to the VTK output files	false	
111	vtk-write-saturations	A Boolean value set to true or false that turns on (true) or off (false) the output of the phase saturations to the VTK output files.	true	
112	vtk-write-temperature	A Boolean value set to true or false that turns on (true) or off (false) the output of the temperature array to the VTK output files.	true	
113	vtk-write-total-mass-fractions	A Boolean value set to true or false that turns on (true) or off (false) the output of the total mass fractions to the VTK output files.	false	
114	vtk-write-total-mole-fractions	A Boolean value set to true or false that turns on (true) or off (false) the output of the total mole fractions to the VTK output files.	false	
115	vtk-write-viscosities	A Boolean value set to true or false that turns on (true) or off (false) the output of the component phase viscosities to the VTK output files.	false	
116	vtk-write-water-formation-volume- factor	A Boolean value set to true or false that turns on (true) or off (false) the output of the oil formation volume factor (Bo) to the VTK output files.	false	

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OPM Flow 2018-10 Command Line Options				
No.	Variable Name	Description	Default	
Not	<u>es:</u>			
I)	ltems shaded in gray are considered t associated with these command line p	to be developer options that should be used with operameters are subject to change.	caution as the values	
2	As per all UNIX and LINUX based system the input is case dependent.			
2)	, p	stem the input is tast dependent		

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Table 2.2: OPM Flow 2018-10 Command Line Options

As per all UNIX and LINUX based system the input is case dependent. The command line parameters must either be entered as lower case or CamelCase, for example:

flow --enable-dry-run=false CASNAME.DATA

flow -- EnableDryRun=false CASNAME.DATA

If a parameter file is used to set one of the above parameters, the leading "--" should not be incorporated in the parameter file, for example on the command line use:

flow --enable-dry-run=false CASNAME.DATA

but in the parameter file use:

ecl-deck-filename=CASENAME.DATA

enable-dry-run=false

or:

or:

EclDeckFileName=CASENAME.DATA

EnableDryRun=false

OPM Flow prints out the command line run time and compile time parameters in CamelCase in both the *.PRT and *.DBG files for reference, one can therefore use this as basis for creating additional run specific parameter files.

Note

It is worth considering having a parameter file for each run of the form CASENAME.param, in order to re-run a case and to use the same parameters for other cases.

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Example

The following example is taken for the Norne project and the comments in the example (preceded with "//") explain the parameter setting used in the file.

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```
// INPUT AND OUTPUT OPTIONS
// -----
// Input File
11
ecl-deck-filename=NORNE_ATW2013.DATA
// Output and Output Directory
ecl-output-dir=OPM
// NEWTON SOLVER PARAMETER
//
//Define Numerical Tolerances
flow-tolerance-mb=1e-5
flow-tolerance-cnv=1e-2
flow-tolerance-wells=1e-2
// Set Min Newtonian Solver iterations to 1 and Max to 15
flow-newton-min-iterations=1
flow-newton-max-iterations=15
```

Notice that the leading "--" have not be incorporated in the parameter file, as per the notes in Table 2.2. In order to use the above parameter file called one would use the following format:

```
flow --parameter-file=CASENAME.PARAM
```

If the above parameter file was called NORNE ATW2013.PARAM, then the command would be:

3 KEYWORD DOCUMENTATION STRUCTURE

The OPM Flow manual is constructed in a manner to enable the reader to reference various parts of the document by using the table contents or by imply by pressing on a link embedded in the text. This automatic cross referencing has been extensively employed to ensure effective documentation of the keywords used by the simulator.

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3.1 Keyword Definitions

Each keyword is defined in it's own section that contains a section header, that contains the keyword name in capital letters followed by a brief description of the keyword's function. This is then followed by Keyword Table Section which defines the status of the keyword and which sections of the input deck the keyword can be utilized. Table 3.1 illustrates a typical Keyword Table Section defining the keyword status with the various OPM Flow sections.



Table 3.1: Example Keyword Table Section

The cells are colored in such a manner as to quickly indicate to the reader the keyword's section availability and function availability, with green colored cells indicating the keyword is available for this section and is mostly or fully implemented. Cells colored gray indicate that keyword cannot be used in that particular section, and cells colored orange show that cell is only partially implemented within OPM Flow, for example OPM Flow may simple just recognize the keyword and ignore the keyword's function, or only part of the keyword's function is implemented. Finally, cells colored red means that keyword is available in the commercial simulator but has not been implemented in OPM Flow. In this scenario the keyword should not be used in OPM Flow as it will result in unpredictable results, including causing the simulator to abort or throw an exception.

3.2 Multi-Section Keywords

As there are numerous keywords that can be used within multiple OPM Flow sections of the input file, for example the *ADD* and *EQUALS* keywords, there is a need to avoid duplication of the keyword definitions but at the same time attempt to define only those keywords for a given section. Thus for multi-section keywords, the keyword is defined in the first available section that the keyword can be found. The *Keyword Table Section* as shown below for the *ADD* keyword below, indicates which sections the keyword can be utilized.

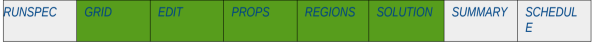


Table 3.2:ADD Keyword Table Section

Here the keyword can be used in the GRID, EDIT, PROPS, REGIONS and SOLUTION sections as indicated by those cells colored green and not for the cells colored in light gray.

A complete list of keywords in alphabetic order is given in section 13 KEYWORD INDEX - ALPHABETIC LISTING and clicking on a specific keyword will take the reader to the keyword definition in a particular section.

3.3 Keyword Formats

All keywords in OPM Flow should be entered in capital case and start in column one, lowercase entry of keywords will be produce errors and keywords not starting in column one will not be recognized. There are three types of keyword format types used by OPM Flow for data input. The description of the three types is given in the next three section together with some examples.

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3.3.1 Keyword Format Type - Comment

Comments in the input deck can occur anywhere in the file are preceded by "--" in columns one and two, for example for the EQUL keyword:

```
--
-- MAX MAX RSVD TVDP TVDP
-- EQLNUM DEPTH NODES TABLE NODES
EQLDIMS
9 1* 20 1* 1* /
```

In addition, comments can be place after "/" that terminates a record entry as shown below;

```
--
-- ARRAY CONSTANT -- --------- BOX -------
-- I1 I2 J1 J2 K1 K2
MULTIPLY
'PERMZ' 0.50000 1* 1* 1* 1* 1* 1* / PERMZ * 0.5
```

3.3.2 Keyword Format Type - Activation

This type of keyword format only consists of the keyword itself and is usually used to invoke a feature or to switch on or off a processing feature. The keyword is document by describing the functionality or action the keyword performs, followed by an example. Examples of this type of keyword include API (to switch on API tracking), GAS (to activate the gas phase in the model), ECHO (to switching echoing of the input file to the output file), and SKIP (for skipping parts of the input deck). For example the GAS keyword in the RUNSPEC section would be described as:

Description

This keyword indicate that the gas phase is present in the model and must be used for oil-gas, gas-water, oil-water-gas input decks that contain the gas phase. The keyword will also invoke data input file checking to ensure that all the required gas phase input parameters are defined in the input deck.

There is no data required for this keyword.

Example

```
-- GAS PHASE IS PRESENT IN THE RUN
-- GAS
```

The above example declares that the gas phase is active in the model.

3.3.3 Keyword Format Type - Vector (Row Vector)

Vector based keywords consist of the keyword followed by a vector of parameters on a separate line and may consists of multiple lines of vectors with each line representing a data set set (see the second example for this type of vector keyword). The vector may contain integer, real and character parameters depending on the keywords requirements. This type of keyword is documented by describing the functionality or action the keyword performs, a table describing the parameters associated with the keyword, followed by one or two examples on how to use the keyword. For example the DIMENS keyword in the RUNSPEC section would be described as:

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Description

DIMENS defines the dimensions of the model entered as integer vector. The keyword can be used for all grid types.

No.	Name	Description	Default
I	NX	The number of grid blocks in the \times direction for Cartesian grids or the number of grid blocks in the r direction for radial grids	None
2	NY	The number of grid blocks in the y direction for Cartesian grids or the number of grid blocks in the theta direction for radial grids.	None
3	NZ	The number of grid blocks in the z direction for both Cartesian and radial grids.	None

Notes:

1) The keyword is terminated by "/".

Table 3.3: DIMENS Keyword Description

Note that NX, NY and NZ are <u>not maximum</u> values but the actual size of the grid. OPM Flow applies these parameters when reading in particular data sets. For example if NX, NY, and NZ are set to 10, 10 and 10 respectively, then for the grid property data like PORO; OPM Flow expects to read in $10 \times 10 \times 10$ or 1,000 porosity values for the PORO array. If the number of porosity values is not equal to 1,000 then OPM Flow will produce an error.

Examples

```
--
-- MAX MAX MAX
-- NDIVIX NDIVIY NDIVIZ
DIMENS
46 112 22 /
```

The above example defines the dimensions for the Norne model of 36 cells in the x direction, 122 cells in the y direction and 22 cells cells in the z direction.

For vector keywords that have parameters associated with units then there is a slightly different table format to that used above to take into account the documenting the defaults for the three sets of units supported by OPM Flow, for example for the ROCK keyword is describe as follows:

Description

ROCK defines the rock compressibility for various regions in the model. The number of ROCK vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the ROCK tables to different grid blocks in the model is done via the PVTNUM keyword in the REGION section. One data set consists of one record or line which is terminated by a "/".

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This keyword must be defined in the OPM Flow input deck.

No.	Name		Default		
		Field	Metric	Laboratory	
I	Pref	Pref is a real number parameters for this data			
		psia barsa atma		atma	
		1.032	1.032	1.032	Default
2	Cf	Cf is a real number de pressure, Cf(Pref) and is	fining the rock compress defined as:	ibility at the reference	
		I/psia I/barsa I/atma			
		0.0	0.0	0.0	Defined

Notes:

The each data set terminated by terminated by "/" at the end of the line, there is no terminator for the keyword.

Table 3.4: ROCK Keyword Description

Examples

The following shows the PVTW keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to one.

```
--- ROCK COMPRESSIBILITY
--- (1) REFERENCE PRESSURE IS TAKEN FROM THE HCPV WEIGHTED RESERVOIR PRESSURE
--- AS THE PORV IS ALREADY AT RESERVOIR CONDITIONS (FLOW USES THE REFERENCE
--- PRESSURE) TO CONVERT THE GIVEN PORV TO RESERVOIR CONDITIONS USING THE DATA
--- ON THE ROCK KEYWORD)
---
ROCK

3566.9 5.0E-06 / ROCK COMPRESSIBILITY REGION 1
3966.9 5.5E-06 / ROCK COMPRESSIBILITY REGION 2
4566.9 6.0E-06 / ROCK COMPRESSIBILITY REGION 3
```

There is no terminating "/" for this keyword.

In this case the example shows a multiple data set entry of the vector format keyword, with three ROCK data sets being defined by the keyword.

3.3.4 Keyword Format Type - Vector (Columnar Vector)

Columnar vector based keywords consist of the keyword followed by a columnar vector of parameters in a separate column for each parameter. The vector may contain integer, real and character parameters depending on the keywords requirements. This type of keyword is documented by describing the functionality or action the keyword performs, a table describing the parameters associated with the keyword, followed by one or two examples on how to use the keyword. For example the SWFN keyword in the PROPS section would be described as:

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Description

The SWFN keyword defines the water relative permeability and water-oil capillary pressure data versus water saturation tables for when water is present in the input deck. This keyword should only be used if water is present in the run.

No.	Name	Description					
		Field	Metric	Laboratory			
I	SWAT		A columnar vector of real monotonically increasing down the column values starting from zero and terminating at one, that defines the water saturation.				
		dimensionless	dimensionless	dimensionless	None		
2	KRW	A columnar vector of rethe column and that are equal to one that define gas saturation. The first value in the column					
		dimensionless	dimensionless	None			
3	PCWO	A columnar vector of rest the column that defines the SWATINIT keywood columnar vector has to be					
		psia	bars	atm	None		

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 3.3: SWFN Keyword Description

Example

```
WATER RELATIVE PERMEABILITY TABLES (SWFN)
SWFN
         SWAT
                     KRW
                                 PCOW
_ _
         FRAC
                     FRAC
                                 PSIA
                                 1*
                  0.00000
0.00050
            0.15
                                   1*
            0.30
            0.40
                     0.00390
                                  1*
             0.50
                     0.01500
                                  1*
            0.60 0.04100
0.65 0.06250
0.70 0.09150
                                   1*
                                   1*
                                   1*
```

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The example defines two SWFN tables for use when water is present in the run. In the tables the water-oil capillary pressure data has been defaulted with "I*" and will be set to zero as there are no other values for the water-oil capillary pressure columns.

3.3.5 Keyword Format Type - Array

This type of keyword defines a property for the grid or an area of the grid using a previously entered BOX keyword to define the area where the property will be defined. For array data a full set of values for each element in the array is required. For example, the documentation for the PORO array would be:

Description

PORO defines the porosity for all the cells in the model via an array. The keyword can be used for all grid types.

No.	Name		Default			
		Field Metric Laboratory				
I	PORO	cell in the model. The nu NY x NZ parameters on	PORO is an array of real numbers assigning the porosity values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*100.0.			
		dimensionless	dimensionless			

Notes:

1) The keyword is terminated by "/".

Table 3.3: PORO Keyword Description

See also the DX, DY and TOPS keywords to fully define a Cartesian Regular Grid.

Examples

```
-- DEFINE GRID BLOCK POROSITY DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- PORO
300*0.300
```

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3.4 INPUT FILE STRUCTURE

OPM Flow input files are similar to commercial simulators that are used in the oil and gas industry, that is the input file is separate into sections in an effort to avoid an engineer's input data errors and a computer programmer's code to interpret the data. OPM Flow has been designed, from an engineer's prospective and input structure to be similar to Schlumberger's industry wide ECLIPSE 1006 simulator. Table 3.4 lists the various section together with a brief description of purpose of the section. As well if section is mandatory or not for a run to form a valid input deck

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Section	Description	Required
Name		Optional
RUNSPEC	This is the first section in the OPM Flow input file and defines the key parameters for the simulator including the dimensions of the model, phases present in the model (oil, gas and water for example), number of tables for a given property and the maximum number of rows for each table, the maximum number of groups, wells and well completions, as well as various options to be invoked by OPM Flow.	Required
GRID	Defines the basic grid properties, including structure, faults and various static rock properties (porosity, permeability etc.). Upon completion of reading this section, the software calculates the pore volume (PORV) for each and the transmissibilities (TRANX, TRANY and TRANX,) between all the cells, as well as calculating the transmissibilities of the Non-Neighbor Connections ("NNC")	Required
EDIT	The properties calculated by OPM Flow in the GRID section are available for editing in this section (PORV,TRANX etc.).	Optional
PROPS	This section defines the fluid properties for all the phases present in the run, for example oil viscosity, oil formation volume factor etc. The section also defines the rock flow properties as per the relative permeabilities and the distribution of the fluids based on the capillary pressure functions.	Required
REGIONS	The REGIONS section allows the engineer to define various regions in the model for reporting purposes and to define how the fluid and rock property defined in the PROPS section are allocated throughout the model.	Required
SOLUTION	Defines the parameters to initialize the model, fluid contacts, reservoir pressures etc., together with the data from the previous sections. This section, if requested, reports the initial in-place volumes for phases present in the model, as well as the average pressure for the various defined regions.	Required
SUMMARY	Defines the time vector property data to be written out to various report output files for reviewing the results of the simulation. This data includes field, group, well and well completion production and injection data, for example field oil rate versus time.	Optional
	Grid block data can also be reported versus time as well, for example grid block pressure versus time ² .	
SCHEDULE	The final section is the SCHEDULE that the defines the field, group and well parameters, targets and constraints that should be applied to the field, group or wells, numerical controls, the operating schedule and reporting requirements.	Required

Notes:

- Although the SUMMARY section is optional, it is nearly always included in order to obtain results from the simulation run.
- 2) The OPM ResInsight three-dimensional visualization software has a feature to display a grid block property change versus time, so it should not be necessary to export the grid block data to the SUMMARY report output files.

Table 3.4: OPM Flow Input Deck Sections

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⁶ ECLIPSE Industry-Reference Reservoir Simulator – Reference Manual 2015.1, Schlumberger.

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4 GLOBAL SECTION KEYWORDS

Keywords used in this section can be used in all input file sections

4.1 Keyword Definitions

4.1.1 DEBUG - Define the Debug Data to be Printed to File

RUNSPEC GRID EDIT PROPS REG	SIONS SOLUTION SUMMARY SCHEDULE
-----------------------------	---------------------------------

Description

This keyword defines the debug data to be written to the debug it is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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4.1.2 ECHO - ACTIVATE ECHOING OF USER INPUT FILES TO THE PRINT FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Turns on echoing of all the input files to the print file; note that this keyword is activated by default and can subsequently be switched off by the NOECHO activation keyword.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- SWITCH OFF ECHOING OF INPUT FILES
NOECHO

-- INCLUDE SIMULATION GRID WITH SLOPING FAULTS
INCLUDE
'./INCLUDE/GRID/IRAP_1005.GRDECL' /

-- SWITCH ON ECHOING OF INPUT FILES
ECHO
```

The examples deactivates the echoing of the input files, reads in the grid geometry data using the INCLUDE keyword, and then activates the echoing of the input files again.

Note

Especially for the large voluminous data sets in the GRID section, it is good practice to deactivate the echoing of the input files when loading this data to avoid the print output file becoming too large to view in a text editor.

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4.1.3 END - Define the End of the Input File

RUNSPEC GRID EDIT PROPS REGION	NS SOLUTION SUMMARY SCHEDULE
--------------------------------	------------------------------

Description

This keyword marks the end of the input file and can occur in any section. Any keywords and data after the END keyword are ignored.

There is no data required for this keyword.

Example

```
-- SCHEDULE SECTION - 2006-01-01
RPTSCHED
'WELLS=2'
            'WELSPECS' 'CPU=2'
                                    'FIP=2'
DATES
         2006 /
 1 JAN
RPTSCHED
'NOTHING'
DATES
 1 APR
         2006 /
         2006 /
   JLY
 1
 1 OCT
         2006 /
ECH0
END
-- SCHEDULE SECTION - 2007-01-01
RPTSCHED
           'WELSPECS' 'CPU=2'
                                   'FIP=2'
'WELLS=2'
DATES
         2007 /
 1 JAN
```

In the above example OPM Flow will process the data up to October I, 2006 only, and then start to run the simulation. All keywords after the END file keyword will not be read or processed.

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4.1.4 ENDING - Define the End of an Include File

RUNSPEC GRID EDIT PROPS REGION	NS SOLUTION SUMMARY SCHEDULE
--------------------------------	------------------------------

Description

This keyword marks the end of an include file specified on the INCLUDE keyword. When the ENDINC keyword is encountered in the INCLUDE file, input data is read from the next keyword in the current file. Any keywords and data after the ENDINC keyword in the INCLUDE file are ignored.

There is no data required for this keyword.

Example

```
-- SCHEDULE SECTION - 2006-01-01
RPTSCHED
         'WELSPECS'
                   'CPU=2'
                              'FIP=2'
'WELLS=2'
DATES
       2006 /
1 JAN
RPTSCHED
'NOTHING'
DATES
 1 APR
        2006 /
        2006 /
 1
  JIY
        2006 /
1
  0CT
ECH0
-- END OF INCLUDE FILE PROCESSING
ENDINC
-- SCHEDULE SECTION - 2007-01-01
RPTSCHED
                   'CPU=2'
'WELLS=2'
         'WELSPECS'
                             'FIP=2'
DATES
       2007 /
1 JAN
```

In the above example OPM Flow will process the data up to October I, 2006 only, and return control to the file that called the INCLUDE keyword, and then continue processing the input files. All keywords after the ENDINC keyword in the INCLUDE FILE will not be read or processed.

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4.1.5 ENDSKIP - DEACTIVATES SKIPPING OF KEYWORDS AND INPUT DATA

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

Turns off skipping of keywords that was activated by the SKIP, SKIP100, or SKIP300 keywords. Note that for each SKIP keyword activated there must be a corresponding ENDSKIP keyword.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

The example skips reading of the of the grid geometry data using the INCLUDE keyword, and then reverts back to reading the input files again.

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4.1.6 EXTRAPMS - ACTIVATE EXTRAPOLATION WARNING MESSAGES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The EXTRAPMS keyword activates extrapolation warning messages for when OPM Flow extrapolates the PVT or VFP tables. Frequent extrapolation warning messages should be investigated and resolved as this would indicate possible incorrect data and may result in the simulator extrapolating to unrealistic values.

No.	Name	Description	Default
I	EXTRAP	Defines a single integer that activates the extrapolation warning message options for PVT and VFP tables. EXTRAP can have the following values:	0
		0) 0 - No warning messages are give (the default).	
		 I – PVT table extrapolation warnings are printed. 	
		2) 2 – VFP table extrapolation warnings are printed.	
		 3 – PVT and VFP table extrapolation warnings are printed. 	
		 4 - PVT and VFP table extrapolation warnings are printed with additional information. 	

Notes:

- 1) In addition extrapolation warnings will also be given for Rs and Rv if options (1), (3), and (4) are requested.
- 2) The keyword is terminated by "/".

Table 4.1: EXTRAPMS Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

-- ACTIVATE EXTRAPOLATION MESSAGES
-- EXTRAPMS
2

,

The above example activates the default the VFP table extrapolation warnings option.

4.1.7 INCLUDE - LOAD ANOTHER DATA FILE AT THE CURRENT POSITION

R	UNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The INCLUDE keyword informs OPM Flow to continue reading input data from the specified INCLUDE file. When the end of the INCLUDE file is reached, or the ENDINC is encountered, input data is read from the next keyword in the current file. Although INCLUDE files can be nested, that is INCLUDE files within INCLUDE files etc., in practice this should be avoided due to the complexity of tracking the files.

No.	o. Name Description					
1	FILEINC	A character string enclosed in quotes that defines a file to read in and be processed by OPM Flow.	None			
Note:	_	s terminated by "/".				

Table 4.2: INCLUDE Keyword Description

Examples

The first example shown below loads the grid file from the same directory as the data file.

```
--

-- LOAD INCLUDE FILE

--

INCLUDE

'NOR-OPM-A00-GRID.inc' /
```

The next example loads the same file one directory above from where the data file is located.

```
--
-- LOAD INCLUDE FILE
--
INCLUDE
'../NOR-OPM-A00-FAULTS.inc' /
```

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4.1.8 MESSAGES - Define Message Print Limits and Stop Limits

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Revision: Rev-I

Description

The MESSAGES keyword defines the print and stops levels for various messages. The "print limits" set the maximum number of messages that will be printed, after which no more messages will be printed and the "stop limits" terminate the run when these limits are exceeded. There are six levels of message that increase in severity from informative all the way programming errors, as outlined in Table 4.3.

No.	Name	Description	Default
I	PRTSMESG	An integer defining the maximum number of MESSAGE type messages to be printed after which this type of message stops printing.	1,000,000
2	PRTSCOMT	An integer defining the maximum number of COMMENT type messages be printed after which this type of message stops printing.	1,000,000
3	PRTSWARN	An integer defining the maximum number of WARNING type messages be printed after which this type of message stops printing.	10,000
4	PRTSPROB	An integer defining the maximum number of PROBLRM type messages to be printed after which this type of message stops printing.	100
5	PRTSERRS	An integer defining the maximum number of ERROR type messages to be printed after which this type of message stops printing.	100
6	PRTSBUGS	An integer defining the maximum number of BUG type messages to be printed after which this type of message stops printing.	100
7	STOPMESG	An integer defining the maximum number of MESSAGE type messages to be printed after which OPM Flow terminates the run. Not used by OPM Flow.	1,000,000
8	STOPCOMT	An integer defining the maximum number of COMMENT type messages to be printed after which OPM Flow terminates the run. Not used by OPM Flow.	1,000,000
9	STOPWARN	An integer defining the maximum number of WARNING type messages to be printed after which OPM Flow terminates the run. Not used by OPM Flow.	10,000
10	STOPPROB	An integer defining the maximum number of PROBLEM type messages to be printed after which OPM Flow terminates the run. Not used by OPM Flow.	100
11	STOPERRS	An integer defining the maximum number of ERROR type messages to be printed after which OPM Flow terminates the run. Not used by OPM Flow.	10
12	STOPBUGS	An integer defining the maximum number of BUG type messages to be printed after which OPM Flow terminates the run. Not used by OPM Flow.	I
13	PRTGRPMS	An integer defining the maximum number of GROUP MESSAGE type messages to be printed after which this type of message stops printing. Not used by OPM Flow.	10

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No.	Name	Description	Default				
Notes	<u>:</u>						
I)	MESSAGE type m	nessages are informative messages.					
2)	COMMENT type messages are probably not data errors.						
3)	WARNING type message are possible data errors and should be investigated.						
4)	PROBLEM type errors messages are associated with numerical problems.						
5)	ERROR type messages are errors are need to be fixed before the simulator can run the input deck.						
6)	BUG type of mes	ssages are potential programming errors.					
7)	The keyword is t	erminated by "/".					

Revision: Rev-I

Table 4.3: MESSAGES Keyword Description

Examples

```
-- MESS COMMT WARN PROBL ERROR BUG MESS COMMT WARN PROBL ERROR BUG
-- LIMIT LIMIT LIMIT LIMIT LIMIT LIMIT STOP STOP STOP STOP STOP
MESSAGES

1* 1* 1* 1500 1* 1* 1* 1* 1* 1000 1* 1* 1* /
```

The above example sets the PROBLEM type message print limit to 1,500 and the stop limit to 1,000.

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4.1.9 NOECHO - DEACTIVATE ECHOING OF USER INPUT FILES TO THE PRINT FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Turns off echoing of all the input files to the print file. Note by default echoing of the inputs files is active. but can subsequently be switched off by the NOECHO activation keyword.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- SWITCH OFF ECHOING OF INPUT FILES
NOECHO

-- INCLUDE SIMULATION GRID WITH SLOPING FAULTS
INCLUDE
    './INCLUDE/GRID/IRAP_1005.GRDECL' /

-- SWITCH ON ECHOING OF INPUT FILES
ECHO
```

The examples deactivates the echoing of the input files, reads in the grid geometry data using the INCLUDE keyword, and then activates the echoing of the input files again.

Note

Especially for the large voluminous data sets in the GRID section, it is good practice to deactivate the echoing of the input files when loading this data to avoid the print output file becoming too large to view in a text editor.

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4.1.10 NOWARN - DEACTIVATE WARNING MESSAGES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Turns off warning messages to be printed to the print file; note that this keyword is deactivated by default and can subsequently be switched off by the WARN activation keyword. The warning messages may be turned on and off using keywords WARN and NOWARN.

It is recommended that WARN should always be used and action taken if necessary. For subsequent runs, the warning messages can be turned off.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- SWITCH OFF WARNING MESSAGES
NOWARN

-- INCLUDE SIMULATION GRID WITH SLOPING FAULTS
INCLUDE
    './INCLUDE/GRID/IRAP_1005.GRDECL' /

-- SWITCH ON WARNING MESSAGES
WARN
```

The examples deactivates the warning messages before reading the grid geometry data using the INCLUDE keyword, and then activates the warning messages after reading the INCLUDE file.

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4.1.11 SKIP - ACTIVATE SKIPPING OF ALL KEYWORDS AND INPUT DATE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SKIP keyword turns on skipping of keywords until the ENDSKIP activation keyword is encountered. All keywords and between the SKIP and ENDSKIP keywords are ignored by OPM Flow. See also the SKIP100 and SKIP300 keywords that skip only the "black-oil" and compositional keywords, respectively.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- SWITCH ON SKIPPING OF ALL KEYWORDS AND DATA
-- SKIP
-- INCLUDE SIMULATION GRID WITH SLOPING FAULTS
-- INCLUDE
'./INCLUDE/GRID/IRAP_1005.GRDECL' /
-- SWITCH ON READING OF ALL KEYWORDS AND DATA
-- ENDSKIP
```

The example skips reading of the of the grid geometry data using the INCLUDE keyword, and then reverts back to reading the input files again.

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4.1.12 SKIP100 - ACTIVATE SKIPPING OF "BLACK-OIL" KEYWORDS AND INPUT DATE

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHED

Revision: Rev-I

Description

This keyword turns on skipping of "black-oil" keywords until the ENDSKIP activation keyword is encountered. All "black-oil" keywords between the SKIP100 and ENDSKIP keywords are ignored by OPM Flow. See also the SKIP and SKIP300 keywords that skip all and only the compositional keywords, respectively. There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
- -
         SWITCH ON SKIPPING OF BLACK-OIL KEYWORDS AND DATA
- -
SKIP100
         INCLUDE BLACK-OIL PVT DATA
INCLUDE
         './INCLUDE/'BLACK-OIL-PVT'
         INCLUDE COMPOSITIONAL PVT DATA
- -
INCLUDE
         './INCLUDE/'COMPOSITION-PVT-EOS'
         SWITCH ON READING OF ALL KEYWORDS AND DATA
- -
ENDSKIP
         WATER PVT TABLE
- -
PVTW
         REF PRES BW
                             CW
                                        VISC
                                                 VISC
- -
                   RB/STB
                            1/PSIA
                                       CPOISE
                                                GRAD
- -
                   -----
                             -----
                                       -----
          4840.0
                                        0.370
                                                             / WATER DATA REGION 1
                  1.019
                             2.7E-6
- -
         OTI
                  WAT
                             GAS
         DENSITY DENSITY
                             DENSITY
- -
DENSITY
          39.0
                   62.37
                             0.04520
                                                             / PVT DATA REGION 1
         ROCK COMPRESSIBILITY
- -
- -
         REF PRES CF
                   1/PSIA
- -
         PSIA
ROCK
                                                             / ROCK COMPRESSIBILITY
          3966.9
                  5.0E-06
```

The examples skips reading of the of the 'BLACK-OIL-PVT' data set and reads the 'COMPOSITION-PVT-EOS' file using the INCLUDE keyword, before proceeding to revert back to reading the input files again.

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4.1.13 SKIP300 - ACTIVATE SKIPPING OF "COMPOSITIONAL" KEYWORDS AND INPUT DATE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Turns on skipping of "compositional" keywords until the ENDSKIP activation keyword is encountered. All "compositional" keywords between the SKIP300 and ENDSKIP keywords are ignored by OPM Flow. See also the SKIP and SKIP100 keywords that skip all and only the "black-oil" keywords, respectively. There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
SWITCH ON SKIPPING OF E300 KEYWORDS AND DATA
- -
SKIP300
         INCLUDE BLACK-OIL PVT DATA
INCLUDE
         './INCLUDE/'BLACK-OIL-PVT'
         INCLUDE COMPOSITIONAL PVT DATA
INCLUDE
         './INCLUDE/'COMPOSITION-PVT-EOS'
         SWITCH ON READING OF KEYWORDS AND DATA
ENDSKIP
        WATER PVT TABLE
- -
PVTW
         REF PRES BW
                            CW
                                       VISC
                                                VISC
                   RB/STB
                            1/PSIA
                                      CPOISE
                                               GRAD
                   -----
                                      -----
         4840.0
                                       0.370
                                                           / WATER DATA REGION 1
                   1.019
                             2.7E-6
         OIL
                  WAT
                             GAS
- -
         DENSITY DENSITY
                             DENSITY
DENSITY
                                                           / PVT DATA REGION 1
         39.0
                   62.37
                             0.04520
         ROCK COMPRESSIBILITY
- -
         REF PRES CF
         PSIA
                   1/PSIA
                  _____
ROCK
                                                           / ROCK COMPRESSIBILITY
          3966.9
                 5.0E-06
```

The examples reads the 'BLACK-OIL-PVT' file using the INCLUDE keyword, then skips reading of the 'COMPOSITION-PVT-EOS' data set, before proceeding to revert back to reading the input files again.

4.1.14 WARN - ACTIVATE WARNING MESSAGES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Turns on warning messages to be printed to the print file; note that this keyword is activated by default and can subsequently be switched off by the NOWARN activation keyword. The warning messages may be turned on and off using keywords WARN and NOWARN. OPM Flow always prints error messages.

It is recommended that WARN should always be used and action taken if necessary for the initial runs, once the run has been "cleaned up" the warning messages can be turned off.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- SWITCH OFF WARNING MESSAGES
NOWARN

-- INCLUDE SIMULATION GRID WITH SLOPING FAULTS
INCLUDE
    './INCLUDE/GRID/IRAP_1005.GRDECL' /

-- SWITCH ON WARNING MESSAGES
WARN
```

The examples deactivates the warning messages before reading the grid geometry data using the INCLUDE keyword, and then activates the warning messages after reading the INCLUDE file.

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5 RUNSPEC SECTION

5.1 INTRODUCTION

This is the first section in the OPM Flow input file and defines the key parameters for the simulator including the dimensions of the model, phases present in the model (oil, gas and water for example), number of tables for a given property and the maximum number of rows for each table, the maximum number of groups, wells and well completions, as well as various options to be invoked by OPM Flow.

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The section is defined by the RUNSPEC keyword and is terminated by the GRID keyword.

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5.2 Keyword Definitions

5.2.1 ACTDIMS - ACTION Keyword DIMENSIONS

|--|

Revision: Rev-I

Description

The ACTION keyword defines the maximum number of properties associated with the ACTION keyword. The ACTION keyword allows the user to enter computational logic and calculation to the simulation run based on the how the simulation run is proceeding.

Although this keyword is read by OPM Flow, the ACTION and UDQ computational logic and calculations have not been fully implemented and therefore this keyword should not be used as it may result in OPM Flow aborting.

No.	Name	Description	Default
I	MXACTNS	A positive integer value that defines the maximum number of ACTION keywords defined in the input deck.	2
2	MXLINES	A positive integer value that defines the maximum number of lines in an ACTION statement.	50
3	MXCHARS	A positive integer value that defines the maximum characters in an ACTION statement.	80
4	MXSTATMS	A positive integer value that defines the maximum number of conditional statements in the ACTION statement.	3
Notes			

Notes:

Table 5.1:ACTDIMS Keyword Description

Although this keyword is read by OPM Flow, the ACTION and UDQ computational logic and calculations have not been fully implemented and therefore this keyword should not be used as it may result in OPM Flow terminating.

Examples

	ACTION	ACTION	ACTION	ACTION	
	MXACTNS	MXLINES	MXCHARS	MXSTATMS	
ACTDIMS					
	2	50	80	3	/

The above example defines the default values for the ACTDIMS keyword.

¹⁾ The keyword is terminated by "/".

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5.2.2 AITS - INTELLIGENT TIME STEPPING ACTIVATION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

Turns on ECLIPSES intelligent time stepping.

This keyword is ignored by OPM Flow and has no effect on the simulation

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5.2.3 API - ACTIVATE API TRACKING

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on API tracking so that the various "oil types" are tracked in the model.

In many reservoirs the initial API gravity of oil varies with depth due to the heavy viscous fractions occupying the deepest part of the reservoir whilst the lighter more mobile fractions will occupy the upper part of the reservoir. As a reservoir is depleted the API gravity of oil in a cell will gradually change as the different fluids mix.

In OPM Flow it is possible to define different PVT regions in a reservoir, as in all finite difference formulated simulators, oil moving from one region to another will suddenly assume the properties of that region it has moved to. The fluid type tracking option allows the smooth change of PVT properties in a cell to be simulated by correlating PVT properties against the API gravity of oil.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

-- ACTVATE THE API TRACKING OPTION
-- API

The above example switches on the API tracking facility.

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5.2.4 AQUDIMS - AQUIFER DIMENSIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The AQDDIMS keyword defines the dimensions of the various aquifer property data. The data is normally entered on a single line and is terminated by a "/".

No.	Name	Description	Default
1	MXAQN	A positive integer value that defines the AQUNUM keyword maximum number of lines associated with this keyword.	I
2	MXNAQC	A positive integer value that defines the AQUCON keyword maximum number of lines of connection data associated with this keyword.	I
3	NIFTBL	A positive integer value that defines the AQUTAB keyword maximum number of Carter-Tracy aquifer tables associated with this keyword.	I
4	NRIFTB	A positive integer value that defines the AQUTAB keyword maximum number of rows in the Carter-Tracy aquifer tables associated with this keyword. NRIFTB must not be less than than 36 in order to accommodate the default infinite acting Carter-Tracy aquifer influence function.	36
5	NANAQ	A positive integer value that defines the AQUFET and AQUCT maximum number of analytical aquifers defined by these two keywords.	I
6	NCAMAX	A positive integer value that defines the maximum number of cells connected to an analytical aquifer	I
7	MXNALI	A positive integer value that defines the maximum number of aquifer lists.	0
8	MXAAQL	A positive integer value that defines the maximum number of analytic aquifers in any single aquifer list as defined by (7).	0

Notes:

Table 5.2:AQDIMS Keyword Description

Examples

		AQF						•	
	MXAQN	MXNAQC	NIFTBL	NRIFTB	NANAQ	NCAMAX	MXNALI	MXAAQL	
AQUDIMS	1*	1*	1*	1*	1*	1*	1*	1*	/

The above example defines the default values for the AQUDIMS keyword.

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¹⁾ The keyword is terminated by "/".

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5.2.5 BLACKOIL - ACTIVATE BLACK OIL PHASES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the "black-oil" formulation, and is equivalent to setting the phases present in the model to be oil, vaporized oil, gas, and dissolved gas. Note if water is present in the model this needs to be explicitly stated via the WATER keyword in the RUNSPEC section (see also the DEADOIL and LIVEOIL keywords in the RUNSPEC section). The keyword is used by the commercial simulator's compositional THERMAL option to set the phases present in the model.

There is no data required for this keyword.

Example

The following example activates the "black-oil" phases in the model.

```
-- ACTIVATE BLACK-OIL PHASES
```

BLACKOIL

Alternatively one could explicitly declare the phases using the following keywords in the RUNSPEC section.

```
-- OIL PHASE IS PRESENT IN THE RUN
-- OIL
-- VAPORIZED OIL IN WET GAS IS PRESENT IN THE RUN
-- VAPOIL
-- GAS PHASE IS PRESENT IN THE RUN
-- GAS
-- DISSOLVED GAS IN LIVE OIL IS PRESENT IN THE RUN
-- DISGAS
```

The above example switches on the "black-oil" phases in the model.

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5.2.6 CPR - ACTIVATE CONSTRAINED PRESSURE RESIDUAL ("CPR") LINEAR SOLVER

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Turns on the Constrained Pressure Residual ("CPR") linear solver. There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

See section 2.2 Running Flow on how to invoke various numerical schemes via the OPM Flow command line interface.

Example

-- ACTIVATE CONSTRAINED PRESSURE RESIDUAL LINEAR SOLVER FOR THE RUN CPR

The above example activates linear solver for the run.

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5.2.7 DEADOIL - ACTIVATE THE DEAD OIL PHASE (NO FREE OR DISSOLVED GAS)

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY S	SCHEDULE
----------------------------------------------------	----------

Revision: Rev-I

Description

This keyword deactivates free and dissolved gas in the model and therefore makes the oil phase dead oil in the "black-oil" formulation, and is equivalent to setting the phases present in the model to be oil and water phases. Note if water is present in the model this needs to be explicitly stated via the WATER keyword in the RUNSPEC section (see also the BLACKOIL and LIVEOIL keywords in the RUNSPEC section). The keyword is used by the commercial simulator's compositional THERMAL option to set the phases present in the model.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

The following example activates the "black-oil" phases in the model.

-- ACTIVATE DEAD-OIL PHASE

DEADOIL

Alternatively one could explicitly declare the phases using the following keywords in the RUNSPEC section.

```
-- OIL PHASE IS PRESENT IN THE RUN
-- OIL
-- WATER PHASE IS PRESENT IN THE RUN
-- WATER
```

The above example switches on the oil and water phases in the model.

⁷ "Live" oil is oil that contains gas in solution, which is normally the case for most conventional oil reservoirs. However, for oil reservoirs classified as heavy oil reservoirs, the in situ dissolved gas may be negligible and oil would then be classified as gasfree oil which is commonly referred to as "dead" oil.



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5.2.8 DIMENS - Define the Dimension of the Model

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DIMENS defines the dimensions of the model entered as integer vector. The keyword can be used for all grid

No.	Name	Description	Default
I	NX	A positive integer value that defines the number of grid blocks in the x direction for Cartesian grids or the number of grid blocks in the r direction for radial grids	None
2	NY	A positive integer value that defines the number of grid blocks in the y direction for Cartesian grids or the number of grid blocks in the theta direction for radial grids.	None
3	NZ	A positive integer value that defines the number of grid blocks in the z direction for both Cartesian and radial grids.	None

Table 5.3: DIMENS Keyword Description

Note that NX, NY and NZ are not maximum values but the actual size of the grid. OPM Flow applies these parameters when reading in particular data sets. For example if NX, NY, and NZ are set to 10, 10 and 10 respectively, then for the grid property data like PORO; OPM Flow expects to read in 10 x 10 x 10 or 1,000 porosity values for the PORO array. If the number of porosity values is not equal to 1,000 then OPM Flow will produce an error.

Examples

MAX MAX MAX NDIVIX NDIVIY NDIVIZ **DIMENS** 46 112 22

The above example defines the dimensions for the Norne model of 46 cells in the x direction, 112 cells in the y direction and 22 cells cells in the z direction.

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¹⁾ The keyword is terminated by "/".

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5.2.9 DISGAS - ACTIVATE THE DISSOLVED GAS PHASE IN THE MODEL

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION	SUMMARY	SCHEDULE
------------------------------------------	---------	----------

Revision: Rev-I

Description

This keyword indicates that dissolved gas is present in live⁸ oil in the model and the keyword should only be used if the there is both oil and gas phases in the model. The keyword may be used for oil-water and oil-water-gas input decks that contain the oil and gas phases. The keyword will also invoke data input file checking to ensure that all the required oil and gas phase input parameters are defined in the input deck.

If the oil has a constant and uniform dissolved gas concentration, Gas-Oil Ratio ("GOR"), and if the reservoir pressure never drops below the saturation pressure pressure (bubble point pressure), then the model can be run more efficiently by omitting the GAS and DISGAS keywords from the RUNSPEC section, treating the oil as a dead oil, and defining a constant Rs (GOR) value with keyword RSCONST or RSCONSTT in the PROPS section. This results in the model being run with as a dead oil problem with no active gas phase. However, OPM Flow takes into account the constant Rs in the calculations and reporting.

There is no data required for this keyword.

Example

... DISSOLVED GAS IN LIVE OIL IS PRESENT IN THE RUN

DISGAS

The above example declares that the dissolved gas in the oil phase is active in the model.

⁹ "Dead" oil is oil that it contains no dissolved gas or a relatively thick oil or residue that has lost its volatile components.



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⁸ "Live" oil is oil that contains gas in solution, which is normally the case for most conventional oil reservoirs. However, for oil reservoirs classified as heavy oil reservoirs, the in situ dissolved gas may be negligible and oil would then be classified as gasfree oil which is commonly referred to as "dead" oil.

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5.2.10 ENDSCALE - ACTIVATE RELATIVE PERMEABILITY END-POINT SCALING OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The ENDSCALE keyword activates OPM Flow's relative permeability end-point scaling option. The relative permeability functions are defined using the either the:

- 1) SWOF, SGOF, SLGOF series of saturation functions, or the
- 2) SWFN, SGFN, SGWFN, SOF2, SOF3, SOF32D series of functions.

And are allocated to the grid cells via the SATNUM keyword.

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWLX, SWLY and SWLZ instead of SWL etc. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWLX, SWLY-, SWLY- SWLZ and SWLZ-, instead of SWL or the SWLX, SWLY and SWLZ set of keywords.

The keyword also defines the number of saturation end-point tables that allows for the re-scaling of the saturation functions to be tion of depth as oppose to being a grid property array. This is accomplished via the ENKRVD and ENPVTD keywords in the PROPS section.

No.	Name	Description	Default
I	DIRECT	A character string that activates or deactivates directional end-point scaling scaling option.	NODIR
		If DIRECT is set to NODIR then directional end-point scaling is switch off and the same saturation function is used in the x, y and z directions (unless activated otherwise by the SATOPS keyword in the RUNSPEC section). In this case the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR and SOGCR saturation grid arrays and the KRG, KROG, KROW and KRW relative permeability grid cell arrays should be use to enter the grid block end-point data.	
		If DIRECT is to DIRECT then directional end-point scaling is switch on and the same saturation function is used in the x, y and z directions (unless activated otherwise by the SATOPS keyword in the RUNSPEC section). Here the directional form of the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR and SOGCR saturation grid arrays and the KRG, KROG, KROW and KRW relative permeability grid cell arrays should be use to enter the grid block end-point data. For example SWLX, SWLY and SWLZ for SWL Activates or deactivates directional end-point scaling.	
		Only the default option is supported by OPM Flow.	

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No.	Name	Description	Default
2	IRREVERS	A character string that activates or deactivates non-reversible end-point scaling option.	REVERS
		If IRREVERS is set to REVER then the end-point scaling is set to reversible and results in the same set of of end-point arrays being used for flow from the x_1 to x_{l+1} direction as for the flow from the x_l to the x_{l-1} for all directions (x, y and z). Here the SWLX, SWLY and SWLZ series of keywords should be used instead of SWL type of keywords.	
		Alternatively, if IRREVERS is set to IRREVERS then the end-point scaling is set to non-reversible and results in different sets of of end-point arrays being applied for flow from the x_1 to x_{1+1} direction and the x_1 to the x_{1-1} direction, for all directions (x, y, z) . in this case the SWLX+, SWLX-, SWLY+, SWLY- SWLZ+ and SWLZ- series of keywords should be utilized instead of SWL or the SWLX, SWLY and SWLZ set of keywords.	
		Only the default option is supported by OPM Flow.	
3 NTENDP	NTENDP	A positive integer that defines the maximum number of saturation end-point depth tables. The end-point depth tables are used to re-scale the saturation tables as a function of depth as oppose to being a grid block property. NTENDP may also be specified on the TABDIMS keyword, and if specified on both here and on the TABDIMS keyword the maximum value of the two is used.	I
		Only the default option is supported by OPM Flow.	
4	NNODES	A positive integer the defines the maximum number entries for saturation end-point depth tables.	20
		Only the default option is supported by OPM Flow.	
5	MODE	A positive integer that activates the options for temperature dependent saturation end-point scaling.	0
		Only the default value, MODE equal to zero, is implemented that means that scaling can only be performed by grid block end-point scaling properties or via saturation end-point depth tables.	

Revision: Rev-I

- Note that the IRREVERS option can only be set to IRREVERS if the DIRECT parameter is set equal to DIRECT.
- The keyword is terminated by "/".

Table 5.4: ENDSCALE Keyword Description

Examples

	DIRC	REVERSE	MAX	MAX
	SCALE	SCALE	TABLES	NODES
ENDSCALE	Ξ			
	NODIR	REVERS	1*	1*

The above example invokes the end-point scaling option with end-point scaling being non-directional and reversible with the default number of saturation end-point depth tables (one) with 20 entries per table.

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5.2.11 EQLDIMS - Define the Equilibration Data Dimensions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The EQLDIMS keyword defines the maximum number of properties associated with equilibrating the model, that is initializing the model. A reservoir grid can be separated into separate regions in order to apply different pressure regimes and/or fluid contacts. Care should be taken that the different regions are not in communication if the pressures or fluid contacts are different for the various regions, as this would lead to an unstable initialization and would also imply errors in the model description as implemented.

Name	Description	Default
NTEQUL	A positive integer value that defines the number of equilibration regions entered using the EQLNUM keyword in the REGIONS section and the number of entries associated with the EQUIL keyword in the SOLUTION section.	I
NPRSVD	A positive integer value setting the number of pressure versus depth entries used by OPM Flow in determining equilibration parameters.	100
	Unless there is a requirement for a very fine equilibration this parameter should be defaulted.	
NDRXVD	A positive integer value that the defines the maximum number of depth entries in equilibration property versus depth tables (RSVD, RVVD, PBVD or PDVD etc.) as defined in the SOLUTION section.	20
NTTRVD	A positive integer that defines the maximum number of TVDP tables that describe the initial tracer concentration versus depth. This option is ignored by OPM Flow.	I
NSTRVD	A positive integer that defines the maximum number of depth entries in the TVDP tables as described in (4).	20
	NTEQUL NPRSVD NDRXVD NTTRVD	NTEQUL A positive integer value that defines the number of equilibration regions entered using the EQLNUM keyword in the REGIONS section and the number of entries associated with the EQUIL keyword in the SOLUTION section. NPRSVD A positive integer value setting the number of pressure versus depth entries used by OPM Flow in determining equilibration parameters. Unless there is a requirement for a very fine equilibration this parameter should be defaulted. NDRXVD A positive integer value that the defines the maximum number of depth entries in equilibration property versus depth tables (RSVD, RVVD, PBVD or PDVD etc.) as defined in the SOLUTION section. NTTRVD A positive integer that defines the maximum number of TVDP tables that describe the initial tracer concentration versus depth. This option is ignored by OPM Flow. NSTRVD A positive integer that defines the maximum number of depth entries in

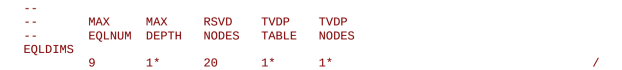
Notes:

- NTEQUL is the exact number of entries must be entered on the EQUIL keyword, otherwise OPM Flow will report an error. It is not a maximum number as for the other region arrays.
- 2) The keyword is terminated by "/".

Table 5.5: EQLDIMS Keyword Description

It is common that the EQLNUM and FIPNUM arrays are identical so that the fluid in-place reporting matches the equilibration regions. Thus, in order to avoid errors in this case, one should just use one array (say the FIPNUM property array) and use the COPY keyword to generate the EQLNUM array.

Example



The above example defines nine equilibration regions the default values for the remaining parameters on the EQLDIMS keyword.

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5.2.12 EQLOPTS - ACTIVATES THE EQUILIBRATION OPTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The EQLOPTS keyword defines the equilibration options by stating the character command to activate an option to be used for initializing the model. Multiple commands may be utilized to activate several equilibration options following the keyword.

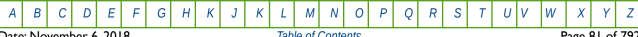
No.	Name	Description	Default
1	MOBILE	A character string that activates the mobile fluid critical saturation end point correction.	None
		If the MOBILE command is stated then this option is activated.	
		This option is not supported and should be defaulted with I^{\ast} on the keyword.	
2	QUIESC	A character string that activates the initial quiescence option that modifies the equilibrium calculated phase pressures to ensure that a steady state solution is obtained. This options ensures that there is no flow potential between the grid blocks in a given region, which is the normal case when block-centered equilibration is used by setting BOINIT on the EQUIL keyword to zero in the SOLUTION section.	None
		If the QUIESC command is stated then this option is activated.	
		This option is ignored by OPM Flow.	
3	THPRES	A character string that activates the inter-region equilibration flow option. This option allows for a threshold pressure variable entered via the THPRES keyword to define a pressure which prevents flow between regions until the THPRES value between regions is exceeded.	None
		If the THPRES command is stated then this option is activated.	
4	IRREVER	A character string that activates the irreversible inter-region equilibration flow option. This option can only be invoked if the THPRES command has been stated. The option allows for different THPRES values for different directions.	None
		If the IRREVER command is stated then this option is activated.	
		This option is not supported and should be defaulted with I* on the keyword.	

Table 5.6: EQLOPTS Keyword Description

Examples

- ACTIVATE EQUILIBRATION OPTIONS MOBILE ENDPOINT(MOBILE) STEADY STATE(QUIESC) THRESHOLD(THPRES) IRREVERSIBLE THRESHOLD(IRREVERS) **EQLOPTS**
 - 'THPRES' 'IRREVERS'

The above example activates the threshold pressure option with different threshold pressure for different directions.



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5.2.13 FAULTDIM - Define the Number of Fault Segments

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The FAULTDIM keyword defines the maximum number of records (or segments) that can be entered with the FAULTS keyword. The FAULTS keyword defines the faults in the grid than can be used for setting (or resetting) transmissibility barriers across the fault planes.

No.	Name	Description	Default
I	MFSEGS	A positive integer value that defines the maximum number of records (segments) for the FAULTS keyword.	0
Notes	<u>s:</u>		
I)	The keyword is	terminated by "/".	

Table 5.7: FAULTDIM Keyword Description

Examples

-- FAULT -- SEGMS

FAULTDIM

10000

The above example defines the maximum number of records that can be entered using the FAULT keyword to be 10,0000 segments.

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5.2.14 FIELD - ACTIVATES THE OIL FIELD SYSTEM OF UNITS FOR THE MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the oil FIELD system of units for the model.

OPM Flow has three sets of units, namely: METRIC, FIELD and LAB and one of these keyword should be invoked in the RUNSPEC section to avoid any ambiguity. Both the simulator input and output units are controlled by including one of the METRIC, FIELD or LAB keywords in the RUNSPEC section of the input file.

There is no data required for this keyword and there is no terminating "/" for this keyword.

Example

-- SWITCH ON THE FIELD SYSTEM OF UNITS FOR BOTH INPUT AND OUTPUT FIELD

The above example switches on the FIELD system of units for the model.

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5.2.15 FMTIN - ACTIVATES THE FORMAT INPUT FILE OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the Format Input Files option for all input files. Similar to the commercial simulator, OPM Flow has various options for reading various input files and writing the resulting OPM Flow output files as described in Table 5.8.

Process	Option	Description	Files
Input	i.e. text files, as oppose to binary files. The input deck type. The option relates to the OPM Flow derived files tha for example when restarting from another case.		*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART
	UNIFIN	If the keyword is omitted then the default is for binary file input. A character string that defines the input files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. For this option a single summary file and a single restart file will be read. If the keyword is omitted then the default is for one file per report time step.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART
Output	FMTOUT	A character string that sets all output files to be formatted as ASCII i.e. text files, as opposed to binary files. The *.PRT, *.LOG and *.DEBUG files are always of this type. The option relates to the OPM Flow output files only. In this case the files will be portable across operating systems, but will also be very large in terms of hard disk space. For this reason it is recommend that the default option is used so that binary files are outputted. If the keyword is omitted then the default is for binary file input.	*.SMSPEC *.SUMMARY *.GRID *.INIT *.RSSPEC *.RESTART
UNIFOUT		A character string that defines the output files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. Here a single summary file and a single restart file will be generated, as opposed to one file per report time step If the keyword is omitted then the default is for one file per report time step input.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART

Notes:

- A binary file is computer-readable but not human-readable. All executable programs are stored in binary files, as are most numeric data files. In contrast, text files are stored in a form (usually ASCII) that is humanreadable.
- 2) For unified files if the run terminates unexpectedly, or there is sufficient disk space, then the last report output is not stored. Their main advantage is that if a number of simulations reside in one directory, their output is organized. There is no limit on the number of reporting steps that a unified file can store.

Table 5.8: FMTOUT Keyword Description

There is no data required for this keyword.

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Revision: Rev-I

Example

-- SWITCH ON THE FORMAT INPUT FILES OPTION $\ensuremath{\mathsf{FMTIN}}$

The above example switches on the format input file option.

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5.2.16 FMTOUT - ACTIVATES THE FORMAT OUTPUT FILE OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the Format Output Files option for all output files. Similar to the commercial simulator, OPM Flow has various options for reading various input files and writing the resulting OPM Flow output files as described in Table 5.9.

Process	Option	Description	Files
Input	FMTIN	A character string that defines the input files to be formatted as ASCII i.e. text files, as oppose to binary files. The input deck file is always of this type. The option relates to the OPM Flow derived files that used as input, for for example when restarting from another case. If the keyword is omitted then the default is for binary file input.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART
	UNIFIN	A character string that defines the input files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. For this option a single summary file and a single restart file will be read. If the keyword is omitted then the default is for one file per report time step.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART
Output	FMTOUT	A character string that sets all output files to be formatted as ASCII i.e. text files, as opposed to binary files. The *.PRT, *.LOG and *.DEBUG files are always of this type. The option relates to the OPM Flow output files only. In this case the files will be portable across operating systems, but will also be very large in terms of hard disk space. For this reason it is recommend that the default option is used so that binary files are outputted. If the keyword is omitted then the default is for binary file input.	*.SMSPEC *.SUMMARY *.GRID *.INIT *.RSSPEC *.RESTART
	UNIFOUT	A character string that defines the output files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. Here a single summary file and a single restart file will be generated, as opposed to one file per report time step. If the keyword is omitted then the default is for one file per report time step input.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART

Notes:

- A binary file is computer-readable but not human-readable. All executable programs are stored in binary
 files, as are most numeric data files. In contrast, text files are stored in a form (usually ASCII) that is humanreadable.
- 2) For unified files if the run terminates unexpectedly, or there is sufficient disk space, then the last report output is not stored. Their main advantage is that if a number of simulations reside in one directory, their output is organized. There is no limit on the number of reporting steps that a unified file can store.

Table 5.9: FMOUT Keyword Description

There is no data required for this keyword.

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Revision: Rev-I

Example

-- SWITCH ON THE FORMAT OUTPUT FILES OPTION $\ensuremath{\mathsf{FMTOUT}}$

The above example switches on the format output file option.

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5.2.17 FULLIMP - ACTIVATES FULLY IMPLICIT SOLUTION OPTION

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY	SCHEDULE
--------------------------------------------------	----------

Revision: Rev-I

Description

FULLIMP keyword activates the Fully Implicit Solution formulation and solution options. OPM Flow users a different numerical scheme which makes this keyword redundant; hence, OPM Flow ignores this keyword. It is documented here for completeness.

There is no data required for this keyword.

See section 2.2 Running Flow on how to invoke various numerical schemes via the OPM Flow command line interface.

Example

```
-- ACTIVATES THE FULLY IMPLICIT SOLUTION OPTION
```

FULLIMP

The above example switches on the fully implicit solution option; however, this has no effect in OPM Flow input decks.

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5.2.18 GAS - ACTIVATE THE GAS PHASE IN THE MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword indicate that the gas phase is present in the model and must be used for oil-gas, gas-water, oilwater-gas input decks that contain the gas phase. The keyword will also invoke data input file checking to ensure that all the required gas phase input parameters are defined in the input deck.

There is no data required for this keyword.

Example

GAS PHASE IS PRESENT IN THE RUN

GAS

The above example declares that the gas phase is active in the model.

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5.2.19 GRIDOPTS - GRID PROCESSING OPTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

GRIDOPTS activates the negative directional dependent transmissibility multipliers option, defines the maximum number of MULTNUM regions and the number of PINCHNUM regions for the model.

No.	Name	Description	Default
ı	TRANMULT	A character string that activates the negative directional dependent transmissibility multipliers option by setting TRANMULT to YES. Setting the value to NO switches off this option.	NO
		OPM Flow users a positive directional dependent transmissibility formulation to describe the flow between two cells, that is for cell (I, J, K) OPM Flow calculates the x face transmissibility between (I, J, K) and (I + I, J, K) cell face. Modification to the transmissibilities in this case is accomplished by the MULTX, MULTY and MULTZ. Keywords.	
		Setting TRANMULT to YES invokes the option to use a negative directional dependent multiplier scheme using the MULTX-, MULTY and MULTZ- keywords. In this case OPM Flow applies the x face transmissibility between (I - I, J, K) and (I, J, K) cell face when using the MULTX-, MULTY and MULTZ- keywords.	
		This option is not supported and should be defaulted with I^{*} or NO on the keyword.	
2	NRMULT	A positive integer value that defines the maximum number of MULTNUM regions for the MULTNUM array.	0
		The MULTNUM array is used in the GRID section to define various interregion transmissibility regions in the model and NRMULT sets the maximum number of regions which is the maximum value of an element in the MULTNUM array.	
		Inter-region MULTNUM transmissibility multipliers can be defined using the MULTREGT and regional pore volumes multipliers can be set using the MULTREGP keyword.	
3	NRPINC	A positive integer value that defines the maximum number of PINCHNUM regions for the PINCHNUM array.	0
		The PINCHNUM array is used in the GRID section to define various regions in the model and NRPINC sets the maximum of regions which is the maximum value of an element in the PINCHNUM array.	
		Each regions criteria for setting the pinch out criteria is set by the PINCHREG keyword.	

Table 5.10: GRIDOPTS Keyword Description

See also the MULTNUM, MULTREGPV, MULTREGT, PINCHNUM, and PINCHREG keywords.

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Revision: Rev-I

Example

--- NEG MAX MAX
-- MULTS MULTNUM PINCHNUM
GRIDOPTS
NO 9 1* /

The above example switches off the negative directional dependent transmissibility multipliers option and defines the maximum of MULTNUM regions to be nine,. The NRPINC parameter is defaulted which means there the maximum number of PINCHREG regions is zero.

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5.2.20 IMPES - ACTIVATES IMPLICIT PRESSURE EXPLICIT SATURATION SOLUTION OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

IMPES keyword activates on Implicit Pressure Explicit Saturation formulation and solution options, commonly know as IMPES.

OPM Flow users a different numerical scheme which makes this keyword redundant; hence, OPM Flow ignores this keyword. It is documented here for completeness.

There is no data required for this keyword.

Example

- -

ACTIVATES THE IMPES SOLUTION OPTION

IMPES

The above example switches on the fully implicit solution option; however, this has no effect in OPM Flow input decks.

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5.2.21 LAB - ACTIVATES THE LABORATORY SYSTEM OF UNITS FOR THE MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the LABORATORY system of units for the model.

OPM Flow has three sets of units, namely: METRIC, FIELD and LAB and one of these keyword should be invoked in the RUNSPEC section to avoid any ambiguity. Both the simulator input and output units are controlled by including one of the METRIC, FIELD or LAB keywords in the RUNSPEC section of the input file

There is no data required for this keyword.

Example

-- SWITCH ON THE LABORATORY SYSTEM OF UNITS FOR BOTH INPUT AND OUTPUT

The above example switches on the LABORATORY system of units for the model.

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5.2.22 LICENSES - Define Required Licenses for Run

RUNSPEC	GRID EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
---------	-----------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

This keyword defines the additional software licenses that are required to invoke various licensed options in the commercial simulator at the start of the run. The commercial simulator requests a license when keywords associated with a licensed option is encountered in the input deck, this may result in the license being unavailable at the time of request and after the simulation has been initiated, resulting in the run terminating. This keyword avoids this scenario by reserving the license at the start of the run.

OPM Flow is an open source project and therefore there is no license management of the various implemented options; hence, this keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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5.2.23 LIVEOIL - ACTIVATE THE LIVE OIL PHASE (OIL WITH FREE AND DISSOLVED GAS)

RUNSPEC GRID EDIT PROPS REGIO	NS SOLUTION SUMMARY SCHEDULE
-------------------------------	------------------------------

Revision: Rev-I

Description

This keyword activates oil, free and dissolved gas in the model and therefore makes the oil phase live oil 10 in the "black-oil" formulation, and is equivalent to setting the phases present in the model to be oil, dissolved gas, gas and water phases. Note if water is present in the model this needs to be explicitly stated via the WATER keyword in the RUNSPEC section (see also the BLACKOIL and DEADOIL keywords in the RUNSPEC section). The keyword is used by the commercial simulator's compositional THERMAL option to set the phases present in the model.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

The following example activates the "black-oil" phases in the model.

-- ACTIVATE DEAD-OIL PHASE

LIVEDOIL

Alternatively one could explicitly declare the phases using the following keywords in the RUNSPEC section.

```
OIL PHASE IS PRESENT IN THE RUN

OIL

DISSOLVED GAS IN LIVE OIL IS PRESENT IN THE RUN

GAS

WATER PHASE IS PRESENT IN THE RUN

WATER
```

The above example switches on the oil, dissolved gas, gas and water phases in the model.

[&]quot;Live" oil is oil that contains gas in solution, which is normally the case for most conventional oil reservoirs. However, for oil reservoirs classified as heavy oil reservoirs, the in situ dissolved gas may be negligible and oil would then be classified as gasfree oil which is commonly referred to as "dead" oil.



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Revision: Rev-I

5.2.24 LGR - Define Local Grid Refinement Parameters

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY	SCHEDULE
--------------------------------------------------	----------

Description

This keyword defines various parameters for the local grid refinement option.

Currently, OPM Flow does not support the local grid refinement feature.

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5.2.25 MEMORY - Define Allocated Memory

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the memory allocation for the run.

OPM Flow users dynamic memory allocation and therefore the keyword has no effect and is ignored by the simulator.

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5.2.26 METRIC - ACTIVATES THE METRIC SYSTEM OF UNITS FOR THE MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the METRIC system of units for the model.

OPM Flow has three sets of units, namely: METRIC, FIELD and LAB and one of these keyword should be invoked in the RUNSPEC section to avoid any ambiguity. Both the simulator input and output units are controlled by including one of the METRIC, FIELD or LAB keywords in the RUNSPEC section of the input file.

There is no data required for this keyword.

Example

-- SWITCH ON THE METRIC SYSTEM OF UNITS FOR BOTH INPUT AND OUTPUT
-- METRIC

The above example switches on the METRIC system of units for the model.

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5.2.27 MISCIBLE - Define Miscibility Todd-Longstaff Parameters

F	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The MISCIBLE keyword defines the options associated with the Todd-Longstaff II mixing parameters used for when polymer flooding or CO₂ EOR simulation cases are being run

No.	Name	Description	Default
I	NTMISC	A positive integer value that declares the number miscible residual oil saturations versus water saturations tables for SORWMIS keyword and the number Todd-Longstaff mixing parameters entries on the TLMIXPAR keyword.	I
2	NSMISC	A positive integer value that sets the maximum number of entries (or rows) for each SORWMIS table defined by the SORWMIS keyword.	20
3	MISOPT	A character string that defines the numerical dispersion control options for the oil and gas relative permeability curves, set to either NONE or TWOPOINT:	NONE
		NONE – standard single point up streaming, that is using the immediate neighbor	
		 TWOPOINT – two-point up streaming, that is using the immediate neighbor plus one cell for better numerical dispersion control but with a higher computational cost. 	
		Only the default value of NONE is supported.	

The keyword is terminated by "/".

Table 5.11: MISCIBLE Keyword Description

Example

NTAB MAX **UPSTRM** NTMISC NSMISC MISOPT **MISCIBLE** 20 NONE

The above example defines the default values for the MISCIBLE keyword, that is one table with a maximum of 20 rows per table using the standard one cell upstream option.

M. R. Todd and W. J Longstaff, The Development, Testing, and Application Of a Numerical Simulator for Predicting Miscible Flood Performance". In: J. Petrol. Tech. 24.7 (1972), pages 874{882.



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5.2.28 MONITOR - ACTIVATES OUTPUT OF THE MONITORING DATA AND FILE

F	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The MONITOR keyword activates the writing out of the run time monitoring information used by post-processing graphics software to display run time information, for example the simulated production and injection rates and cumulative values. OPM Flow does not have this functionality.

Hence, OPM Flow ignores this keyword but it is documented here for completeness.

There is no data required for this keyword.

Example

- -

ACTIVATES MONITORING OUTPUT DATA AND FILES

MONITOR

The above example switches on the output required for run time monitoring required by post-processing graphics software to review the simulation results in real time as the run progresses; however, this has no effect in OPM Flow input decks.

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5.2.29 MSGFILE - ACTIVE OR DEACTIVATE MESSAGE FILE OUTPUT

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

MSGFILE keyword activates or deactivates the message file output used by pre- and post-processing software. Note that message file processing is not available in OPM Flow.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

No.	Name	Description	Default
I	MSGOPT	A positive integer set to 0 for to deactivate message file output or 1 to activate message file output.	I
Notes			
I)	The keyword is t	rerminated by "/".	

Table 5.12: MSGFILE Keyword Description

Example

- -

-- OUTPUT -- OPTN

MSGFILE

0

/

Revision: Rev-I

The above example deactivates the message file output, but the keyword is ignored by OPM Flow.

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5.2.30 NOCASC - ACTIVATE LINEAR SOLVER TRACER ALGORITHM

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

NOCASC keyword activates the linear solver tracer algorithm for single phase tracers.

OPM Flow users a different numerical scheme which makes this keyword redundant; hence, OPM Flow ignores this keyword. It is documented here for completeness.

There is no data required for this keyword.

Example

- -

- TRACER SOLVER OPTION

NOCASC

The above example switches on the linear solver tracer algorithm; however, this has no effect in OPM Flow input decks.

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5.2.31 NOINSPEC - DEACTIVATE OUTPUT OF THE INIT INDEX FILE

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY	SCHEDULE
--------------------------------------------------	----------

Revision: Rev-I

Description

The NOINSPEC keyword deactivates the writing out of the INIT index file (*.INSPEC). The initialization data (or static data) is written out to two files one file contains the data, *.INIT, and the second file contains an index of the data i(*.INSPEC) stored in the *.INIT file. This functionality is redundant as most post-processing software require the *.INSPEC file to load the *.INIT data set.

Hence, OPM Flow ignores this keyword. It is documented here for completeness.

There is no data required for this keyword.

Example

```
-- DEACTIVATES OUTPUT OF THE INIT INDEX FILE *.INSPEC -- NOINSPEC
```

The above example switches off the writing of the INIT index file (*.INSPEC); however, this has no effect in OPM Flow input decks.

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5.2.32 NOMONITO - DEACTIVATES OUTPUT OF THE MONITORING DATA AND FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The NOMONITO keyword deactivates the writing out of the run time monitoring information used by post-processing graphics software to display run time information, for example the simulated production and injection rates and cumulative values. OPM Flow does not have this functionality.

Hence, OPM Flow ignores this keyword but it is documented here for completeness.

There is no data required for this keyword.

Example

_ _

DEACTIVATES MONITORING OUTPUT DATA AND FILES

NOMONITO

The above example switches off the output required for run time monitoring required by post-processing graphics software to review the simulation results in real time as the run progresses; however, this has no effect in OPM Flow input decks.

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5.2.33 NONNC - DEACTIVATES NON-NEIGHBOR CONNECTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The NONNC keyword deactivates non-neighbor connections ("NNCs") in the current run. NNCs create off-diagonal elements in the Jacobi matrix that impact the numerical efficiency of the solution algorithms, and thus if the run does not contain NNC's then there is the potential for greater computation efficiency. Unfortunately, nearly all models, except for the most simple models, generate NNCs via for example:

- 1) aquifer connections,
- 2) faults, and
- 3) manually entered NNCs, including those automatically generate by pre-processing software.

Due to the limited application of this option, the feature has not been implemented in OPM Flow and hence OPM Flow ignores this keyword. It is documented here for completeness.

There is no data required for this keyword.

Example

- -

-- DEACTIVATES NON-NEIGHBOR CONNECTIONS

- -

NONNC

The above example switches off the NNCs; however, this has no effect in OPM Flow input decks.

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5.2.34 NORSSPEC - DEACTIVATE OUTPUT OF THE RESTART INDEX FILE

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY	SCHEDULE
--------------------------------------------------	----------

Revision: Rev-I

Description

The NORSSPEC keyword deactivates the writing out of the RESTART index file (*.RSSPEC). The restart data (pressure, saturations etc. through time for each active cell) is written out to two files one file contains the data, *.UNRST for example, and the second file contains an index of the data (*.RSSPEC) stored in the *.UNRST file. This functionality is redundant as most post-processing software require the *.RSSPEC file to load the *.UNRST data set.

Hence, OPM Flow ignores this keyword. It is documented here for completeness.

There is no data required for this keyword.

Example

-- DEACTIVATES OUTPUT OF THE RESTART INDEX FILE *.RSSPEC -- NORSSPEC

The above example switches off the writing of the restart index file (*.RSSPEC); however, this has no effect in OPM Flow input decks.

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5.2.35 NOSIM - ACTIVATES THE NO SIMULATION MODE FOR DATA FILE CHECKING

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

NOSIM switches the mode of OFM Flow to data input checking mode. In this mode the input file is read and all messages and print instructions are sent to the respective output files. The SCHEDULE section is read but the simulation is not performed.

There is no data required for this keyword and there is no terminating "/" for this keyword.

Example

The example below switches OPM Flow to no simulation mode for data checking of the input deck.

-- SWITCH NO SIMULATION MODE FOR DATA CHECKING COMMENT OUT TO RUN THE MODEL NOSIM

And the next example shows how to commented out the NOSIM activation keyword so that the simulation will proceed.

-- SWITCH NO SIMULATION MODE FOR DATA CHECKING COMMENT OUT TO RUN THE MODEL
-- NOSIM

Note

Simulation input decks are complex and are therefore prone to typing errors, thus before submitting a run that will take over 15 minutes or so, it is a good idea to run the model with the NOSIM option. If no errors are found then the NOSIM keyword should be commented out by placing "--" before the keyword, and then re-running the model.

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5.2.36 NSTACK - Define the Stack Length for the Iterative Linear Solver

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The NSTACK keyword defines the maximum number of previous search directions stored by the linear solver. Increasing the value of NSTACK may improve the efficiency of the solver on difficult problems, but will increase the memory requirements of the simulator. The default value of 10 should be sufficient for most problems; however, if OPM Flow is having issues with the convergence of the linear questions then increasing NSTACK and LITMAX on the TUNING keyword may improve performance.

OPM Flow users a different numerical scheme which makes this keyword redundant; hence, OPM Flow ignores this keyword. It is documented here for completeness.

No.	Name	Description	Default
I	NSTACK	A positive integer that defines the maximum number of previous search directions stored by the linear solver.	10

Notes:

- I) NSTACK and LITMAX on the TUNING keyword are related such that NSTACK should always be less than or equal to LITMAX.
- 2) The keyword is terminated by "/".

Table 5.13: NSTACK Keyword Description

Example

_ _

SET STACK SIZE FOR LINEAR SOLVER

--

NSTACK

30 /

The above example sets maximum number of previous search directions stored by the linear solver to 30, this has no effect in OPM Flow input decks.

Note

If the run is suffering from linear convergence problems, then check the data first for any data issues before manipulating the numerical control parameters. For example, if OPM Flow has written some WARNING messages with respect to endpoint scaling, etc., then resolve these messages first before adjusting the numerical controls.

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5.2.37 NUMRES - Define the Number of Reservoir Grids

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The NUMRES keyword defines the number of reservoir grids (COORD data sets) that the simulator should process. Currently, this should only be set to one in OPM Flow

No.	Name	Description	Default
I	NUMRES	A positive integer greater than one that defines the maximum number COORD data sets to be processed by OPM Flow. This should be set to one.	I
Notes	•		
I)	The keyword is t	erminated by "/".	

Table 5.14: NUMRES Keyword Description

Example

-- DEFINE THE NUMBER OF RESERVOIR GRIDS (COORD DATA SETS)
-- NUMRES
1

The above example sets the maximum number of COORD data sets to be processed to one, this is the only value that can currently be used in OPM Flow.

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5.2.38 NUPCOL - Define the Number of Newtonian Iterations Used to Update WELL TARGETS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The NUPOL keyword defines the maximum number of Newtonian iterations within a time step that may be used to update the well production and injection targets, after which the well targets will be frozen until the time step calculations have converged and the time step is complete.

Wells under group control may suffer from some dependency with other wells in the same group that are under group control. This may cause some oscillation in the production and injection well rates within the group. In order to avoid this, after the number Newtonian iterations within a time step surpasses NUPCOL, the group well rates are frozen until the time step has converged. Reducing the potential of well rate oscillations within the time step may result in the group targets and limits not being exactly being met in this case. Increasing the value of NUPCOL to greater than the default value of three, will improve the accuracy of the group targets and limits at the expense of computational efficiency.

OPM Flow users a different numerical scheme which makes this keyword redundant; hence, OPM Flow ignores this keyword. It is documented here for completeness. See section 2.2 Running Flow on how to set the numerical control parameters for OPM Flow.

No.	Name	Description	Default
1	NUPCOL	A positive integer that defines the maximum number of Newtonian iterations used to update well targets within a time step.	3
Notes	-	terminated by "/".	1

Table 5.15: NUPCOL Keyword Description

Example

```
DEFINE THE NUMBER OF ITERATIONS TO UPDATE WELL FLOW TARGETS
NUPCOL
         3
```

The above example sets the default NUPCOL value; however, this has no effect in OPM Flow input decks.

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5.2.39 OIL - ACTIVATE THE OIL PHASE IN THE MODEL

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
-----------------------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword indicate that the oil phase is present in the model and must be used for oil-gas, oil-water, oil-water-gas input decks that contain the oil phase. The keyword will also invoke data input file checking to ensure that all the required oil phase input parameters are defined in the input deck.

There is no data required for this keyword.

Example

```
-- OIL PHASE IS PRESENT IN THE RUN
-- OIL
```

The above example declares that the oil phase is active in the model.

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5.2.40 OPTIONS - ACTIVATE VARIOUS PROGRAM OPTIONS

RUNSP	EC GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The OPTIONS keyword activates various OPM Flow program options; however only the options known to be available and tested are documented out of the over 200 activation operations available in the commercial simulator.

No.	Description	Default
I - 273	Undefined.	0
Notes:		
I) The	keyword is terminated by "/".	

Table 5.16: OPTIONS Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

```
--
-- SKIP ACTIVATE
-- OPTIONS OPTION
OPTIONS
77*0 1
```

The above example activates the use of scratch files for pre-processing grid geometry data for non-neighbor connections. Note if multiple options are required then one can just repeat the format of the example to activate multiple options as the keyword does not overwrite previous entries. So for example:

 OPTIONS	SKIP OPTIONS	ACTIVATE OPTION	
OFITONS	7*0	1	,
		-	/
	SKIP	ACTIVATE	
	OPTIONS	OPTION	
OPTIONS			
	77*0	1	/
	SKIP	ACTIVATE	
	OPTIONS	OPTION	
OPTIONS			
	177*0	1	/

Could be used to activate the 8, 78 and 178 options if they were available.

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5.2.41 PARALLEL - Define Run Configuration

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PARALLEL keyword defines the run to use parallel processing and sets the domain decomposition options. See Section 2.2 on how to run OPM Flow in parallel.

No.	Name	Description	Default
I	NPROCS	A positive integer that defines the number of domains or parallel processors to use for this run.	I
2	RTYPE	A character string set to either SERIAL to run the parallel code in serial mode for testing the code, or DISTRIBUTED to full utilize parallel processing.	PARALLEL
Notes	•		l

The keyword is terminated by "/". I)

Table 5.17: PARALLEL Keyword Description

OPM Flow users a different numerical scheme which makes this keyword redundant; hence, OPM Flow ignores this keyword. It is documented here for completeness.

See section 2.2 Running Flow on how to run OPM Flow in parallel mode.

Example

PARALLEL MULTI-CORE OPTIONS MACHINE TYPE NDMAIN

PARALLEL

2 DISTRIBUTED

The above example sets the number of domains (or processor)s to two and for the simulation to run in parallel mode. this has no effect in OPM Flow input decks.

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5.2.42 PATHS - Define FILENAME DIRECTORY PATH ALIASES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PATHS allows the user to define alias directory filenames to avoid long filenames with the INCLUDE, IMPORT, RESTART or GDFILE. To use the alias, should be prefixed with the \$ symbol.

OPM Flow currently does not support this feature and ignores this keyword. It is documented here for completeness.

No.	. Name Description			
- 1	NPROCS	A character string enclosed in quotes defining the alias.	None	
2	RTYPE	A character string enclosed in quotes defining the directory filename.	None	
NI-4			1	

Notes:

- 1) Multiple entries must be on separate lines (see the example).
- 2) The keyword is terminated by "/".

Table 5.18: PATHS Keyword Description

Examples

```
--
-- PATH PATH
-- ALIAS DIRECTORY FILENAME
PATHS

'GRID' '/DISK1/NORNE/2017/GRID-INCLUDES' /
'SCHD' '/DISK1/NORNE/2017/SCHD-INCLUDES' /
```

The above example defines "GRID" and "SCHD" aliases in the RUNSPEC section than can be used in the GRID and SCHEDULE sections of the input deck. The next example shows how to use the "GRID" alias with the INCLUDE keyword in the GRID section.

```
--
-- LOAD INCLUDE FILES
--
INCLUDE
   '$GRID/PORO.INC'

INCLUDE
   '$GRID/PERMX.INC'

INCLUDE
   '$GRID/NTG.INC'

/
```

Here the porosity, permeability and net-to-gross arrays are loaded in the GRID section using the directory filename aliases declared in the RUNSPEC section.

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5.2.43 POLYMER - ACTIVATE THE POLYMER PHASE IN THE MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword indicate that the polymer phase is present in the model and to activate the polymer flooding mode. I The keyword will also invoke data input file checking to ensure that all the required polymer phase input parameters are defined in the input deck.

There is no data required for this keyword.

Example

POLYMER

-- POLYMER PHASE IS PRESENT IN THE RUN

The above example declares that the polymer phase is active in the model.

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5.2.44 PIMTDIMS - Define Well Productivity Scaling Table Dimensions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PIMTDIMS keyword defines the maximum number of PIMULTAB tables and the maximum number of entries (or rows) per PIMULTAB table. The PIMULTAB keyword is used to define a well's productivity index factor as a function of a well's producing water cut, and the PIMULTAB keyword defined in the SCHEDULE section of the input deck.

No.	Name	Description	Default
I	NTPIMT	A positive integer value that defines the maximum number of PIMULTAB keywords defined in the input deck.	0
2	NRPIMT	A positive integer value defining the maximum number of entries (rows) in the PIMULTAB keyword.	0

Notes:

1) The keyword is terminated by "/".

Table 5.19: PIMTDIMS Keyword Description

Example

- -

-- MAX MAX -- TABLES ENTRIES

PIMTDIMS

1 51

The above example defines that there is one PIMULTAB table with a maximum number of 51 rows.

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5.2.45 RADIAL - RADIAL GRID ACTIVATION OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

RADIAL¹² activates the radial grid geometry option for the model, if this keyword is omitted then Cartesian geometry is assumed by OPM Flow.

Although this keyword is read by OPM Flow, radial grids have not been fully implemented and therefore this type of grid should not be used.

Radial grids are not currently implemented in this version of OPM Flow but are expected to be incorporated in a future release.



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5.2.46 REGDIMS - Define the Maximum Number of Regions for a Region Array

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The REGDIMS keyword defines the maximum number of regions for various region arrays used in the model. Note that the maximum number of FIPNUM regions can be defined both on this keyword and the TABDIMS keyword, if it set in both locations the maximum value is used. The reason for this type of inconsistency is due to the commercial simulator evolving with time as new features were added, but at the same time having to maintain backward input deck compatibility.

No.	Name	Description	Default
I	NTFIP	A positive integer defining the maximum number of regions in the FIPNUM region array. Note that this parameter may also be set on the TABDIMS keyword as well. If NTFIP is set in both places then the maximum value is used.	I
2	NMFIPR	A positive integer defining the total maximum number of regions. The FIPNUM regions are defined by (I). if additional sets of fluid in-place regions are required, as per the FIPxxx series of fluid in-place region keywords, then these are to be defined here by adding to the value in (I). So for example, if NTFIP equals 5 and the number of distinct FIPxx regions is three, then the value to enter for NMFIPR is eight.	I
3	NRFREG	A positive integer defining the maximum number of independent reservoir regions in the ISOLNUM region array.	0
4	MXNFLX	A positive integer defining the maximum number of flux regions in the FLUXNUM region array. MXNFLN can also be defined on the TABDIMS keywords as well. If MXNFLX is defined both here and on the TABDIMS keyword then the maximum value of the two is used.	0
5	NUSREG	A positive integer defining the maximum user defined regions in a commercial simulator's compositional model. This parameter is included for compatibility and should be defaulted as it is not used in OPM Flow.	0
6	NTCREG	A positive integer defining the maximum number of regions in the COALNUM region array.	I
7	NOPREG	A positive integer defining the maximum number of regions in the OPERNUM region array.	0
8	NWKDREG	A positive integer defining the maximum maximum of real double-precision work arrays for use with the OPERATE and OPERATER keywords	0
9	NWKIREG	A positive integer defining the maximum number of integer work arrays for use with the OPERATE and OPERATER keywords	0
10	NPLMIX	A positive integer defining the maximum number of regions in the PLMIXNUM region array.	I

1) The keyword is terminated by "/".

Table 5.20: REGDIMS Keyword Description

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Revision: Rev-I

Exampl	e
--------	---

	MAX	T0TAL	INDEP	FLUX	TRACK	CBM	OPERN	WORK	WORK	P0LY	
 REGDIMS	FIPNUM	REGNS	REGNS	REGNS	REGNS	REGNS	REGNS	REAL	INTG	REGNS	3
KEGDING	9	12	1*	1*	1*	1*	1*	1*	1*	1*	/

The above example defines the number of FIPNUM regions to be nine and the number of FIPxxx type of regions to be three (12-9), the rest of the region sizes are set to the default values.

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5.2.47 ROCKCOMP - ACTIVATE ROCK COMPACTION

RUNSPEC GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
--------------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

The ROCKCOMP keyword activates rock compaction and defines various rock compaction options for the run. By default OPM Flow models rock compaction via pore volume compressibility as entered on the ROCK keyword in the PROPS section. This keyword enables pressure dependent pore volume and transmissibility multipliers for rock compaction that are entered in the PROPS section using the ROCKTAB keyword. Currently OPM Flow only supports the default options for rock compaction.

No.	. Name Description			
I	ROCKOPT	A character string that defines the rock compaction option based on one of the following character strings:	REVERS	
		 REVERS: Rock compaction is reversible with increasing pressure. The rock compaction multipliers should be entered via the ROCKTAB keyword in the PROPS section. Note that this is the default value and is the only option currently supported by OPM Flow. 		
		 IRREVERS: Rock compaction is irreversible, that is the rock expansion does not occur when the pressure subsequently decreases. 		
		3) HYSTER: Invokes the hysteresis rock compaction option.		
		 BOBERG: Rock compaction hysteresis is modeled using the Boberg formulation 13. 		
		5) REVLIMIT: Activates the reversible hysteresis rock compaction option that limits the pore volume subject to reversibility based on the minimum pressure in a grid block and the initial water saturation. This option is only intended to be used with the water induced compaction model, neither of which are currently supported by OPM Flow		
		6) PALM-MAN: Rock compaction hysteresis is modeled using the Palmer-Mansoori ¹⁴ formulation for coal bed methane reservoirs, neither of which are supported by OPM Flow.		
		 NONE: Deactivates rock compaction, unless the water induced compaction model has been invoked. 		
		Only the default option is supported by OPM Flow.		
2	NTROCC	A positive integer that defines the number of rock compaction tables, that is the number of ROCKTAB tables to be used by OPM Flow.	I	
3	WATINOPT	A character string that states if the water induced rock compaction option should be used (YES) or not (NO). Only the default option is supported by OPM Flow	NO	

¹⁴ Clarkson, C.R., Pan, Z., Palmer, I. and Harpalani, S. "Predicting Sorption-Induced Strain and Permeability Increase With Depletion for Coalbed-Methane Reservoirs", SPE 114778-PA, SPE Journal (2010) 15, No. 1, 152–159.

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Beattie, C.I., Boberg, T.C., and McNab, G.S. "Reservoir Simulation of Cyclic Steam Stimulation in the Cold Lake Oil Sands," paper SPE 18752, Society of Petroleum Engineers Journal, (1991) 6, No. 2, 200-206.

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No.	Name	Description	Default
4	PORTXROP	A character string that specifies the model to be used for the porosity dependence transmissibility dependence for when transmissibility is dependent on porosity, should be set to either:	*
		 EXP: An exponential porosity-transmissibility relationship should be used. 	
		 CZ: The Carmen-Kozeny¹⁵· 16 and 17 porosity-transmissibility relationship should be used. 	
		This option is currently ignored by OPM Flow.	
Note:	<u>s:</u>		
I)	The keyword is	s terminated by "/".	

Revision: Rev-I

Table 5.21: ROCKCOMP Keyword Description

Example

-- ROCK NUMBER WAT POR-TRAN
-- OPTN TABLES INDUCE OPTION
ROCKCOMP

NO

1*

REVERS 5

The latest the state of the positional state of the state

The above example defines the default values for the ROCKCOMP keyword with five rock compaction tables.

¹⁵ J. Kozeny, "Ueber kapillare Leitung des Wassers im Boden." Sitzungsber Akad. Wiss., Wien, 136(2a): 271-306, 1927.

P.C. Carman, "Fluid flow through granular beds." Transactions, Institution of Chemical Engineers, London, 15: 150-166, 1937.

¹⁷ P.C. Carman, "Flow of gases through porous media." Butterworths, London, 1956

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5.2.48 RPTRUNSP - ACTIVATES RUNSPEC REPORTING

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates reporting of all the RUNSPEC options utilized in the run. There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

Example

- -

SWITCH ON RUNSPEC SECTION REPORTING

- -

RPTRUNSP

The above example switches on RUNSPEC reporting; however, this has no effect in OPM Flow input decks.

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5.2.49 RUNSPEC -Define the Start of the RUNSPEC Section of Keywords

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The RUNSPEC activation keyword marks the start of the RUNSPEC section that defines the key parameters for the simulator including the dimensions of the model, phases present in the model (oil, gas and water for example), number of tables for a given property and the maximum number of rows for each table, the maximum number of groups, wells and well completions, as well as various options to be invoked by OPM Flow.

Apart from COMMENTS this keyword should be the first keyword in the input deck.

There is no data required for this keyword and there is no keyword terminating "/"

Example



The above example marks the start of the RUNSPEC section in the OPM Flow data input file.

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5.2.50 SATOPTS - ACTIVATE RELATIVE PERMEABILITY ASSIGNMENT OPTIONS

F	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SATOPTS keyword activates OPM Flow's relative permeability assignment options. The relative permeability functions are defined using the either the:

- 1) SWOF, SGOF, SLGOF series of saturation functions, or the
- 2) SWFN, SGFN, SGWFN, SOF2, SOF3, SOF32D series of functions.

The allocation of the relative permeability tables to the grid cells is dependent on the options selected on this keyword (SATOPTS).

If the DIRECT option as been activated and the IRREVERS <u>has not been invoked</u> on the SATOPTS keyword, then <u>different relative permeability functions are used for each x, y, and z directions.</u> Here the KRNUMX, KRNUMX and KRNUMZ keywords are used for Cartesian grids to allocate the relative permeability tables to the cells. For Radial grids the KRNUMR, KRNUMT and KRNUMZ keywords should be used. This results in the same relative permeability curves being used in both the x_i to x_{i-1} and the x_i to x_{i+1} flow directions. Similarly for the y direction the same curves are used for the y_i to y_{i-1} and the y_i to y_{i+1} flow directions. And again for the z direction, the same relative permeability function is used for flow in the z_i to z_{i-1} and the z_i to z_{i+1} flow directions.

If the DIRECT option as been activated and the IRREVERS <u>has been invoked</u> on the SATOPTS keyword, then KRNUMX, KRNUMY and KRNUMZ keywords are used for Cartesian grids to allocate the relative permeability tables in the x_i to x_{i+1} , y_i to y_{i+1} , z_i to z_{i+1} , flow directions, respectively. For Radial grids the KRNUMR, KRNUMT and KRNUMZ keywords should be used. For flow in the x_i to x_{i-1} flow directions, etc., the KRNUMX-, KRNUMY- and KRNUMZ- keywords are used for Cartesian grids and the KRNUMR-, KRNUMT- and KRNUMZ- are used for radial grids.

The HYSTER option activates the relative permeability hysteresis option of the non-wetting phases (liquid and vapour) may show hysteretic behavior in that their values depend on whether the non-wetting phase saturation is increasing or decreasing. For this option the user specifies two sets of saturation functions, one for a drainage process (decreasing wetting phase saturation) and one for an imbibition process (increasing wetting phase saturation). For a process starting at the maximum wetting phase saturation, on the drainage curve, with the wetting phase saturation decreasing, the drainage curve is followed. Similarly for a process starting at the minimum wetting phase saturation with the wetting phase saturation increasing, the imbibition curve is followed. If the drainage or imbibition process is reversed at some point, then the data does not necessarily run back over its previous values. In OPM Flow the Carlson is used to describe relative permeability hysteresis and the Killough of the second of the carlson of the car

If the DIRECT option as been activated and the IRREVERS has.not.been.invoked on the SATOPTS keyword, then the same set are keywords as for the DIRECT only option are used to assign the drainage relative permeability curves, that is: KRNUMX, KRNUMY, etc., plus the IMBNUMX, IMBNUMY, and IMBNUMZ, keywords for the imbibition curves. If the DIRECT option as been activated and the IRREVERS has.been.invoked on the SATOPTS keyword, then the same set are keywords as for the DIRECT and IRREVERS option are used to assign the drainage relative permeability curves, that is: KRNUMX, KRNUMX-, etc., plus the IMBNUMX, IMBNUMY, MBNUMZ, IMBNUMX-, IMBNUMY-, IMBNUMZ- keywords for the imbibition curves. See Table 5.23 for the various relative permeability table allocation keywords for the various combination of DIRECT, IRREVERS and HYSTER command options.

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¹⁸ Carlson, F. M. (1981) SPE 10157, presented at the 56th Annual SPE Fall Meeting, San Antonio, 1981

¹⁹ Killough, J. E. "Reservoir Simulation with History-dependent Saturation Functions," paper SPE 5106, Society of Petroleum Engineers Journal (1976) 16, No. 1, 37-48.

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The keyword should be followed by one or more of the following keyword options.

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No.	Name	Description	Default				
I	DIRECT	A character string that activates the directional relative permeability assignment option. If the DIRECT command is stated then directional relative permeability assignment is activated and different relative permeability function are assigned to the x, y and z directions. In this case the KRNUMX, KRNUMY and KRNUMZ keywords are used for Cartesian grids to allocate the relative permeability tables. For Radial grids the KRNUMR, KRNUMT and KRNUMZ keywords should be used. Only the default option is supported by OPM Flow.					
2	IRREVERS	A character string the activates reversible directional relative permeability assignment option. If IRREVERS is stated then the relative permeability assignment is set to non-reversible and results in different sets of relative permeability tables being applied for flow from the x_i to x_{i+1} direction and the x_i to the x_{i-1} direction, for all directions (x,y,z) . in this case the KRNUMX, KRNUMY and KRNUMZ keywords are used for Cartesian grids to allocate the relative permeability tables in the x_i to x_{i+1} flow directions etc For Radial grids the KRNUMR, KRNUMT and KRNUMZ keywords should be used. For flow in the x_i to x_{i-1} flow directions, etc., the KRNUMX-, KRNUMY- and KRNUMZ- keywords are used for Cartesian grids and the KRNUMR-, KRNUMT- and KRNUMZ- are used for radial grids.	None				
3	HYSTER	A character string that activates the hysteresis option. If the HYSTER and DIRECT options have activated and the IRREVERS has not been invoked on the SATOPTS keyword, then different relative permeability functions are used for each x, y, and z directions and for the drainage and imbibition processes. Here the drainage relative permeability curves are allocated via the KRNUMX, KRNUMX and KRNUMZ keywords for Cartesian grids and the KRNUMR, KRNUMT and KRNUMZ keywords for radial grids. The imbibition relative permeability curves are allocated via the IMBNUMX, IMBNUMY and IMBNUMZ keywords for Cartesian grids and the IMBNUMR, IMBNUMT and IMBNUMZ keywords for radial grids. If the HYSTER, DIRECT and IREVERS options have activated, then different relative permeability functions are used for each x, y, and z directions, flow direction and for the drainage and imbibition processes. Then in addition to aforementioned relative permeability curves allocation keywords for the x _i to x _{i+1} flow direction etc., the x _i to x _{i-1} flow directions keywords, KRNUMX-, KRNUMY- and KRNUMZ- are used for Cartesian grids and the KRNUMR-, KRNUMT- and KRNUMZ- are used for radial grids. The imbibition relative permeability curves are allocated via the IMBNUMX-, IMBNUMY- and IMBNUMZ keywords for Cartesian grids and the IMBNUMR, IMBNUMT and IMBNUMZ keywords for radial grids. Only the default option is supported by OPM Flow.	None				
4	SURFTENSA	A character string that activates the capillary pressure surface tension pressure dependency option. Only the default option is supported by OPM Flow.	None				

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No.	Name	Name Description					
Notes	<u> </u>						
I)	 Note that the IRREVERS command can only been activated if the DIRECT command is activated at the same time. 						
2)	See Table 5.23 fo	r the various relative permeability table allocation keywords.					
3)	The keyword is to	erminated by "/".					

Revision: Rev-I

Table 5.22: SATOPTS Keyword Description

For clarity the following table outlines the keywords that should be used in allocating the relative permeability tables for the various SATOPTS options.

Option	Car	tesian	R	adial
DIRECT Flow in all directions	KRNUMX		KRNUMR	
	KRNUMY		KRNUMT	
	KRNUMZ		KRNUMZ	
DIRECT and IRREVERS				
Flow in the i to i +1 directions.	KRNUMX,		KRNUMR	
	KRNUMY		KRNUMT	
	KRNUMZ		KRNUMZ	
Flow in the i to i - I directions.	KRNUMX-		KRNUMR-	
	KRNUMY-		KRNUMT-	
	KRNUMZ-		KRNUMZ-	
DIRECT and HYSTER	<u>Drainage</u>	<u>Imbibition</u>	<u>Drainage</u>	<u>Imbibition</u>
Flow in all directions.	KRNUMX	IMBNUMX	KRNUMR	IMBNUMR
	KRNUMY	IMBNUMY	KRNUMT	IMBNUMT
	KRNUMZ	IMBNUMZ	KRNUMZ	IMBNUMZ
DIRECT, IRREVERS and HYSTER	<u>Drainage</u>	Imbibition	<u>Drainage</u>	Imbibition
Flow in the i to i +1 directions.	KRNUMX	IMBNUMX	KRNUMR	IMBNUMR
	KRNUMY	IMBNUMY	KRNUMT	IMBNUMT
	KRNUMZ	IMBNUMZ	KRNUMZ	IMBNUMZ
Flow in the i to i - I directions.	KRNUMX-,	IMBNUMX-	KRNUMR-	IMBNUMR-
	KRNUMY-	IMBNUMY-	KRNUMT-	IMBNUMT-
	KRNUMZ-	IMBNUMZ-	KRNUMZ-	IMBNUMZ-
	<u>'</u>		1	'

Notes:

 Note the drainage and imbibition classification is related to the wetting phase, that may be either oil or water; however, water is normally assumed in most cases but there are exceptions to this, especially for heavy oils.

Table 5.23: SATOPTS Relative Permeability Function Allocation Keywords.

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Examples

The first example actives the directional relative permeability assignment option only and hence the following keywords are used to allocate the relative permeability arrays for Cartesian grids: KRNUMX, KRNUMY, and KRNUMZ.

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```
-- ACTIVATE RELATIVE PERMEABILITY ASSIGNMENT HYSTERESIS OPTIONS
-- DIRECTTIONAL(DIRECT) IRREVERSIBLE(IRREVERS) HYSTERESIS(HYSTER)
SATOPTS
'DIRECT' //
```

The next example actives the directional irreversible relative permeability assignment options, and hence the following keywords are used to allocate the relative permeability arrays for Cartesian grids: KRNUMX, KRNUMY, KRNUMZ-, KRNUMY- and KRNUMZ-.

```
-- ACTIVATE RELATIVE PERMEABILITY ASSIGNMENT HYSTERESIS OPTIONS
-- DIRECTTIONAL(DIRECT) IRREVERSIBLE(IRREVERS) HYSTERESIS(HYSTER)
SATOPTS
'DIRECT' 'IRREVERS' //
```

Finally, the last option invokes all three assignment options.

```
-- ACTIVATE RELATIVE PERMEABILITY ASSIGNMENT HYSTERESIS OPTIONS
-- DIRECTTIONAL(DIRECT) IRREVERSIBLE(IRREVERS) HYSTERESIS(HYSTER)
SATOPTS
'DIRECT' 'IRREVERS' 'HYSTER'
```

In this case the drainage relative permeability curves are allocated by the KRNUMX, KRNUMY, KRNUMZ, KRNUMX-, KRNUMY-, KRNUMZ- keywords, and the imbibition relative permeability curves are allocated by the IMBNUMX, IMBNUMY, IMBNUMZ, IMBNUMX-, IMBNUMY-, IMBNUMZ- keywords.

Note

This keyword activates how relative permeability curves are assigned in the model. The ENDSCALE keyword allows the end-point scaling also to vary with direction, flow direction and hysteresis process. Resulting in a great deal of flexibility.

Whether or not all these features should be used though is another question.

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5.2.51 SAVE - ACTIVATES OUTPUT OF A SAVE FILE FOR FAST RESTARTS

RUNSP	EC GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates output of a SAVE file for fast restarts. There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

Example

WRITE OUT SAVE FILE FOR FAST RESTARTS

SAVE

The above example requests that a SAVE file be written out; however, this has no effect in OPM Flow input decks.

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5.2.52 SMRYDIMS - Define Maximum Number of Summary Vectors to be Written

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The SMRYDIMS keyword defines the maximum number of summary vectors to be written out to the SUMMARY file (*.SUMMARY).

OPM Flow users dynamic memory allocation and therefore the keyword has no effect and is ignored by the simulator, but is documented here for completeness.

No.	Name Description								
I	NSUMMX	A positive integer that defines the maximum number of summary vectors to be written out to the SUMMARY file (*.SUMMARY).	10000						
Notes:									
I)	The keyword is	terminated by "/".							

Table 5.24: SMRYDIMS Keyword Description

Example

-- SET THE MAXIMUM NUMBER OF SUMMARY VECTORS THAT CAN BE WRITTEN OUT
-- SMRYDIMS 10000 //

The above example sets maximum number of summary vectors that can be written out to the SUMMARY file to the default value of 10,000; however, this has no effect in OPM Flow input decks.

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5.2.53 SOLVENT - ACTIVATE THE SOLVENT PHASE IN THE MODEL

RUNSPEC GRID EDIT PRO	OPS REGIONS	SOLUTION	SUMMARY	SCHEDULE
-----------------------	-------------	----------	---------	----------

Revision: Rev-I

Description

This keyword indicates that the solvent phase is present in the model and to activate the four component solvent model for this run. In addition to this keyword, the oil, water and gases phases should also be declared for the run using the OIL, WATER and GAS keywords. The keyword will also invoke data input file checking to ensure that all the required Solvent phase input parameters are defined in the input deck.

There is no data required for this keyword.

Example

_ _

-- SOLVENT PHASE IS PRESENT IN THE RUN

- -

SOLVENT

The above example declares that the solvent phase is active in the model.

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5.2.54 START - SIMULATION START DATE

	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
- 1		-						

Revision: Rev-I

Description

This keyword sets the start date for the simulation switches. If the DATES keyword is to be used during the simulation, then a start date should be entered.

No.	Name	Description	Default
I	DAY	A positive integer that defines the day of the month, the value should be greater than or equal to one and less than or equal to 31.	None
2	MONTH	Character string for the month and should be one of the following 'JAN', 'FEB', 'MAR', 'APR', 'MAY', 'JUN', 'JUL' (or 'JLY'), 'AUG', 'SEP', 'OCT', 'NOV', or 'DEC'	None
3	YEAR	A positive four digit integer value of the start year, which must be specified fully by four digits, that is 1986.	None

Notes:

1) The keyword is terminated by "/".

Table 5.25: START Keyword Description

Example

-- DEFINE THE START DATE FOR THE RUN
-- START
01 'JAN' 2014

The above example sets the start date for the run to be January 1, 2014.

<u>Note</u>

Whenever possible it is a good idea to always set the start date to be at the beginning of the year as per the example. As like most simulators, OPM Flow reports are always stated at the number of days from the start date (and sometimes at a given date). If the start date is at the beginning of the year, then calculating the actual date is relatively straight forward and simple.

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5.2.55 TABDIMS - Define the Number of Tables and the Table Dimensions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The TABDIMS keyword defines the maximum number of tables for a given table type dataset and the maximum number of entries for the various tables. The commercial simulator combines both the "black-oil" and compositional simulator variables on this keyword; however, although all the parameters are explained below only the "black-oil" parameters are used by OPM Flow.

No.	Name	Description	Default			
I	NTSFUN	A positive integer that defines the maximum number of relative permeability table sets defined in the input deck. The tables are allocated to different parts of the grid by the SATNUM keyword.	I			
2	NTPVT	A positive integer that defines the maximum number of fluid property table sets defined in the input deck. The tables are allocated to different parts of the grid by the PVTNUM keyword.	I			
3	NSSFUN	A positive integer that defines the maximum number of saturation entries in the relative permeability tables defined in the input deck.	20			
4	NPPVT	A positive integer that defines the maximum number of pressure entries in the PVT tables.				
5	NTFIP A positive integer defining the maximum number of regions in the FIPNUM region array. Note that this parameter may also be set on the REGDIMS keyword as well. If NTFIP is set in both places then the maximum value is used.					
6	NRPVT	A positive integer that defines the maximum number of Rs and Rv entries in the PVT tables. If the DISGAS and VAPOIL options have notr been activated then this	20			
		parameter is ignored.				
7	NRVPVT	A positive integer that defines the maximum number of Rv entries in the PVT tables for the compositional commercial simulator.	 *			
8	NTENDP A positive integer that defines the maximum number of saturation endpoint depth tables. The end-point depth tables are used to re-scale the saturation tables as a function of depth as oppose to being a grid block property. NTENDP may also be specified on the ENDSCALE keyword, and if specified on both here and on the ENDSCALE keyword the maximum value of the two is used.		I			
9	NMEOSR	A positive integer that defines the maximum number of reservoir equations of states for the compositional commercial simulator.	I			
10	NMEOSS	A positive integer that defines the maximum number of separator or surface equations of states for the compositional commercial simulator.	I			
П	MXNFLN	A positive integer defining the maximum number flux regions in the FLUXNUM region array. MXNFLN can also be defined on the REGDIMS keywords as well. If MXNFLX is defined both here and on the REGDIMS keyword then the maximum value of the two is used.	10			
12	MXNTHR	A positive integer that defines the maximum number of thermal regions for the compositional commercial simulator.	I			

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No.	Name	Description	Default		
13	NTROCC A positive integer that defines the maximum number of rock compressibility entries enter by the ROCK keyword defined in the input deck. The tables are allocated to different parts of the grid by the PVTNUM keyword.				
14	MXNPMR	A positive integer that defines the maximum number of pressure maintenance regions for the compositional commercial simulator.	0		
15	NTABKT	A positive integer that defines the maximum number of temperature dependent K-value tables for the when the thermal option is activated in the compositional commercial simulator.	0		
16	NTALPHA	A positive integer that defines the maximum number of transport coefficient tables for the compositional commercial simulator.	0		
17	NASPKA	A positive integer that defines the maximum number of maximum number of entries in the ASPKDAM keyword tables for the compositional commercial simulator.	0		
18	MXRAWG	A positive integer that defines the maximum number of entries in the ASPREWG keyword tables for the compositional commercial simulator.	0		
19	MXRASO	A positive integer that defines the maximum number of pressure maintenance regions for the compositional commercial simulator.	0		
20		Not Used	 *		
21	MCASPP A positive integer that defines the maximum number of column entries in the ASPPW2D keyword tables for the compositional commercial simulator.		0		
22	MRASPP	A positive integer that defines the maximum number of row entries in the ASPPW2D keyword tables for the compositional commercial simulator.	0		
23	MXRATF	A positive integer that defines the maximum number of entries in the ASPWETF table for the compositional commercial simulator.	0		
24	MXNKVT	A positive integer that defines the maximum number of composition dependent K-value tables for the compositional commercial simulator.	0		
25	RESVED	Not Used	*		

Revision: Rev-I

Notes:

- Table sets are groups of keywords that need to be defined to complete a table set. For example if NTPVT is set to to three, then there must be three PVTO tables and three PVTG tables to complete the three PVT data set.
- 2) The keyword is terminated by "/".

Table 5.26:TABDIMS Keyword Description

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Examples

	MAX	MAX	MAX	MAX	MAX	MAX	E300	
	NTSFUN	NTPVT	NSSFUN	NPPVT	NTFIP	NRPVT	BLANK	NTEND
TABDIMS								
	15	a	40	30	1*	1*	1*	1

Revision: Rev-I

The above example defines maximum number of relative permeability tables to be 15 with a maximum number or rows of 40, and the maximum number of PVT tables to be nine with a maximum number or rows of 30.

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5.2.56 TEMP - ACTIVATE THE TEMPERATURE MODELING OPTION

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY S	SCHEDULE
----------------------------------------------------	----------

Revision: Rev-I

Description

This keyword activates the temperature modeling option. There is no data required for this keyword.

The temperature option (TEMP keyword) and the thermal option (THERMAL keyword) are two separate modeling facilities in the commercial simulator, although some keywords can be used by both options, for example the RTEMP keyword. OPM Flow's thermal implementation is based on solving the energy equation fully coupled with the "black-oil" equations so the results are not directly equivalent to the commercial simulator's "black-oil" TEMP or compositional THERMAL formulations. To activate OPM Flow's thermal implementation use the THERMAL keyword in the RUNSPEC section.

The reservoir temperature can be set to a constant or to be varying with depth utilizing the keywords in the following table:

Input Section	Constant Temperature	Temperature Variation With Depth		
PROPS	RTEMP and RTEMPA			
SOLUTION	RTEMP and RTEMPA	RTEMPVD and TEMPVD		

Notes:

1) The TEMP option is not implemented in OPM Flow; however, some of the above keywords can be used with OPM Flow's THERMAL option.

Table 5.27: Reservoir Temperature Keywords

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness. See the THERMAL keyword in the RUNSPEC section to activate OPM Flow's THERMAL option instead.

Example

ACTIVATE THE TEMPERATURE MODELING OPTION

TEMP

The above example activates the temperature modeling option, although the keyword is ignored by OPM Flow and has no effect on the simulation.

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5.2.57 THERMAL - ACTIVATE THE THERMAL MODELING OPTION

F	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates the thermal modeling option. There is no data required for this keyword. The energy "black-oil" implementation in OPM Flow is a mixture of the commercial simulators "black-oil" and the commercial simulators "compositional thermal" keywords, as well as some OPM Flow specific keywords.

The temperature option (TEMP keyword) and the thermal option (THERMAL keyword) are two separate modeling facilities in the commercial simulator, although some keywords can be used by both options, for example the RTEMP keyword. OPM Flow's thermal implementation is based on solving the energy equation fully coupled with the "black-oil" equations so the results are not directly equivalent to commercial simulator's "black-oil" TEMP or compositional THERMAL formulations.

Keywords specifically associated with both OPM Flow's THERMAL and the commercial simulators TEMP and THERMAL options are listed in Table 5.28 for easy of reference.

Section	Keyword	Function	OPM Flow	Commercial	
			THERMAL	TEMP	THERMAL
	HEATCR	Rock Heat Capacity.			
	HEATCRT	Rock Heat Capacity Temperature.			
	THCGAS	Gas Phase Thermal Conductivity.			
	THCOIL	Oil Phase Thermal Conductivity.			
GRID	THCONR	Thermal Conductivity of liquids and reservoir rock.			
	THCONSF	Thermal Conductivity of liquids and reservoir rock scaling factor applied to THCONR to account for gas saturation.			
	THCROCK	Rock Thermal Conductivity.			
	THCSOLID	Solid Phase Thermal Conductivity.			
	THCWATER	Water Thermal Conductivity.			
PROPS	GASDENT	Gas Density Temperature Coefficients (OPM Flow keyword).			
	GASVISCT	Gas Viscosity versus Temperature Functions (OPM Flow keyword).			
	OILDENT	Oil Density Temperature Coefficients (OPM Flow keyword).			
	OILVISCT	Oil Viscosity versus Temperature Functions (OPM Flow keyword).			
	RTEMP	Constant Initial Reservoir Temperature.			
	RTEMPA	Constant Initial Reservoir Temperature.			
	RTEMPVD	Initial Reservoir Temperature versus Depth.			
	TEMPVD	Initial Reservoir Temperature versus Depth.			
	SPECHEAT	Specific Heat of Oil, Water and Gas			
	SPECROCK	Specific Heat of the Reservoir Rock			

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S ection	Keyword	Function	OPM Flow	Commercial	
			THERMAL	TEMP	THERMAL
	WATDENT	Oil Density Temperature Coefficients.			
	WATVISCT	Oil Viscosity versus Temperature Function.			
REGION	THERMNUM	Thermal Region Numbers.			
	RTEMP	Constant Initial Reservoir Temperature.			
SOLUTION	RTEMPA	Constant Initial Reservoir Temperature.			
	TEMPI	Initial Reservoir Temperature for All Cells.			
SCHEDULE	WTEMP	Set An Injection Well's Fluid Temperature			
	WINJTEMP	Define Injection Fluid Thermal Properties			

Revision: Rev-I

Notes:

Table 5.28:THERMAL Option Associated Keywords

Example

- -

- ACTIVATE THE THERMAL MODELING OPTION

_ _

THERMAL

The above example activates the thermal modeling option.

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¹⁾ The list is focused on the OPM Flow implementation of the energy and "black-oil" formulation and therefore does not necessary include all the commercial simulator's keywords.

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5.2.58 TITLE - Define the Title for the Input Deck

GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE	PEC GRID EDIT	DIT PROPS REGIO	SOLUTION SUMMARY
---------------------------------------------------	---------------	-----------------	------------------

Revision: Rev-I

Description

The TITLE keyword defines the title for the input deck. The title text will be printed on all reports so as to act as a reference for the run.

No.	Name	Description	Default
I	TITLE	A character string that defines the TITLE for the input deck	None

Notes:

- All the characters on the line are processed as a string and therefore there is no need to enclose the TITLE in quotes.
- 2) There is no terminator '/' for the keyword.

Table 5.29:TITLE Keyword Description

Note

It is good practice to include the name of the input file in the tittle (without the extension) for when cross checking results from multiple cases.

Examples

```
-- DEFINE THE TITLE FOR THE RUN
TITLE
SPE01-THEM01-OPM1810-R01 - OPM THERMAL OPTION RUN
```

The above example defines the title for the run to be "SPE01-THEM01-OPM1810-R01 - OPM THERMAL OPTION RUN".

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5.2.59 TRACERS - ACTIVATE TRACER OPTIONS AND SET TRACER ARRAY DIMENSIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The TRACER keyword defines the number of tracers in the model and the various passive tracer tracking options.

No.	Name	Description	Default
I	MXOILTR	A positive integer defining the maximum number of passive oil tracers defined using the TRACER keyword.	0
2	MXWATTR	A positive integer defining the maximum number of passive water tracers defined using the TRACER keyword.	0
3	MXGASTR	A positive integer defining the maximum number of passive gas tracers defined using the TRACER keyword.	0
4	MXENVTR	A positive integer defining the maximum number of passive environmental tracers defined using the TRACER keyword.	0
5	DIFFOPT	A character string defining the numerical diffusion option for tracer tracking runs that should be set to:	NODIFF
		DIFF activates the numerical diffusion control options.	
		2) NODIFF deactivates the numerical diffusion control options.	
6	MXITRTR	A positive integer defining the maximum number of non-linear iterations to be used when the tracer option is activated.	12
7	MNITRTR	A positive integer defining the minimum number of non-linear iterations to be used when the tracer option is activated.	I
8	NONLIN	A character string stating if passive tracers as should be linear (NO) or non-linear (YES).	No
9	LNCONFAC	A real value defining the initial linear convergence factor. The default value of I* means the parameter will not be utilized.	*
10	NLCONFAC	A real value defining the initial non-linear convergence factor. The default value of I* means the parameter will not be utilized.	*
11	CONFAC	A real value defining the LNCONFAC and NLCONFAC convergence factors to be used after the initial convergence factor has been applied.	1.0
12	NUMCONF	A positive integer defining the maximum number of times CONFAC can be used.	0

Table 5.30:TRACERS Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Revision: Rev-I

Example

-- NO OIL NO WAT NO GAS NO ENV DIFF MAX MIN TRACER
-- TRACERS TRACERS TRACERS CONTL NONLIN NONLIN NONLIN
TRACERS

0 7 1 0 'NODIFF' 1* 1* 1* /

The above example defines seven tracers in the water phase and one tracer in the gas phase.

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5.2.60 UDADIMS - Define the Dimensions of the User Defined Arguments

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the dimensions of the User Defined Arguments ("UDA") used by OPM Flow that can be applied to various connection, group, and well keywords in the SCHEDULE section. UDAs are defined by the UDQ keyword that is used to specify values to be constants, SUMMARY variables, as defined in SUMMARY section, or a formula using various mathematical functions together with constants and SUMMARY variables.

Although this keyword is read by OPM Flow, the ACTION and UDQ computational logic and calculations have not been fully implemented and therefore this keyword should not be used as it may result in OPM Flow aborting.

No.	Name	Description	Default
ı	NMUDA	NMUDA is a positive integer that defines the number of augments in a keyword that are replaced by numeric UDQ values.	0
2	NULL	Not Used	*
3	MXUDA	MXUDA is a positive integer that defines the maximum number of <u>unique</u> <u>augments</u> in a keyword that are replaced numeric UDQ values.	100
		Note that MXUDA differs from NMUDA, for example:	
		 If only the oil rate argument of, say the WCONPROD keyword is specified by a UDQ, then both NMUDA and NMUDA equal one. 	
		 However, if a second WCONPROD uses a different UDQ, then NMUDA equals two, but MXUDA would can still be one. 	
		 Finally, if the same UDQ is used separately in two lines of WCONPROD data, then both NMUDA and MXUDA must be set to two. 	
		As MXUDA's default value is 100 then this only needs to be increased where the same UDQ is used as a UDA more than 100 times.	
Notes	<u>s:</u>		
I)	The keyword is	s terminated by "/".	

Table 5.31: UDADIMS Keyword Description

Note that OPM Flow has a more restricted UDQ feature set than the commercial simulator, so not all options and functions are available.

Example

```
USER DEFINED ARGUMENT DIMENSIONS
                          TOTAL
         NO.
                 NOT
         ARGS
                 USED
                          UDQ
UDADIMS
         10
                  1*
                          10
```

In the above example both NMUDA and MXUDA are set equal to ten.

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5.2.61 UDQDIMS - Define the Dimensions of the User Defined UDQ Feature

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

This keyword defines the dimensions associated with the UDQ keyword used in OPM Flow to calculate various user defined values in the SCHEDULE section. The UDQ keyword defined variables can be constants, SUMMARY variables, as defined in the SUMMARY section, or a formula using various mathematical functions together with constants and SUMMARY variables.

Although this keyword is read by OPM Flow, the ACTION and UDQ computational logic and calculations have not been fully implemented and therefore this keyword should not be used as it may result in OPM Flow aborting.

No.	Name	Description	Default
T	MXFUNS	A positive integer that defines the maximum number of functions that can be included when defining a UDQ definition. This should also include any brackets that will be used in the UDQ definition.	16
2	MXITEMS	MXITEMS is a positive integer that defines the maximum number of ITEMS allowed in an UDQ definition.	16
3	MXUDC	MXUDC is a positive integer that defines the maximum number of user defined CONNECTION quantities allowed in an UDQ definition.	0
4	MXUDF	MXUDF is a positive integer that defines the maximum number of user defined FIELD quantities allowed in an UDQ definition.	0
5	MXUDG	MXUDG is a positive integer that defines the maximum number of user defined GROUP quantities allowed in an UDQ definition.	0
6	MXUDR	MXUDR is a positive integer that defines the maximum number of user defined REGION quantities allowed in an UDQ definition.	0
7	MXUDS	MXUDS is a positive integer that defines the maximum number of user defined SEGMENT quantities allowed in an UDQ definition.	0
8	MXUDW	MXUDW is a positive integer that defines the maximum number of user defined WELL quantities allowed in an UDQ definition.	0
9	MXUDA	MXUDA is a positive integer that defines the maximum number of user defined AQUIFER quantities allowed in an UDQ definition.	0
10	MXUDB	MXUDB is a positive integer that defines the maximum number of user defined BLOCK quantities allowed in an UDQ definition.	0
П	RSEED	RSEED is a character string that determines if a new random number seed should be generated for restart runs for use in the UDQ functions RANDN, RANDU RRNDN and RRNDU. If RSEED is set to Y than a new seed will be generated and if set to the default value of N or I* then the same seed of the "base" simulation will be employed.	N
		See also the RSEED integer variable on the UDQPARAM keyword in the RUNSPEC section to set the random number seed for the current run.	
		This feature is not supported by OPM Flow.	

Table 5.32: UDQDIMS Keyword Description

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Note that OPM Flow has a more restricted UDQ feature set than the commercial simulator, so not all options and functions are available.

Revision: Rev-I

Example

	USER DE	FINED AR	GUMENT	DIMENSI	ONS FAC	ILITY						
	MAX	MAX	MAX	MAX	MAX	MAX	MAX	MAX	MAX	MAX	RA	ND
	FUNCS	ITEMS	CONNS	FIELD	GROUP	REGS	SEGTM	WELL	AQUF	BLCKS	0P	Т
UDQDIMS												
-	50	25	0	50	50	0	0	0	0	0	N	/

In this case the maximum number of functions that can be included when defining a UDQ definition is set to 50, maximum number of items allowed in an UDQ definition is 25, the maximum number of user defined field quantities allowed in an UDQ definition is 50, and the maximum number of user defined group quantities allowed in an UDQ definition is also 50. All other parameters are defaulted including the RSEED variable (the same seed of the "base" simulation will be employed).

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5.2.62 UDQPARAM - Define Parameters for the User Defined Quantity Feature

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the dimensions of the User Defined Arguments ("UDA") used by OPM Flow that can be applied to various connection, group, and well keywords in the SCHEDULE section. UDAs are defined by the UDQ keyword that is used to specify values to be constants, SUMMARY variables, as defined in SUMMARY section, or a formula using various mathematical functions together with constants and SUMMARY variables.

Although this keyword is read by OPM Flow, the ACTION and UDQ computational logic and calculations have not been fully implemented and therefore this keyword should not be used as it may result in OPM Flow aborting.

No.	Name	Description	Default
I	RSEED	RSEED is a positive integer greater than zero that sets a new random number seed for use in the UDQ functions RANDN, RANDU RRNDN and RRNDU.	1
		See also the RSEED character variable on the UDQDIMS keyword in the RUNSPEC section to default the random number seed for a restart run.	
		This feature is not supported by OPM Flow.	
2	RANGE	RANGE is areal positive value greater than or equal to one and less than or equal to 1.0×10^{20} , that sets the absolute range for user define quantities.	I × 10 ²⁰
		The default value of 1 \times 10 ²⁰ sets the range from -1 \times 10 ²⁰ to +1 \times 10 ²⁰ .	
3	DEFAULT	DEFAULT is real value that is the default numerical value given to undefined UDQ variables and should be in the same range as RANGE.	0.0
4	TOLUDQ	TOLUDQ a real positive number greater than zero and less than one that defines the tolerance used to determine if two real values are equal.	I x 10 ⁻⁴
		Floating-point numbers (as implemented in computers) are never exact, one cannot compare floating point numbers for exact equality. Thus, TOLUDQ defines a tolerance. For example, the default value of $I \times 10^{-4}$ means that if the difference between two real values is less than $I \times 10^{-4}$ then the values are considered equal.	

Table 5.33: UDQPARAM Keyword Description

Note that OPM Flow has a more restricted UDQ feature set than the commercial simulator, so not all options and functions are available.

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Revision: Rev-I

Example

```
-- USER DEFINED DEFAULT VALUES
-- SEED RANGE UNDEFINED COMPARISON
-- INTG -AND+ VALUE TOLERANCE
UDQPARAM
1 1.0E20 0.0 1.0E-4
```

The example explicitly sets the default values for all four variables on the UDAPARAM keyword, namely the random seed to one, the range to 1×10^{20} , the undefined UDQ variables to zero, and the comparison tolerance to 1.0×10^{-4} .

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5.2.63 UNIFIN - ACTIVATES THE UNIFIED INPUT FILE OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the Unified Input Files option for all input files. Similar to the commercial simulator, OPM Flow has various options for reading various input files and writing the resulting OPM Flow output files as described in Table 5.34.

Process	Option	Description	Files
Input	FMTIN	A character string that defines the input files to be formatted as ASCII i.e. text files, as oppose to binary files. The input deck file is always of this type. The option relates to the OPM Flow derived files that used as input, for for example when restarting from another case. If the keyword is omitted then the default is for binary file input.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART
	UNIFIN	A character string that defines the input files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. For this option a single summary file and a single restart file will be read. If the keyword is omitted then the default is for one file per report time step.	*.SMSPEC *.SUMMARY *.RSSPEC *. RESTART
Output	FMTOUT	A character string that sets all output files to be formatted as ASCII i.e. text files, as opposed to binary files. The *.PRT, *.LOG and *.DEBUG files are always of this type. The option relates to the OPM Flow output files only. In this case the files will be portable across operating systems, but will also be very large in terms of hard disk space. For this reason it is recommend that the default option is used so that binary files are outputted. If the keyword is omitted then the default is for binary file input.	*.SMSPEC *.SUMMARY *.GRID *.INIT *.RSSPEC *. RESTART
	UNIFOUT	A character string that defines the output files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. Here a single summary file and a single restart file will be generated, as opposed to one file per report time step. If the keyword is omitted then the default is for one file per report time step input.	*.SMSPEC *.SUMMARY *.RSSPEC *. RESTART

Notes:

- A binary file is computer-readable but not human-readable. All executable programs are stored in binary
 files, as are most numeric data files. In contrast, text files are stored in a form (usually ASCII) that is humanreadable.
- 2) For unified files if the run terminates unexpectedly, or there is sufficient disk space, then the last report output is not stored. Their main advantage is that if a number of simulations reside in one directory, their output is organized. There is no limit on the number of reporting steps that a unified file can store.

Table 5.34: UNIFIN Keyword Description

There is no data required for this keyword.

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Revision: Rev-I

Example

- -

SWITCH ON THE UNIFIED INPUT FILES OPTION

UNIFIN

The above example switches on the unified input file option.

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5.2.64 UNIFOUT - ACTIVATES THE UNIFIED OUTPUT FILE OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the Unified Output Files option for all output files. Similar to the commercial simulator, OPM Flow has various options for reading various input files and writing the resulting OPM Flow output files as described in Table 5.35.

Process	Option	Description	Files
Input	FMTIN	A character string that defines the input files to be formatted as ASCII i.e. text files, as oppose to binary files. The input deck file is always of this type. The option relates to the OPM Flow derived files that used as input, for for example when restarting from another case. If the keyword is omitted then the default is for binary file input.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART
	UNIFIN	A character string that defines the input files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. For this option a single summary file and a single restart file will be read. If the keyword is omitted then the default is for one file per report time step.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART
Output	FMTOUT	A character string that sets all output files to be formatted as ASCII i.e. text files, as opposed to binary files. The *.PRT, *.LOG and *.DEBUG files are always of this type. The option relates to the OPM Flow output files only. In this case the files will be portable across operating systems, but will also be very large in terms of hard disk space. For this reason it is recommend that the default option is used so that binary files are outputted. If the keyword is omitted then the default is for binary file input.	*.SMSPEC *.SUMMARY *.GRID *.INIT *.RSSPEC *.RESTART
	UNIFOUT	A character string that defines the output files to be unified files, as opposed to non-unified multiple files. A unified file is a single file containing output for each reporting time step. Here a single summary file and a single restart file will be generated, as opposed to one file per report time step. If the keyword is omitted then the default is for one file per report time step input.	*.SMSPEC *.SUMMARY *.RSSPEC *.RESTART

Notes:

- A binary file is computer-readable but not human-readable. All executable programs are stored in binary files, as are most numeric data files. In contrast, text files are stored in a form (usually ASCII) that is humanreadable.
- 2) For unified files if the run terminates unexpectedly, or there is sufficient disk space, then the last report output is not stored. Their main advantage is that if a number of simulations reside in one directory, their output is organized. There is no limit on the number of reporting steps that a unified file can store.

Table 5.35: UNIFOUT Keyword Description

There is no data required for this keyword.

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Revision: Rev-I

Example

- -

SWITCH ON THE UNIFIED OUTPUT FILES OPTION

--

UNIFOUT

The above example switches on the unified output file option.

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5.2.65 VAPOIL - ACTIVATE THE VAPORIZE OIL IN WET GAS PHASE IN THE MODEL

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
-----------------------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword indicates that vaporized oil (more commonly referred to as condensate) is present in wet²⁰ gas in the model and the keyword should only be used if the there is both oil and gas phases in the model. The keyword may be used for gas-water and oil-water-gas input decks that contain the oil and gas phases. The keyword will also invoke data input file checking to ensure that all the required oil and gas phase input parameters are defined in the input deck.

If the gas has a constant and uniform vaporized oil concentration, Condensate-Gas Ratio ("CGR"), and if the reservoir pressure never drops below the saturation pressure pressure (dew point pressure), then the model can be run more efficiently by omitting the OIL and VAPOIL keywords from the RUNSPEC section, treating the gas as a dry gas²¹, and defining a constant Rv (CGR) value with keyword RVCONST or RVCONSTT in the PROPS section. This results in the model being run with as a dry gas problem with no active oil (condensate) phase. However, OPM Flow takes into account the constant Rv in the calculations and reporting.

There is no data required for this keyword.

Example

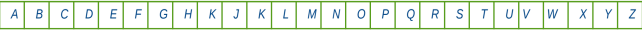
- -

VAPORIZED OIL IN WET GAS IS PRESENT IN THE RUN

VAP0IL

The above example declares that the vaporized oil, i.e. condensate, in the gas phase is active in the model.

Natural gas that occurs in the absence of condensate or liquid hydrocarbons, or gas that had condensable hydrocarbons removed, is called dry gas. It is primarily methane with some intermediates. The hydrocarbon mixture is solely gas in the reservoir and there is no liquid (condensate surface liquid) formed either in the reservoir or at surface. The term dry indicates that the gas does not contain heavier hydrocarbons to form liquids at the surface conditions. Dry gas typically has GOR's greater than 100,000 scf/stb or 18,000 Sm3/m3.



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Natural gas that contains significant heavy hydrocarbons such as propane, butane and other liquid hydrocarbons is known as wet gas or rich gas. The general rule of thumb is if the gas contains less methane (typically less than 85% methane) and more ethane, and other more complex hydrocarbons, it is labeled as wet gas. Wet gas normally has GOR's less than 100,000 scf/stb or 18,000 Sm3/m3, with the condensate having a gravity greater than 50 °API.

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5.2.66 VFPIDIMS - INJECTION VERTICAL FLOW PERFORMANCE TABLE DIMENSIONS

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
-----------------------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

VFPIDIMS keyword defines the maximum dimensions of the injection well Vertical Lift Performance ("VFP") tables defined by VFPINJ keyword. The VFP tables for the producing wells are defined by the VFPPDIMS keyword.

No.	Name	Description	Default
I	MXMFLO	A positive integer that defines the maximum number of injection rate entries for the VFPINJ keyword.	0
2	MXMTHP	A positive integer that defines the maximum number of THP entries for the VFPINJ keyword.	0
3	MXVFPTAB	A positive integer that defines the maximum number of VFPINJ tables entered through the VFPINJ keyword.	0

Notes:

Table 5.36:VFPIDIMS Keyword Description

Example

```
INJECTING VFP TABLES
         VFP
                 VFP
                          VFP
         MXMFLO MXMTHP
                         NMMVFT
VFPIDIMS
         10
                 10
                          12
```

The above example defines that the maximum number of injection rates and THP entries on the VFPINJ

keyword is 10, and the maximum number of BHP entries is 12.

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¹⁾ The keyword is terminated by "/".

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5.2.67 VFPPDIMS - PRODUCTION VERTICAL FLOW PERFORMANCE TABLE DIMENSIONS

RUNSPEC GRID EDIT PRO	OPS REGIONS	SOLUTION	SUMMARY	SCHEDULE
-----------------------	-------------	----------	---------	----------

Revision: Rev-I

Description

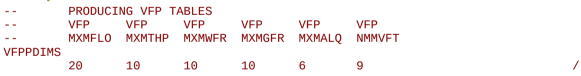
VFPPDIMS keyword defines the maximum dimensions of the production well Vertical Lift Performance ("VFP") tables defined by VFPPROD keyword. The VFP tables for the injection wells are defined by the VFPIDIMS keyword.

No.	Name	Description	Default
I	MXMFLO	A positive integer that defines the maximum number of production flow rate entries for the VFPPROD keyword.	0
2	MXMTHP	A positive integer that defines the maximum number of THP entries for the VFPPROD keyword.	0
3	MXMWFR	A positive integer that defines the maximum number of water fraction entries (WOR, WCUT, GWR etc.) entries for the VFPPROD keyword.	0
4	MXMGFR	A positive integer that defines the maximum number of gas fraction entries (GOR, GLR, OGR etc.) entries for the VFPPROD keyword.	0
5	MXMALQ	A positive integer that defines the maximum number of artificial lift quantity entries entries for the VFPPROD keyword.	0
6	MXVFPTAB	A positive integer that defines the maximum number of VFPPROD tables entered through the VFPPROD keyword.	0

Notes:

Table 5.37:VFPPDIMS Keyword Description

Example



Here the example shows that there are a maximum of 20 flow rates, 10 THP entries, 10 water and gas fraction entries, and six artificial lift entries for the nine VFPPROD VFP production tables.

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¹⁾ The keyword is terminated by "/".

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5.2.68 WATER - ACTIVATE THE WATER PHASE IN THE MODEL

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Revision: Rev-I

Description

This keyword indicate that the water phase is present in the model and must be used for gas-water, oil-gas, oil-water, oil-water-gas input decks that contain the water phase. The keyword will also invoke data input file checking to ensure that all the required water phase input parameters are defined in the input deck.

There is no data required for this keyword.

Example

WATER PHASE IS PRESENT IN THE RUN

WATER

The above example declares that the oil phase is active in the model.

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5.2.69 WELLDIMS - DEFINE THE WELLS AND GROUP DIMENSIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

WELLDIMS defines various well and group dimensions for the run. The commercial simulator combines both the "black-oil" and compositional simulator variables on this keyword; however, although all the parameters are explained below only the "black-oil" parameters are used by OPM Flow.

No.	Name	Description	Default
I	MXWELS	A positive integer defining the maximum number of wells for this model.	0
2	MXCONS	A positive integer defining the maximum number of grid block connections per well for this model.	0
3	MXGRPS	A positive integer defining the maximum number of groups for this model.	0
4	MXGRPW	A positive integer defining the maximum number of wells that can belong to a group in the model.	0
5	MXSTAGE	A positive integer defining the maximum number of stages per separator for this model.	5
		This option is ignored by OPM Flow.	
6	MXSTRMS	A positive integer defining the maximum number of well streams for this model.	10
		This option is ignored by OPM Flow.	
7	MXMIXS	A positive integer defining the maximum number of mixtures for this model.	5
		This option is ignored by OPM Flow.	
8	MXSEPS	A positive integer defining the maximum number of separators for this model.	4
		This option is ignored by OPM Flow.	
9	MXCOMPS	A positive integer defining the maximum number of mixture components in a mixture for the model.	3
		This option is ignored by OPM Flow.	
10	MXDOCOMP	A positive integer defining the maximum number of well completions that can cross a parallel run domain boundary when the PARALLEL option has been activated.	0
		This option is ignored by OPM Flow.	
11	MXWSLIST	A positive integer defining the maximum number of well lists that a well may be concurrent belong to at one time for this model.	I
		This option is ignored by OPM Flow.	
12	MXWLISTS	A positive integer defining the maximum number of dynamic well lists for this model.	I
		This option is ignored by OPM Flow.	
13	MXWSECD	A positive integer defining the maximum number of secondary wells for this model.	10
		This option is ignored by OPM Flow.	

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No.	Name	Description	Default
14	MXNGPP	A positive integer defining the maximum number of entries per completion in the generalized pseudo-pressure tables used for to calculate the blocking factor associated with condensate drop-out in gas condensate reservoirs.	201
		If the generalized pseudo-pressure option has not been activated then this is ignored.	
		This option is ignored by OPM Flow.	

Revision: Rev-I

Notes:

- 1) Only parameters (1) to (4) are used by OPM Flow.
- 2) The keyword is terminated by "/".

Table 5.38:WELLDIMS Keyword Description

Example

-- WELL WELL GRUPS GRUPS
-- MXWELS MXCONS MXGRPS MXGRPW
WELLDIMS
60 110 18 40

The above example defines the maximum number of wells to be 60 with 110 completions per well, and maximum number of groups to be 18 with maximum number of wells per group of 40. All other parameters are defaulted.

A B C D E F G H K J K L M N O P Q R S T U V W X Y Z

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5.2.70 WSEGDIMS - Define Multi-Segment Well Dimensions

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
-----------------------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The WSEGSDIMS keyword defines the multi-segment well dimensions for the multi-segment well model and the keyword is obligatory if multi-segment wells are being employed in the model.

No.	Name	Description	Default
I	MXWELS	A positive integer defining the maximum number of multi segment wells for this model.	0
2	MXSEGS	A positive integer defining the maximum number of segments per well for this model.	I
3	MXBRAN	A positive integer defining the maximum number of branches per multi- segment well, including the main branch.groups for this model.	I
4	MXLINKS	A positive integer defining the maximum number of segment links per multi-segment well.	0

Table 5.39:WESEGDIMS Keyword Description

Example

```
WELL
                 WELL
                          BRANCH
                                  SEGMENT
         MXWELS MXSEGS
                         MXBRAN
                                  MXLINKS
WSEGDIMS
                 100
                          10
                                  10
```

The above example defines the maximum number of multi-segment wells to be five with up to 100 segments per multi-segment well, a maximum number of 10 branches per multi-segment well, and up to 10 segment links per multi-segment well.

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The keyword is terminated by "/".

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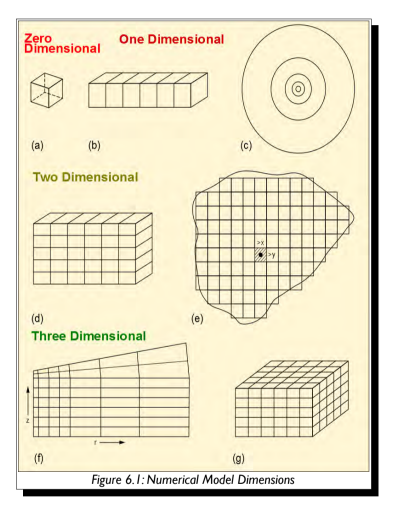
6 GRID SECTION

6.1 Introduction

The GRID section defines the basic grid properties, including structure, faults and various static rock properties (porosity, permeability etc.). The information in this section will be used by the software to calculate the pore volume (PORV) for each cell, the cell mid-point depths, and the regular transmissibilities (TRANX, TRANY and TRANX) between all the cells, as well as across faults. The OPM Flow calculated parameters can then be edited in the EDIT section.

All models can be classified by the number of dimensions as show in Figure 6.1 (after Mattax²²). The zero and one dimension models are employed in analytical modeling, while the higher dimensions are used in numerical modeling. The term 4D modeling refers to a 3D model with the fourth dimension being the time domain derived from time-lapse seismic, that is the comparison of 3D seismic surveys at two or more points in time.

OPM Flow enables the user to define ID, 2D and 3D models using three types of grids: Cartesian Regular Grid, Radial Grid²³, and Irregular Corner-Point Grids. The first two type of grids are rather limited in their ability to describe the structural complexity of oil and gas reservoirs; however, this simplicity allows the engineer to quickly build simple models to investigate reservoir performance. Indeed in the early days of numerical modeling back in the late 1970's twodimensional cross-section and radial models were the main models used to predict reservoir performance due to limited computer resources at the time. That is not to say that full field models were not developed, but that



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these full field models were very coarse in comparison to what is designed and built today using static earth modeling software.

A brief introduction to the three types of grids and the data requirements to fully defined the structural element of the grid together with the rock properties necessary to complete the GRID section data requirements is outlined in the following section. This is then followed by the keyword definitions applicable to this section.

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Mattax, C.C. and Dalton R.L. 1990. Reservoir Simulation. Society of Petroleum Engineers, Henry L. Doherty Series, Monograph Vol. 13

²³ Radial grids are not currently implemented in OPM Flow version 2018-04 but is expected to be incorporated in the next release.

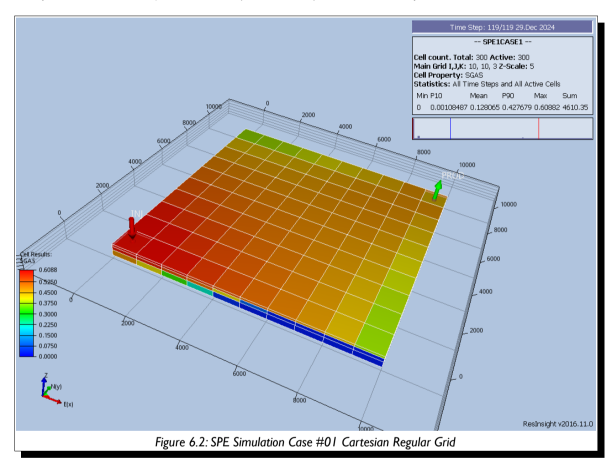
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6.2 DATA REQUIREMENTS

6.2.1 CARTESIAN REGULAR GRID

This type of grid defines a regular orthogonal grid based on defining the x, y and z dimensions of all the cells and is normally employed when a complex structural model is not required. Figure 6.2 shows the SPE Comparative Solution Project Number I ("SPE-CSP01") as documented by Odeh²⁴.

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The model consists of a simple $10 \times 10 \times 3$ (NX, NY, NY) grid and is defined using the following GRID section keywords to define the grid geometry:

```
-- DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX x NY x NZ = 300)
-- (There Are In Total 300 Cells With Length 1000ft In X-Direction)

DX

300*1000

-- DEFINE GRID BLOCK Y DIRECTION CELL SIZE (BASED ON NX x NY x NZ = 300)
-- (There Are In Total 300 Cells With Length 1000ft In Y-Direction)

DY

300*1000

-- DEFINE GRID BLOCK Z DIRECTION CELL SIZE (BASED ON NX x NY x NZ = 300)
-- (The Layers Are 20, 30 And 50 Ft Thick, In Each Layer There Are 100 Cells)

DZ

100*20.0 100*30.0 100*50.0
```

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Odeh, A. "Comparison of Solutions to a Three Dimensional Black-Oil Reservoir Simulation Problem." JPT 33 (1981):13-25.

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```
DEFINE GRID BLOCK TOPS FOR THE TOP LAYER (BASED ON NX = 100, NY = 100)
   (Layer 2 and 3 TOPS Calculated by Simulator)
TOPS
25*3100 25*3105 25*3110
The rock property data required to complete the GRID section is as follows:
   DEFINE GRID BLOCK POROSITY DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
- -
POR0
300*0.300
-- DEFINE GRID BLOCK PERMY DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
PFRMX
100*500.0
              100%50.0
                         100*200.0
-- DEFINE GRID BLOCK PERMY DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
PFRMY
100*500.0
              100%50.0
                         100*200.0
-- DEFINE GRID BLOCK PERMZ DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- (Not Defined in Original Paper So Assume That PERMX = PERMY = PERMZ)
PERMZ
 100*500.0
              100%50.0
                         100*200.0
```

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The above keywords define all the properties required for the GRID section for this type of grid geometry.

6.2.2 RADIAL GRID

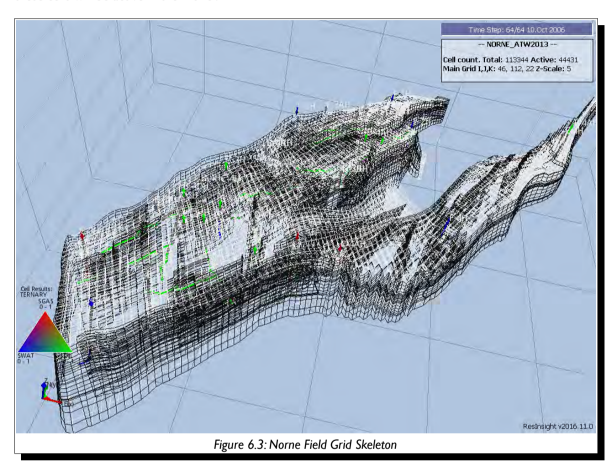
To be written in a future release of the manual.

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6.2.3 IRREGULAR CORNER-POINT GRIDS

This type of grid is an industry standard grid used to formulate the structure of complex reservoirs. Here static modeling software is used to build the model which is then exported and imported into a numerical model. Figure 6.3 illustrates the skeleton grid for the Norne Field which has dimensions of $46 \times 112 \times 22$ in the x, y and z dimensions respectively. This results in a total number of cells of 113,344 although not all of these cells will be active in the model.

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Similar to Cartesian Regular Grid the grid geometry must be defined for each cell and the properties for each cell defined. The formulation of the grid geometry is based on *corner-point geometry*, basically coordinate lines or pillars are given, then top and bottom surfaces for the cell are given by specifying the depth (z-coordinates) of the cell's corner points along each of the four adjacent pillars. The cell then forms an irregular hexahedron as depicted in Figure 6.4. Note that the figure shows a corner-point cell which is more or less orthogonal, which is ideally is what we want to minimize grid orientation effects.

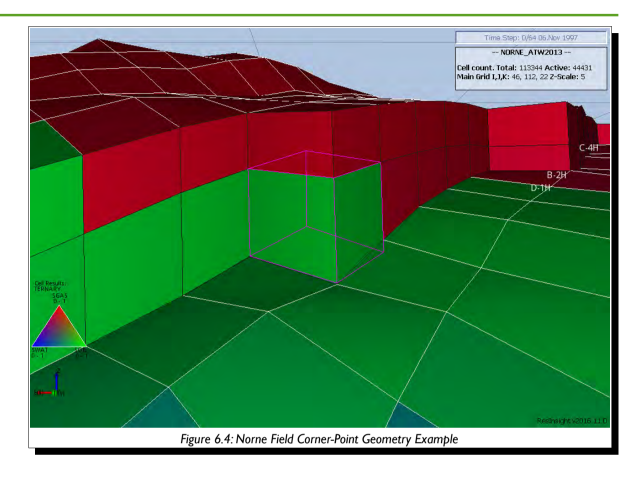
The data required to define this type of grid consists of the SPECGRID to define the dimensions of the grid, that is:

	MAX	MAX	MAX	MAX	GRID
	NDIVIX	NDIVIY	NDIVIZ	NUMRES	TYPE
SPECGRID	46	112	22	1	F

A portion of the coordinate line data defined by the COORD keyword from the Norne model is shown on the next page.

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COORD					
X1	Y1	Z1	X2	Y2	Z2
453114.000	7319921.000	3037.473	453114.000	7319921.000	3132.831
453155.031	7319840.000	2983.933	453142.750	7319864.000	3173.572
453196.094	7319759.000	3005.969	453171.500	7319807.500	3215.836
453237.156	7319678.000	3000.265	453200.250	7319751.000	3217.250
453278.188	7319597.000	2989.348	453229.031	7319694.000	3213.951
453319.250	7319516.500	2995.680	453257.781	7319637.500	3215.323
453356.250	7319443.500	3000.855	453308.750	7319537.000	3220.549
453393.250	7319370.500	3005.252	453359.688	7319436.500	3210.393
453423.969	7319310.000	3030.862	453394.219	7319368.500	3203.438
453454.688	7319249.500	3036.870	453428.719	7319300.500	3190.770
453485.406	7319189.000	3038.017	453463.219	7319232.500	3190.660
453516.125	7319128.000	3045.027	453497.750	7319164.500	3188.813
453546.844	7319067.500	3055.410	453532.250	7319096.500	3185.966
453577.562	7319007.000	3066.541	453566.750	7319028.500	3184.325
453608.281	7318946.500	3076.624	453601.250	7318960.500	3183.584
453639.000	7318886.000	3086.938	453635.781	7318892.500	3184.057
453669.719	7318825.500	3096.153	453670.281	7318824.500	3185.988
453700.438	7318765.000	3104.703	453704.781	7318756.500	3188.598
453731.156	7318704.500	3097.016	453739.281	7318688.500	3180.484
453761.875	7318644.000	3088.539	453773.812	7318620.500	3177.091
453780.000	7318608.000	3098.118	453796.562	7318575.500	3176.401
453798.125	7318572.500	3096.691	453819.344	7318530.500	3172.299

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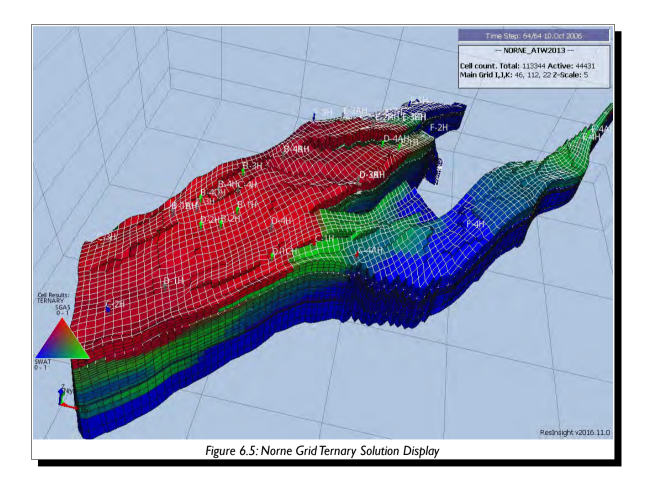
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The final keyword to define an Irregular Corner-Point geometry grid is the ZCORN keyword that defines the depths of the cell corners. A portion of the ZCORN data from the Norne model is shown below.

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ZCORN					
3037.473	2983.933	2983.933	3005.969	3005.969	3000.265
3000.265	2989.348	2989.348	2995.680	2995.680	3000.855
3000.855	3005.252	3005.252	3030.862	3030.862	3036.870
3036.870	3038.017	3038.017	3045.027	3045.027	3055.410
3055.410	3066.541	3066.541	3076.624	3076.624	3086.938
3086.938	3096.153	3096.153	3104.703	3104.703	3097.016
3097.016	3088.539	3088.539	3098.118	3098.118	3096.691
3096.691	3093.886	3093.886	3085.393	3085.393	3081.957
3081.957	3080.645	3080.645	3115.021	3115.021	3130.474
3130.474	3204.674	3204.674	3193.187	3193.187	3169.512
3169.512	3101.928	3101.928	3044.277	3044.277	3023.930
3023.930	2964.244	2964.244	2900.178	2900.178	2875.715
2875.715	2864.913	2864.913	2855.256	2855.256	2841.119
2841.119	2826.261	2826.261	2806.556	2806.556	2781.052
2781.052	2791.720	2791.720	2817.940	2817.940	2813.308
2813.308	2788.492				

The rock property data required to complete the GRID section is the same as for a Cartesian Regular grid, as defined in **Section 6.2.1** and the data is defined using the same keywords. The resulting Norne model showing the ternary solution variable is displayed in Figure 6.5.





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6.2.4 ROCK PROPERTIES

Irrespective of the grid type used to define the structural component of the model various static properties need to be defined in order for the model to have a complete grid definition, these properties include the identification of active and inactive grid blocks, porosity, permeability, and the reservoir quality via the net-to-gross fraction ("NTG"). These parameters must be set for each cell in the model

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Property	Description	Cartesian And Irregular Corner-Point Grids Keywords	Radial Grid Keywords
Active and Inactive cells	Defines if a cell in the model is active by setting the ACTNUM property for a cell to either one or inactive by setting the value to zero.	ACTN	ML
	Cells that are inactive in the model are ignored computationally and can act as barriers to flow. Thus, a shale in a conventional reservoir is normally treated as non-reservoir and is made inactive either by setting the ACTNUM, PORO, or NTG to zero for the cells representing the shale.		
Porosity	Porosity is a measure of the space in a reservoir rock. It is defined as the fraction of the total bulk volume of the rock not occupied by solids, that is it is the fraction of the cell that is porous and contains the reservoir fluids.	PORC)
Reservoir Quality	Reservoir quality of the cell in terms of the gross volume derived from the structural grid and the net volume available for fluid flow in the model expressed as a fraction from zero to one. A zero values means the cell does contribute to flow and therefore is made inactive. A value of one means the gross and net volumes are identical for the cell	NTG	
Permeability	Permeability is a measure of the ease with which a fluid will flow through a porous medium. In numerical models permeability is dependent on the direction of flow, that is x, y and z directions in Cartesian and Irregular Corner-Point Grids, and the radial, theta and z directions in radial grids.	PERMX PERMY PERMZ	PERMR PERMTHT PERMZ
	There are various formulations for permeability, absolute permeability, effective permeability, gas permeability, liquid permeability etc. and the values are saturation dependent.		
	Thus, values entered should be consistent with the relative permeability entered in the PROPS section. Normally Kair (Sg=1.0) should be entered for the cell permeability and the values may or not be corrected for overburden or humidity drying effects. Correcting for liquid flow and saturation end points etc., is accomplished by the relative permeability curves.		
	For example, if Kair (Sg=1.0) has been entered for the cell permeability when Krg (Sg=1-Swc) should be less than one.		

Table 6.1: Key Static Grid Properties



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Note

Static grid properties are frequently generated from a static earth model using petrophysical evaluation of the well logs and propagated through the model based on a variety of geostatistical techniques.

Petrophysical evaluations are conducted in either in the "Total" or the "Effective" porosity domain, and is important that all the rock property data is entered into the model is of the same basis. It is not important which porosity domain is used, as long as all the data is in the same domain.

Pore volume and transmissibility are common terms in the reservoir simulation vernacular. Pore volume is self-explanatory, that is, given the grid property data the pore volume for each cell is calculated using:

$$PV = Cell\ Gross\ Volume \times PORO \times NTG \times ACTNUM$$
 (6.1)

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Where

PV = the pore volume of a cell,

Cell Gross Volume = the gross volume (or bulk volume) calculated from the structural

parameters of the cell,

PORO = cell porosity,

NTG = cell net-to-gross ratio, and ACTNUM = active and inactive cell indicator.

Any cell with a pore volume equal to zero is made inactive automatically in the model. However, there may be some cells that have small pore volumes than may negatively impact computational performance of the model. If this is the case then the MINPV keyword in the GRID section can be used to make these cells inactive.

There has been a trend in the industry in recent years to not apply petrophysical cut-offs in static models. This results in large models with numerous cells with very low porosity values (less than 0.01 for example) and corresponding very low permeabilities. The theory behind this approach is that the numerical model will determine the effective (or net) reservoir. This may be appropriate in unconventional reservoirs as all the cells in the model will have similar values of porosity and permeability, but in conventional reservoirs this methodology will lead to severe computational issues when attempting to run the model due to very tight cells being next to relative high permeability cells. Again, the MINPV keyword can be used to resolve this issue.

Transmissibility on the other hand is more complex as it relates the flow from one cell face to another cell face and is a function of the area open to flow, the direction of flow, the permeability, saturation and viscosity of the phases flowing between the cells. For a single phase flow in a Cartesian grid the x-direction transmissibility is of the form:

$$T_{x_{i+1/2, j}} = \left[\frac{k_x h \left(\Delta y\right)}{\mu \left(\Delta x\right)}\right]_{i+1/2, j}$$
(6.2)

As transmissibility is a property of the flow between two cell faces, not a block centered grid cell property like porosity or permeability, then the nomenclature for transmissibility is different. In OPM Flow, the transmissible of cell face $T_x(i, j, k)$ is the transmissibility between cells (i, j, k) and (i+1, j, k). In some simulators it would be between (i, j, k) and (i-1, j, k). This is important to note if manual modifications to cell connections are to be made in the model.

Note that modifications to grid property data can only be done in the GRID section, thereafter only the calculated pore volumes and transmissibilities are available for adjustment.

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6.3 Keyword Definitions

6.3.1 ACTNUM - SET THE STATUS OF A GRID BLOCK TO ACTIVE OR INACTIVE

RUNS	SPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
------	------	------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

ACTNUM specifies which grid blocks are active or inactive. A value of I indicates the block is active and a value of 0 indicates the block is inactive. A grid block is inactive if its pore volume is less than the value entered using keyword MINPV. This keyword can be used to make blocks with a pore volume greater than MINPV inactive. Note that a value must be entered for each grid block in the model. Grid blocks are ordered with the I index cycling fastest, followed by the J and K indices. As for all array data repeat counts may be used, for example 100*I; however the full array must be specified.

Note that the a cells activity can also be set using the EQUALS keyword by selection only those cells that are required to be made inactive.

No.	Name	Description	Default
I	ACTNUM	An array of integers that define the activity of a cell by setting it to 1 for being active or 0 for inactive, for each grid block in the model.	 *

Notes:

- I) A total of NX \times NY \times NZ integer values, as defined by the DIMENS in the RUNSPEC section, must be specified for the array.
- 2) The keyword is terminated by "/".

Table 6.2:ACTNUM Keyword Description

Examples

The example below sets several cells to be inactive for a $4 \times 5 \times 2$ model.

ACTNUM

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

```
CONSTANT --
                         ----- BOX -----
-- -- ARRAY
                         I1 I2 J1 J2
                                          K1 K2
EQUALS
                         1*
                                 1*
                                     1*
                                          1* 1* / SET ACTIVE CELLS
   'ACTNUM'
             1.0000
  'ACTNUM'
                                            1 / SET INACTIVE CELLS
             0.0000
                             2
                                 1
                                    2
                                          1
                         1
  'ACTNUM'
                                              2 / SET INACTIVE CELLS
             0.0000
                                     4
```

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6.3.2 ADD - ADD A CONSTANT TO A SPECIFIED ARRAY

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The ADD keyword adds a constant to a specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the ADD keyword is being applied.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	An integer or real value to be added to the ARRAY in the same units as the ARRAY property.	0
3	II	A positive integer that defines the lower bound of the array in the I-direction to be modified must be greater than or equal I and less than or equal to I2 and NX.	I
4	12	A positive integer that defines the upper bound of the array in the I-direction to be modified must be greater than or equal to II and less than or equal to NX	NX
5	JI	A positive integer that defines the lower bound of the array in the J-direction to be modified must be greater than or equal I and less than or equal to J2 and NY.	I
6	J2	A positive integer that defines the upper bound of the array in the J-direction to be modified must be greater than or equal to JI and less than or equal to NY.	NY
7	KI	A positive integer that defines the lower bound of the array in the K-direction to be modified must be greater than or equal to one and less than or equal to K2 and NZ.	I
8	K2	A positive integer that defines the upper bound of the array in the K-direction to be modified must be greater than or equal to KI and less than or equal to NZ.	NZ

Notes:

- Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.3:ADD Keyword Description

The applicable arrays for each section are defined in Table 6.4 on the following page.



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		ADD Keyword	d and Variable O	ptions by Sectio	n	
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
DX	DEPTH	SWL	ENDNUM	PRESSURE		
DY	PORV	SWCR	EQLNUM	SWAT		
DZ	TRANX	SWU	FIPNUM	SGAS		
PERMX	TRANY	SGL	IMBNUM	RV		
PERMY	TRANZ	SGCR	MISCNUM	RS		
PERMZ	DIFFX	SGU	PVTNUM	TBLK		
MULTX	DIFFY	KRW	ROCKNUM	GI		
MULTY	DIFFZ	KRO	SATNUM	OILAPI		
MULTZ	TRANR	KRG	WH2NUM	SALT		
DR	TRANTHT	PCG		GASCONC		
THETA	DIFFR	PCW		SOLVCONC		
PERMR	DIFFTHT			SOLVFRAC		
PERMTHT				SFOAM		
DZNET				SPOLY		
PORO						
NTG						
FLUXNUM						
MULTNUM						
MPANUM						
DIFFX						
DIFFY						
DIFFZ						
DIFFR						
DIFFTHT						

Table 6.4:ADD Keyword Applicable Arrays by Section

Example

The above example ADDS 20 units to the PERMX array in the GRID section to all grid blocks in the model.

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6.3.3 ADDREG - ADD A CONSTANT TO AN ARRAY BASED ON A REGION NUMBER

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

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Description

The ADDREG keyword adds a constant to a specified array or part of an array based on cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the ADDREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the ADDREG keyword is being used.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	An integer or real value to be added to the ARRAY in the same units as the ARRAY property for a given REGION	0
3	REGION NUMBER	REGION NUMBER is a positive integer representing the region for which the CONSTANT in (2) should be applied	None
4	REGION ARRAY	The REGION ARRAY to use for applying the CONSTANT in (2) based on the REGION NUMBER in (3). REGION ARRAY can have the following values:	М
		F for the FLUXNUM array	
		2) M for the MULTNUM array	
		3) O for the OPERNUM array	

Notes:

- Where the REGION NUMBER should be less than or equal to the maximum number of regions as defined on the REGDIMS keyword for the FIPNUM and OPERNUM arrays or the GRIDOPTS keyword for the MULTNUM array in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.5:ADDREG Keyword Description

The applicable arrays for each section are defined in Table 6.6 on the following page.

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	Al	DDREG Keywo	ord and Variable	Options by Sec	tion	
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
DX	DEPTH	SWL	ENDNUM	PRESSURE		
DY	PORV	SWCR	EQLNUM	SWAT		
DZ	TRANX	SWU	FIPNUM	SGAS		
PERMX	TRANY	SGL	IMBNUM	RV		
PERMY	TRANZ	SGCR	MISCNUM	RS		
PERMZ	DIFFX	SGU	PVTNUM	TBLK		
MULTX	DIFFY	KRW	ROCKNUM	GI		
MULTY	DIFFZ	KRO	SATNUM	OILAPI		
MULTZ	TRANR	KRG	WH2NUM	SALT		
DR	TRANTHT	PCG		GASCONC		
THETA	DIFFR	PCW		SOLVCONC		
PERMR	DIFFTHT			SOLVFRAC		
PERMTHT				SFOAM		
DZNET				SPOLY		
PORO						
NTG						
FLUXNUM						
MULTNUM						
MPANUM						
DIFFX						
DIFFY						
DIFFZ						
DIFFR						
DIFFTHT						

Table 6.6: ADDREG Keyword Applicable Arrays by Section

Examples

```
-- FIRST DEFINE THE PROPERTY ARRAYS AND MULTNUM ARRAYS FOR 10 X 10 X 20 MODEL
-- -- ARRAY CONSTANT -- ------ BOX -----
                          I1 I2 J1 J2 K1 K2
EQUALS
                                   1*
1*
                                                   / PORO TO 0.20 IN MODEL
/ PERMX TO 0.10 IN MODEL
   'PORO'
              0.2000
                           1*
                              1*
                                       1*
                                            1* 1*
                                            1*
   'PERMX'
                           1*
                             1*
                                       1*
                                               1*
              100.00
                                       1*
                           1*
                                   1*
                              1*
                                            1* 1* / MULTNUM IN MODEL
  'MULTNUM'
              1
  'MULTNUM'
                              5
                                   1
                                       5
                                            6
                                                6
                                                   / MULTNUM IN MODEL
   'MULTNUM'
                                       1*
                              1*
                                            10 10 / MULTNUM IN MODEL
-- NOW RESET PORO AND PERMX BASED ON THE MULTNUM REGION NUMBER
-- ADD A CONSTANT TO AN ARRAY BASED ON A REGION NUMBER
                                REGION ARRAY
     ARRAY
               CONSTANT REGION
--
               VALUE
                        NUMBER
                                M / F / O
ADDREG
     'PORO'
              0.050
              0.100 2
    'P0R0'
              -0.050
     'PORO'
                      3
                      1
     'PERMX'
              25.00
                                  М
     'PERMX'
               100.0
     'PERMX'
              -50.00
```

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The example first defines the PORO and PERMX property arrays for the model and then sets the MULTNUM array to I for all cells in the model, after which selected areas of model are assigned various MULTNUM integer values. The ADDREG can then be invoked to add or subtract constant values from the PORO and PERMX arrays for the various MULTNUM regions.

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6.3.4 AQUANCON - Define Analytical Connections to the Grid

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

AQUANCON keyword defines how analytical aquifers are connected to the simulation grid., this includes Carter-Tracy and Fetkovich analytical aquifers, although the Carter-Tracy aquifer is the only aquifer currently implemented in OPM Flow.

No.	Name		Default					
	Field Metric Laboratory							
I	AQUNUM	AQUNUM is a positive integer greater than or equal to one and less than the maximum number of analytical aquifers as defined by the NANAQ variable on the AQUDIMS keyword in the RUNSPEC section, that defines the aquifer to be connected to the grid.						
2	П		defines the lower bouned to the grid and must be all to 12 and NX.		I			
3	12		efines the upper bound of ed to the grid and must be to NX		NX			
4	JI		defines the lower bouned to the grid and must be lal to J2 and NY.		I			
5	J2	direction to be connected	A positive integer that defines the upper bound of the cells in the J-direction to be connected to the grid and must be greater than or equal to JI and less than or equal to NY.					
6	KI	A positive integer that defines the lower bound of the cells in the K-direction to be to be connected to the grid and must be greater than or equal to one and less than or equal to K2 and NZ.						
7	K2	direction to be connected	A positive integer that defines the upper bound of the cells in the K-direction to be connected to grid and must be greater than or equal to KI and less than or equal to NZ.					
8	AQUFACE	declared by this record a I) X+, Y+, or Z+ negative direct 2) I+, J+, or K+	 AQUFACE is a character string that sets the connection "face" of the cells declared by this record and should be set to one of the following: I) X+, Y+, or Z+ for the positive direction, or X-, Y- or Z- for the negative direction transmissibilities. 2) I+, J+, or K+ for the positive direction, or I-, J- or K- for the negative direction transmissibilities. 					
9	AQUFLUX	AQUFLUX is a positive real value that sets the fraction of the total influx between the aquifer and the defined cells declared on this keyword. If defaulted the cell face for each cell is applied and if a values is declared then then this values is applied to all cells declared by this record.						
		ft²	m²	cm ²	! *			
10	AQUCOEF		sitive values that scales the cells declared on this					
		dimensionless	dimensionless	dimensionless	1.0			

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No.	Name		Description		Default
		Field	Metric	Laboratory	
П	AQUOPT	AQUOPT is a character should be set to one of	string that sets the cell fac the following:	ce connection and	NO
		connections ins	nnections <u>can adjoin</u> to acti ide the reservoir grid. It is out thoroughly checking th	not recommended to use	
			nnections <u>cannot adjoin</u> to ide the reservoir grid. This e.		

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Notes:

- Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.7: AQUANCON Keyword Description

Example

The following example defines aquifer number one connected to the I+ face of various cells in the model.

```
ANALYTIC AQUIFER CONNECTION
- -
                                              CONNECT AQF
                                                                         ADJOIN
                    ----- BOX -----
                                                               AQF
         NUMBER I1 I2
                          J1 J2
                                    K1 K2
                                              FACE
                                                       INFLX MULTI
                                                                         CELLS
AQUANCON
                                              'I+'
                                                              1*
                                                                        'NO'
                57 57
                         28
                             36
                                   46 58
         1
              111 111
                         38
                                              'I+'
                                                      1*
                                                              1*
                                                                        ' NO '
                             41
                                   22
                                       31
         1
                                              'I+'
                                                      1*
                                                              1*
         1
                96
                    96
                         44
                             49
                                   22
                                       31
                                                                        'NO'
                                                              1*
                                              'I+'
                                                      1*
                                   54
                                                                        'NO'
                             35
         1
                43
                    43
                         28
                                       58
                                              'I+'
                                                      1*
                                                              1*
                                                                        'NO'
         1
                98
                    98
                         38 42
                                   32 40
                                              'I+'
                                                              1*
                                                                        'NO'
         1
                79
                    79
                         41
                             67
                                   5
                                      11
                                                              1*
                                              'I+'
                                                                        'NO'
                                   12 17
         1
                61
                   61
                         48
                             72
```

See the AQUCT keyword in the GRID section for a complete example on defining and connecting a Carter-Tracy aquifer to a simulation grid.

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6.3.5 AQUCON - Define Numerical Aquifer Connections to the Grid

SPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY	D EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDUL
-----------------------------------------------	-----------------------------------------------

Revision: Rev-I

Description

AQUCON keyword defines how numerical aquifers are connected to the simulation grid.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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6.3.6 AQUCT - Define Carter-Tracy Analytical Aquifers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The AQUCT keyword defines a Carter Tracy aquifer, the properties of the aquifer, including the Carter-Tracy aquifer influence function associated with the aquifer, defined by the AQUTAB keyword in the PROPS section.

Each row entry in the AQUCT keyword defines one Carter-Tracy aquifer.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	AQUID		r than or equal to one an DIMS keyword in the Raquifer number.		1
2	DATUM	DATUM is a single positi datum depth for PRESS.	ive value that defines the	Carter-Tracy reference	
		feet	m	cm	None
3	PRESS	PRESS is a single posit DATUM.	ive value that defines th	ne aquifer pressure at	
			hen the simulator will s be in equilibrium with th		
		psia	barsa	atma	I *
4	PERM	PERM is a real positive nu	umber that assigns the per	meability to the aquifer.	
		mD	mD	mD	None
5	PORO	PORO is a real positive to one that assigns the po	number greater than zero	and less than or equal	
		dimensionless	dimensionless	dimensionless	None
6	RCOMP	RCOMP is a real nu compressibility (Ct) at th	umber defining the tot e DATUM pressure.	al (rock and water)	
		I/psia	I/barsa	I/atma	None
7	RE	RE is a real positive numb	per that defines the Carter	r-Tracy aquifer external	
		feet	m	cm	None
8	DZ	DZ is a real positive num net thickness.	ber that defines the Carte	er-Tracy aquifer average	
		feet	m	cm	None
9	ANGLE	the angular connection be reservoir. A value of 360°	number that defines the a etween the aquifer and the degrees, the default value ands the hydrocarbon reser	e hydrocarbon , indicates that the	
		degrees	degrees	degrees	360.0

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No.	Name		Description		Default
		Field	Metric	Laboratory	
10	PVTNUM	PVTNUM is positive inte variable on the TABDIMS the PVTW table allocated	keyword in the RUNSF	PEC section, that defines	I
П	AQUTAB	AQUTAB is positive integration variable as declared on the that defines the AQUTAB	e AQUDIMS keyword in	the RUNSPEC section,	I
		The default value of one saquifer influence table no PROPS section The first the table number two.	t the first table in the AQ	UTAB keyword in the	
12	SALT	SALT is a real positive nutring the aquifer. This variable is ignored by		tial salt concentration in	
		lb/stb	kg/sm³	gm/scc	0.0
13	TEMP	TEMP is a real positive nu aquifer at DATUM.	imber that defines the in	itial temperature of the	
		This variable is ignored by	OPM Flow.		
		°F	°C	°C	*

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Notes:

- The keyword is followed by up to NANAQ records as defined on the AQUDIMS keyword in the RUNSPEC section
- 2) Each record is terminated by a "/" and the keyword should be terminated by a "/".

Table 6.8: AQUCT Keyword Description

Note

OPM Flow includes the infinite acting Carter-Tracy aquifer influence table as a default for table number one; thus data entered on AQUTAB keyword starts from table number two.

In order to full define a Carter-Tracy aquifer one has to define the aquifer properties via the AQUCT keyword, the Carter-Tracy influence functions via the AQUTAB keyword in the PROPS section, if the default infinite acting table is not being employed, and how the aquifer is connected to the reservoir using the AQUANCON keyword in the GRID or SOLUTION sections.

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Example

```
Given the following grid and aquifer dimensions in the RUNSPEC section:
```

```
MAX
                MAX
                        MAX
- -
        NDIVIX NDIVIY NDIVIZ
DIMENS
                        5
        20
                1
        AQF
                AQF
                        AQF
                                AQF
                                        AQF
                                                AQF
                                                       AQF
                                                              AQF
- -
                MXNAQC NIFTBL NRIFTB NANAQ
        MXAQN
                                                NCAMAX MXNALI MXAAQL
AQUDIMS
        1*
                                                1*
                                                       1*
                                                              1*
                                100
                                        1
```

And AQUTAB in the PROPS section

1000.00

666.982 /

```
-- CARTER-TRACY AQUIFER INFLUENCE TABLES
-- (STARTS FROM TABLE NO. 2, AS DEFAULT IS TABLE NO. 1)
-- AQUITAB
-- DIMLESS DIMLESS
-- TIME PRESSURE
```

DIMLESS TIME	DIMLESS PRESSURE
0.01	0.112
0.05	0.229
0.10	0.315
0.15	0.376
0.20	0.424
0.22	0.443
0.24	0.459
0.26	0.476
0.28	0.492
0.30	0.507
0.32	0.522
0.34	0.536
0.36	0.551
0.38	0.565
0.40	0.579
0.42	0.593
0.44	0.607
0.46	0.621
0.48	0.634
0.50	0.648
0.60	0.715
0.70	0.782
0.80	0.849
0.90	0.915
1.00	0.982
2.00	1.649
3.00	2.316
5.00	3.649
10.00	6.982
20.00	13.649
30.00	20.316
50.00	33.649
100.00	66.982
200.00	133.649
300.00	200.316
500.00	333.649

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```
The Carter-Tracy aquifer is defined in the GRID or SOLUTION sections as:
                         CARTER-TRACY AQUIFER DESCRIPTION
- -
        ID
              DATUM
                      AQF
                              AQF
                                     AQF
                                             AQF
                                                     AQF
                                                                  INFL
                                                                          PVT
                                                                               AQU
        NUM
             DEPTH
                      PRESS PERM
                                             RCOMP
                                     POR<sub>0</sub>
                                                     RET
                                                            DΖ
                                                                  ANGLE NUM
                                                                               TAB
AQUCT
             2000.0 269
                              100.0 0.30
                                                            10.0 360.0
                                                                                2
         1
                                             3.0e-5 330
                                                                           1
And the connection of the aquifer is set in the GRID or SOLUTION sections as:
                         ANALYTIC AQUIFER CONNECTION
_ _
                                              CONNECT AOF
         ID
                 ----- BOX -----
                                                               AOF
                                                                         ADJOIN
         NUMBER I1 I2
                         J1 J2
                                  K1 K2
                                              FACE
                                                       INFLX
                                                              MULTI
                                                                         CELLS
AQUANCON
                                                                         'NO'
                     1
                               1
                                        1
                                               J-
                                                        1.0
         1
                 1
                          1
                                    1
                                                               1.0
```

Here one Carter-Tracy aquifer is connected to a single cell (I, I, I) at the J- face (or X- face) of the cell.

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6.3.7 AQUNUM - NUMERICAL AQUIFER ASSIGNMENT

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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6.3.8 BOX - Define a Range of Grid Blocks to Enter Property Data

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

BOX defines a range of grid blocks for which subsequent data is assigned for all the cells in the defined BOX. Note that the BOX grid is reset by the keyword ENDBOX by resetting the current defined BOX to be the whole grid. The keyword can be used for any array and for all grid types.

No.	Name	Description	Default
I	П	A positive integer that defines the lower bound of the array in the I-direction to be modified must be greater than or equal I and less than or equal to I2 and NX.	I
2	12	A positive integer that defines the upper bound of the array in the I-direction to be modified must be greater than or equal to II and less than or equal to NX	NX
3	JI	A positive integer that defines the lower bound of the array in the J-direction to be modified must be greater than or equal I and less than or equal to J2 and NY.	ı
4	J2	A positive integer that defines the upper bound of the array in the J-direction to be modified must be greater than or equal to JI and less than or equal to NY.	NY
5	KI	A positive integer that defines the lower bound of the array in the K-direction to be modified must be greater than or equal to one and less than or equal to K2 and NZ.	I
6	K2	A positive integer that defines the upper bound of the array in the K-direction to be modified must be greater than or equal to KI and less than or equal to NZ.	NZ

Notes:

- Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) The keyword is terminated by "/".

Table 6.9: BOX Keyword Description

See also the ADD, COPY, ENDBOX, EQUALS, and MULTIPLY keywords can also be used to enter data in a subset of the model.

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Examples

```
DEFINE A BOX GRID FOR THE BOTTOM LAYER OF A 100 X 100 X 20 MODEL
     ----- BOX -----
    I1 I2
            J1 J2 K1 K2
B<sub>0</sub>X
    1* 1*
              1* 1*
                       20 20 / SELECT THE BOTTOM LAYER
   DEFINE THE POROSITY AND OTHER PROPERTIES ON THE BOX GRID
POR<sub>0</sub>
  10000*0.300
PERMX
 5000*100.0 5000*75.0
NTG
  10000*0.500
   RESET THE INPUT BOX TO BE THE FULL MODEL
ENDBOX
```

The above example set the BOX grid to be the last layer in the model which means that 100×100 , that is 10,000 data elements need to entered for each property.

Alternatively, one could use the EQUALS keyword to accomplish the same thing.

```
CONSTANT --
                        ----- BOX -----
-- -- ARRAY
                         I1 I2 J1 J2
                                        K1 K2
EQUALS
  'PORO'
             0.3000
                         1* 1*
                                1* 1*
                                         20 20 / PORO TO 0.30 IN LAYER 20
                         1 50 1* 1*
  'PERMX'
             0.1000
                                         20 20 / PERMX TO 100. IN LAYER 20
                         50 100 1* 1*
  'PERMX'
             0.1000
                                         20 20 / PERMX TO 75.0 IN LAYER 20
  'NTG'
             0.0500
                           1*
                                 1* 1*
                                         20 20 / NRT TO 0.50 IN LAYER 20
```

Note

It is good practice to always use the ENDBOX keyword to reset the input back to the full grid when all the modifications for a sub-grid have been completed.

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6.3.9 CARFIN - Define a Cartesian Local Grid Refinement

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

CARFIN defines a Cartesian local grid refinement ("LGR") in a cell or a group of cells in the main model.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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6.3.10 CIRCLE - COMPLETION OF RADIAL GRID CIRCLE ACTIVATION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

CIRCLE activates the completion of the circle for a radial grid by generating non-neighboring connections between the first and last cells in the radial plane and for all layers in the model. This is only activated if the radial grid extends 360 degrees in the radial plane. The RADIAL keyword in the RUNSPEC section should be activated to indicate that radial geometry is being used.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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6.3.11 COALNUM - Define the Coal Region Numbers

RUNSPEC	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The COALNUM keyword defines the coal region numbers for each grid block used with the coal bed methane option. OPM Flow does not have a coal bed methane option and therefore this keyword is ignored by OPM Flow; however, it is documented here for completeness.

No.	Name	Description	Default
- 1	COALNUM	COALNUM defines an array of positive integers assigning a grid cell to a particular coal region.	I
		The maximum number of COALNUM regions is set by the NTCREG variable on REGDIMS keywords in the RUNSPEC section.	

Notes:

- 1) A total of NX x NY x NZ integer values, as defined by the DIMENS in the RUNSPEC section, must be specified for the array.
- 2) If cell is not assigned a COALNUM region then the default value of I will be used.
- 3) COALNUM value of 0 sets the cell be a non-coal region.
- 4) The keyword is terminated by "/".

Table 6.10: COALNUM Keyword Description

Example

The example below sets three COALNUM regions for a 4 x 5 x 2 model.

COALNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
```

The above will no effect in an OPM Flow input deck.

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6.3.12 COORD - Define a Set of Coordinates Lines for a Reservoir Grid

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

COORD defines a set of coordinate lines or pillars for a reservoir grid via an array. A total of (NX+I) x (NY+I) lines must be specified for each coordinate data set.

For Cartesian geometry, each line is defined by the (x, y, z) coordinates of two distinct points on the line. The lines are entered with I cycling fastest then J. For radial geometry, each line is defined by the (r, theta) coordinates of two distinct points on the line. The lines are entered with R cycling fastest then THETA.

The keyword can only be used with Irregular Corner-Point Grids.

No.	Name		Description					
		Field	Metric	Laboratory	-			
I	XI-Column	Top X coordinate						
2	YI-Column	Top Y coordinate			-			
3	ZI-Column	Top Z coordinate			-			
4	X2-Column	Base X coordinate			-			
5	Y2-Column	Base Y coordinate			=			
6	Z2-Column	Base Z coordinate			1			
		feet	metres	cm	None			

Notes:

- 1) THETA values are in degrees for all units.
- 2) Regular Cartesian and Radial grid keywords cannot be used with this keyword, that is: DR, DRV, DTHETA, DTHETAV, DX, DXV, DY, DYV, DZ, INRAD, and TOPS.
- 3) The keyword is terminated by "/".

Table 6.11: COORD Keyword Description

See also the SPECGRID, COORDSYS and ZCORN keywords to fully define an Irregular Corner-Point Grid.

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Example

```
-- SPECIFY VERTICAL COORDINATE LINES FOR A REGULAR 3 x 2 GRID
--(DX = 100 \text{ and } DY = 200)
-- X1
           Y1
                 Z1
                                  Y2
                                         Z2
__ __
                                         _ _ _ _
COORD
     0
             0
                 1000
                            0
                                    0
                                         5000
   100
             0
                 1000
                           100
                                    0
                                         5000
   200
                 1000
                          200
                                         5000
            0
                                    0
   300
             0
                 1000
                          300
                                    0
                                         5000
           200
                                  200
                                         5000
     0
                 1000
                            0
   100
           200
                 1000
                          100
                                  200
                                         5000
   200
           200
                 1000
                          200
                                  200
                                         5000
   300
           200
                 1000
                          300
                                  200
                                         5000
                                         5000
     0
           400
                 1000
                            0
                                  400
   100
           400
                 1000
                           100
                                  400
                                         5000
   200
           400
                 1000
                           200
                                  400
                                         5000
   300
           400
                 1000
                          300
                                  400
                                         5000
```

The above example defines vertical coordinate lines for a regular 3 by 2 grid with DX equal to 100 and DY equal to 200.

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6.3.13 COORDSYS - Define Coordinate Grid Options

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

This keyword sets various options for when multiple grid system are being used.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.14 COPY - Copy Array Data to Another Array

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The COPY keyword copies an array (or part of an array) to another array or part of an array. The arrays can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the COPY keyword is being used.

No.	Name	Description	Default
I	ARRAY-I	The name of the array to be copied from.	None
		This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	
2	ARRAY-2	The name of the array to be copied to.	None
		This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	
3	П	A positive integer that defines the lower bound of the array in the I-direction to be modified must be greater than or equal I and less than or equal to I2 and NX.	I
4	12	A positive integer that defines the upper bound of the array in the I-direction to be modified must be greater than or equal to II and less than or equal to NX	NX
5	JI	A positive integer that defines the lower bound of the array in the J-direction to be modified must be greater than or equal I and less than or equal to J2 and NY.	I
6	J2	A positive integer that defines the upper bound of the array in the J-direction to be modified must be greater than or equal to JI and less than or equal to NY.	NY
7	KI	A positive integer that defines the lower bound of the array in the K-direction to be modified must be greater than or equal to one and less than or equal to K2 and NZ.	I
8	K2	A positive integer that defines the upper bound of the array in the K-direction to be modified must be greater than or equal to KI and less than or equal to NZ.	NZ

Notes:

- Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.12: COPY Keyword Description

The applicable arrays for each section are defined in Table 6.13 on the following page.

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	(COPY Keywor	d and Variable C	Options by Section	on	
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
DX	DEPTH	SWL	ENDNUM	PRESSURE		
DY	PORV	SWCR	EQLNUM	SWAT		
DZ	TRANX	SWU	FIPNUM	SGAS		
PERMX	TRANY	SGL	IMBNUM	RV		
PERMY	TRANZ	SGCR	MISCNUM	RS		
PERMZ	DIFFX	SGU	PVTNUM	TBLK		
MULTX	DIFFY	KRW	ROCKNUM	GI		
MULTY	DIFFZ	KRO	SATNUM	OILAPI		
MULTZ	TRANR	KRG	WH2NUM	SALT		
DR	TRANTHT	PCG		GASCONC		
THETA	DIFFR	PCW		SOLVCONC		
PERMR	DIFFTHT			SOLVFRAC		
PERMTHT				SFOAM		
DZNET				SPOLY		
PORO						
NTG						
FLUXNUM						
MULTNUM						
MPANUM						
DIFFX						
DIFFY						
DIFFZ						
DIFFR						
DIFFTHT						

Table 6.13: COPY Keyword Applicable Arrays by Section

Example

```
-- -- SOURCE
              DESTIN. --
                           ----- BOX -----
                          I1 I2 J1 J2 K1 K2
COPY
              'PERMY'
                         1* 1* 1* 1* 1* 1* 1* / CREATE PERMY
1* 1* 1* 1* 1* 1* / CREATE PERMZ
  'PERMX'
              'PERMZ'
  'PERMX'
-- -- ARRAY
              CONSTANT -- ----- BOX -----
                          I1 I2 J1 J2 K1 K2
MULTIPLY
   'PERMZ'
              0.50000
                          1* 1* 1* 1* 1* 1* / PERMZ * 0.5
```

The above example copies PERMX array to the PERMY and PERMZ arrays in the GRID section for all grid blocks in the model. The PERMZ array is then multiplied by 0.5 for all grid blocks in the model.

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6.3.15 COPYREG - COPY AN ARRAY TO ANOTHER ARRAY BASED ON A REGION NUMBER

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

6.3.16 DESCRIPTION

The COPYREG keyword copies a specified array or part of an array based on cells with a specific region number to another array. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the COPYREG keyword is read by the simulator. The property arrays can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the COPYREG keyword is being used.

No.	Name	Description	Default
I	ARRAY-I	The name of the array to be copied from.	None
		This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	
2	ARRAY-2	The name of the array to be copied to.	None
		This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	
3	REGION NUMBER	Integer REGION NUMBER is the region for which the array data in (I) should be copied to array data in (2).	None
4	REGION ARRAY	The REGION ARRAY to use for selecting the REGION NUMBER in (3) for selecting the data to be copied. REGION ARRAY can have the following values:	М
		F for the FLUXNUM array	
		2) M for the MULTNUM array	
		3) O for the OPERNUM array	

Notes:

- Where the REGION NUMBER should be less than or equal to the maximum number of regions as defined on the REGDIMS keyword for the FIPNUM and OPERNUM arrays or the GRIDOPTS keyword for the MULTNUM array in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.14: COPYREG Keyword Description

The applicable arrays for each section are defined in Table 6.15 on the following page.

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	cc	PYREG Keyw	ord and Variable	Options by Sec	tion	
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
DX	DEPTH	SWL	ENDNUM	PRESSURE		
DY	PORV	SWCR	EQLNUM	SWAT		
DZ	TRANX	SWU	FIPNUM	SGAS		
PERMX	TRANY	SGL	IMBNUM	RV		
PERMY	TRANZ	SGCR	MISCNUM	RS		
PERMZ	DIFFX	SGU	PVTNUM	TBLK		
MULTX	DIFFY	KRW	ROCKNUM	GI		
MULTY	DIFFZ	KRO	SATNUM	OILAPI		
MULTZ	TRANR	KRG	WH2NUM	SALT		
DR	TRANTHT	PCG		GASCONC		
THETA	DIFFR	PCW		SOLVCONC		
PERMR	DIFFTHT			SOLVFRAC		
PERMTHT				SFOAM		
DZNET				SPOLY		
PORO						
NTG						
FLUXNUM						
MULTNUM						
MPANUM						
DIFFX						
DIFFY						
DIFFZ						
DIFFR						
DIFFTHT						

Table 6.15: COPYREG Keyword Applicable Arrays by Section

Example

```
-- COPY AN ARRAY TO ANOTHER ARRAY BASED ON A REGION NUMBER
                ARRAY
                          REGION
                                   REGION ARRAY
--
      ARRAY
      FROM
                          NUMBER
                                    M / F / O
COPYREG
     'PERMX'
                'PERMY'
                                     М
                                                      / COPY PERMX TO PERMY
                          1
     'PERMX'
                'PERMZ'
                                                      / COPY PERMX TO PERMZ
-- NOW RESET PERMZ BASED ON THE MULTNUM REGION NUMBER
-- MULTIPLY AN ARRAY BY A CONSTANT BASED ON A REGION NUMBER
      ARRAY
                CONSTANT REGION
                                   REGION ARRAY
- -
                VALUE
                          NUMBER
                                    M / F / O
MULTIREG
     'PERMZ'
                 0.95
                          1
                                     М
```

The above example first copies the PERMX property array for region number one to the PERMY and PERMZ property arrays for region one using the MULTNUM array to define the region numbers. After which PERMZ property array for region one is multiplied by 0.5 using the MULTIREG keyword.

FLOW DOCUMENTATION MANUAL (2018-10)

6.3.17 DR - Define the Size of Grid Blocks in the R Direction for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DR defines the size of all grid blocks in the R direction via an array for each cell in a Radial Grid model.

No.	Name		Description						
		Field	Metric	Laboratory					
I	DR	DR is an array of real nu for each cell in the mode							
		Repeat counts may be us							
		feet	m	cm	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.16: DR Keyword Description

See also the DRV, DTHETAV, DZ and TOPS keywords in the GRID section to fully define a radial grid model.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

Given the dimensions of the grid in the RUNSPEC section to be 10, 1, 8 for NX, NY and NZ respectively, then DR should be defined as:

```
INNER RADIUS OF FIRST GRID BLOCK IN THE RADIAL DIRECTION
- -
INRAD
 0.25
         DEFINE GRID BLOCK R DIRECTION CELL SIZE
DR
 1.75
                                  50.55
       2.32 5.01 10.84
                          23.39
                                         109.21
                                                 235.92 509.68
                                                                 1101.0
 1.75
             5.01 10.84
                          23.39
                                 50.55
                                                 235.92 509.68
                                                                 1101.0
       2.32
                                         109.21
 1.75
                                 50.55
                                                        509.68
       2.32
             5.01 10.84
                          23.39
                                         109.21
                                                 235.92
                                                                 1101.0
 1.75
       2.32
             5.01
                   10.84
                          23.39
                                  50.55
                                         109.21
                                                 235.92
                                                         509.68
                                                                 1101.0
 1.75
       2.32
             5.01
                   10.84
                           23.39
                                  50.55
                                         109.21
                                                 235.92
                                                         509.68
                                                                 1101.0
 1.75
       2.32
                   10.84
                           23.39
                                  50.55
                                                 235.92
             5.01
                                         109.21
                                                         509.68
                                                                 1101.0
 1.75
       2.32
             5.01
                   10.84
                           23.39
                                  50.55
                                         109.21
                                                 235.92
                                                         509.68
                                                                 1101.0
 1.75
       2.32
             5.01
                   10.84
                           23.39
                                  50.55
                                         109.21
                                                 235.92
                                                         509.68
                                                                 1101.0
```

The above example defines the size of the cells in the R direction based on 80 cells in the model as defined by the DIMENS keyword in the RUNSPEC section. Note the INRAD keyword to define the inner radius of the radial grid.

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6.3.18 DRV - Define the Size of Grid Blocks in the R Direction via a Vector

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

DRV²⁵ defines the size of grid blocks in the R direction via a vector as opposed to defining the property for each cell for a Radial Grid. The RADIAL keyword in the RUNSPEC section should be activated to indicate that radial geometry is being used.

No.	Name			Default	
		Field	Metric	Laboratory	
I	DRV	DRV is a vector of real noin the R direction in a rac			
		Repeat counts may be us			
		feet	m	cm	None

Notes:

- The number of entries should correspond to the NX parameter on the DIMENS keyword in the RUNSPEC section
- 2) The keyword is terminated by "/".

Table 6.17: DRV Keyword Description

See also the DR, DTHETAV, DZ and TOPS keywords in the GRID section to fully define a radial grid model.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- INNER RADIUS OF FIRST GRID BLOCK IN THE RADIAL DIRECTION
-- INRAD
0.25
/--
-- DEFINE GRID BLOCK SIZES IN THE R DIRECTION
--
DRV
1.75 2.32 5.01 10.84 23.39 50.55 109.21 235.92 509.68 1101.0 /
```

The above example defines the size of the cells in the R direction based on NX equals 10 on the DIMENS keyword in the RUNSPEC section. Note the INRAD keyword to define the inner radius of the radial grid.

²⁵ Radial grids are not currently implemented in OPM Flow version 2018-04 but is expected to be incorporated in the next release.



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6.3.19 DTHETA - Define the Size of GRID BLOCKS IN THE THETA DIRECTION FOR ALL CELLS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

DTHETA defines the size of all grid blocks in the R direction via an array for each cell in a Radial Grid model.

No.	Name			Default						
		Field	Field Metric Laboratory							
I	DR	DTHETA is an array of redirection in radial grids for								
		Repeat counts may be us	Repeat counts may be used, for example 10*25.0							
		degrees	degrees	degrees	None					

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/". 2)

Table 6.18: DTHETA Keyword Description

See also the DRV, DTHETAV, DZ and TOPS keywords in the GRID section to fully define a radial grid model.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

Given the dimensions of the grid in the RUNSPEC section to be 10, 6, I for NX, NY and NZ respectively, then DTHETA should be defined as:

```
DEFINE GRID BLOCK SIZES IN THE THETA DIRECTION
DTHETA
    10*60.0
    10*60.0
    10*60.0
    10*60.0
    10*60.0
    10*60.0
```

The above example defines the size of the cells in the R direction based on 60 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

FLOW DOCUMENTATION MANUAL (2018-10)

6.3.20 DTHETAV - Sets the Size of Grid Blocks in THETA Direction via a Vector

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

DTHETAV26 defines the size of grid blocks in the THETA direction via a vector as opposed to defining the property for each cell for a Radial Grid. The RADIAL keyword in the RUNSPEC should be activated to indicate that radial geometry is being used.

No.	Name		Default					
		Field	Metric	Laboratory				
I	DTHETAV	DTHETAV is a vector of blocks in the THETA dire						
		Repeat counts may be us	Repeat counts may be used, for example 10*100.0.					
		degrees	degrees	degrees	None			

Notes:

- The number of entries should correspond to the NY parameter of the DIMENS keyword in the RUNSPEC section.
- 2) The keyword is terminated by "/".

Table 6.19: DTHETA Keyword Description

See also the DRV, DZV and TOPS keywords to fully define a radial grid model.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- DEFINE GRID BLOCK SIZES IN THE THETA DIRECTION (BASED ON NY = 6)
-- DTHETAV
60.0 60.0 60.0 60.0 60.0 60.0
```

The above example defines the size of the cells in the THETA direction based on NY equals six in the DIMENS keyword in the RUNSPEC section.

Radial grids are not currently implemented in OPM Flow version 2018-04 but is expected to be incorporated in the next release.



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6.3.21 DUMPFLUX - ACTIVATE WRITING OUT OF A FLUX FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates the writing out of a full field (the full grid) FLUX file for later processing in a Flux Boundary run. The Flux Boundary feature allows for the segmentation of the full grid into flux boundary areas which allow for a sub-area of the grid to be run and at the same time model the flux across the boundary derived from the main grid. The object of this feature is to be able to investigate the performance of various areas of the model without having to run the full field, thus improving computational efficiency and run times, but still obtain "reasonable" results due to the incorporation of the fluxes across the boundary.

This feature is not available in OPM Flow; however it is documented here for completeness.

There is no data required for this keyword.

Example

- -

-- ACTIVATE WRITING OUT OF A FLUX FILE DUMPFLUX

The above example switches on the writing of the FLUX output file; the keyword has no effect and is ignored by the simulator.

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6.3.22 DX - Define the Size of Grid Blocks in the X Direction for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DX defines the size of all grid blocks in the X direction via an array for each cell in a Cartesian Regular Grid model.

No.	Name		Description						
		Field	Metric	Laboratory					
I	DX	for each cell in the mode							
		Repeat Counts may be us	Repeat counts may be used, for example 10*100.0.						
		feet	m	cm	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.20: DX Keyword Description

See also the DY, DZ and TOPS keywords to fully define a Cartesian Regular Grid.

Example

```
-- DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX x NY x NZ = 300)
-- DX
300*1000
/
```

The above example defines the size of the cells in the X direction based on 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.23 DXV - Define the Size of Grid Blocks in the X Direction via a Vector

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DXV defines the size of grid blocks in the X direction via a vector as opposed to defining the X direction cell size for each cell for a Cartesian Regular Grid.

No.	Name		Description					
		Field	Field Metric Laboratory					
I	DXV	DXV is a vector of rea	DXV is a vector of real numbers describing the cell size for the grid blocks in the X direction.					
		Repeat counts may be us	Repeat counts may be used, for example 10*100.0.					
		feet	m	cm	None			

Notes:

- The number of entries should correspond to the NX parameter on the DIMENS keyword in the RUNSPEC section.
- 2) The keyword is terminated by "/".

Table 6.21: DXV Keyword Description

See also the DYV, DZV and TOPS keywords to fully define a Cartesian Regular Grid.

Example

```
-- DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX = 5)
-- DXV
5*100 /
```

The above example defines the size of the cells in the X direction based on NX equals 5 on the DIMENS keyword in the RUNSPEC section.

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6.3.24 DY - Define the Size of Grid Blocks in the Y Direction for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DY defines the size of all grid blocks in the Y direction via an array for each cell in a Cartesian Regular Grid model.

No.	Name		Description						
		Field	Metric	Laboratory					
I	DY	DY is an array of real nu for each cell in the mode	DY is an array of real numbers describing the cell size in the Y direction or each cell in the model.						
		Repeat counts may be us	Repeat counts may be used, for example 10*100.0.						
		feet	m	cm	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.22: DY Keyword Description

See also the DX, DZ and TOPS keywords to fully define a Cartesian Regular Grid.

Example

```
-- DEFINE GRID BLOCK Y DIRECTION CELL SIZE (BASED ON NX x NY x NZ = 300)
-- DY 300*1000
```

The above example defines the size of the cells in the Y direction based on 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.25 DYV - Define the Size of Grid Blocks in the Y Direction via a Vector

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DYV defines the size of grid blocks in the Y direction via a vector as opposed to defining the Y direction cell size for each cell for a Cartesian Regular Grid.

No.	Name		Description					
		Field	Metric	Laboratory				
I	DYV	DYV is a vector of real notine Y direction.	DYV is a vector of real numbers describing the cell size for the grid blocks in the Y direction.					
		Repeat counts may be us	Repeat counts may be used, for example 10*100.0.					
		feet	feet m cm					

Notes:

- The number of entries should correspond to the NY parameter on the DIMENS keyword in the RUNSPEC section.
- The keyword is terminated by "/". 2)

Table 6.23: DYV Keyword Description

See also the DXV, DZV and TOPS keywords to fully define a Cartesian Regular Grid.

Example

```
DEFINE GRID BLOCK Y DIRECTION CELL SIZE (BASED ON NY = 5)
DYV
  5*100
```

The above example defines the size of the cells in the Y direction based on NY equals 5 on the DIMENS keyword in the RUNSPEC section.

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6.3.26 DZ - Define the Size of Grid Blocks in the Z Direction for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DZ defines the size of all grid blocks in the Z direction via an array for each cell in a Cartesian Regular Grid model.

No.	Name		Description						
		Field	Metric	Laboratory					
I	DZ	DZ is an array of real nu for each cell in the mode	Z is an array of real numbers describing the cell size in the Z direction or each cell in the model.						
		Repeat counts may be us	Repeat counts may be used, for example 10*100.0.						
		feet	m	cm	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.24: DZ Keyword Description

See also the DX, DY and TOPS keywords to fully define a Cartesian Regular Grid.

Example

```
-- DEFINE GRID BLOCK Z DIRECTION CELL SIZE (BASED ON NX x NY x NZ = 300)
-- DZ
-- 100*20.0 100*30.0 100*50.0
```

The above example defines the size of the cells in the Z direction based on 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.27 DZV - DEFINE THE SIZE OF GRID BLOCKS IN THE Z DIRECTION VIA A VECTOR

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DZV defines the size of grid blocks in the Z direction via a vector as opposed to defining the thickness property for each cell. The keyword is used for both Cartesian Regular Grids and Radial Grids.

No.	Name		Default							
		Field	Metric	Laboratory						
I	DZV	blocks in the Z direction.	DZV is a vector of real numbers describing the cell size for the grid locks in the Z direction. Repeat counts may be used, for example 10*20.0.							
		feet	m	cm	None					

Notes:

- The number of entries should correspond to the NZ parameter on the DIMENS keyword in the RUNSPEC section..
- he keyword is terminated by "/". 2)

Table 6.25: DZV Keyword Description

See also the DXV, DXY and TOPS keywords for a Cartesian Regular Grid and DRV, DTHETAV and TOPS keywords to fully define a Radial Grid model.

Example

```
DEFINE GRID BLOCK SIZES IN THE Z DIRECTION (BASED ON NZ = 20)
DZV
                         5.0 15*3.0
                   2.0
3.0
       5.0
            3.0
```

The above example defines the size of the cells in the Z direction based on NZ equals 20 on the DIMENS keyword in the RUNSPEC section.

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6.3.28 ENDBOX - Define the End of the BOX Defined Grid

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword marks the end of a previously defined BOX sub-grid as defined by a previously entered BOX keyword. The keyword resets the input grid to be the full grid as defined by the NX, NY, and NZ variables on the DIMENS keyword in the RUNSPEC section.

There is no data required for this keyword.

Example

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
        ----- BOX -----
- -
        I1 I2 J1 J2 K1 K2
B0X
                                                       / DEFINE BOX AREA
        10 10
                1
                    6
                         1
                             1
        DEFINE GRID BLOCK PERMZ DATA FOR THE INPUT BOX
PERMZ
 6*0.01
        DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The above example defines a subset of the grid and sets the cells PERMZ values to 0.01 for that area. After which the ENDBOX keyword resets the input to be the full grid.

Note

It is good practice to always use the ENDBOX keyword to reset the input back to the full grid when all the modifications for a sub-grid have been completed.

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6.3.29 ENDFIN - END THE DEFINITION OF A LOCAL GRID REFINEMENT

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHED	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
--------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

ENDFIN defines the end of a Cartesian or radial local grid refinement ("LGR") definition and a LGR property definition data set.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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6.3.30 EQUALREG - Sets an Array to a Constant by Region Number

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

6.3.31 DESCRIPTION

The EQUALREG keyword sets a specified array to a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the EQUALREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the EQUALREG keyword is being used.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	An integer or real value to be assigned to the ARRAY in the same units as the ARRAY property for a given REGION	0
3	REGION NUMBER	REGION NUMBER is a positive integer representing the region for which the CONSTANT in (2) should be applied	None
4	REGION ARRAY	The REGION ARRAY to use for applying the CONSTANT in (2) based on the REGION NUMBER in (3). REGION ARRAY can have the following values:	М
		F for the FLUXNUM array	
		2) M for the MULTNUM array	
		3) O for the OPERNUM array	

Notes:

- Where the REGION NUMBER should be less than or equal to the maximum number of regions as defined on the REGDIMS keyword for the FIPNUM and OPERNUM arrays or the GRIDOPTS keyword for the MULTNUM array in the RUNSPEC section.
- Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.26: EQUALREG Keyword Description

The applicable arrays for each section are defined in Table 6.27 on the following page.

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	EQUALREG Keyword and Variable Options by Section										
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE					
DX	DEPTH	SWL	ENDNUM	PRESSURE							
DY	PORV	SWCR	EQLNUM	SWAT							
DZ	TRANX	SWU	FIPNUM	SGAS							
PERMX	TRANY	SGL	IMBNUM	RV							
PERMY	TRANZ	SGCR	MISCNUM	RS							
PERMZ	DIFFX	SGU	PVTNUM	TBLK							
MULTX	DIFFY	KRW	ROCKNUM	GI							
MULTY	DIFFZ	KRO	SATNUM	OILAPI							
MULTZ	TRANR	KRG	WH2NUM	SALT							
DR	TRANTHT	PCG		GASCONC							
THETA	DIFFR	PCW		SOLVCONC							
PERMR	DIFFTHT			SOLVFRAC							
PERMTHT				SFOAM							
DZNET				SPOLY							
PORO											
NTG											
FLUXNUM											
MULTNUM											
MPANUM											
DIFFX											
DIFFY											
DIFFZ											
DIFFR											
DIFFTHT											

Table 6.27: EQUALREG Keyword Applicable Arrays by Section

Examples

```
-- FIRST DEFINE MULTNUM ARRAYS FOR 10 X 10 X 20 MODEL
-- -- ARRAY
              CONSTANT --
                          ----- BOX -----
                          I1 I2 J1 J2 K1 K2
EQUALS
                              1*
                                  1* 1*
                                          1* 1* / MULTNUM IN MODEL
   'MULTNUM'
                          1*
                              1*
                                  1* 1*
                                           6 6
   'MULTNUM'
                                                   / MULTNUM IN MODEL
                                  1* 1*
   'MULTNUM'
                                           10 10 / MULTNUM IN MODEL
-- NOW SET PORO AND PERMX BASED ON THE MULTNUM REGION NUMBER
-- SETS A CONSTANT TO AN ARRAY BASED ON A REGION NUMBER
- -
- -
     ARRAY
               CONSTANT REGION
                                 REGION ARRAY
                                M / F / O
                        NUMBER
               VALUE
EQUALREG
     'PORO'
              0.200
     'PORO'
              0.150
                      3
     'PORO'
              0.120
     'PERMX'
              100.00
                       1
                                  Μ
    'PERMX'
              75.00
                        2
                                  Μ
    'PERMX'
               50.00
```

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The example first defines the MULTNUM array to I for all cells in the model, after which selected areas of model are assigned various MULTNUM integer values. The EQUALREG can then be invoked to set a constant values for the PORO and PERMX arrays for the various MULTNUM regions.

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6.3.32 EQUALS - Sets a Specified Array to a Constant

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
---------	------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

6.3.33 DESCRIPTION

The EQUALS keyword sets a specified array or part of an array to a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the EQUALS keyword is being used.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	An integer or real value that the ARRAY will be set to in the same units as the ARRAY property.	None
3	П	A positive integer that defines the lower bound of the array in the I-direction to be modified must be greater than or equal to one and less than or equal to I2 and NX.	I
4	12	A positive integer that defines the upper bound of the array in the I-direction to be modified must be greater than or equal to II and less than or equal to NX	NX
5	JI	A positive integer that defines the lower bound of the array in the J-direction to be modified must be greater than or equal to one and less than or equal to J2 and NY.	I
6	J2	A positive integer that defines the upper bound of the array in the J-direction to be modified must be greater than or equal to JI and less than or equal to NY.	NY
7	KI	A positive integer that defines the lower bound of the array in the K-direction to be modified must be greater than or equal to one and less than or equal to K2 and NZ.	I
8	K2	A positive integer that defines the upper bound of the array in the K-direction to be modified must be greater than or equal to KI and less than or equal to NZ.	NZ

Notes:

- Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.28: EQUALS Keyword Description

The applicable arrays for each section are defined in Table 6.29 on the following page.

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EQUALS Keyword and Variable Options by Section GRID **EDIT** PROPS **REGIONS** SOLUTION **SUMMARY SCHEDULE** DX DEPTH SWL ENDNUM **PRESSURE** DY **PORV SWCR EQLNUM SWAT** DΖ TRANX SWU **FIPNUM** SGAS PERMX TRANY SGL **IMBNUM** RV PERMY TRANZ SGCR **MISCNUM** RS PERMZ PVTNUM **DIFFX** SGU **TBLK** MULTX DIFFY KRW ROCKNUM GI MULTY SATNUM **OILAPI** DIFFZ KRO MULTZ KRG WH2NUM TRANR **SALT** DR **TRANTHT PCG GASCONC** THETA PCW SOLVCONC DIFFR **SOLVFRAC** PERMR **DIFFTHT** PERMTHT **SFOAM DZNET SPOLY** PORO NTG

Revision: Rev-I

Table 6.29: EQUALS Keyword Applicable Arrays by Section

Examples

FLUXNUM MULTNUM MPANUM DIFFX DIFFY DIFFZ DIFFR DIFFTHT

The above example resets the PERMX, PERMY and PERMZ, arrays to 0.10, 0.10, and 0.01 for all cells in layer five, respectively.

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6.3.34 FAULTS - Define Faults in the Grid Geometry

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The FAULTS keyword defines the faults in the grid geometry and the keyword is normally exported with the grid geometry COORD and ZCORN data sets from static earth modeling software. Note that the FAULT keyword is not required to describe the structural geometry as this is already accounted for in the COORD and ZCORN data sets, but instead lists the fault traces with respect to the grid. Once the fault traces have been defined with the FAULTS keyword then the fault transmissibilities can be modified by the MULTFLT keyword. Note that without the FAULTS keyword one would still get proper cross-fault transmissibilities but they would not be modifiable using MULTFLT keyword.

No.	Name	Description	Default
I	FLTNAME	FLTNAME is a character string enclosed in quotes with a maximum length of eight characters, that defines the name of the fault.	None
2	П	The lower bound of the fault's I-direction range must be greater than or equal to one and less than or equal to I2 and NX.	None
3	12	The upper bound of the fault's I-direction range must be greater than or equal to II and less than or equal to NX	None
4	JI	The lower bound of the fault's J-direction range must be greater than or equal to one and less than or equal to J2 and NY.	None
5	J2	The upper bound of the fault's J-direction range must be greater than or equal to JI and less than or equal to NY.	None
6	KI	The lower bound of the fault's K-direction range must be greater than or equal to one and less than or equal to K2 and NZ.	None
7	K2	The upper bound of the fault's K-direction range must be greater than or equal to KI and less than or equal to NZ.	None
8	FLTFACE	FLTFACE is a character string enclosed in quotes with a maximum length of two characters, that classifies the fault face.	None
		 If TRANMULT on the GRIDOPTS keyword in the RUNSPEC section is set to NO then FLTFACE can have values of X, Y, or Z. 	
		 Alternatively, if TRANMULT on the GRIDOPTS keyword in the RUNSPEC section is set to YES then FLTFACE can have values of X, Y, or Z for the positive direction, or X-, Y- or Z- for the negative direction transmissibilities. 	

Notes:

- Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) The FAULTDIM keyword in the RUNSPEC defines the maximum number of records (or segments) that can be entered with the FAULTS keyword.
- 3) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.30: FAULTS Keyword Description

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Example

The example below defines two fault traces, the first being the 'M_WEST' fault and the second the 'BC' fault trace.

DEFINE FAUL	TS IN	THE G	RID G	EOMETR	RY			
 FALLE			E 4.1	U T TD	ACE			
FAULT	T-1	TO		JLT TR				
NAME	I1	12	J1	J2	K1	K2	FACE	
FAULTS	-	-	2	2	4	22	'X'	,
'M_WEST'	5	5 5	3	3	1	22	'X'	/
'M_WEST'	5		4	4	1	22		/
'M_WEST'	5	5	5	5	1	22	'X'	/
'M_WEST'	5	5	6	6	1	22	'X'	/
'M_WEST'	5	5	7	7	1	22	'X'	/
'M_WEST'	5	5	8	8	1	22	'X'	/
'M_WEST'	5	5	9	9	1	22	'X'	/
'M_WEST'	5	5	10	10	1	22	'X'	/
'M_WEST'	5	5	11	11	1	22	'X'	/
'BC'	43	43	8	8	1	22	'Y'	/
'BC'	42	42	9	9	1	22	'X'	/
'BC'	44	44	8	8	1	22	'Y'	/
'BC'	45	45	8	8	1	22	'Y'	/
'BC'	46	46	8	8	1	22	'Y'	/
'BC'	31	31	9	9	1	22	'Y'	/
'BC'	30	30	10	10	1	22	'X'	/
'BC'	32	32	9	9	1	22	'Y'	/
'BC'	33	33	9	9	1	22	'Y'	/
'BC'	34	34	9	9	1	22	'Y'	/
'BC'	35	35	9	9	1	22	'Y'	/
'BC'	36	36	9	9	1	22	'Y'	/
'BC'	37	37	9	9	1	22	ΙΫ́Ι	,
'BC'	38	38	9	9	1	22	'Ÿ'	,
'BC'	39	39	9	9	1	22	ΙΫ́Ι	,
'BC'	40	40	9	9	1	22	ΙÝΙ	,
			Ŭ	Ŭ	_		•	,

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6.3.35 FLUXNUM - Define the Flux Regions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The FLUXNUM keyword defines the flux region numbers for each grid block, as such there must be one entry for each cell in the model. The array is used with the flux boundary option to define the various flux regions²⁷. In addition, the array can be used with the EQUALREG, ADDREG, COPYREG, MULTIREG, MULTREGP and MULTREGT keywords in calculating various grid properties in the GRID section.

No.	Name	Description	Default
I	FLUXNUM	FLUXNUM defines an array of positive integers assigning a grid cell to a particular flux region.	I
		The maximum number of flux regions is set by the MXNFLX variable on the REGDIMS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.31: FLUXNUM Keyword Description

Examples

The example below sets three FLUXNUM regions for a 4 x 5 x 2 model.

FLUXNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
1 1 1 1
```

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

```
-- -- ARRAY
             CONSTANT --
                          ----- BOX -----
                         I1 I2
                                  J1 J2
                                          K1 K2
EQUALS
                                     1*
   'FLUXNUM'
                                  1*
                                             1* / SET REGION 1
                                     2
   'FLUXNUM'
                         1
                                  1
                                          1
                                              1 / SET REGION 2
   'FLUXNUM'
                                              2 / SET REGION 3
```

²⁷ Although the FLUXNUM keyword is supported, OPM Flow currently does not support the flux boundary option.

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6.3.36 GDFILE - LOAD A GRID FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The GDFILE keyword loads a GRID file that contains the structural data for the grid as a set of topological cuboidal cells, and EGRID files that contain structural and property data. Note OPM Flow only supports reading in EGRID files at this time.

No.	Name	Description	Default
I	GRIDFILE	A character string enclosed in quotes that defines the GRID or EGRID file to be read in and be processed by OPM Flow.	None
		Again, OPM Flow only supports reading in EGRID files.	
2	FMTOPT	A defined character string that defines the format of the GRID or EGRID file to be read and should be set to one of the following:	U
		 FORMATTED: If the file is formatted as ASCII i.e. a text file, as oppose to a binary file. The option can be abbreviated to just the letter F. 	
		2) UNFORMATTED: If the file is in binary format, note this option can be abbreviated to just the letter U. This type of file is operating system dependent, that is a Microsoft Windows generated binary file cannot be read in by a Linux based system unlike formatted files.	
		If the keyword is omitted then the default is for binary file input.	
Note	<u>s:</u>		

The keyword is terminated by "/".

Table 6.32: GDFILE Keyword Description

See also the GRIDFILE keyword in the GRID section.

Examples

The first example shown below loads the NOR-OPM-A00-GRID.EGRID file in binary format from the same directory as the data file.

```
LOAD A GRID FILE
INCLUDE
         'NOR-OPM-A00-GRID.EGRID'
```

The next example loads the same EGRID file one directory above from where the data file is located.

```
LOAD a GRID FILE
INCLUDE
         '../NOR-OPM-A00-GRID.EGRID' /
```

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Revision: Rev-I

6.3.37 GDORIENT - Define GRID ORIENTATION PARAMETERS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

This keyword defines the grid orientation parameters for post-processing applications.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.38 GRID - Define the Start of the GRID Section of Keywords

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The GRID activation keyword marks the end of the RUNSPEC section and the start of the GRID section that defines the key grid property data for the simulator including the grid structure, porosity, permeability and other relevant grid property data.

There is no data required for this keyword.

Example



The above example marks the end of the RUNSPEC section and the start of the GRID section in the OPM Flow data input file.

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6.3.39 GRIDFILE - SET THE GRID FILE OUTPUT OPTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword controls the output of a standard GRID or extended GRID file, as well as the extensible EGRID file for post-processing applications. The extended and extensible GRID formats are comparable; however, the extensible GRID format is more compact and is the only format supported by OPM Flow.

No.	Name	Description	Default	
I	I NGRID	A positive integer that controls the output of the of the GRID geometry file:	0	
		0 - for no GRID file to be written out.		
		I - for the standard GRID file to be written out.		
		2 - for the extended GRID file to be written out.		
		Only the default value of zero is supported.		
2	NEGRID	A positive integer that controls the output of the of the EGRID geometry file:	I	
		0 - for no extensible GRID file to be written out.		
		I - for the extensible GRID file to be written out.		
		Only the default value of one is supported.		

The keyword is terminated by "/".

Table 6.33: GRIDFILE Keyword Description

Example

```
-- GRID FILE OUTPUT OPTIONS
-- GRID EGRID
-- OPTN OPTN
GRIDFILE
0 1
```

The above example defines that no GRID file will be written out and that the extensible GRID (that is the EGRID geometry format) file will be produced. This is the only configuration that OPM Flow supports

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6.3.40 GRIDUNIT - Define the Grid Units

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The GRIDUNIT keyword defines the units of the grid data. It is usually output by pre-processing software when exporting the grid geometry. The data is not used by OPM Flow intrinsically, but is merely written to the output EGRID file, as specified by the GRIDFILE keyword, for the use of post-processing software like OPM ResInsight.

No.	Name	Description	Default
I	GRIDUNIT	A character string that defines the units of the coordinates stated on the MAPAXES keyword, and should be set to:	METRES
		I) FIELD for field units	
		2) METRES for metric units, or	
		3) LAB for laboratory units	
2	MAPOPT	A character string that defines if the grid data are measured relative to the map, or relative to the origin as stated on the MAPAXES keyword. MAPOPT should either be left blank (the default) indicating the origin is relative to the origin on the MAPAXES keyword, or set equal to MAP measured relative to the map.	*

Notes:

- 1) Note the alternative spelling METRES, that is METERS is not recognized.
- 2) The keyword is terminated by "/".

Table 6.34: GRIDUNIT Keyword Description

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6.3.41 HEATCR - DEFINE RESERVOIR ROCK HEAT CAPACITY FOR ALL CELLS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The HEATCR keyword defines the reservoir rock volumetric heat capacity for all cells for when OPM Flow's thermal calculation is activated by the THERMAL keywords in the RUNSPEC section.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description						
		Field	Field Metric Laboratory						
I	HEATCR		HEATCR is an array of real positive numbers that define reservoir rock volumetric heat capacity of a grid block.						
		Repeat counts may be us	Repeat counts may be used, for example 3000*25.0						
		Btu/ft³/°R	kJ/m³/K	J/cm³/K	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.35: HEATCR Keyword Description

Note this keyword is incompatible with SPECROCK keyword in the PROPS section.

Example

```
-- DEFINE GRID BLOCK RESERVOIR ROCK HEAT CAPACITY
-- FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- HEATCR
```

300*32.0

The above example defines the reservoir rock volumetric heat capacity of 32.0 for each cell in the 300 grid block model.

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6.3.42 HEATCRT - Define Reservoir Rock Heat Capacity Temperature Dependence for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The HEATCRT keyword defines the reservoir rock volumetric heat capacity temperature dependence for all cells for when OPM Flow's thermal calculation is activated by the THERMAL keywords in the RUNSPEC section.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description						
		Field	Field Metric Laboratory						
I	HEATCRT	volumetric heat capacity	HEATCRT is an array of real positive numbers that define reservoir rock volumetric heat capacity temperature dependence of a grid block.						
		Repeat counts may be us							
		Btu/ft³/°R²	kJ/m³/K²	J/cm ³ /K ²	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.36: HEATCRT Keyword Description

Note this keyword is incompatible with SPECROCK keyword in the PROPS section.

The data for this keyword and the HEATCR keyword are use to calculate the reservoir rock volumetric heat capacity temperature dependence using the following relationship:

Heat Capacity of Rock = HEATCR
$$(Temp - Temp_{ref}) + \frac{\text{HEATCRT}(Temp - Temp_{ref})^2}{2}$$
 (6.3)

Example

```
-- DEFINE RESERVOIR ROCK HEAT CAPACITY TEMPERATURE DEPENDENCE
-- FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- HEATCRT
300*0.05
```

The above example defines the reservoir rock volumetric heat capacity temperature dependence of 0.05 for each cell in the 300 grid block model.

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6.3.43 INIT - ACTIVATES THE INIT FILE OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on the writing of the INIT file that contains the static data specified in the GRID, PROPS and REGIONS sections. For example, the PORO, PERM and NTG arrays from the GRID section. The data is used in post-processing software, for example ResInsight, to visualize the static grid properties.

The INIT file can either be written out in formatted form as ASCII i.e. text files, if the FMTOUT keyword has been activated, or binary format if the FMTOUT keyword has <u>not</u> been activated. Normally, this option is always activated and when activated the binary form of the file is used

There is no data required for this keyword.

Example

- -

ACTIVATE WRITING THE INIT FILE FOR POST-PROCESSING

INIT

The above example switches on the writing of the INIT file for post-processing in ResInsight.

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6.3.44 INRAD - Define the Inner Radius of a Radial Grid

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

INRAD²⁸ defines the inner radius of the reservoir model for a radial grid geometry. The RADIAL keyword in the RUNSPEC should be activated to indicate that radial geometry is being used.

	Field	Metric	Laboratory					
NRAD	A single real positive num	single real positive number defining the inner radius of a radial grid.						
	feet	m	cm	None				
1	NRAD							

Table 6.37: INRAD Keyword Description

See also the DR, DRV, DTHETA, DTHETAV and TOPS keywords to fully define a Radial Grid.

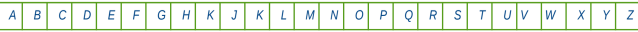
This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

```
-- INNER RADIUS OF FIRST GRID BLOCK IN THE RADIAL DIRECTION
-- INRAD
0.25
```

The above example defines the inner radius of a radial grid to be 0.25 feet.

Radial grids are not currently implemented in OPM Flow version 2018-04 but is expected to be incorporated in the next release.



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6.3.45 ISOLNUM - Define the Independent Reservoir Regions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The ISOLNUM keyword defines areas of the grid that consists of isolated reservoirs where the only form of communication between the reservoirs is via wellbore connections This enables the reservoir flow equations to be solved independently for greater computational efficiency.

No.	Name	Description	Default
1	ISOLNUM	ISOLNUM defines an array of positive integers assigning a grid cell to a particular isolated reservoir region.	I
		The maximum number of ISOLNUM regions is set by the NRFREG variable on the REGDIMS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) Setting ISOLNUM for a cell to zero makes the cell inactive.
- 3) The keyword is terminated by "/".

Table 6.38: ISOLNUM Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

The example below defines three separate independent reservoirs; the first reservoir covers the whole grid and layers I to 50, reservoir two cover the whole grid and layers 52 to 150, and finally the third reservoir again covers the whole grid but with layers 152 to 300. The layers 51 and 151 are shale layers made inactive by setting ISOLNUM to zero.

```
-- ARRAY
                    CONSTANT --
                                ----- BOX -----
                                I1 I2 J1 J2 K1 K2
EQUALS
        'ISOLNUM'
                                            1*
                                                 1
                                                     50 / DEFINED RESERVOIR 1
                                         1*
                                            1*
                                    1*
                                                 51 51 / DEFINED A SHALE
                                1*
        'ISOLNUM'
                                         1* 1*
                                    1*
                                                52 150 / DEFINED RESERVOIR 2
        'ISOLNUM'
                                    1*
                                         1*
                                            1*
        'ISOLNUM'
                                                151 151 / DEFINED A SHALE
        'ISOLNUM'
                                                152 300 / DEFINED RESERVOIR 3
```

Note the above example has no effect as the keyword is ignored by the simulator.

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6.3.46 JFUNC - ACTIVATES THE LEVERETT J-FUNCTION OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

JFUNC keyword activates Leverett-J-Function²⁹ option which is a commonly used technique to normalize capillary pressure base on laboratory measured core plugs porosity and permeability values and the resulting capillary pressure data.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
I	JFOPT		defines which capillary d to, based on the following		вотн				
		I) WATER: apply pressure data of	the J-Function option to only.	the water-oil capillary					
		2) GAS: apply the data only.	J-Function option to the g	gas-oil capillary pressure					
		3) BOTH: apply t oil capillary pre	he J-Function option to thessure data.	ne water-oil and the gas-					
2	OWSTEN		A positive real number that defines oil-water surface tension used to denormalized J-Function data entered in the PROPS section						
		dynes/cm	dynes/cm	dynes/cm	None				
3	OGSTEN		A positive real number that defines oil-gas surface tension used to denormalized J-Function data entered in the PROPS section						
		dynes/cm	dynes/cm	dynes/cm	None				
4	ALPHA	porosity term in the J-Fr	A positive real value that defines an alternative power value for the porosity term in the J-Function equation, that is instead of $\sqrt{\frac{k}{\phi}}$ use $\frac{k^{0.5}}{\phi^{\alpha}}$ instead in the transformation.						
5	BETA	A positive real number that defines an alternative power value for the permeability term in the J-Function equation, that is instead of $\sqrt{\frac{k}{\phi}}$ use $\frac{k^{\beta}}{\phi^{0.5}}$ instead in the transformation.							

²⁹ Leverett, M. C.; "Capillary Behaviour in Porous Solids", Trans. AIME (1941) 142, 152-168.



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No.	Name	Description					
6	PERM	PERM is a character string that sets the permeability array to be used in the transform, based on the following options:	XY				
		I) X: use the PERMX array.					
		2) XY: use the average of the PERMX and PERMY arrays.					
		3) Y: use the PERMY array.					
		4) Z: use the PERMZ array.					

Notes:

1) The keyword is terminated by "/".

Table 6.39: JFUNC Keyword Description

Just like the relative permeability data capillary pressure data are measured on core plugs with varying quality and perhaps from different reservoirs. It is therefore necessary to determine averaged data, before employing the data in engineering calculations. This is commonly done by using the Leverett J-function³⁰, which is defined as:

$$J(S_w) = \frac{P_{c,res}(S_w) \sqrt{\frac{k}{\varphi}}}{\sigma}$$
(6.4)

Revision: Rev-I

Where:

J (S_w) = dimensionless function of water saturation

P_c (S_w) = capillary pressure (kPa) k = permeability, (m²)

 $\varphi = \text{permeability, (iii)}$ $\varphi = \text{porosity (fraction)}$

 σ = interfacial tension (mN/m)

 Θ = contact angle

Sometimes the equation is stated with the $\cos \theta$ term included, that is:

$$J(S_w) = \frac{P_{c,res}(S_w) \sqrt{\frac{k}{\varphi}}}{\sigma \cos \Theta}$$
 (6.5)

Since the above function is just a normalizing function, then units are not important, as long as when we denormalize the *average* curve we use the same unit set. Secondly, if all the capillary pressure data has been converted to reservoir conditions, we actually ignore the denominator as it is a constant, and we can therefore just use:

$$J(S_w) = P_{c,res}(S_w) \sqrt{\frac{k}{\varphi}}$$
 (6.6)

However, in the simulator it is necessary to use the formal definition as outlined in equation (6.4). In addition to the standard the equation the keyword allows for de-normalizing the curve to use alternative power functions instead of the standard 0.5 used in equation (6.4), that is:

³⁰ Leverett, M. C.; "Capillary Behaviour in Porous Solids", Trans. AIME (1941) 142, 152-168.



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 $J(S_w) = \frac{P_{c,res}(S_w) \left(\frac{k^{\beta}}{\varphi^{\alpha}}\right)}{\sigma}$ (6.7)

Revision: Rev-I

Where:

 $J(S_w)$ = dimensionless function of water saturation

 $P_c(S_w)$ = capillary pressure (kPa) k = permeability, (m²) ϕ = porosity (fraction)

 σ = interfacial tension (mN/m)

 Θ = contact angle

α = porosity power valueβ = permeability value

The JFUNC keyword allows the data entered as capillary pressure in the saturation tables, for example, by using the SGFN and SWFN keywords in the PROPS section to be treated as J-functions instead, and to denormalize these curves for each active cell in the model using the options and values defined with the JFUNC keyword combined with a cells porosity and permeability values.

Example

```
DEFINE LEVERETT J-FUNCTION PARAMETERS
                 OILWAT
                         GASOIL PORO
                                           PERM
                                                   PERM
         OPTN
                 SDENS
                          SDEN
                                  ALPHA
                                           BETA
                                                   OPTN
JFUNC
         WATER
                 22.5
                          1*
                                  0.5
                                           0.5
                                                   XY
```

The above example results in the oil-water capillary pressure data entered on the SWFN keyword in the PROPS section being treated a J-Functions, and that the J-Function s should be de-normalized using an oil-water surface density of 22.5 dynes/cm, using the default power values and the average of the PERMX and PERMY values for each grid block.

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6.3.47 MAPAXES- DEFINE THE MAP ORIGIN INPUT DATA

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

MAPAXES specifies the origin of the map used to create the grid. It is usually output by pre-processing software when exporting the grid geometry. The data is not used by OPM Flow intrinsically, but is merely written to the output EGRID file, as specified by the GRIDFILE keyword, for the use of post-processing software like OPM ResInsight.

No.	Name	Description						
		Field (feet)	Metric (metres)	Laboratory (metres)				
ı	ΧI	XI is a real number that	defines the x co-ordinate	e of a point on the y-axis.	None			
2	ΥI	YI is a real number that of	YI is a real number that defines the y co-ordinate of a point on the y-axis.					
3	X2	X2 is a real number that	X2 is a real number that defines the x co-ordinate of the origin.					
4	Y2	Y2 is a real number that of	defines the y co-ordinate	of the origin.	None			
5	X3	X3 is a real number that axis.	X3 is a real number that defines the x co-ordinate of a point on the x-axis.					
6	Y3	Y3 is a real number that o	Y3 is a real number that defines the y co-ordinate of a point on the x-axis.					

Notes:

Table 6.40: MAPAXES Keyword Description

Example

```
----- MAPAXES ------
     X1
         Y1 X2
                      X3 Y3
MAPAXES
     0.0
                   0.0
          100.0 0.0
                       100.0 0.0 /
```

The above example defines the map axes to be exported to the grid file for use by post-processing software.

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¹⁾ The keyword is terminated by "/".

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6.3.48 MAPUNITS - Define the Map Axes Units

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The MAPUNITS keyword defines the units of the coordinates stated on the MAPAXES keyword. It is usually output by pre-processing software when exporting the grid geometry. The data is not used by OPM Flow intrinsically, but is merely written to the output EGRID file, as specified by the GRIDFILE keyword, for the use of post-processing software like OPM ResInsight.

No.	Name	Description	Default
1	MAPUNITS	A character string that defines the units of the coordinates stated on the MAPAXES keyword, and should be set to:	METRES
		FIELD for field units	
		2) METRES for metric units, or	
		3) LAB for laboratory units	

Notes:

- 1) Note the alternative spelling of METRES, that is METERS is not recognized.
- 2) The keyword is terminated by "/".

Table 6.41: MAPUNITS Keyword Description

Example

- -

SET THE MAP UNITS FOR THE MAPAXES KEYWORD

MAPUNITS

METRES

The above example specifies the units on the MAPAXES to be the default METRES.

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6.3.49 MAXVALUE - Sets a Maximum Value for an Array Element

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDUL	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
----------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

6.3.50 DESCRIPTION

The MAXVALUE keyword sets a maximum value for the specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the MAXVALUE keyword is being used.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	CONSTANT is a positive integer or positive real value that an ARRAY element will be reset to if an element in the defined input BOX, as defined by items (3) to (8), is grater than CONSTANT. CONSTANT has in the same units as the ARRAY property.	None
3	П	The lower bound of the array in the I-direction to be modified must be greater than or equal to one and less than or equal to I2 and NX.	I
4	12	The upper bound of the array in the I-direction to be modified must be greater than or equal to II and less than or equal to NX	NX
5	JI	The lower bound of the array in the J-direction to be modified must be greater than or equal to one and less than or equal to J2 and NY.	1
6	J2	The upper bound of the array in the J-direction to be modified must be greater than or equal to JI and less than or equal to NY.	NY
7	KI	The lower bound of the array in the K-direction to be modified must be greater than or equal to one and less than or equal to K2 and NZ.	I
8	K2	The upper bound of the array in the K-direction to be modified must be greater than or equal to KI and less than or equal to NZ.	NZ

Notes:

- 1) Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.42: MAXVALUE Keyword Description

The applicable arrays for each section are defined in Table 6.43 on the following page.

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MAXVALUE Keyword and Variable Options by Section GRID **EDIT** PROPS **REGIONS** SOLUTION **SUMMARY SCHEDULE** DX DEPTH SWL DY **PORV SWCR** DΖ TRANX SWU PERMX TRANY SGL PERMY TRANZ SGCR PERMZ SGU **DIFFX** MULTX DIFFY KRW MULTY DIFFZ KRO MULTZ KRG TRANR DR TRANTHT **PCG** THETA DIFFR PCW PERMR **DIFFTHT** PERMTHT **DZNET** PORO NTG **FLUXNUM** MULTNUM **MPANUM DIFFX** DIFFY DIFFZ **DIFFR**

Revision: Rev-I

Table 6.43: MAXVALUE Keyword Applicable Arrays by Section

Example

DIFFTHT

The above example resets the maximum values for the PERMX, PERMY and PERMZ, arrays to 100.0, 100.0 and 10.0, respectively, for all cells in layer five.

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6.3.51 MINPV - SET A MINIMUM GRID BLOCK PORE VOLUME THRESHOLD FOR ALL CELLS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

MINPV defines a minimum threshold pore volume that makes all grid blocks whose pore volume is below this value inactive in the mode (inactive cells are not used in OPM Flow calculations). Note this keyword is different to the MINPVV keyword in the GRID section that sets a minimum threshold pore volume for individual cells in the model.

No.	Name		Description						
		Field							
I	MPVTHRES		MPVTHRES is a real positive number that defines the minimum pore volume for a cell to be active in the model.						
		rb	rm³	rcc					
		1.0e-6	1.0e-6 1.0e-6 1.0e-6						
Notes	s:								

Table 6.44: MINPV Keyword Description

See also the PINCH keyword for the treatment of inactive grid cells and pinch-outs.

Example

```
MINIMUM PORE VOLUME FOR ACTIVE CELLS
MINPV
500.0
```

The above example defines 500 rb (or m³) as the minimum pore volume for a cell to be active in the model.

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¹⁾ The keyword is terminated by "/".

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6.3.52 MINPVV - SET A MINIMUM GRID BLOCK PORE VOLUME THRESHOLD FOR INDIVIDUAL CELLS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

MINPVV is an array that defines the minimum threshold pore volume for each cell, that makes grid blocks whose pore volume is below this value inactive in the model (inactive cells are not used in OPM Flow calculations). Note this keyword is different to the MINPV keyword in the GRID section that sets a constant minimum threshold pore volume for all cells in the model.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

No.	Name			Default		
		Field	Field Metric Laboratory			
I	MPVTHRES		f real positive numbers the			
		rb				
		1.0e-6	1.0e-6	1.0e-6	Defined	

Notes:

- 1) The number of entries should correspond to the NX \times NY \times NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.45: MINPVV Keyword Description

See also the PINCH keyword for the treatment of inactive grid cells and pinch-outs.

Example

```
DEFINE A BOX GRID FOR THE BOTTOM TWO LAYERS OF A 100 X 100 X 20 MODEL
           ----- BOX -----
- -
         I1 I2
                  J1 J2
                           K1 K2
B<sub>0</sub>X
                      1*
                            19 20 / SELECT THE BOTTOM LAYER
             1*
- -
         MINIMUM PORE VOLUME FOR INDIVIDUAL CELLS TO BE ACTIVE
MINPVV
         10000*500.0
                        10000*750.0
         RESET THE INPUT BOX TO BE THE FULL MODEL
- -
ENDBOX
```

The above example defines 500 rb (or m^3) as the minimum pore volume for all cells in layer 19 to be active in the model, and 750 rb (or m^3) as the minimum pore volume for all cells in layer 20.

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6.3.53 MINVALUE - SET A MINIMUM VALUE FOR AN ARRAY ELEMENT

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

6.3.54 DESCRIPTION

The MINVALUE keyword sets a minimum value for the specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the MINVALUE keyword is being used.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	CONSTANT is a positive integer or positive real value that an ARRAY element will be reset to if an element in the defined input BOX, as defined by items (3) to (8), is less than CONSTANT. CONSTANT has in the same units as the ARRAY property.	None
3	П	A positive integer that defines the lower bound of the array in the I-direction to be modified must be greater than or equal to one and less than or equal to I2 and NX.	I
4	12	A positive integer that defines the upper bound of the array in the I-direction to be modified must be greater than or equal to II and less than or equal to NX	NX
5	JI	A positive integer that defines the lower bound of the array in the J-direction to be modified must be greater than or equal to one and less than or equal to J2 and NY.	I
6	J2	A positive integer that defines the upper bound of the array in the J-direction to be modified must be greater than or equal to JI and less than or equal to NY.	NY
7	KI	A positive integer that defines the lower bound of the array in the K-direction to be modified must be greater than or equal to one and less than or equal to K2 and NZ.	I
8	K2	A positive integer that defines the upper bound of the array in the K-direction to be modified must be greater than or equal to KI and less than or equal to NZ.	NZ

Notes:

- I) Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.46: MINVALUE Keyword Description

The applicable arrays for each section are defined in Table 6.47 on the following page.



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	EQUALS Keyword and Variable Options by Section								
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE			
DX	DEPTH	SWL							
DY	PORV	SWCR							
DZ	TRANX	SWU							
PERMX	TRANY	SGL							
PERMY	TRANZ	SGCR							
PERMZ	DIFFX	SGU							
MULTX	DIFFY	KRW							
MULTY	DIFFZ	KRO							
MULTZ	TRANR	KRG							
DR	TRANTHT	PCG							
THETA	DIFFR	PCW							
PERMR	DIFFTHT								
PERMTHT									
DZNET									
PORO									
NTG									
FLUXNUM									
MULTNUM									
MPANUM									
DIFFX									
DIFFY									
DIFFZ									
DIFFR									
DIFFTHT									

Table 6.47: MINVALUE Keyword Applicable Arrays by Section

Example

The above example resets the minimum values for the PERMX, PERMY and PERMZ, arrays to 1.0, 1.0 and 0.1, respectively, for all cells in layer five.

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6.3.55 MULTFLT - MULTIPLY THE TRANSMISSIBILITY OF A DEFINED FAULT BY A CONSTANT

RUNSPEC GRID EDIT PROPS REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------------------------------	----------	---------	----------

Revision: Rev-I

Description

MULTFLT enables the transmissibilities across defined faults, as declared by the FAULTS keyword, to be modified. They keyword allows for the re-scaling of the existing fault transmissibilities calculated by OPM Flow, for example setting a fault to be completely sealing by setting the multiplier to zero..

No.	Name	Description	Default
I	FLTNAME	FLTNAME is a character string enclosed in quotes with a maximum length of eight characters, that defines the name of the fault that FLTMULT will be applied to. FLTNAME must have previously been defined using the FAULTS keyword in GRID section	None
2	FLT-TRS	A positive real number that sets the transmissible multiplier to be applied to the FLTNAME transmissibilities positive real number that sets the transmissible multiplier to be applied to the FLTNAME transmissibilities.	1.0
3	FLT-DIF	A positive real number that sets the diffusitivity multiplier to be applied to the FLTNAME diffusivities.	1.0
		This option should only be used if the Diffusion option has been made activate by the DIFFUSE keyword in the RUNSPEC section.	
		OPM Flow does not support the Diffusion option.	

Notes:

- Repeated entries of the same FLTNAME will result in all but the last entry being overwritten. I)
- Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.48: MULTFLT Keyword Description

Example

```
MODIFY THE TRANSMISSIBILITES ACROSS DEFINED FAULTS
         FAULT
                           TRANS
                                            DIFUSS
         NAME
                           MULTIPLIER
                                            MULTIPLIER
MULTFLT
         'FAULT01'
                           0.0
                                                                 / FAULT MULTIPLIERS
         'FAULT02'
                           0.0
                                                                 / FAULT MULTIPLIERS
         'FAULT03'
                                                                 / FAULT MULTIPLIERS
                           0.0
```

The above example sets the fault transmissibility multiplier for defined faults named FAULT01, FAULT02, and FAULT03 to zero making the faults sealing in the model.

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6.3.56 MULTIPLY - MULTIPLY A SPECIFIED ARRAY BY A CONSTANT

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

6.3.57 DESCRIPTION

The MULTIPLY keyword multiplies a specified array or part of an array by a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the keyword is being used.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	An integer or real value that the ARRAY will be set to in the same units as the ARRAY property.	None
3	П	A positive integer that defines the lower bound of the array in the I-direction to be modified must be greater than or equal to one and less than or equal to I2 and NX.	I
4	12	A positive integer that defines the upper bound of the array in the I-direction to be modified must be greater than or equal to II and less than or equal to NX	NX
5	JI	A positive integer that defines the lower bound of the array in the J-direction to be modified must be greater than or equal to one and less than or equal to J2 and NY.	I
6	J2	A positive integer that defines the upper bound of the array in the J-direction to be modified must be greater than or equal to JI and less than or equal to NY.	NY
7	KI	A positive integer that defines the lower bound of the array in the K-direction to be modified must be greater than or equal to one and less than or equal to K2 and NZ.	I
8	K2	A positive integer that defines the upper bound of the array in the K-direction to be modified must be greater than or equal to KI and less than or equal to NZ.	NZ

Notes:

- Where NX, NY and NZ are the dimensions of the model as defined on the DIMENS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.49: MULTIPLY Keyword Description

The applicable arrays for each section are defined in Table 6.50 as shown on the next page.



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	MULTIPLY Keyword and Variable Options by Section									
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE				
DX	DEPTH	SWL	ENDNUM	PRESSURE						
DY	PORV	SWCR	EQLNUM	SWAT						
DZ	TRANX	SWU	FIPNUM	SGAS						
PERMX	TRANY	SGL	IMBNUM	RV						
PERMY	TRANZ	SGCR	MISCNUM	RS						
PERMZ	DIFFX	SGU	PVTNUM	TBLK						
MULTX	DIFFY	KRW	ROCKNUM	GI						
MULTY	DIFFZ	KRO	SATNUM	OILAPI						
MULTZ	TRANR	KRG	WH2NUM	SALT						
DR	TRANTHT	PCG		GASCONC						
THETA	DIFFR	PCW		SOLVCONC						
PERMR	DIFFTHT			SOLVFRAC						
PERMTHT				SFOAM						
DZNET				SPOLY						
PORO										
NTG										
FLUXNUM										
MULTNUM										
MPANUM										
DIFFX										
DIFFY										
DIFFZ										
DIFFR										
DIFFTHT										

Table 6.50: MULTIPLY Keyword Applicable Arrays by Section

Example

The above example multiples the PERMZ property array by 0.5 throughout the model.

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6.3.58 MULTIREG - MULTIPLY AN ARRAY BY A CONSTANT BASED ON A REGION NUMBER

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

6.3.59 DESCRIPTION

The MULTIREG keyword multiplies an array or part of an array by a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTIREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the MULTIREG keyword is being used.

No.	Name	Description	Default
I	ARRAY	The name of the array to be modified. This is the keyword name identifying the property and is up to eight characters in length and enclosed in quotes.	None
2	CONSTANT	An integer or real value to multiply the ARRAY by in the same units as the ARRAY property for a given REGION.	0
3	REGION NUMBER	REGION NUMBER is a positive integer representing the region for which the CONSTANT in (2) should be applied	None
4	REGION ARRAY	The REGION ARRAY to use for applying the CONSTANT in (2) based on the REGION NUMBER in (3). REGION ARRAY can have the following values:	М
		F for the FLUXNUM array	
		2) M for the MULTNUM array	
		3) O for the OPERNUM array	

Notes:

- Where the REGION NUMBER should be less than or equal to the maximum number of regions as defined on the REGDIMS keyword for the FIPNUM and OPERNUM arrays or the GRIDOPTS keyword for the MULTNUM array in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.51: MULTIREG Keyword Description

The applicable arrays for each section are defined in Table 6.52 as shown on the following page.

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Revision: Rev-I

	MU	JLTREG Keyw	ord and Variable	Options by Sec	tion	
GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
DX	DEPTH	SWL	ENDNUM	PRESSURE		
DY	PORV	SWCR	EQLNUM	SWAT		
DZ	TRANX	SWU	FIPNUM	SGAS		
PERMX	TRANY	SGL	IMBNUM	RV		
PERMY	TRANZ	SGCR	MISCNUM	RS		
PERMZ	DIFFX	SGU	PVTNUM	TBLK		
MULTX	DIFFY	KRW	ROCKNUM	GI		
MULTY	DIFFZ	KRO	SATNUM	OILAPI		
MULTZ	TRANR	KRG	WH2NUM	SALT		
DR	TRANTHT	PCG		GASCONC		
THETA	DIFFR	PCW		SOLVCONC		
PERMR	DIFFTHT			SOLVFRAC		
PERMTHT				SFOAM		
DZNET				SPOLY		
PORO						
NTG						
FLUXNUM						
MULTNUM						
MPANUM						
DIFFX						
DIFFY						
DIFFZ						
DIFFR						
DIFFTHT						

Table 6.52: MULTIREG Keyword Applicable Arrays by Section

Example

```
-- FIRST DEFINE THE PROPERTY ARRAYS AND MULTNUM ARRAYS FOR 10 X 10 X 20 MODEL
-- -- ARRAY CONSTANT -- ------ BOX -----
                          I1 I2 J1 J2 K1 K2
EQUALS
                                   1*
1*
                                                   / PORO TO 0.20 IN MODEL
/ PERMX TO 0.10 IN MODEL
   'PORO'
              0.2000
                           1*
                              1*
                                       1*
                                            1* 1*
                                            1*
   'PERMX'
                           1*
                             1*
                                               1*
                                       1*
              100.00
                                       1*
                           1*
                              1*
                                   1*
                                            1* 1* / MULTNUM IN MODEL
  'MULTNUM'
              1
  'MULTNUM'
                              5
                                   1
                                       5
                                            6
                                                6
                                                    / MULTNUM IN MODEL
                                   1*
                                       1*
   'MULTNUM'
                              1*
                                            10 10 / MULTNUM IN MODEL
-- NOW RESET PORO AND PERMX BASED ON THE MULTNUM REGION NUMBER
-- MULTIPLY AN ARRAY BY A CONSTANT BASED ON A REGION NUMBER
                                REGION ARRAY
     ARRAY
               CONSTANT REGION
- -
               VALUE
                        NUMBER
                                M / F / O
MULTIREG
     'PORO'
               1.050
               1.100 2
    'P0R0'
    'PORO'
               0.950 3
                       1
     'PERMX'
               1.25
                                  М
     'PERMX'
                1.30
     'PERMX'
                0.90
```

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The example first defines the PORO and PERMX property arrays for the model and then sets the MULTNUM array to I for all cells in the model, after which selected areas of model are assigned various MULTNUM integer values. The MULTIREG can then be invoked to multiple the PORO and PERMX arrays by a constant for the various MULTNUM regions.

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6.3.60 MULTNUM - Define the Multiple Transmissibility Regions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The MULTNUM keyword defines the inter-region transmissibility region numbers for each grid block, as such there must be one entry for each cell in the model. The array can be used with the EQUALREG, ADDREG, COPYREG, MULTREGP and MULTREGT keywords in calculating various grid properties in the GRID section.

No.	Name	Description	Default
I	MULTNUM	MULTNUM defines an array of positive integers assigning a grid cell to a particular inter-region transmissibility region.	I
		The maximum number of MULTNUM regions is set by the NRMULT variable on the GRIDOPTS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.53: MULTNUM Keyword Description

Examples

The example below sets three MULTNUM regions for a $4 \times 5 \times 2$ model.

MULTNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
```

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

```
CONSTANT --
                           ----- BOX -----
-- -- ARRAY
                           I1 I2
                                  J1 J2
                                            K1 K2
EQUALS
                           1*
                              1*
                                       1*
                                            1*
   'MULTNUM'
                                    1*
                                                1* / SET REGION 1
              1
   'MULTNUM'
              2
                               2
                                       2
                                                1 / SET REGION 2
                           1
                                   1
                                            1
   'MULTNUM'
                                                2 / SET REGION 3
```

One can then increase PERMX by 25% in region three only.

```
-- MULTIPLY AN ARRAY BY A CONSTANT BASED ON A REGION NUMBER
-- ARRAY CONSTANT REGION REGION ARRAY
-- VALUE NUMBER M / F / O
MULTIREG 'PERMX' 1.25 3 M /
```

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6.3.61 MULTPV - MULTIPLY CELL PORE VOLUMES BY A CONSTANT

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

MULTPV multiples the pore volumes of a cell by a real positive constant for all the cells in the model via an array. An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTPV keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name	Description	Default
I	MULTPV	MULTPV is an array of real positive numbers assigning the pore volume multipliers for each cell in the model.	1.0
		Repeat counts may be used, for example 20*100.0.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.54: MULTPV Keyword Description

See also the MULTREGP for scaling the cell pore volumes by region numbers.

Examples

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
- -
         ----- BOX -----
- -
         I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
                                                             / DEFINE BOX AREA
             10
                  1
                       6
                            1
                                3
         SET MULTX+ TRANSMISSIBILITY MULTIPLIERS
- -
MULTPV
18*0.0500
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
_ _
ENDBOX
```

The above example defines a 0.05 scaling multiplier for the 18 cells defined by the preceding BOX statement. The ENDBOX keyword resets the input box to the full grid.

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6.3.62 MULTREGP- MULTIPLY PORE VOLUMES BASED ON REGION NUMBER

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

6.3.63 DESCRIPTION

The MULTREGP keyword multiplies the pore volume of a cell by a constant for all cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTREGP keyword is read by the simulator. The constant should be a real number.

No.	Name	Description	Default
I	REGION NUMBER	REGION NUMBER is a positive integer representing the region for which the CONSTANT in (2) should be applied.	None
2	CONSTANT	A real value to multiply the pore volume by for a given REGION.	I
3	REGION ARRAY	The REGION ARRAY to use for applying the CONSTANT in (2) based on the REGION NUMBER in (I). REGION ARRAY can have the following values:	М
		I) F for the FLUXNUM array	
		2) M for the MULTNUM array	
		3) O for the OPERNUM array	

Notes:

- Where the REGION NUMBER should be less than or equal to the maximum number of regions as defined on the REGDIMS keyword for the FIPNUM and OPERNUM arrays or the GRIDOPTS keyword for the MULTNUM array in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.55: MULTREGP Keyword Description

Example

```
-- RESET PORE VOLUME FOR DIFFERENT REGIONS
      REGION
                PORV
                               REGION ARRAY
      NUMBER
                MULT
                               M / F / O
- -
MULTREGP
                 1.0456573
                                                Fault Block 1
                                                Fault Block 2
         2
                 0
                               М
         3
                 0.9756715
                                                Fault Block 3
                                                 Inactive Blocks
```

The above example re-scales the pore volumes for MULTNUM regions one and three and makes regions two and four inactive by setting their pore volumes to zero.

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6.3.64 MULTREGT- MULTIPLY TRANSMISSIBILITIES BETWEEN REGIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

6.3.65 DESCRIPTION

The MULTREGT keyword multiplies the transmissibility between two regions by a constant. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTREGT keyword is read by the simulator. The constant should be a real number.

No.	Name	Description	Default
I	REGIONI	A positive integer value that defines the from REGION number for which the CONSTANT in (2) should be applied.	None
2	REGION2	A positive integer value that defines the to REGION number for which the CONSTANT in (2) should be applied.	None
3	CONSTANT	A real value to multiply the transmissibility between REGION1 and REGION2.	0
4	DIR	A character string that defines the direction to apply the transmissibility multiplier between the two regions, should be set to one of the following X, Y, Z, XY, YX, XZ, or XYZ.	XYZ
	TYPE	 A character string that defines the type of connections the transmissibility multiplier should be applied to, should be one of the following: NNC – Only apply the transmissibility multiplier between REGION1 and REGION2 to non-neighbor connections. NONNC – Do not apply the transmissibility multiplier between REGION1 and REGION2 to non-neighbor connections. ALL - Apply the transmissibility multiplier between REGION1 and REGION2 to all connections. 	ALL
4	REGION ARRAY	The REGION ARRAY to use for applying the CONSTANT in (2) based on the REGION NUMBER in (1). REGION ARRAY can have the following values: 1) F for the FLUXNUM array 2) M for the MULTNUM array	М
		3) O for the OPERNUM array	

Notes:

- Where the REGION NUMBER should be less than or equal to the maximum number of regions as defined on the REGDIMS keyword for the FIPNUM and OPERNUM arrays or the GRIDOPTS keyword for the MULTNUM array in the RUNSPEC section.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.56: MULTREGT Keyword Description

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Revision: Rev-I

Example

```
-- SET TRANSMISSIBILITES ACROSS DIFFERENT RESERVOIRS TO ZERO TO ISOLATE
RESERVOIRS
     REGION
              REGION
                       TRANS
                               DIREC NNC
                                             REGION ARRAY
     FROM
                       MULT
                               0PT
                                       0PTS
                                             M / F / O
MULTREGT
     1*
              1*
                       0.0
                               1*
                                      'ALL'
                                                         / ALL REGIONS SEALED
```

The above example isolates all regions from one another by setting the transmissibility for the MULTNUM regions to zero for all directions and for connections types.

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6.3.66 MULTX - MULTIPLY CELL TRANSMISSIBILITY IN THE +X DIRECTION

RUNSPEC GRID EDIT PROPS REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------------------------------	----------	---------	----------

Revision: Rev-I

Description

MULTX multiples the transmissibility between two cell faces in the +X direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, J, K) between the cells (I, J, K) and (I+I, J, K).

An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTX keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name	Description	Default
I	MULTX+	MULTX+ is an array of real positive numbers assigning the transmissibility multipliers in the +X direction to each cell face in the model.	1.0
		Repeat counts may be used, for example 20*100.0.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.57: MULTX Keyword Description

See also the MULTX-, MULTY, MULTZ- and MULTZ- keywords for scaling transmissible between grid cells.

Examples

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
         I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
         10 10
                  1
                      6
                           1
                               3
                                                           / DEFINE BOX AREA
         SET MULTX+ TRANSMISSIBILITY MULTIPLIERS
MULTX
18*0.300
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The above example defines a 0.3 scaling multiplier for the 18 cells defined by the preceding BOX statement. The ENDBOX keyword resets the input box to the full grid.

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6.3.67 MULTX- - MULTIPLY CELL TRANSMISSIBILITY IN THE -X DIRECTION

RUNSPEC GRID EDIT PROPS REGIONS	SOLUTION	SUMMARY	
---------------------------------	----------	---------	--

Revision: Rev-I

Description

MULTX- multiples the transmissibility between two cell faces in the -X direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I-I, I, K) and (I, J, K).

An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTX- keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name	Description	Default
I	MULTX-	MULTX- is an array of real positive numbers assigning the transmissibility multipliers in the -X direction to each cell face in the model.	1.0
		Repeat counts may be used, for example 20*100.0.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/". 2)

Table 6.58: MULTX- Keyword Description

See also the MULTX, MULTY, MULTY-, MULTZ and MULTZ- keywords for scaling transmissible between grid cells.

Examples

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
         I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
                                                           / DEFINE BOX AREA
         10 10
                  1
                      6
                           1
                               1
         SET MULTX TRANSMISSIBILITY MULTIPLIERS CELLS
MULTX-
6*0.500
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The above example defines a 0.5 scaling multiplier for the six cells defined by the preceding BOX statement. The ENDBOX keyword resets the input box to the full grid.

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6.3.68 MULTY - MULTIPLY CELL TRANSMISSIBILITY IN THE +Y DIRECTION

RUNSPEC GRID EDIT PROPS REGIONS	SOLUTION	SUMMARY	
---------------------------------	----------	---------	--

Revision: Rev-I

Description

MULTY multiples the transmissibility between two cell faces in the +Y direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, J, K) between the cells (I, J, K) and (I, J+I, K).

An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTY keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name	Description	Default
I	MULTY+	MULTY+ is an array of real positive numbers assigning the transmissibility multipliers in the +Y direction to each cell face in the model.	1.0
		Repeat counts may be used, for example 20*100.0.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.59: MULTY Keyword Description

See also the MULTY-, MULTX-, MULTZ and MULTZ- keywords for scaling transmissible between grid cells.

Examples

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
         I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
                                                           / DEFINE BOX AREA
         10 10
                  1
                      6
                           1
                               3
         SET MULTX+ TRANSMISSIBILITY MULTIPLIERS
MULTY
18*0.300
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The above example defines a 0.3 scaling multiplier for the 18 cells defined by the preceding BOX statement. The ENDBOX keyword resets the input box to the full grid.

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6.3.69 MULTY- - MULTIPLY CELL TRANSMISSIBILITY IN THE -Y DIRECTION

RUNSPEC GRID EDIT PROPS REGIONS	SOLUTION	SUMMARY	
---------------------------------	----------	---------	--

Revision: Rev-I

Description

MULTY- multiples the transmissibility between two cell faces in the -Y direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, J, K) between the cells (I, J-I, K) and (I, J, K).

An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTY- keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name	Description	Default
I	MULTY-	MULTY- is an array of real positive numbers assigning the transmissibility multipliers in the -Y direction to each cell face in the model.	1.0
		Repeat counts may be used, for example 20*100.0.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.60: MULTY- Keyword Description

See also the MULTY, MULTX-, MULTZ and MULTZ- keywords for scaling transmissible between grid cells.

Examples

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
         I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
                                                           / DEFINE BOX AREA
         10 10
                  1
                      6
                           1
                               1
         SET MULTX TRANSMISSIBILITY MULTIPLIERS CELLS
MULTY-
6*0.500
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The above example defines a 0.5 scaling multiplier for the six cells defined by the preceding BOX statement. The ENDBOX keyword resets the input box to the full grid.

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6.3.70 MULTZ - MULTIPLY CELL TRANSMISSIBILITY IN THE +Z DIRECTION

RUNSPEC GRID EDIT PROPS REGIONS	SOLUTION	SUMMARY	
---------------------------------	----------	---------	--

Revision: Rev-I

Description

MULTZ multiples the transmissibility between two cell faces in the $\pm Z$ direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, J, K) between the cells (I, J, K) and (I, J, K+1).

An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTZ keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name	Description	Default
I	MULTZ+	MULTZ+ is an array of real positive numbers assigning the transmissibility multipliers in the +Z direction to each cell face in the model.	1.0
		Repeat counts may be used, for example 20*100.0.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.61: MULTZ Keyword Description

See also the MULTZ-, MULTX-, MULTY and MULTY- keywords for scaling transmissible between grid cells.

Examples

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
         I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
                                                           / DEFINE BOX AREA
         10 10
                  1
                      18
                          1
                              1
         SET MULTX+ TRANSMISSIBILITY MULTIPLIERS
MULTZ
18*0.300
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The above example defines a 0.3 scaling multiplier for the 18 cells defined by the preceding BOX statement. The ENDBOX keyword resets the input box to the full grid.

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6.3.71 MULTZ- - MULTIPLY CELL TRANSMISSIBILITY IN THE -Z DIRECTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	
---------	------	------	-------	---------	----------	---------	--

Revision: Rev-I

Description

MULTZ- multiples the transmissibility between two cell faces in the -Z direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I, K-I) and (I, J, K).

An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTZ- keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name	Description	Default
I	MULTZ-	MULTZ- is an array of real positive numbers assigning the transmissibility multipliers in the -X direction to each cell face in the model.	1.0
		Repeat counts may be used, for example 20*100.0.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/". 2)

Table 6.62: MULTZ- Keyword Description

See also the MULTZ, MULTX, MULTX-, MULTY and MULTY- keywords for scaling transmissible between grid cells.

Examples

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
         I1 I2 J1 J2
                         K1 K2
B<sub>0</sub>X
                                                           / DEFINE BOX AREA
         10 10
                  1
                      6
                           1
                               1
         SET MULTX TRANSMISSIBILITY MULTIPLIERS CELLS
MULTZ-
6*0.500
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The above example defines a 0.5 scaling multiplier for the six cells defined by the preceding BOX statement. The ENDBOX keyword resets the input box to the full grid.

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6.3.72 NEWTRAN - ACTIVATE IRREGULAR CORNER-POINT GRID TRANSMISSIBILITIES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on Irregular Corner-Point Grid geometry transmissibility calculation, which is the default option for this type of grid. Grids defined with the COORD and ZCORN keywords will always invoke this option by default.

For Cartesian Regular Grids defined by the DX, DY, and DZ series of keywords the block center geometry transmissibility calculations should be activated via the OLDTRAN keyword. Again this is automatically invoked if this type of grid is being employed.

There is no data required for this keyword.

Example

- -- ACTIVATE IRREGULAR CORNER-POINT GRID TRANSMISSIBILITIES
- **NEWTRAN**

The above example manually activates Irregular Corner-Point Grid transmissibility calculations.

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6.3.73 NNC - Define Non-Neighbor Connections Between Cells Manually

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

NNC enables Non-Neighbor Connections ("NNC") to be manually defined. This keyword is normally generated by static modeling software as opposed to be manually entered in the OPM Flow input deck due to the verbosity and complexity of calculating the required parameters for this keyword.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
1	II	joined in a non-neighbor	A positive integer that defines the first grid block in the I-direction to be joined in a non-neighbor connection, must be greater than or equal to one and less than or equal to NX on the DIMENS in the RUNSPEC section.					
2	JI	joined in a non-neighbor	efines the first grid block connection, must be great NY on the DIMENS in th	er than or equal to one	None			
3	KI	joined in a non-neighbor	efines the first grid block i connection, must be great NZ on the DIMENS in th	er than or equal to one	None			
4	12	be joined in a non-neight	A positive integer that defines the second grid block in the I-direction to be joined in a non-neighbor connection, must be greater than or equal to one and less than or equal to NX on the DIMENS in the RUNSPEC section.					
5	J2	A positive integer that defines the second grid block in the J-direction to be joined in a non-neighbor connection, must be greater than or equal to one and less than or equal to NY on the DIMENS in the RUNSPEC section.						
6	K2	A positive integer that defines the second grid block in the K-direction to be joined in a non-neighbor connection, must be greater than or equal to one and less than or equal to NZ on the DIMENS in the RUNSPEC section.						
7	TRANSNNC		e real number greater tha cy between the first grid b K2).	•				
		The default value of zero zero.	sets the transmissibility b	etween the two cells to				
		cP.rb/day/psia	cP.rm³/day/bars	cP.rcc/hr/atm	0.0			
8	ISATNUMI	(relative permeability tab	ISATNUMI is a positive integer defining which saturation table number (relative permeability table) to be used for flow from the first grid block to the second grid block.					
		The default value of zero means the existing saturation table allocated to the upstream cell (II,JI,KI).						
9	ISATNUM2		nteger defining which satu le) to be used for flow fro ck.		0			
		The default value of zero the downstream cell (12,	means the existing satura [2,K2].	tion table allocated to				

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No.	Name		Description		Default				
		Field	Metric	Laboratory					
10	IPRSNUMI	IPRSNUMI is a positive in (PVT table) to be used fo grid block.			0				
		The default value of zero upstream cell (II,JI,KI).	The default value of zero means the existing PVT table allocated to the upstream cell (II,JI,KI).						
11	IPRSNUM2		IPRSNUM2 is a positive integer defining which pressure table number (PVT table) to be used for flow from the second grid block to the first grid block.						
			The default value of zero means the existing PVT table allocated to the downstream cell (12,12,K2).						
12	FACEI	the first grid block to the	FACEI is a character string that defines the face associated with flow from the first grid block to the second grid block, where FACEI can have values of: X+, X-, Y+, Y-, Z+, or Z						
13	FACE2	FACE2 is a character string the second grid block to of: X+, X-, Y+, Y-, Z+, or Z	None						
14	DIFFNNC	DIFFNNC is a positive re the first grid block (II, JI,							
		feet	meters	cm	0.0				
15	DISPNINC	DISPNNC s a positive res 1 (Areax Porosity) the second grid block (12,	between the first grid	block (II, JI, KI) and					
		ft ⁻²	m ⁻²	cm ⁻²	0.0				
16	AREANNC		AREANNC is a positive real number that defines the area associated with the connection between the first grid block (II, JI, KI) and the second grid block (I2, J2, K2).						
		ft²	m²	cm ²	None				
17	PERMNNC	AREANNC is a positive r associated with the conne the second grid block (12,	ection between the first g	rid block (II, JI, KI) and					
		mD	mD	mD	None				

Revision: Rev-I

Notes:

- Only functionality defined by items (1) to (7) are activated in OPM Flow.
- 2) Each record must be terminated by a "/" and the keyword is terminated by "/".

Table 6.63: NNC Keyword Description

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Note that although items (8) to (17) for this keyword are not available in OPM Flow, even if they were, it is strongly recommended that these items are defaulted if the data is being entered manually, as opposed to being generated by a pre-processing software.

Revision: Rev-I

Care should be taken that cells in different PVTNUM regions (see the PVTNUM keyword in the REGIONS section) are not connected, since the fluid properties are associated with a cell. If for example, a rbbl or a rm3 of oil flows from PVTNUM region I to PVTNUM region 2, then the oil properties of that oil will change from the PVT I data set to the PVT data set 2. This will result in material balance errors, that may or may not cause numerical issues.

If the transmissibility across a fault needs to be modified see the FAULTS and MULTFLT keywords in the GRID section for an alternative and less complicated method to modifying fault transmissibilities. Transmissibility between reservoir regions can be modified by using MULTREGT keyword, provided MULTNUM has been used to define the inter-region transmissibility region numbers for each grid block. Finally, the MULTX, MULTY and MULTZ series of keywords can be used to modify transmissibility between various cells. All the aforementioned keywords are described in the GRID section.

Example

```
MANUALLY DEFINE NON-NEIGHBOR CONNECTIONS
               ---- BOX -----
                                     -- TRANSNCC --
           J1 K1
                      I2 J2 K2
        I1
NCC
                                        0.2500
                                                  / SET NNC FOR FAULT
                 2
                      1
                           1
                                3
            1
                                        0.2500
                                                  / SET NNC FOR FAULT
        1
                                        0.2500
                                                  / SET NNC FOR FAULT
```

The above example defines the transmissibility between cells (1, 1, 1) and (1, 1, 2), (1, 1, 2) and (1, 1, 3) and finally between (1, 1, 3) and (1, 1, 4) to be 0.2500.

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6.3.74 NOGGF - DEACTIVATE OUTPUT OF GRID GEOMETRY FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword deactivates the output of a standard GRID or extended GRID file, as well as the extensible EGRID file for post-processing applications.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.75 NTG - Define the Net-to-Gross Ratio for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

NTG defines the Net-to-Gross Ratio ("NTG") for all the cells in the model via an array. The keyword can be used for all grid types.

No.	Name		Description						
		Field	Field Metric Laboratory						
I	NTG	than or equal to one, the each cell in the model. To NX x NY x NZ paramet	NTG is an array of real numbers greater than or equal to zero and less than or equal to one, that are assigned the net-to-gross ratio values for each cell in the model. The number of entries should correspond to the NX \times NY \times NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 200 * 0.850.						
		dimensionless	dimensionless dimensionless						

Notes:

- Setting a cells NTG value to zero will make the cell inactive, similar to setting the cells ACTNUM property to zero.
- 2) The keyword is terminated by "/".

Table 6.64: NTG Keyword Description

See also the PORO, PERMX, PERMY and PERMX keywords to fully define a grid's properties.

Example

```
-- DEFINE GRID BLOCK NTG DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- NTG
100*1.000 100*0.850 100*0.500 /
```

The above example defines a constant NTG of 1.00 for the first 100 cells, then 0.85 for the second 100 hundred cells, and finally 0.500 for the last 100 cell, for the 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.76 OLDTRAN - ACTIVATE CARTESIAN REGULAR GRID TRANSMISSIBILITIES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword switches on Cartesian Regular Grids geometry transmissibility calculation (or block centered transmissibility calculations), which is the default option for this type of grid. Grids defined by the DX, DY, and DZ series of keywords will always invoke this option by default.

For Irregular Corner-Point Grids defined by the COORD and ZCORN keywords Irregular Corner-Point Grid geometry transmissibility calculations should be activated via the NEWTRAN keyword. Again this is automatically invoked if this type of grid is being employed.

There is no data required for this keyword.

Example

-- ACTIVATE CARTESIAN REGULAR GRID TRANSMISSIBILITIES

OLDTRAN

The above example manually activates Cartesian Regular Grid transmissibility calculations.

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6.3.77 OPERATE - Define Mathematical Operations on Arrays

|--|

Revision: Rev-I

Description

This keyword defines the mathematical operations on arrays in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.78 OPERATER - Define Mathematical Operations on Arrays by Region

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the mathematical operations on arrays for specific regions in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.79 OPERNUM - Define Regions for Mathematical Operations on Arrays

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the OPERATE region numbers for each grid block. The OPERATE keyword defines mathematical operations on arrays in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords combined with the MULTNUM region array.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.80 OUTRAD - DEFINE THE OUTER RADIUS OF A RADIAL GRID

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

OUTRAD31 defines the OUTER radius of the reservoir model for a radial grid geometry. The RADIAL keyword in the RUNSPEC should be activated to indicate that radial geometry is being used.

No.	Name		Description						
		Field	Field Metric Laboratory						
I	OUTRAD	A single real positive n radius of a radial grid.	A single real positive number greater than INRAD defining the outer radius of a radial grid.						
		feet	m	cm	None				
Notes	<u>s:</u>	1	1	1	I				

I) The keyword is terminated by "/".

Table 6.65: OUTRAD Keyword Description

The keyword allows for an alternative method of entering the size of the R direction grid cells instead of entering the data using the DR or DRV keywords in the GRID section. Given the internal radius set by the INRAD keyword, the external radius set by the OUTRAD keyword and the number of grid cells in the R direction set by the NX variable on the DIMENS keyword in the RUNSPEC section, the R direction cells sizes are computed automatically on a geometric spacing, as defined by:

$$\frac{R_i}{R_{i-1}} = \left(\frac{OUTRAD}{R_{i,-1}}\right)^{\frac{1}{(NX-i_j+1)}}$$
(6.8)

Revision: Rev-I

or

$$R_{i} = \left(R_{i_{j}-1}\right) \left(\frac{OUTRAD}{R_{i_{j}-1}}\right)^{\frac{(i-i_{j}+1)}{(NX-i_{j}+1)}}$$
(6.9)

and the DR value for the ith cell, that is the value that can also be manually entered on the DR keyword in the GRID section, is given by:

$$DR_i = R_i - R_{i-1} {(6.10)}$$

Where:

 DR_i = DR value for the i^{th} cell

 R_i = current total radius for the i radii. R_{i-1} = total radius for the i-1 radii.

NX (NR) = number of radial grid cells excluding the inner radius

OUTRAD = the outer radius of the radial grid, the value includes the inner radius.

For example, given an inner radius set to 0.25, an outer radius of 2,050 and the number of cells in the R direction set to ten, then Table 6.66 shows the grid size calculations.

³¹ Radial grids are not currently implemented in OPM Flow version 2018-04 but is expected to be incorporated in the next release.



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OUTRAD Radial Grid Example									
INRAD	0.25								
OUTRAD	2050.0								
NX	10								
NX	Ri	DR	Ratio						
0	0.250	0.250							
I	0.616	0.366	1.463						
2	1.516	0.900	2.463						
3	3.733	2.217	2.463						
4	9.193	5.460	2.463						
5	22.638	13.445	2.463						
6	55.748	33.109	2.463						
7	137.281	81.533	2.463						
8	338.058	200.777	2.463						
9	832.477	494.420	2.463						
10	2050.000	1217.523	2.463						
Total		2050.000							
	// OLITOAD I	D - 4: -1 C -: 4 F							

Revision: Rev-I

Table 6.66: OUTRAD Radial Grid Example

See also the DR, DRV, DTHETA, DTHETAV and TOPS keywords to fully define a Radial Grid.

Example

```
-- INNER RADIUS OF FIRST GRID BLOCK IN THE RADIAL DIRECTION
-- INRAD
0.25
/ --
-- OUTER RADIUS OF FIRST GRID BLOCK IN THE RADIAL DIRECTION
--
OUTRAD
2050.0
```

The above example defines the inner radius to be 0.25 and the outer radius to be 2,050 feet. Note that the outer radius includes the inner radius.

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6.3.81 PERMR - Define the Permeability for Each Cell in the R Direction

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PERMR³² sets the permeability for each cell in the R direction in a radial geometry grid. The RADIAL keyword in the RUNSPEC should be activated to indicate that radial geometry is being used.

Although this keyword is read by OPM Flow, radial grids have not been fully implemented and therefore this type of grid should not be used.

Radial grids are not currently implemented in this version of OPM Flow but are expected to be incorporated in a future release.

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6.3.82 PERMTHT - Define the Permeability for Each Cell in the THETA Direction

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

PERMTHT³³ sets the permeability for each cell in the THETA direction in a radial geometry grid. The RADIAL keyword in the RUNSPEC should be activated to indicate that radial geometry is being used.

Although this keyword is read by OPM Flow, radial grids have not been fully implemented and therefore this type of grid should not be used.

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Radial grids are not currently implemented in this version of OPM Flow but are expected to be incorporated in a future release.

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6.3.83 PERMX - Define the Permeability in the X Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PERMX defines the permeability in the X direction for all the cells in the model via an array. The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name		Description						
		Field	Metric	Laboratory					
I	PERMX	the X direction to each o	ERMX is an array of real positive numbers assigning the permeability in the X direction to each cell in the model. Repeat counts may be used, for example 20*100.0.						
		mD	None						

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.67: PERMX Keyword Description

See also the PERMY and PERMZ keywords to fully define the permeability for the model.

Example

```
--
-- DEFINE GRID BLOCK PERMX DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
--
PERMX
100*500.0 100*50.0 100*200.0
```

The above example defines the PERMX to be 500.0, 50.0, and 200.0 for the first, second and third layers in the model for all 300 cells, as defined by the DIMENS keyword in the RUNSPEC section.

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Revision: Rev-I

6.3.84 PERMXX - Define the Permeability Tensor in the XX Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Description

This keyword defines the permeability tensor off-diagonal values for the XX direction.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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Revision: Rev-I

6.3.85 PERMXY - Define the Permeability Tensor in the XY Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Description

This keyword defines the permeability tensor off-diagonal values for the XY direction.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.86 PERMY - Define the Permeability in the Y Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PERMY defines the permeability in the Y direction for all the cells in the model via an array. The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name		Description						
		Field	Metric	Laboratory					
I	PERMY	the Y direction to each co	ERMY is an array of real positive numbers assigning the permeability in the Y direction to each cell in the model.						
		Repeat counts may be us							
		mD	mD	mD	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.68: PERMY Keyword Description

See also the PERMX and PERMZ keywords to fully define the permeability for the model.

Example

```
--
-- DEFINE GRID BLOCK PERMY DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
--
PERMY
100*500.0 100*50.0 100*200.0
```

The above example defines the PERMY to be 500.0, 50.0, and 200.0 for the first, second and third layers in the model for all 300 cells, as defined by the DIMENS keyword in the RUNSPEC section.

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/ Dipperson ---- A...

Revision: Rev-I

6.3.87 PERMYY - Define the Permeability Tensor in the YY Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Description

This keyword defines the permeability tensor off-diagonal values for the YY direction.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.88 PERMYZ – Define the Permeability Tensor in the YZ Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the permeability tensor off-diagonal values for the YZ direction.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.89 PERMZ - Define the Permeability in the Z Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PERMZ defines the permeability in the Z direction for all the cells in the model via an array. The keyword can be used for all grid types, except for the Radial Grid geometry.

No.	Name			Default				
		Field						
I	PERMZ		PERMZ is an array of real positive numbers assigning the permeability in the Z direction to each cell in the model.					
		Repeat counts may be us	Repeat counts may be used, for example 200*50.0.					
		mD	mD mD mD					

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.69: PERMZ Keyword Description

Note

Although PERMX and PERMZ are commonly set to be equal, PERMZ is typically not equal to either PERMX or PERMY. Normally PERMZ is set as a fraction of PERMX with typical values ranging from 0.1 to 0.5 times PERMX.

See also the PERMX and PERMY keywords to fully define the permeability for the model.

Example

```
--
-- DEFINE GRID BLOCK PERMZ DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
--
PERMZ
100*50.0 100*5.0 100*20.0
```

The above example defines the PERMZ to be 50.0, 5.0, and 20.0 for the first, second and third layers in the model for all 300 cells, as defined by the DIMENS keyword in the RUNSPEC section.

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Revision: Rev-I

6.3.90 PERMZX - Define the Permeability Tensor in the ZX Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Description

This keyword defines the permeability tensor off-diagonal values for the ZX direction.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.91 PERMZZ – Define the Permeability Tensor in the ZZ Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the permeability tensor off-diagonal values for the ZZ direction.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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6.3.92 PINCH - DEFINE PINCH-OUT LAYER OPTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PINCH keyword defines the parameters used to control the generation of Non-Neighbor Connections ("NNCs") in the vertical (K) direction due to layers pinching out. This keyword is applied to all layers in the model as opposed to the PINCHREG keyword that offers more flexibility by applying the pinch-out controls to various regions in the model defined by the PINCHNUM keyword.

OPM Flow will automatically generate connections between non neighbor cells in the vertical direction based on the parameters on this keyword.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
I	PINCHTHK	_	A real number defining the pinch-out threshold thickness for any cell. NNCs are generated across inactive cells having a vertical thickness less than PINCHTHK.						
		ft.	ft. m cm						
		0.001	0.001	0.001	Defined				
2	PINCHOPT	MINPV keyword has bee	A character string controlling the generation of pinch-outs when the MINPV keyword has been used to deactive cells with small pore volumes. PINCHOPT can either be set to:						
		I) GAP to allow t made inactive greater than PI							
		NOGAP to e threshold whee the MINPV key							
3	PINCHGAP		he maximum "empty" thic rid layers for a non-zero						
		ft.	m	cm					
		1.0E20	1.0E20	1.0E20	Defined				
4	PINCHCAL		ontrolling the calculation on the calculation of th		ТОРВОТ				
		from the half-o	 TOPBOT results in the pinch-out transmissibility being calculated from the half-cell Z-direction transmissibilities of the active cells on either side of the pinched-out layers. 						
		the Z-direction	the pinch-out transmissibi n transmissibilities harmon tive cells on either side of	ic average of all the cells					

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No.	Name	Description							
		Field	Metric	Laboratory					
5	PINCHMUL	transmissibilities when	A character string controlling the calculation of the pinch-out transmissibilities when adjustments have been made by the MULTZ keyword. PINCHMUL can either be set to:						
		,	TOP results in the pinch-out transmissibility being calculated from the active cell at the top of the pinch-out.						
		the minimum v		lity being calculated from e active cell at the top of in the pinch-out vertical					
		en PINCHMUL is reset CHMUL.							

Revision: Rev-I

1) The keyword is terminated by "/".

Table 6.70: PINCH Keyword Description

Examples

The first example below will create NNCs between the cells above and below any cell having vertical thickness less than 0.01 in either feet or metres.

```
-- SET PINCH-OUT PARAMETERS FOR CALCULATING PINCH-OUT PROPERTIES
-- PINCH
-- THRESHOLD GAP EMPTY TRANS
-- THICKNESS NO GAP GAP CALC
1* 1* 1* 1* /
```

For the second example, the MINPV keyword is used to set the minimum pore volume to 500 m^3 (metric units) and then the PINCH keyword is invoked with PINCHGAP set equal to GAP, as follows:

```
--
-- MINIMUM PORE VOLUME FOR ACTIVE CELLS
--
MINPV
1000.0 /
--
-- SET PINCH-OUT CRITERIA FOR THE MODEL
--
PINCH
-- THRESHOLD GAP EMPTY TRANS
-- THICKNESS NO GAP GAP CALC
0.1 GAP 1* 1* /
```

In the above example the MINPV keyword will deactivate all cells with pore volumes less than 500 m³. These deactivated cells are inactive in the model and therefore are not included in the flow calculations; however, by default they will result in no-flow barriers but may not be thin enough for PINCH to create NNCs across them. By setting PINCHGAP equal to GAP on the PINCH keyword (the default setting), then OPM Flow generates NNCs across the cells that have been deactivated by the MINPV keyword. However, in this case there may be grid blocks in the model with a pore volume greater than MINPV but a thickness less than the pinch-out threshold. These cells will not be deactivated by the PINCH keyword.

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6.3.93 PINCHNUM - Define Pinch-Out Regions for the PINCHREG Keyword

RUNSPEC	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PINCHNUM keyword defines the pinch-out region numbers for each grid block, as such there must be one entry for each cell in the model. The array is used with the PINCHREG keyword to set the pinch-out options and threshold thickness for each region.

No.	Name	Description	Default
1	PINCHNUM defines an array of positive integers assigning a grid cell to a particular PINCHNUM region.		I
		The maximum number of PINCHNUM regions is set by the NRPINC variable on the GRIDOPTS keyword in the RUNSPEC section.	
Notes	<u>:</u>		
I)	the RUNSPEC se	ntries should correspond to the NX \times NY \times NZ parameters on the DIMENS ection, unless the BOX keyword defines a sub area of the grid, in which case the should correspond to the number of cells defined by the BOX statement.	

The keyword is terminated by "/".

Table 6.71: PINCHNUM Keyword Description

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

Examples

The example below sets defines three PINCHNUM regions for various layers in a model based on the model's layering.

One can then set the pinch-out criteria for each region.

```
SET PINCH-OUT CRITERA VIA THE PINCHNUM REGION
- -
PINCHREG
         THRESHOLD
                     GAP
                              EMPTY
                                       TRANS
         THICKNESS
                     NO GAP
                              GAP
                                       CALC
         0.1
                     1*
                              1*
                                       1*
                                                           / PINCHNUM 01
                     1*
                                       1*
         1.0
                              10
                                                           / PINCHNUM 02
                     NOGAP
                              20
                                       1*
                                                           / PINCHNUM 03
         1.0
```

The above example sets the default pinch-out criteria for grid blocks defined as region one via the PINCHNUM array and different criteria for regions two and three.

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6.3.94 PINCHREG - DEFINE PINCH-OUT REGION OPTIONS

RUNSPEC	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PINCHREG keyword defines the parameters used to control the generation of Non-Neighbor Connections ("NNCs") in the vertical (K) direction due to layers pinching out in combination with the PINCHNUM keyword. This allows different regions in the model to use different criteria in controlling the how pinch-outs are generated. The keyword should contain NRPINC records defining the criteria for each pinch-out region defined with the PINCHNUM keyword. NRPINC is the maximum number of PINCHNUM regions defined via the GRIDOPTS keyword in the RUNSPEC section.

An alternative method to set the pinch-out criteria is to use the PINCH keyword, that applies the criteria to the whole model.

OPM Flow will automatically generate connections between non neighbor cells in the vertical direction based on the parameters on this keyword.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
I	PINCHTHK	A real number defining NNCs are generated act than PINCHTHK.							
		ft.							
		0.001	0.001	0.001	Defined				
2	PINCHOPT A character string controlling the generation of pinch-outs when the MINPV keyword has been used to deactivate cells with small pore volumes. PINCHOPT can either be set to:								
		made inactive	 GAP to allow the generation of NNCs across cells that have been made inactive with the MINPV keyword when the thickness is greater than PINCHTHK threshold. 						
		NOGAP to e threshold when the MINPV key							
3	PINCHGAP		he maximum "empty" thic rid layers for a non-zero						
		ft.	m	cm					
		1.0E20	1.0E20	1.0E20	Defined				
4	PINCHCAL		ontrolling the calculation		TOPBOT				
		 TOPBOT results in the pinch-out transmissibility being calculated from the half-cell Z-direction transmissibilities of the active cells on either side of the pinched-out layers. 							
		the Z-direction	the pinch-out transmissibi n transmissibilities harmon tive cells on either side of	ic average of all the cells					

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No.	Name	Description					
		Field	Field Metric Laboratory				
5	PINCHMUL	A character string controlling the calculation of the pinch-out transmissibilities when adjustments have been made by the MULTZ keyword. PINCHMUL can either be set to: 1) TOP results in the pinch-out transmissibility being calculated from the active cell at the top of the pinch-out.					
		2) ALL results in the minimum v	2) ALL results in the pinch-out transmissibility being calculated from the minimum value of the MULTZ of the active cell at the top of the pinch-out and all the inactive cells in the pinch-out vertical				
			s been set equal to ALL th the entered value for PINO				

Revision: Rev-I

Notes:

- The keyword should contain NRPINC records defining the criteria for each pinch-out region defined with the PINCHNUM keyword. NRPINC is the maximum number of PINCHNUM regions defined via the GRIDOPTS keyword in the RUNSPEC section.
- 2) Each record must be terminated by a "/" there is no keyword terminating "/".

Table 6.72: PINCHREG Keyword Description

Example

```
SET PINCH-OUT CRITERA VIA THE PINCHNUM REGION
PINCHREG
         THRESHOLD
                      GAP
                               EMPTY
                                        TRANS
                      NO GAP
         THICKNESS
                               GAP
                                        CALC
         0.1
                      1*
                                                             / PINCHNUM 01
                      1*
                                        1*
                               10
                                                              / PINCHNUM 02
         1.0
         1.0
                      NOGAP
                                20
                                        1*
                                                              / PINCHNUM 03
```

The above example sets the default pinch-out criteria for grid blocks defined as region one via the PINCHNUM array and different values for regions two and three.

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6.3.95 PORO - Define the Porosity Values for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PORO defines the porosity for all the cells in the model via an array. The keyword can be used for all grid types.

No.	Name		Description					
		Field	Field Metric Laboratory					
1	PORO	to zero and less than or cell in the model.	PORO is an array of real positive numbers that are greater than or equal to zero and less than or equal to one that are the porosity values for each cell in the model. Repeat counts may be used, for example 3000*0.15					
		dimensionless	dimensionless	dimensionless	None			

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.73: PORO Keyword Description

See also the NTG, PERMX, PERMY and PERMX keywords to fully define a grid's properties

Example

```
-- DEFINE GRID BLOCK POROSITY DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- PORO 300*0.300
```

The above example defines a constant porosity of 0.300 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.96 RADFIN - DEFINE A RADIAL LOCAL GRID REFINEMENT WITH ONE COLUMN

RUNSPEC	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the a radial local grid refinement using one columns Local grid refinement is currently not supported by OFM Flow.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

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6.3.97 RADFIN4 - Define a Radial Local Grid Refinement with Four Columns

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the a radial local grid refinement using four columns. Local grid refinement is currently not supported by OFM Flow.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Revision: Rev-I

6.3.98 REFINE - START THE DEFINITION OF A LOCAL GRID REFINEMENT

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

The REFINE keyword defines the start of a Cartesian or radial Local Grid Refinement ("LGR") definition that sets the properties of the selected LGR. The keyword is then followed by the property keywords associated with the section where the keyword is being invoked. For example, if the REFINE keyword is used in the GRID section then most of the keywords in that section can be used to set the grid properties for the LGR.

The ENDFIN keyword is used to terminate the LGR definition.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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6.3.99 RPTGRID - Define GRID Section Reporting

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the data in the GRID section that is to be printed to the output print file in human readable format. The keyword has two distinct forms, the first of which consists of the keyword followed by a series of integers on the next line indicating the data to be printed (see the first example). This is the original formal in the commercial simulator and was subsequently superseded by the second format. The second format consists of the keyword followed by a series of character strings that indicate the data to be printed. In most cases the character string is the keyword used to load the data in the OPM Flow input deck, for example PORO for the porosity array. Its is anticipated that OPM Flow will eventually support the functionality of the second format only, the first format although recognized will be completely ignored.

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

No.	Name	Description	Default
ı	ALLNCC	Print all the non-neighbor connections.	N/A
2	COORD	Print the coordinate lines.	N/A
3	COORDYS	Print the coordinate systems.	N/A
4	DEPTH	Print grid cells center depths.	N/A
			N/A
NI - 4			1

Notes:

I) The keyword is terminated by "/".

Table 6.74: RPTGRID Keyword Description

Note

This keyword has the potential to produce very large print files that some text editors may have difficulty loading, coupled with the fact that reviewing the data in this format is very cumbersome.

A more efficient solution is to load the *.INIT file into OPM ResInsight to view the data graphically, this also has the benefit of being able to filter the grid based on I, J, K ranges and grid properties.

Examples

The first example shows the original format of this keyword; although the keyword and format are recognized by OPM Flow, the format is ignored and is unlikely to be implemented in in the simulator.

```
-- DEFINE GRID SECTION REPORT OPTION (ORIGINAL FORMAT)
-- RPTGRID
1 2*0 1 3*1 /
```

The next example shows the second format of the keyword which may be supported in a future release of OPM Flow.

```
-- DEFINE GRID SECTION REPORT OPTIONS
--
RPTGRID
DX DY DZ DEPTH PORO PERMX /
```



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6.3.100 SPECGRID- Define the Dimensions of a Corner-Point Grid

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SPECGRID defines the dimensions of a corner-point grid in the x, y, and z planes as well as the number of reservoirs, where each reservoir has it's own set of corner-point geometry data.

The keyword can only be used with Irregular Corner-Point Grids.

No.	Name	Description	Default
I	NDIVIX	A positive integer value that defines the number of cells in the \boldsymbol{X} or \boldsymbol{R} direction	I
2	NDIVIY	A positive integer value that defines the number of cells in the Y or THETA direction	I
3	NDIVZ	A positive integer value that defines the number of cells in the Z direction	I
4	NUMRES	A positive integer values that defines number of coordinate data sets, or independent reservoirs in the model.	I
		OPM Flow currently only accepts a single data set, that is the default value of one.	
5	TYPE	A character string set to either T of F that defines the type of grid to be defined by subsequent keywords:	F
		T = Radial grid with radial coordinates	
		2) F = Cartesian grid	

Notes:

- 1) The keyword is terminated by "/".
- 2) The dimensions are also entered on the DIMENS section in the RUNSPEC section and the two sets of numbers should be consistent.

Table 6.75: SPECGRID Keyword Description

See also the COORD, COORDSYS and ZCORN keywords to fully define an Irregular Corner-Point Grid.

Example



The above example defines the a 46 x 112 x 22 grid with one set of irregular corner-point data.

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6.3.101 SWATINIT - Define the Initial Water Saturation Array for Capillary Pressure Scaling

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

SWATINIT defines the initial water saturation for all the cells in the model via an array. The keyword can be used for all grid types. SWATINIT is used to initialize the model by setting each grid block's initial water saturation ("Sw"). If the array is present in the input deck, then OPM Flow will re-scale the water-oil capillary pressure curves entered via the SWFN saturation functions in the PROPS section, so that the resulting initialized Sw matches the values in the SWATINIT array.

Normally the SWATINIT array is generated in the static earth model when calculating the hydrocarbons inplace. volumes using Saturation Height Functions ("SHF") derived from capillary pressure functions. Static earth models do not directly use capillary pressure in these type of calculations as individual cell pressures are not required. There is therefore some potential for inconsistencies to arise between the two sets of formulations. This is normally manifested by extreme scaling in the scaled capillary pressure values calculated by the simulator. If this is the case then the PPCMAX keyword can be used to set a maximum scaled capillary pressure value. Note that as large values of scaled capillary pressures can result in numerical issues, a more technically sound approach would be to resolve these inconsistencies before continuing with the model build.

No.	Name		Description						
		Field							
I	SWATINIT	equal to zero and less the saturation values to each	f real positive numbers than or equal to one, that cell in the model. ed, for example 3000*0.15	define the initial water					
		dimensionless	dimensionless dimensionless						

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.76: SWATINIT Keyword Description

See also the PPCMAX to control the re-scaling of the capillary pressure entries on the SWFN saturation function keyword in the PROPS section.

Example

```
--
-- DEFINE GRID BLOCK INITIAL SW DATA FOR ALL CELLS
-- (BASED ON NX x NY x NZ = 300)
--
SWATINIT
300*0.300
```

The above example defines a constant initial water saturation of 0.300 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.102 THCGAS - Define Gas Phase Thermal Conductivity for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The THCGAS keyword defines the gas phase thermal conductivity for when the thermal calculation is activated by the THERMAL keyword in the RUNSPEC section, and should be used in conjunction with THCROCK keyword in the GRID section.

This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name			Default				
		Field						
I	THCGAS		THCGAS is an array of real positive numbers that define the thermal conductivity of the gas phase in each grid block.					
		Repeat counts may be us						
		Btu/ft/day/°R	kJ/m/day/K	J/cm/hr/K	None			

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.77: THCGAS Keyword Description

The THCGAS data is used in conjunction with the other thermal conductivity arrays to calculate the porosity weighted thermal conductivity of a grid block using:

```
Average Thermal Conductivity = \frac{PORO \times (THCOIL + THCGAS + THCWATER + THCSOLID)}{NUMBER OF PHASES IN THE MODEL} \times (1 - PORO) \times THCROCK  (6.11)
```

See also the THCOIL, THCWATER, THROCK and THCSOLID keywords in the GRID section

Example

```
-- DEFINE GRID BLOCK GAS PHASE THERMAL CONDUCTIVITY
- FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- THCGAS
300*20.0
```

The above example defines the gas phase thermal conductivity of 20.0 for each cell in the 300 grid block model as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.103 THCOIL - Define Oil Phase Thermal Conductivity for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The THCOIL keyword defines the oil phase thermal conductivity for when the thermal calculation is activated by the THERMAL keyword in the RUNSPEC section, and should be used in conjunction with THCROCK keyword in the GRID section.

This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name			Default				
		Field						
I	THCOIL		THCOIL is an array of real positive numbers that define the thermal conductivity of the oil phase in each grid block.					
		Repeat counts may be us						
		Btu/ft/day/°R	kJ/m/day/K	J/cm/hr/K	None			

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.78: THCOIL Keyword Description

The THCOIL data is used in conjunction with the other thermal conductivity arrays to calculate the porosity weighted thermal conductivity of a grid block using:

```
Average Thermal Conductivity=\frac{PORO\times [THCOIL+THCGAS+THCWATER+THCSOLID]}{NUMBER OF PHASES IN THE MODEL}\times (1-PORO)\times THCROCK (6.12)
```

See also the THCGAS, THCWATER, THROCK and THCSOLID keywords in the GRID section

Example

```
-- DEFINE GRID BLOCK OIL PHASE THERMAL CONDUCTIVITY
- FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- THCOIL
300*20.0
```

The above example defines the oil phase thermal conductivity of 20.0 for each cell in the 300 grid block model, as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.104 THCONR - Define Rock and Fluid Thermal Conductivity for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The THCONR keyword defines the reservoir rock plus fluid thermal conductivity for all cells for when the thermal calculation is activated by the THERMAL keywords in the RUNSPEC section.

Note that the THCROCK and THCONR keywords are mutually exclusive.

This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description					
		Field	Field Metric Laboratory					
I	THCONR		THCONR is an array of real positive numbers that define the combined rock and fluid conductivity of a grid block.					
		Repeat counts may be us	Repeat counts may be used, for example 3000*25.0					
		Btu/ft/day/°R	Btu/ft/day/°R kJ/m/day/K J/cm/hr/K					

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.79: THCONR Keyword Description

Example

```
-- DEFINE GRID BLOCK ROCK-FLUID THERMAL CONDUCTIVITY
-- FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
```

THCONR

300*25.0

The above example defines the combined rock and fluid thermal conductivity of 25.0 for each cell in the 300 grid block model, as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.105 THCONSF - Define Gas Saturation Dependent Thermal Conductivity Scaling Factor for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The THCONSF keyword defines a gas saturation dependent scaling factor to the fluid and reservoir rock thermal conductivities entered via the THCONR keyword in the GRID section, for when the thermal calculation is activated by the THERMAL keyword in the RUNSPEC.

Note that the THCROCK and THCONR keywords are mutually exclusive.

This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description					
		Field	Field Metric Laboratory					
I	THCONSF	than or equal to one, t factor that is applied to	THCONSF is an array of real positive numbers, greater than zero and less han or equal to one, that define the gas saturation dependent scaling actor that is applied to the THCONR data, entered via the THCONR teyword, to adjust the thermal conductivity of the reservoir cells in each grid block.					
		Repeat counts may be us						
		dimensionless	dimensionless	dimensionless	None			

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.80: THCROCK Keyword Description

The THCONSF keyword defines a scaling factor which is a function of the gas saturation that scales a cells total thermal conductivity (reservoir fluids plus reservoir rock) entered via the THCONR keyword in the GRID section. This combination of keywords, THCONSF and THCONR implies that the oil and water phase thermal conductivities are saturation independent with respect to the liquid phase, and that only the gas saturation influences a cell's thermal conductivity as entered via the THCONR keyword. Thus, THCONSF scales the THCORNR values via a multiplier Ω , by:

$$\Omega_{i,j,k} = (1 - \text{THCONSF x Gas Saturation})_{i,j,k}$$
 (6.13)

See also the THCGAS, THCOIL, THCWATER and THCSOLID keywords in the GRID section, for an alternative way to enter the thermal conductivity properties. However, the THCONSF keyword cannot be used with the THCGAS, THCOIL, THCWATER and THCSOLID keywords. Secondly, the solid phase is not supported by OPM Flow and therefore neither is the THCSOLID keyword.

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Revision: Rev-I

Example

-- DEFINE GRID SGAS DEPENDENT SCALING FACTOR FOR THE THCONR ARRAY
- FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- THCONSF
300*0.12

The above example defines the gas saturation thermal conductivity scaling factor to be applied to the THCONR to be 0.12 for all 300 cells in the model, as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.106 THCROCK - Define Reservoir Rock Thermal Conductivity for All **C**ELLS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The THCROCK keyword defines the reservoir rock thermal conductivity for when the thermal calculation is activated by the THERMAL keyword in the RUNSPEC section.

Note that the THCROCK and THCONR keywords are mutually exclusive.

This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description					
		Field	Field Metric Laboratory					
I	THCROCK		THCROCK is an array of real positive numbers that define the thermal conductivity of the reservoir rock in each grid block.					
		Repeat counts may be us	Repeat counts may be used, for example 3000*20.0					
		Btu/ft/day/°R	Btu/ft/day/°R kJ/m/day/K J/cm/hr/K					

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/".

Table 6.81: THCROCK Keyword Description

The THCROCK data is used in conjunction with the other thermal conductivity arrays to calculate the porosity weighted thermal conductivity of a grid block using:

```
Average Thermal Conductivity = \frac{PORO \times [THCOIL + THCGAS + THCWATER + THCSOLID]}{NIJMBER OF PHASES IN THE MODEL} \times (1 - PORO) \times THCROCK
                                                                                                                                                  (6.14)
                                                     NUMBER OF PHASES IN THE MODEL
```

See also the THCGAS, THCOIL, THCWATER and THCSOLID keywords in the GRID section

Example

```
DEFINE GRID BLOCK RESERVOIR ROCK THERMAL CONDUCTIVITY
         FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
THCROCK
```

300*20.0

The above example defines the reservoir rock thermal conductivity of 20.0 for each cell in the 300 grid block model, as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.107 THCSOLID - Define Solid Phase Thermal Conductivity for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The THCSOLID keyword defines the solid phase thermal conductivity for when the thermal calculation is activated by the THERMAL keyword in the RUNSPEC section in the commercial simulator. THCSOLID should be used in conjunction with THCROCK keyword in the GRID section.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate.

This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section in the commercial simulator.

No.	Name		Default					
		Field	Field Metric Laboratory					
1	THCSOLID		THCSOLID is an array of real positive numbers that define the thermal conductivity of the solid phase in each grid block.					
		Repeat counts may be use	Repeat counts may be used, for example 3000*20.0					
		Btu/ft/day/°R	Btu/ft/day/°R kJ/m/day/K J/cm/hr/K					

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 6.82: THCSOLID Keyword Description

The THSOLID data is used in conjunction with the other thermal conductivity arrays to calculate the porosity weighted thermal conductivity of a grid block using:

```
 \label{eq:average} \textbf{Average Thermal Conductivity} = \frac{PORO \times (THCOIL + THCGAS + THCWATER + THCSOLID)}{\text{NUMBER OF PHASES IN THE MODEL}} \times (1 - PORO) \times THCROCK
```

See also the THCGAS, THCOIL, THCWATER and THCROCK keywords in the GRID section.

Example

```
DEFINE GRID BLOCK RESERVOIR SOLID PHASE THERMAL CONDUCTIVITY
         FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
THSOLID
         300*20.0
```

The above example defines the solid phase thermal conductivity of 20.0 for each cell in the 300 grid block model, as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.108 THCWATER - Define Water Phase Thermal Conductivity for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The THCWATER keyword defines the water phase thermal conductivity for when the thermal calculation is activated by the THERMAL keyword in the RUNSPEC section, and should be used in conjunction with THCROCK keyword in the GRID section.

This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description					
		Field						
I	THCWATER	,	THCWATER is an array of real positive numbers that define the thermal conductivity of the water phase in each grid block.					
		Repeat counts may be us	Repeat counts may be used, for example 3000*20.0					
		Btu/ft/day/°R	kJ/m/day/K	J/cm/hr/K	None			

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/".

Table 6.83: THCWATER Keyword Description

The THCWATER data is used in conjunction with the other thermal conductivity arrays to calculate the porosity weighted thermal conductivity of a grid block using:

```
 \label{eq:average} \textbf{Average Thermal Conductivity} = \frac{PORO \times (THCOIL + THCGAS + THCWATER + THCSOLID)}{\text{NUMBER OF PHASES IN THE MODEL}} \times (1 - PORO) \times THCROCK
```

See also the THCGAS, THCOIL, THROCK and THCSOLID keywords in the GRID section

Example

```
DEFINE GRID BLOCK WATER PHASE THERMAL CONDUCTIVITY
        FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
THCWATER
 300*20.0
```

The above example defines the water phase thermal conductivity of 20.0 for each cell in the 300 grid block model, as defined by the DIMENS keyword in the RUNSPEC section.

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6.3.109 TOPS - Define the Depth at the Center of the Top Face for Each Cell

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

TOPS defines the depth of the top face of each cell in the model.

It can only be used with the Cartesian Regular Grid or Radial Grid models.

No.	Name		Description		Defaul		
		Field	Metric	Laboratory			
1	TOPS	each cell in the model. Clayer only based on NX remaining TOPS based on TOPS may be entered for	I numbers defining the decome can either just enter x NY entries and OPM neither DZ or DZV. Alterner each cell in the model.	the TOPS for the first Flow will calculate the natively NX x NY x NZ			
		Repeat counts may be used, for example 10*5201.0.					
		feet m cm					

Table 6.84:TOPS Keyword Description

See also the DEPTHS keyword to define the structural depth for the cells.

Examples

The example below defines the TOPS of the cells for each cell for NX = 5, NY = 5 and NZ = 3 model, as well as the X and Y direction cells sizes.

```
DEFINE GRID BLOCK TOPS FOR ALL LAYERS (BASED ON NX = 5, NY = 5, NZ = 3)
TOPS
 25*3100 25*3105 25*3110
    DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX = 5)
\mathsf{DXV}
  5*100
    DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NY = 5)
DYV
  5*100
```

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The next example defines the same grid as before but with the TOPS keyword only defining the top layer and DZV keyword defining the cells thickness.

Revision: Rev-I

```
-- DEFINE GRID BLOCK TOPS FOR THE TOP LAYER (BASED ON NX = 5, NY = 5, NZ = 3)
-- TOPS
25*3100 //
-- DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX = 5)
-- DZV
3*5.0 //
-- DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX = 5)
-- DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX = 5)
-- DZV
5*100 //
-- DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NY = 5)
-- DYV
5*100 //
```

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6.3.110 ZCORN - Define the Depth of Each Corner-Point of a Grid Block

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

ZCORN defines the depth of each corner point of a grid block on the pillars defining the reservoir grid.

A total of $8 \times NX \times NY \times NZ$ values are needed to fully define all the depths in the model. The depths specifying the top of the first layer are entered first with one point for each pillar for each grid block. The points are entered with the X axis cycling fastest. Next come the depths of the bottom of the first layer. The top of layer two follows etc.

The keyword can be only used be uses with Irregular Corner-Point Grids.

No.	Name		Default		
		Field Metric Laboratory			
I	ZCORN	An array of depths with 8 x NZ entries	8 depths for each cell, for	a total of 8 x Nx x NY	
		feet	metres	cm	None

Notes:

- I) Regular Cartesian and Radial grid keywords cannot be used with this keyword, that is: DR, DRV, DTHETA, DTHETAV, DX, DXV, DY, DYV, DZ, INRAD, and TOPS.
- 2) The keyword is terminated by "/".

Table 6.85: ZCORN Keyword Description

See also the SPECGRID, COORD and COORDSYS keywords to fully define an Irregular Corner-Point Grid.

Example

```
-- SPECIFY CORNER-POINT DEPTHS FOR A 3 x 2 x 2 GRID,
  WITH CONSTANT SLOPE IN THE X AND Y DIRECTIONS
-- SUCH THAT ALL CORNER POINTS OF NEIGHBOURING BLOCKS ALIGN
ZCORN
  1450
         1500
                 1500
                         1550
                                 1550
                                        1600 # top of layer 1
  1500
         1550
                 1550
                         1600
                                 1600
                                        1650
  1500
         1550
                 1550
                         1600
                                 1600
                                        1650
  1550
         1600
                 1600
                         1650
                                 1650
                                        1700
  1460
                                        1610 # bottom of layer 1
         1510
                 1510
                                 1560
                         1560
  1510
         1560
                 1560
                         1610
                                 1610
                                        1660
  1510
         1560
                 1560
                         1610
                                 1610
                                        1660
  1560
         1610
                 1610
                         1660
                                 1660
                                        1710
  1460
                                        1610 # top of layer 2
         1510
                 1510
                         1560
                                 1560
  1510
                 1560
         1560
                         1610
                                 1610
                                        1660
  1510
         1560
                 1560
                         1610
                                 1610
                                        1660
  1560
         1610
                 1610
                         1660
                                 1660
                                        1710
                                        1620 # bottom of layer 2
  1470
         1520
                 1520
                                 1570
                         1570
  1520
         1570
                 1570
                         1620
                                 1620
                                        1670
  1520
         1570
                 1570
                         1620
                                 1620
                                        1670
  1570
                 1620
                                 1670
                                        1720
         1620
                         1670
```

The above example defines depths of the vertical coordinate lines for a regular 3 by 2 by 2 grid with a constant slope in the x and y directions such that all the corner points of neighboring blocks are aligned.

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7 EDIT SECTION

7.1 Introduction

This section enables user defined changes to be applied after OPM Flow has processed the data in the GRID section, that is the resulting pore volume (PORV) and transmissibility arrays (TRANX, TRANY and TRANX). The entered primary static arrays (PORO, PERMX etc.) in the GRID section are no longer available and all modifications are applied to the pore volume and transmissibility arrays.

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Historically the intention of this section was allow for the editing of the processed data; however, the features available in this section have, through time, migrated to the GRID section. For example the array operator keywords like ADD, COPY, MULTIPLY, etc. are available in the GRID section and thus enabling editing of the primary static arrays (PORO, PERMX etc.). This increased in capability in the GRID section has therefore made the EDIT section some what redundant.

Nevertheless the section is sometimes used by users to incorporate history matching parameter changes and by static earth modeling software packages to import directly the pore volumes and transmissibilities calculated in the static model directly into the numerical model via the EDIT section. Although the latter work flow is not very common.

7.2 DATA REQUIREMENTS

As the primary purpose of this section is to modify the simulator's calculated pore volumes and transmissibilities, then the properties used to define these arrays must have been fully defined in the GRID section. The arrays available for modification in the EDIT section are listed in Table 7.1 together with the associated GRID arrays used to generate the EDIT property array.

	regular Corner-Point Grids Keywords	Radial	Grid Keywords
GRID	EDIT	GRID	EDIT
TOPS	DEPTH	TOPS	DEPTH
DX		DR	
DY		THETA	
DZ	PORV	DZ	PORV
DZNET	PORV	DZNET	FORV
PORO		PORO	
NTG		NTG	
PERMX	TRANX	PERMR	TRANR
MULTX		MULTR	IKAINK
PERMY	TRANY	PERMTHT	TRANTHT
MULTY	IRANI	MULTTHT	INAMINI
PERMZ	TDANIZ	PERMZ	TRANZ
MULTZ	TRANZ	MULTZ	INAINZ

Notes:

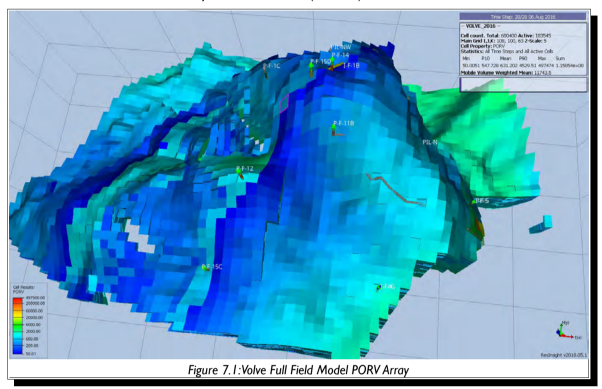
- 1) Currently Radial Grids have not been implemented in OPM Flow.
- 2) The GRID property association to the EDIT property is only indicitive as several variables, DZNET and NTG for example, are also used in the transmissibility calculations.

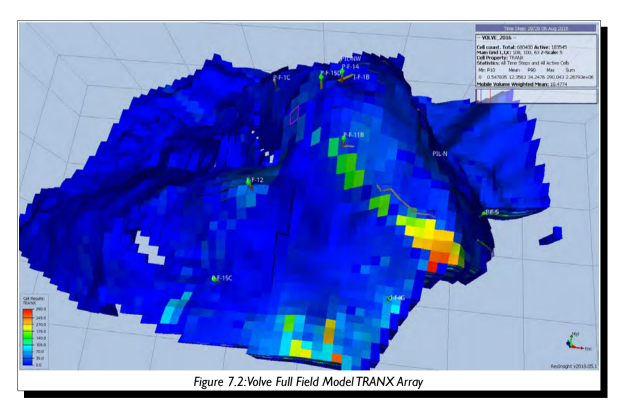
Table 7.1: EDIT Section Arrays Available for Modification

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An example pore volume array (PORV property) from the Volve³⁴ field is shown in Figure 7.1 and Figure 7.2 illustrates the model's transmissibility in the x-direction (TRANX).





³⁴ The Volve Data was approved for data sharing in 2018 by the initiative of the last Operating company, Equinor and approved by the license partners ExxonMobil E&P Norway AS and Bayerngas Norge AS in the end of 2017.

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7.3 Keyword Definitions

7.3.1 ADD - ADD A CONSTANT TO A SPECIFIED ARRAY

The ADD keyword adds a constant to a specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the ADD keyword is being applied.

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See ADD – Add a Constant to a Specified Array in the GRID section for a full description.

7.3.2 ADDREG - ADD A CONSTANT TO AN ARRAY BASED ON A REGION NUMBER

The ADDREG keyword adds a constant to a specified array or part of an array based on cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the ADDREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the ADDREG keyword is being used.

See ADDREG - Add a Constant to an Array based on a Region Number in the GRID section for a full description.

7.3.3 BOX - Define a Range of Grid Blocks to Enter Property Data

BOX defines a range of grid blocks for which subsequent data is assigned for all the cells in the defined BOX. Note that the BOX grid is reset by the keyword ENDBOX by resetting the current defined BOX to be the whole grid. The keyword can be used for any array and for all grid types.

See BOX - Define a Range of Grid Blocks to Enter Property Data in the GRID section for a full description.

7.3.4 COPY - Copy Array Data to Another Array

The COPY keyword copies an array (or part of an array) to another array or part of an array. The arrays can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the COPY keyword is being used.

See COPY - Copy Array Data to Another Array in the GRID section for a full description.

7.3.5 COPYREG - COPY AN ARRAY TO ANOTHER ARRAY BASED ON A REGION NUMBER

The COPYREG keyword copies a specified array or part of an array based on cells with a specific region number to another array. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the COPYREG keyword is read by the simulator. The property arrays can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the COPYREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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7.3.6 DEPTH - EDITS THE DEPTH AT THE CENTER OF EACH CELL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The DEPTH keywords modifies the depth at the center of selected cells in the model. The cells DEPTH are calculated by OPM Flow at the end of the GRID section and this keyword allows the user to adjust the calculated depths in the EDIT section. The area to be modified can be defined via the various grid selection keywords, ADD, BOX, EQUALS, etc., and areas that are not selected remain unchanged.

No.	Name		Default						
		Field Metric Laboratory							
1	DEPTH		DEPTH is an array of real numbers defining the depth at the center of each cell in the model. Only the values in						
		Repeat counts may be us	Repeat counts may be used, for example 30*5201.0.						
		feet	feet m cm						

Notes:

1) The keyword is terminated by "/".

Table 7.2:TOPS Keyword Description

See also the TOPS keyword to define the top structural depth for the cells.

Examples

The example below defines the TOPS of the cells for each cell for NX = 5, NY = 5 and NZ = 3 model, as well as the X and Y direction cells sizes.

```
----- BOX -----
    I1 I2
             J1 J2 K1 K2
B<sub>0</sub>X
                     20 20 / SET BOX AREA TO BE MODIFIED
             11 11
   DEFINE GRID BLOCK X DIRECTION CELL SIZE (BASED ON NX = 5)
DEPTH
 10*3500.0
     ----- BOX -----
- -
    I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
    1* 1*
                1*
                      1* 1* / RESET BOX DEFAULTS
```

Alternatively the EQUALS keyword can be used to perform the same edit.

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7.3.7 EDIT - Define the Start of the EDIT Section of Keywords

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

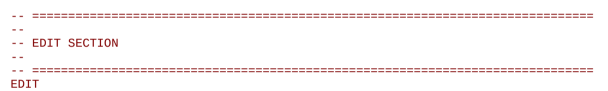
Revision: Rev-I

Description

The EDIT activation keyword marks the end of the GRID section and the start of the EDIT section that enables modifications to the OPM Flow calculated properties derived from the data entered in the GRID section, for example grid block pore volumes via the PORV array and the transmissibilities via the TRANX, TRANY and TRANZ family of keywords.

There is no data required for this keyword.

Example



The above example marks the end of the GRID section and the start of the EDIT section in the OPM Flow data input file.

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7.3.8 EDITNIC - Scale Non-Neighbor Connections Between Cells Manually

RUNSPEC GRID EDIT PROPS F	REGIONS SOLUTION SUMMARY SCHEDULI	
---------------------------	-----------------------------------	--

Revision: Rev-I

Description

EDITNNC enables Non-Neighbor Connections ("NNC"), entered via the NNC keyword or calculated by the simulator, to be multiplied (re-scaled) by a constant. For example, if the existing transmissibility between non-neighbor connections is $T_{\rm old}$ and the multiplier is C, then the resulting transmissibility, $T_{\rm new}$, will be $T_{new} = C \ x \ T_{old}$. Only previously defined NNC's entered via the NNC keyword or calculated by the simulator can be edited, otherwise an error will occur.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	II	joined in a non-neighbor	efines the first grid block connection, must be great NX on the DIMENS in th	er than or equal to one	None
2	JI	joined in a non-neighbor	efines the first grid block connection, must be great NY on the DIMENS in the	er than or equal to one	None
3	KI	joined in a non-neighbor	efines the first grid block i connection, must be great NZ on the DIMENS in th	er than or equal to one	None
4	12	A positive integer that defines the second grid block in the I-direction to be joined in a non-neighbor connection, must be greater than or equal to one and less than or equal to NX on the DIMENS in the RUNSPEC section.			None
5	J2	A positive integer that defines the second grid block in the J-direction to be joined in a non-neighbor connection, must be greater than or equal to one and less than or equal to NY on the DIMENS in the RUNSPEC section.			None
6	K2	A positive integer that de be joined in a non-neight one and less than or e section.	None		
7	TRANSMUL	defines a constant that	e real number greater that scales the transmissibility second grid block (12, J2, K	between the first grid	
		The default vale of one m	neans no scaling will be app	olied.	
		dimensionless	dimensionless	dimensionless	ı
8	ISATNUMI		nteger defining which satule) to be used for flow fro		0
		The default value of zero the upstream cell (11,J1,K	means the existing satura l).	tion table allocated to	

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No.	Name		Description		Default			
9	ISATNUM2	ISATNUM2 is a positive ir (relative permeability table block to the first grid block	e) to be used for flow fro		0			
			The default value of zero means the existing saturation table allocated to the downstream cell (I2,J2,K2).					
10	IPRSNUMI		IPRSNUMI is a positive integer defining which pressure table number (PVT table) to be used for flow from the first grid block to the second grid block.					
		The default value of zero means the existing PVT table allocated to the upstream cell (II,JI,KI).						
11	IPRSNUM2		IPRSNUM2 is a positive integer defining which pressure table number (PVT table) to be used for flow from the second grid block to the first grid block.					
			The default value of zero means the existing PVT table allocated to the downstream cell (12,12,K2).					
12	FACEI	FACEI is a character strin the first grid block to the of: X+, X-, Y+, Y-, Z+, or Z	second grid block, where		None			
13	FACE2	FACE2 is a character strin the second grid block to t of: X+, X-, Y+, Y-, Z+, or Z	he first grid block, where		None			
14	DIFFNNC	scales the diffusivity between	DIFFNNC is a positive real number greater than or equal to zero that scales the diffusivity between the first grid block (II, JI, KI) and the second grid block (I2, J2, K2).					
		dimensionless	dimensionless	dimensionless	0.0			

Revision: Rev-I

Table 7.3: EDITNNC Keyword Description

Note that although items (8) to (14) for this keyword are not available in OPM Flow, even if they were, it is strongly recommended that these items are defaulted if the data is being entered manually, as opposed to being generated by pre-processing software.

Care should be taken that cells in different PVTNUM regions (see the PVTNUM keyword in the REGIONS section) are not connected, since the fluid properties are associated with a cell. If for example, a rbbl or a rm³ of oil flows from PVTNUM region I to PVTNUM region 2, then the oil properties of that oil will change from the PVT I data set to the PVT data set 2. This will result in material balance errors, that may or may not cause numerical issues.

If the transmissibility across a fault needs to be modified see the FAULTS and MULTFLT keywords in the GRID section for an alternative and less complicated method to modifying fault transmissibilities. Transmissibility between reservoir regions can be modified by using MULTREGT keyword, provided MULTNUM has been used to define the inter-region transmissibility region numbers for each grid block. Finally, the MULTX, MULTY and MULTZ series of keywords can be used to modify transmissibility between various cells. All the aforementioned keywords are described in the GRID section.

See also the EDITNNCR keyword in the EDIT section that resets an existing NNC to a user defined value.

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Revision: Rev-I

Example

```
MANUALLY RESCALE NON-NEIGHBOR CONNECTIONS
         ----- BOX -----
                                        -- TRANSMUL --
        I1 J1 K1 I2 J2 K2
EDITNCC
                                                   / SET NNC FOR FAULT
/ SET NNC FOR FAULT
/ SET NNC FOR
                        1
                             1
                                  2
                                           0.2000
         1
             1
                  1
         1
             1
                   2
                        1
                             1
                                  3
                                            0.2000
                                                      / SET NNC FOR FAULT
             1
                   3
                                  4
                                            0.2000
```

The above example multipliers the transmissibility between cells (1, 1, 1) and (1, 1, 2), (1, 1, 2) and (1, 1, 3) and finally between (1, 1, 3) and (1, 1, 4) by 0.200.

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7.3.9 EDITNNCR - RESET NON-NEIGHBOR CONNECTIONS BETWEEN CELLS MANUALLY

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

EDITNNCR enables Non-Neighbor Connections ("NNC"), entered via the NNC keyword or calculated by the simulator, to be reset to a user defined value. Only previously defined NNC's entered via the NNC keyword or calculated by the simulator can be edited, otherwise an error will occur. See also the EDITNNC keyword in the EDIT section that scales an existing NNC.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	II	joined in a non-neighbor	efines the first grid block connection, must be great NX on the DIMENS in th	er than or equal to one	None
2	ון	A positive integer that d joined in a non-neighbor and less than or equal to	None		
3	KI	A positive integer that de joined in a non-neighbor and less than or equal to	None		
4	12	A positive integer that d be joined in a non-neight one and less than or e section.	None		
5	J2	A positive integer that d be joined in a non-neight one and less than or e section.	None		
6	K2	A positive integer that do be joined in a non-neight one and less than or e section.	None		
7	TRANSNNC	TRANSNNC is a positive defines the transmissibility second grid block (12, J2, This value cannot be default.)			
		cP.rb/day/psia	cP.rm³/day/bars	cP.rcc/hr/atm	None
8	ISATNUMI	ISATNUMI is a positive i (relative permeability tab to the second grid block.	0		
		The default value of zero the upstream cell (II,JI,K	means the existing satura l).	tion table allocated to	

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No.	Name		Description		Default				
9	ISATNUM2		nteger defining which satu e) to be used for flow fro ck.		0				
		The default value of zero the downstream cell (I2,	means the existing satura (2,K2).	tion table allocated to					
10	IPRSNUMI		nteger defining which pres r flow from the first grid b		0				
		The default value of zero upstream cell (II,JI,KI).	he default value of zero means the existing PVT table allocated to the pstream cell (II,JI,KI).						
-11	IPRSNUM2	IPRSNUM2 is a positive in (PVT table) to be used for grid block.	0						
			The default value of zero means the existing PVT table allocated to the downstream cell (I2,J2,K2).						
12	FACEI		ng that defines the face ass second grid block, where Z		None				
13	FACE2		ng that defines the face ass the first grid block, where Z		None				
14	DIFFNNC		ral number greater than or een the first grid block (H K2).						
		The default value is the value	The default value is the value calculated in the GRID section.						
		feet	*						
Notes	<u>s:</u>								
I)	Each record mus	st be terminated by a "/" an	d the keyword is terminat	ed by "/".					

Revision: Rev-I

Table 7.4: EDITNNCR Keyword Description

Note that although items (8) to (14) for this keyword are not available in OPM Flow, even if they were, it is strongly recommended that these items are defaulted if the data is being entered manually, as opposed to being generated by pre-processing software.

If the transmissibility across a fault needs to be modified see the FAULTS and MULTFLT keywords in the GRID section for an alternative and less complicated method to modifying fault transmissibilities. Transmissibility between reservoir regions can be modified by using MULTREGT keyword, provided MULTNUM has been used to define the inter-region transmissibility region numbers for each grid block. Finally, the MULTX, MULTY and MULTZ series of keywords can be used to modify transmissibility between various cells. All the aforementioned keywords are described in the GRID section.

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Example

```
MANUALLY RESET NON-NEIGHBOR CONNECTIONS
         ----- BOX -----
                                        -- TRANSNNC --
         I1 J1 K1 I2 J2 K2
EDITNCCR
                                                   / SET NNC FOR FAULT
/ SET NNC FOR FAULT
/ SET NNC FOR
                         1
                              1
                                   2
                                            0.2500
              1
                   1
         1
         1
              1
                   2
                         1
                              1
                                   3
                                            0.2500
                                                       / SET NNC FOR FAULT
              1
                   3
                                   4
                                            0.2500
```

The above example res-sets the transmissibility between cells (1, 1, 1) and (1, 1, 2), (1, 1, 2) and (1, 1, 3) and (1, 1, 3) and (1, 1, 4) to be 0.2500.

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7.3.10 ENDBOX - Define the End of the BOX Defined Grid

This keyword marks the end of a previously defined BOX sub-grid as defined by a previously entered BOX keyword. The keyword resets the input grid to be the full grid as defined by the NX, NY, and NZ variables on the DIMENS keyword in the RUNSPEC section.

Revision: Rev-I

See ENDBOX - Define the End of the BOX Defined Grid in the GRID section for a full description.

7.3.11 ENDFIN - END THE DEFINITION OF A LOCAL GRID REFINEMENT

NDFIN defines the end of a Cartesian or radial local grid refinement ("LGR") definition and a LGR property definition data set.

See ENDFIN - End the Definition of a Local Grid Refinement in the GRID section for a full description.

7.3.12 EQUALREG - Sets an Array to a Constant by Region Number

The EQUALREG keyword sets a specified array to a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the EQUALREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the EQUALREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

7.3.13 EQUALS - Sets a Specified Array to a Constant

The EQUALS keyword sets a specified array or part of an array to a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the EQUALS keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

7.3.14 MAXVALUE - Sets a Maximum Value for an Array Element

The MAXVALUE keyword sets a maximum value for the specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the MAXVALUE keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

7.3.15 MINVALUE - SET A MINIMUM VALUE FOR AN ARRAY ELEMENT

The MINVALUE keyword sets a minimum value for the specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the MINVALUE keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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7.3.16 MULTFLT - MULTIPLY THE TRANSMISSIBILITY OF A DEFINED FAULT BY A CONSTANT

MULTFLT enables the transmissibilities across defined faults, as declared by the FAULTS keyword, to be modified. They keyword allows for the re-scaling of the existing fault transmissibilities calculated by OPM Flow, for example setting a fault to be completely sealing by setting the multiplier to zero..

Revision: Rev-I

See MULTFLT - Multiply the Transmissibility of a Defined Fault by a Constant in the GRID section for a full description.

7.3.17 MULTIPLY - MULTIPLY A SPECIFIED ARRAY BY A CONSTANT

The MULTIPLY keyword multiplies a specified array or part of an array by a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

7.3.18 MULTIREG - MULTIPLY AN ARRAY BY A CONSTANT BASED ON A REGION NUMBER

The MULTIREG keyword multiplies an array or part of an array by a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTIREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the MULTIREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

7.3.19 MULTPV - MULTIPLY CELL PORE VOLUMES BY A CONSTANT

MULTPV multiples the pore volumes of a cell by a real positive constant for all the cells in the model via an array. An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTPV keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

See MULTPV - Multiply Cell Pore Volumes by a Constantin the GRID section for a full description.

7.3.20 MULTREGP- MULTIPLY PORE VOLUMES BASED ON REGION NUMBER

The MULTREGP keyword multiplies the pore volume of a cell by a constant for all cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTREGP keyword is read by the simulator. The constant should be a real number.

See Error: Reference source not found in the GRID section for a full description.

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7.3.21 MULTREGT- Multiply Transmissibilities Between Regions

The MULTREGT keyword multiplies the transmissibility between two regions by a constant. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTREGT keyword is read by the simulator. The constant should be a real number.

Revision: Rev-I

See Error: Reference source not found in the GRID section for a full description.

7.3.22 MULTX - MULTIPLY CELL TRANSMISSIBILITY IN THE +X DIRECTION

MULTX multiples the transmissibility between two cell faces in the +X direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I, K) and (I+I, J, K).

See MULTX - Multiply Cell Transmissibility in the +X Direction in the GRID section for a full description.

7.3.23 MULTX- - MULTIPLY CELL TRANSMISSIBILITY IN THE -X DIRECTION

MULTX- multiples the transmissibility between two cell faces in the -X direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I-I, I, K) and (I, I, K).

See MULTX- - Multiply Cell Transmissibility in the -X Direction in the GRID section for a full description.

7.3.24 MULTY - MULTIPLY CELL TRANSMISSIBILITY IN THE +Y DIRECTION

MULTY multiples the transmissibility between two cell faces in the +Y direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I, K) and (I, J+I, K).

See MULTY - Multiply Cell Transmissibility in the +Y Direction in the GRID section for a full description.

7.3.25 MULTY- - MULTIPLY CELL TRANSMISSIBILITY IN THE -Y DIRECTION

MULTY- multiples the transmissibility between two cell faces in the -Y direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I-I, K) and (I, J, K).

See MULTY- - Multiply Cell Transmissibility in the -Y Direction in the GRID section for a full description.

7.3.26 MULTZ - MULTIPLY CELL TRANSMISSIBILITY IN THE +Z DIRECTION

MULTZ multiples the transmissibility between two cell faces in the +Z direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I, K) and (I, J, K+1).

See MULTZ - Multiply Cell Transmissibility in the +Z Direction in the GRID section for a full description.

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7.3.27 MULTZ- - MULTIPLY CELL TRANSMISSIBILITY IN THE -Z DIRECTION

MULTZ- multiples the transmissibility between two cell faces in the -Z direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, J, K) between the cells (I, J, K-I) and (I, J, K).

Revision: Rev-I

See MULTZ- - Multiply Cell Transmissibility in the -Z Direction in the GRID section for a full description.

7.3.28 OPERATE - Define Mathematical Operations on Arrays

This keyword defines the mathematical operations on arrays in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

See OPERATE - Define Mathematical Operations on Arrays in the GRID section for a full description.

7.3.29 OPERATER - Define Mathematical Operations on Arrays by Region

This keyword defines the mathematical operations on arrays for specific regions in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

See OPERATER - Define Mathematical Operations on Arrays by Region in the GRID section for a full description.

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7.3.30 PORV - Define the Pore Volumes for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PORV defines the pore volumes for all the cells in the model via an array. The keyword can be used for all grid types, except for the Radial Grid geometry. The keyword effectively overwrites previously entered and calculated data.

No.	Name		Default				
		Field					
I	PORV	PORV is an array of real cell in the model.					
		Repeat counts may be us					
		rb	rb rm³ rcc				

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) Values not reset by this keyword remain unaltered.
- 3) The keyword is terminated by "/".

Table 7.5: PORV Keyword Description

Example

Here the BOX statement is used to define the input grid for the PORV keyword, which overwrites the pore volume previously calculated with pore volume values of zero, resulting in a no-flow boundary in that part of the field between layers 19 and 21, since layer 20 is deactivated. The ENDBOX keyword resets the input box to the full grid.

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Revision: Rev-I

7.3.31 REFINE - START THE DEFINITION OF A LOCAL GRID REFINEMENT

The REFINE keyword defines the start of a Cartesian or radial local grid refinement ("LGR") definition that sets the properties of the selected LGR. The keyword is then followed by the property keywords associated with the section where the keyword is being invoked. For example, if the REFINE keyword is used in the GRID section then most of the keywords in that section can be used to set the grid properties for the LGR.

See REFINE – Start the Definition of a Local Grid Refinement in the GRID section for a full description.

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7.3.32 TRANX - Define the Transmissibility in the X Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

TRANX defines the transmissibility in the X direction for all the cells in the model via an array. The keyword can be used for all grid types, except for the Radial Grid geometry. The keyword effectively overwrites previously entered and calculated data. The transmissibility overwritten is the +X face transmissibility of each grid block, that is for cell (I, I, K) the transmissibility between cells (I, I, K) and (I+I, I, K).

No.	Name		Default						
		Field							
I	TRANX		TRANX is an array of real positive numbers assigning the tranmissibility in the X direction to each cell in the model.						
		Repeat counts may be us							
		cP.rb/day/psia	cP.rb/day/psia cP.rm³/day/bars cP.rcc/hr/atm						

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in I) the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- Values not reset by this keyword remain unaltered.
- The keyword is terminated by "/".

Table 7.6:TRANX Keyword Description

See also the TRANY and TRANYZ keywords to modify the transmissibilities in the other directions.

Example

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
- -
        I1 I2 J1 J2 K1 K2
B0X
                 10 10
                          1 120
                                                        / DEFINE BOX AREA
        SET TRANX+ TRANSMISSIBILITY
- -
TRANX
120*0.00
        DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
--
ENDBOX
```

Here the BOX statement is used to define the input grid for the TRANX keyword, which overwrites the transmissibility previously calculated with transmissibility values of zero, resulting in a no-flow boundary in that part of the field. The ENDBOX keyword resets the input box to the full grid.

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7.3.33 TRANY - Define the Transmissibility in the Y Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

TRANY defines the transmissibility in the Y direction for all the cells in the model via an array. The keyword can be used for all grid types, except for the Radial Grid geometry. The keyword effectively overwrites previously entered and calculated data. The transmissibility overwritten is the +Y face transmissibility of each grid block, that is for cell (I, I, K) the transmissibility between cells (I, I, K) and (I, I+I, K).

No.	Name		Default					
		Field						
I	TRANY	in the Y direction to each	TRANY is an array of real positive numbers assigning the transmissibility in the Y direction to each cell in the model. Repeat counts may be used, for example 20*100.0.					
		cP.rb/day/psia	None					

Notes:

- I) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- Values not reset by this keyword remain unaltered.
- The keyword is terminated by "/".

Table 7.7:TRANY Keyword Description

See also the TRANX and TRANZ keywords to modify the transmissibilities in the other directions.

Example

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
- -
        I1 I2 J1 J2 K1 K2
B0X
                 10 10
                          1 120
                                                        / DEFINE BOX AREA
        SET TRANY+ TRANSMISSIBILITY
_ _
TRANY
120*0.00
        DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
--
ENDBOX
```

Here the BOX statement is used to define the input grid for the TRANY keyword, which overwrites the transmissibility previously calculated with transmissibility values of zero, resulting in a no-flow boundary in that part of the field. The ENDBOX keyword resets the input box to the full grid.

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7.3.34 TRANZ - Define the Transmissibility in the Z Direction for All the Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

TRANX defines the transmissibility in the z direction for all the cells in the model via an array. The keyword can be used for all grid types, except for the Radial Grid geometry. The keyword effectively overwrites previously entered and calculated data. The transmissibility overwritten is the +Z face transmissibility of each grid block, that is for cell (I, I, K) the transmissibility between cells (I, I, K) and (I, I, K+1).

No.	Name		Default					
		Field						
I	TRANZ		TRANZ is an array of real positive numbers assigning the transmissibility in the Z direction to each cell in the model.					
		Repeat counts may be us						
		cP.rb/day/psia	None					

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in I) the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- Values not reset by this keyword remain unaltered.
- The keyword is terminated by "/".

Table 7.8:TRANZ Keyword Description

See also the TRANX and TRANY keywords to modify the transmissibilities in the other directions.

Example

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS
         ----- BOX -----
- -
        I1 I2 J1 J2 K1 K2
B0X
        1* 100
                 1* 100
                          20 20
                                                          / DEFINE BOX AREA
        SET TRANZ+ TRANSMISSIBILITY
_ _
TRAN7
1000*0.00
        DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

Here the BOX statement is used to define the input grid for the TRANZ keyword, which overwrites the transmissibility previously calculated with transmissibility values of zero, resulting in a no-flow boundary in that part of the field between layers 20 and 21. The ENDBOX keyword resets the input box to the full grid.

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8 PROPS SECTION

8.1 Introduction

The PROPS section is the section that contains the fluid property keywords used to define the PVT behavior of the fluids in the model and therefore the data is fluid type dependent. For example, if the oil phase has been activated in the RUNSPEC section via the OIL keyword then one of the oil PVT keywords needs to be defined in this section. The other main type of data required for the PROPS section is the saturation tables that govern how the various fluids flow in the model relative to the other phases. Again, the saturation tables are fluid type dependent.

Revision: Rev-I

8.2 DATA REQUIREMENTS

For all phases activated in the model a complete description of the PVT behavior (PVT tables) and fluid flow behavior (saturations tables) is required. As the data is fluid type dependent, the next two sections attempt to define the appropriate keywords for the various activated fluid types.

8.2.1 Fluid Property Tables

Table 8.1 outlines the various fluid types that can be active in the model, together with the associated RUNSPEC section keywords that activate the phases, versus the PVT keywords that can be used to define the PVT behavior.

Fluid Property Keywords Versus Fluid Type							
Item	Oil		Gas		Water	Polymer Solvent	
Fluid Type	Dead Oil	Live Oil	Dry Gas	Wet Gas	Water	Polymer	Solvent
RUNSPEC Keywords	OIL	OIL DISGAS	GAS	GAS VAPOIL	WATER	POLYMER	SOLVENT
Pressure Dependent PVT	PVCDO PVDO	PVCO PVTO	PVDG PVZD	PVTG	PVTW		PVDS
Surface Density	DENSITY GRAVITY						SDENSITY
Polymer						PLYADS PLYROCK PLYVISC PLMIXPAR PLYMAX PLYSHLOG	

Notes:

- 1) The PVTZD keyword cannot be used in OPM Flow.
- 2) When two keywords are stated for a given fluid type then either one can be used to define the PVT behavior for the given phase.

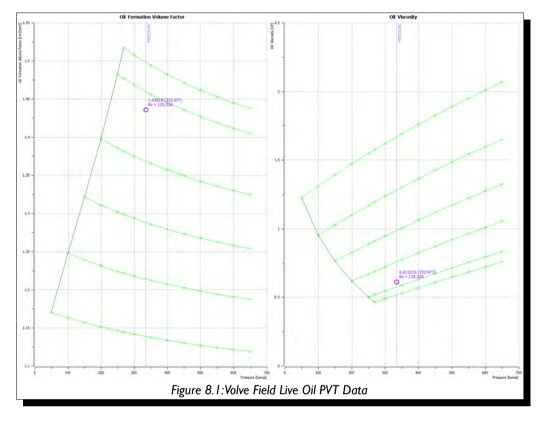
Table 8.1: Fluid Property Keywords versus Fluid Type

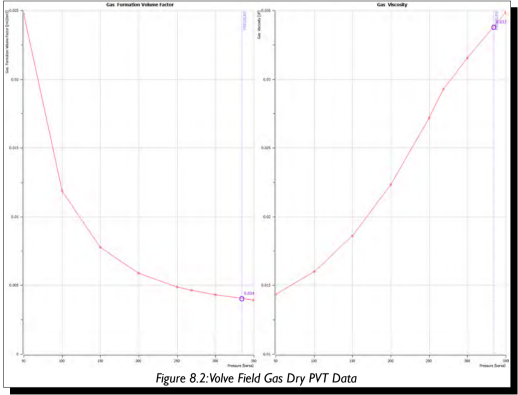
In addition to the above the ROCK keyword should be used to define the rock compressibility.

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Typical live oil and dry gas PVT data is from the Volve³⁵ field is shown in Figure 8.1 and Figure 8.2, respectively.





The Volve Data was approved for data sharing in 2018 by the initiative of the last Operating company, Equinor and approved by the license partners ExxonMobil E&P Norway AS and Bayerngas Norge AS in the end of 2017.

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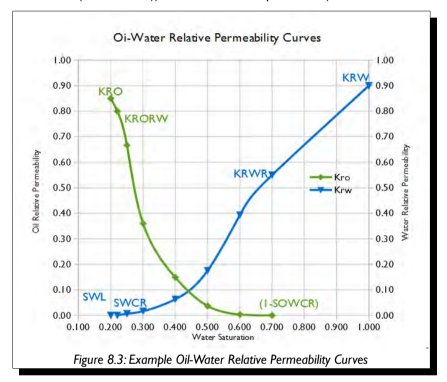
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8.2.2 SATURATION TABLES (RELATIVE PERMEABILITY AND CAPILLARY PRESSURE TABLES)

Saturation tables contain the relative permeability and capillary pressure data as a function of fluid saturation and are used to both initialize the model and to describe multi-phase flow in the reservoir. Multiple saturation tables can be entered and allocated to various areas in the model, based on rock typing. Alternatively, a limited number of saturation tables may be entered and allocated by region and combined with end-point scaling option to enable a more robust reservoir rock characterization.

Revision: Rev-I

A typical oil-water relative permeability set of curves is shown in Figure 8.3 indicating the oil end-point data (KRO, KRORW and (I - SOWCR)) and the water end-point data (KRWR, KRW. SWL and SWCR).



The associated oil-water end-point definitions are outlined in the following table:

Туре	End-Point Keyword	Oil-Water End-Point Definitions			
Saturation	SWL	Connate water saturation, that is the smallest water saturation in a water saturation function table.			
	SWCR	Critical water saturation, that is the largest water saturation for which the water relative permeability is zero.			
	SOWCR	Critical oil-in-water saturation, that is the largest oil saturation for which the oil relative permeability is zero in an oil-water system.			
	KRW	Relative permeability of water at the maximum water saturation (normally the maximum water saturation is one).			
Relative	KRO	Relative permeability of oil at the maximum oil saturation.			
Permeability	KRWR	Relative permeability of water at the residual oil saturation or the residual gas saturation in a gas-water run.			
	KRORW	Relative permeability of oil at the critical water saturation.			

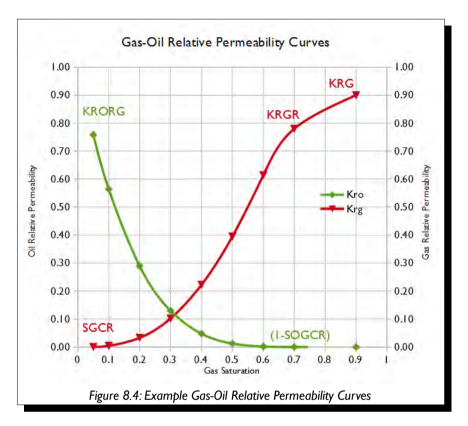
Table 8.2: Oil-Water Relative Permeability End-Point Data Definitions

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Similarly for gas-oil systems, Figure 8.4 illustrates a typical gas-oil relative permeability set of curves indicating the oil end-point data (KRORG and (I - SOGCR)) and the gas end-point data (KRGR, KRG and SGCR).

Revision: Rev-I



The gas-oil end-point definitions are outlined in the following table:

Туре	End-Point Keyword	Gas-Oil End-Point Definitions
Saturation	SGCR	Critical gas saturation, that is the largest gas saturation for which the gas relative permeability is zero.
	SOGCR	Critical oil-in-gas saturation, that is the largest oil saturation for which the oil relative permeability is zero in an oil-gas-connate water system.
Relative Permeability	KRG	Relative permeability of gas at the maximum gas saturation.
	KRGR	Relative permeability of gas at the residual oil saturation or the critical water saturation in a gas-water run.
	KRORG	Relative permeability of oil at the critical gas saturation.

Table 8.3: Gas-Oil Relative Permeability End-Point Data Definitions

End-point scaling is activated in the RUNSPEC section with the ENDSCALE keyword and the data used to apply end-point scaling is entered in the PROPS section using the end-point keywords defined in Table 8.2 and Table 8.3 to define each grid block's end-point data. There is also direction dependent versions of the keywords for when directional end-point scaling has been activated. For example for critical water saturation, SWCR is used with non-direction end-point scaling and the SWCRX±, SWCRX± and SWCRX± series of keyword is used for when directional end-point scaling has been activated. In addition, there is also the facility to incorporate end-point scaling based on the drainage and / or imbibition process which again can be either non-directional or directional.

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Saturation functions can be entered via several keywords consisting of two format types as depicted in the following table:

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Format Type One				Format Type Two			
Keyword	Oil	Gas	Water	Keyword	Oil	Gas	Water
SGOF	Pcog			SGFN ¹		Pcog	
SLGOF	Pcog			SGWFN		Pcgw	
SWOF	Pcwo		Pcwo	SOF2 ²	No Pc		
				SOF3 ³	No Pc		
				SOF32D	No Pc		
				SWFN			Pcwo

Notes:

- In gas-water systems, the gas-water Pcgw data should be entered on the SWFN keyword and the Pcog on the SGFN keyword should be set to zero.
- 2) The SOF2 defines the relative permeability in oil-gas and oil-water runs only, and the miscible hydrocarbon in SOVENT runs. This keyword should not be used to define the oil relative permeability when oil, gas and water are present.
- 3) Defines oil in relative with respect to water and oil relative permeability with respect gas.

Table 8.4: Saturation Table Formats and Phases

Note that only format type can be used in a run, that is one must either use format type one relative permeability keywords to define the required saturation functions, or format two. One cannot combine the keywords from the different format types in the same input deck.

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8.3 Keyword Definitions

8.3.1 ADD - ADD A CONSTANT TO A SPECIFIED ARRAY

The ADD keyword adds a constant to a specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the ADD keyword is being applied.

Revision: Rev-I

See ADD – Add a Constant to a Specified Array in the GRID section for a full description.

8.3.2 ADDREG - ADD A CONSTANT TO AN ARRAY BASED ON A REGION NUMBER

The ADDREG keyword adds a constant to a specified array or part of an array based on cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the ADDREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the ADDREG keyword is being used.

See ADDREG - Add a Constant to an Array based on a Region Number in the GRID section for a full description.

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8.3.3 ADSALNOD - SALT CONCENTRATION BASED ON SATNUM ARRAY

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

ADSALNOD defines the salt concentration value based on a cells SATNUM number. The ADSALNOD property is used in the calculation of a polymer viscosity when the polymer and the salt options has been activated by the POLYMER and BRINE keywords in the RUNSPEC section. In the RUNSPEC section the number of SATNUM functions is declared by the NTSFUN variable on the TABDIMS keyword and allocated to individual cells by the SATNUM property array in the REGIONS section. NSSFUN on the TABDIMS keyword in the RUNSPEC section defines the maximum number of rows (or saturation values) in the relative permeability saturation tables and also sets the maximum number of entries for each ADSALNOD data set. The number of values for each data set must correspond to the number of polymer solution adsorption entries on the PLYADSS keyword. For example, if there are three sets of relative permeability tables and four values on the PLYADSS keyword, then three ADSALNOD data sets with four values of salt concentrations need to be entered.

The salt concentrations within each data set should be positive and monotonically increasing and each ADSALNOD data set is delimited by "/" including the last data set.

No.	Name		Description						
- 1	SALTCON	Field	Field Metric Laboratory						
		A real positive columnar vector that sets the salt concentrations for the given relative permeability saturation tables.							
		lb/stb	kg/sm³	gm/scc	None				
Notes	<u>s:</u>	'							

1) Each data set must be terminated by a "/" including the last data set.

Table 8.5: ADSALNOD Keyword Description

An alternative manner of entering the salt concentrations is by utilizing the PVTNUM region array by using the SALTNODE keyword in the PROPS section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Example

Given three sets of relative permeability tables and four values on the PLYADSS keyword, then the data salt concentration should be entered as follows:

Revision: Rev-I

```
-- SETS SALT CONCENTRATION FOR POLYMER SOLUTION ADSORPTION
-- VIA SATNUM ARRAY ALLOCATION
      SALT
ADSALNOD
      1.0
      5.0
      10.5
      25.0
                / SATNUM TABLE NO. 01
      1.0
      3.0
      7.5
      15.0
                / SATNUM TABLE NO. 02
      1.0
      7.5
      20.5
                / SATNUM TABLE NO. 03
      35.0
```

See also the SALTNODE keyword.

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8.3.4 APIGROUP - Define API Tracking Number of Grouped Oil PVT Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

The APIGROUP keyword defines the maximum number of groups of oil PVT tables when the API tracking option has been activated via the API keyword in the RUNSPEC section.

This keyword is ignored by OPM Flow and has no effect on the simulation

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8.3.5 ASPKDAM - Define Asphaltene Permeability Damage

RUNSPEC	GRID	EDIT	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the data required to model permeability damage, that is permeability reduction, due to the deposit of asphaltene coming out solution. This modeling option is not implemented in OPM Flow as this feature is normally incorporated in compositional simulators.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate.

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8.3.6 ASPPW2D - Define Asphaltene Two Parameters Precipitation Data

RUNSPEC	GRID	EDIT	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

This keyword defines the data required to model asphaltene precipitation case where the precipitation is a function of pressure and temperature based on the percentage molar weight of one or more specified components. The characterization is specified by the first item of the ASPHALTE keyword, and the component range is specified by the ASPFLOC keyword. This modeling option is not implemented in OPM Flow as this feature is normally incorporated in compositional simulators.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate.

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8.3.7 ASPREWG - Define Asphaltene as Percentage Weight

RUNSPEC	GRID	EDIT	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

This keyword defines the data required to model asphaltene precipitation case where the precipitation is a function of pressure based on the percentage molar weight of one or more specified components. The characterization is specified by the first item of the ASPHALTE keyword, and the component range is specified by the ASPFLOC keyword. This modeling option is not implemented in OPM Flow as this feature is normally incorporated in compositional simulators.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate.

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8.3.8 ASPWETF - Define Asphaltene Wettability Factor Data

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDUL

Description

This keyword defines the wettability factor as a function of asphaltene deposition. This modeling option is not implemented in OPM Flow as this feature is normally incorporated in compositional simulators.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate.

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8.3.9 AQUCT - Define Carter-Tracy Analytical Aquifers

The AQUCT keyword defines a Carter Tracy aquifer, the properties of the aquifer, including the Carter-Tracy aquifer influence function associated with the aquifer, defined by the AQUTAB keyword in the PROPS section.

See AQUCT - Define Carter-Tracy Analytical Aquifers in the GRID section for a full description

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8.3.10 AQUTAB - Define Carter-Tracy Aquifer Influence Functions

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

The AQUCT keyword defines additional Carter-Tracy³⁶ aquifer functions to be used in the model. Carter-Tracy representation of the aquifer influx is via a q_w term in the nonlinear aquifer influence function Q(t). It allows the water influx from the aquifer to be represented in the simulator by assuming that there is a constant water influx rate over finite time periods. It is derived from the superposition methods of van Everdingen and Hurst³⁷, whose superposition methods are not suitable for implementation in reservoir simulation software, although they are very useful in interpreting aquifer response. The storage requirements and calculation complexity of handling the resulting superposition formulas can be largely eliminated by use of the Carter-Tracy approximate water influx method.

No.	Name		Description					
		Field	Metric	Laboratory				
I	TD	Dimensionless Time						
		dimensionless	dimensionless	dimensionless	None			
2	PD	Dimensionless Pressure	Dimensionless Pressure					
		dimensionless	dimensionless	dimensionless	None			

Notes:

- 1) The keyword is followed by NIFTBL tables as declared on the AQUDIMS keyword in the RUNSPEC section.
- Each table must contain at least two complete rows with a maximum of NRIFTB rows as declared on the AQUDIMS keyword in the RUNSPEC section. Note that NRIFTB must not be less than than 36 in order to accommodate the default infinite acting Carter-Tracy aquifer influence function.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.6:AQUTAB Keyword Description

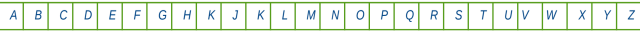
Note

OPM Flow includes the infinite acting Carter-Tracy aquifer influence table as a default for table number one; thus data entered on this keyword starts from table number two.

Table 8.7 to Table 8.10 outline the standard finite tables derived by van Everdingen and Hurst that are used for the Carter-Tracy analytical aquifers and are taken from Bradely³⁸ table 38-6 on page 38-12. In the tables

 r_D is defined as the ratio of the aquifer external radius divided by hydrocarbon radius, that is: $r_D = \frac{r_e}{r_o}$.

³⁸ Bradley Howard B., et. al., Petroleum Engineering Handbook, Society of Petroleum Engineers (June 1989), ISBN:9781555630102.



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³⁶ Carter, R. D., and Tracy, G. W. "An Improved Method for Calculating Water Influx." Transactions of AIME, Vol. 219 (1060), pp 415-417.

³⁷ Van Everdingen, A. F., and Hurst, W. "The Application of the Laplace Transform to Flow Problems in Reservoirs." Transactions of AIME, Vol. 186 (1949), pp. 305-324.

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	Carter-Tracy Aquifer Influence Functions										
No.		= 1.5 nsionless		= 2.0 nsionless	_	= 2.5 nsionless	r _D = 3.0 Dimensionless				
	t _D	P□	t _D	P□	t _D	P□	t _D	P□			
I	0.0600	0.2510	0.2200	0.4430	0.4000	0.5650	0.5200	0.6270			
2	0.0800	0.2880	0.2400	0.4590	0.4200	0.5760	0.5400	0.6360			
3	0.1000	0.3220	0.2600	0.4760	0.4400	0.5870	0.5600	0.6450			
4	0.1200	0.3550	0.2800	0.4920	0.4600	0.5980	0.6000	0.6620			
5	0.1400	0.3870	0.3000	0.5070	0.4800	0.6080	0.6500	0.6830			
6	0.1600	0.4200	0.3200	0.5220	0.5000	0.6180	0.7000	0.7030			
7	0.1800	0.4520	0.3400	0.5360	0.5200	0.6280	0.7500	0.7210			
8	0.2000	0.4840	0.3600	0.5510	0.5400	0.6380	0.8000	0.7400			
9	0.2200	0.5160	0.3800	0.5650	0.5600	0.6470	0.8500	0.7580			
10	0.2400	0.5480	0.4000	0.5790	0.5800	0.6570	0.9000	0.7760			
П	0.2600	0.5800	0.4200	0.5930	0.6000	0.6660	0.9500	0.7910			
12	0.2800	0.6120	0.4400	0.6070	0.6500	0.6880	1.0000	0.8060			
13	0.3000	0.6440	0.4600	0.6210	0.7000	0.7100	1.2000	0.8650			
14	0.3500	0.7240	0.4800	0.6340	0.7500	0.7310	1.4000	0.9200			
15	0.4000	0.8040	0.5000	0.6480	0.8000	0.7520	1.6000	0.9730			
16	0.4500	0.8840	0.6000	0.7150	0.8500	0.7720	2.0000	1.0760			
17	0.5000	0.9640	0.7000	0.7820	0.9000	0.7920	3.0000	1.3280			
18	0.5500	1.0440	0.8000	0.8490	0.9500	0.8120	4.0000	1.5780			
19	0.6000	1.1240	0.9000	0.9150	1.0000	0.8320	5.0000	1.8280			
20			1.0000	0.9820	2.0000	1.2150					
21			2.0000	1.6490	3.0000	1.5960					
22			3.0000	2.3160	4.0000	1.9770					
23			5.0000	3.6490	5.0000	2.3580					

Table 8.7: Carter-Tracy Aquifer Influence Functions ($R_D = 1.5, 2.0, 2.5$ and 3.0)

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		Ca	rter-Tracy	Aquifer In	fluence Fun	ctions		
No.		: 3.5 sionless	r _D = Dimens		r _D = Dimens		r _D = 5.0 Dimensionless	
	t _D	P□	t _D	р₀	t _D	P□	t _D	p _D
I	1.0000	0.8020	1.5000	0.9270	2.0000	1.0230	3.0000	1.1670
2	1.1000	0.8300	1.6000	0.9480	2.1000	1.0400	3.1000	1.1800
3	1.2000	0.8570	1.7000	0.9680	2.2000	1.0560	3.2000	1.1920
4	1.3000	0.8820	1.8000	0.9880	2.3000	1.0720	3.3000	1.2040
5	1.4000	0.9060	1.9000	1.0070	2.4000	1.0870	3.4000	1.2150
6	1.5000	0.9290	2.0000	1.0250	2.5000	1.1020	3.5000	1.2270
7	1.6000	0.9510	2.2000	1.0590	2.6000	1.1160	3.6000	1.2380
8	1.7000	0.9730	2.4000	1.0920	2.7000	1.1300	3.7000	1.2490
9	1.8000	0.9940	2.6000	1.1230	2.8000	1.1440	3.8000	1.2590
10	1.9000	1.0140	2.8000	1.1540	2.9000	1.1580	3.9000	1.2700
П	2.0000	1.0340	3.0000	1.1840	3.0000	1.1710	4.0000	1.2810
12	2.2500	1.0830	3.5000	1.2550	3.2000	1.1970	4.2000	1.3010
13	2.5000	1.1300	4.0000	1.3240	3.4000	1.2220	4.4000	1.3210
14	2.7500	1.1760	4.5000	1.3920	3.6000	1.2460	4.6000	1.3400
15	3.0000	1.2210	5.0000	1.4600	3.8000	1.2690	4.8000	1.3600
16	4.0000	1.4010	5.5000	1.5270	4.0000	1.2920	5.0000	1.3780
17	5.0000	1.5790	6.0000	1.5940	4.5000	1.3490	5.5000	1.4240
18	6.0000	1.7570	6.5000	1.6600	5.0000	1.4030	6.0000	1.4690
19			7.0000	1.7270	5.5000	1.4570	6.5000	1.5130
20			8.0000	1.8610	6.0000	1.5100	7.0000	1.5560
21			9.0000	1.9940	7.0000	1.6150	7.5000	1.5980
22			10.0000	2.1270	8.0000	1.7190	8.0000	1.6410
23					9.0000	1.8230	9.0000	1.7250
24					10.0000	1.9270	10.0000	1.8080
25					11.0000	2.0310	11.0000	1.8920
26					12.0000	2.1350	12.0000	1.9750
27					13.0000	2.2390	13.0000	2.0590
28					14.0000	2.3430	14.0000	2.1420
29					15.0000	2.4470	15.0000	2.2250

Table 8.8: Carter-Tracy Aquifer Influence Functions ($R_D = 3.5, 4.0, 4.5$ and 5.0)

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	Carter-Tracy Aquifer Influence Functions									
No.	r _D = Dimens		r _D = Dimens			r _D = 8.0 Dimensionless		9.0 ionless		
	t _D	р₀	t _D	р₀	t _D	р₀	t _D	P□		
I	4.0000	1.2750	6.0000	1.4360	8.0000	1.5560	10.0000	1.6510		
2	4.5000	1.3220	6.5000	1.4700	8.5000	1.5820	10.5000	1.6730		
3	5.0000	1.3640	7.0000	1.5010	9.0000	1.6070	11.0000	1.6930		
4	5.5000	1.4040	7.5000	1.5310	9.5000	1.6310	11.5000	1.7130		
5	6.0000	1.4410	8.0000	1.5590	10.0000	1.6530	12.0000	1.7320		
6	6.5000	1.4770	8.5000	1.5860	10.5000	1.6750	12.5000	1.7500		
7	7.0000	1.5110	9.0000	1.6130	11.0000	1.6970	13.0000	1.7680		
8	7.5000	1.5440	9.5000	1.6380	11.5000	1.7170	13.5000	1.7860		
9	8.0000	1.5760	10.0000	1.6630	12.0000	1.7370	14.0000	1.8030		
10	8.5000	1.6070	11.0000	1.7110	12.5000	1.7570	14.5000	1.8190		
П	9.0000	1.6380	12.0000	1.7570	13.0000	1.7760	15.0000	1.8350		
12	9.5000	1.6680	13.0000	1.8010	13.5000	1.7950	15.5000	1.8510		
13	10.0000	1.6980	14.0000	1.8450	14.0000	1.8130	16.0000	1.8670		
14	11.0000	1.7570	15.0000	1.8880	14.5000	1.8310	17.0000	1.8970		
15	12.0000	1.8150	16.0000	1.9310	15.0000	1.8490	18.0000	1.9260		
16	13.0000	1.8730	17.0000	1.9740	17.0000	1.9190	19.0000	1.9550		
17	14.0000	1.9310	18.0000	2.0160	19.0000	1.9860	20.0000	1.9830		
18	15.0000	1.9880	19.0000	2.0580	21.0000	2.0510	22.0000	2.0370		
19	16.0000	2.0450	20.0000	2.1000	23.0000	2.1160	24.0000	2.0900		
20	17.0000	2.1030	22.0000	2.1840	25.0000	2.1800	26.0000	2.1420		
21	18.0000	2.1600	24.0000	2.2670	30.0000	2.3400	28.0000	2.1930		
22	19.0000	2.2170	26.0000	2.3510	35.0000	2.4990	30.0000	2.2440		
23	20.0000	2.2740	28.0000	2.4340	40.0000	2.6580	34.0000	2.3450		
24	25.0000	2.5600	30.0000	2.5170	45.0000	2.8170	38.0000	2.4460		
25	30.0000	2.8460					40.0000	2.4960		
26							45.0000	2.6210		
27							50.0000	2.7460		

Table 8.9: Carter-Tracy Aquifer Influence Functions ($R_D = 6.0, 7.0, 8.0$ and 9.0)

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Carter-Tracy Aquifer Influence Functions													
			Linear sionless										
t _D	р₀	t _D	р₀										
12.0000	1.7320	0.0050	0.0798										
12.5000	1.7500	0.0100	0.1130										
13.0000	1.7680	0.0200	0.1596										
13.5000	1.7840	0.0600	0.2764										
14.0000	1.8010	0.0800	0.3192										
14.5000	1.8170	0.1000	0.3568										
15.0000	1.8320	0.1200	0.3909										
15.5000	1.8470	0.1600	0.4515										
16.0000	1.8620	0.2000	0.5052										
17.0000	1.8900	0.2400	0.5544										
18.0000	1.9170	0.3000	0.6228										
19.0000	1.9430	0.4000	0.7294										
20.0000	1.9680	0.6000	0.9328										
22.0000	2.0170	0.8000	1.1333										
24.0000	2.0630	1.0000	1.3333										
26.0000	2.1080												
28.0000	2.1510												
30.0000	2.1940												
32.0000	2.2360												
34.0000	2.2780												
36.0000	2.3190												
38.0000	2.3600												
40.0000	2.4010												
50.0000	2.6040												
60.0000	2.8060												
70.0000	3.0080												
	r _D = Dimens t _D 12.0000 12.5000 13.0000 13.5000 14.0000 14.5000 15.5000 16.0000 17.0000 18.0000 20.0000 24.0000 24.0000 24.0000 28.0000 30.0000 32.0000 34.0000 36.0000 38.0000 38.0000 40.0000 50.0000	r _D = 10.0 Dimensionless t _D p _D 12.0000 1.7320 12.5000 1.7500 13.0000 1.7680 13.5000 1.7840 14.0000 1.8010 14.5000 1.8170 15.0000 1.8320 15.5000 1.8470 16.0000 1.8620 17.0000 1.8900 18.0000 1.9170 19.0000 1.9430 20.0000 1.9680 22.0000 2.0170 24.0000 2.0630 26.0000 2.1080 28.0000 2.1510 30.0000 2.1940 32.0000 2.2360 34.0000 2.3190 38.0000 2.3190 38.0000 2.4010 50.0000 2.6040 60.0000 2.8060	r _D = 10.0 Dimensionless Finite Dimensionless t _D p _D t _D 12.0000 1.7320 0.0050 12.5000 1.7500 0.0100 13.0000 1.7680 0.0200 13.5000 1.7840 0.0600 14.0000 1.8010 0.0800 14.5000 1.8170 0.1000 15.0000 1.8470 0.1600 15.5000 1.8470 0.1600 16.0000 1.8620 0.2000 17.0000 1.8900 0.2400 18.0000 1.9170 0.3000 19.0000 1.9430 0.4000 20.0000 2.0170 0.8000 24.0000 2.0630 1.0000 26.0000 2.1080 1.0000 28.0000 2.1510 30.0000 32.0000 2.2780 36.0000 34.0000 2.3600 40.0000 2.4010 50.0000 2.6040 60.0000 2.8060 60.0000										

Table 8.10: Carter-Tracy Aquifer Influence Functions ($R_D = 10$ and Finite Linear)

For the finite linear Carter-Tracy influence function in Table 8.10 set the inner radius of the aquifer to the $= \frac{360 \times Width}{(2 \times \pi \times Length)}$ on the AQUCT keyword in the length of linear aquifer and the angle of influence to grid section.

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For reference Table 8.11 outlines the content terminal rate case for an infinite aquifer derived by van Everdingen and Hurst that is the default table number one used for the Carter-Tracy analytical aquifers and is taken from Bradely³⁹ table 38-3 on page 38-6.

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Ca	rter-Tracy I	nfinite Ra unction (ence
No.	Infini Dimensi		No.		nite sionless
	T _D	PD		T₀	P _D
I	1.0 x 10 ⁻²	0.112	19	4.0	1.275
2	5.0 x 10 ⁻²	0.229	20	5.0	1.362
3	1.0 x 10 ⁻¹	0.315	21	6.0	1.436
4	1.5 x 10 ⁻¹	0.376	22	7.0	1.500
5	2.0 x 10 ⁻¹	0.424	23	8.0	1.556
6	2.5 x 10 ⁻¹	0.469	24	9.0	1.604
7	3.0 x 10 ⁻¹	0.503	25	10.0	1.651
8	4.0 x 10 ⁻¹	0.564	26	15.0	1.829
9	5.0 x 10 ⁻¹	0.616	27	20.0	1.960
10	6.0 x 10 ⁻¹	0.659	28	25.0	2.067
11	7.0 x 10 ⁻¹	0.702	29	30.0	2.147
12	8.0 x 10 ⁻¹	0.735	30	40.0	2.282
13	9.0 x 10 ⁻¹	0.772	31	50.0	2.388
14	1.0	0.802	32	60.0	2.476
15	1.5	0.927	33	70.0	2.550
16	2.0	1.020	34	80.0	2.615
17	2.5	1.101	35	90.0	2.672
18	3.0	1.169	36	100.0	2.723

Table 8.11: Carter-Tracy Infinite Radial Aquifer Influence Function (Default)

For an overview of analytical aquifers see Dake⁴⁰.

Dake, L.P. Fundamentals of Reservoir Engineering, Amsterdam, The Netherlands, Elsevier Science BV (1978) Chapter 9.

Α	В	С	D	Ε	F	G	Н	K	J	Κ	L	М	Ν	0	Р	Q	R	S	Т	U	V	W	Χ	Υ	Z
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³⁹ Bradley Howard B., et. al., Petroleum Engineering Handbook, Society of Petroleum Engineers (June 1989), ISBN:9781555630102.

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Example CARTER-TRACY AQUIFER INFLUENCE TABLES (STARTS FROM TABLE NO. 2, AS DEFAULT IS TABLE NO. 1) AQUTAB TD PD _ _ 0.06 0.251 0.08 0.288 0.322 0.10 0.12 0.355 0.14 0.387 0.16 0.420 0.18 0.452 0.20 0.484 0.22 0.516 0.24 0.548 0.26 0.580 0.28 0.612 0.30 0.644 0.35 0.724 0.40 0.804 0.45 0.884 0.50 0.964 0.55 1.044 / RD=1.5 TABLE NO. 02 0.60 1.124 PDTD 0.22 0.443 0.24 0.459 0.26 0.476 0.28 0.492 0.30 0.507 0.32 0.522 0.34 0.536 0.36 0.551 0.38 0.565 0.40 0.579 0.42 0.593 0.44 0.607 0.46 0.621 0.48 0.634 0.50 0.648 0.6 0.715 0.7 0.782 0.8 0.849 0.9 0.915 1.0 0.982 2.0 1.649

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/ RD=2.0 TABLE NO. 03

The above example defines tables two and three Carter-Tracy aquifer influence tables.

2.316

3.649

3.0

5.0

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8.3.11 BOX - DEFINE A RANGE OF GRID BLOCKS TO ENTER PROPERTY DATA

BOX defines a range of grid blocks for which subsequent data is assigned for all the cells in the defined BOX. Note that the BOX grid is reset by the keyword ENDBOX by resetting the current defined BOX to be the whole grid. The keyword can be used for any array and for all grid types.

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See BOX - Define a Range of Grid Blocks to Enter Property Data in the GRID section for a full description.

8.3.12 COPY - COPY ARRAY DATA TO ANOTHER ARRAY

The COPY keyword copies an array (or part of an array) to another array or part of an array. The arrays can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the COPY keyword is being used.

See COPY - Copy Array Data to Another Array in the GRID section for a full description.

8.3.13 COPYREG - COPY AN ARRAY TO ANOTHER ARRAY BASED ON A REGION NUMBER

The COPYREG keyword copies a specified array or part of an array based on cells with a specific region number to another array. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the COPYREG keyword is read by the simulator. The property arrays can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the COPYREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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8.3.14 DENSITY - Define the Surface Oil, Water Gas Densities for the Fluids

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

DENSITY defines the oil, water and gas surface densities for the fluids for various regions in the model. The number of DENSITY vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the DENSITY data sets to different grid blocks in the model is done via the PVTNUM keyword in the REGION section. One data set consists of one record or line which is terminated by a "/". This surface density or gravity must be entered using either the DENSITY or GRAVITY keywords irrespective of which phases are active in the model.

No.	Name		Description		Default
		Field	Metric	Laboratory	
ı	OILDEN	OILDEN is a real number	r defining the density of oi	l at surface conditions.	
		lb/ft³	kg/m³	gm/cc	
		600	600	600	Defined
2	WATDEN	WATDEN is a real null conditions.			
		lb/ft³	kg/m³	gm/cc	
		999.014	999.014	999.014	Defined
3	GASDEN	GASDEN is a real nu conditions.	mber defining the dens	sity of gas at surface	
		lb/ft³	kg/m³	gm/cc	
		1.000	1.000	1.000	Defined

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.12: DENSITY Keyword Description

According to the SPE SI standard⁴¹, **Relative Density** (γ) replaces **Specific Gravity** as the term used to define the ratio of the density of a known material to the density of reference material, at standard conditions of pressure and temperature. Standard conditions vary throughout the world, but for oil field units one normally uses I4.7 psia and 60 °F, while for SI units some areas use I0I.325 kPa and I5 °C.

See also the GRAVITY keyword.

The SI Metric System of Units and SPE Metric Standard, Adopted for Use as a Voluntary Standard by the SPE Board of Directors, June 1983, Society of Petroleum Engineers.



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Examples

The following shows the DENSITY keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to one.

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The next example shows the DENSITY keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to three.

```
- -
        OIL
                WAT
                           GAS
        DENSITY DENSITY
- -
                           DENSITY
DENSITY
        38.0
                                                        / PVT DATA REGION 1
                62.30
                          0.04500
        39.0
                           0.04520
                                                        / PVT DATA REGION 2
                 62.37
                62.40
                           0.04800
                                                        / PVT DATA REGION 3
```

There is no terminating "/" for this keyword.

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8.3.15 EHYSTR - Define Hysteresis Model and Parameters

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

The EHYSTR keyword defines the hysteresis model and associated parameters when the hysteresis option has been activated by the HYSTER variable on the SATOPTS keyword in the RUNSPEC section. Both the Carlson⁴² and Killough⁴³ models are available.

No.	Name		Description		Default						
I	HYSTRCP		s a positive real value that defi or capillary pressure hysteresis mo		0.1						
		The value sh	ould range from 0.05 to 0.10.								
		This option i	s ignored by OPM Flow.								
2	HYSTMOD	model to be	An integer value that determines the relative permeability hysteresis model to be used depending on the phase and the wettability of the system. HYSTMOD should be set to one of the following values:								
			Water Wet Hysteresis N	Models							
		HYSMOD	Non-Wetting Phases	Wetting Phase							
		-1	prating the model with the nd running the model with								
		This option implies no hysteresis.									
		0	Carlson Hysteresis Model	SATNUM							
		I	Carlson Hysteresis Model	IMBNUM							
		2	Killough Hysteresis Model	SATNUM							
		3	Killough Hysteresis Model	IMBNUM							
		4	Killough Hysteresis Model	Killough Hysteresis Model							
			Oil Wet to Water Wet Wa	ater Wet							
		5	Carlson Non- Wetting Modeling for Gas and Water	SATNUM							
		6	Killough Non- Wetting Modeling for Gas and Water	SATNUM							
		7	Killough Non- Wetting Modeling for Gas and Water	Killough Non- Wetting Modeling for the Wetting Oil Phase							

⁴³ Killough, J. E. "Reservoir Simulation with History-dependent Saturation Functions," paper SPE 5106, Society of Petroleum Engineers Journal (1976) 16, No. 1, 37-48.

Α	В	С	D	Ε	F	G	Н	K	J	Κ	L	М	N	0	Р	Q	R	S	Т	U	V	W	Χ	Υ	Z
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⁴² Carlson, F. M. "Simulation of Relative Permeability Hysteresis to the Non-Wetting Phase," paper SPE 10157, presented at the SPE Annual Technical Conference & Exhibition, San Antonio, Texas, USA (October 5-7, 1981).

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No.	Name	Description	Default
3	HYSTREL	HYSTREL is a positive real number that defines the Killough's wetting phase relative permeability curvature parameter. This parameter is only applicable if HYSMOD is set to either 4 or 7.	1.0
		This option is ignored by OPM Flow.	
4	HYSTSGR	HYSTSGR is a positive real number that sets a scaling parameter for the trapped non-wetting phase saturation in the Killough model.	0.1
		This option is ignored by OPM Flow.	
5	HYSTOPT	A character string that determines if the hysteresis model should be activated for relative permeability, capillary pressure curves, or both. And should be set to one of the following:	вотн
		 BOTH: apply hysteresis modeling to both relative permeability, and capillary pressure curves. 	
		2) PC: apply hysteresis modeling to capillary pressure curves only.	
		3) KR: apply hysteresis modeling to relative permeability curves only.	
		For all cases HYSTMOD defines the model to be used for relative permeability hysteresis modeling (if applicable). Capillary pressure hysteresis always users the Killough capillary pressure model.	
		Note only the default value of BOTH is supported by OPM Flow.	
6	HYSTSCAN	A character string that determines the shape of Killough capillary pressure scanning curves when secondary reversal curves, that is for a drainage, imbibition, drainage cycle.	RETR
		RETR: Secondary drainage curves re-traverses the same scanning curve.	
		 NEW: Secondary drainage curves follows a new scanning curve and further reversals also generate a new scanning curve. 	
		This option is ignored by OPM Flow.	
7	HYSTMOB	A character string that determines how to apply the mobility control correction invoked by the MOBILE variable on the EQLOPTS keyword in the RUNSPEC section. HYSTMOB should be set to one of the following:	DRAIN
		DRAIN: Only the drainage curve end-points are modified.	
		 BOTH: Both the drainage and imbibition curve end-points are modified. 	
		The Mobility Control option is not supported in OPM Flow so this parameter has no effect.	
8	HYSTWET	A character string that sets the wetting phase in three phase systems to either oil or gas and should be set to one of the following:	None
		 OIL: Oil is set as the wetting phase and the oil-gas relative permeability curves are determined by HYSTMOD for the wetting phase. 	
		 GAS: Oil is set as the non-wetting phase and the oil-gas relative permeability curves are determined by HYSTMOD for the non- wetting phase. 	
		Note for all the above cases the gas relative permeability curves are always treated as as a non-wetting phase.	
		This option is ignored by OPM Flow.	
9		Not used	
10		Not used	

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No.	Name	Description	Default
-11		Not used	
12		Not used	
13		Not used	
Notes			

Revision: Rev-I

1) The keyword is terminated by "/".

Table 8.13: EHYSTR Keyword Description

Example

```
-- HYSTERESIS MODEL AND PARAMETERS
-- PC-CUR MODEL RELPERM TRAPPED OPTION SHAPE MOBILIT WET
-- HYSTRCP HYSTMOD HYSTREL HYSTSGR HYSTOPT HYSTSCAN HYSTMOB HYSTWET
EHYSTR

0.1 0 0.1 1* KR 1* 1* 1* /
```

The above example defines the hysteresis model and parameters used in the Norne model. Here the default value is used for the Killough curvature parameter for capillary pressure hysteresis mode, the Carlson hysteresis model is used for the non-wetting phase and SATNUM for the wetting phase, 0.1 is used for Killough's wetting phase relative permeability curvature parameter (this parameter is ignored because the Carlson model has been selected), the default values for the trapped non-wetting phase saturation in the Killough mode (again, this parameter is ignored because the Carlson model has been selected, and the hysteresis modeling is only applied to relative permeability curves.

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8.3.16 ENDBOX - Define the End of the BOX Defined Grid

This keyword marks the end of a previously defined BOX sub-grid as defined by a previously entered BOX keyword. The keyword resets the input grid to be the full grid as defined by the NX, NY, and NZ variables on the DIMENS keyword in the RUNSPEC section.

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See ENDBOX - Define the End of the BOX Defined Grid in the GRID section for a full description.

8.3.17 ENDFIN - END THE DEFINITION OF A LOCAL GRID REFINEMENT

ENDFIN defines the end of a Cartesian or radial local grid refinement ("LGR") definition and a LGR property definition data set.

See ENDFIN - End the Definition of a Local Grid Refinement in the GRID section for a full description.

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8.3.18 ENKRVD - Define Relative Permeability End-Points versus Depth

FUNCTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the maximum oil, gas, and water relative permeability versus depth for the three phases. This functionality is not supported in OPM Flow.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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8.3.19 ENPTVD - Define Relative Permeability Saturation End-Points versus Depth

RUNS	SPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the variation of the relative permeability saturation end-points (SWL, SWCR, etc.) for all three phases versus depth. This functionality is not supported in OPM Flow.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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8.3.20 EQUALREG - Sets an Array to a Constant by Region Number

The EQUALREG keyword sets a specified array to a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the EQUALREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the EQUALREG keyword is being used.

Revision: Rev-I

See Error: Reference source not found in the GRID section for a full description.

8.3.21 EQUALS - Sets a Specified Array to a Constant

The EQUALS keyword sets a specified array or part of an array to a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the EQUALS keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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8.3.22 FILLEPS - ACTIVATE SATURATION END-POINT EXPORT TO THE INIT FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword switches on the export of the saturation end-point data (SWL, SWCR, SOWCR array etc.) to the *.INIT file so that the data can be viewed in post-processing software

There is no data required for this keyword.

This keyword is not supported by OPM Flow but is documented here for completeness.

Example

-- ACTIVATES SATURATION END-POINT EXPORT TO THE INIT FILE
-- FILLEPS

The above example switches on the export of the end-point saturation data to the *.INIT file.

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8.3.23 GASDENT - Define Gas Density Temperature Coefficients

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

GASDENT defines the gas density as a function of temperature coefficients for when OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC. The reference pressure for this table is given by the VISCREF keyword in the PROPS section. Note this is an OPM Flow keyword used with OPM Flow's "black-oil" thermal model that is not available in the commercial simulator's "black-oil" thermal formulation.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	TEMP	TEMP is a real positive value greater than zero that defines the absolute reference temperature used with TEXP1 and TEXP2 to estimate the change in gas density with respect to temperature.					
		°R	°K	°K			
		527.67	293.15	293.15	Defined		
2	TEXPI	TEXPI is a real positive value greater than zero that defines the gas thermal expansion coefficient of the first order.					
		I/°R	I/°K	I/°K			
		1.67 x 10 ⁻⁴	3.0×10^{-4}	3.0 x 10 ⁻⁴	Defined		
3	TEXP2 is a real positive value greater than zero that defines the gas thermal expansion coefficient of the second order.						
		I/°R²	I/°K²	I/°K²			
		9.26 x 10 ⁻⁷	3.0×10^{-6}	3.0 × 10 ⁻⁶	Defined		

Notes:

- 1) The keyword is followed by NTPVT records as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each data set is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.14: GASDENT Keyword Description

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Example

The following example shows the GASDENT keyword using the default values, for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC section and for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to two.

Revision: Rev-I

```
- -
         GAS DENSITY TEMPERATURE COEFFICIENTS (OPM FLOW THERMAL KEYWORD)
         GAS
                    DENSITY
                               DENSITY
_ _
         TEMP
                    COEFF1
                               COEFF2
GASDENT
         1*
                    1*
                               1*
                                                              / TABLE NO. 01
         1*
                    1*
                               1*
                                                              / TABLE NO. 02
```

There is no terminating "/" for this keyword.

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8.3.24 GASVISCT - Define Gas Viscosity versus Temperature Functions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

GASVISCT defines the gas viscosity as a function of temperature for when OPM Flow's thermal option has been activated by the THERMAL keywords in the RUNSPEC. The reference pressure for this table is given by the VISCREF keyword in the PROPS section. Note this is an OPM Flow keyword used with OPM Flow's "black-oil" thermal model that is not available in the commercial simulator's "black-oil" thermal formulation.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	TEMP	A columnar vector of revalues that defines the te	real monotonically increa mperature values.	sing down the column	
		°F	°C	°C	None
2	VIS	A columnar vector of reather gas viscosity for the contraction			
		VIS should be given at the on the VISCREF keyword	ned by the PRS variable		
		сР	сР	сР	None

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.15: GASVISCT Keyword Description

Example

The following example shows the GASVISCT keyword for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC section and for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set equal to one.

```
GAS VISCOSITY VERSUS TEMPERATURE TABLES (OPM FLOW EXTENSION KEYWORD)
         GAS
                    GAS
         TEMP
                    VISC
GASVISCT
           100.0
                    0.0500
           110.0
                    0.0550
           120.0
                    0.0580
           150.0
                    0.0620
           165.0
                    0.0625
                                                             / TABLE NO. 01
```

There is no terminating "/" for this keyword.

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8.3.25 GRAVITY- Define the Surface Oil, Water Gas Gravities for the Fluids

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

GRAVITY defines the oil API gravity and water and gas surface specific gravities for the fluids for various regions in the model. The number of GRAVITY vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the GRAVITY data sets to different grid blocks in the model is done via the PVTNUM keyword in the REGION section. One data set consists of one record or line which is terminated by a "/".

This surface density or gravity must be entered using either the DENSITY or GRAVITY keywords irrespective of which phases are active in the model.

No.	Name		Default						
		Field	Metric	Laboratory					
I	OILAPI	OILAPI is a real number	defining the density of oil	at surface conditions.					
			n Institute (API) classifies es API (°API), the relatio I gravity (γ _{API}) is given by:						
		$\gamma_{\scriptscriptstyle AI}$	$\gamma_{API} = \frac{141.5}{\gamma_o} - 131.5$						
		°API	°API	°API	None				
2	WATGRAV	WATGRAV is a real nu conditions.	imber defining the densi	ty of water at surface					
		(water = 1.0)	(water =1.0)	(water =1.0)					
		0.7773	0.7773	0.7773	Defined				
3	GRAVGAS	GRAVGAS is a real nuconditions.	GRAVGAS is a real number defining the density of gas at surface conditions.						
		(air =1.0)	(air = 1.0) (air = 1.0) (air = 1.0)						
		1.000	1.000	1.000	Defined				

Notes:

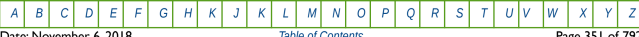
- The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section. I)
- The each data set terminated by terminated by "/" at the end of the line, there is no "/" terminator for the keyword.

Table 8.16: GRAVITY Keyword Description

According to the SPE SI standard⁴⁴, **Relative Density** (y) replaces **Specific Gravity** as the term used to define the ratio of the density of a known material to the density of reference material, at standard conditions of pressure and temperature. Standard conditions vary throughout the world, but for oil field units one normally uses I4.7 psia and 60 °F, while for SI units some areas use I0I.325 kPa and I5 °C.

This keyword is not supported by OPM Flow but is documented here for completeness; however, the density data can be entered using the DENSITY keyword.

The SI Metric System of Units and SPE Metric Standard, Adopted for Use as a Voluntary Standard by the SPE Board of Directors, June 1983, Society of Petroleum Engineers.



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Examples

The following shows the GRAVITY keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to one.

Revision: Rev-I

```
-- OIL WAT GAS
-- GRAVITY GRAVITY GRAVITY
-- GRAVITY
39.0 1.012 0.650 / GRAVITY PVT DATA REGION 1
```

The next example shows the DENSITY keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to three.

	OIL	WAT	GAS	
	GRAVITY	GRAVITY	GRAVITY	
GRAVITY				
	37.0	1.012	0.650	/ GRAVITY PVT DATA REGION 1
	38.0	1.012	0.646	/ GRAVITY PVT DATA REGION 2
	39.0	1.012	0.640	/ GRAVITY PVT DATA REGION 3

There is not terminating "/" for this keyword.

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8.3.26 IMKRVD - Imbibition Relative Permeability End-Points versus Depth Functions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the maximum imbibition oil, gas, and water relative permeability versus depth for the three phases. This functionality is not supported in OPM Flow.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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8.3.27 IMPTVD - Imbibition Relative Permeability Saturation End-Points versus Depth

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the variation of the imbibition relative permeability saturation end-points (SWL, SWCR, etc.) for all three phases versus depth. This functionality is not supported in OPM Flow.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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8.3.28 IPCG - End-Point Scaling of Grid Cell Gas Capillary Pressure (Imbibition)

RUNS	PEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
------	-----	------	------	-------	---------	----------	---------	----------	--

Description

IPCG defines the maximum <u>imbibition</u> gas-oil capillary pressure values for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. <u>In addition, the HYSTER option on the SATOPTS keyword in the RUNSPEC section has to be activated to invoke the hysteresis option</u>. The keyword can be used for all grid types.

The capillary pressure for a grid block is scaled by:

$$P_c = P_{c_{TABLE}} \left(\frac{IPCG}{P_{c_{TABLE-MAX}}} \right)$$
 (8.1)

Revision: Rev-I

Where:

P_c = the resulting imbibition gas-oil capillary pressure for a grid cell.

IPCG = the maximum capillary pressure from the IPCG array for a given cell.

 $P_{c_{\mathit{TABLE}}}$ = the capillary pressure in the inhibition capillary pressure table

allocated to the grid block.

= the maximum capillary pressure in the inhibition capillary pressure table

allocated to the grid block at $S_a = 1 - S_{wco}$.

No.	Name		Description						
		Field	Field Metric Laboratory						
I	IPCG		IPCG is an array of positive real numbers assigning the maximum imbibition gas capillary pressure values for each cell in the model.						
		Repeat counts may be us	Repeat counts may be used, for example 30*100.0.						
		psia	bars	atm	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 8.17: IPCG Keyword Description

See also the PCG keyword for the equivalent drainage functionality.

Example

```
--
-- DEFINE GRID BLOCK IPCG DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
--
IPCG
100*50.0 100*75.0 100*125.0
```

The above example defines the a IPCG for 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.29 IPCW - End-Point Scaling of Grid Cell Water Capillary Pressure (Imbibition)

RUNS	PEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
------	-----	------	------	-------	---------	----------	---------	----------	--

Description

IPCW defines the maximum <u>imbibition</u> water-oil or water-gas capillary pressure values for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable endpoint scaling and the use of this keyword. In addition, the HYSTER option on the SATOPTS keyword in the RUNSPEC section has to be activated to invoke the hysteresis option. The keyword can be used for all grid types.

The capillary pressure for a grid block is scaled by:

$$P_c = P_{c_{TABLE}} \left(\frac{IPCW}{P_{c_{TABLE}-MAX}} \right)$$
 (8.2)

Revision: Rev-I

Where:

P_c = the resulting imbibition water capillary pressure for a grid cell. IPCW = the maximum capillary pressure from the IPCW array for a given cell.

 $P_{c_{\textit{TABLE}}}$ = the capillary pressure in the inhibition capillary pressure table

allocated to the grid block.

 $P_{c_{\textit{TABLE-MAX}}}$ = the maximum capillary pressure in the inhibition capillary pressure table allocated to the grid block (that is at the connate water saturation).

No.	Name	Description					
		Field	Metric	Laboratory			
I	IPCW	IPCW is an array of imbibition water capillary					
		Repeat counts may be used, for example 30*100.0.					
		psia	bars	atm	None		

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 8.18: IPCW Keyword Description

See also the PCW keyword for the equivalent drainage functionality.

Example

```
--
-- DEFINE GRID BLOCK IPCW DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
--
IPCW
100*50.0 100*75.0 100*125.0
```

The above example defines the a IPCW for 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.30 ISGCR - End-Point Scaling of Grid Cell Critical Gas Saturation (Imbibition)

RUNS	PEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
------	-----	------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

ISGCR defines the <u>imbibition</u> critical gas saturation for all the cells in the model via an array when the endpoint scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section and <u>the hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section</u>. The critical gas saturation is defined as the maximum gas saturation for which the gas relative permeability is zero in a two-phase relative permeability table.

The keyword can be used for all grid types.

No.	Name		Default					
		Field	Field Metric Laboratory					
I	ISGCR	values to each cell in the to the NX x NY x NZ pa	ISGCR is an array of real numbers assigning the critical gas saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.03					
		dimensionless	dimensionless	dimensionless	permeability table.			

Notes:

- Note this the non-direction dependent version of the critical gas saturation used with the end-point scaling option. If directional end-point scaling has been activated then the ISGCRX± , ISGCRX± and ISGCRX± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.19: ISGCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISGCRX, ISGCRY and ISGCRZ instead of ISGCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISGCRX, ISGCRX-, ISGCRY, ISGCRY, ISGCRY, ISGCRZ-, instead of the ISGCR keyword.

Example

```
--
-- DEFINE GRID BLOCK END-POINT ISGCR DATA FOR ALL CELLS (NX x NY x NZ = 300)
--
ISGCR
300*0.050
```

The above example defines a constant critical gas saturation of 0.05 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.31 ISGL - End-Point Scaling of Grid Cell Connate Gas Saturation (Imbibition)

RUNS	PEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
------	-----	------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

ISGL defines the <u>imbibition</u> connate gas saturation for all the cells in the model via an array when the endpoint scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section and <u>the hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section</u>. The connate gas saturation is defined as the minimum gas saturation in a two-phase gas relative permeability table.

The keyword can be used for all grid types.

No.	Name	Description							
		Field Metric Laboratory							
I	ISGL	values to each cell in the to the NX x NY x NZ pa	ISGL is an array of real numbers assigning the connate gas saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.03 dimensionless						
		dimensionless	dimensionless	dimensionless	permeability table.				

Notes:

- Note this the non-direction dependent version of the connate gas saturation array used with the end-point scaling option. If directional end-point scaling has been activated then the ISGLX±, ISGLX± and SGZ± series of keyword should be used.
- 2) If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.20: ISGL Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISGLX, ISGLY and ISGLZ instead of ISGL. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISGLX, ISGLY-, ISGLY-, ISGLZ and ISGLZ-, instead of the ISGL keyword.

Example

```
-- DEFINE GRID BLOCK END-POINT ISGL DATA FOR ALL CELLS (NX x NY x NZ = 300)
-- ISGL 300*0.030
```

The above example defines a constant connate gas saturation of 0.03 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.32 ISGU - End-Point Scaling of Grid Cell Maximum Gas Saturation (Imbibition)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

ISGU defines the <u>imbibition</u> maximum gas saturation for all the cells in the model via an array when the endpoint scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section and <u>the hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section.</u> The maximum gas saturation is defined as the maximum gas saturation in a two-phase gas relative permeability table.

The keyword can be used for all grid types.

No.	Name	Description						
		Field	Metric	Laboratory				
I	ISGU	ISGU is an array of real numbers assigning the maximum gas saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.70						
		dimensionless	dimensionless	dimensionless	permeability table.			

Notes:

- Note this the non-direction dependent version of the maximum gas saturation array used with the end-point scaling option. If directional end-point scaling has been activated then the ISGUX± , ISGUX± and ISGU± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.21: ISGU Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISGUX, ISGUY and ISGUZ instead of ISGU. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISGUX, ISGUY-, ISGUY-, ISGUZ and ISGUZ-, instead of the ISGU keyword.

Example

```
--
-- DEFINE GRID BLOCK END-POINT ISGU DATA FOR ALL CELLS (NX x NY x NZ = 300)
--
ISGU
300*0.700
```

The above example defines a constant connate gas saturation of 0.70 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.33 ISOGCR - End-Point Scaling of Grid Cell Critical Oil Saturation to Gas (Imbibition)

RUNS	PEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
------	-----	------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

ISOGCR defines the <u>imbibition</u> critical oil saturation for all the cells in the model via an array when the endpoint scaling option has been invoked via the ENDSCALE in the RUNSPEC section and <u>the hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section</u>. The critical oil saturation with respect to gas is defined as the maximum oil saturation for which the oil relative permeability is zero in a two-phase gas-oil relative permeability table.

The keyword can be used for all grid types.

No.	Name		Description						
		Field	Field Metric Laboratory						
I	ISOGCR	with respect to gas values should correspond to the keyword.	SOGCR is an array of real numbers assigning the critical oil saturation with respect to gas values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS seyword.						
		dimensionless	dimensionless dimensionless dimensionless						

Notes:

- Note this the non-direction dependent version of the critical gas saturation used with the end-point scaling option. If directional end-point scaling has been activated then the ISOGCRX±, ISOGCRX± and ISOGCRX± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.22: ISOGCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISOGCRX, ISOGCRY and ISOGCRZ instead of ISOGCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISOGCRX, ISOGCRX-, ISOGCRY-, ISOGCRZ and ISOGCRZ-, instead of the ISOGCR keyword.

Example

```
-- DEFINE GRID BLOCK END-POINT ISOGCR DATA FOR ALL CELLS (NX x NY x NZ = 300)
-- ISOGCR
300*0.200
```

The above example defines a constant critical gas saturation of 0.20 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.34 ISOWCR - END-POINT SCALING OF GRID CELL CRITICAL OIL SATURATION TO WATER (IMBIB.)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

ISOWCR defines the imbibition critical oil saturation for all the cells in the model via an array when the endpoint scaling option has been invoked via the ENDSCALE in the RUNSPEC section and the hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section. The critical oil saturation with respect to water is defined as the maximum oil saturation for which the oil relative permeability is zero in a two-phase oil-water relative permeability table.

The keyword can be used for all grid types.

No.	Name		Description					
		Field	Field Metric Laboratory					
I	ISOWCR	with respect to water v	real numbers assigning the alues to each cell in the nd to the NX x NY x led, for example 30*0.30	model. The number of	Taken from cell allocated relative permeability table.			
		dimensionless	dimensionless	dimensionless				

Notes:

- Note this the non-direction dependent version of the critical gas saturation used with the end-point scaling option. If directional end-point scaling has been activated then the ISOWCRX±, ISOWCRX± and ISOWCRX± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability 2) table.
- The keyword is terminated by "/".

Table 8.23: ISOGCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISOWCRX, ISOWCRY and ISOWCRZ instead of ISOWCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISOWCRX, ISOWCRX-, ISOWCRY, ISOWCRY-, ISOWCRZ and ISOWCRZ-, instead of the ISOWCR keyword.

Example

```
DEFINE GRID BLOCK END-POINT ISOWCR DATA FOR ALL CELLS (NX x NY x NZ = 300)
ISOWCR
  300*0.200
```

The above example defines a constant critical gas saturation of 0.20 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section

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8.3.35 ISWCR - End-Point Scaling of Grid Cell Critical Water Saturation (Imbibition)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

ISWCR defines the <u>imbibition</u> critical water saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE in the RUNSPEC section and <u>the hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section</u>. The critical water saturation is defined as the maximum water saturation for which the water relative permeability is zero in a two-phase relative permeability table.

The keyword can be used for all grid types.

No.	Name		Description			
		Field	Metric	Laboratory		
I	ISWCR is an array of real numbers assigning the critical water saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.20					
		dimensionless	dimensionless	dimensionless	permeability table.	

Notes:

- Note this the non-direction dependent version of the critical water saturation used with the end-point scaling option. If directional end-point scaling has been activated then the ISWCRX±, ISWCRX± and ISWCRX± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.24: ISWCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISWCRX, ISWCRY and ISWCRZ instead of ISWCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISWCRX, ISWCRX-, ISWCRY, ISWCRY, ISWCRY, ISWCRZ-, instead of the ISWCR keyword.

Example

```
--
-- DEFINE GRID BLOCK END-POINT ISWCR DATA FOR ALL CELLS (NX x NY x NZ = 300)
--
ISWCR
300*0.200
```

The above example defines a constant critical water saturation of 0.20 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.36 ISWL - End-Point Scaling of Grid Cell Connate Water Saturation (Imbibition)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

ISWL defines the <u>imbibition</u> connate water saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE in the RUNSPEC section and <u>the hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section</u>. The connate water saturation is defined as the minimum water saturation in a two-phase water relative permeability table.

The keyword can be used for all grid types.

No.	Name		Description		Default
		Field	Metric	Laboratory	-
1	ISWL	values to each cell in the	numbers assigning the comodel. The number of enterameters on the DIMENS ed, for example 30*0.15	tries should correspond	Taken from cell allocated relative permeability
		dimensionless	dimensionless	dimensionless	table.

Notes:

- Note this the non-direction dependent version of the connate gas saturation array used with the end-point scaling option. If directional end-point scaling has been activated then the ISWLX±, ISWLX± and SWZ± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.25: ISWL Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISWLX, ISWLY and ISWLZ instead of ISWL. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISWLX, ISWLX-, ISWLY, ISWLY-, ISWLZ and ISWLZ-, instead of the ISWL keyword.

Example

```
-- DEFINE GRID BLOCK END-POINT ISWL DATA FOR ALL CELLS (NX x NY x NZ = 300)
-- ISWL 300*0.150
```

The above example defines a constant connate water saturation of 0.15 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.37 ISWU - END-POINT SCALING OF GRID CELL MAXIMUM WATER SATURATION (IMBIBITION)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

ISWU defines the <u>imbibition</u> maximum water saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE in the RUNSPEC section and <u>the hysteresis</u> model option has been activated on the SATOPTS keyword in the RUNSPEC section. The maximum water saturation is defined as the maximum water saturation in a two-phase water relative permeability table.

The keyword can be used for all grid types.

No.	Name		Description		Default
		Field	Metric	Laboratory	
1	ISWU	values to each cell in the	numbers assigning the man model. The number of en arameters on the DIMENS ed, for example 30*0.70	tries should correspond	Taken from cell allocated relative permeability
		dimensionless	dimensionless	dimensionless	table.

Notes:

- Note this the non-direction dependent version of the maximum water saturation array used with the endpoint scaling option. If directional end-point scaling has been activated then the ISWUX±, ISWUX± and ISWU± series of keyword should be used.
- 2) If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.26: ISWU Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the ISWL, ISWCR, ISWU, ISGL, ISGCR, ISGU, ISOWCR, and ISOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is ISWUX, ISWUY and ISWUZ instead of ISWU. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is ISWUX, ISWUX-, ISWUY-, ISWUY-, ISWUZ and ISWUZ-, instead of the ISWU keyword.

Example

```
-- DEFINE GRID BLOCK END-POINT ISWU DATA FOR ALL CELLS (NX x NY x NZ = 300)
-- ISWU 300*0.700
```

The above example defines a constant connate gas saturation of 0.70 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.38 KRG - End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

KRG defines the scaling parameter at the maximum drainage gas relative permeability value (SGU), normally SGU is equal to $1.0 - S_{wc}$, for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The SCALCERS keyword in the PROPS section defines the options used in the re-scaling process, the options are two point scaling and three point scaling.

No.	Name		Description		Default
		Field	Metric	Laboratory	
1	KRG	less than or equal to 1.0, cell in the model.	ve real numbers which ar that are the assigned scali ed, for example 50*0.400.	ing KRG values for each	Taken from cell allocated relative permeability
		dimensionless	dimensionless	dimensionless	table.

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 8.27: KRG Keyword Description

For the two point scaling option and for the KRGR gas relative permeability array NOT present in the input deck the k_{rg} value for a grid block is scaled by:

$$k_{rg} = k_{rg_{TABLE}} \left(\frac{KRG}{k_{rg_{TABLE-MAX}}} \right)$$
 (8.3)

Where:

 k_{rg} = the resulting k_{rg} value for a grid cell.

KRG = the scaling gas relative permeability value from the KRG array for a given

cell.

 $k_{rg_{\tiny TABLE}}$ = the gas relative permeability from a grid block's gas-oil table at the grid

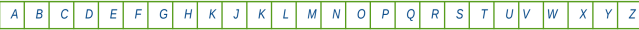
blocks gas saturation.

 $k_{rg_{TABLE-MAX}}$ = the maximum gas relative permeability from a grid block's gas-oil table, that is at the connate water saturation (S_{wc}).

If the KRGR keyword is present in the input deck then the scaling matches the relative permeability at the critical saturation of the displacing phase.

If three point scaling option has been selected via the SCALECRS keyword in the PROPS section the critical displacing phase is defined as:

No	Phases Present	Critical Saturation
I	Gas-Oil	S _{critical} = 1.0 – SOGCR - SWL
2	Gas-Oil-Water	S _{critical} = 1.0 – SOGCR - SWL



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No	Phases Present	Critical Saturation
3	Gas-Water	S critical = 1.0 - SWCR

Revision: Rev-I

Table 8.28: Critical Displacement Relationships

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY, SWUY-, SWUZ and SWUZ-, instead of the SWU keyword.

End-point scaling also allows the entered relative permeability functions to be scale on the relative permeability values using the KRG, KRGR, KRO, KRORG, KRORW, KRW and KRWR relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is KRGX, KRGY and KRGZ instead of KRG, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is KRGX, KRGX-, KRGY, KRGY-, KRGZ and KRGZ-, instead of the KRG keyword.

If hysteresis hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section, then the equivalent imbibition arrays suffixed with the letter I, for example IKRG, can be used to define the KRG for the relative permeability imbibition tables.

Examples

The first example defines an input box for the whole grid and for layers one to three, for layer one KRG is set equal to 0.550, for layer two KRG equals 0.575, and for layer three KRG equals 0.600.

```
- -
         DEFINE INPUT BOX FOR EDITING INPUT ARRAYS (NX=100, NY=100)
- -
         ----- BOX -----
         I1 I2
                 J1 J2
                          K1 K2
B<sub>0</sub>X
                  1*
                      1*
             1*
         1*
                                                            / DEFINE BOX AREA
                           1
                               3
         SET KRG VALUES FOR THREE LAYERS IN THE MODEL
- -
KRG
1000*0.555 1000*0.575 1000.0.600
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
ENDBOX
The next example does exactly the same thing using the EQUALS keyword instead.
-- -- ARRAY
               CONSTANT --
                            ----- BOX -----
                            I1 I2 J1 J2
                                              K1 K2
EQUALS
                                         1*
   'KRG'
               0.5550
                                                   1 / KRG FOR LAYER 1
                                               1
                                1*
                                     1*
                                         1*
   'KRG'
               0.5750
                                               2
                                                   2 / KRG FOR LAYER 2
   'KRG'
                                1*
                                         1*
                                               3
                                                   3 / KRG FOR LAYER 3
               0.6000
```

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8.3.39 KRGR - End-Point Scaling of Grid Cell Krgr (1-Soggr) (Drainage)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

KRGR defines the scaling parameter at the relative permeability of gas at residual oil saturation (I -SOGCR), or critical water saturation in a gas-water run (S_{wc}), for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The SCALCERS keyword in the PROPS section defines the options used in the re-scaling process, the options are two point scaling and three point scaling.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	KRGR	and less than or equal to		ed scaling KRGR values	Taken from cell allocated relative permeability table.
		dimensionless	dimensionless	dimensionless	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/".

Table 8.29: KRGR Keyword Description

When the KRGR keyword is present in the input deck then the scaling matches the relative permeability at the critical saturation of the displacing phase (oil or water).

If three point scaling option has been selected via the SCALECRS keyword in the PROPS section the critical displacing phase is defined as:

No	Phases Present	Critical Saturation
I	Gas-Oil	S _{critical} = 1.0 – SOGCR - SWL
2	Gas-Oil-Water	S _{critical} = 1.0 – SOGCR - SWL
3	Gas-Water	S _{critical} = I.0 – SWCR

Table 8.30: Critical Displacement Relationships

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY, SWUY-, SWUZ and SWUZ-, instead of the SWU keyword.

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End-point scaling also allows the entered relative permeability functions to be scale on the relative permeability values using the KRG, KRGR, KRO, KRORG, KRORW, KRW and KRWR relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is KRGRX, KRGRY and KRGRZ instead of KRGR, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is KRGRX, KRGRY, KRGRY, KRGRY, KRGRZ and KRGRZ-, instead of the KRGR keyword.

Revision: Rev-I

If hysteresis hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section, then the equivalent imbibition arrays suffixed with the letter I, for example IKRGR, can be used to define the KRG for the relative permeability imbibition tables.

Examples

The first example defines an input box for the whole grid and for layers one to three, for layer one KRRG is set equal to 0.500, for layer two KRGR equals 0.570, and for layer three KRGR equals 0.580.

```
_ _
        DEFINE INPUT BOX FOR EDITING INPUT ARRAYS (NX=100, NY=100)
- -
         ----- BOX -----
        I1 I2
                J1 J2
                        K1 K2
BOX
                     1*
                                                          / DEFINE BOX AREA
        SET KRGR VALUES FOR THREE LAYERS IN THE MODEL
- -
KRGR
1000*0.500 1000*0.570 1000.0.580
        DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
ENDBOX
```

The next example does exactly the same thing using the EQUALS keyword instead.

```
CONSTANT --
                          ----- BOX -----
-- -- ARRAY
                         I1 I2 J1 J2 K1 K2
EQUALS
                         1*
                                  1*
   'KRGR'
             0.5500
                             1*
                                     1*
                                          1
                                              1 / KRGR FOR LAYER 1
                                  1*
                                              2 / KRGR FOR LAYER 2
                          1*
                            1*
   'KRGR'
             0.5700
                                     1*
                                          2
                         1*
                                  1* 1*
                                              3 / KRGR FOR LAYER 3
                            1*
   'KRGR'
             0.5800
                                          3
```

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8.3.40 KRO - End-Point Scaling of Grid Cell Kro(Swl) (Drainage)

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

KRO defines the scaling parameter for the drainage oil relative permeability value at the connate water saturation (SWL), for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The SCALCERS keyword in the PROPS section defines the options used in the re-scaling process, the options are two point scaling and three point scaling.

No.	Name	Description			Default		
		Field	Field Metric Laboratory				
I	KRO	less than or equal to 1.0, cell in the model.	Repeat counts may be used for example 50*0.500.				
		dimensionless dimensionless dimensionless					

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 8.31: KRO Keyword Description

For the two point scaling option and for the KRORW or KRORG oil relative permeability arrays NOT being present in the input deck the k_{ro} value for a grid block is scaled by:

$$k_{ro} = k_{ro_{TABLE}} \left(\frac{KRO}{k_{ro_{TABLE}-MAX}} \right)$$
 (8.4)

Where:

 k_{ro} = the resulting k_{ro} value for a grid cell.

KRO = the scaling oil relative permeability value from the KRO array for a given

cell.

 $k_{ro_{TABLE}}$ = the oil relative permeability from a grid block's oil relative permeability

table at the grid blocks oil saturation.

 $k_{\scriptscriptstyle TO_{\scriptscriptstyle TABLE-MAX}}$ = the maximum oil relative permeability from a grid block's oil relative table,

that is at the critical water saturation (S_{wcr}).

If the KRORW or KRORG keywords are present in the input deck then the scaling matches the relative permeability at the critical saturation of the displacing phase.

If three point scaling option has been selected via the SCALECRS keyword in the PROPS section the critical displacing phase is defined as:

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No	Keywords Present	Critical Saturation
I	KRORW	S critical = 1.0 - SWCR - SGL
2	KRORG	S _{critical} = I.0 – SGCR - SWL

Revision: Rev-I

Table 8.32: Critical Displacement Relationships

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY-, SWUY-, SWUZ and SWUZ-, instead of the SWU keyword.

End-point scaling also allows the entered relative permeability functions to be scale on the relative permeability values using the KRG, KRGR, KRO, KRORG, KRORW, KRW and KRWR relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is KROX, KROY and KROZ instead of KRO, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is KROX, KROY-, KROY-, KROZ and KROZ-, instead of the KRO keyword.

If hysteresis hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section, then the equivalent imbibition arrays suffixed with the letter I, for example IKRO, can be used to define the KRO for the relative permeability imbibition tables.

Examples

The first example defines an input box for the whole grid and for layers one to three, for layer one KRO is set equal to 0.850, for layer two KRO equals 0.875, and for layer three KRO equals 0.900.

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS (NX=100, NY=100)
- -
          ----- BOX -----
         I1 I2 J1 J2
                         K1 K2
B<sub>0</sub>X
                  1* 1*
                                                            / DEFINE BOX AREA
         SET KRO VALUES FOR THREE LAYERS IN THE MODEL
KR0
1000*0.855 1000*0.875 1000.0.900
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
- -
The next example does exactly the same thing using the EQUALS keyword instead.
               CONSTANT --
                            ----- BOX -----
-- -- ARRAY
                            I1 I2 J1 J2
                                             K1 K2
EQUALS
                                     1*
                            1*
                                1*
                                         1*
   'KR0'
                                                   1 / KRO FOR LAYER 1
               0.8550
                                               1
   'KOG'
               0.8750
                            1*
                                1*
                                     1*
                                         1*
                                                  2 / KRO FOR LAYER 2
                                               2
                                         1*
   'KR0'
               0.9000
                                                   3 / KRO FOR LAYER 3
```

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8.3.41 KRORG - End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

KRORG defines the scaling parameter the drainage relative permeability of oil at the critical gas saturation (SGCR), for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The SCALCERS keyword in the PROPS section defines the options used in the re-scaling process, the options are two point scaling and three point scaling.

No.	Name		Description			
		Field	Field Metric Laboratory			
I	KRORG	and less than or equal to	KRORG is an array of positive real numbers which are greater than zero and less than or equal to 1.0, that are the assigned scaling KRORG values for each cell in the model. Repeat counts may be used, for example 50*0.850.			
	dimensionless dimensionless dimensionless				permeability table.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 8.33: KRORG Keyword Description

When the KRORG keyword is present in the input deck then the scaling matches the relative permeability at the critical saturation of the displacing phase.

If three point scaling option has been selected via the SCALECRS keyword in the PROPS section the critical displacing phase is defined as:

No	Keywords Present	Critical Saturation
I	KRORW	S _{critical} = I.0 – SWCR - SGL
2	KRORG	S _{critical} = 1.0 – SGCR - SWL

Table 8.34: Critical Displacement Relationships

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY, SWUY-, SWUZ and SWUZ-, instead of the SWU keyword.

End-point scaling also allows the entered relative permeability functions to be scale on the relative permeability values using the KRG, KRGR, KRORG, KRORGRG, KRORGRW, KRW and KRWR relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be

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directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is KRORGX, KRORGY and KRORGZ instead of KRORG, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the nonreversible versions of the aforementioned arrays should be used, that is KRORGX, KRORGX-, KRORGY, KRORGY-, KRORGZ and KRORGZ-, instead of the KRORG keyword.

Revision: Rev-I

If hysteresis hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section, then the equivalent imbibition arrays suffixed with the letter I, for example IKRORG, can be used to define the KRORG for the relative permeability imbibition tables.

Examples

The first example defines an input box for the whole grid and for layers one to three, for layer one KRORG is set equal to 0.750, for layer two KRORG equals 0.775, and for layer three KRORG equals 0.800.

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS (NX=100, NY=100)
- -
- -
          ----- BOX -----
         I1 I2
                 J1 J2
                           K1 K2
B<sub>0</sub>X
         1*
             1*
                       1*
                   1*
                                                               / DEFINE BOX AREA
                             1
                                 3
- -
         SET KRORG VALUES FOR THREE LAYERS IN THE MODEL
KRORG
1000*0.755 1000*0.775 1000.0.800
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
ENDBOX
The next example does exactly the same thing using the EQUALS keyword instead.
```

```
-- -- ARRAY
             CONSTANT --
                          ----- BOX -----
                         I1 I2 J1 J2 K1 K2
EQUALS
                         1*
                             1*
                                  1*
                                      1*
   'KRORG'
             0.7550
                                          1
                                              1 / KRORG FOR LAYER 1
                                  1*
                         1*
                                              2 / KRORG FOR LAYER 2
   'KRORG'
                            1*
                                     1*
             0.7750
                                          2
                         1*
                                  1* 1*
                                              3 / KRORG FOR LAYER 3
                            1*
   'KRORG
             0.8000
                                          3
```

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8.3.42 KRORW - End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

KRORW defines the scaling parameter the drainage relative permeability of oil at the critical water saturation (SWCR), for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The SCALECRS keyword in the PROPS section defines the options used in the re-scaling process, the options are two point scaling and three point scaling.

No.	Name	Description			Default	
		Field Metric Laboratory				
I	KRORW	and less than or equal to for each cell in the mode	KRORW is an array of positive real numbers which are greater than zero and less than or equal to 1.0, that are the assigned scaling KRORW values for each cell in the model. Repeat counts may be used, for example 50*0.850			
		dimensionless	dimensionless	dimensionless	permeability table.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 8.35: KRORW Keyword Description

When the KRORW keyword is present in the input deck then the scaling matches the relative permeability at the critical saturation of the displacing phase.

If three point scaling option has been selected via the SCALECRS keyword in the PROPS section the critical displacing phase is defined as:

No	Keywords Present	Critical Saturation
I	KRORW	S _{critical} = I.0 – SWCR - SGL
2	KRORG	S _{critical} = 1.0 – SGCR - SWL

Table 8.36: Critical Displacement Relationships

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY-, SWUY-, SWUZ and SWUZ-, instead of the SWU keyword.

End-point scaling also allows the entered relative permeability functions to be scale on the relative permeability values using the KRG, KRGR, KRORW, KRORWRG, KRORWRW, KRW and KRWR relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be

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directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is KRORWX, KRORWY and KRORWZ instead of KRORW, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is KRORWX, KRORWX-, KRORWY-, KRORWY-, KRORWZ and KRORWZ-, instead of the KRORW keyword.

Revision: Rev-I

If hysteresis hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section, then the equivalent imbibition arrays suffixed with the letter I, for example IKRORW, can be used to define the KRORW for the relative permeability imbibition tables.

Examples

The first example defines an input box for the whole grid and for layers one to three, for layer one KRORW is set equal to 0.750, for layer two KRORW equals 0.775, and for layer three KRORW equals 0.800.

```
DEFINE INPUT BOX FOR EDITING INPUT ARRAYS (NX=100, NY=100)
- -
- -
         ----- BOX -----
         I1 I2
                J1 J2
                          K1 K2
B<sub>0</sub>X
         1*
             1*
                     1*
                  1*
                                                             / DEFINE BOX AREA
                           1
                                3
- -
         SET KRORW VALUES FOR THREE LAYERS IN THE MODEL
KRORW
1000*0.755 1000*0.775 1000.0.800
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
ENDBOX
The next example does exactly the same thing using the EQUALS keyword instead.
-- -- ARRAY
               CONSTANT --
                             ----- BOX -----
                            I1 I2 J1 J2 K1 K2
```

```
EQUALS
                           1*
                               1*
                                    1*
                                        1*
   'KRORW'
              0.7550
                                             1
                                                 1 / KRORW FOR LAYER 1
                                    1*
                           1*
   'KRORW'
                              1*
                                       1*
                                                 2 / KRORW FOR LAYER 2
              0.7750
                                             2
                           1*
                                    1* 1*
                                                 3 / KRORW FOR LAYER 3
                              1*
   'KRORW
              0.8000
                                             3
```

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8.3.43 KRW - End-Point Scaling of Grid Cell Krw(Sw = 1.0) (Drainage)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

KRW defines the scaling parameter at the maximum drainage oil relative permeability value (SWU), that is for S_w = 1.0, for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The SCALCERS keyword in the PROPS section defines the options used in the re-scaling process, the options are two point scaling and three point scaling.

No.	Name	Name Description Field Metric Laboratory			Default		
1	KRW	less than or equal to 1.9 each cell in the model.	Repeat counts may be used for example 50*1,000.				
	dimensionless dimensionless dimensionless				permeability table.		

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/".

Table 8.37: KRW Keyword Description

For the two point scaling option and for the KRWR water relative permeability array NOT present in the input deck the k_{rw} value for a grid block is scaled by:

$$k_{rw} = k_{rw_{TABLE}} \left(\frac{KRW}{k_{rw_{TABLE}-MAY}} \right)$$
 (8.5)

Where:

= the resulting KRW value for a grid cell.

= the scaling water relative permeability value from the KRW array for a given cell.

= the water relative permeability from a grid block's oil relative permeability

table at the grid blocks water saturation.

= the maximum water relative permeability from a grid block's water

relative table, that is at the maximum water saturation.

If the KRWR keyword is present in the input deck then the scaling matches the relative permeability at the critical saturation of the displacing phase.

If three point scaling option has been selected via the SCALECRS keyword in the PROPS section the critical displacing phase is defined as:

No	Phases Present	Critical Saturation
1	Gas-Oil	S critical = I.0 - SOWCR - SGL



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No	Phases Present	Critical Saturation
2	Gas-Oil-Water	S critical = 1.0 - SOWCR - SGL
3	Gas-Water	S _{critical} = I.0 – SGCR

Revision: Rev-I

Table 8.38: Critical Displacement Relationships

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY-, SWUY-, SWUZ and SWUZ-, instead of the SWU keyword.

End-point scaling also allows the entered relative permeability functions to be scale on the relative permeability values using the KRG, KRGR, KRW, KRORG, KRORW, KRW and KRWR relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is KRWX, KRWY and KRWZ instead of KRW, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is KRWX, KRWX-, KRWY, KRWY-, KRWZ and KRWZ-, instead of the KRW keyword.

If hysteresis hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section, then the equivalent imbibition arrays suffixed with the letter I, for example IKRW, can be used to define the KRW for the relative permeability imbibition tables.

Examples

The first example defines an input box for the whole grid and for layers one to three, for layer one KRW is set equal to 0.850, for layer two KRW equals 0.875, and for layer three KRW equals 0.900.

The next example does exactly the same thing using the EQUALS keyword instead.

```
-- -- ARRAY
               CONSTANT --
                            ----- BOX -----
                            I1
                               12
                                     J1
                                         J2
                                              K1
                                                  K2
EQUALS
   'KRW'
               0.8550
                            1*
                                1*
                                     1*
                                         1*
                                                     / KRW FOR LAYER 1
                                              1
                                                  1
                                         1*
                                1*
                                     1*
   'KRW'
               0.8750
                                              2
                                                  2 / KRW FOR LAYER 2
                                1*
   'KRW'
                                     1*
                                         1*
               0.9000
                                              3
                                                  3 / KRW FOR LAYER 3
```

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8.3.44 KRWR - End-Point Scaling of Grid Cell KRWR(Sw = 1.0) (Drainage)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

KRWR defines the scaling parameter at the maximum drainage oil relative permeability value (SWU), that is for $S_w = 1.0$, for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The SCALCERS keyword in the PROPS section defines the options used in the re-scaling process, the options are two point scaling and three point scaling.

No.	Name		Description				
		Field	Metric	Laboratory			
1	KRWR	and less than or equal to	sitive real numbers which o I.0, that are the assigne I. ed, for example 50*1.000.	ed scaling KRWR values	Taken from cell allocated relative permeability		
		dimensionless	table.				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 8.39: KRWR Keyword Description

When the KRWR keyword is present in the input deck then the scaling matches the relative permeability at the critical saturation of the displacing phase.

If three point scaling option has been selected via the SCALECRS keyword in the PROPS section the critical displacing phase is defined as:

No	Phases Present	Critical Saturation		
I	Gas-Oil	S critical = I.0 - SOWCR - SGL		
2	Gas-Oil-Water	S critical = I.0 - SOWCR - SGL		
3	Gas-Water	S critical = I.0 – SGCR		

Table 8.40: Critical Displacement Relationships

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY-, SWUY-, SWUZ and SWUZ-, instead of the SWU keyword.

End-point scaling also allows the entered relative permeability functions to be scale on the relative permeability values using the KRG, KRGR, KRWR, KRORG, KRORW, KRWR and KRWRR relative

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permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is KRWRX, KRWRY and KRWRZ instead of KRWR, There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is KRWRX, KRWRX-, KRWRY, KRWRY-, KRWRZ-, instead of the KRWR keyword.

Revision: Rev-I

If hysteresis hysteresis model option has been activated on the SATOPTS keyword in the RUNSPEC section, then the equivalent imbibition arrays suffixed with the letter I, for example IKRWR, can be used to define the KRWR for the relative permeability imbibition tables.

Examples

The first example defines an input box for the whole grid and for layers one to three, for layer one KRWR is set equal to 0.750, for layer two KRWR equals 0.775, and for layer three KRWR equals 0.800.

```
- -
         DEFINE INPUT BOX FOR EDITING INPUT ARRAYS (NX=100, NY=100)
- -
         ----- BOX -----
         I1 I2 J1 J2 K1 K2
B<sub>0</sub>X
            1*
                 1* 1*
                           1
                               3
                                                           / DEFINE BOX AREA
         SET KRWR VALUES FOR THREE LAYERS IN THE MODEL
KRWR
1000*0.755 1000*0.775 1000.0.800
         DEFINE END OF INPUT BOX EDITING OF INPUT ARRAYS
ENDBOX
```

The next example does exactly the same thing using the EQUALS keyword instead.

```
-- -- ARRAY
             CONSTANT --
                          ----- BOX -----
- -
                         I1 I2 J1 J2 K1 K2
EQUALS
                         1*
                                  1*
                                     1*
   'KRWR'
             0.7550
                             1*
                                          1
                                              1 / KRWR FOR LAYER 1
                                  1* 1*
                         1* 1*
   'KRWR'
                                              2 / KRWR FOR LAYER 2
             0.7750
                                          2
                         1* 1*
                                 1*
                                     1*
                                         3 3 / KRWR FOR LAYER 3
   'KRWR'
             0.8000
```

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8.3.45 MAXVALUE - Sets a Maximum Value for an Array Element

The MAXVALUE keyword sets a maximum value for the specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the MAXVALUE keyword is being used.

Revision: Rev-I

See Error: Reference source not found in the GRID section for a full description.

8.3.46 MINVALUE - SET A MINIMUM VALUE FOR AN ARRAY ELEMENT

The MINVALUE keyword sets a minimum value for the specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the MINVALUE keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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8.3.47 MISC - Define Solvent Miscibility-Immiscibility Transform Functions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

MISC defines the transformation between the miscible and immiscible relative permeability models, for when the MISCIBLE and SOLVENT keywords in the RUNSPEC section has been activated. The keyword can only be used with the MISCIBLE option and for when the oil, water, gas and solvent phases are active in the model.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	SSOL	values starting from zero	A columnar vector of real monotonically increasing down the column values starting from zero and terminating at one, that defines the solvent fraction with respect to the solvent and gas saturation, and is defined by:				
		Where Sg is the gas satur					
		Note that the first entry last entry should be one					
		dimensionless	dimensionless	dimensionless			
2	MISC	A columnar vector of rethat are greater than or corresponding miscibility	None				
		The first entry in the co should be one to fully de					
		dimensionless	dimensionless	dimensionless			

Notes:

- 1) The keyword is followed by NTMISC tables as declared on the MISCIBLE keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSMISC rows as declared on the MISCIBLE keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.41: MISC Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Example	e						
 	SOLVENT	MISCIBILITY-	IMMISCIBLITY TRA	ANSFORM TABLE			
SGCWMIS							
	SS0L	MISC					
	FRAC	FRAC					
	0.0000	0.0000					
	0.2000	0.2500					
	0.5000	0.7500					
	1.0000	1.0000			/ TABLE	NO. 01	
	SS0L	MISC					
	FRAC	FRAC					
	0.0000	0.0000					
	0.3000	0.2500					
	0.6000	1.0000					
	1.0000	1.0000			/ TABLE	NO. 02	

The above example defines two solvent miscible-immiscible transform tables assuming NTMISC equals two and NSMISC is greater than or equal to four on the MISCIBLE keyword in the RUNSPEC section.

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8.3.48 MSFN - MISCIBLE NORMALIZED RELATIVE PERMEABILITY TABLES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The MSFN keyword defines the miscible <u>normalized</u> relative permeability tables for when the MISCIBLE and or SOLVENT options have been activated in the RUNSPEC section using the respective keyword. The MISCIBLE keyword invokes a three component formulation (oil, water and solvent gas or an oil, water and solvent oil). Why the SOLVENT keyword results in a four component model (oil, water and gas plus a solvent). This keyword should only be used if the MISCIBLE and or SOLVENT options have been activated.

No.	Name	Description	Default
I	SGAS	A columnar vector of real monotonically increasing down the column values starting from zero and terminating at one, that defines the gas plus solvent saturation.	None
2	KRSG	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the gas plus solvent relative permeability multiplier.	None
3	KRO	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the oil relative permeability multiplier.	None

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/

Table 8.42: MSFN Keyword Description

Examples

	MISCIBLE	NORMALIZED	RELATIVE PERMEABILITY TABLES	
MSFN				
	SGAS	KRSG	KR0	
	FRAC			
	0.0000	0.0000	1.0000	
	1.0000	1.0000	0.0000	/ TABLE NO. 01
	SGAS	KRSG	KR0	
	FRAC			
	0.0000	0.0000	1.0000	
	0.2000	0.2000	0.8000	
	0.4000	0.3000	0.7000	
	0.6000	0.4000	0.6000	
	0.8000	0.5000	0.4000	
	1.0000	1.0000	0.0000	/ TABLE NO. 02

The above example defines two MSN tables for use the MISCIBLE and SOLVENT options.

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8.3.49 MULTIPLY - MULTIPLY A SPECIFIED ARRAY BY A CONSTANT

The MULTIPLY keyword multiplies a specified array or part of an array by a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the keyword is being used.

Revision: Rev-I

See Error: Reference source not found in the GRID section for a full description.

8.3.50 MULTIREG - MULTIPLY AN ARRAY BY A CONSTANT BASED ON A REGION NUMBER

The MULTIREG keyword multiplies an array or part of an array by a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTIREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the MULTIREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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8.3.51 OILDENT - DEFINE OIL DENSITY TEMPERATURE COEFFICIENTS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

OILDENT defines the oil density as a function of temperature coefficients for when OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC. The reference pressure for this table is given by the VISCREF keyword in the PROPS section. Note this is an OPM Flow keyword used with OPM Flow's "black-oil" thermal model that is not available in the commercial simulator's "black-oil" thermal formulation.

This keyword can only be used if OPM Flow's thermal thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	TEMP	reference temperature	value greater than zero the used with TEXP1 and To respect to temperature.		Defined		
		°R	°K	°K			
		527.67	293.15	293.15			
2	TEXPI		TEXPI is a real positive value greater than zero that defines the oil thermal expansion coefficient of the first order.				
		I/°R	I/°K	I/°K			
		1.67 x 10 ⁻⁴	3.0 x 10 ⁻⁴	3.0 x 10 ⁻⁴			
3	TEXP2	·	TEXP2 is a real positive value greater than zero that defines the oil thermal expansion coefficient of the second order.				
		I/°R²	I/°K²	I/°K²			
		9.26 x 10 ⁻⁷	3.0 x 10 ⁻⁶	3.0 x 10 ⁻⁶			

Notes:

- 1) The keyword is followed by NTPVT records as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each data set is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.43: OILDENT Keyword Description

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Example

The following example shows the OILDENT keyword using the default values, for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC section and for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to two.

Revision: Rev-I

```
-- OIL DENSITY TEMPERATURE COEFFICIENTS (OPM FLOW THERMAL KEYWORD)
-- OIL DENSITY DENSITY
-- TEMP COEFF1 COEFF2
-- OILDENT

1* 1* 1* 1* / TABLE NO. 01
1* 1* 1* 1* / TABLE NO. 02
```

There is no terminating "/" for this keyword.

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8.3.52 OILVISCT - Define Oil Viscosity versus Temperature Functions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

OILVISCT defines the oil viscosity as a function of temperature for when OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. The reference pressure and solution gasoil ratio of the oil for this table is given by the VISCREF keyword in the PROPS section. Note this is an OPM Flow keyword used with OPM Flow's "black-oil" thermal model that is not available in the commercial simulator's "black-oil" thermal formulation.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Default			
		Field	Metric	Laboratory		
I	TEMP	A columnar vector of r	None			
		°F	°C	°C		
2	VIS	A columnar vector of rea	None			
		VIS should be given at th defined by PRS and RS va				
	cP cP cP					

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.44: OILVISCT Keyword Description

Example

The following example shows the OILVISCT keyword for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC section and for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set equal to one.

```
OIL VISCOSITY VERSUS TEMPERATURE TABLES (OPM FLOW EXTENSION KEYWORD)
         OIL
                    OIL
         TEMP
                    VISC
OILVISCT
           100.0
                    0.600
           110.0
                    0.650
           120.0
                    0.680
           150.0
                    0.720
           165.0
                    0.725
                                                             / TABLE NO. 01
```

There is no terminating "/" for this keyword.

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8.3.53 OPERATE - Define Mathematical Operations on Arrays

This keyword defines the mathematical operations on arrays in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

Revision: Rev-I

See OPERATE - Define Mathematical Operations on Arrays in the GRID section for a full description.

8.3.54 OPERATER - Define Mathematical Operations on Arrays by Region

This keyword defines the mathematical operations on arrays for specific regions in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

See OPERATER – Define Mathematical Operations on Arrays by Region in the GRID section for a full description.

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8.3.55 PCG - End-Point Scaling of Grid Cell Maximum Gas Capillary Pressure (Drainage)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Description

PCG defines the maximum drainage gas-oil capillary pressure values for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable end-point scaling and the use of this keyword. The keyword can be used for all grid types.

The capillary pressure for a grid block is scaled by:

$$P_c = P_{c_{TABLE}} \left(\frac{PCG}{P_{c_{TABLE-MAX}}} \right)$$
 (8.6)

Revision: Rev-I

Where:

 P_c = the resulting drainage gas-oil capillary pressure for a grid cell.

PCG = the maximum capillary pressure from the PCG array for a given cell.

 $\mathcal{O}_{C_{TABLE}}$ = the capillary pressure in the drainage capillary pressure table allocated to the grid block.

 $P_{c_{\mathit{TABLE-MAX}}}$ = the maximum capillary pressure in the drainage capillary pressure table

allocated to the grid block at $S_a = 1 - S_{wco}$.

No.	Name		Default					
		Field						
I	PCG		PCG is an array of positive real numbers assigning the maximum drainage gas-oil capillary pressure values for each cell in the model.					
		Repeat counts may be us	Repeat counts may be used, for example 30*100.0.					
		psia	bars	atm				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If the HYSTER on the SATOPTS keyword in the RUNSPEC section has been activated to invoke hysteresis then PCG scales the drainage curve and IPCG scales the imbibition curve.
- 3) The keyword is terminated by "/".

Table 8.45: PCG Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness. See also the IPCG keyword for the equivalent imbibition functionality.

Example

```
-- DEFINE GRID BLOCK PCG DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
-- PCG
100*50.0 100*75.0 100*125.0
```

The above example defines the a PCW for 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.56 PCW - END-POINT SCALING OF GRID CELL WATER CAPILLARY PRESSURE (Drainage)

RUNS	SPEC	GRID	EDIT		REGIONS	SOLUTION	SUMMARY	SCHEDULE
------	------	------	------	--	---------	----------	---------	----------

Description

PCW defines the maximum drainage water-oil or water-gas capillary pressure values for all the cells in the model via an array. The ENDSCALE keyword in the RUNSPEC section should be activated to enable endpoint scaling and the use of this keyword. The keyword can be used for all grid types. The capillary pressure for a grid block is scaled by:

$$P_c = P_{c_{TABLE}} \left(\frac{PCW}{P_{c_{TABLE-MAX}}} \right)$$
 (8.7)

Revision: Rev-I

Where:

= the resulting drainage water capillary pressure for a grid cell.

PCW = the maximum capillary pressure from the PCW array for a given cell.

 $P_{c_{\mathit{TABLE}}}$ = the capillary pressure in the drainage capillary pressure table allocated

to the grid block.

 $P_{c_{\scriptscriptstyle TARIF-MAX}}$ = the maximum capillary pressure in the drainage capillary pressure table

allocated to the grid block (that is at the connate water saturation).

No.	Name		Default					
		Field Metric Laboratory						
T.	PCW		PCW is an array of positive real numbers assigning the maximum drainage water capillary pressure values for each cell in the model.					
		Repeat counts may be us	Repeat counts may be used, for example 30*100.0.					
		psia	bars	atm				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- If the HYSTER on the SATOPTS keyword in the RUNSPEC section has been activated to invoke hysteresis then PCW scales the drainage curve and IPCW scales the imbibition curve.
- The keyword is terminated by "/".

Table 8.46: PCW Keyword Description

See also the IPCW keyword for the equivalent imbibition functionality.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

Example

```
DEFINE GRID BLOCK PCW DATA FOR ALL CELLS (BASED ON NX x NY x NZ = 300)
PCW
  100*50.0 100*75.0 100*125.0
```

The above example defines the a PCW for 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.57 PLMIXPAR - Define the Polymer Todd-Longstaff Mixing Parameters

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PLMIXPAR keyword defines the Todd-Longstaff⁴⁵ mixing parameters for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section. This keyword must be present in the input deck if the POLYMER keyword has been activated.

Note that this keyword is used only for the polymer option, if the MISCIBLE keyword in the RUNSPEC section has been invoked then in addition the TLMIXPAR keyword is also required to define the Todd-Longstaff mixing parameters for the MISCIBLE option.

No.	Name		Default						
		Field							
I	PLMVIS		A real positive value that is greater than or equal to zero and less than or equal to one, that defines the viscosity Todd-Longstaff mixing parameter for each polymer region.						
		dimensionless	None						

Notes:

- The keyword is followed by NPLMIX tables as declared on the REGDIMS keyword in the RUNSPEC section, for each rock region. There should be only one row per table.
- 2) Each entry is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.47: PLMIXPAR Keyword Description

Example

The above example defines three polymer Todd-Longstaff mixing parameter data sets, based on the NPLMIX variable on the REGDIMS keyword in the RUNSPEC section being equal to three.

Todd, M. and Longstaff, W. "The Development, Testing and Application of a Numerical Simulator for Predicting Miscible Flood Performance," paper SPE 3484, Journal of Canadian Petroleum Technology (1972) 24, No. 7, 874-882.



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8.3.58 PLYADS - Define Polymer Rock Adsorption Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PLYADS keyword defines the rock polymer adsorption tables for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section. Alternatively, the functions can be entered via the PLYADSS keyword in the PROPS section for when salt sensitivity is to be considered.

No.	Name	ne Description				
		Field	Metric	Laboratory		
I	POLCON	A columnar vector of real monotonically increasing down the column values that defines the polymer concentration in the solution surrounding the rock.				
		The first entry should be				
		lb/stb	kg/sm³	gm/scc		
2	POLRATIO	A columnar vector of real increasing down the column values that defines the mass of adsorbed polymer per unit mass of rock of the saturated concentration of polymer adsorbed by the rock.				
		The first entry should concentration.				
		lb/lb	kg/kg	gm/gm		

Notes:

- The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.48: PLYADS Keyword Description

See also the PLYADSS keyword in the PROPS section to also define rock polymer adsorption tables when the polymer concentration is a function of salinity.

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Example POLYMER ROCK ADSORPTION TABLE **PLYADS** POLYMER POLYMER - -POLCON POLRATIO _____ _____ 0.0 0.00000 2.0 0.00003 4.0 0.00005 6.0 0.00007 8.0 0.00009 10.0 0.00011 12.0 0.00012 / TABLE NO. 01 14.0 0.00015 POLYMER POLYMER POLCON POLRATIO _____ 0.0 0.00000 3.0 0.00004 5.0 0.00006 7.0 0.00008 8.0 0.00009 10.0 0.00011 / TABLE NO. 02

The above example defines two polymer rock adsorption tables assuming NTSFUN equals two and NSSFUN is greater than or equal to eight on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.59 PLYADSS - Define Polymer Rock Adsorption with Salt Dependence **T**ABLES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PLYADSS keyword defines the rock polymer adsorption tables for when the polymer and the salt options has been activated by the POLYMER and BRINE keywords in the RUNSPEC section.

Note that the BRINE option is not currently supported by OPM Flow; however, the polymer rock adsorption functions without salt dependence may be entered via the PLYADS keyword in the PROPS section, for when salt sensitivity is not to be considered.

No.	Name Description					
		Field	Metric	Laboratory		
I	POLCON	A columnar vector of real monotonically increasing down the column values that defines the polymer concentration in the solution surrounding the rock.				
		The first entry should concentration data set.	be zero to define a no	polymer and no salt		
		POLCON should onl POLCON/POLRATIO POLCON/POLRATIO ta				
		lb/stb	kg/sm³	gm/scc		
2	POLRATIO	A columnar vector of real increasing down the column values that defines the mass of adsorbed polymer per unit mass of rock of the saturated concentration of polymer adsorbed by the rock for a given POLCON and the salt concentration given by SALTCON on the ADSALNOD keyword in the PROPS section.				
		The first table data set en no salt concentration date				
		Subsequent POLRATIC combination for a given order) by the SALTCOI PROPS section.				
		Each POLCON/POLRATIO/SALT data sets should be terminated by a "/"				
		lb/lb	kg/kg	gm/gm		

Notes:

- The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section. I)
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.49: PLYADS Keyword Description

See also the PLYADSS keyword in the PROPS section to also define rock polymer adsorption tables when the polymer concentration is a function of salinity.

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```
Example
         SETS SALT CONCENTRATION FOR POLYMER SOLUTION ADSORPTION
- -
         VIA SATNUM ARRAY ALLOCATION
         SALT
ADSALNOD
         1.0
         5.0
         10.5
         25.0
                   / SATNUM TABLE NO. 01
- -
         POLYMER ROCK ADSORPTION WITH SALT DEPENDANCY TABLE
- -
PLYADS
         POLYMER
                    POLYMER
- -
         POLCON
                    POLRATIO
- -
         _____
                    _____
             0.0
                     0.00000
                     0.00000
                     0.00000
                     0.00000
                                                             / TABLE NO. 01
         POLYMER
                    POLYMER
         POLCON
                    POLRATIO
- -
         -----
                     -----
             1.0
                     0.00002
                     0.00003
                     0.00004
                                                             / TABLE NO. 02
                     0.00005
         POLYMER
                    POLYMER
         POLCON
                    POLRATIO 1
             2.0
                     0.00003
                     0.00004
                     0.00005
                                                             / TABLE NO. 03
                     0.00006
         POLYMER
                    POLYMER
         POLCON
                    POLRATIO
             3.0
                     0.00004
                     0.00005
                     0.00006
                     0.00007
                                                             / TABLE NO. 04
```

The above example defines four polymer rock adsorption tables for four salt concentration on the ADSALNOD keyword, assuming NTSFUN equals one and NSSFUN is greater than or equal to four on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.60 PLYDHFLF - Define Polymer Thermal Degradation Half-Life Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PLYDHFLF keyword defines the polymer thermal degradation half-life with respect to temperature functions for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

No.	Name		Default						
		Field Metric		Laboratory					
I	TEMP		A columnar vector of real monotonically increasing down the column values that defines the polymer thermal degradation temperature.						
		°F	°C	°C					
2	POLHFLF	A columnar vector of re half-life.	A columnar vector of real values that defines the corresponding polymer half-life.						
		days	days	hours					

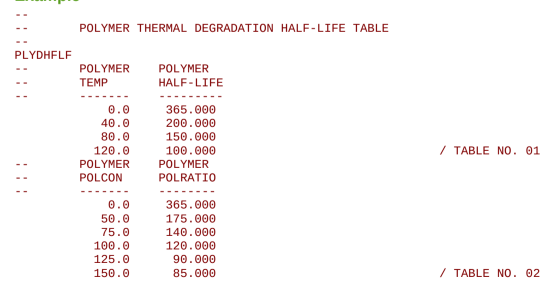
Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.50: PLYDHFLF Keyword Description

This keyword is not supported by OPM Flow but is documented here for completeness, as the thermal option is currently under development.

Example



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The example defines two polymer thermal degradation half-life tables, based on the NTPVT variable on the TABDIMS keyword in the RUNSPEC section being equal to two and NPPVT variable on the same keyword being greater than or equal to six.

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8.3.61 PLYMAX - Define Polymer-Salt Viscosity Mixing Concentrations

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PLYMAX keyword defines maximum polymer and salt concentrations that are to be used in the mixing parameter calculation of the fluid component viscosities, for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

Note that If the BRINE option has not be activated by the BRINE keyword in the RUNSPEC section, then the salt concentrations in the second column are ignored.

Currently the BRINE option is not implemented in OPM Flow and therefore SALTCON is ignored.

No.	Name		Description					
		Field	Metric	Laboratory				
I	POLCON		A real value that defines the polymer concentration in the solution which is used to calculate maximum polymer fluid component viscosity.					
		lb/stb	kg/sm³	gm/scc				
2	SALTCON		s the salt concentration in the polymer fluid compone		None			
		keyword in the RUNSPE	Note that If the BRINE option has not been activated by the BRINE keyword in the RUNSPEC section, then this variable is ignored; however, there should still be dummy entries in this case.					
		This variable is ignored as the BRINE option is not implemented in OPM Flow.						
		lb/stb	kg/sm³	gm/scc				

Notes:

- The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section, for each rock region. There should be only one row per table.
- Each table is terminated by "/" and there is no "/" terminator for the keyword. 2)

Table 8.51: PLYDMAX Keyword Description

Example

```
POLYMER-SALT VISCOSITY MIXING CONCENTRATIONS
PLYMAX
         POLYMER
                    SALT
- -
         POLCON
                    SALTCON
                                                              / TABLE NO. 01
         0.0100
                     0.0500
         0.0075
                     0.0400
                                                               TABLE NO. 02
         0.0050
                     0.0300
                                                              / TABLE NO. 03
```

The above example defines three polymer-salt viscosity mixing concentrations, based on the NPLMIX variable on the REGDIMS keyword in the RUNSPEC section being equal to three.

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8.3.62 PLYROCK - Define Polymer-Rock Properties

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PLYROCK keyword defines rock properties for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	PSPACE	A real positive value that maximum water saturati space for this rock type.					
		dimensionless	dimensionless	dimensionless	None		
2	PERMFAC		at is greater than or equ meability to the water ph een adsorbed.				
		dimensionless	dimensionless	dimensionless	None		
3	DENSITY	A real value that define conditions.					
		lb/rtb	kg/rm³	gm/rcc	None		
4	ADINDX	A positive integer of I or 2 that defines desorption option.					
		 then polymer desorption may occurs by retracing the polymer adsorption isotherm when the local polymer concentration in the solution decreases. 					
		2) then no polyme	er desorption may occurs				
		Dimensionless	Dimensionless	Dimensionless			
		I	I	I	Defined		
5	POLMAX	A real positive non-zero value that defines the maximum polymer adsorption to be used in the calculation of the resistance factor for the water phase.					
		lb/lb	kg/kg	gm/gm	None		

Notes:

- The keyword is followed by NPLMIX tables as declared on the REGDIMS keyword in the RUNSPEC section, for each polymer flooding region. There should be only one row per table.
- 2) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.52: PLYROCK Keyword Description

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1.9500

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0.1500

Revision: Rev-I

/ TABLE NO. 03

Example POLYMER-ROCK PROPERTIES PLYR0CK P0RE PERM INSITU DESORP MAX FACTOR DENSITY --SPACE OPTN P0LY --------------------1.7500 1 0.00012 2 0.00015 1 0.00014 / TABLE NO. 01 0.1200 1800.0 0.1300 1.8500 1980.0 / TABLE NO. 02

2005.0

The above example defines three polymer-rock tables, based on the NTSFUN variable on the REGDIMS keyword in the RUNSPEC section being equal to three.

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8.3.63 PLYSHEAR - ACTIVATE AND DEFINE POLYMER SHEARING PARAMETERS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PLYSHEAR keyword activates and the defines the polymer shear thinning-thickening option for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

No.	Name	Description						
		Field	Metric	Laboratory				
I	VELOCITY	values that defines the wa	A columnar vector of real monotonically increasing down the column values that defines the water-polymer flow velocity. The VELOCITY value for the first row in the table should be zero.					
		feet/day	m/day	cm/hour				
2	VISFAC	effective water and polynof the polymer occurs.	, ,					
		Normally VISFAC value for the first row in the table should be one. dimensionless dimensionless dimensionless						

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.53: PLYSHEAR Keyword Description

Example

ACTIVATE AND DEFINE POLYMER SHEARING PARAMETERS PLYSHEAR WAT-POLY **VISCOSITY** VELOCITY FACTOR 0.01.000 1.0 0.900 3.0 0.800 / TABLE NO. 01 6.0 0.700 WAT-POLY **VISCOSITY VELOCITY FACTOR** 0.0 1.000 1.0 0.900 2.0 0.800 4.0 0.750 6.0 0.700 / TABLE NO. 02 8.0 0.650

The above example activates the polymer shear thinning-thickening option and defines two polymer shear thinning-thickening tables, based on the NTPVT variable on the TABDIMS keyword in the RUNSPEC section being equal to two and NPPVT variable on the same keyword being greater than or equal to six.

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8.3.64 PLYSHLOG - ACTIVATE AND DEFINE THE POLYMER SHEARING LOGARITHMIC

PARAMETERS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword activates and defines the parameters for the logarithm-based polymer shear thinning/thickening option.

No.	Name			Default				
		Field	Metric	Laboratory				
1-1	POLCON	A real positive value that the VELOCITY and VISFA	defines the reference pol C data for this keyword.	ymer concentration for				
		lb/stb	kg/sm³	gm/scc	None			
1-2	SALTCON	A real positive value that VELOCITY and VISFAC of						
		Note that If the BRINE keyword in the RUNSPE						
		This variable is ignored a Flow.	This variable is ignored as the BRINE option is not implemented in OPM Flow.					
		lb/stb	kg/sm³	gm/scc	None			
1-3	TEMP		A real positive value defines the reference polymer temperature for the VELOCITY and VISFAC data for this keyword.					
		Note that If the TEMP keyword in the RUNSPEC						
		This variable is ignored a is not implemented in Ol						
		°F	°C	°C	None			
1-4	1	Record terminated by a '	·(₁),		Not Applicable			
2-1	VELOCITY		real monotonically increa water-polymer flow velo SALTCON and TEMP.					
		The VELOCITY value for the first row in the table should be a very small value that is greater than zero and less than 1×10^{-6} .						
		feet/day	m/day	cm/hour	None			
2-2	VISFAC	A columnar vector of real positive values that define the dimensionless shear effect multiplier for the given VELOCITY entry for the reference conditions of POLCON, SALTCON and TEMP.						
		Normally VISFAC value for						
		dimensionless	dimensionless	dimensionless	None			
1-4	1	Record terminated by a '	·//"		Not Applicable			

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No.	Name		Description		Default
		Field	Metric	Laboratory	

Revision: Rev-I

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must be entered with two records, with entries I-I, I-2, I-3 and I-4 representing record number one and 2-I, 2-2 and 2-3 representing record number two in the "No." column in this table.
- 3) Each of the records are terminated by a "/" and is explicitly shown in the above rows.
- 4) For record number two a minimum of two rows and a maximum of NPPVT rows, as declared on the TABDIMS keyword in the RUNSPEC section, are required.
- 5) There is no "/" terminator for the keyword.

Table 8.54: PLYSHLOG Keyword Description

See the PLYSHEAR keyword for the alternative polymer shear thinning/thickening option that is also implemented in OPM Flow.

Example

The following example show how to enter two PLYSHLOG tables given that the NTPVT variable on the TABDIMS keyword in the RUNSPEC section is set equal to two.

```
- -
         POLYMER SHEARING LOGARITHMIC PARAMETERS
- -
PLYSHLOG
         REF
                     REF
                                 REF
         POLCON
                     SALTCON
                                 TFMP
         0.5
         VELOCITY
                     VISFAC
         0.000001
                      1.00
         0.000001
                      1.10
         0.0001
                      1.30
         0.001
                      1.47
         0.01
                      1.67
         0.1
                      2.00
         1.0
                      2.20
         10.0
                      2.30
         100.0
                      2.40
         1000.0
                      2.40
                                                             / TABLE NO. 01
         REF
                     REF
                                 REF
         POLCON
                                 TEMP
                     SALTCON
         0.5
         VELOCITY
                     VISFAC
- -
                      1.00
         0.0000001
         0.000001
                      1.10
         0.0001
                      1.35
         0.001
                      1.57
         0.01
                      1.87
         0.1
                      2.20
         1.0
                       2.40
```

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10.02.60100.02.651000.02.65

/ TABLE NO. 02

Revision: Rev-I

The example activates the polymer logarithmic shear thinning-thickening option and defines two polymer shear thinning-thickening tables, based on the NTPVT variable on the TABDIMS keyword in the RUNSPEC section being equal to two and NPPVT variable on the same keyword being greater than or equal to ten.

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8.3.65 PLYVISC - Define Polymer Viscosity Scaling Factors

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDUL	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
----------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

PLYSVISC defines the polymer viscosity scaling factors used to determine the relationship of pure water viscosity with respect to increasing polymer saturation within a grid block. The polymer option must be activated by the POLYMER keyword in the RUNSPEC section in order to use this keyword.

The BRINE option in the RUNSPEC should be deactivated if this keyword is to be used.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	POLCON		real monotonically increa olymer concentration in t		None			
		The first entry should be	The first entry should be zero to define a no polymer concentration.					
		lb/stb	kg/sm³	gm/scc				
2	VISFAC		A columnar vector of real increasing or equal values that defines a factor that scales the effective viscosity of the solution for the given POLCON entry.					
		Normally VISFAC value for	Normally VISFAC value for the first row in the table should be one.					
		dimensionless	dimensionless	dimensionless				

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.55: PLYVISC Keyword Description

Example

- -POLYMER VISCOSITY SCALING FACTOR TABLES **PLYVISC** POLYMER **VISCOSITY** POLCON **VISFAC** 0.0000 1.000 0.0002 10.000 0.0004 20.000 / TABLE NO. 01 0.0008 40.000 POLYMER **VISCOSITY** POLCON **VISFAC** 0.0000 1.000 0.0003 10.000 0.0005 20.000 0.0007 40.000 0.0009 45.000 0.0011 55.000 / TABLE NO. 02

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The example defines two polymer viscosity scaling factor tables, based on the NTPVT variable on the TABDIMS keyword in the RUNSPEC section being equal to two and NPPVT variable on the same keyword being greater than or equal to six.

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8.3.66 PMISC - Define Miscibility versus Pressure Tables

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

PMISC defines the transition between immiscible and miscible displacement as a function of oil pressure tables, for when the MISCIBLE keyword in the RUNSPEC section has be activated. If this keyword is absent from the input deck and MISCIBLE keyword in the RUNSPEC keyword has been activated, then miscibility is independent of the oil phase pressure.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
I	PRS		A columnar vector of real monotonically increasing down the column values that defines the oil phase pressure.						
		psia	barsa	atma					
2	MISC		A columnar vector of real equal or increasing down the column values that defines the corresponding miscibility factor.						
		· ·	MISC is a scaling that should lie be zero and one, where zero means no miscibility and one means full miscibility.						
	dimensionless dimensionless dimensionless								

Notes:

- The keyword is followed by NTMISC tables as declared on the MISCIBLE keyword in the RUNSPEC section. I)
- Each table must contain a minimum of two rows and a maximum of NSMISC rows as declared on the MISCIBLE keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword. 3)

Table 8.56: PMISC Keyword Description

Example

MISCIBILITY VERSUS PRESSURE TABLES **PMISC** OIL MISCIBILE - -PRESS **FACTOR** 1000.0 0.000 2000.0 0.250 3000.0 1.000 / TABLE NO. 01 1.000 4000.0 MISCIBILE OIL PRESS **FACTOR** -----1500.0 0.000 2000.0 0.000 2500.0 0.250 3000.0 0.350 3500.0 1.000 4000.0 1.000 / TABLE NO. 02

The above example defines two miscibility versus pressure tables assuming NTMISC equals two and NSMISC is greater than or equal to six on the MISCIBLE keyword in the RUNSPEC section.

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8.3.67 **PPCWMAX** DEFINE **SWATINIT** CALCULATED CAPILL ARY **P**RESSURE

CONSTRAINTS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The PPCWMAX keyword defines the maximum capillary pressure allowed when scaling the capillary pressure tables to match the inputted SWATINIT array. This is primary used for when the SWATINIT array has values of water saturation above the connate water saturation significantly outside than capillary pressure transition zone, that is high on the structure. In this case OPM Flow may generate large values for the capillary pressure which may result in numerical converge problems. This keyword sets the maximum allowable calculated capillary pressure and how the water saturation should be treated when the limit is exceeded.

No.	Name		Description					
		Field	Metric	Laboratory				
I	PCWO		A columnar vector of real values that defines the maximum allowable capillary pressure for each SATNUM region.					
		The default value of infini						
		psia barsa atma						
2	OPTN	A columnar vector of cha	No					
		 NO: To ignore the SWATINIT value for the offending cell for when PCWO is exceeded. In this cases the capillary pressure for the block is set to the maximum (PCWO) and the water saturation is re-calculated based on PCWO. 						
		 YES: To set the SWATINIT value to the connate water saturation for the offending cell for when PCWO is exceeded. In this case the capillary pressure is set to the maximum value of the appropriate SATNUM table and the initial water saturation is calculated to be consistent with the tables maximum capillary pressure. This results in the capillary pressures not being re-scale for the offending cell. 						

Notes:

- 1) The keyword is followed by NTSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each row show contain two values representing PCWO and OPTN values.
- Each row is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.57: PPCWMAX Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Note

Using this keyword to limit the re-scaled grid block capillary pressure values will effect the fluids inplace when the simulator has to re-calculate values due to the capillary pressure limit being exceeded.

In addition the high grid block capillary pressures may be indicative of an inconsistency between the tabular SATNUM capillary pressure values and the provided SWATINIT array. This inconsistency may be a result of the SWATINIT array being derived using a saturation height function, as is customary in static modeling software, and the numerical models tabulated capillary pressure.

Example

```
SET MAXIMUM PC FOR SWATINIT INITIALIZATION
                MATCH
        MAX
        PC
                  SWATINIT
         ------
                  _____
PPCWMAX
        100.0
                  YES
                                                         / TABLE NO 01
        125.0
                  YES
                                                         / TABLE NO 02
        135.0
                  YES
                                                         / TABLE NO 03
```

The above example sets the maximum capillary pressure for three saturation regions to 100, 125 and 135 with SWATINIT reset to the connate water saturation for when the capillary pressure limit is exceeded.

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8.3.68 PROPS - Define the Start of the PROPS Section of Keywords

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

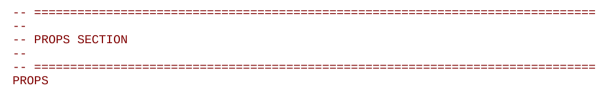
Revision: Rev-I

Description

The PROPS activation keyword marks the end of the EDIT section and the start of the PROPS section that defines the key fluid and rock property data property data for the simulator

There is no data required for this keyword.

Example



The above example marks the end of the EDIT section and the start of the PROPS section in the OPM Flow data input file.

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8.3.69 PVCDO - OIL PVT Properties for Dead OIL (Constant Compressibility)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PVCDO defines the oil PVT properties for dead oil 46 with constant compressibility. If the oil has a constant and uniform dissolved gas concentration, Gas-Oil Ratio ("GOR"), and if the reservoir pressure never drops below the saturation pressure pressure (bubble point pressure), then the model can be run more efficiently by omitting the GAS and DISGAS keywords from the RUNSPEC section, treating the oil as a dead oil, and defining a constant Rs (GOR) value with keyword RSCONST or RSCONSTT in the PROPS section. This results in the model being run as a dead oil problem with no active gas phase. However, OPM Flow takes into account the constant Rs in the calculations and reporting.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	PRS	PRS is a real positive va	alue defining the oil refer s data set.	rence pressure for the			
		psia	barsa	atma	None		
2	OFVF	-	OFVF is a real positive value defining the oil formation volume factor (Bo) at the reference pressure.				
		rb/stb	rm³/sm³	rcc/scc	None		
3	OCOMP	OCOMP is a real positive value defining the oil compressibility (Co) at the oil reference pressure and is defined as: $C_o \ = \ -\frac{1}{B_o} \bigg(\frac{dB_o}{dP} \bigg)$					
		I/psia	I/barsa	I/atma	None		
4	OVISC	OVISC is a real positive reference pressure.	e value defining the oil v	iscosity (μ_o) at the oil			
		СР	СР	СР	None		
5	OVISCOMP		itive value defining the oil wc(Pref) and is defined as:	viiscosibility (μ_{oc}) at the			
		1	$\mu_{oc} = -\frac{1}{\mu_o} \left(\frac{d\mu_o}{dP} \right)$				
		I/psia	I/barsa	I/atma	None		

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.58: PVCDO Keyword Description

See also the RSCONST and RSCONSTT keywords to define the constant Rs for dead oil and PVDO as an alternative keyword to enter the dead oil properties.

[&]quot;Dead" oil is oil that it contains no dissolved gas or a relatively thick oil or residue that has lost its volatile components.





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Exampl	le								
	OIL PVT T	ABLE FOR D	EAD WITH C	ONSTANT CO	OMPRESSIBI	LITY			
PVCD0									
	REF PRES	B0	CO	VISC	VISC				
	PSIA	RB/STB	1/PSIA	CPOISE	GRAD				
	3840.0	1.080	1.5E-6	1.750	0.0	/	TABLE	NO.	01
	3840.0	1.100	1.5E-6	1.050	0.0	/	TABLE	NO.	02
	3840.0	1.120	1.6E-6	0.950	0.0	/	TABLE	NO.	03
	3840.0	1.140	1.7E-6	0.850	0.0	/	TABLE	NO.	04
	3840.0	1.160	1.7E-6	0.800	0.0	/	TABLE	NO.	05

The above example defines five dead oil PVT tables with constant compressibility and viscosity, and assumes that NTPVT equals five on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.70 PVDG - OIL PVT Properties for Dead OIL (Constant Compressibility)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PVDG defines the gas PVT properties for dry gas 47. If the gas has a constant and uniform vaporized oil concentration, Condensate-Gas Ratio ("CGR"), and if the reservoir pressure never drops below the saturation pressure pressure (dew point pressure), then the model can be run more efficiently by omitting the OIL and VAPOIL keywords from the RUNSPEC section, treating the gas as a dry gas, and defining a constant Rv (CGR) value with keyword RVCONST or RVCONSTT in the PROPS section. This results in the model being run with as a dry gas problem with no active oil (condensate) phase. However, OPM Flow takes into account the constant Rv in the calculations and reporting.

No.	Name		Default			
		Field	Metric	Laboratory		
I	PRS	A columnar vector of real monotonically increasing down the column values that defines the gas phase pressure.				
		psia	barsa	atma	None	
2	GFVF		al decreasing down the colume fact			
		rb/Mscf	rm³/sm³	rcc/scc	None	
3	GVISC	A columnar vector of rea				
		сР	сР	сР	None	

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.59: PVDG Keyword Description

See also the RVCONST and RVCONSTT keywords to define the constant Rv for dry gas.

Natural gas that occurs in the absence of condensate or liquid hydrocarbons, or gas that had condensable hydrocarbons removed, is called dry gas. It is primarily methane with some intermediates. The hydrocarbon mixture is solely gas in the reservoir and there is no liquid (condensate surface liquid) formed either in the reservoir or at surface. The term dry indicates that the gas does not contain heavier hydrocarbons to form liquids at the surface conditions. Dry gas typically has GOR's greater than 100,000 scf/stb or 18,000 Sm3/m3.



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Exampl	les		
	CAS DVT	TABLE FOR D	IDV CAS
	GAS PVI	TABLE FUR D	IKT GAS
PVDG			
	PRES	BG	VISC
	PSIA	RB/MSCF	CPOISE
	14.7	197.8092	0.0129
	50.0	65.9364	0.0130
	100.0	31.6495	0.0130
	230.0	13.8813	0.0131
	460.0	6.8210	0.0132
	690.0	4.4703	0.0135
	920.0	3.2968	0.0138
	1150.0	2.6113	0.0141
	1380.0	2.1560	0.0145
	1610.0	1.8316	0.0150
	1840.0	1.5952	0.0155
	2070.0	1.4129	0.0161
	2300.0	1.2700	0.0167
	2372.0	1.2305	0.0169
	2530.0	1.1551	0.0174
	2760.0	1.0621	0.0181
	2990.0	0.9841	0.0189
	3220.0	0.9190	0.0196
	3450.0	0.8638	0.0204
	4500.0	0.6910	0.0242
	6000.0	0.5616	0.0293
	PRES	BG	VISC
	PSIA	RB/MSCF	CPOISE
	14.7	265.0126	0.0133
	50.0	66.2531	0.0133
	100.0	33.1266	0.0133
	230.0	14.4552	0.0134
	460.0	7.0357	0.0136
	690.0	4.6493	0.0138
	920.0	3.4417	0.0140
	1150.0	2.7227	0.0144
	1380.0	2.2522	0.0147
	1610.0	1.9158	0.0151
	1840.0	1.6702	0.0156
	2070.0	1.4805	0.0162
	2300.0	1.3317	0.0167
	2372.0	1.2927	0.0169
	2530.0	1.2119	0.0173
	2760.0	1.1135	0.0180
	2990.0	1.0325	0.0187
	3220.0	0.9637	0.0194
	3450.0	0.9055	0.0201
	4500.0	0.7228	0.0236
	6000.0	0.5837	0.0285

The above example defines two dry PVT tables assuming NTPVT equals two and NPPVT is greater than or equal to 22 on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.71 PVDO - OII PVT PROPERTIES FOR DEAD OII

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PVDO defines the oil PVT properties for dead oil 48. If the oil has a constant and uniform dissolved gas concentration, Gas-Oil Ratio ("GOR"), and if the reservoir pressure never drops below the saturation pressure pressure (bubble point pressure), then the model can be run more efficiently by omitting the GAS and DISGAS keywords from the RUNSPEC section, treating the oil as a dead oil, and defining a constant Rs (GOR) value with keyword RSCONST or RSCONSTT in the PROPS section. This results in the model being run as a dead oil problem with no active gas phase. However, OPM Flow takes into account the constant Rs in the calculations and reporting.

No.	Name	Description					
		Field	Metric	Laboratory			
I	PRS	A columnar vector of real monotonically increasing down the column values that defines the oil phase pressure.					
		psia	barsa	atma	None		
2	OFVF		al decreasing down the colume factors				
		rb/stb	rm³/sm³	rcc/scc	None		
3	OVISC	A columnar vector of reathe corresponding oil pha					
		cP	cP	cP	None		

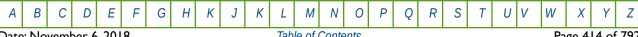
Notes:

- The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section. I)
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword. 3)

Table 8.60: PVDO Keyword Description

See also the RSCONST and RSCONSTT keywords to define the constant Rs for dead oil and PVCDO as an alternative keyword to enter the dead oil properties.

[&]quot;Dead" oil is oil that it contains no dissolved gas or a relatively thick oil or residue that has lost its volatile components.



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Exam	ples		
	OIL PVT T	ABLE FOR	DEAD OIL
PVD0			
	PSAT	В0	VISC
	PSIA	RB/STB	CPOISE
	400	4 0400	
	400	1.0102	1.16
	1200 2000	1.0040 0.9960	1.164 1.167
	2800	0.9880	1.172
	3600	0.9802	1.177
	4400	0.9724	1.181
	5200	0.9646	1.185
	5600	0.9607	1.19
	800	1.0255	1.14
	1600	1.0172	1.14
	2400	1.0091	1.14
	3200	1.0011	1.14
	4000 4800	0.9931 0.9852	1.14 1.14
	5600	0.9832	1.14
	3000	0.3114	1.14

The above example defines two dead oil PVT tables with variable viscosity and compressibility with respect to pressure, and assumes that NTPVT equals two and NPPVT is greater than or equal to eight on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.72 PVDS - SOLVENT PVT PROPERTIES FOR THE SOLVENT MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PVDS defines the solvent PVT properties for use with SOLVENT option. The solvent is treated as an additional dry gas phase within the model. This keyword should only be used if the SOLVENT model has been invoked in the RUNSPEC section.

No.	Name		Default				
		Field	Metric	Laboratory			
I	PRS	A columnar vector of r	None				
		psia barsa atma					
2	GFVF	A columnar vector of real decreasing down the column values that defines the corresponding solvent phase formation volume factor.					
		rb/Mscf	rm³/sm³	rcc/scc			
3	GVISC	A columnar vector of rea	None				
		cP	сР	сР			

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.61: PVDS Keyword Description

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Examp	les		
	GAS SOLV	VENT PVT TAB	SLE
PVTS	DDEO	B0	\/T00
	PRES	BG BR/MSCE	VISC CPOISE
	PSIA	RB/MSCF	CPUISE
-	700.0	4.4703	0.0135
	920.0	3.2968	0.0138
	1150.0	2.6113	0.0141
	1380.0	2.1560	0.0145
	1610.0	1.8316	0.0150
	1840.0	1.5952	0.0155
	2070.0	1.4129	0.0161
	2300.0	1.2700	0.0167
	2372.0	1.2305	0.0169
	2530.0	1.1551	0.0174
	2760.0	1.0621	0.0181
	2990.0	0.9841	0.0189
	3220.0	0.9190	0.0196
	3450.0	0.8638	0.0204
	4500.0	0.6910	0.0242
	6000.0	0.5616	0.0293
	PRES	BG	VISC
	PSIA	RB/MSCF	CPOISE
	700.0	4.6493	0.0138
	920.0	3.4417	0.0140
	1150.0	2.7227	0.0144
	1380.0	2.2522	0.0147
	1610.0	1.9158	0.0151
	1840.0	1.6702	0.0156
	2070.0	1.4805	0.0162
	2300.0	1.3317	0.0167
	2372.0	1.2927	0.0169
	2530.0	1.2119	0.0173
	2760.0	1.1135	0.0180
	2990.0	1.0325	0.0187
	3220.0	0.9637	0.0194
	3450.0	0.9055	0.0201
	4500.0	0.7228	0.0236
	6000.0	0.5837	0.0285

The above example defines two solvent PVT tables assuming NTPVT equals two and NPPVT is greater than or equal to 16 on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.73 PVTG - GAS PVT Properties for Wet GAS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PVTG defines the gas PVT properties for wet gas⁴⁹. This keyword should be used when the VAPOIL keyword has be declared in the RUNSPEC section indicating that that vaporized oil (more commonly referred to as condensate) is present in wet gas in the model. The keyword may be used for gas-water and oil-water-gas input decks that contain the oil and gas phases.

No.	Na	ıme		Description		Default		
			Field Metric Laboratory					
I	PRS		gas phase pressure. the ("CGR") or Rv, the gas the corresponding pressure.	reasing down the column at defines the saturated formation volume factor a are for the stated saturate	I condensate-gas ratio and the gas viscosity for d RVS.			
			gas viscosity with respect as a sub table under R	ibility of the gas formation t to the under-saturated F VU, FVFU and VISU colo for each sub table entry. I ""."	Rv is optionally included umns, that is it is not			
		The under saturated Rv entries are optional, except for perhaps the last PRS entry to define the PVT properties above the initial saturation pressure.						
			psia	barsa	atma	None		
2	RVS	RVU	A columnar vector of reand under saturated (RVI	eal positive number for bo J) Rv sub table entries.	oth the saturated (RVS)			
			The RVS entry on the main table is the saturated CGR at the pressure indicated by PRS and may be increasing or decreasing in value as PRS varies.					
			Subsequent under-saturated Rvs for a sub table at the given PRS, as defined by RVU, are monotonically decreasing for entries in a given sub table.					
			stb/Mscf	sm³/sm³	rcc/scc	None		
3	FVFS	FVFU	A columnar vector of real decreasing down the column values that defines the corresponding gas phase formation volume factor for a given pressure (PRS) and for a given Rv (either RVS or RVU).					
			rb/Mscf	rm³/sm³	rcc/scc	None		
4	VISS	VISS VISU VISS a columnar vector of real increasing down the column values that defines the corresponding gas phase viscosity for a given pressure (PRS) and for a given RVS.						
	VISU a columnar vector of real decreasing from VISS down the column values that defines the corresponding gas phase viscosity for a given pressure (PRS) and for a given RVU.							
			cP	cP	cP	None		

Natural gas that contains significant heavy hydrocarbons such as propane, butane and other liquid hydrocarbons is known as wet gas or rich gas. The general rule of thumb is if the gas contains less methane (typically less than 85% methane) and more ethane, and other more complex hydrocarbons, it is labeled as wet gas. Wet gas normally has GOR's less than 100,000 scf/stb or 18,000 Sm3/m3, with the condensate having a gravity greater than 50 °API.

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No.	Name		Description		Default	
		Field	Field Metric La			

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Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NPPVT rows and NRPVT Rv entries as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Apart from the PRES data there must be same number of entries for each column.
- 4) Each sub table defining the under saturated gas properties must be terminated by "/".
- 5) Each main table is terminated by a single "/" on a separate line and there is no "/" terminator for the keyword.

Table 8.62: PVTG Keyword Description

Examples

```
GAS PVT TABLE FOR WET GAS
PVTG
         PRES
                     RV
                                   BG
                                                 VISC
                   STB/MSCF
- -
         PSIA
                                 RB/MSCF
                                                CPOISE
                   _____
                                 -----
                                                _ _ _ _ _
          300
                   0.000132
                                  0.042340
                                                 0.01344
                   0
                                  0.042310
                                                 0.01389
          600
                   0.000124
                                  0.020460
                                                 0.01420
                   0
                                  0.020430
                                                 0.01450
          900
                   0.000126
                                  0.013280
                                                 0.01526
                   0
                                  0.013250
                                                 0.01532
         1200
                   0.000135
                                  0.009770
                                                 0.01660
                                  0.009730
                                                 0.01634
                   0
         1500
                   0.000149
                                  0.007730
                                                 0.01818
                   0
                                  0.007690
                                                 0.01752
         1800
                   0.000163
                                  0.006426
                                                 0.01994
                   0
                                  0.006405
                                                 0.01883
                   0.000191
         2100
                                  0.005541
                                                 0.02181
                   0
                                  0.005553
                                                 0.02021
         2400
                   0.000225
                                  0.004919
                                                 0.02370
                                  0.004952
                   0
                                                 0.02163
                                                               / TABLE NO. 1
         PRFS
                     RV
                                                 VISC
                                   BG
                   STB/MSCF
                                 RB/MSCF
                                                CPOISE
         PSTA
                                 _ _ _ _ _ _
          300
                   0.000132
                                  0.042340
                                                 0.01344
          600
                   0.000124
                                  0.020460
                                                 0.01420
          900
                   0.000126
                                  0.013280
                                                 0.01526
         1200
                   0.000135
                                  0.009770
                                                 0.01660
         1500
                   0.000149
                                  0.007730
                                                 0.01818
         1800
                   0.000163
                                  0.006426
                                                 0.01994
                                  0.005541
         2100
                   0.000191
                                                 0.02181
                                  0.004919
         2400
                   0.000225
                                                 0.02370
                                  0.004952
                                                 0.02163
                                                               / TABLE NO. 2
```

The above example defines two wet PVT tables assuming NTPVT equals two, NPPVT is greater than or equal to eight, and NRPVT greater than or equal to two on the TABDIMS keyword in the RUNSPEC section.

Notice that there is no terminating "/" for this keyword only for a table and a sub table.

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8.3.74 PVTO - OIL PVT PROPERTIES FOR LIVE OIL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PVTO defines the oil PVT properties for live⁵⁰ and the keyword should only be used if the there is both oil and gas phases in the model. This keyword should be used when the DISGAS keyword has be declared in the RUNSPEC section indicating that that dissolved gas (more commonly referred to as solution gas) is present in the oil. The keyword may be used for oil-water and oil-water-gas input decks.

No.	Na	me			Default				
			Field	Metric	Laboratory				
I	RS		saturated gas-oil ratio (reasing down the column "GOR") or Rs, that de oil viscosity for the ta	fines the oil formation				
	For a given RS the variability of the oil formation volume factor and the oil viscosity with respect to the saturated RS and pressure is optionally included as a sub table under PRSU, FVFU and VISU columns, that is it is not necessary to repeat RS for each sub table entry. However, each sub table must be terminated by a "/".								
			The under-saturated PRI last RS entry to define pressure.						
			If there are no following row should be terminat entries then the last PRSI						
			Mscf/stb	sm³/sm³	scc/scc	None			
2	PRSS	PRSU	column values that define pressure), that defines the	vector of real monotonica es the oil phase saturation e oil formation volume fac SS pressure for a given sa	pressure (bubble-point tor and the oil viscosity				
			column values that defir defines the oil formation	lumnar vector of real monotonically increasing down the last defines the oil phase under-saturated pressure that ormation volume factor and the oil viscosity for the LSU pressure for a given saturated RS.					
			Note that PRSU should b	e greater than PRSS.					
			psia	barsa	atma	None			
3	FVFS	FVFU	FVFS is a columnar vector defines the corresponding a given pressure (PRSS) a						
			FVFU is a columnar vector defines the corresponding factor for a given pressure.						
			rb/stb	rm³/sm³	rcc/scc	None			

[&]quot;Live" oil is oil that contains gas in solution, which is normally the case for most conventional oil reservoirs. However, for oil reservoirs classified as heavy oil reservoirs, the in situ dissolved gas may be negligible and oil would then be classified as gasfree oil which is commonly referred to as "dead" oil.

Α	В	С	D	Ε	F	G	Н	Κ	J	Κ	L	М	N	0	Р	Q	R	S	Τ	U	V	W	Χ	Υ	Z
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No.	Na	ıme		Description						
			Field	Metric	Laboratory					
4	VISS	VISU VISS a columnar vector of real increasing down the column values that defines the corresponding oil phase saturated viscosity for a given pressure (PRSS) and for a given RS. If this is the only entry for a given RS and PRSS then the record should be terminate by a "/".								
			values that defines the confor a given pressure (PRS	ISU a columnar vector of real decreasing from VISS down the column alues that defines the corresponding oil phase under-saturated viscosity or a given pressure (PRSU) and for a given RS. If this is the only entry for given RS and PRSU then the record should be terminate by a "/".						
			cР	cP	cP	None				

Revision: Rev-I

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NPPVT rows and NRPVT Rs entries as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Apart from the RS data there must be same number of entries for each column.
- 4) Each sub table defining the under saturated gas properties must be terminated by "/".
- 5) Each main table is terminated by a single "/" on a separate line and there is no "/" terminator for the keyword.

Table 8.63: PVTO Keyword Description

Examples

The following example defines live oil PVT tables assuming NTPVT equals two, NPPVT is greater than or equal to eight, and NRPVT greater than or equal to two on the TABDIMS keyword in the RUNSPEC section.

```
OIL PVT TABLE FOR LIVE OIL
PVT0
                    PSAT
                                          VISC
         RS
                               B0
         MSCF/STB PSIA
                               RB/STB
                                          CPOISE
                    _____
                               _____
          0.0010
                      14.7
                               1.05340
                                         1.7230
          0.0890
                      500.0
                               1.08890
                                          1.1670
          0.2060
                     1000.0
                                          0.8570
                               1.13850
          0.3360
                    1500.0
                               1.19640
                                          0.6840
          0.4050
                    1750.0
                               1.22800
                                          0.6240
          0.4750
                     2000.0
                               1.26110
                                          0.5750
          0.5480
                     2250.0
                               1.29570
                                          0.5340
          0.6220
                     2500.0
                               1.33160
                                          0.5000
                     2750.0
          0.6980
                               1.36890
                                          0.4700
          0.7750
                     3000.0
                               1.40740
                                          0.4450
          0.8530
                     3250.0
                               1.44710
                                          0.4220
          0.9330
                     3500.0
                               1.48790
                                          0.4020
          1.0140
                     3750.0
                               1.52980
                                          0.3840
          1.0960
                     4000.0
                               1.57280
                                          0.3680
          1.1800
                     4258.0
                               1.61760
                                          0.3530
          1.2630
                     4500.0
                               1.66190
                                          0.3400
          1.3480
                     4750.0
                               1.70780
                                          0.3280
          1.4340
                     5000.0
                               1.75480
                                          0.3170
          1.6060
                     5500.0
                               1.85020
                                          0.2980
                     6242.0
                               1.83040
                                          0.3186
                                                  / TABLE NO. 1
```

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 RS	PSAT	В0	VISC	
 MSCF/STB	PSIA	RB/STB	CPOISE	
0.0010	14.7	1.05340	1.7230	/
0.0390	250.0	1.06830	1.4220	/
0.0890	500.0	1.08890	1.1670	/
0.1460	750.0	1.11250	0.9850	/
0.2060	1000.0	1.13850	0.8570	/
0.2700	1250.0	1.16660	0.7590	/
0.3360	1500.0	1.19640	0.6840	/
0.4050	1750.0	1.22800	0.6240	/
0.4750	2000.0	1.26110	0.5750	/
0.5480	2250.0	1.29570	0.5340	/
0.6220	2500.0	1.33160	0.5000	/
0.6980	2750.0	1.36890	0.4700	/
0.7750	3000.0	1.40740	0.4450	/
0.8530	3250.0	1.44710	0.4220	/
0.9330	3500.0	1.48790	0.4020	/
1.0140	3750.0	1.52980	0.3840	/
1.0960	4000.0	1.57280	0.3680	/
1.1800	4258.0	1.61760	0.3530	/
1.2630	4500.0	1.66190	0.3400	/
1.3480	4750.0	1.70780	0.3280	/
1.4340	5000.0	1.75480	0.3170	/
1.6060	5500.0	1.85020	0.2980	
	6242.0	1.83040	0.3186	/
				/ TABLE NO.

Notice that there is no terminating "/" for this keyword only for a table and a sub table.

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8.3.75 PVTW - Define Water Fluid Properties for Various Regions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PVTW defines the water properties for various regions in the model. The number of PVTW vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the PVTW tables to different grid blocks in the model is done via the PVTNUM keyword in the REGION section. One data set consists of one record or line which is terminated by a "/". If the water phase is active in the model, which is normally the case, then this keyword must be defined in the OPM Flow input deck.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	PRES	PRES is a real number d other parameters for this	efining the water references data set.	ce pressure (P) for the				
		psia	barsa	atma	None			
2	WFVF	WFVF is a real number of at the water reference pr	defining the water format essure.	ion volume factor (Bw)				
		rb/stb	rm³/sm³	rcc/scc				
		1.0	1.0	1.0	Defined			
3	WCOMP	WCOMP is a real number defining the water compressibility (Cw) at the water reference pressure and is defined as: $C_{_W} = -\frac{1}{B_{_W}} \left(\frac{dB_{_W}}{dP} \right)$						
		I/psia	I/barsa	I/atma				
		0.00004	0.00004	0.00004	Defined			
4	WVISC	WVISC is a real number reference pressure	r defining the water visco	osity (μ_w) at the water				
		СР	СР	СР				
		0.50	0.50	0.50	Defined			
5	WVISCOMP		mber defining the water v , Uwc(Pref) and is defined					
		$\mu_{wc} = -\frac{1}{\mu_w} \left(\frac{d \mu_w}{dP} \right)$						
		I/psia	I/barsa	I/atma				
		0.0	0.0	0.0	Defined			

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.64: PVTW Keyword Description

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Examples

The following shows the PVTW keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to one.

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```
WATER PVT TABLE
--
PVTW
--
        REF PRES BW
                           CW
                                     VISC
                                             VISC
                 RB/STB
                           1/PSIA
        PSIA
                                     CPOISE
                                              GRAD
                                     _____
        4840.0
                 1.019
                           2.7E-6
                                     0.370
                                              1*
                                                        / TABLE NO. 01
```

The next example shows the PVTW keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to three.

PVTW						
	REF PRES	BW	CW	VISC	VISC	
	PSIA	RB/STB	1/PSIA	CPOISE	GRAD	
	4640.0	1.008	2.5E-6	0.350	1*	/ TABLE NO. 01
	4840.0	1.019	2.7E-6	0.370	1*	/ TABLE NO. 02
	4940.0	1.030	2.8E-6	0.390	1*	/ TABLE NO. 03

The above example defines three water PVT tables and assumes that NTPVT equals three on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.76 REFINE - START THE DEFINITION OF A LOCAL GRID REFINEMENT

The REFINE keyword defines the start of a Cartesian or radial local grid refinement ("LGR") definition that sets the properties of the selected LGR. The keyword is then followed by the property keywords associated with the section where the keyword is being invoked. For example, if the REFINE keyword is used in the GRID section then most of the keywords in that section can be used to set the grid properties for the LGR.

See REFINE – Start the Definition of a Local Grid Refinement in the GRID section for a full description.

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8.3.77 RKTRMDIR - ACTIVATE ROCKTAB KEYWORD DIRECTIONAL TRANSMISSIBILITY

MULTIPLIERS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword activates the directional transmissibility multipliers for the ROCKTAB keyword. This results in two additional columns being inputted on the ROCKTAB keyword. This feature is currently not supported in OPM Flow.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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8.3.78 ROCK - Define the Rock Compressibility for Various Regions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

ROCK defines the rock compressibility for various regions in the model. The number of ROCK vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the ROCK tables to different grid blocks in the model is done via the PVTNUM keyword in the REGION section. One data set consists of one record or line which is terminated by a "/".

This keyword must be defined in the OPM Flow input deck.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
I	PRES	PRES is a real number de parameters for this data	efining the rock reference set.	pressure for the other	Default				
		psia	barsa	atma					
		1.032	1.032	1.032					
2	RCOMP	RCOMP is a real numb	er defining the rock con and is defined as:	npressibility (Cf) at the	Defined				
			$C_f = -\frac{1}{V} \left(\frac{dV}{dP} \right)$						
		I/psia I/barsa I/atma							
		0.0	0.0	0.0					

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.65: ROCK Keyword Description

Examples

The following shows the ROCK keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to one.

The next example shows the ROCK keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to three.

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-- ROCK COMPRESSIBILITY (1) REFERENCE PRESSURE IS TAKEN FROM THE HCPV WEIGHTED RESERVOIR PRESSURE AS THE PORV IS ALREADY AT RESERVOIR CONDITIONS (OPM FLOW USES THE REFERENCE PRESSURE) TO CONVERT THE GIVEN PORV TO RESERVOIR CONDITIONS USING THE DATA ON THE ROCK KEYWORD) - --- REF PRES CF -- PSIA 1/PSIA **ROCK** 3566.9 5.0E-06 / ROCK COMPRSSIBILITY REGION 1 3966.9 5.5E-06 / ROCK COMPRSSIBILITY REGION 2 / ROCK COMPRSSIBILITY REGION 3 4566.9 6.0E-06

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The above example defines three ROCK tables and assumes that NTPVT equals three on the TABDIMS keyword in the RUNSPEC section.

There is no terminating "/" for this keyword.

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8.3.79 ROCKOPTS - DEFINE ROCK COMPACTION AND COMPRESSIBILITY OPTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The ROCKOPTS keyword defines various option with respect to rock compaction and rock compressibility.

No.	Name	Description	Default
I	ROCKOPTI	ROCKOPTI is a character string that defines the treatment of how the overburden pressures supplied by the OVERBURD keyword are applied to the tabulated pressures in the ROCKTAB keywords:	PRESSURE
		3) STRESS: Use this option if the overburden pressures on the OVERBURD keyword are greater than the fluid pressure which results in the effective fluid pressure being negative. To avoid the rock compaction tables being entered with negative pressure values use this option. In this case the pore volume and transmissibility multipliers will be tabulated against the effective overburden pressure	
		4) PRESSURE: In this case the pore volume and transmissibility multipliers should be effective pressure. This the default value.	
		ROCKOPTI should be set to PRESSURE if the OVERBURD is not used in the input deck.	
2	ROCKOPT2	ROCKOPT2 is a character string that sets the reference pressure option:	NOSTORE
		 STORE: Copies the initial calculated grid block pressures into the overburden pressure array, resulting in the pore volumes being referenced at the initial pressures instead of the reference pressures as per the ROCKTAB keyword. 	
		 NOSTORE: This option results in the pore volumes being referenced as per the ROCKTAB keyword. This is the default value. 	
		Note that STORE option should not be used with the OVERBURD keywords as the OVERBURD data will be overwritten.	
3	ROCKOPT3	ROCKOPT3 is a character string that defines which region array should be used to allocate the various ROCK and ROCKTAB tabless. ROCKOPT3, should be set to ROCKNUM, SATNUM or PVTNUM.	PVTNUM
4	ROCKOPT4	ROCKOPT4 is a character string that sets the initial conditions for the HYSTER and BOBERG options:	DEFLATION
		 DEFLATION: This option defines the reservoir rock to be fully compacted and the deflation curve is used to calculated the initial pore volume and transmissibility multipliers. This is the default value. 	
		 ELASTIC: This option sets the pore volume and transmissibility multipliers to one, as the reservoir rock is set to lie on the elastic curve. 	

The keyword is terminated by "/".

Table 8.66: ROCKOPTS Keyword Description

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This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Revision: Rev-I

Example

-- ROCKOPT1 ROCKOPT2 ROCKOPT3 ROCKOPT3
-- PRS/STRE NO/STORE ARRAY
-- ROCKOPTS
PRESSURE NOSTORE PVTNUM DEFLATION / ROCK COMP OPTIONS

The above example defines the default values for the ROCKOPTS keyword.

FLOW DOCUMENTATION MANUAL (2018-10)

8.3.80 ROCKTAB - ROCK COMPACTION TABLES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The ROCKTAB keyword defines the rock compaction attributes to be applied for when the rock compaction option has been invoked by the ROCKCOMP keyword in the RUNSPEC section. ROCKTAB defines pore volume and transmissibility multipliers versus pressure that are used in the compaction calculations. If the RKTRMDIR has been activated in the PROPS section, then the transmissibility multiplier is directional dependent and two additional columns are used to define the y and z direction transmissibility multipliers.

This keyword should only be used if compaction option has been enabled.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
ı	PRS	If the ROCKOPTI variab keyword in the PROPS s real monotonically incre reference pressure for w If ROCKOPTI has been vector of real monotonic					
		psia	bars	atm	None		
2	PORV	A columnar vector of increasing down the colufor a given PRS.					
		dimensionless	dimensionless	dimensionless	None		
3	TRANS	columnar vector of real	bsent from the input dopositive values that are endefine the x, y, and z dir ponding PRS.	ther equal or increasing			
		If the RKTRMDIR is pres vector of real positive va column that define only the corresponding PRS.	or increasing down the				
		dimensionless	dimensionless	dimensionless	None		
4	TRANSY	If the RKTRMDIR is absent from the input deck, then TRANSY is ignored.					
		If the RKTRMDIR is p columnar vector of real down the column that multipliers for the corres	ther equal or increasing				
		dimensionless	dimensionless	dimensionless	None		

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No.	Name		Description					
		Field	Field Metric Laboratory					
5	TRANSZ	If the RKTRMDIR is a ignored.	If the RKTRMDIR is absent from the input deck, then TRANSZ is ignored.					
		columnar vector of real down the column that	If the RKTRMDIR is present in the input deck, then TRANSZ is a columnar vector of real positive values that are either equal or increasing down the column that define only the z directional transmissibility multipliers for the corresponding PRS.					
		dimensionless	dimensionless	dimensionless	None			

Revision: Rev-I

Notes:

- The keyword is followed by NTROCC tables as declared on the ROCKCOMP keyword in the RUNSPEC I) section.
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the 2) TABDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.67: ROCKTAB Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

The example below defines two rock compaction tables, assuming NTROCC is equal to two on the ROCKCOMP keyword and NPPVT is greater than or equal to four on the TABDIMS keyword and that the RKTRMDIR keyword is present in the input deck.

	ROCK COM	MPACTION T	TABLES			
ROCKTAB						
	PRESS	PORV	TX(YZ)	TY	TZ	
		MULT	MULT	MULT	MULT	
	1000.0	0.9600	0.9650	0.9650	0.9650	
	1500.0	0.9800	0.9850	0.9850	0.9500	
	3000.0	0.9900	0.9950	0.9950	0.9950	
	4500.0	1.0000	1.0000	1.0000	1.0000	
	4750.0	1.0100	1.0100	1.0100	1.0100	/ TABLE NO. 01
	PRESS	P0RV	TX(YZ)	TY	TZ	
		MULT	MULT	MULT	MULT	
	1000.0	0.9600	0.9650	0.9650	0.9650	
	1500.0	0.9800	0.9850	0.9850	0.9500	
	3000.0	0.9900	0.9950	0.9950	0.9950	
	4500.0	1.0000	1.0000	1.0000	1.0000	
	4750.0	1.0100	1.0100	1.0100	1.0100	/ TABLE NO. 02

As the x, y and z directional transmissibility multipliers are identical in the above example, we could eliminate the RKTRMDIR keyword from the input deck and enter the data in the three column format, as shown on the next page.

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--ROCK COMPACTION TABLES **ROCKTAB PRESS** PORV TX(YZ) --MULT MULT - -_____ _____ 1000.0 0.9600 0.9650 1500.0 0.9800 0.9850 3000.0 0.9900 0.9950 4500.0 1.0000 1.0000 / TABLE NO. 01 4750.0 1.0100 1.0100 PRESS PORV TX(YZ) MULT MULT ____ -----1000.0 0.9600 0.9650 1500.0 0.9800 0.9850

Revision: Rev-I

/ TABLE NO. 02

The net result of the two examples in this case is identical.

0.9900

1.0000

1.0100

0.9950

1.0000

1.0100

3000.0

4500.0

4750.0

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8.3.81 RPTPROPS - Define PROPS Section Reporting

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the data in the PROPS section that is to be printed to the output print file in human readable format. The keyword has two distinct forms, the first of which consists of the keyword followed by a series of integers on the next line indicating the data to be printed (see the first example). This is the original formal in the commercial simulator and was subsequently superseded by the second format. The second format consists of the keyword followed by a series of character strings that indicate the data to be printed. In most cases the character string is the keyword used to load the data in the OPM Flow input deck, for example PVDG for the dry gas PVT tables. Its is anticipated that OPM Flow will eventually support the functionality of the second format only, the first format although recognized will be completely ignored.

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

No.	Name	Description	Default
I	PVTDG	Print dry gas PVT tables	N/A
2	PVTG	Print wet gas PVT tables	N/A
3	SGFN	Print gas relative permeability saturation function tables.	N/A
4	SGL	Print connate gas saturation array.	N/A
			N/A
		1	1

Notes:

I) The keyword is terminated by "/".

Table 8.68: RPTPROPS Keyword Description

Note

Except for tabular like data, PVTDG etc., this keyword has the potential to produce very large print files that some text editors may have difficulty loading. A more efficient solution for array type data is to load the *.INIT file into OPM ResInsight to view the data graphically, this also has the benefit of being able to filter the grid based on I, J, K ranges and grid properties.

Examples

The first example shows the original format of this keyword; although the keyword and format are recognized by OPM Flow, the format is ignored and is unlikely to be implemented in in the simulator.

```
DEFINE PROPS SECTION REPORT OPTION (ORIGINAL FORMAT)

RPTPROPS

1 2*0 1 3*1 /
```

The next example shows the second format of the keyword which may be supported in a future release of OPM Flow.

```
-- DEFINE PROPS SECTION REPORT OPTIONS
-- RPTPROPS
PVDO SOF2 SGFN SWFN /
```

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8.3.82 RSCONST - DEFINE CONSTANT GOR FOR DEAD OIL PVT FILLIDS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

RSCONST defines a constant Gas-Oil Ratio ("GOR"), for all dead oil⁵¹ PVT fluids. If the oil has a constant and uniform dissolved gas concentration, GOR, and if the reservoir pressure never drops below the saturation pressure pressure (bubble point pressure), then the model can be run more efficiently by omitting the GAS and DISGAS keywords from the RUNSPEC section, treating the oil as a dead oil, and defining a constant Rs (GOR) value with keywords RSCONST or RSCONSTT in the PROPS section. This results in the model being run as a dead oil problem with no active gas phase. However, OPM Flow takes into account the constant Rs in the calculations and reporting.

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

	Description						
Field	Metric	Laboratory					
A real positive value that the model	·						
Mscf/stb	sm³/sm³	scc/scc	None				
		pressure (bubble point					
psia	barsa	atma	None				
<u> </u>	Dai Sa	auna					
	A real positive value that the model Mscf/stb A real positive value that pressure) for all the oil P	Field Metric A real positive value that defines the dead oil GOR the model Mscf/stb sm³/sm³ A real positive value that defines that saturation pressure) for all the oil PVT tables in the model. psia barsa	Field Metric Laboratory A real positive value that defines the dead oil GOR for all oil PVT tables in the model Mscf/stb sm³/sm³ scc/scc A real positive value that defines that saturation pressure (bubble point pressure) for all the oil PVT tables in the model. psia barsa atma				

Table 8.69: RSCONST Keyword Description

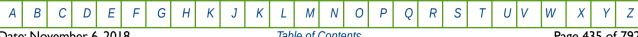
See also the RSCONSTT keyword to define a different constant Rs to the various dead oil PVT tables and the PVDO and PVCDO keywords to enter the dead oil properties. All of the aforementioned keywords are in the PROPS section.

Examples

The example sets the dead oil GOR to 5 scf/stb and the bubble point pressure to 14.7 psia.

```
DEAD OIL PVT CONSTANT GOR AND SATURATION PRESSURE
- -
RSCONST
                    PSAT
- -
         MSCF/STB PSIA
          0.0050
                     14.7
```

[&]quot;Dead" oil is oil that it contains no dissolved gas or a relatively thick oil or residue that has lost its volatile components.



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8.3.83 RSCONSTT - Define Constant GOR for Dead Oil PVT Fluids

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

RSCONSTT defines a constant Gas-Oil Ratio ("GOR"), for <u>each</u> dead oil⁵² PVT fluid in the model. If the oil has a constant and uniform dissolved gas concentration, GOR, and if the reservoir pressure never drops below the saturation pressure pressure (bubble point pressure), then the model can be run more efficiently by omitting the GAS and DISGAS keywords from the RUNSPEC section, treating the oil as a dead oil, and defining a constant Rs (GOR) value with keywords RSCONST or RSCONSTT in the PROPS section. This results in the model being run as a dead oil problem with no active gas phase. However, OPM Flow takes into account the constant Rs in the calculations and reporting.

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

No.	Name		Description					
		Field	Metric	Laboratory				
I	RS	A real positive columnar PVT table in the model	A real positive columnar vector that defines the dead oil GOR for each oil PVT table in the model					
		Mscf/stb	sm³/sm³	scc/scc	None			
2	PRS	•	A real positive columnar vector that defines the saturation pressure (bubble point pressure) for each the oil PVT table in the model.					
		psia	barsa	atma	None			

Notes:

- 1) The keyword is followed by NTPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each row must contain two values representing the RS and PRS variables.
- 3) Each row is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.70: RSCONSTT Keyword Description

See also the RSCONST keyword to define a constant Rs to all the various dead oil PVT tables and the PVDO and PVCDO keywords to enter the dead oil properties. All of the aforementioned keywords are in the PROPS section.

Examples

The example sets the dead oil GOR to 5, 6.5 and 8.0 scf/stb for PVT tables one, two and three, respectively and the bubble point pressure to 14.7 psia. for all three tables.

-- DEAD OIL PVT CONSTANT GOR AND SATURATION PRESSURE

RSCONSTT

 RS	PSAT	
 MSCF/STB	PSIA	
0.0050	14.7	/ TABLE NO. 01
0.0065	14.7	/ TABLE NO. 02
0.0080	14.7	/ TABLE NO. 03

⁵² "Dead" oil is oil that it contains no dissolved gas or a relatively thick oil or residue that has lost its volatile components.

FLOW DOCUMENTATION MANUAL (2018-10)

8.3.84 RTEMP - Define the Initial Reservoir Temperature for the Model

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEL	RUNSPEC	SPEC GRID L	EDIT PRO	OPS REGIONS	SOLUTION	SUMMARY	SCHEDULE
--------------------------------------------------------	---------	-------------	----------	-------------	----------	---------	----------

Description

This keyword defines the reservoir temperature for when OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. The RTEMPA keyword is alias for RTEMP; however the former is ignored by OPM Flow.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

		Description					
	Field	Laboratory					
RTEMP	Single real positive value model.						
	°F	°C	°C	None			
ξ-	TEMP	model.	model.	model.			

Notes:

1) The keyword is terminated by "/".

Table 8.71: RTEMP Keyword Description

See also the RTEMPVD keyword in SOLUTION section to define the reservoir temperature as a function of depth.

Example

-

RESERVOIR TEMPERATURE

----RTEMP

190.0

/ RESERVOIR TEMPERATURE

Revision: Rev-I

The above example defines the reservoir temperature to be 190 °F.

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8.3.85 RTEMPA - DEFINE THE INITIAL RESERVOIR TEMPERATURE FOR THE MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

/ RESERVOIR TEMPERATURE

Description

This keyword defines the reservoir temperature for when temperature or thermal options has been activated by the either the TEMP or THERMAL keywords in the RUNSPEC section in the commercial simulator.

The RTEMP keyword is alias for RTEMPA; however the latter is ignored by OPM Flow.

No.	Name		Default		
		Field	Metric	Laboratory	
I	RTEMPA	Single real positive value model.	None		
		°F	°C	°C	
Note	<u>s:</u>				

The keyword is terminated by "/".

Table 8.72: RTEMPA Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness. Use the RTEMP keyword instead to set the reservoir temperature.

Example

RESERVOIR TEMPERATURE RTEMPA 190.0

The above example defines the reservoir temperature to be 190 °F.

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8.3.86 SALNODE - SALT CONCENTRATION BASED PVTNUM ARRAY

	RUNSPEC	GRID	EDIT		REGIONS	SOLUTION	SUMMARY	SCHEDULE
--	---------	------	------	--	---------	----------	---------	----------

Revision: Rev-I

Description

SALNODE defines the salt concentration value based on a cells PVTNUM number. The SALNODE property is used in the calculation of a polymer viscosity when the polymer and the salt options has been activated by the POLYMER and BRINE keywords in the RUNSPEC section. In the RUNSPEC section the number of PVTNUM functions is declared by NTPVT variable on the TABDIMS keyword and allocated to individual cells by the PVTNUM property array in the REGIONS section. NPPVT on the TABDIMS keyword in the RUNSPEC section defines the maximum number of rows (or pressure values) in the PVT tables and also sets the maximum number of entries for each SALNODE data set. The number of values for each data set must correspond to the number of polymer solution adsorption entries on the PLYADSS keyword. For example if there are three sets of PVT tables and four values on the PLYADSS keyword, then three SALNODE data sets with four values of salt concentrations need to be entered.

No.	Name		Default		
		Field	Metric	Laboratory	
1	SALTCON	A real monotonically inc concentration for a given	None		
		lb/stb	kg/sm³	gm/scc	
Notes	<u>s:</u>				I

1) Each table is terminated by "/" including the last table; however, there is no "/" terminator for the keyword.

An alternative manner of entering the salt concentrations is by utilizing the PVTNUM region array by using the ADSALNOD keyword in the PROPS section.

Table 8.73: SALNODE Keyword Description

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

Example

Given three sets of relative permeability tables and four values on the PLYADSS keyword and two SALNODE data sets with four values of salt concentrations then the data should be entered as follows:

```
-- SETS SALT CONCENTRATION FOR POLYMER SOLUTION ADSORPTION
-- VIA PVTNUM ARRAY ALLOCATION
-- SALT
-- SALNODE

1.0
5.0
10.5
25.0 / PVTNUM TABLE NO. 01
1.0
3.0
7.5
15.0 / PVTNUM TABLE NO. 02
```

See also the ADSALNOD keyword.

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8.3.87 SCALECRS - SET END-POINT SCALING OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SCALECRS keyword sets the end-point scaling option to be either two-point or three-point scaling. This determines which end-points on the relative permeability curves are used for scaling based on the supplied end-point arrays (SGCR, SWCR, etc.).

No.	Name	Description	Default
1	SCALEOPT	SCALEOPT is a character string that sets the endpoint scaling option and should be set to either NO or YES:	NO
		NO: Activates two-point end-point scaling.	
		2) YES: Activates three-point end-point	

Notes:

1) The keyword is terminated by "/".

Table 8.74: SCALECRS Keyword Description

The end-point scaled for each option and the arrays used in the end-point scaling are summarized in the following table:

Option	Phases	Relative Permeability End-Point	Minimum Saturation End-Point	Middle Saturation End-Point	Maximum Saturation End-Point
Two-	Water	KRW	SWCR		SWU
Point	Oil	KRG	SGCR		SGU
	Oil-Water	KRORW	SOWCR		(1.0 – SWL - SGL)
	Oil-Gas	KRORG	SOGCR		(1.0 – SWL - SGL)
Three-	Water	KRW	SWCR	(I.0 – SOWCR - SGL)	SWU
Point	Oil	KRG	SGCR	(I.0 - SOGCR-SWL)	SGU
	Oil-Water	KRORW	SOWCR	(1.0 – SWCR - SGL)	(1.0 – SWL - SGL)
	Oil-Gas	KRORG	SOGCR	(1.0 – SGCR - SGL)	(1.0 – SWL - SGL)
		Tv	vo Phase Gas-Wa	ter Simulations	
	Water	KRW	SWCR	(1.0 - SGCR)	SWU
	Gas	KRG	SGCR	(1.0 -SWCR)	SGU

Table 8.75: End-Point Arrays Used in the End-Point Scaling Options

Example

- -

-- TWO-POINT END-POINT SCALING IS NO THREE POINT IS YES

-- SCALEOPT -- SCALECRS

YES

ES / SCALING OPTION

The above example activates three-point end-point scaling of the relative permeability curves.



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8.3.88 SDENSITY - Define the Miscible or Solvent Surface Gas Density

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SDENSITY keyword defines density at surface conditions of either the miscible injection gas for when the MISCIBLE option has been invoked in the RUNSPEC section, or the solvent for when the SOLVENT option has been invoked in the RUNSPEC section. This keyword must be invoked if either the MISCIBLE or SOLVENT options have been activated in the RUNSPEC section.

No.	Name		Description					
		Field Metric Laboratory						
I	SOLDEN	SOLDEN is a real po-	SOLDEN is a real positive number defining the density at surface conditions of either:					
		I) the miscible injuint invoked in the I						
		2) the solvent for the RUNSPEC						
		lb/ft³	kg/m³	gm/cc				

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.76: SDENSITY Keyword Description

In addition to this keyword, the surface density or gravity of the in-place natural gas must be entered using either the DENSITY or GRAVITY keywords.

Examples

The following shows the SDENSITY keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to one.

```
-- MIS-SOL

-- DENSITY

-- SDENSITY

0.04520 / MIS-SOL DENSITY
```

The next example shows the SDENSITY keyword for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to three.

```
-- MIS-SOL

-- DENSITY

-- SDENSITY

0.04520 / MIS-SOL DENSITY 1

0.05520 / MIS-SOL DENSITY 2

0.06420 / MIS-SOL DENSITY 3
```

There is no terminating "/" for this keyword.

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8.3.89 SGCR - END-POINT SCALING GRID CELL CRITICAL GAS SATURATIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SGCR defines the critical gas saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The critical gas saturation is defined as the maximum gas saturation for which the gas relative permeability is zero in a two-phase relative permeability table.

The keyword can be used for all grid types.

No.	Name		Description						
		Field	Field Metric Laboratory						
1	SGCR	values to each cell in the to the NX x NY x NZ pa	SGCR is an array of real numbers assigning the critical gas saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example $30*0.03$						
	dimensionless dimensionless dimensionless								

Notes:

- Note this the non-direction dependent version of the critical gas saturation used with the end-point scaling option. If directional end-point scaling has been activated then the SGCRX±, SGCRX± and SGCRX± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability 2)
- The keyword is terminated by "/". 3)

Table 8.77: SGCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SGCRX, SGCRY and SGCRZ instead of SGCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the nonreversible versions of the aforementioned arrays should be used, that is SGCRX, SGCRX-, SGCRY, SGCRY-, SGCRZ and SGCRZ-, instead of the SGCR keyword.

Example

```
DEFINE GRID BLOCK END-POINT SGCR DATA FOR ALL CELLS (FOR NX x NY x NZ = 300)
SGCR
  300*0.050
```

The above example defines a constant critical gas saturation of 0.05 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.90 SGCWMIS - MISCIBLE CRITICAL GAS VERSUS WATER SATURATION FUNCTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

SGCWMIS defines the dependency between the miscible critical gas saturation and the water saturation, for when the MISCIBLE keyword in the RUNSPEC section has been activated. The keyword can only be used with the MISCIBLE option and for when the oil, water and gas phases are active in the model.

No.	Name		Description Field Metric Laboratory				
		Field					
1	SWAT		real monotonically increa o and terminating atone,	S	None		
		dimensionless	dimensionless	dimensionless			
2	SGCMIS	that are greater than or	A columnar vector of real equal or increasing down the column values that are greater than or equal to zero and less than one, that define the corresponding miscible gas critical gas saturation for the corresponding water saturation SWAT.				
		dimensionless	dimensionless	dimensionless			

Notes:

- The keyword is followed by NTMISC tables as declared on the MISCIBLE keyword in the RUNSPEC section. I)
- Each table must contain a minimum of two rows and a maximum of NSMISC rows as declared on the 2) MISCIBLE keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword. 3)

Table 8.78: SGCWMIS Keyword Description

Example

MISCIBLE CRITICAL GAS VERSUS WATER SATURATION TABLE **SGCWMIS SWAT SGCRMIS FRAC** FRAC 0.0000 0.0000 0.2000 0.0300 1.0000 0.0300 / TABLE NO. 01 **SWAT SGCRMIS FRAC** FRAC 0.0000 0.0000 0.3000 0.0500 1.0000 0.0500 / TABLE NO. 02

The above example defines two miscible critical gas saturation versus water saturation tables assuming NTMISC equals two and NSMISC is greater than or equal to three on the MISCIBLE keyword in the RUNSPEC section.

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8.3.91 SGFN - Gas Saturation Tables (Format Type 2)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SGFN keyword defines the gas relative permeability and oil-gas capillary pressure data versus gas saturation tables for when gas is present in the input deck. This keyword should only be used if the gas is present in the run.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
1	SGAS		real monotonically increa o and terminating at on	S	None	
		dimensionless	dimensionless	dimensionless		
2	KRG	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the gas relative permeability.				
		dimensionless	dimensionless	dimensionless		
3	PCOG		tor of real values that are either equal or increasing down defines the oil-gas relative capillary pressure.			
		psia	bars	atm		

Notes:

- The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section. I)
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.79: SGFN Keyword Description

Example

- -GAS RELATIVE PERMEABILITY TABLES (SGFN) **SGFN SGAS** KRG PCG0 **FRAC PSIA** 1* 0.00 0.0000 0.0002 0.20 1* / TABLE NO. 01 0.85 0.4450 1* _____ 0.00 0.0000 1* 0.20 0.0002 0.0062 1* 0.30 0.45 0.0450 1* 1* 0.50 0.0707 0.60 1* 0.1412 0.70 0.2412 1* 0.4450 1* / TABLE NO. 02 0.85

The example defines two SGFN tables for when gas is present in the input deck.

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8.3.92 SGL - End-Point Scaling Grid Cell Connate Gas Saturations

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SGL defines the connate gas saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The connate gas saturation is defined as the minimum gas saturation in a two-phase gas relative permeability table.

The keyword can be used for all grid types.

No.	Name		Default						
		Field	Field Metric Laboratory						
1	SGL	to each cell in the model NX x NY x NZ paramet	of real numbers assigning the connate gas saturation values he model. The number of entries should correspond to the parameters on the DIMENS keyword. may be used, for example 30*0.03						
		dimensionless	permeability table.						

Notes:

- Note this the non-direction dependent version of the connate gas saturation array used with the end-point scaling option. If directional end-point scaling has been activated then the SGLX± , SGLX± and SGZ± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.80: SGL Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SGLX, SGLY and SGLZ instead of SGL. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SGLX, SGLY-, SGLY-, SGLZ and SGLZ-, instead of the SGL keyword.

Example

```
--
DEFINE GRID BLOCK END-POINT SGL DATA FOR ALL CELLS (FOR NX x NY x NZ = 300)
--
SGL
300*0.030
```

The above example defines a constant connate gas saturation of 0.03 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.93 SGOF - Gas-Oil Saturation Tables versus Gas (Format Type 1)

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Revision: Rev-I

Description

The SGOF keyword defines the oil and gas relative permeability and oil-gas capillary versus gas saturation tables for when oil and gas are present in the input deck. This keyword should only be used if the gas is present in the run.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	SGAS	A columnar vector of real monotonically increasing down the column values starting from zero and terminating at one, that defines the gas saturation.					
		dimensionless	dimensionless	dimensionless			
2	KRG	the column and that are	al values that are either ed greater than or equal to the gas relative permeabil	zero and less than or	None		
		dimensionless	dimensionless	dimensionless			
3	KRO	A columnar vector of real values that are either equal or decreasing down the column and that are greater than or equal to zero and less than or equal to one that defines the oil relative permeability with respect to gas and connate water saturation.					
		When water is active in the run, the first entry the column, that is at $krog(Sg = 0)$, must be the same as the first entry in the corresponding SWOF table, that is at $krow(So = I - Swco)$.					
		The last value in the colu	mn should be zero.				
		dimensionless	dimensionless	dimensionless			
4	PCOG		al values that are either ed he oil-gas relative capillary		None		
		psia	bars	atm			

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.81: SGOF Keyword Description

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Revision: Rev-I

Examp	le				
 SG0F	GAS-OIL F	RELATIVE PE	RMEABILITY	TABLES (SGOF)	
	SG FRAC	KRG	KROG	PCOG PSIA	
	0.00000 0.03000 0.80000	0.000000 0.000000 0.900000	0.90000 0.82500 0.00000	0.0000 0.0000 0.0000	/ TABLE No. 01
	0.00000 0.03000 0.04420 0.05850 0.07270 0.08700 0.10120 0.11550 0.12970 0.14390 0.15820 0.17240 0.18670 0.20090 0.21520 0.22940 0.24360 0.25790 0.27210 0.28640 0.3060 0.31480 0.31480 0.32910 0.34330 0.35760 0.31480 0.35760 0.37180 0.37180 0.4030 0.41450 0.42880 0.44300 0.45730 0.47150 0.48580	0.000000 0.000000 0.004200 0.048500 0.072700 0.097000 0.121200 0.145500 0.169700 0.193900 0.218200 0.242400 0.266700 0.290900 0.315200 0.339400 0.363600 0.387900 0.412100 0.46600 0.46600 0.599100 0.557600 0.557600 0.557600 0.66400 0.6630300 0.654500 0.678800 0.727300 0.727300 0.751500 0.775800	0.90000 0.82500 0.80000 0.775000 0.75000 0.72500 0.70000 0.67500 0.65000 0.62500 0.55000 0.555000 0.555000 0.47500 0.45000 0.42500 0.40000 0.47500 0.37500 0.37500 0.32500 0.32500 0.22500 0.22500 0.215000 0.17500 0.15000 0.17500 0.15000 0.17500 0.15000 0.15000 0.10000 0.07500 0.05000 0.05000	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	
	0.50000 0.80000	0.800000 0.900000	0.00000 0.00000	0.0000 0.0000	/ TABLE No. 02

The example defines two SGOF tables for use when oil, gas and water are present in the run.

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FLOW DOCUMENTATION MANUAL (2018-10)

8.3.94 SGU - END-POINT SCALING GRID CELL GAS SATURATION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SGU defines the maximum gas saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The maximum gas saturation is defined as the maximum gas saturation in a two-phase gas relative permeability table.

The keyword can be used for all grid types.

No.	Name		Description Field Metric Laboratory					
		Field						
I	SGU	SGU is an array of real numbers assigning the maximum gas saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.70						
	dimensionless dimensionless dimensionless							

Notes:

- Note this the non-direction dependent version of the maximum gas saturation array used with the end-point scaling option. If directional end-point scaling has been activated then the SGUX±, SGUX± and SGU± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.82: SGU Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SGUX, SGUY and SGUZ instead of SGU. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SGUX, SGUX-, SGUY, SGUY-, SGUZ and SGUZ-, instead of the SGU keyword.

Example

```
--
-- DEFINE GRID BLOCK END-POINT SGU DATA FOR ALL CELLS (FOR NX x NY x NZ = 300)
--
SGU
300*0.700
```

The above example defines a constant connate gas saturation of 0.70 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.95 SGWFN - Gas-Water Saturation Tables (Format Type 2)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SGWFN keyword defines the gas and water relative permeability and gas-water capillary pressure data versus gas saturation tables for when gas and water are present in the input deck. This keyword should only be used if the gas and water are present in the run.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
I	SGAS		real monotonically increa o and terminating at on		None	
		dimensionless	dimensionless	dimensionless		
2	KRG	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the gas relative permeability. Note that the first entry in the column must be zero.				
		dimensionless	dimensionless	dimensionless		
3	KRW	the column and that are equal to one that defines gas saturation.	al values that are either eq greater than or equal to s the water relative perm	zero and less than or	None	
		The last value in the colu				
		dimensionless	dimensionless	dimensionless		
4	PCGW		al values that are either ed he gas-water relative capil		None	
		psia	bars	atm		

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.83: SGWFN Keyword Description

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Revision: Rev-I

Examp	le				
 SGWFN	GAS-WATER	RELATIVE P	PERMEABILI	TY TABLES (SGWFN)	
	SG FRAC	KRG	KRW	PCOW PSIA	
	0.000000 0.200000 0.699099 0.700000	0.0000 0.0002 0.4973 1.0000	0.9000 0.7664 0.0000 0.0000	0.000000 0.000000 0.000000 0.000000	/ TABLE NO. 01
	0.000000 0.200000 0.245309 0.261989 0.303091 0.368269 0.435026 0.486387 0.522283 0.550683 0.575342 0.599076 0.621294 0.642171 0.658984 0.671123 0.679268	0.0000 0.0002 0.0004 0.0010 0.0044 0.0191 0.0519 0.0940 0.1339 0.1725 0.2115 0.2542 0.2991 0.3458 0.3868 0.4183 0.4403	0.9000 0.7664 0.7443 0.6907 0.5671 0.3962 0.2528 0.1643 0.1137 0.0803 0.0559 0.0367 0.0223 0.0120 0.0061 0.0030 0.0015	0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000 0.000000	, INDEL NO. CI
	0.684963 0.688893 0.692025 0.694641 0.696976 0.699099 0.700000	0.4562 0.4674 0.4765 0.4841 0.4910 0.4973 1.0000	0.0008 0.0004 0.0002 0.0001 0.0000 0.0000	0.000000 0.000000 0.000000 0.000000 0.000000	/ TABLE NO. 02

The example defines two SGWFN tables for use when oil, gas and water are present in the run.

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8.3.96 SHRATE - ACTIVATE AND DEFINE THE POLYMER SHEARING LOGARITHMIC

PARAMETERS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Description

This keyword activates the logarithm-based polymer shear thinning/thickening option and defines the shear rate constant. This keyword can only be used in conjunction with the PLYSHLOG in the PROPS section

This keyword is ignored by OPM Flow and has no effect on the simulation.

See the PLYSHEAR keyword for the alternative polymer shear thinning/thickening option that is implemented in OPM Flow.

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8.3.97 SLGOF - Gas-Oil Saturation Tables versus Gas (Format Type 1)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SLGOF keyword defines the oil and gas relative permeability and oil-gas capillary pressure versus liquid saturation tables for when oil and gas are present in the input deck. This keyword should only be used if both oil and gas are present in the run.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	SLIQ		real monotonically increa to and terminating at on		None		
			orrespond to residual liq be 1.0 to correspond to a				
	dimensionless	dimensionless	dimensionless				
2	KRG	the column and that are	vector of real values that are either equal or decreasing down and that are greater than or equal to zero and less than or that defines the gas relative permeability				
	dimensionless	dimensionless	dimensionless				
3	KRO	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the oil relative permeability with respect to gas and connate water saturation.					
		krog(Sg = 0), must be the	When water is active in the run, the last entry the column, that is at $rog(Sg = 0)$, must be the same as the first entry in the corresponding WOF table, that is at krow(So = I - Swco).				
		The first value in the colu	ımn should be zero.				
		dimensionless	dimensionless	dimensionless			
4	PCOG		al values that are either eq he oil-gas relative capillary		None		
		psia	bars	atm			

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.84: SLGOF Keyword Description

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Revision: Rev-I

Exampl	e				
 SL0F	GAS-0IL	RELATIVE PE	RMEABILIT	Y TABLES (SLGOF)	
	SLIQ FRAC	KRG	KROG	PCOG PSIA	
	0.30060 0.31480 0.32910 0.34330 0.35760 0.37180 0.38610 0.40030 0.41450 0.42880 0.44300 0.45730 0.47150 0.48580 0.50000 0.80000 0.30060 0.31480 0.32910 0.34330 0.35760 0.37180	0.55000 0.42500 0.35000 0.27500 0.25000 0.22500 0.20000 0.17500 0.15000 0.12500 0.05000 0.05000 0.02500 0.00000 0.02500 0.055000 0.055000 0.25000 0.25000 0.25000 0.25000	0.0000 0.2848 0.3091 0.4333 0.5576 0.5818 0.6061 0.6303 0.6545 0.6788 0.7030 0.7273 0.7515 0.7758 0.8000 0.9000 0.0000 0.2848 0.3091 0.4333 0.5576 0.5818	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	/ TABLE No. 01
	0.38610 0.40030 0.41450 0.42880 0.44300 0.45730 0.47150 0.48580 0.50000	0.20000 0.17500 0.15000 0.12500 0.10000 0.07500 0.05000 0.02500 0.00000	0.6061 0.6303 0.6545 0.6788 0.7030 0.7273 0.7515 0.7758 0.8000	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	
	0.80000	0.00000	0.9000	0.0000	/ TABLE No. 02

The example defines two SGOF tables for use when oil, gas and water are present in the run.

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8.3.98 SOF2 - OIL SATURATION TABLES WITH RESPECT TO GAS OR WATER (FORMAT Type 2)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SOF2 keyword defines the oil relative permeability versus oil saturation tables for when oil and gas or oil and water are present in the input deck. The keyword is also used to define the relative permeability of the miscible hydrocarbon phase in SOLVENT runs This keyword should only be used if the oil is present in

No.	Name		Description					
		Field	Metric	Laboratory	l			
I	SOIL	values starting from zero the hydrocarbon solvent For two phase runs the o SOLVENT option has be						
		dimensionless	1					
3	KRO	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the oil relative permeability with respect to gas and connate water saturation.						
	For two phase runs the oil relative permeability should be entered and for when the SOLVENT option has been activated in the RUNSPEC section the relative permeability of the miscible hydrocarbon phase with respect to water. The last value in the column should be zero.							
		dimensionless	İ					

Notes:

- The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section. I)
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the 2) TABDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.85: SOF2 Keyword Description

Not this keyword should only be used for when the SOLVENT keyword in the RUNSPEC section has been activated. It should not be use for two-phase oil-water runs.

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Example	9		
	OIL RELATIVE	PERMEABILITY TABLES (SOF2)	
S0F2			
	SOIL	KR0	
	FRAC	FRAC	
	0.00	0.000000	
	0.05	1.197e-5	
	0.10	0.000191	
	0.15	0.000969	
	0.20	0.003065	
	0.25	0.007483	
	0.30	0.015517	
	0.35	0.028747	
	0.40	0.049041	
	0.45 0.56	0.078555 0.119730	
	0.55	0.175297	
	0.60	0.248272	
	0.65	0.341961	
	0.70	0.459956	
	0.75	0.606134	
	0.80	0.784664	
	0.85	1.000000	/ TABLE NO. 01
	0.00	0.000000	
	0.05	1.197e-5	
	0.10	0.000191	
	0.15 0.20	0.000969 0.003065	
	0.25	0.007483	
	0.30	0.015517	
	0.35	0.028747	
	0.40	0.049041	
	0.45	0.078555	
	0.56	0.119730	
	0.55	0.175297	
	0.60	0.248272	
	0.65	0.341961	
	0.70	0.459956	
	0.75	0.606134	
	0.80	0.784664	/ TABLE NO. 02
	0.85	1.000000	/ IADLE NU. UZ

The example defines two SOF2 tables for when oil and gas or oil and water are present in the input deck.

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8.3.99 SOF3 - OIL SATURATION TABLES WITH RESPECT TO GAS AND WATER (FORMAT TYPE 2)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SOF3 keyword defines the oil relative permeability versus oil saturation tables for when oil, gas and water are present in the input deck. The keyword should only be used if oil, gas and water are present in the input deck.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	SOIL		real monotonically increa o and terminating at one, saturation.		None			
		The final entry should be	The final entry should be at the connate water saturation, that is 1- Swc.					
		dimensionless	dimensionless	dimensionless				
3	KROW	A columnar vector of reathe column and that are equal to one that define and water saturation. The first value in the column	None					
		dimensionless	dimensionless	dimensionless				
4	KROG	the column and that are equal to one that defines	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the oil relative permeability with respect to oil, gas and connate water saturation.					
		The first value in the column should be zero.						
	dimensionless dimensionless dimensionless							

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.86: SOF3 Keyword Description

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Revision: Rev-I

Example	е				
	OIL RELATIVE	PERMEABILITY	TABLES	(S0F3)	
S0F3					
30F3 	SOIL	KR0	KROG		
	FRAC	FRAC	FRAC		
	0.00		0.00000		
	0.05		0.00000		
	0.10		0.00000		
	0.15		0.00000		
	0.20 0.25		0.00000		
	0.30		0.05932		
	0.35		0.13158		
	0.40		0.21082		
	0.45		0.29960		
	0.56		0.40095		
	0.55	0.175297	0.51818		
	0.60	0.248272	0.65476		
	0.65	0.341961	0.81420		
	0.70		1.00000		
	0.75		1.00000		
	0.80		1.00000		/ TABLE NO. 4
	0.85	1.000000	1.00000		/ TABLE NO. 1
	0.00		0.00000		
	0.05		0.00000		
	0.10		0.00000		
	0.15		0.00000		
	0.20	0.003065	0.00000		
	0.25	0.007483	0.00000		
	0.30		0.05932		
	0.35		0.13158		
	0.40		0.21082		
	0.45		0.29960		
	0.56		0.40095		
	0.55 0.60		0.51818 0.65476		
	0.65		0.81420		
	0.70		1.00000		
	0.75		1.00000		
	0.80		1.00000		
	0.85	1.000000	1.00000		/ TABLE NO. 2

The example defines two SOF3 tables for when oil, gas and water are present in the input deck.

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8.3.100 SOGCR - END-POINT SCALING GRID CELL CRITICAL OIL SATURATION WITH RESPECT TO GAS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Description

SOGCR defines the critical oil saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The critical oil saturation with respect to gas is defined as the maximum oil saturation for which the oil relative permeability is zero in a two-phase gas-oil relative permeability table.

The keyword can be used for all grid types.

No.	Name		Default					
		Field						
I	SOGCR	with respect to gas value should correspond to the keyword.	SOGCR is an array of real numbers assigning the critical oil saturation with respect to gas values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.30 dimensionless dimensionless dimensionless dimensionless					
		dimensionless						

Notes:

- Note this the non-direction dependent version of the critical gas saturation used with the end-point scaling option. If directional end-point scaling has been activated then the SOGCRX±, SOGCRX± and SOGCRX± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability
- The keyword is terminated by "/".

Table 8.87: SOGCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SOGCRX, SOGCRY and SOGCRZ instead of SOGCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the nonreversible versions of the aforementioned arrays should be used, that is SOGCRX, SOGCRX, SOGCRY, SOGCRY-, SOGCRZ and SOGCRZ-, instead of the SOGCR keyword.

Example

DEFINE GRID BLOCK END-POINT SOGCR DATA FOR ALL CELLS (FOR NX x NY x NZ = 300) SOGCR 300*0.200

The above example defines a constant critical gas saturation of 0.20 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.101 SORWMIS - MISCIBLE RESIDUAL OIL VERSUS WATER SATURATION FUNCTIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SORWMIS defines the dependency between the miscible residual oil saturation and the water saturation, for when the MISCIBLE keyword in the RUNSPEC section has been activated. The keyword can only be used with the MISCIBLE option and for when the oil, water and gas phases are active in the model.

No.	Name		Description					
		Field	Metric	Laboratory				
I	SWAT		columnar vector of real monotonically increasing down the column alues starting from zero and terminating at one, that defines the water attraction.					
		dimensionless	dimensionless	dimensionless				
2	SORMIS	that are greater than or	A columnar vector of real equal or increasing down the column values that are greater than or equal to zero and less than one, that define the corresponding miscible residual oil saturation for the corresponding water saturation SWAT.					
		dimensionless	dimensionless	dimensionless				

Notes:

- 1) The keyword is followed by NTMISC tables as declared on the MISCIBLE keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSMISC rows as declared on the MISCIBLE keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.88: SORWMIS Keyword Description

Example

MISCIBLE RESIDUAL OIL VERSUS WATER SATURATION TABLE **SORWMIS SWAT SORWMIS FRAC** FRAC 0.0000 0.0000 0.2000 0.0000 1.0000 0.0000 / TABLE NO. 01 **SWAT SORWMIS FRAC FRAC** 0.0000 0.0000 0.3000 0.1000 0.7500 0.1500 / TABLE NO. 02

The above example defines two miscible residual oil versus water saturation tables assuming NTMISC equals two and NSMISC is greater than or equal to three on the MISCIBLE keyword in the RUNSPEC section.

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8.3.102 SOWCR - END-POINT SCALING GRID CELL CRITICAL OIL SATURATION WITH RESPECT TO WATER

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

SOWCR defines the critical oil saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The critical oil saturation with respect to water is defined as the maximum oil saturation for which the oil relative permeability is zero in a two-phase oil-water relative permeability table.

The keyword can be used for all grid types.

No.	Name		Default		
		Field			
I	SOWCR	with respect to water v	real numbers assigning the alues to each cell in the nd to the NX x NY x led, for example 30*0.30	model. The number of	Taken from cell allocated relative permeability table.
		dimensionless	dimensionless	dimensionless	

Notes:

- Note this the non-direction dependent version of the critical gas saturation used with the end-point scaling option. If directional end-point scaling has been activated then the SOWCRX± , SOWCRX± and SOWCRX± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability
- The keyword is terminated by "/".

Table 8.89: SOGCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SOWCRX, SOWCRY and SOWCRZ instead of SOWCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SOWCRX, SOWCRX-, SOWCRY, SOWCRY-, SOWCRZ and SOWCRZ-, instead of the SOWCR keyword.

Example

DEFINE GRID BLOCK END-POINT SOWCR DATA FOR ALL CELLS (FOR NX x NY x NZ = 300) SOWCR 300*0.200

The above example defines a constant critical gas saturation of 0.20 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.103 SPECHEAT - Define the Specific Heat of Oil, Water and Gas

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SPECHEAT defines the specific heat of the oil, water and gas phases for various PVT regions in the model for when the THERMAL option has been activated in the RUNSPEC section. The number of SPECHEAT vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the SPECHEAT data sets to different grid blocks in the model is done via the PVTNUM keyword in the REGION section.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
I	TEMP	A columnar vector of real monotonically increasing down the column values that define the temperature for the corresponding oil, water and gas specific heat values.				
		°F	°C	°C		
2	OILSHEAT	OILSHEAT is a columnar vector of positive real numbers defining the specific heat of oil at the corresponding temperature, TEMP.				
		Btu/lb/°R	kJ/kg/K	J/gm/K		
3	WATSHEAT	WATSHEAT is a columnar vector of positive real numbers defining the specific heat of water at the corresponding temperature, TEMP.				
		Btu/lb/°R	kJ/kg/K	J/gm/K		
4	GASSHEAT	GASHEAT is a columnar vector of positive real numbers defining the specific heat of gas at the corresponding temperature, TEMP.				
		Btu/lb/°R	kJ/kg/K	J/gm/K		

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.90: SPECHEAT Keyword Description

See also the SPECROCK keyword to define the reservoir rock specific heat.

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Revision: Rev-I

Example

The example below defines three fluid phases specific heat versus temperature tables assuming NTPVT equals three and NPPVT is greater than or equal to two on the TABDIMS keyword in the RUNSPEC section.

	SPECIFIC H	HEAT OF OIL,	WATER AND	GAS TABLE	
SPECHEAT					
	TEMP	SPECHEAT	SPECHEAT	SPECHEAT	
		OIL	WATER	GAS	
	0.000	0.5000	1.5000	0.5000	
	250.000	0.5000	1.5000	0.5000	/ TABLE NO. 01
	TEMP	SPECHEAT	SPECHEAT	SPECHEAT	
		OIL	WATER	GAS	
	0.000	0.5500	1.5000	0.5000	
	260.000	0.5500	1.5000	0.5000	/ TABLE NO. 02
	TEMP	SPECHEAT	SPECHEAT	SPECHEAT	
		OIL	WATER	GAS	
	0.000	0.5500	1.5500	0.5000	
	270.000	0.6000	1.5500	0.5000	/ TABLE NO. 03

There is no terminating "/" for this keyword.

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8.3.104 SPECROCK - DEFINE THE SPECIFIC HEAT OF THE RESERVOIR ROCK

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SPECROCK defines the specific heat of the reservoir rock for various PVT regions in the model for when the THERMAL option has been activated in the RUNSPEC section. The number of SPECROCK vector data sets is defined by the NTSFUN parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the SPECROCK data sets to different grid blocks in the model is done via the SATNUM keyword in the REGION section.

This keyword can only be used if OPM's Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	TEMP		real monotonically increa emperature for the corre		None
		°F	°C	°C	
2	ROCKHEAT	ROCKHEAT is a column specific heat of the rock a	nar vector of positive rea		None
		Btu/ft³/°R	kJ/m³/K	J/cc/K	

Notes:

- The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section. I)
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.91: SPECROCK Keyword Description

See also the SPECHEAT keyword to define the specif heat relationships for the oil, water and gas phases.

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Revision: Rev-I

Example

The example below defines three rock specific heat versus temperature tables assuming NTSFUN equals three and NSSFUN is greater than or equal to two on the TABDIMS keyword in the RUNSPEC section.

SPECIFIC HEAT OF ROCK		
TEMP	SPECHEAT	
	ROCK	
0.000	20.000	
250.000	20.000	
0.000	21.000	
260.000	21.000	
0.000	23.000	
270.000	23.000	
	TEMP 0.000 250.000 0.000 260.000 0.000	

There is no terminating "/" for this keyword.

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8.3.105 SSFN - Solvent and Gas Relative Permeability Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SSFN keyword defines the miscible <u>normalized</u> relative permeability tables for when the SOLVENT option has been activated in the RUNSPEC section using the respective keyword. The MISCIBLE keyword invokes a three component formulation (oil, water and solvent gas or an oil, water and solvent oil). Why the SOLVENT keyword results in a four component model (oil, water and gas plus a solvent). This keyword should only be used if the SOLVENT options have been activated.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
I	SGAS	A columnar vector of real monotonically increasing down the column values starting from zero and terminating at one, that defines the gas plus solvent saturation ration which is defined as either:				
	$\overline{(S_g)}$	$\frac{S_g}{+S_s}$ or $\frac{S_s}{\left(S_g + \right)}$	$\overline{S_s}$			
		Where Sg is the gas satu	ration and Ss is the solven	t saturation.		
		dimensionless	dimensionless	dimensionless		
2	KRG ^t	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the gas relative permeability. The resulting gas relative permeability is calculated from: $k_{rg} \ = \ k_{rgt} \big(S_g \ + \ S_s \big) k_{rg}^{\ t}$ where krgt is the data in this column and krgt is the gas relative permeability from the SGFN keyword				
		dimensionless	dimensionless	dimensionless		
3	KRS ^t	the column and that are	al values that are either ed e greater than or equal to es the solvent relative per ility is calculated from:	zero and less than or	None	
		k_{rs}	$= k_{rgt}(S_g + S_s)k_s$	t rs		
		where krSt is the data permeability from the SG	in this column and kr GFN keyword	gt is the gas relative		
		dimensionless	dimensionless	dimensionless		

Notes:

- The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.92: SSFN Keyword Description

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Revision: Rev-I

Exampl	е					
 SSFN	SOLVENT R	ELATIVE PERM	BEABILITY TABLES			
 	SGAS FRAC	KRGT	KRST			
	0.0000 1.0000	0.0000 1.0000	1.0000	/	TABLE NO. 0	1
	0.0000 0.2000 0.4000 0.6000 0.8000 1.0000	0.0000 0.2000 0.3000 0.4000 0.5000	0.0000 0.3000 0.5000 0.7000 0.7500	/	TABLE NO. 0	12

The above example defines two SSFN tables for use with the MISCIBLE and SOLVENT options.

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8.3.106 SWCR - END-POINT SCALING GRID CELL CRITICAL WATER SATURATION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SWCR defines the critical water saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The critical water saturation is defined as the maximum water saturation for which the water relative permeability is zero in a two-phase relative permeability table.

The keyword can be used for all grid types.

No.	Name		Default					
		Field	Metric	Laboratory	-			
1	SWCR	values to each cell in the to the NX x NY x NZ pa	SWCR is an array of real numbers assigning the critical water saturation values to each cell in the model. The number of entries should correspond to the NX \times NY \times NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.20					
	dimensionless dimensionless dimensionless							

Notes:

- Note this the non-direction dependent version of the critical water saturation used with the end-point scaling option. If directional end-point scaling has been activated then the SWCRX±, SWCRX± and SWCRX± series of keyword should be used.
- 2) If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.93: SWCR Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWCRX, SWCRY and SWCRZ instead of SWCR. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWCRX, SWCRX-, SWCRY, SWCRY-, SWCRZ and SWCRZ-, instead of the SWCR keyword.

Example

```
-- DEFINE GRID BLOCK END-POINT SWCR DATA FOR ALL CELLS (FOR NX x NY x NZ = 300)
-- SWCR
300*0.200
```

The above example defines a constant critical water saturation of 0.20 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.107 SWFN - WATER SATURATION TABLES (FORMAT TYPE 2)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SWFN keyword defines the water relative permeability and water-oil capillary pressure data versus water saturation tables for when water is present in the input deck. This keyword should only be used if water is present in the run.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	SWAT	A columnar vector of revalues starting from zero saturation.	None				
		dimensionless	dimensionless	dimensionless			
2	KRW	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the water relative permeability with respect to gas saturation. The first value in the column should be zero.					
		dimensionless	dimensionless	dimensionless			
3	PCWO	A columnar vector of real values that are either equal or increasing down the column that defines the water-oil relative capillary pressure. If the SWATINIT keyword has been used to initialize the model then					
		columnar vector has to b					
		psia	bars	atm			

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.94: SWFN Keyword Description

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0.80

0.85

0.90

0.95

1.00

Example WATER RELATIVE PERMEABILITY TABLES (SWFN) SWFN KRW **PCOW** - -**SWAT** - -**FRAC FRAC PSIA** ____ 0.15 0.00000 1* 1* 0.20 6.25e-6 1* 0.25 0.00010 0.30 0.00050 1* 1* 0.00160 0.35 0.40 0.00390 1* 0.45 0.00810 1* 1* 0.50 0.01500 1* 0.55 0.02560 1* 0.60 0.04100 1* 0.06250 0.65 0.70 0.09150 1* 1* 0.75 0.12960 1* 0.80 0.17850 0.85 0.24010 1* 0.90 0.31640 0.95 0.40960 1* 1.00 0.52200 1* / TABLE NO. 1 _ _ _ _ _ _ _ _ -----_ _ _ _ _ 1* 0.15 0.00000 1* 0.20 6.25e-6 1* 0.00010 0.25 0.30 0.00050 1* 1* 0.35 0.00160 1* 0.40 0.00390 0.45 0.00810 1* 1* 0.50 0.01500 0.55 0.02560 1* 0.60 0.04100 1* 1* 0.65 0.06250 1* 0.70 0.09150 1* 0.75 0.12960

Revision: Rev-I

The example defines two SWFN tables for use when water is present in the run. In the tables the water-oil capillary pressure data has been defaulted with "I*" and will be set to zero as there are no other values for the water-oil capillary pressure columns.

/ TABLE NO. 2

1*

1* 1*

1*

0.17850

0.24010

0.31640

0.40960

0.52200

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8.3.108 SWL - End-Point Scaling Grid Cell Connate Water Saturation

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SWL defines the connate water saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The connate water saturation is defined as the minimum water saturation in a two-phase water relative permeability table.

The keyword can be used for all grid types.

No.	Name		Default					
		Field	Metric	Laboratory				
I	SWL	values to each cell in the to the NX x NY x NZ pa	SWL is an array of real numbers assigning the connate water saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.15					
dimensionless dimensionless dimens					permeability table.			

Notes:

- Note this the non-direction dependent version of the connate gas saturation array used with the end-point scaling option. If directional end-point scaling has been activated then the SWLX±, SWLX± and SWZ± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.95: SWL Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWLX, SWLY and SWLZ instead of SWL. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWLX, SWLY-, SWLY-, SWLY-, SWLZ and SWLZ-, instead of the SWL keyword.

Example

```
--
-- DEFINE GRID BLOCK END-POINT SWL DATA FOR ALL CELLS (FOR NX x NY x NZ = 300)
--
SWL
300*0.150
```

The above example defines a constant connate water saturation of 0.15 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.109 SWOF - WATER-OIL SATURATION TABLES (FORMAT TYPE 1)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SWOF keyword defines the water and oil relative permeability and water-oil capillary pressure data versus water saturation tables for when water and oil are present in the input deck. This keyword should only be used if water and oil present in the run.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
I	SWAT	A columnar vector of real monotonically increasing down the column values starting from zero and terminating at one, that defines the water saturation.				
		The first entry is the coshould be 1.0.	onnate water saturation S	Swc and the last entry		
		dimensionless	dimensionless	dimensionless		
2	KRW	A columnar vector of real values that are either equal or increasing down the column and that are greater than or equal to zero and less than or equal to one that defines the water relative permeability with respect to gas saturation. The first value in the column should be zero.				
		dimensionless	dimensionless	dimensionless		
3	KRO	A columnar vector of real values that are either equal or decreasing down the column and that are greater than or equal to zero and less than or equal to one that defines the oil relative permeability with respect to oil and water saturation.				
		When gas is active in the run, the first entry the column, that is at $krow(So = I-Swc)$, must be the same as the first entry in the corresponding SGOF or SLGOF table, that is at $krog(Sg = 0)$.				
		The first value in the colu	ımn should be one.			
		dimensionless	dimensionless	dimensionless		
4	PCWO	A columnar vector of real values that are either equal or increasing down the column that defines the water-oil relative capillary pressure.				
		If the SWATINIT keyword has been used to initialize the model then columnar vector has to be strictly monotonically increasing.				
		psia	bars	atm		

Notes:

- 1) The keyword is followed by NTSFUN tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NSSFUN rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.96: SWOF Keyword Description

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Revision: Rev-I

Exampl	е				
	WATER-OIL	RELATIVE F	PERMEABILI	ΓΥ TABLES (SWOF)	
 SWOF					
	SWAT	KRW	KROW	PCOW	
	FRAC			PSIA	
	0.200000	0.0000	0.9000	0.000000	
	0.238616	0.0002	0.7664	0.000000	
	0.245309	0.0004	0.7443	0.000000	
	0.261989 0.303091	0.0010 0.0044	0.6907	0.000000	
	0.368269	0.0191	0.5671 0.3962	0.000000 0.000000	
	0.435026	0.0519	0.2528	0.000000	
	0.486387	0.0940	0.1643	0.000000	
	0.522283	0.1339	0.1137	0.000000	
	0.550683	0.1725	0.0803	0.000000	
	0.575342	0.2115	0.0559	0.000000	
	0.599076	0.2542	0.0367	0.000000	
	0.621294	0.2991	0.0223	0.000000	
	0.642171	0.3458	0.0120	0.000000	
	0.658984	0.3868	0.0061	0.000000	
	0.671123	0.4183	0.0030	0.000000	
	0.679268	0.4403	0.0015	0.000000	
	0.684963	0.4562	0.0008	0.000000	
	0.688893	0.4674	0.0004	0.000000	
	0.692025	0.4765	0.0002	0.000000	
	0.694641	0.4841	0.0001	0.000000	
	0.696976	0.4910	0.0000	0.000000	
	0.699099	0.4973	0.0000	0.000000	
	0.700000 1.000000	0.5000 0.9000	0.0000 0.0000	0.000000 0.000000	/ TABLE NO. 01
	1.000000	0.9000	0.0000	0.000000	/ TABLE NO. 01
	0.200000	0.0000	0.9000	0.000000	
	0.238616	0.0002	0.7664	0.000000	
	0.245309	0.0004	0.7443	0.000000	
	0.261989	0.0010	0.6907	0.000000	
	0.303091	0.0044	0.5671	0.000000	
	0.368269	0.0191	0.3962	0.000000	
	0.435026	0.0519	0.2528	0.000000	
	0.486387	0.0940	0.1643	0.000000	
	0.522283	0.1339	0.1137	0.000000	
	0.550683	0.1725	0.0803	0.000000	
	0.575342	0.2115	0.0559	0.000000	
	0.599076	0.2542	0.0367	0.000000	
	0.621294	0.2991	0.0223	0.000000	
	0.642171	0.3458	0.0120	0.000000	
	0.658984	0.3868	0.0061	0.000000	
	0.671123 0.679268	0.4183	0.0030 0.0015	0.000000	
	0.684963	0.4403 0.4562	0.0013	0.000000 0.000000	
	0.688893	0.4502	0.0004	0.000000	
	0.692025	0.4765	0.0004	0.000000	
	0.694641	0.4841	0.0002	0.000000	
	0.696976	0.4910	0.0000	0.000000	
	0.699099	0.4973	0.0000	0.000000	
	0.700000	0.5000	0.0000	0.000000	
	1.000000	0.9000	0.0000	0.000000	/ TABLE NO. 01
The exami	ale defines two	SWFN table	s for use wh	en water and oil are r	present in the run. In the table

The example defines two SWFN tables for use when water and oil are present in the run. In the tables the water-oil capillary pressure data has been set to zero.

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8.3.110 SWU - END-POINT SCALING GRID CELL GAS SATURATION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

SWU defines the maximum water saturation for all the cells in the model via an array when the end-point scaling option has been invoked via the ENDSCALE keyword in the RUNSPEC section. The maximum water saturation is defined as the maximum water saturation in a two-phase water relative permeability table.

The keyword can be used for all grid types.

No.	Name		Default					
		Field						
1	SWU	values to each cell in the to the NX x NY x NZ pa	SWU is an array of real numbers assigning the maximum water saturation values to each cell in the model. The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword. Repeat counts may be used, for example 30*0.70					
		dimensionless	dimensionless	dimensionless	table.			

Notes:

- Note this the non-direction dependent version of the maximum water saturation array used with the endpoint scaling option. If directional end-point scaling has been activated then the SWUX±, SWUX± and SWU± series of keyword should be used.
- If the value for a cell has been defaulted then OPM Flow users the value from the cell's relative permeability table.
- 3) The keyword is terminated by "/".

Table 8.97: SWU Keyword Description

End-point scaling allows the entered relative permeability functions to be re-scaled based on the saturation end-points allocated to each cell by the SWL, SWCR, SWU, SGL, SGCR, SGU, SOWCR, and SOGCR saturation grid arrays for the saturation end-points, and the KRG, KROG, KROW and KRW relative permeability grid cell arrays for the relative permeability end-point data. In addition end-point scaling may be directional dependent in which case the directional dependent versions of the aforementioned arrays should be used, that is SWUX, SWUY and SWUZ instead of SWU. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is SWUX, SWUX-, SWUY-, SWUY and SWUZ-, instead of the SWU keyword.

Example

```
--
-- DEFINE GRID BLOCK END-POINT SWU DATA FOR ALL CELLS (FOR NX x NY x NZ = 300)
--
SWU
300*0.700
```

The above example defines a constant connate gas saturation of 0.70 to all 300 cells in the model as defined by the DIMENS keyword in the RUNSPEC section.

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8.3.111 THERMEXI - Define Component Thermal Expansion Coefficients

RUNSPEC GRIE) EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
--------------	--------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

THERMEXI defines the thermal expansion coefficients for all components in the model for when the EOS and THERMAL options have been invoked by the EOS and THERMAL keywords in the RUNSPEC section.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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8.3.112 TLMIXPAR - Define the Miscible Todd-Longstaff Mixing Parameters

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The TLMIXPAR keyword defines the Todd-Longstaff⁵³ mixing parameters, for when either the miscible or solvent options have been activated by the MISCIBLE or SOLVENT keyword in the RUNSPEC section. This keyword must be present in the input deck if the MISCIBLE or SOLVENT keywords have been activated.

Note that If the POLYMER option has been activated by the POLYMER keyword in the RUNSPEC section, then this keyword is ignored and the mixing parameters are taken from the PLMIXPAR keyword instead.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	TLMVIS	A real positive value that is greater than or equal to zero and less than or equal to one, that defines the viscosity Todd-Longstaff mixing parameter for each miscibility region.						
		dimensionless	dimensionless	dimensionless	None			
2	TLMDEN		A real positive value that is greater than or equal to zero and less than or equal to one, that defines the density Todd-Longstaff mixing parameter for each miscibility region.					
		dimensionless	entered for TLMVIS					

Notes:

- The keyword is followed by NTMISC tables as declared on the MISCIBLE keyword in the RUNSPEC section, for each rock region. There should be only one row per table.
- 2) Each entry is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.98:TLMIXPAR Keyword Description

Example

TODD-LONGSTAFF MIXING PARAMETERS - -**TLMIXPAR** TLM TIM VISCOS **DENSITY** _____ _____ 0.3500 0.3500 / TABLE NO. 01 0.2500 1* / TABLE NO. 02 0.7500 / TABLE NO. 03 0.6500

The above example defines three Todd-Longstaff mixing parameter data sets, based on the NTMISC variable on the MISCIBLE keyword in the RUNSPEC section being equal to three.

Todd, M. and Longstaff, W. "The Development, Testing and Application of a Numerical Simulator for Predicting Miscible Flood Performance," paper SPE 3484, Journal of Canadian Petroleum Technology (1972) 24, No. 7, 874-882.



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8.3.113 TOLCRIT - Define The Critical Saturation Tolerance

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Critical fluid saturations are determine from the relative permeability tables, that is the last saturation in a relative permeability table where the relative permeability of a phase is set equal to zero. Since floating-point numbers (as implemented in computers) are never exact, one cannot compare floating point numbers for exact equality. Thus, this keywords defines a value below which is considered equivalent to zero in determining the critical saturation for a phase.

No.	Name		Description		Default				
		Field Metric Laboratory							
I	TOLCRIT	TOLCRIT is a real positive number greater than zero and less than one that defines the critical saturation tolerance used to determine the critical saturation of a fluid in the relative permeability tables.							
	The default value of 1 \times 10 ⁻⁶ means that saturation values less than this value will be treated as being equal to zero.								
		dimensionless							

I) The keyword is terminated by "/".

Table 8.99:TOLCRIT Keyword Description

See also section 8.2.2 Saturation Tables (Relative Permeability and Capillary Pressure Tables) for a description of the relative permeability tables and the various end-point definitions, including oil, water and gas critical saturations.

Example

```
--- SET THE CRITICAL SATURATION TOLERANCE
--
TOLCRIT
1.0E-6
```

The above example defines the critical saturation tolerance to be the default value of 1×10^{-6} .

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8.3.114 TRACER - Define Passive Tracer Variables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The TRACER keyword defines a series of passive tracers that are associated with a phase (oil, water, or gas) in the model. The maximum number of tracers for each phase are declared on the TRACER keyword in the RUNSPEC section.

No.	Name		Description		Default		
		Field	Metric	Laboratory	_		
1	NAME	A three letter character	string defining the tracer's	name.	None		
			ames beginning with the le g issues in post-processing				
2	PHASE		A three letter character string that defines the tracer given by NAME to a particular fluid phase. The character should be set to OIL, WAT or GAS.				
3	UNITS	The units for the tracer model.	This should be the sam	e as the PHASE in the			
		Liquid: stb	Liquid: sm³	Liquid: scc	Same as the phases in		
		Gas: Mscf	Gas: sm³	Gas: scc	the model		
4	SOLPHASE	A three or four letter of solution phase. The char MULT. Note that SOLPHASE of option has been activate section.	None				
5	KPNUM		used with the partitioned and TRACERKM keyword		None		
			y needs to be defined if with the PARTTRAC key				
6	PARPHASE	calculation for when th	string defining the phase e MULT option has bee e set to OIL, WAT, GAS o	n for SOLPHASE. The	None		
		Note that PARPHASE only needs to be defined if the partitioned tracer option has been activate with the PARTTRAC keyword in the RUNSPEC section.					

1) Each record (or row) should be terminated by a "/" and the keyword is terminated by "/".

Table 8.100:TRACER Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

'WAT'

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Example

'Iw2'

Revision: Rev-I

/ WAT INJECTOR 2

```
DEFINE TRACER NAMES
         TRACER
                   TRACER
         NAME
                   PHASE
--
                   _____
TRACER
        'IGS'
                   'GAS'
                                                               / GAS INJECTOR
        'DGS'
                   'GAS'
                                                               / DISOLVED GAS
        'IW1'
                   'WAT'
                                                               / WAT INJECTOR 1
```

The above example defines four passive tracers one for a gas injection well, one for tracking the dissolved gas, and two to track the injected water from two water injection wells.

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8.3.115 TREF - Define Component Fluid Densities Reference Temperatures

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

TREF defines the fluid component densities in-situ reference temperatures in the compositional commercial simulator or the live oil component in-situ reference temperature in the "black-oil" commercial simulator with the THERMAL option activated via the THERMAL keyword in the RUNSPEC section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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8.3.116 TREFS - Define Component Fluid Densities Reference Temperature at Surface

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

TREFS defines the fluid component densities surface reference temperatures in the compositional commercial simulator

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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8.3.117 VISCREF - Define Viscosity-Temperature Reference Conditions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

VISCREF defines the reference conditions for the viscosity-temperature tables, GASVISCT, OILVISCT and WATVISCT, for when the thermal option has been activated by THERMAL keyword in the RUNSPEC section. This keyword can only be used if the thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name	Description					
		Field	Metric	Laboratory			
I	PRES	PRES is a real positive viscosity and temperature					
		psia	barsa	atma	None		
2	RS	•	RS is a real positive number defining the reference gas-oil ratio for when the model contains gas dissolved as activated by the DISGAS keyword in the RUNSPEC section				
		Mscf/stb	sm³/sm³	scc/scc	None		
3	API	has been invoked by the	API is a real number defining the oil API for when the API tracking option has been invoked by the API keyword in the RUNSPEC section. Note that OPM Flow does not support API tracking, and therefore this variable is ignored.				
		°API	°API	°API	None		

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.101:VISCREF Keyword Description

OPM Flow currently does not support API tracking and therefore item (3) of this keyword is ignored. See also the OILVISCT, GASVISCT and WATVISCT keywords in the PROPS section.

Example

The following example shows the VISCREF keyword for when the thermal option has been activated by the TEMP keyword in the RUNSPEC section and for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to five.

```
- -
         REF
                     REF
                                REF
- -
         PRESSURE
                     GOR
                                API
                     _____
VISCREF
          3000.0
                     0.500
                                                               / TABLE NO. 01
                     0.550
                                                               / TABLE NO. 02
          3200.0
          3300.0
                     0.580
                                                               / TABLE NO. 03
          3400.0
                     0.620
                                                                 TABLE NO. 04
          3500.0
                     0.625
                                                               / TABLE NO. 05
```

There is no terminating "/" for this keyword.

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8.3.118 WATDENT - Define Water Density Temperature Coefficients

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

WATDENT defines the water density as a function of temperature coefficients for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC. The reference pressure for this table is given by the VISCREF keyword in the PROPS section.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description	Description				
		Field	Metric	Laboratory				
I	TEMP	reference temperature		greater than zero that defines the absolute with TEXP1 and TEXP2 to estimate the aspect to temperature.				
		°R	°K	°K				
		527.67	293.15	293.15				
2	TEXPI	TEXPI is a real positive value greater than zero that defines the water thermal expansion coefficient of the first order.						
		I/°R	I/°K	I/°K				
		1.67 x 10 ⁻⁴	3.0×10^{-4}	3.0 x 10 ⁻⁴				
3	TEXP2	TEXP2 is a real positive thermal expansion coeffic			Defined			
		I/°R²	I/°K²	I/°K²				
		9.26 x 10 ⁻⁷	3.0×10^{-6}	3.0 × 10 ⁻⁶				

Notes:

- 1) The keyword is followed by NTPVT records as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each data set is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.102:WATDENT Keyword Description

Example

The following example shows the WATDENT keyword using the default values, for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC section and for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to two.

```
WATER DENSITY TEMPERATURE COEFFICIENTS (OPM FLOW EXTENSION KEYWORD)
- -
- -
         WATER
                     DENSITY
                                DENSITY
- -
         TEMP
                     COEFF1
                                C0EFF2
WATDENT
                     1*
                                1*
         1*
                                                               / TABLE NO. 01
                                                               / TABLE NO. 02
```

There is no terminating "/" for this keyword.

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8.3.119 WATVISCT - Define Water Viscosity versus Temperature Functions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

WATVISCT defines the water viscosity as a function of temperature for when thermal option has been activated by the THERMAL keywords in the RUNSPEC. The reference pressure for this table is given by the VISCREF keyword in the PROPS section.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description					
		Field	Metric	Laboratory				
I	TEMP	A columnar vector of revalues that defines the te	None					
		°F	°C	°C				
2	VIS		al increasing down the col		None			
	VIS should be given at the reference pressure defined by the PRS variable on the VISCREF keyword.							
		сР	сР	cP				

Notes:

- 1) The keyword is followed by NTPVT tables as declared on the TABDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NPPVT rows as declared on the TABDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 8.103:WATVISCT Keyword Description

Example

The following example shows the WATVISCT keyword for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC section and for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set equal to one.

```
WATER VISCOSITY VERSUS TEMPERATURE TABLES
         WATER
                    WATER
         TEMP
                    VISC
WATVISCT
           100.0
                    0.500
           110.0
                    0.550
           120.0
                    0.580
           150.0
                    0.620
           165.0
                    0.625
                                                              / TABLE NO. 01
```

There is no terminating "/" for this keyword.

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9 REGIONS SECTION

9.1 Introduction

The REGIONS section defines how various properties in the PROPS and SOLUTION sections are allocated to individual cells within the model, as well as defining various fluid in-place reporting regions. This is accomplished by assigning an integer value to each cell that represents the data set of the property to be assigned to the grid block.

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9.2 DATA REQUIREMENTS

OPM Flow, like most numerical modeling software, users a default value of one for the various region arrays and thus if there is only one PVT data set for example, then there is no need to define the region array associated with allocating the PVT tables (PVTNUM), as all cells will be allocated PVT table number one. However, if there more than one PVT table entered in the PROPS section and PVTNUM is not defined in the model then PVT tables greater than one will not be used and there will be no warning message indicting the fact.

Properties Section	Property Allocation	REGIONS Section Keyword
SOLUTION	Equilibrium region allocation based on the EQUIL keyword records.	EQLNUM
REGION	Fluid In-Place reporting via the FIPNUM array that divides the model into different fluid in-place reporting regions.	FIPNUM
PROPS	PVT table allocation of the DENSITY, PVDG, PVDO, PVTG, PVTO, PVCO, PVTW and ROCK tables	PVTNUM
PROPS	Saturation (relative permeability) table allocation of the SGFN, SWFN, SOF2, SOF3, SGOF, and SWOF tables.	SATNUM
	Function Specific Regions	
PROPS	ENPTVD and ENKRVD versus depth table allocation for when ENDSCALE option has been activated in the RUNSPEC section.	ENDNUM
PROPS	Imbibition saturation table allocation of the SWFN, SOF2, SOF3 or SWOF imbibition tables.	IBMNUM
PROPS	Miscible regions based on the TLMIXPAR records when the MISCIBLE or SOLVENT keywords have been activated in the RUNSPEC section.	MISNUM
PROPS	Rock compaction table allocation for when the ROCKCOMP keyword as been activated in the RUNSPEC section, that allocates the ROCKTAB series of tables to a cell.	ROCKNUM
PROPS	Surfactant saturation (relative permeability) tables allocation allocating the SWFN, SOF2, SOF3 or SWOF as miscible tables.	SURFNUM
PROPS	High salinity water wet saturation table allocation using the high salinity water wet saturation SWFN and SOFN tables.	SURFWNUM

Notes:

- I) Only EQLNUM, FIPNUM, PVTNUM, SATNUM, IMBUM and MISNUM are available in OPM Flow.
- Note that is common to set the FIPNUM array to be equal to the EQLNUM to have fluid in-place
 reporting for each equilibrium region, this can be done by using the COPY keyword to copy the EQLNUM
 array to the FIPNUM array.

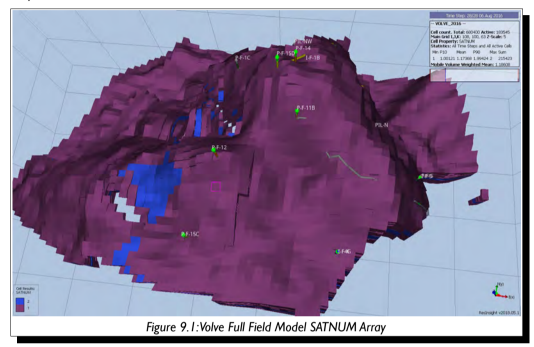
Table 9.1: REGION Section Allocation Array Summary

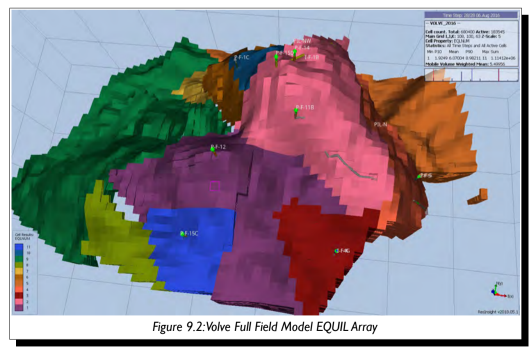
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The allocation is based on a complete property data set, that is all the property data associated with a given data set is allocated to the cell. For example, if the fluid properties for the model are the same, (for example, PVTO and PVDG keyword data) but the rock compressibility is varying with depth resulting in, say three different ROCK keyword records, then there has to be three complete data sets in order to allocate the three ROCK records. This would mean that the PVTO and PVDG keywords, in this instance, would have to be repeated three times to match the three ROCK keyword records.

Example SATNUM and EQUIL arrays from the Volve⁵⁴ field are displayed in Figure 9.1 and Figure 9.2, respectively.





The Volve Data was approved for data sharing in 2018 by the initiative of the last Operating company, Equinor and approved by the license partners ExxonMobil E&P Norway AS and Bayerngas Norge AS in the end of 2017.

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9.3 Keyword Definitions

9.3.1 ADD - ADD A CONSTANT TO A SPECIFIED ARRAY

The ADD keyword adds a constant to a specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the ADD keyword is being applied.

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See ADD – Add a Constant to a Specified Array in the GRID section for a full description.

9.3.2 ADDREG - ADD A CONSTANT TO AN ARRAY BASED ON A REGION NUMBER

The ADDREG keyword adds a constant to a specified array or part of an array based on cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the ADDREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the ADDREG keyword is being used.

See ADDREG - Add a Constant to an Array based on a Region Number in the GRID section for a full description.

9.3.3 BOX - Define a Range of Grid Blocks to Enter Property Data

BOX defines a range of grid blocks for which subsequent data is assigned for all the cells in the defined BOX. Note that the BOX grid is reset by the keyword ENDBOX by resetting the current defined BOX to be the whole grid. The keyword can be used for any array and for all grid types.

See BOX - Define a Range of Grid Blocks to Enter Property Data in the GRID section for a full description.

9.3.4 COPY - Copy Array Data to Another Array

The COPY keyword copies an array (or part of an array) to another array or part of an array. The arrays can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the COPY keyword is being used.

See COPY - Copy Array Data to Another Array in the GRID section for a full description.

9.3.5 COPYREG - COPY AN ARRAY TO ANOTHER ARRAY BASED ON A REGION NUMBER

The COPYREG keyword copies a specified array or part of an array based on cells with a specific region number to another array. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the COPYREG keyword is read by the simulator. The property arrays can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the COPYREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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Revision: Rev-I

9.3.6 ENDBOX - Define the End of the BOX Defined Grid

This keyword marks the end of a previously defined BOX sub-grid as defined by a previously entered BOX keyword. The keyword resets the input grid to be the full grid as defined by the NX, NY, and NZ variables on the DIMENS keyword in the RUNSPEC section.

See ENDBOX - Define the End of the BOX Defined Grid in the GRID section for a full description.

9.3.7 ENDFIN - END THE DEFINITION OF A LOCAL GRID REFINEMENT

Description

ENDFIN defines the end of a Cartesian or radial local grid refinement ("LGR") definition and a LGR property definition data set.

See ENDFIN – End the Definition of a Local Grid Refinement in the GRID section for a full description.

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9.3.8 ENDNUM - Define the End-Point Scaling Depth Region Numbers

RUNSPEC GRID EDIT PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
-------------------------	---------	----------	---------	----------

Revision: Rev-I

Description

The ENDNUM keyword defines the end-point scaling depth table region numbers for each grid block. The end-point scaling depth tables for various regions are defined by the ENPVTD55 and the ENKRVD56 keywords in the PROPS section. In the RUNSPEC section the NTENDP variable on the ENDSCALE keyword defines the maximum number of depth tables.

No.	Name	Description	Default
I	ENDNUM	ENDNUM defines an array of positive integers assigning a grid cell to a particular end-point scaling depth table region.	I
		The maximum number of ENDNUM regions is set by the NTENDP variable on the ENDSCALE keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a ENDNUM region number then the default value of one will be used.
- 3) The keyword is terminated by "/".

Table 9.2: ENDNUM Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

The example below sets three ENDNUM regions for a 4 x 5 x 2 model.

ENDNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
```

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

```
----- BOX -----
-- -- ARRAY
              CONSTANT --
                          I1 I2 J1 J2
                                         K1 K2
EQUALS
                                  1*
   'ENDNUM'
                                      1*
                                           1* 1* / SET REGION 1
   'ENDNUM'
                              2
                                  1
                                      2
                                              1 / SET REGION 2
                          1
                                           1
   'ENDNUM'
                                               2 / SET REGION 3
```

This keyword is ignored by OPM Flow and has no effect on the simulation

This keyword is ignored by OPM Flow and has no effect on the simulation

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9.3.9 EQLNUM - Define the Equilibration Region Numbers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The EQLNUM keyword defines the equilibration region numbers for each grid block. The equilibration data for various regions are defined in the SOLUTION section. For example, the EQUIL keyword in the SOLUTION defines the initial pressures and fluid contacts for each equilibration region identified by the EQLNUM region array.

No.	Name	Description	Default
I	EQLNUM	EQLNUM defines an array of positive integers assigning a grid cell to a particular fluid in-place region.	I
		The maximum number of EQLNUM regions is set by the NTEQUIL variable on the EQLDIMS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The EQULNUM and PVTNUM arrays need to be consistent, that is the all cells with the same EQLNUM can only belong to one PVTNUM region.
- 3) If cell is not assigned a EQLNUM region number then the default value will be used.
- 4) The keyword is terminated by "/".

Table 9.3: EQLNUM Keyword Description

Examples

The example below sets three EQLNUM regions for a 4 x 5 x 2 model.

EQLNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
1 1 1 1
```

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

```
-- -- ARRAY CONSTANT -- I1 I2 J1 J2 K1 K2

EQUALS

'EQLNUM' 1 1* 1* 1* 1* 1* 1* / SET REGION 1
'EQLNUM' 2 1 2 1 2 1 1 / SET REGION 2
'EQLNUM' 3 1 2 1 2 2 2 / SET REGION 3
```

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9.3.10 EQUALREG - Sets an Array to a Constant by Region Number

The EQUALREG keyword sets a specified array to a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the EQUALREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the EQUALREG keyword is being used.

Revision: Rev-I

See Error: Reference source not found in the GRID section for a full description.

9.3.11 EQUALS - Sets a Specified Array to a Constant

The EQUALS keyword sets a specified array or part of an array to a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the EQUALS keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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9.3.12 FIPNUM - Define the Fluid In-Place Region Numbers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The FIPNUM keyword defines the fluid in-place region numbers for each grid block. The simulator can print out summaries of the fluid in-place in each region, the current flow rates between regions, and the cumulative flows between regions.

Note that the total number of FIPNUM and FIP regions must be defined by the NMFIPR variable on the REGDIMS keyword in the RUNSPEC section.

No.	Name	Description	Default
I	FIPNUM	FIPNUM defines an array of positive integers greater than or equal to one, that assigns a grid cell to a particular fluid in-place region.	I
		The maximum number of FIPNUM regions is set by the NTFIP variable on either REGDIMS or TABDIMS keywords in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a FIPNUM region then the default value will be used.
- 3) The keyword is terminated by "/".

Table 9.4: FIPNUM Keyword Description

Examples

The example below sets three FIPNUM regions for a 4 x 5 x 2 model.

FIPNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
```

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

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Revision: Rev-I

Note

In most simulation models the FIPNUM array is used to define various regions in the model for fluid in-place reporting and to identify (or report) the flow between the different regions. When calibrating a model's in-place volumes it would be useful to use the FIPNUM array combined with the MULTREGP keyword to accomplish this. However, the FIPNUM array cannot be used in the GRID section.

A work around is to:

- I) Use the FIPNUM array but change the keyword to MULTNUM and incorporate this keyword or INCLUDE file in the GRID section.
- 2) Use the MULTREGP to calibrate the fluid in-place volumes for the various regions.
- 3) In the REGIONS section, use the COPY keyword to copy the MULTNUM array to the FIPNUM array.

The above work flow will ensure that both arrays and the reporting of fluid in-place regions are consistent.

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9.3.13 FIPOWG - ACTIVATE OIL, GAS, AND WATER FIP ZONE REPORTING

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The FIPOWG keyword activates automatic fluid in-place reporting based on the initial oil, gas and water zones defined by the initial equilibration. The fluid contacts on the EQUIL keyword in the SOLUTION section determine the reporting fluid category a grid cell belongs to. For example all grid cells with depths above the gas-oil contact on the EQUIL keyword will be assigned to the gas zone and reported accordingly. Similarly, grid cells with depths between the gas-oil contact and the water-oil contact will be assigned to the oil zone. And finally, grid cells with depths below the oil-water contact will be assigned to the water zone. The simulator can print out summaries of the fluid in-place in each region, the current flow rates between regions, and the cumulative flows between regions.

Note that the total number of FIP and FIPNUM regions must be defined by the NMFIPR variable on the REGDIMS keyword in the RUNSPEC section.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

-- ACTIVATE OIL, GAS, AND WATER FIP ZONE REPORTING
-FIPOWG

The above example switches on automatic fluid in-place reporting based on the initial oil, gas and water zones defined by the initial equilibration.

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9.3.14 IMBNUM - Define the Imbibition Saturation Table Region Numbers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The IMBNUM keyword defines the imbibition saturation tables (relative permeability and capillary pressure tables) region numbers for each grid block, as such there must be one entry for each cell in the model. The region number specifies which set of relative permeability tables (SGFN, SWFN, SOF2, SOF3, SOF32D, SGOF, SLGOF and SWOF) are used to calculate the relative permeability and capillary pressure in a grid block.

No.	Name	Description			
I	IMBNUM	IMBNUM defines an array of positive integers assigning a grid cell to a particular saturation table region.	I		
		The maximum number of IMBNUM regions is set by the NTSFUN variable on the TABDIMS keyword in the RUNSPEC section.			

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a IMBNUM region then the default value of one will be used.
- 3) The keyword is terminated by "/".

Table 9.5: IMBNUM Keyword Description

In addition saturation table assignment may be may be directional dependent in which case the directional dependent versions of the aforementioned array should be used, that is IMBNUMX, IMBNUMY and IMBNUMZ instead of IMBNUM. There is also the facility to make the directional endpoint scaling reversible or non-reversible and if the non-reversible option is selected the non-reversible versions of the aforementioned arrays should be used, that is IMBNUMX, IMBNUMX-, IMBNUMY, IMBNUMY-, IMBNUMZ and IMBNUMZ-, instead of the IMBNUM keyword.

Example

The example below sets three IMBNUM regions for a 4 x 5 x 2 model using the EQUALS keyword.

```
CONSTANT --
                              ----- BOX -----
-- -- ARRAY
                              I1 I2
                                                 K1 K2
                                      J1 J2
EQUALS
                                       1*
   'IMBNUM'
                                           1*
                                                 1* 1* / SET REGION 1
                1
                                                     1 / SET REGION 2
2 / SET REGION 3
   'IMBNUM'
                2
                              1
                                       1
                                            2
                                                 1
   'IMBNUM'
```

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9.3.15 MISNUM - Define the Miscibility Region Numbers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The MISNUM keyword defines the miscibility region number mixing tables as defined by the TLMIXPAR keyword in the PROPS section, for when the miscibility option has been activated by the MISCIBLE keyword in the RUNSPEC section. MISNUM also allocates miscible residual oil saturation versus water saturation tables (SORWMIS keyword in the PROPS section) used to calculate the relative permeability and PVT properties for a grid cell.

Note that although this keyword can only be used when the miscibility option is active, it is not necessary to use this keyword even if the MISCIBLE keyword in the RUNSPEC has been activated as the default value of one will be applied to all grid blocks. Secondly, a value of zero for a grid cell results in immiscible fluids in that grid cell.

No.	Name	Description	Default
I	MISNUM	MISNUM defines an array of positive integers greater than or equal to zero, that assign a grid cell to a particular table of mixing parameters as defined by the TLMIXPAR and SORWMIS keywords.	I
		A value of zero sets the fluids within a grid cell to be immiscible.	
		The maximum number of MISNUM regions is set by the NTMIS variable on the MISCIBLE keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a MISNUM region then the default value of one will be used.
- 3) The keyword is terminated by "/".

Table 9.6: MISNUM Keyword Description

See also the TLMIXPAR and SORWMIS keyword in the PROPS section.

Example

The example below sets three MISNUM regions in the model on a layer by layer basis, using the EQUALS keyword.

```
-- -- ARRAY
             CONSTANT --
                         ----- BOX -----
                         I1 I2 J1 J2 K1 K2
EQUALS
                                1*
                                    1*
                         1* 1*
                                            12 / SET REGION 1
  'MISNUM'
                                         1
                                 1*
                                    1*
                                         13 55 / SET REGION 2
  'MISNUM'
                         1* 1*
                                 1* 1*
  'MISNUM'
                                         56 120 / SET REGION 3
```

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9.3.16 MULTIPLY - MULTIPLY A SPECIFIED ARRAY BY A CONSTANT

The MULTIPLY keyword multiplies a specified array or part of an array by a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the keyword is being used.

Revision: Rev-I

See Error: Reference source not found in the GRID section for a full description.

9.3.17 MULTIREG - MULTIPLY AN ARRAY BY A CONSTANT BASED ON A REGION NUMBER

The MULTIREG keyword multiplies an array or part of an array by a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTIREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the MULTIREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

9.3.18 OPERATE - Define Mathematical Operations on Arrays

This keyword defines the mathematical operations on arrays in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

See OPERATE - Define Mathematical Operations on Arrays in the GRID section for a full description.

9.3.19 OPERNUM - Define Regions for Mathematical Operations on Arrays

This keyword defines the OPERATE region numbers for each grid block. The OPERATE keyword defines mathematical operations on arrays in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords combined with MULTNUM region array.

See OPERNUM - Define Regions for Mathematical Operations on Arrays in the GRID section for a full description.

9.3.20 OPERATER - Define Mathematical Operations on Arrays by Region

This keyword defines the mathematical operations on arrays for specific regions in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

See OPERATER - Define Mathematical Operations on Arrays by Region in the GRID section for a full description.

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9.3.21 PLMIXNUM - Define the Polymer Region Numbers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The PLMIXNUM keyword defines the polymer region number for each grid block that is used to assign the mixing tables as well as the maximum polymer and salt concentrations, as defined by the PLMIXPAR and PLYMAX keywords in the PROPS section, for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

The maximum polymer concentration and the associated salt concentration are declared on the PLYMAX keyword.

No.	Name	Description	Default
I	PLMIXNUM	PLMIXNUM defines an array of positive integers greater than or equal to one, that assign a grid cell to a particular table of mixing parameters as defined by the PLMIXPAR and PLYMAX keywords.	I
		The maximum number of PLMIXNUM regions is set by the NPLMIX variable on the REGDIMS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a PLMIXNUM region then the default value of one will be used.
- 3) The keyword is terminated by "/".

Table 9.7: PLMIXNUM Keyword Description

See also the PLYADS, PLYADSS, PLYDHLF, PLYMAX, PLYROCK, PLYSHEAR, PLYSHLOG and PLYVISC keywords in the PROPS section.

Example

The example below sets three PLMIXNUM regions in the model on a layer by layer basis, using the EQUALS keyword.

```
CONSTANT -- ----- BOX -----
-- -- ARRAY
                             I1 I2 J1 J2 K1 K2
EQUALS
                                               1 12 / SET REGION 1
13 55 / SET REGION 2
                                 1*
                                      1*
   'PLMIXNUM'
                 1
                                          1*
   'PLMIXNUM'
                                 1*
                                      1*
                                          1*
                                          1*
   'PLMIXNUM'
                                               56 120 / SET REGION 3
```

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9.3.22 PVTNUM - Define the PVT Regions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PVTNUM keyword defines the PVT region numbers for each grid block, as such there must be one entry for each cell in the model. The region number specifies which set of PVT tables (DENSITY, PVDG, PVDO, PVTG, PVTO, PVCO, PVTW and ROCK) are used to calculate the PVT properties in grid block.

No.	Name	Description	Default
I	PVTNUM	PVTNUM defines an array of positive integers assigning a grid cell to a particular PVT region.	I
		The maximum number of PVTNUM regions is set by the NTPVT variable on the TABDIMS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword,in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The EQULNUM and PVTNUM arrays need to be consistent, that is the all cells with the same PVTNUM can only belong to one EQLNUM region.
- 3) If cell is not assigned a PVTNUM region then the default value will be used.
- 4) The keyword is terminated by "/".

Table 9.8: PVTNUM Keyword Description

Note

Care should be taken that cells in different PVTNUM regions are not in communication, since the fluid properties are associated with a cell. If for example, a rbbl or a rm³ of oil flows from PVTNUM region I to PVTNUM region 2, then the oil properties of that oil will change from the PVT I data set to the PVT data set 2. This will result in material balance errors, that may or may not cause numerical issues.

To avoid this one should use the MULTNUM (or FLUXNUM, or OPERNUM) array with the MULTREGT array to ensure that the various PVTNUM regions are not in communication.

Examples

The example below sets three PVTNUM regions for a $4 \times 5 \times 2$ model.

PVTNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
```

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Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

There third example shows how to ensure the various PVT regions are isolated. First of all define the MULTNUM array in the GRID section and ensure all the regions are isolated.

Revision: Rev-I

```
- -
-- GRID SECTION
GRID
-- -- ARRAY
          CONSTANT -- ----- BOX -----
                  I1 I2 J1 J2 K1 K2
EQUALS
  'MULTNUM'
          1
                  1* 1*
                        1* 1*
                              1* 1* / SET REGION 1
  'MULTNUM'
                               1 1 / SET REGION 2
                     2
                        1
                           2
  'MULTNUM'
                           2
                                 2 / SET REGION 3
/
-- SET TRANSMISSIBILITES ACROSS DIFFERENT RESERVOIRS TO ZERO TO ISOLATE
-- RESERVOIRS
                         NNC
   REGION
          REGION TRANS
                     DIREC
                               REGION ARRAY
- -
                              M / F / O
   FROM
                MULT
                     0PT
                          0PTS
MULTREGT
   1*
          1*
                     1*
                          'ALL'
                                       / ALL REGIONS SEALED
                0.0
```

Then in the REGIONS section copy the MULTNUM array to the PVTNUM array.

```
-- REGIONS SECTION
REGIONS
-- COPY AN ARRAY TO ANOTHER ARRAY BASED ON A REGION NUMBER
     ARRAY
             ARRAY
                     REGION
                             REGION ARRAY
    FROM
             TO
                     NUMBER
                            M / F / O
COPYREG
    'MULTNUM'
             'PVTNUM' 1
                                           / COPY MULT TO PVT 1
                             М
                                           / COPY MULT TO PVT 2
/ COPY MULT TO PVT 3
    'MULTNUM'
             'PVTNUM'
                     2
                             Μ
             'PVTNUM'
    'MULTNUM'
                     3
                             М
```

All the separate PVT regions are now isolated.

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Revision: Rev-I

9.3.23 REFINE - START THE DEFINITION OF A LOCAL GRID REFINEMENT

The REFINE keyword defines the start of a Cartesian or radial local grid refinement ("LGR") definition that sets the properties of the selected LGR. The keyword is then followed by the property keywords associated with the section where the keyword is being invoked. For example, if the REFINE keyword is used in the GRID section then most of the keywords in that section can be used to set the grid properties for the LGR.

See REFINE – Start the Definition of a Local Grid Refinement in the GRID section for a full description.

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9.3.24 REGIONS - Define the Start of the REGIONS Section of Keywords

PEC GRID EDIT PROPS	REGIONS SOLUTION	SUMMARY SCHEDULE
---------------------	------------------	------------------

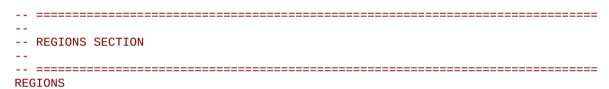
Revision: Rev-I

Description

The REGIONS activation keyword marks the end of the PROPS section and the start of the REGIONS section that defines how various fluid and rock property data defined in the PROPS section are allocated to the individual cells in the model.

There is no data required for this keyword.

Example



The above example marks the end of the PROPS section and the start of the REGIONS section in the OPM Flow data input file.

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9.3.25 ROCKNUM - Define Rock Compaction Table Region Numbers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The ROCKNUM keyword defines the rock compaction table region numbers for each grid block, as such there must be one entry for each cell in the model. The region number specifies which set of rock compaction tables defined by the ROCKTAB keyword are used to calculate the rock compaction in a grid block.

No.	Name	Description	Default
I	ROCKNUM	ROCKNUM defines an array of positive integers assigning a grid cell to a particular rock compaction table region.	I
		The maximum number of ROCKNUM regions is set by the NTROCC variable on the ROCKCOMP keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a ROCKNUM region then the default value will be used.
- 3) The keyword is terminated by "/".

Table 9.9: ROCKNUM Keyword Description

Examples

The example below sets three ROCKNUM regions for a 4 x 5 x 2 model.

ROCKNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
```

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

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9.3.26 RPTREGS - Define REGIONS Section Reporting

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the data in the REGIONS section that is to be printed to the output print file in human readable format. The keyword has two distinct forms, the first of which consists of the keyword followed by a series of integers on the next line indicating the data to be printed (see the first example). This is the original formal in the commercial simulator and was subsequently superseded by the second format. The second format consists of the keyword followed by a series of character strings that indicate the data to be printed. In most cases the character string is the keyword used to load the data in the OPM Flow input deck, for example FIPNUM for the fluid in-place array. Its is anticipated that OPM Flow will eventually support the functionality of the second format only, the first format although recognized will be completely ignored.

No.	Name	Description	Default
- 1	EQLNUM	Print the equilibration region array.	N/A
2	FIPNUM	Print the fluid in-place array.	N/A
3	PVTNUM	Print the PVT table assignment array.	N/A
4	SATNUM	Print the saturation function (relative permeability) assignment array.	N/A
			N/A

Notes:

The keyword is terminated by "/".

Table 9.10: RPTREGS Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

Note

This keyword has the potential to produce very large print files that some text editors may have difficulty loading, coupled with the fact that reviewing the data in this format is very cumbersome.

A more efficient solution is to load the *.INIT file into OPM ResInsight to view the data graphically, this also has the benefit of being able to filter the grid based on I, J, K ranges and grid properties.

Example

The first example shows the original format of this keyword; although the keyword and format are recognized by OPM Flow, the format is ignored and is unlikely to be implemented in in the simulator.

```
-- DEFINE REGIONS SECTION REPORT OPTION (ORIGINAL FORMAT)
-- RPTREGS
1 2*0 1 3*1 /
```

The next example shows the second format of the keyword which may be supported in a future release of OPM Flow.

```
-- DEFINE REGIONS SECTION REPORT OPTIONS
--
RPTREGS
DX DY DZ DEPTH PORO PERMX /
```



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9.3.27 SATNUM - Define the Saturation Table Region Numbers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SATNUM keyword defines the saturation tables (relative permeability and capillary pressure tables) region numbers for each grid block, as such there must be one entry for each cell in the model. The region number specifies which set of relative permeability tables (SGFN, SWFN, SOF2, SOF3, SOF32D, SGOF, SLGOF and SWOF) are used to calculate the relative permeability and capillary pressure in a grid block.

No.	Name	Description	Default
I	SATNUM	SATNUM defines an array of positive integers assigning a grid cell to a particular saturation table region.	I
		The maximum number of SATNUM regions is set by the NTSFUN variable on the TABDIMS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a SATNUM region then the default value will be used.
- 3) The keyword is terminated by "/".

Table 9.11: SATNUM Keyword Description

Examples

The example below sets three SATNUM regions for a $4 \times 5 \times 2$ model.

SATNUM

```
2 2 1 1 # layer 1
2 2 1 1
1 1 1 1
1 1 1 1
3 3 1 1 # layer 2
3 3 1 1
1 1 1 1
1 1 1 1
```

Alternatively the EQUALS keyword could be employed to accomplish the same task, that is:

```
-- -- ARRAY
              CONSTANT --
                           ----- BOX -----
                           I1 I2
                                   J1
                                      J2
                                            K1
EQUALS
   'SATNUM'
                                   1*
                                       1*
                                           1*
                                               1* / SET REGION 1
   'SATNUM'
                                               1 / SET REGION 2
                          1
                                   1
                                       2
                                           1
   'SATNUM'
                                       2
                                               2 / SET REGION 3
```

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9.3.28 TNUM - Define Passive Tracer Concentration Regions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The TNUM keyword defines the regions associated with the series of tracers associated with a phase (oil, water, or gas) in the model. The maximum number of tracers for each phase are declared on the TRACER keyword in the RUNSPEC section. Unlike other keywords, the TNUM keyword must be concatenated with the phase and the name of the tracer declared by TRACER keyword in the PROPS section. The following table outlines the format of the TNUM keyword name.

No.	Name	Description	Default
I	TNUM	A four letter character equal equal to TNUM that is the root keyword name for this data set array.	None
2	PHASE	A one letter character string that must be equal to F or S, that is concatenated to TNUM.	None
		The letter F states that the tracer is for the "free" phase, for example oil or water, as well as gas cap gas.	
		The letter S signifies that the tracer is a "solution" phase tracer, for example gas dissolved in oil (as activated by the DISGAS keyword in the RUNSPEC section), or condensate (vaporized oil) in the gas (as per the VAPOIL keyword in the RUNSPEC section).	
		Note tracers that are defined by the letter S to be in the "solution" phase, must also be initialized by the "free" phase as well.	
3	NAME	A three letter character string defining the tracer's name, which is concatenate to TNUM and PHASE to given the full name of the keyword	None
		Note it is best to void names beginning with the letters F, S. and T as these names may great naming issues in post-processing software.	

Table 9.12:TNUM Keyword Name Format

Following the declaration of the full keyword name, TNUMPHASENAME, the keyword is followed by the data as outlined below.

No.	Name	Description	Default
- 1	TNUMDATA	TUNDATA defines an array of positive integers assigning a grid cell to a particular tracer table region.	I
		The maximum number of TNUMDATA regions is set by the NTTRVD variable on the EQLDIMS keyword in the RUNSPEC section.	

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) If cell is not assigned a TNUMDATA region then the default value will be used.
- 3) The keyword is terminated by "/".

Table 9.13:TNUM Keyword Data Description

See also the TRACER keyword in the PROPS section and the TBLK keyword in the SOLUTION section.

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This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

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Examples

First define four passive tracers one for a free gas, one for dissolved gas, one for oil and one to track the water.

```
_ _
          DEFINE TRACER NAMES
- -
          TRACER
                     TRACER
- -
          NAME
                     PHASE
- -
          _ _ _ _ _
TRACER
         'GCG'
                     'GAS'
                                                                      / GAS CAP GAS
         'DGS'
                     'GAS'
                                                                      / DISOLVED GAS
         'OTI'
                     'OTI'
                                                                      / OIL
         'WAT'
                     'WAT'
                                                                      / WAT
```

Given a $100 \times 100 \times 5$ grid with DISGAS activated in the RUNSPEC section, then the following TNUM keywords define the various tracer regions given that NTTRVD equals four on the EQLDIMS keyword in the RUNSPEC section.

```
DEFINE PASSIVE TRACER CONCENTRATION REGIONS
_ _
TNUMFGCG
1000*1
1000*2
1000*2
1000*2
1000*2
TNUMSDGS
1000*1
1000*1
1000*1
1000*1
1000*1
TNUMFOIL
1000*3
1000*3
1000*3
1000*3
1000*3
TNUMFWAT
1000*4
1000*4
1000*4
1000*4
1000*4
```

The keyword name is derived from the TNUM keyword, plus either F or S, plus the tracer name declared in the TRACER keyword. For example for the gas cap (free gas) this would be TNUM+F+GAS to give the TNUMFGAS keyword. And for the dissolved (solution) gas this would be TNUM+S+DGS resulting in the TNUMSDGS keyword.

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10 SOLUTION SECTION

10.1 Introduction

To be written in a future release of the manual.

10.2 DATA REQUIREMENTS

To be written in a future release of the manual.

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10.3 Keyword Definitions

10.3.1 ADD - ADD A CONSTANT TO A SPECIFIED ARRAY

The ADD keyword adds a constant to a specified array or part of an array. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the ADD keyword is being applied.

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See ADD – Add a Constant to a Specified Array in the GRID section for a full description.

10.3.2 ADDREG - ADD A CONSTANT TO AN ARRAY BASED ON A REGION NUMBER

The ADDREG keyword adds a constant to a specified array or part of an array based on cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the ADDREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the ADDREG keyword is being used.

See ADDREG - Add a Constant to an Array based on a Region Number in the GRID section for a full description.

10.3.3 AQUANCON - Define Analytical Connections to the Grid

AQUANCON keyword defines how analytical aquifers are connected to the simulation grid., this includes Carter-Tracy and Fetkovich analytical aquifers, although the Carter-Tracy aquifer is the only aquifer currently being implemented in OPM Flow.

See AQUANCON – Define Analytical Connections to the Grid in the GRID section and AQUCT keyword in the SOLUTION section for a complete example on defining and connecting a Carter-Tracy aquifer to a simulation grid.

10.3.4 AQUCT - Define Carter-Tracy Analytical Aquifers

The AQUCT keyword defines a Carter Tracy aquifer, the properties of the aquifer, including the Carter-Tracy aquifer influence function associated with the aquifer, defined by the AQUTAB keyword in the PROPS section.

See AQUCT - Define Carter-Tracy Analytical Aquifers in the GRID section for a full description

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10.3.5 AQUFETP - FETKOVICH AQUIFER DEFINITION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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10.3.6 BOX - Define a Range of Grid Blocks to Enter Property Data

BOX defines a range of grid blocks for which subsequent data is assigned for all the cells in the defined BOX. Note that the BOX grid is reset by the keyword ENDBOX by resetting the current defined BOX to be the whole grid. The keyword can be used for any array and for all grid types.

Revision: Rev-I

See BOX - Define a Range of Grid Blocks to Enter Property Data in the GRID section for a full description.

10.3.7 COPY - COPY ARRAY DATA TO ANOTHER ARRAY

The COPY keyword copies an array (or part of an array) to another array or part of an array. The arrays can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the COPY keyword is being used.

See COPY - Copy Array Data to Another Array in the GRID section for a full description.

10.3.8 COPYREG - Copy an Array to Another Array based on a Region Number

The COPYREG keyword copies a specified array or part of an array based on cells with a specific region number to another array. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the COPYREG keyword is read by the simulator. The property arrays can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the COPYREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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10.3.9 DATUM - DEFINE THE DATUM DEPTH FOR THE MODEL

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

The DATUM keyword defines the datum depth for the model. This allows for all grid block pressures and potentials to be calculated at a common depth.

No.	Name	Description					
		Field	Metric	Laboratory			
T	DATUM	DATUM is a single positive value that defines the datum depth for the model.					
		feet	m	cm	None		
Notes:							
The keyword is terminated by "/".							

Table 10.1: DATUM Keyword Description

See also the DATUMR keyword that defines the datum for each fluid in-place region.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

-- DATUM -- DEPTH -- -----DATUM 5000.0

/ DATUM DEPTH FOR REPORTING

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The above example defines the datum for the model to be 5000.0

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10.3.10 DATUMR - DEFINE DATUM DEPTHS FOR THE FIPNUM REGIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SUMMARY	SCHEDULE

Description

The DATUMR keyword defines the datum depth for each fluid in-place region (FIPNUM) declared in the model. This allows for all grid block pressures and potentials to be calculated at a common depth within a FIPNUM region.

No.	Name		Default		
		Field	Metric	Laboratory	
1	DATUMR	DATUMR is a vector of each fluid in-place region.			
		feet	m	cm	None

Notes:

- 1) The keyword is followed by FIPNUM values as defined on REGDIMS keyword in the RUNSPEC section.
- 2) The keyword is terminated by "/".

Table 10.2: DATUMR Keyword Description

See also the DATUM keyword that defines the datum depth for the model.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

Examples

```
-- DATUM
-- DEPTH
-- DATUMR
4800.0
4900.0
5000.0
```

/ DATUM DEPTH FOR REPORTING

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The above example defines the datum depth for three FIPNUM regions, for when FIPNUM has been set equal to three on the REGDIMS keyword in the RUNSPEC section.

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10.3.11 ENDBOX - Define the End of the BOX Defined Grid

This keyword marks the end of a previously defined BOX sub-grid as defined by a previously entered BOX keyword. The keyword resets the input grid to be the full grid as defined by the NX, NY, and NZ variables on the DIMENS keyword in the RUNSPEC section.

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See ENDBOX - Define the End of the BOX Defined Grid in the GRID section for a full description.

10.3.12 ENDFIN - END THE DEFINITION OF A LOCAL GRID REFINEMENT

ENDFIN defines the end of a Cartesian or radial local grid refinement ("LGR") definition and a LGR property definition data set.

See ENDFIN - End the Definition of a Local Grid Refinement in the GRID section for a full description.

10.3.13 EQUALREG - Sets an Array to a Constant by Region Number

The EQUALREG keyword sets a specified array to a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the EQUALREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the EQUALREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

10.3.14 EQUALS - Sets a Specified Array to a Constant

The EQUALS keyword sets a specified array or part of an array to a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the EQUALS keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

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10.3.15 EQUIL - Define the Equilibration Initialization Data

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the parameters used to initialize the model for when equilibration is calculated by OPM Flow. This is the standard methodology to initialize a model, the non-standard formulation of entering the pressures and saturations for each grid cell is seldom employed in the industry. The keyword can be used for all grid types.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	DATUM	DATUM is a single posit for PRESS.	ive value that defines the	reference datum depth				
		feet m cm						
2	PRESS	PRESS is a single positive	value that defines the pre	ssure at DATUM.				
		respect to the gas phase.	If the DATUM depth lies above the GOC then PRESS is the pressure with respect to the gas phase. If the DATUM depth is below OWC then PRESS refers to the water phase pressure. Otherwise, PRESS refers to the oil phase pressure.					
		psia	barsa	atma	0.0			
3	WATCONT		runs containing oil, gas ar oil-water contact (OWC).	d water WATCONT is				
			uns containing oil and warer contact (OWC).	ater WATCONT is the				
	For two phase runs containing gas and water WATCONT is the depth of the gas-water contact (GWC).							
		feet	m	cm	0.0			
4	WATCAP	 For three phase runs containing oil, gas and water WATCAP is the oil-water capillary pressure at the OWC. 						
		For two phase runs containing oil and water WATCAP is the oil- water capillary pressure at the OWC.						
			phase runs containing gas and water WATCAP is the gaspillary pressure at the GWC					
		psia	barsa	atma	0.0			
5	GASCONT		runs containing oil, gas an oil-water contact (OWC).	nd water GASCONT is	0.0			
			where there is no gas old be set to a value shallo	,				
			ere is initially no oil zone, the GASCONT should b					
		For two phase ignored.	runs containing oil and	water GASCONT is				
		For two phase ignored.	runs containing gas and	water GASCONT is				

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No.	Name		Description		Default		
		Field	Metric	Laboratory			
		feet	m	cm			
6	GASCAP		I) For three phase runs containing oil, gas and water GASCAP is the gas-oil capillary pressure at the GWC.				
		2) For two phase rur	ns containing oil and water	GASCAP is ignored.			
		3) For two phase rur	ns containing gas and wate	r GASCAP is ignored.			
		psia	barsa	atma	0.0		
7	EQLOPTI		value that sets the initializent the run, as activated by t				
		calculated from da bubble-point pres (gas-oil ratio vers	of EQLOPTI results in ata entered on the PBVD sure versus depth table) sus depth table). If this of SVD keywords must be particular to the property of the surface of the s	(saturation pressure or or the RSVD keyword option is selected than			
			location of multiple PBV through the EQLNUM d.				
		the saturated gas-	QLOPT1 results in the g oil ratio at the GOC. In NT and the PBVD and R	this case DATUM must			
		 A negative value of EQLOPT1 results in the same option for when EQLOPT1 is zero. 					
		EQLOPTI is ignored if there is no dissolved gas in the run.					
		dimensionless	dimensionless	dimensionless	0		
8	EQLOPT2	_	value that sets the initializete) is present in the ru	-			
		being calculated pressure or dew keyword (condens	of EQLOPT2 results in the from data entered on point pressure versus deplosate-gas ratio versus dept er PDVD or RVVD keywo	the PDVD (saturation oth table) or the RVVD h table). If this option is			
			ocation of multiple PDV through the EQLNUM d.				
		set to the saturate	QLOPT2 results in the condensate-gas ratio as equal GASCONT and omitted.	t the GOC. In this case			
		3) A negative value of EQLOPT2 is zero.	of EQLOPT2 results in the	e same option for when			
		EQLOPT2 is ignored if th	ere is no vaporized oil in	the run.			

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0

dimensionless

dimensionless

dimensionless

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No.	Name		Description		Default			
		Field	Metric	Laboratory				
9	EQLOPT3	EQLOPT3 is an integer value that sets the initialization accuracy options for the equilibration calculation.						
		saturations at th calculation. This r potentially less a	e center of the grid bleesults in a stable initialize	PM Flow using the fluid lock in the equilibration ation at the expense of a calculation, especially for in the block.				
	grid cell into equilibration calc calculation at the	$2\left N\right +1$ horizont ulation. This results in a expense of initialization	ne simulator dividing each cal sub-blocks for the can accurate fluid in-place stability, that is there may is no production at the					
		Increasing the value of N increases the accuracy of the calculation, with the maximum value of N being set to 20 by OPM Flow.						
		3) A positive value of EQLOPT3 results in the same option for when EQLOPT3 is negative, except that tilted fault blocks are used in the calculation. Again, increasing the value of N increases the accuracy of the calculation, with the maximum value of N being set to 20 by OPM Flow.						
		Note this option	should be used with Irre	gular Corner-Point Grids.				
		EQLOPT3 is ignored for	Radial Grids.					
		Only EQLOPT3 equal to	o zero is supported by O	PM Flow.				
		dimensionless	dimensionless	dimensionless	0			
10		Not used.	1					
П		Not used						

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Notes:

- The keyword is followed by NTEQL records as declared on the EQLDIMS keyword in the RUNSPEC I)
- Each record is terminated by a "/" and there is no "/" terminator for the keyword.

Table 10.3: EQUIL Keyword Description

See also the PRESSURE, SGAS, SOIL and SWAT keywords in the SOLUTION section to initialize the model using the non-standard formulation of entering the pressures and saturations for each grid cell.

Examples

	DATUM	DATUM	OWC	PCOW	GOC	PCG0	RS	RV	N	
		PRESS						OPT		
	DEPTH	PKE55	DEPTH		DEPTH		UPI	UPI	UPI	
EQUIL										
	3650.0	1560.0	3712.0	0.00	1000.0	0.00	1	0	-5	/
	3650.0	1560.0	3741.0	0.00	1000.0	0.00	1	0	-5	/
	3650.0	1560.0	3741.0	0.00	1000.0	0.00	1	0	-5	/

The above example defines three equilibration records for when NTEQL equals three on the EQLDIMS keyword in the RUNSPEC section. Here there is no gas cap and the GOC has been set to a value above the reservoirs (1000.0), and the default value of EQLOPT (-5) has been explicitly stated.

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10.3.16 MULTIPLY - MULTIPLY A SPECIFIED ARRAY BY A CONSTANT

The MULTIPLY keyword multiplies a specified array or part of an array by a constant. The constant can be real or integer depending on the array type; however, the arrays that can be operated on is dependent on which section the keyword is being used.

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See Error: Reference source not found in the GRID section for a full description.

10.3.17 MULTIREG - Multiply an Array by a Constant based on a Region Number

The MULTIREG keyword multiplies an array or part of an array by a constant for cells with a specific region number. The region number array can be FLUXNUM, MULTNUM or OPERNUM and these arrays must be defined and be available before the MULTIREG keyword is read by the simulator. The constant can be real or integer depending on the property array type; however, the property arrays that can be operated on is dependent on which section the MULTIREG keyword is being used.

See Error: Reference source not found in the GRID section for a full description.

10.3.18 OPERATE - Define Mathematical Operations on Arrays

This keyword defines the mathematical operations on arrays in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

See OPERATE - Define Mathematical Operations on Arrays in the GRID section for a full description.

10.3.19 OPERATER - DEFINE MATHEMATICAL OPERATIONS ON ARRAYS BY REGION

This keyword defines the mathematical operations on arrays for specific regions in the commercial simulator and is currently not supported by OPM Flow. However, similar functionality is provided by the ADD and MULTIPLY keywords.

See OPERATER – Define Mathematical Operations on Arrays by Region in the GRID section for a full description.

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10.3.20 PBUB - Define the Initial Equilibration Bubble-Point Pressure for All GRID BLOCKS

RUNSPEC GRID EDIT	PROPS REGIONS	SOLUTION SUMMARY	SCHEDULE
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Revision: Rev-I

Description

The PBUB keyword defines the initial equilibration buble-point saturation pressures values for all grid cells in the model and should be used in conjunction with the PDEW, PRESSURE, RS, RV, SGAS, SOIL and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if dissolved gas has been activated in the model via the DISGAS keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

No.	Name		Default	
		Field		
1	PBUB	PBUB is an array of real bubble-point saturation p		
		Repeat counts may be us		
		psia barsa atma		None

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/". 2)

Table 10.4: PBUB Keyword Description

See also the PBVD, PDEW, PRESSURE, RV, SGAS, SOIL and SWAT keywords to fully define the initial state of the model.

Example

```
DEFINE INITIAL EQUILIBRATION PSAT VALUES FOR ALL CELLS IN THE MODEL
         BASED ON NX = 100, NY = 100 AND NZ = 3
- -
PBUB
         1000*3500.0
                        1000*3525.0
                                        1000*0.3535.0
```

The above example defines the initial equilibration bubble-point saturation pressure values to be 3500.0 for all the cells in the first layer, 3525.0 for all the cells in the second layer, and finally 3535.0 for all the cells in the third layer.

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10.3.21 PBVD - Equilibration Bubble-Point versus Depth Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PBVD keyword defines the bubble-point pressure versus depth tables for each equilibration region that should be used when there is dissolved gas in the model (DISGAS has been activated in the RUNSPEC section) and the EQLOPTI variable has been set to a positive integer on the EQUIL keyword in the SOLUTION section.

No.	Name		Default		
		Field	Metric	Laboratory	
I	DEPTH	A columnar vector of ralues that defines the calues, PBVALS.			
		feet	m	cm	None
2	PBVALS	A columnar vector of reat the corresponding DE			
		psia	None		

Notes:

- The keyword is followed by NTEQL records as declared on the EQLDIMS keyword in the RUNSPEC I) section.
- 2) Each table must contain a minimum of two rows and a maximum of NDRXVD rows as declared on the EQLDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 10.5: PBVD Keyword Description

Alternatively, the dissolved gas-oil ratio versus depth tables may be entered using the RSVD keyword in the SOLUTION section instead of this keyword. See also the RSVD and EQUIL keywords in the SOLUTION section.

Example

Given NTEQL equals three and NDRXVD is greater than or equal to two on the EQLDIMS keyword in the RUNSPEC section, then the following example defines the bubble-point versus depth functions.

```
- -
         DEPTH
                   PSAT
_ _
                   PRESS
          _ _ _ _ _ _
                    _____
PBVD
          3000.0
                   3000.0
                                                         / PSAT VS DEPTH EQUIL REGN 01
         8000.0
                   3025.0
         3000.0
                   3100.0
                                                         / PSAT VS DEPTH EQUIL REGN 02
          8000.0
                   3125.0
                   3200.0
          3000.0
                                                         / PSAT VS DEPTH EQUIL REGN 03
          8000.0
                   3225.0
```

Here three tables are entered and each table is terminated by "/" and there is no keyword terminating "/".

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10.3.22 PDEW - Define the Initial Equilibration Dew-Point Pressure for All GRID BLOCKS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
---------	------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

The PDEW keyword defines the initial equilibration dew-point pressure values for all grid cells in the model and should be used in conjunction with the PBUB, PRESSURE, RS, RV, SGAS, SOIL and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if vaporized oil been activated in the model via the VAPOIL keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

No.	Name			Default					
		Field	Metric Laboratory						
1	PDEW		PDEW is an array of real positive numbers assigning the initial equilibration dew-point pressure values to each cell in the model.						
		Repeat counts may be us							
		psia	psia barsa atma						

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- The keyword is terminated by "/".

Table 10.6: PDEW Keyword Description

See also the PBUB, PRESSURE, RS, RV, SGAS, SOIL and SWAT keywords to fully define the initial state of the model.

Example

```
DEFINE INITIAL EQUILIBRATION PSAT VALUES FOR ALL CELLS IN THE MODEL
         BASED ON NX = 100, NY = 100 AND NZ = 3
PDEW
         1000*3500.0
                        1000*3525.0
                                       1000*0.3535.0
```

The above example defines the initial equilibration dew-point saturation pressure values to be 3500.0 for all the cells in the first layer, 3525.0 for all the cells in the second layer, and finally 3535.0 for all the cells in the third layer.

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10.3.23 PDVD - Define Equilibration Dew-Point versus Depth Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The PDVD keyword defines the dew-point pressure versus depth tables for each equilibration region that should be used when there is vaporized oil in the model (VAPOIL has been activated in the RUNSPEC section) and the EQLOPT2 variable has been set to a positive integer on the EQUIL keyword in the SOLUTION section.

No.	Name		Default		
		Field	Metric	Laboratory	
I	DEPTH	A columnar vector of radius that defines the values, PDVALS.			
		feet	m	cm	None
2	PDVALS	A columnar vector of re the corresponding DEPT			
		psia	barsa	atma	None

Notes:

- The keyword is followed by NTEQL records as declared on the EQLDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NDRXVD rows as declared on the EQLDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 10.7: PDVD Keyword Description

Alternatively, the vaporized oil-gas ratio (condensate-gas ratio) versus depth tables may be entered using the RVVD keyword in the SOLUTION section instead of this keyword.

See also the RVVD and EQUIL keywords in the SOLUTION section.

Example

Given NTEQL equals three and NDRXVD is greater than or equal to two on the EQLDIMS keyword in the RUNSPEC section, then the following example defines the bubble-point versus depth functions.

DEPTH	PSAT							
	PRESS							
3000.0	2000.0							
8000.0	2025.0	/	PSAT	VS	DEPTH	EQUIL	REGN	01
3000.0	2100.0							
8000.0	3125.0	/	PSAT	VS	DEPTH	EQUIL	REGN	02
3000.0	2200.0							
8000.0	2225.0	/	PSAT	VS	DEPTH	EQUIL	REGN	03
	3000.0 8000.0 3000.0 8000.0 3000.0	PRESS 3000.0 2000.0 8000.0 2025.0 3000.0 2100.0 8000.0 3125.0 3000.0 2200.0	PRESS 3000.0 2000.0 8000.0 2025.0 / 3000.0 2100.0 8000.0 3125.0 / 3000.0 2200.0	PRESS 3000.0 2000.0 8000.0 2025.0 / PSAT 3000.0 2100.0 8000.0 3125.0 / PSAT 3000.0 2200.0	PRESS 3000.0 2000.0 8000.0 2025.0			

Here three tables are entered and each table is terminated by "/" and there is no keyword terminating "/".

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10.3.24 PRESSURE - Define the Initial Equilibration Pressures for All Grid Blocks

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The PRESSURE keyword defines the initial equilibration pressures for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, RS, RV, SGAS, SOIL and SWAT keywords etc., to fully describe the initial state of the model.

The keyword is used by the Enumeration Initialization method to initialize the model, as opposed to the Equilibration Initialization method that utilizes the EQUIL keyword in the SOLUTION section. This is the non-standard formulation to initialize the model and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name		Default				
		Field	Metric Laboratory				
I	PRESS	pressures to each cell in	positive numbers assigning the model. ed, for example 20*4200.0	,			
		psia	barsa	None			

Notes:

- 1) The number of entries should correspond to the NX \times NY \times NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.8: PRESSURE Keyword Description

See also the PBUB, PDEW, RS, RV, SGAS, SOIL and SWAT keywords to fully define the initial state of the model.

Example

```
-- DEFINE INITIAL EQUILIBRATION PRESSURES FOR ALL CELLS IN THE MODEL
-- BASED ON NX = 100, NY = 100 AND NZ = 3
--
PRESSURE
1000*4500.0 1000*4510.0 1000*4520.0 /
```

The above example defines the initial equilibration pressures to be 4500.0 for all the cells in the first layer, 4510.0 for all the cells in the second layer, and finally 4520.0 for all the cells in the third layer.

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10.3.25 REFINE - START THE DEFINITION OF A LOCAL GRID REFINEMENT

The REFINE keyword defines the start of a Cartesian or radial local grid refinement ("LGR") definition that sets the properties of the selected LGR. The keyword is then followed by the property keywords associated with the section where the keyword is being invoked. For example, if the REFINE keyword is used in the GRID section then most of the keywords in that section can be used to set the grid properties for the LGR.

See REFINE – Start the Definition of a Local Grid Refinement in the GRID section for a full description.

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10.3.26 RESTART - RESTART RUN FROM AN EXISTING RESTART FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The RESTART keyword defines the parameters to restart the simulation from a previous run that has written a RESTART file out to disk. Only restarting from RESTART files is permitted by OPM Flow; restarting from SAVE files is not implemented.

No.	Name	Description	Default
I	RSNAME	The RSNAME variable is a character string that defines the root name of the RESTART file to be read into the current input deck.	None
2	RSNUM	A positive integer that defines the restart point on the RESTART file to be read and to be used to initialize the model.	None
		When OPM Flow writes a restart point a message is printed to the *.PRT file indicating the time step the restart was written out.	
3	RSTYPE	Not used.	None
4	RSFORMAT	Not used.	None
Note	<u>s:</u>		

The keyword is terminated by "/".

Table 10.9: RESTART Keyword Description

The most direct way to start a restart run is to:

- Copy the existing data file that created the RESTART file and give it a new name. For example if
 the RESTART file is from a case named NOR-OPM-A01DATA, then the copied data file could be
 named NOR-OPM-A01-R1.DATA.
- Edit the copied data file (NOR-OPM-A01-R1.DATA) and delete all equilibration keywords (EQUIL, RSVD, etc.) or the enumeration keywords used to initialize the model (PRESSURE, SGAS, SOIL. SWAT, etc.) in the SOLUTION section.
- 3) In the SOLUTION section of NOR-OPM-A01-R1.DATA file insert the RESTART keyword, using NOR-OPM-A01 as RSNAME and the required RSNUM value for the time step to restart from.
- 4) In the SCHEDULE section of NOR-OPM-A01-R1.DATA file insert the SKIPREST keyword at the very beginning of the SCHEDULE section. The SKIPREST keyword causes the simulator to only read in data it requires for restarting the run up to the RESTART point (RSNUM). Note that certain keywords always need to be present in a restart run in the SCHEDULE section as the data is not stored on the RESTART file, for example the VFP tables (VFPPROD and VFPINJ keywords). The SKIPREST keyword automatically processes the input deck and reads the required data.
- 5) In the SCHEDULE section of NOR-OPM-A01-R1.DATA file after the RESTART point make any required changes, save the file and run the NOR-OPM-A01-R1.DATA with OPM Flow.

See also RPTRST, RPTSCHED and SKIPREST keywords.

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Examples

The example below defines a restart from the previously run NOR-OPM-A01 case at time step number 40.

```
-- SOLUTION SECTION
FLEXIBLE RESTART FROM PREVIOUS SIMULATION RUN
              RESTART
                   RESTART
    FTLF
    NAME
              NUMBER
                   TYPE
                       FORMAT
RESTART
    'NOR-OPM-A01'
                   1*
                        1*
              40
```

In addition in the SCHEDULE section the SKIPREST keyword should be used to correctly read in the schedule data up to the RESTART point.

```
-- SCHEDULE SECTION
-- SCHEDULE SECTION
-- SCHEDULE
-- ACTIVATE SKIPREST OPTION TO AVOID MODIFYING SCHEDULE SECTION
-- SKIPREST
```

Note is is advisable to place the SKIPREST keyword at the very beginning of the SCHEDULE section.

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10.3.27 RPTRST - Define Data to be Written to the RESTART File

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the data and frequency of the data to be written to the RESTART file at each requested restart point. In addition to the solution data arrays required to restart a run and the frequency of the data to be written, the user may request additional data to be written to to the restart file for visualization in OPM ResInsight.

The format consists of the keyword followed by a series of character strings that indicate the data to be printed. In most cases the character string is the keyword used to load the data in the OPM Flow input deck, for example KRG for the gas relative permeability of each grid block at the requested times. Its is anticipated that OPM Flow will support additional functionality as development progresses.

No.	Name	Description	Array Name
1	ALLPROPS	An alias for DEN, KRG, KRO, KRW, and VISC restart variable names combined that writes all the properties associated with these keywords.	
2	BASIC	BASIC defines the frequency at which the restart data for restarting a run and the additional requested data is written to the RESTART file. The paarameter is assigned a value, OPTION, using the form BASIC = OPTION, where OPTION is an integer variable set to: 1) OPTION = I then the restart files are written at every report time, but only the last one in the run is kept. This minimizes the restart file size but only the final results are stored, limiting the visualization in OPM ResInsight. 2) OPTION = 2 then the restart files are written at every report time step until this switch is reset and all the restarts are kept. 3) OPTION = 3 then the restart files are written every nth report time step with the frequency determined by the mnemonic "FREQ=n". This feature is not currently supported by OPM Flow. 4) OPTION = 4 then the restart files are written at the first report step of each year. 5) OPTION = 5 then the restart files are written at the first report step of each month. 6) OPTION = 6 then the restart files are written at every time step.	
		In addition for OPTION equal to 3, 4, and 5 the data may be written every n th report time with the frequency determined by the mnemonic "FREQ=n". However, this feature is currently not currently supported in OPM Flow.	
3	DEN	Oil, gas and water fluid phases in-situ densities.	OIL_DEN GAS_DEN WAT_DEN
4	KRG	Gas relative permeability at the grid blocks gas saturation.	GASKR
5	KRO	Oil relative permeability at the grid blocks oil saturation.	OILKR
6	KRW	Water relative permeability at the grid blocks water saturation.	WATKR

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No.	Name	Description	Array Name
7	RSSAT	Saturated dissolved gas-oil ratio for each grid block to enable restarts.	RSSAT
8	RVSAT	Saturated vaporized oil-gas ratio for each grid block to enable restarts.	RVSAT
9	VISC	Oil, gas and water fluid phases in-situ grid block viscosity data.	OIL_VISC GAS_VISC WAT_VISC

Revision: Rev-I

Notes:

1) The keyword is terminated by "/".

Table 10.10: RPTRST Keyword Description

Note that OPM Flow automatically writes out all the data required to make a restart run as outlined in the table below:

No.	Restart Variable Name	Variable Description	Variable Array Name
1	KRG	Gas relative permeability at the grid blocks gas saturation.	GASKR
2	KRNSW_GO	Gas-oil relative permeability scaling factor array used when the hysteresis option has been activated by the HYSTER variable on the SATOPTS keyword in the RUNSPEC section, to enable restarts.	KRNSWGO
3	KRNSW_OW	Oil-water relative permeability scaling factor array used when the hysteresis option has been activated by the HYSTER variable on the SATOPTS keyword in the RUNSPEC section, to enable restarts.	KRNSW_OW
4	KRO	Oil relative permeability at the grid blocks oil saturation.	OILKR
5	PCSWM_GO	Gas-oil capillary pressure scaling factor array used when the hysteresis option has been activated by the HYSTER variable on the SATOPTS keyword in the RUNSPEC section, to enable restarts.	PCSWM_GO
6	PCSWM_OW	Oil-Water capillary pressure scaling factor array used when the hysteresis option has been activated by the HYSTER variable on the SATOPTS keyword in the RUNSPEC section, to enable restarts.	PCSWM_OW
7	POLY	Polymer concentration for each grid block to enable restarts for when the POLYMER option has been activated in the RUNSPEC section	CPOLYMER
8	PRESS	Pressure data for each grid block to enable restarts.	PRESSURE
9	RS	Dissolved gas-oil ratio for each grid block to enable restarts.	RS
10	RV	Vaporized oil-gas ratio for each grid block to enable restarts.	RVS
11	SGAS	Gas saturation for each grid block to enable restarts.	SGAS
12	SOIL	Oil saturation each grid block to enable restarts.	SOIL
12	SOMAX	Maximum oil saturation used in determining the rate of oil vaporization in the presence of undersaturated gas and the rate at which the remaining oil gets "heavier" via the reduction in the solution gas-oil ratio ("Rs").	SOMAX
14	SSOL	Solvent saturation for each grid block to enable restarts for when the SOLVENT option has been activated in the RUNSPEC section	SSOL

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No.	Restart Variable Name	Variable Description	Variable Array Name
15	SWAT	Water ratio for each grid block to enable restarts.	SWAT
16	TEMP	Temperature of each grid block, currently not used in this version of OPM Flow.	Not Used
17	TERNARY	Three phase ternary saturation data is data array is calculated by OPM ResInsight when the RESTART file is loaded into OPM ResInsight,	TERNARY (Calculated)
18	KRW	Water relative permeability at the grid blocks water saturation.	WATKR

Revision: Rev-I

Notes:

 Only items (1) to (14) that are necessary to restart a run are written to the restart file, for example if the niether the POLYMER and SOLVENT options have not been invoked in the RUNSPEC section then the CPOLYMER and SSOL arrays will not be written to the restart file.

Table 10.11: Data Sets Automatically Written to the RESTART File

Examples

The first example request that the standard restart data be written out every month.

```
-- RESTART CONTROL BASIC = 4 (YEARLY) 5 (MONTHLY)
--
RPTRST
BASIC=4
```

The next example requests that the standard restart data be written at every report time step until this switch is reset and all the restarts are kept. In addition to the standard the data the the gas, oil and water relative permeability data will also be written out at each report time step.

```
-- RESTART CONTROL BASIC = 4 (YEARLY) 5 (MONTHLY)
-- RPTRST
BASIC=4 KRG KRO KRW //
```

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10.3.28 RPTSOL - Define SOLUTION Section Reporting

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the data in the SOLUTION section that is to be printed to the output print file in human readable format. The keyword has two distinct forms, the first of which consists of the keyword followed by a series of integers on the next line indicating the data to be printed (see the first example). This is the original formal in the commercial simulator and was subsequently superseded by the second format. The second format consists of the keyword followed by a series of character strings that indicate the data to be printed. In most cases the character string is the keyword used to load the data in the OPM Flow input deck, for example PVDG for the dry gas PVT tables. Its is anticipated that OPM Flow will eventually support the functionality of the second format only, the first format although recognized will be completely ignored.

This keyword is ignored by OPM Flow and has no effect on the simulation, but is documented here for completeness.

No.	Name	Description	Default
1	DENO	Print the oil reservoir density array	N/A
2	EQUIL	Print the equilibration report.	N/A
3	FIP	Print the fluid in-place report.	N/A
4	FIPRESV	Print the reservoir volumes in-place report.	N/A
			N/A

Notes:

I) The keyword is terminated by "/".

Table 10.12: RPTSOL Keyword Description

Note

Except for non-array like data, FIP etc., this keyword has the potential to produce very large print files that some text editors may have difficulty loading. A more efficient solution for array type data is to load the *.INIT and *.RESTART files into OPM ResInsight to view the data graphically, this also has the benefit of being able to filter the grid based on I, J, K ranges and grid properties.

Examples

The first example shows the original format of this keyword; although the keyword and format are recognized by OPM Flow, the format is ignored and is unlikely to be implemented in in the simulator.

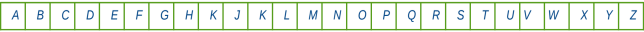
```
DEFINE SOLUTION SECTION REPORT OPTION (ORIGINAL FORMAT)

RPTSOL

1 2*0 1 3*1 /
```

The next example shows the second format of the keyword which may be supported in a future release of OPM Flow.

```
-- DEFINE SOLUTION SECTION REPORT OPTIONS
-- RPTSOL FIP=2 FIPRESV RESTART=3
```



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10.3.29 RS - Define the Initial Equilibration GOR (Rs) for All Grid Blocks

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The RS keyword defines the initial equilibration gas-oil ratio values for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, PRESSURE, RV, SGAS, SOIL and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if dissolved gas has been activated in the model via the DISGAS keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name		Description						
		Field	Metric	Laboratory					
I	RS	gas-oil ratio values to eac	RS is an array of real positive numbers assigning the initial equilibration gas-oil ratio values to each cell in the model.						
		Repeat counts may be us	Repeat counts may be used, for example 20*1.30.						
		Mscf/stb	sm³/sm³	scc/scc	None				

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.13: RS Keyword Description

See also the PBUB, PDEW, PRESSURE, RV, SGAS, SOIL and SWAT keywords to fully define the initial state of the model.

Example

```
DEFINE INITIAL EQUILIBRATION GOR VALUES FOR ALL CELLS IN THE MODEL
BASED ON NX = 100, NY = 100 AND NZ = 3

RS
1000*1.3500 1000*1.3010 1000*1.3000
```

The above example defines the initial equilibration GOR values to be 1.3500 for all the cells in the first layer, 1.3010 for all the cells in the second layer, and finally 1.3000 for all the cells in the third layer.

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10.3.30 RSVD - Equilibration Dissolved Gas-Oil Ratio (Rs) versus Depth Tables

|--|

Revision: Rev-I

Description

The RSVD keyword defines the dissolved gas-oil ratio (Rs) versus depth tables for each equilibration region that should be used when there is dissolved gas in the model (DISGAS has been activated in the RUNSPEC section) and the EQLOPTI variable has been set to a positive integer on the EQUIL keyword in the SOLUTION section.

No.	Name		Default				
		Field	Metric	Laboratory			
I	DEPTH		,				
		feet m cm					
2	RSVALS		A columnar vector of real values that defines the dissolved gas-oil ratio values at the corresponding DEPTH.				
		Mscf/stb	sm³/sm³	scc/scc	None		

Notes:

- The keyword is followed by NTEQL records as declared on the EQLDIMS keyword in the RUNSPEC I) section.
- Each table must contain a minimum of two rows and a maximum of NDRXVD rows as declared on the EQLDIMS keyword in the RUNSPEC section.
- Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 10.14: RSVD Keyword Description

Alternatively, the oil bubble-point pressure versus depth tables may be entered using the PBVD keyword in the SOLUTION section instead of this keyword.

See also the PBVD and EQUIL keywords in the SOLUTION section.

Example

Given NTEQL equals three and NDRXVD is greater than or equal to two on the EQLDIMS keyword in the RUNSPEC section, then the following example defines the bubble-point versus depth functions.

	DEPTH	RS							
		MSCF/STB							
RSVD									
	3000.0	1.400							
	8000.0	1.400	/	RS	VS	DEPTH	EQUIL	REGN	01
	3000.0	1.400							
	8000.0	1.400	/	RS	VS	DEPTH	EQUIL	REGN	02
	3000.0	1.400							
	8000.0	1.400	/	RS	VS	DEPTH	EQUIL	REGN	03

Here three tables are entered with a constant GOR versus depth relationship.

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10.3.31 RTEMP - Define the Initial Reservoir Temperature for the Model

This keyword defines the reservoir temperature for when a temperature option has been activated by the either the TEMP or THERMAL keywords in the RUNSPEC section. The RTEMPA keyword is alias for RTEMP

See RTEMP - Define the Initial Reservoir Temperature for the Model in the PROPS section for a full description.

10.3.32 RTEMPA - Define the Initial Reservoir Temperature for the Model

This keyword defines the reservoir temperature for when a temperature option has been activated by the either the TEMP or THERMAL keywords in the RUNSPEC section. The RTEMP keyword is alias for RTEMPA

See RTEMPA - Define the Initial Reservoir Temperature for the Model in the PROPS section for a full description.

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10.3.33 RTEMPVD - Define the Initial Reservoir Temperature versus Depth Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the reservoir temperature versus depth tables for when the temperature or thermal options has been activated by either the TEMP or THERMAL keywords in the RUNSPEC section in the commercial simulator. For OPM Flow the THERMAL keyword must be used to invoke the thermal option. The RTEMPVD keyword is an alias for the TEMPVD keyword; however, the latter is ignored by OPM Flow.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name	Description						
		Field	Metric	Laboratory				
I	DEPTH		A real monotonically increasing down the column values that defines the depth for corresponding reservoir temperature parameter TEMP.					
		feet m cm						
2	TEMP		A real monotonically increasing down the column values that defines the corresponding reservoir temperature for the given depth.					
		°F	°C	°C	None			

Notes:

- The keyword is followed by NTEQUIL tables as declared on the EQLDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NDRXVD rows as declared on the EQIDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 10.15: RTEMPVD Keyword Description

See also the RTEMP keyword in the PROPS section.

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Exampl	е									
	INITIAL	RESERVOIR	TEMPERATURE	VERSUS	DEPTH	TABLE				
RTEMPVD										
	DEPTH	TEMPERATU	JRE							
	FEET	DEG F								
	1000.0	90.000								
	2000.0	100.000								
	3000.0	130.000								
	4000.0	160.000					/	TABLE	NO.	01
	1000.0	90.000								
	2000.0	100.000								
	3000.0	130.000								
	4000.0	160.000					/	TABLE	NO.	02
	1000.0	90.000								
	2000.0	100.000								
	3000.0	130.000								
	4000.0	160.000					/	TABLE	NO.	03

The above example defines three identical reservoir depth versus temperature tables for the three NTEQUIL regions defined on the EQLDIMS keyword in the RUNSPEC section.

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10.3.34 RV - Define the Initial Equilibration CGR (Rv) for All Grid Blocks

|--|

Revision: Rev-I

Description

The RV keyword defines the initial equilibration vaporized oil-gas ratio values for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, PRESSURE, RS, SGAS, SOIL and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if vaporized oil been activated in the model via the VAPOIL keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name		Default			
		Field	Field Metric Laboratory			
I	RV	RV is an array of real povaporized oil-gas ratio va				
		Repeat counts may be us				
		stb/Mscf	None			

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.16: RV Keyword Description

See also the PBUB, PDEW, PRESSURE, RS, SGAS, SOIL and SWAT keywords to fully define the initial state of the model.

Example

```
-- DEFINE INITIAL EQUILIBRATION CGR VALUES FOR ALL CELLS IN THE MODEL
-- BASED ON NX = 100, NY = 100 AND NZ = 3
-- RV
1000*0.00720 1000*0.00725 1000*0.00730
```

The above example defines the initial equilibration GOR values to be 0.00720 for all the cells in the first layer, 0.00725 for all the cells in the second layer, and finally 0.00730 for all the cells in the third layer.

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10.3.35 RVVD - Equilibration Vaporized Oil-Gas Ratio (Rv) versus Depth Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The RVVD keyword defines the vaporized oil-gas ratio (Rv) versus depth tables for each equilibration region that should be used when there is vaporize oil in the model (VAPOIL has been activated in the RUNSPEC section) and the EQLOPT2 variable has been set to a positive integer on the EQUIL keyword in the SOLUTION section.

No.	Name		Default	
		Field		
I	DEPTH	A columnar vector of r values that defines the d gas ratio values, RVVALS.		
		feet	None	
2	RVVALS	A columnar vector of revalues, values at the corr		
		stb/Mscf	None	

Notes:

- The keyword is followed by NTEQL records as declared on the EQLDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NDRXVD rows as declared on the EQLDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 10.17: RVVD Keyword Description

Alternatively, the gas dew-point pressure versus depth tables may be entered using the PDVD keyword in the SOLUTION section instead of this keyword.

See also the PDVD and EQUIL keywords in the SOLUTION section.

Example

Given NTEQL equals three and NDRXVD is greater than or equal to two on the EQLDIMS keyword in the RUNSPEC section, then the following example defines the bubble-point versus depth functions.

	DEPTH	RS							
		STB/MSCF							
RVVD									
	3000.0	0.00725							
	8000.0	0.00725	/	RV	VS	DEPTH	EQUIL	REGN	01
	3000.0	0.00730							
	8000.0	0.00730	/	RV	VS	DEPTH	EQUIL	REGN	02
	3000.0	0.00750							
	8000.0	0.00750	/	RV	VS	DEPTH	EQUIL	REGN	03

Here three tables are entered with a constant CGR versus depth relationship for each equilibration region.

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10.3.36 SGAS - Define the Initial Equilibration Gas Saturation for All Grid Blocks

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The SGAS keyword defines the initial equilibration gas saturation values for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, PRESSURE, RS, RV, SOIL and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if the gas phase has been activated in the model via the GAS keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name			Default				
		Field	Metric	Laboratory				
1	SGAS	to zero and less than or saturation values to each	SGAS is an array of real positive numbers that are greater than or equal to zero and less than or equal to one assigning the initial equilibration gas saturation values to each cell in the model. Repeat counts may be used, for example 20*0.600.					
		dimensionless	dimensionless	dimensionless	None			

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.18: SGAS Keyword Description

Note for two phase runs it is only necessary to enter one saturation array of one of the phases present in the run (SGAS, SOIL, or SWAT), as the simulator will calculate the other phases by difference. Similarly for three phase runs it is only necessary to enter the array data for two of the phases, as the third saturation will again be calculated by the simulator.

See also the PBUB, PDEW, PRESSURE, RS, RV, SOIL and SWAT keywords to fully define the initial state of the model.

Example

```
--- DEFINE INITIAL EQUILIBRATION GAS SAT VALUES FOR ALL CELLS IN THE MODEL
-- BASED ON NX = 100, NY = 100 AND NZ = 3
--
SGAS
1000*0.7000 1000*0.6500 1000*0.6000 /
```

The above example defines the initial equilibration gas saturation values to be 0.7000 for all the cells in the first layer, 0.6500 for all the cells in the second layer, and finally 0.6000 for all the cells in the third layer.

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10.3.37 SOIL - DEFINE THE INITIAL EQUILIBRATION OIL SATURATION FOR ALL GRID BLOCKS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The SOIL keyword defines the initial equilibration oil saturation values for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, PRESSURE, RS, RV, SGAS and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if the oil phase has been activated in the model via the OIL keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name			Default				
		Field	Metric	Laboratory				
1	SOIL	zero and less than or estaturation values to each	COIL is an array of real positive numbers that are greater than or equal to zero and less than or equal to one assigning the initial equilibration oil saturation values to each cell in the model. Repeat counts may be used, for example 20*0.600.					
		dimensionless	dimensionless	dimensionless	None			

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.19: SOIL Keyword Description

Note for two phase runs it is only necessary to enter one saturation array of one of the phases present in the run (SGAS, SOIL, or SWAT), as the simulator will calculate the other phases by difference. Similarly for three phase runs it is only necessary to enter the array data for two of the phases, as the third saturation will again be calculated by the simulator.

See also the PBUB, PDEW, PRESSURE, RS, RV, SGAS and SWAT keywords to fully define the initial state of the model.

Example

```
-- DEFINE INITIAL EQUILIBRATION OIL SAT VALUES FOR ALL CELLS IN THE MODEL
-- BASED ON NX = 100, NY = 100 AND NZ = 3
-- SOIL
1000*0.7000 1000*0.6500 1000*0.6000 /
```

The above example defines the initial equilibration oil saturation values to be 0.7000 for all the cells in the first layer, 0.6500 for all the cells in the second layer, and finally 0.6000 for all the cells in the third layer.

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10.3.38 SPOLY - Define the Initial Equilibration Polymer Concentration for All Grid Blocks

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The SPOLY keyword defines the initial equilibration polymer <u>concentration</u> values for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, PRESSURE, RS, RV, SGAS, SGAS and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if the polymer phase has been activated in the model via the POLYMER keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name		Description					
		Field	Metric	Laboratory				
I	SPOLY	SPOLY is an array of rea initial equilibration poly model.						
		Repeat counts may be us						
		lb/stb	kg/sm³	gm/scc	None			

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.20: SPOLY Keyword Description

See also the PBUB, PDEW, PRESSURE, RS, RV, SGAS, SOIL and SWAT keywords to fully define the initial state of the model.

Example

```
-- DEFINE INITIAL EQUILIBRATION GAS SAT VALUES FOR ALL CELLS IN THE MODEL
-- BASED ON NX = 100, NY = 100 AND NZ = 3
-- SOIL
1000*0.0000 1000*0.0000 1000*15.000
```

The above example defines the initial equilibration polymer concentration values to be 0.0000 for all the cells in the first and second layers and finally 15.000 for all the cells in the third layer.

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10.3.39 SOLUTION - Define the Start of the SOLUTION Section of Keywords

UNSPEC GRID EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
------------------	-------	---------	----------	---------	----------

Revision: Rev-I

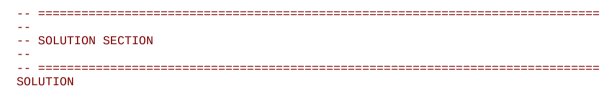
Description

The SOLUTION activation keyword marks the end of the REGIONS section and the start of the SOLUTION section that defines the initialized parameters used to initialized the model, by:

- I) defining fluid contacts and pressures, or
- 2) defining pressures and fluid saturations for all cells in the model, or
- 3) by restarting from a previously run OPM Flow completed run.

There is no data required for this keyword.

Example



The above example marks the end of the REGIONS section and the start of the SOLUTION section in the OPM Flow data input file.

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10.3.40 SSOL - Define the Initial Equilibration Solvent Saturation for All Grid Blocks

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The SSOL keyword defines the initial equilibration solvent saturation values for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, PRESSURE, RS, RV, SGAS, SOIL and SWAT keywords etc., to fully describe the initial state of the model. The keyword should only be used if the solvent phase has been activated in the model via the SOLVENT keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name			Default	
		Field	Metric	Laboratory	
1	SSOL	zero and less than or equ saturation values to each	positive numbers that are positive numbers that are possible to one assigning the inicial cell in the model. The ed, for example 20*0.000.		
		dimensionless	dimensionless	dimensionless	None

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.21: SSOL Keyword Description

See also the PBUB, PDEW, PRESSURE, RS, RV, SGAS, SOIL, and SWAT keywords to fully define the initial state of the model.

Example

```
-- DEFINE INITIAL EQUILIBRATION GAS SAT VALUES FOR ALL CELLS IN THE MODEL
-- BASED ON NX = 100, NY = 100 AND NZ = 3
-- SSOL
1000*0.0000 1000*0.0000 1000*0.0000
```

The above example defines the initial equilibration solvent saturation values to be 0.0 for all the cells in the in the model.

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10.3.41 STONE1 – ACTIVATE STONE'S FIRST THREE PHASE OIL RELATIVE PERMEABILITY MODEL

TROTO TEO OND TEODOLE TREGIONS SOLUTION SOMMANT SOLUTION	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
----------------------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword activates Stone's 57 first three phase oil relative permeability model as modified by Aziz and Settari 58. If the STONE1 and STONE2 keywords are not present in the input deck then the default three phase oil relative permeability model is employed. The STONE1 keyword should only be used in three phase runs containing the oil, gas and water phases.

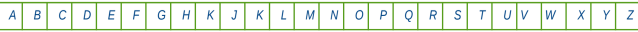
There is no data required for this keyword.

Example

-- ACTIVATE STONE'S FIRST THREE PHASE RELATIVE PERMEABILITY MODEL
-- STONE1

The above example switches on the Modified Stone three phase relative permeability model.

Aziz, K. and Settari, A. Petroleum Reservoir Simulation, London, UK, Applied Science Publishers (1979), page 398.



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⁵⁷ Stone, H. L. "Probability Model for Estimating Three-Phase Relative Permeability," paper SPE 2116, Journal of Canadian Petroleum Technology (1973) 22, No. 2, 214-218.

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10.3.42 STONE1EX - Define Stone's First Three Phase Oil Relative Permeability Parameter

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the exponent used in Stone's ⁵⁹ first three phase oil relative permeability model as modified by Aziz and Settari⁶⁰. The STONE IEX keyword should only be used in three phase runs containing the oil, gas and water phases and when the STONE I keyword in the SOLUTION section has been used to activate Stone's first three phase oil relative permeability model.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	STONEPARI	A real positive value that Stone first three phase of			
		dimensionless	dimensionless	dimensionless	None

Notes:

- The keyword is followed by NTSFUN records as declared on the TABDIMS keyword in the RUNSPEC section.
- 2) Each record must contain only one value and is terminated by "/"
- 3) There is no "/" terminator for the keyword.

Table 10.22: STONE I EX Keyword Description

If the STONE1 and STONE2 keywords are not present in the input deck then the default three phase oil relative permeability model is employed.

Example

Given NTSFUN equals five on the TABDIMS keyword in the RUNSPEC section, then:

Defines the exponents to be used in the Modified Stone first three phase oil relative permeability model, for each of the five saturation tables.

⁶⁰ Aziz, K. and Settari, A. Petroleum Reservoir Simulation, London, UK, Applied Science Publishers (1979), page 398.



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⁵⁹ Stone, H. L. "Probability Model for Estimating Three-Phase Relative Permeability," paper SPE 2116, Journal of Canadian Petroleum Technology (1973) 22, No. 2, 214-218.

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10.3.43 STONE2 - ACTIVATE STONE'S SECOND THREE PHASE OIL RELATIVE PERMEABILITY MODEL

RUNSPEC GRID EDIT	PROPS	REGIONS		SUMMARY	SCHEDULE
-------------------	-------	---------	--	---------	----------

Revision: Rev-I

Description

This keyword activates Stone's 61 second three phase oil relative permeability model as modified by Aziz and Settari62. If the STONE1 and STONE2 keywords are not present in the input deck then the default three phase oil relative permeability model is employed. The STONE2 keyword should only be used in three phase runs containing the oil, gas and water phases.

There is no data required for this keyword.

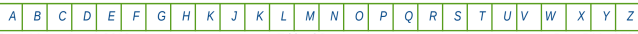
OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

Example

-- ACTIVATE STONE'S SECOND THREE PHASE RELATIVE PERMEABILITY MODEL
-- STONE2

The above example switches on the Modified Stone three phase relative permeability model

⁶² Aziz, K. and Settari, A. Petroleum Reservoir Simulation, London, UK, Applied Science Publishers (1979), page 398.



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⁶¹ Stone, H. L. "Estimation of Three-Phase Relative Permeability and Residual Oil Data," Journal of Canadian Petroleum Technology (1973) 12, No. 4, 53-61.

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10.3.44 SWAT - Define the Initial Equilibration Water Saturation for All Grid Blocks

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The SWAT keyword defines the initial equilibration water saturation values for all grid cells in the model and should be used in conjunction with the PBUB, PDEW, PRESSURE, RS, RV, SGAS and SOIL keywords etc., to fully describe the initial state of the model. The keyword should only be used if water been activated in the model via the WATER keyword in the RUNSPEC section.

This is the non-standard method to initialize the model via enumeration and is seldom employed in the industry. The standard methodology is for OPM Flow to initialize a model using the parameters on the EQUIL keyword combined with other keywords to fully describe the initial state of the model. The keyword can be used for all grid types.

No.	Name			Default					
		Field	Metric	Laboratory					
1	SWAT	to zero and less than o water saturation values to	SWAT is an array of real positive numbers that are greater than or equal to zero and less than or equal to one assigning the initial equilibration water saturation values to each cell in the model. Repeat counts may be used, for example 20*0.300.						
		dimensionless	dimensionless dimensionless						

Notes:

- 1) The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.23: SWAT Keyword Description

Note for two phase runs it is only necessary to enter one saturation array of one of the phases present in the run (SGAS, SOIL, or SWAT), as the simulator will calculate the other phases by difference. Similarly for three phase runs it is only necessary to enter the array data for two of the phases, as the third saturation will again be calculated by the simulator.

See also the PBUB, PDEW, PRESSURE, RS, RV, SGAS and SOIL keywords to fully define the initial state of the model.

Example

```
--
--
--
DEFINE INITIAL EQUILIBRATION GAS SAT VALUES FOR ALL CELLS IN THE MODEL
--
BASED ON NX = 100, NY = 100 AND NZ = 3
--
SWAT
1000*0.2000 1000*0.2500 1000*0.4500 /
```

The above example defines the initial equilibration water saturation values to be 0.2000 for all the cells in the first layer, 0.2500 for all the cells in the second layer, and finally 0.4500 for all the cells in the third layer.

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10.3.45 TEMPI - Define the Initial Temperature Values for All Cells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

TEMPI defines the initial temperature for all the cells in the model via an array for when the THERMAL option has been activated in the RUNSPEC section. This keyword is used to explicitly define the initial reservoir temperature via the Enumeration Initialization method rather than using the Equilibration Initialization method that utilizes the EQUIL keyword in the SOLUTION section.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

The keyword can be used for all grid types.

No.	Name		Description						
		Field	Field Metric Laboratory						
I	TEMPI	•	TEMPI is an array of real positive numbers assigning the initial temperature to each cell in the model.						
		Repeat counts may be us	Repeat counts may be used, for example 20*100.0.						
		°F	°F °C °C						

Notes:

- The number of entries should correspond to the NX x NY x NZ parameters on the DIMENS keyword in the RUNSPEC section, unless the BOX keyword defines a sub area of the grid, in which case the total number of entries should correspond to the number of cells defined by the BOX statement.
- 2) The keyword is terminated by "/".

Table 10.24:TEMPI Keyword Description

See also the RTEMP and RTEMPA keywords in the PROPS section and the RTEMPVD in the SOLUTION section for alternative ways to initialize the model's initial temperature.

Example

```
-- DEFINE GRID BLOCK TEMPERATURE FOR ALL CELLS
- (BASED ON NX x NY x NZ = 300)
-- TEMPI
100*212.0 100*215.0 100*220.0
```

The above example defines the initial temperature to be 212.0, 215.0, and 220.0 °F for the first, second and third layers in the model for all 300 cells, as defined by the DIMENS keyword in the RUNSPEC section.

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10.3.46 TEMPVD - Define the Initial Reservoir Temperature versus Depth Tables

	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
--	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the reservoir temperature versus depth tables for when the temperature or thermal options has been activated by either the TEMP or THERMAL keywords in the RUNSPEC section in the commercial simulator. For OPM Flow the THERMAL keyword must be used to invoke the thermal option. The TEMPVD keyword is an alias for the RTEMPVD keyword; however, the former is ignored by OPM Flow.

No.	Name		Description						
		Field Metric Laboratory							
1	DEPTH		A real monotonically increasing down the column values that defines the depth for corresponding reservoir temperature RTEMP.						
		feet	m	cm	None				
2	RTEMP		A real monotonically increasing down the column values that defines the corresponding reservoir temperature for the given depth						
		°F	°C	°C	None				

Notes:

- The keyword is followed by NTEQUIL tables as declared on the EQLDIMS keyword in the RUNSPEC section.
- Each table must contain a minimum of two rows and a maximum of NDRXVD rows as declared on the EQ:DIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 10.25:TEMPVD Keyword Description

See also the RTEMP keyword in the PROPS section and RTEMPVD keyword in the SOLUTION section..

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Exampl	е								
	INITIAL	RESERVOIR	TEMPERATURE	VERSUS	DEPTH TA	ABLE			
RTEMPVD									
	DEPTH	TEMPERATU	JRE						
	FEET	DEG F							
	1000.0	90.000							
	2000.0	100.000							
	3000.0	130.000							
	4000.0	160.000					/ TABLE	NO.	01
	1000.0	90.000							
	2000.0	100.000							
	3000.0	130.000							
	4000.0	160.000					/ TABLE	NO.	02
	1000.0	90.000							
	2000.0	100.000							
	3000.0	130.000							
	4000.0	160.000					/ TABLE	NO.	03

The above example defines three identical reservoir depth versus temperature tables for the three NTEQUIL regions defined on the EQLDIMS keyword in the RUNSPEC section.

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10.3.47 THPRES - Define Equilibration Region Threshold Pressures

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The THPRES defines the threshold pressure between various equilibration regions that have been defined by the EQLNUM keyword in the REGIONS section. The threshold pressure defines the potential difference between two regions which must be exceeded before flow can occur between the two regions. Once flow occurs the potential between the two regions is reduced by the threshold pressure.

This option must be activated by THPRES variable on EQLOPTS keyword in the RUNSPEC section in order to utilize this feature. Note that the irreversible option, as defined by IRREVER variable on EQLOPTS keyword in the RUNSPEC section, is not supported.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	EQLNUMI	than or equal to NTEQ	ve integer that is greater of UL on the EQLDIMS key 'from'' equilibration region	word in the RUNSPEC				
		dimensionless	dimensionless	dimensionless	None			
2	EQLNUM2	EQLNUMI is an a positive than or equal to NTEQ section, that defines the	word in the RUNSPEC					
		dimensionless dimensionless dimensionless						
3	THPRES	THPRES defines the threand from EQLNUM2 to	eshold pressure from EQ	LNUMI to EQLNUM2				
		The default value of I* sets the threshold pressure to a value that initially prevents flow between the two equilibration regions. Any subsequent production or injection in either of the two equilibration regions will therefore result in flow between the two regions. Thus, this default initially isolates the two equilibration regions.						
		If a equilibration region n keyword the THPRES is s						
		psia	barsa	atma	 *			

Notes:

- 1) The keyword is followed by any number of records with each record terminated by "/".
- 2) The keyword is terminated by "/".

Table 10.26:THPRES Keyword Description

See also the MULTREGT keyword in the GRID section that uses the transmissibility between the MULTNUM, FLUXNUM or OPERNUM region arrays to control the flow between various regions within the model.

Note

Care should be taken that cells in different EQLNUM regions are not in communication, as this will result in in an unstable initial equilibration.

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Examples

Given NTEQUL is equal to six on the EQLDIMS keyword in the RUNSPEC section,

```
EQLNUM EQLNUM
                         THPRES
         FROM
                 T0
                          VALUE
THPRES
                          0.588031
                                                              / REGN 1 TO REGN 2
         1
         2
                 1
                          0.588031
                                                              / REGN 2 TO REGN 1
         1
                 3
                          0.787619
                                                              / REGN 1 TO REGN 3
         3
                 1
                          0.787619
                                                              / REGN 3 TO REGN 1
         1
                 4
                          7.000830
                                                              / REGN 1 TO REGN 4
         4
                 1
                          7.000830
                                                              / REGN 4 TO REGN 1
```

Revision: Rev-I

The above example defines the threshold pressures between equilibration regions one and two, one and three and one and four. As the threshold pressures between regions one and five and one and six (as well as other combinations), have not been explicitly set in the example, the threshold pressures for these combinations are set to zero.

However, as the irreversible option, as defined by IRREVER variable on EQLOPTS keyword in the RUNSPEC section, is not supported, then example can be simplified to:

```
EOLNUM EOLNUM
                         THPRES
         FROM
                 T0
                         VALUE
THPRES
         1
                         0.588031
                                                             / REGN 1 AND REGN 2
         1
                 3
                         0.787619
                                                             / REGN 1 AND REGN 3
                 4
                         7.000830
                                                             / REGN 1 AND REGN 4
         1
```

Again, as the threshold pressures between regions one and five and one and six (as well as other combinations), have not been explicitly set in the example, the threshold pressures for these combinations are set to zero.

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10.3.48 TVDP - Define the Initial Equilibration Tracer Saturation versus Depth Functions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Description

This keyword defines the tracer name as the subsequent four characters after TVDP characters of the keyword, and then defines the tracer saturation as a function of depth.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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10.3.49 VAPPARS - OIL VAPORIZATION PARAMETERS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

VAPPARS defines the rate of oil vaporization in the presence of undersaturated gas and the rate at which the remaining oil gets "heavier" via the reduction in the solution gas-oil ratio ("Rs"). This keyword should only be used if the OIL, GAS, DISGAS and VAPOIL keywords in the RUNSPEC section have been invoked to allow oil, gas, dissolved gas and vaporized oil to be present in the model.

No.	Name		Description		Default					
		Field	Metric	Laboratory						
I	VAPPARI	VAPPARI is a real positive dimensionless number that defines the rate at which oil vaporizes into the available undersaturated gas in a grid block. The default value of zero invokes the standard black-oil formulation in which all oil vaporizes into the available undersaturated phase in a grid cell. Increasing this parameter decrease the rate of vaporization.								
		Typical values for VAPPAR	Typical values for VAPPAR1 range from zero and five.							
		dimensionless dimensionless								
2	VAPPAR2		ve dimensionless number ining oil in a grid cell decr							
		which the remaining oil's available undersaturated	The default value of zero invokes the standard black-oil formulation in which the remaining oil's Rs does not change as the oil vaporizes into the available undersaturated gas in a grid cell. Increasing this parameter increases the difference between the remaining oil and the vaporized oil							
		Typical values for VAPPAR	R2 are less than one.							
		dimensionless	dimensionless	dimensionless	0					

Table 10.27:VAPPARS Keyword Description

Note this keyword is normally used in history matching field performance to control the availability of the vaporized oil phase.

See also the DRSDT and DRVDT keywords in the SCHEDULE section that control the rate at which the solution gas-oil ratio and the vaporized oil-gas ratio increase within a grid block, respectively.

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Examples

The first example sets the black-oil default parameters

```
-- OIL VAPORIZATION PARAMETERS
-- OIL-VAP RS-INCS
-- VAPPAR1 VAPPAR2
VAPPARS
0 0 /
```

And the second example decreases the rate at which the oil vaporizes into the available undersaturated gas and increases the difference between the grid block oil saturation Rs and the vaporized oil Rs within a grid cell.

Revision: Rev-I

```
--
-- OIL VAPORIZATION PARAMETERS
--
-- OIL-VAP RS-INCS
-- VAPPAR1 VAPPAR2
VAPPARS
1.5 0.150 /
```

Again, the keyword is normally used in history matching field performance to control the availability of the vaporized oil phase.

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11 SUMMARY SECTION

11.1 Introduction

The SUMMARY section defines the variables to be written to the summary files that are used to generate line graphs of properties such as oil flow rate versus time, grid plot pressure versus time, etc. The properties to be stored on the summary file are entered in the SUMMARY section of the input file. The items requested are written to the summary file at the end of each successful time step. The SUMMARY section is terminated by the SCHEDULE keyword.

To generate a summary vector, the appropriate keyword must be entered in the SUMMARY section of the input data file. The SUMMARY keywords are listed below.

To be written in a future release of the manual.

11.2 DATA REQUIREMENTS

To be written in a future release of the manual.

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Revision: Rev-I

11.3 Keyword Definitions

11.3.1 ALL - EXPORT STANDARD SUMMARY VARIABLE VECTORS TO FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

This keyword activates the writing out of a standard set of summary production and injection data vectors for the field, group and well objects

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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11.3.2 DATE - ACTIVATE THE DATE OPTION FOR THE SUMMARY FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Description

This keyword activates the writing out of the date of each time step to the SUMMARY file. Normally only the time in days and decimal years are written out to the SUMMARY, activating the DATE option also results in the DATE being written out to the SUMMARY file as well. This option is normally used when RUNSUM keyword in the SUMMARY section has been activated to produce a RSM file.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

The following example shows an example RSM file output when the DATE option has NOT been activated.

SUMMARY OF RUN NO-DATE-KEYWORD											
TIME DAYS	YEARS YEARS	FPR PSIA	FOEW	FOPR STB/DAY	FOPT STB						
<u>0</u>		4467.125	0	0	0						
1.000000 31.00000	0.002738 0.084873	4466.943 4464.476	0.000239 0.007407	3235.662 3230.117	3235.662 100256.4						
60.00000	0.164271	4462.717	0.014291	3193.902	193421.5						
91.00000	0.249144	4460.813 4458.909	0.021523 0.028362	3127.557 3055.878	291306.3 383879.7						
152.0000	0.416153	4456.914 Y file DATE option	0.035262	2982.212	477271.4						

And the activating the SUMMARY file DATE option with:

-- ACTIVATE DATE SUMMARY FILE OPTION

DATE

Results in the following example RSM file output.

Results in the fo	ollowing exam	ipie KS	or file ou	tput.				
SUMMARY OF	RUN WITH-D	ATE-K	EYWORD					
DATE	YEARS YEARS	DAY	MONTH	YEAR	FPR PSIA	FOEW	FOPR STB/DAY	
1-JAN-98		 19	10	 1992	4467.125	 0		
2-JAN-98	0.002738	20	10	1992	4466.943	0.000239	3235.662	
31-JAN-98	0.084873	21	10	1992	4464.476	0.007407	3230.117	
28-FEB-98	0.164271	24	10	1992	4462.717	0.014291	3193.902	
31-MAR-98	0.249144	28	10	1992	4460.813	0.021523	3127.557	
30-APR-98	0.331280	3	11	1992	4458.909	0.028362	3055.878	
31-MAY-98	0.416153	14	11	1992	4456.914	0.035262	2982.212	

Note currently OPM Flow does not write out RSM files.

Α	В	С	D	Ε	F	G	Н	K	J	Κ	L	М	N	0	Р	Q	R	S	Τ	U	V	W	Χ	Υ	Z
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11.3.3 EXCEL - ACTIVATE THE EXCEL OPTION FOR THE SUMMARY FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates the writing out of the RSM file data in a format that can easily be loaded into Microsoft's EXCEL spreadsheet program or LibreOffice's CALC spreadsheet program. The RSM file output is activated by the RUNSUM keyword in the SUMMARY section.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

EXCEL

```
--
-- ACTIVATE EXCEL SUMMARY FILE OPTION
--
```

The above example activates the SUMMARY file EXCEL option for directly loading the RSM file into either Microsoft's EXCEL or LibreOffice's CALC spreadsheet programs

Note currently OPM Flow does not write out RSM files.

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11.3.4 FWSET - EXPORT WELL STATUS VECTORS FOR THE FIELD TO FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates the writing out of a set of data vectors that give the production and injections status of all the wells in the model.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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11.3.5 GMWSET - EXPORT WELL STATUS VECTORS BY GROUP TO FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates the writing out of a set of data vectors that give the production and injections status of all the wells within a group.

There is no data required for this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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11.3.6 MONITOR - ACTIVATES OUTPUT OF THE MONITORING DATA AND FILE

The MONITOR keyword activates the writing out of the run time monitoring information used by post-processing graphics software to display run time information, for example the simulated production and injection rates and cumulative values. OPM Flow does not have this functionality.

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See MONITOR - Activates Output of the Monitoring Data and File in the RUNSPEC section for a full description.

11.3.7 NOMONITO - DEACTIVATES OUTPUT OF THE MONITORING DATA AND FILE

The NOMONITO keyword deactivates the writing out of the run time monitoring information used by post-processing graphics software to display run time information, for example the simulated production and injection rates and cumulative values. OPM Flow does not have this functionality.

See NOMONITO – Deactivates Output of the Monitoring Data and File in the RUNSPEC section for a full description.

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11.3.8 RPTONLY - ACTIVATE THE REPORT TIME STEPS ONLY OPTION FOR THE SUMMARY FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword activates the writing out of the SUMMARY file and RSM file data, if the RSM file option has been requested by the RUNSUM keyword in the SUMMARY section, at report time steps only. The default is for all time steps to be written out to the files. This keyword reduces the file size at the expense of lower resolution in the time domain. There is no data required for this keyword.

The option can be deactivated by the RPTONLYO keyword in the SUMMARY section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

- -

ACTIVATE REPORT TIME STEPS ONLY SUMMARY FILE OPTION

RPTONLY

Note currently OPM Flow does not write out RSM files.

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11.3.9 RPTONLYO - DEACTIVATE THE REPORT TIME STEPS ONLY OPTION FOR THE SUMMARY FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword deactivates the writing out of the SUMMARY file and RSM file data, if the RSM file option has been requested by the RUNSUM keyword in the SUMMARY section, at report time steps only, and switches on writing out all the time steps to the files. This option is the default behavior for when RPTONLY has not been activated. There is no data required for this keyword.

The option can be activated by the RPTONLY keyword in the SUMMARY section that will switch on writing the data at every report time step instead of every time step..

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

- -

DEACTIVATE REPORT TIME STEPS ONLY SUMMARY FILE OPTION

RPTONLYO

Note currently OPM Flow does not write out RSM files.

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11.3.10 RPTSMRY - ACTIVATE OR DEACTIVATE SUMMARY LIST REPORT

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates or deactivates a listing of all the summary variables that are going to be written to the SUMMARY file and RSM file, if the RSM file option has been requested by the RUNSUM keyword in the SUMMARY section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

No.	Name		Description					
		Field	Metric	Laboratory				
1	RPTSMRY	An integer value set to ze	An integer value set to zero for no report, or one to produce the report.					
Notes	<u>:</u>				1			
I)	The keyword is	terminated by "/".						

Table 11.1: RPTSMRY Keyword Description

Examples

```
-- ACTIVATE OR DEACTIVATE SUMMARY LIST REPORT
-- RPTSMRY
1
```

The example switches on the summary list report.

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11.3.11 RUNSUM - ACTIVATE RSM FILE OUTPUT OF THE SUMMARY DATA

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates the writing out of the SUMMARY file date in a columnar format to the PRT file. Normally the SEPARATE keyword in the SUMMARY section is invoked in the same run to direct the data stream to a separate RSM file for easy loading into other programs, for example, Microsoft's EXCEL or LibreOffice's CALC spreadsheet programs.

There is no data required for this keyword.

See also the EXCEL, RPTONLY and SEPARATE keywords in the SUMMARY section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

-- ACTIVATE COLUMNAR SUMMARY DATA REPORTING OPTION

RUNSUM

Note currently OPM Flow does not write out RSM files.

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11.3.12 SEPARATE - ACTIVATE THE SEPARATE RSM FILE OUTPUT OPTION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates the writing out of the SUMMARY file date in a columnar format to the RSM file, if the RUNSUM keyword has been has also been activated in the SUMMARY section. Both the SEPARATE and the RUNSUM keywords need to be invoked. If the SEPARATE option is not activated then the RSM output is directed to the end of the PRT file. Normally the both the SEPARATE and RUNSUM keywords are invoked in the same run to enable easy loading of the data into Microsoft's EXCEL or LibreOffice's CALC spreadsheet programs.

There is no data required for this keyword.

See also the EXCEL, RPTONLY and RUNSUM keywords in the SUMMARY section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

- -

-- ACTIVATE SUMMARY DATA RSM FILE OUTPUT OPTION

_ _

SEPARATE

Note currently OPM Flow does not write out RSM files.

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11.3.13 SUMMARY - Define the Start of the SUMMARY Section of Keywords

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

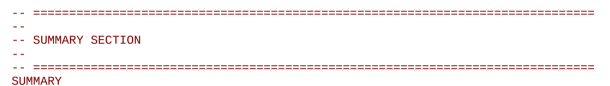
Revision: Rev-I

Description

The SUMMARY activation keyword marks the end of the SOLUTION section and the start of the SUMMARY section that defines the variables to be written out to the SUMMARY file for reporting and plotting of grid block data, production data, etc.

There is no data required for this keyword.

Example



The above example marks the end of the SOLUTION section and the start of the SUMMARY section in the OPM Flow data input file.

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11.3.14 SUMTHIN - Define SUMMARY DATA Reporting Time Steps

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines a time interval for writing out the SUMMARY data to the SUMMARY file and the RSM file, if the RUNSUM keyword has been has also been activated in the SUMMARY section. Only the data for the first time step in the time interval is written out and the other time steps are skipped until the next time interval. This enable the size of the SUMMARY files to be reduced depending on the size of the time interval. However, the keyword will produce irregular time steps reports of the SUMMARY data.

No.	Name		Description		Default					
		Field Metric Laboratory								
I	SUMSTEP	which the first time step	SUMSTEP is a real positive number that defines the time interval for which the first time step of data will be written to the SUMMARY file (and RSM file if RSM output has been activated).							
		steps of 0, 5, 10, 16, 24,	For example, if SUMSTEP is set to 30 days, and the simulator takes time steps of 0, 5, 10, 16, 24, 30, 40, 45, 60, 90 days. Then the SUMMARY data will be written out at time steps 0, 30, 40 and 60 days.							
		days	days	hours	None					

i) The keyword is terminated by 7.

See also the RPTONLY keyword in the SUMMARY section that forces the SUMMARY data to be only written out at report time steps, as oppose to all time steps or SUMSTEPS time intervals.

Table 11.2: SUMTHIN Keyword Description

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Examples

```
-- DEFINE SUMMARY DATA REPORTING TIME STEP INTERVAL
-- SUMSTEP
SUMTHIN 30.0 /
```

The above example defines the SUMMARY file time step interval to be 30 days for both field and metric units.

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12 SCHEDULE SECTION

12.1 Introduction

To be written in a future release of the manual.

12.2 DATA REQUIREMENTS

To be written in a future release of the manual.

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12.3 Keyword Definitions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

12.3.1 ACTION - Define Action Conditions and Command Processing (Field)

Revision: Rev-I

The ACTION keyword defines a series of conditions that invoke run time processing of ACTION functions and is similar to executing a run time script for conditions and variables at the field level.

This keyword is ignored by OPM Flow and has no effect on the simulation; use the ACTIONX keyword instead.

12.3.2 ACTIONG - Define Action Conditions and Command Processing (Groups)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

The ACTIONG keyword defines a series of conditions that invoke run time processing of ACTION functions and is similar to executing a run time script for conditions and variables at the group level

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate; use the ACTIONX keyword instead.

12.3.3 ACTIONR - Define Action Conditions and Command Processing (Regions)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

The ACTIONR keyword defines a series of conditions that invoke run time processing of ACTION functions and is similar to executing a run time script for conditions and variables at the region level

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate; use the ACTIONX keyword instead.

12.3.4 ACTIONS - Define Action Conditions and Command Processing (Well Segments)

RUNSPEC GRID EDIT PROPS	REGIONS SOLU	ITION SUMMARY SCHEDULE
-------------------------	--------------	------------------------

The ACTIONS keyword defines a series of conditions that invoke run time processing of ACTION functions and is similar to executing a run time script for conditions and variables associated with well segments.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate; use the ACTIONX keyword instead.

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12.3.5 ACTIONW - Define Action Conditions and Command Processing (Wells)

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

The ACTIONW keyword defines a series of conditions that invoke run time processing of ACTION functions and is similar to executing a run time script for conditions and variables at the well level.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate; use the ACTIONX keyword instead.

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12.3.6 ACTIONX - Define Action Conditions and Command Processing

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

The ACTIONX keyword defines a series of conditions that invoke run time processing of ACTION functions and is similar to executing a run time script. This is the general purpose version of the ACTION series of keywords that can apply Boolean conditional tests to variables at the field, group, region, well segment and well levels. The ACTION, ACTIONG, ACTIONR, ACTIONS and ACTIONW keywords are not implemented in OPM Flow are are unlikely to be so, as the ACTIONX keyword implements their functionality with greater flexibility.

Although this keyword is read by OPM Flow, the ACTION and UDQ computational logic and calculations have not been fully implemented and therefore this keyword should not be used as it may result in OPM Flow aborting.

This keyword starts the definition of a ACTIONX section that stipulates the Boolean conditions to test and the resulting SCHEDULE keywords to be executed if the Boolean condition evaluates to true. An ACTIONX Definition Section is terminated by an ENDACTIO keyword on a separate single line.

No.	Name		Description		Default
ACTIONX		Define the start of ACTIONX Definition Section. This is then followed on a new line by any number of ACTIONX records that define the conditions for which the defined action will be executed and the various operations to be performed if the conditions are satisfied.			
1-1	ACTNAME	ACTNAME is a character sting of up to length eight that defines the name of this action definition.			None
1-2	ACTNSTEP	ACTNSTEP is a positive integer that defines the number times that the ACTNAME definition is executed. ACTIONX definitions are activated at the end of a time step and this parameter is used to set how many time steps the ACTNAME definition will be invoked. The default value of one means that the definition will be executed only once. Use a large value, for example 10,000 for the definition to be executed at every time step			I
1-3	ACTDELTA	ACTDELTA is a real positive value that stipulates the duration of time that the conditions defined on the second record to be satisfied before the ACTIONX action are executed. For example, if ACTDELTA is defaulted the actions will be executed at the end of the time step for which the conditions are met. If set to say 30, then a minimum of 30 days will pass before the actions are executed (assuming field or metric units).			
		days	days	hours	0.0
1-4	1	Record terminated by a "/"			Not Applicable
2-1	ACTLHS	ACTLHS is a series of olength, that defines a covariable on the left hand	Not Applicable		
		The format for ACTLHS Aquifer, Block, Field, Gr Local Grid Refinement SUMMARY variables, an U used. The format for the			

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No.	Name	Description	Default
2-2	ACTTEST	ACTTEST is a defined character string that the states the Boolean operator and must be set to one of the following Boolean conditionals: 1) >: Greater than. 2) <: Less than. 3) >=: Greater than or equal to. 4) <=: Less than or equal to. 5) =: Equals to. 6) !=: Not equal to For example to test if the field's gas production rate is less than 600 MMscf/d then one would use: ACTIONX PHASE2 GGPR 'FIELD' < 600E3 /	Not Applicable
2-3	ACTRHS	ENDACTIO ACTRHS is a numeric value or a series of character strings, each up to eight characters in length, that defines a constant, an UDQ defined value, or a SUMMARY variable on the right hand side of a Boolean conditional test, as outlined in Table 12.2 (see also ACTLHS).	Not Applicable
2-4	ANDOR	An optional defined character string that stipulates a Boolean operator that must be set to either AND or OR if included on this record, that links this record with additional records of this type. For example, to test if the field's gas production rate is less than 600 MMscf/d after 2020 then one would use: ACTIONX PHASE2 GGPR 'FIELD' < 600E3 AND / YEAR > 2020 / / ENDACTIO This item should be left blank if not required.	Not Applicable
2.5	1	Termination of an ACTIONX Boolean condition record. Note that multiple numbers of records of this type can be entered with each record terminated by a "/", as illustrated above.	Not Applicable
3-1	1	The Boolean condition section of the ACTIONX keyword is terminated by an empty line with a single "/".	Not Applicable

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		Description	Default					
		The next section contains any number of standard SCHEDULE keywords that will be executed if the Boolean expression evaluates to true. For example, to test if the field's gas production rate is less than 600 MMscf/d after 2020 and to open up additional wells if this occurs, then one would use:	Not Applicable					
		ACTIONX PHASE2 1 / GGPR 'FIELD' < 600E3 AND / YEAR > 2020 / WELL PRODUCTION STATUS						
		WELL WELLLOCATION COMPLETION NAME STAT I J K FIRST LAST WELOPEN GP10 OPEN / GP11 OPEN / ENDACTIO						
		Most SCHEDULE keyword can used in an ACTIONX Define Section here, except for the time stepping keywords, i.e, TSTEP and DATES.						
ENDACTIO Define the end of ACTIONX Definition Section.								

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There is no terminating "/" for this keyword, instead the ENDACTIO keyword terminates the keyword.
 Table 12.1:ACTIONX Keyword Description

The variable types and the associated definitions that are available for use with Boolean conditionals are outlined in Table 12.2.

Variable	Description
Туре	
AQUIFER	AQUIFER variable consists of two parameters the:
	I) Aquifer SUMMARY variable; for example, Analytical Aquifer Influx Rate, AAQR, and the
	 Aquifer number consisting of a positive integer greater than zero that defines the aquifer to be used.
BLOCK	BLOCK variable consists of four parameters:
	I) Block SUMMARY variable; for example Block Oil Saturation, BOSAT.
	 Block I location which should be a positive integer greater than or equal to zero and less than or equal to NX that defines the connection location in the I-direction.
	 Block J location which should be a positive integer greater than or equal to zero and less than or equal to NY that defines the connection location in the J-direction.
	4) Block K location which should be a positive integer greater than or equal to one and less than or equal to NZ that defines the connection location in the K-direction.
	The NX, NY, and NZ parameters are defined on the DIMENS keyword in the RUNSPEC section.

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Variable	Description
Туре	
CONSTANTS	CONSTANTS can be any numerical value and can also include an integer constant as a counter secondary variable. This type of variable has the following form:
	I) Any numerical value.
	2) ACTIONX counter as defined by ACTNSTEP in the ACTIONX Definition Section. The default value is zero, but this parameter can be any integer value. For example, if one wanted to activate the action after the third time the Boolean condition was passed then setting ACTNSTEP to one and this parameter to -2 would accomplish this.
FIELD	The FIELD variable consists of any field SUMMARY variable; for example the Field average Pressure, as shown below:
	ACTIONX WIPHASE 1 / FPR < 2500 /
	ENDACTIO
	The above would action a set of SCHEDULE keywords if the field average pressure fell below 2,500 psia for a run using FIELD units.
GROUP	GROUP variable definition consists of:
	I) Group SUMMARY variable; for example, Group Oil Production Rate, GOPR.
	 Group Name which is a character string of up to eight characters in length that defines an existing group, note that the group named FIELD is the top most group.
	To enable an action for when the field's oil production rate drops below 20,000 stb/d then one could use.
	ACTIONX OILMIN 1 /
	GOPR 'FIELD' < 20.0E3 /
	/
	ENDACTIO
REGION	REGION variable definition consists of:
	 Region SUMMARY variable; selected from one of the following only: RPR, RGSAT, ROSAT, RWSAT, RGIP, ROIP, and RWIP. No other region summary fields are permitted in the expressions.
	2) Fluid In-Place region number which is a positive integer greater than or equal to zero that defines the region number. The value should less than or equal to the NTFIP variable on either REGDIMS or TABDIMS keywords in the RUNSPEC section. Note that a zero value indicates the whole model.
	3) Fluid In-Place region family (not used by OPM Flow).

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Variable	Description
Туре	·
TIME	TIME variables consists of one parameter that can have three values:
	DAY for the current simulation day of the month, MNTH for the current simulation month, and YEAR for the current simulation year.
	Thus to set an action for January 1, 2025 one would: use
	ACTIONX DAY = 1 AND /
	MNTH = 'JAN' AND / YEAR = 2025 /
	/ ENDACTIO
WELL	WELL variable definition consists of:
	I) Well SUMMARY variable; for example, Well Oil Production Rate, WOPR.
	2) Well Name which is a character string of up to eight characters in length that defines the well, which must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.
	To reduce the tubing head pressure constraint for when any of the oil producers' oil rate drop below 100 stb/d then one could use. ACTIONX WOILMIN 1 /
	WOPR 'OP*' < 100.0 /
	FLOW WELLS THROUGH LOW PRESSURE SEPARATOR
	WELL WELL TARGET NAME TARG VALUE
	WELTARG 'OP* ' THP 150 /
	ENDACTIO
WELL	WELL CONNECTION variable definition is comprised of:
CONNECTION	I) Well connection SUMMARY variable; for example, Connection Oil Flow Rate, COFR.
	2) Well Name which is a character string of up to eight characters in length that defines the well, which must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.
	 I- Connection: A positive integer greater than or equal to zero and less than or equal to NX that defines the connection location in the I-direction
	 J-Connection: A positive integer greater than or equal to zero and less than or equal to NY that defines the connection location in the J-direction.
	 K- Connection: A positive integer greater than or equal to one and less than or equal to NZ that defines the connection location in the K-direction.
	The NX, NY, and NZ parameters are defined on the DIMENS keyword in the RUNSPEC section.

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Variable Type	Description
WELL LOCAL	WELL LOCAL GRID REFINEMENT CONNECTION variable definition is comprised of:
GRID REFINEMENT CONNECTION	 Well local grid refinement connection SUMMARY variable; for example, the Local Grid Refinement Connection Oil Flow Rate, LCOFR.
CONNECTION	Well Name which is a character string of up to eight characters in length that defines the well, which must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.
	3) Local Grid Refinement Name which is a character string of up to eight characters in length that defines the Local Grid Refinement ("LGR"), which must have been declared previously using the CARFIN or RADFIN keywords in the GRID section, otherwise an error may occur.
	 I- Connection: A positive integer greater than or equal to zero and less than or equal to NX that defines the connection location in the I-direction within the LGR.
	 J-Connection: A positive integer greater than or equal to zero and less than or equal to NY that defines the connection location in the J-direction within the LGR.
	 K- Connection: A positive integer greater than or equal to one and less than or equal to NZ that defines the connection location in the K-direction within the LGR.
	The NX, NY, and NZ parameters are defined on either the CARFIN or RADFIN keywords in the GRID section depending upon whether a Cartesian or radial local grid refinement is being utilized.
	Note Local Grid Refinements are currently not implemented in OPM Flow.
WELL SEGMENT	WELL SEGMENT variable definition consists of:
	Well Segment SUMMARY variable; for example, Segment Oil Flow Rate, SOFR.
	2) Multi-Segment Well which is a character string of up to eight characters in length that defines the well name which must have been declared previously using the WELSPECS and WELSEGS keywords in the SCHEDULE section, otherwise an error may occur.
	 Segment Number, which is a positive integer greater than or equal to two and less than or equal to MXSEGS on WSEGDIMS keyword in the RUNSPEC section that defines the segment.
	Note that the total number of wells should be defined via the WELLSDIMS keyword and the number of multi-segment wells should be declared on the WSEGDIMS keyword, both keywords are in the RUNSPEC section.

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Table 12.2: ACTIONX Variable Definitions

See also the ACTDIMS and UDADIMS keyword in the RUNSPEC section to define the dimensions for the ACTIONX keyword and associated variables.

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Examples

The first example users the UDQ keyword to sort the oil wells from high water cut to low, via the WU_WLIST variable, and then use the ACTIONX keyword to shut-in the worst offending well when the field's water production is greater than 30,000 stb/d.

```
-- DEFINE START OF USER DEFINED QUANTITY SECTION
UDQ
-- OPERATOR VARIABLE EXPRESSION
                      1/(WWCT 'OP*')
            WU WCUT
                                                         / WELL WWCT LIST
DEETNE
DEFINE
            WU LIST
                      SORT(WU WCUT)
                                                         / WELL WWCT LIST SORTED
                                                           END OF UDQ SECTION
-- DEFINE START OF ACTIONX SECTION
- -
ACTIONX
      WSHUTIN
        GWPR 'FIELD' > 30E3 AND
        WU_LIST 'OP*' > 1
                              AND
- -
         DEFINE WELL AND WELL CONNECTIONS FLOWING STATUS
                --LOCATION--
                              COMPLETION
   WELL WELL
                         K FIRST LAST
   NAME STAT
                 I
                    J
WELOPEN
         SHUT
121
         SHUT
                      0
                           0
                                 0
```

ENDACTIO

Apart from checking that the field's water production rate is greater than 30,000 stb/d the Boolean conditional also checks that there is more than one well in the sorted well list. Notice also the use of '?' symbol as a substitution of the well name and that the ACTIONX WSHUTIN series of commands will be executed a total of ten times.

The second example checks to see if the field's gas rates is below 600 MMscf/d and if the simulation time is greater that January 1, 2030. If it is, then compression is installed by re-setting all the gas producing well's THP and BHP pressures to 450 psia and 300 psia respectively. In addition all gas wells currently shut-in are tested to see if they can be opened up under the new THP and BHP constraints.

```
-- START ACTIONX FIELD PHASE-3 AUTOMATIC COMPRESSION
ACTIONX
      PHASE-3
                      1
        GGPR 'FIELD' < 600E3 AND
                      = 1
        DAY
                               AND
                      ='JAN'
        MNTH
                               AND
        YEAR
                      = 2030
/
-- INSTALL COMPRESSION AND RESET WELL THP AND BHPS
    WELL
            WELL
                    TARGET
- -
   NAME
            TARG
                   VALUE
WELTARG
'GP*
            THP
                     450
'GP*
            BHP
                     300
```

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```
-- TEST AND OPEN ALL WELLS UNDER COMPRESSION CONSTRAINTS
-- WELL TEST CLOSE NO. START
-- NAME INTV CHECK CHECK TIME
WTEST
'GP* ' 1.0 PE 1 3 //
-- END OF ACTIONX FIELD PHASE-3 AUTOMATIC COMPRESSION DEFINITION
-- ENDACTIO
```

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12.3.7 BOX - Define a Range of Grid Blocks to Enter Property Data

BOX defines a range of grid blocks for which subsequent data is assigned for all the cells in the defined BOX. Note that the BOX grid is reset by the keyword ENDBOX by resetting the current defined BOX to be the whole grid. The keyword can be used for any array and for all grid types.

See BOX - Define a Range of Grid Blocks to Enter Property Data in the GRID section for a full description.

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12.3.8 CECON - Define Well Connections Economic Limit Criteria

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Revision: Rev-I

Description

CECON set the economic cut-off criteria for a well's connection to the simulation grid.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.9 COMPDAT - Define Well Connections to the Grid

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The COMPDAT keyword defines how a well is connected to the reservoir by defining or modifying existing well connections. Ideally the connections should be declared in the correct sequence, starting with the connection nearest the well head and then working along the wellbore towards the bottom or toe of the well, however this may not be possible or convenient, for example when connections are added or removed from a well during the simulation (see COMPORD in the SCHEDULE section for options regarding connection connection ordering).

No.	Name		Description		Default					
		Field	Metric	Laboratory						
I	WELNAME		A character string of up to eight characters in length that defines the well name for which the well connection data are being defined.							
		previously using the W	Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.							
2	I	A positive integer greater than or equal to zero and less than or equal to NX that defines the connection location in the I-direction.								
		If set to zero or defaulte location I-direction value section.								
3	J	A positive integer greater than or equal to zero and less than or equal to NY that defines the connection location in the J-direction.								
		If set to zero or defaulted with I^{*} the location is taken from the wellhead location J-direction value on the WELSPECS keyword in the SCHEDULE section.								
4	KI	A positive integer greater than or equal to one and less than or equal to NZ that defines the UPPER connection location in the K-direction.								
5	K2	A positive integer greater than or equal to KI and less than or equal to NZ that defines the LOWER connection location in the K-direction.								
6	STATUS	A character string of length four that defines the connections' operational status, STATUS should be set to one of the following character strings:								
	OPEN: the connections are open to flow.									
		2) SHUT: the conne	ctions are closed to flow (shut-in).						
	3) AUTO: the connection are initially closed, but may be opened automatically if an economic limit is violated. Currently this option is not supported by OPM Flow									

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No.	Name	Name Description						
		Field	Metric	Laboratory				
7	SATNUM	declared on the TABDII saturation table number	An integer greater than or equal to zero and less than NTSFUN as declared on the TABDIMS keyword in the RUNSPEC, that defines the saturation table number to be used for flow between the reservoir gird block and the well connections.					
		If SATNUM is set to zero	or defaulted with 1* the	n:				
			on table allocated to the relocated within is used.	he grid block that the				
		 If the hysteresis option has been activated via the HYSTER variable on the SATOPTS keyword in the RUNSPEC section, then both the imbibition and drainage saturation tables allocated to the grid block that the connections are located within are used. The imbibition table allocation can be changed by the COMPIMB keyword in the RUNSPEC section, provided it is entered after the COMPDAT keyword. 						
8	CONFACT	A real value greater than or equal to zero that defines the transmissibility connection factor between the well bore and the reservoir grid block.						
		If set to zero or defaulted with $1*$ then items (9) through (13) are used to calculate CONFACT.						
		cP.rb/day/psia	cP.rm³/day/bars	cP.rcc/hr/atm				
		0	0	0	Defined			
9	RW	A real positive value that defines the well bore <u>diameter</u> of the connections for the well.						
		RW is used in calculating a well's productivity or injectivity index; however the value will be ignored in calculating the connections CONFACT value if CONFAC has been directly entered.						
		feet	m	cm	None			
10	КН	A real value that defines the effective KH (permeability x length) for the connections.						
		If less than or equal to zero or defaulted by I* then KH is calculated from the connected grid blocks. KH is ignored if CONFAC has been directly entered.						
	mD.ft mD.m mD.cm							
П	SKIN A real value that defines the connections dimensionless skin factor.							
		SKIN is used in calculating a well's productivity or injectivity index; however, the value will be ignored in calculating the connections CONFACT value if CONFAC has been directly entered.						
		dimensionless	dimensionless	dimensionless	0.0			
12	DFACT	A real value that defines	the non-Darcy D factor c	oefficient for gas wells.				
			lefaulted with I* and th Ils defined via the WE					
		Currently this option is r	ot supported by OPM Flo	ow.				
	Currently this option is not supported by OPM Flow. day/Mscf day/m³ hour/sc							

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No.	Name		Description					
		Field						
13	DIRECT	connections and should connections also determined	A one letter character string that defines the orientation of the connections and should be set to either X, Y, or Z. The direction of connections also determines the length of the connection used to calculate the connection factor if CONFAC has not been entered directly.					
		The default value is for a to Z.	The default value is for a vertical connection, that is DIRECT is defaulted					

Revision: Rev-I

Notes:

- The keyword is followed by up to MXCONS records as declared on the WELLDIMS keyword in the RUNSPEC section.
- 2) Each record is terminated by a "/" and the keyword should be terminated by a "/".

Table 12.3: COMPDAT Keyword Description

Multiple grid block connections can be defined on one record for vertical wells by assigning different values to K1 and K2, for deviated and horizontal wells this may not be possible and therefore each grid block connection must be separately defined by using one record per connection, setting K1 equal to K2 in each record.

See also the WELSPECS keyword to define wells, the COMPIMB to reset the imbibition relative permeability table allocation, and the COMPORD to re-order the completions along the well trajectory. In addition, the COMPLUMP keyword groups well connections together to form well completions for a well. All the aforementioned keywords are described in the SCHEDULE section.

Note

The term well connection is used to describe individual connections from the wellbore to the reservoir grid, as opposed to well completions. A well completion is used to describe a set of connections, for example, a well may consist of several completions with each completion consisting of multiple connections.

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Example

The following example defines two vertical oil wells using the WELSPECS keyword and their associated connection data.

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```
WELL SPECIFICATION DATA
- -
-- WELL
         GROUP
                    LOCATION
                               BHP
                                      PHASE
                                              DRAIN
                                                     INFLOW
                                                              OPEN
                                                                    CROSS
                                                                            PRESS
-- NAME
         NAME
                      Ι
                           J
                               DEPTH
                                      FLUID
                                              AREA
                                                     EQUANS
                                                              SHUT
                                                                    FLOW
                                                                            TABLE
WELSPECS
                                               1*
0P01
         PLATFORM
                     14
                          13
                                1*
                                       OIL
                                                      STD
                                                              SHUT
                                                                     NO
                                                                             1* /
                     28
                                1*
                                               1*
0P02
         PLATFORM
                          96
                                       OIL
                                                      STD
                                                              SHUT
                                                                     NO
_ _
         WELL CONNECTION DATA
                                           CONN
-- WELL
         --- LOCATION ---
                             OPEN
                                    SAT
                                                  WELL
                                                          KΗ
                                                                SKIN
                                                                        D
                                                                              DIR
          II JJ K1 K2
                                           FACT
                                                          FACT
-- NAME
                             SHUT
                                    TAB
                                                                FACT
                                                                        FACT
                                                                              PEN
                                                  DIA
COMPDAT
                                    1*
                                                                        1*
              1*
                                           1*
                                                          1*
                                                                               'Z' /
          1*
                   20 56
                             OPEN
                                                 0.708
                                                                0.0
0P01
          1*
              1*
                                    1*
                                           1*
                                                          1*
                                                                        1*
                                                                              'Z' /
0P01
                   75 100
                             SHUT
                                                 0.708
                                                                0.0
                                           1*
0P02
             96 75 100
                             OPEN
                                                 0.708
                                                                0.0
```

Well OP01 has two sets of connections; the first one connects grid cells (14, 13, 20) to (14, 13, 56) to the well and is open to flow and the second connecting grid cells (14, 13, 75) to (14, 13, 100) is shut. Well OP02 has only one open connection from cells (35, 96, 75) to cells (35, 96, 100).

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12.3.10 COMPDATL - Define Well Connections to a LGR Grid

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The COMPDATL keyword defines how a well in a Local Grid Refinement ("LGR") is connected to the reservoir by declaring the LGR and defining or modifying existing well connections. Ideally the connections should be declared in the correct sequence, starting with the connection nearest the well head and then working along the wellbore towards the bottom or toe of the well, however this may not be possible or convenient, for example when connections are added or removed from a well during the simulation (see COMPORD in the SCHEDULE section for options regarding connection connection ordering).

This keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.11 COMPIMB - Assign Imbibition Saturation Tables to Well Connections

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The COMPIMB keyword assigns imbibition saturation tables to well connections. The COMPDAT keyword in the SCHEDULE section also assigns imbibition saturation tables to connections, but in this case the table number is the same as for the drainage curve. If this is not the required assignment then the COMPIMB keyword can be used to reset the imbibition saturation table number. For this to be effective the COMPIMB keyword must precede the COMPDAT keyword, otherwise it will have no effect.

The COMPIMB keyword should only be used if the hysteresis option has been activated via the HYSTER variable on the SATOPTS keyword in the RUNSPEC section.

No.	Name			Default	
		Field	Laboratory		
1	WELNAME		to eight characters in leng connection data is being d		None
			ame (WELNAME) must ELSPECS keyword in th occur.		
2	I		r than or equal to zero an nection location in the I-di		0
			ed with I* then all connect K2 criteria are assigned t		
3	J	A positive integer greate NY that defines the conn	0		
		If set to zero or defaulte that also satisfy I, KI and table number.			
4	KI		r than or equal to one ar ER connection location in		0
		If set to zero or defaulte well is used.	d with I* then the upper	most connection in the	
5	K2		r than or equal to KI an VER connection location i	•	0
		If set to zero or defaulted well is used.	most connection in the		
6	IMBNUM	An integer greater than declared on the TABDII imbibition saturation tal reservoir gird block and it	0		
		If IMBNUM is set to a saturation table allocate located within is used.			
		If I, J, K1, K2 are all set allocated to all connection	t to zero or defaulted to ns in the well.	o I*, then IMBNUM is	

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No.	Name		Description		Default
		Field	Metric	Laboratory	

Revision: Rev-I

Notes:

- The keyword is followed by up to MXCONS records as declared on the WELLDIMS keyword in the RUNSPEC section.
- 2) Each record is terminated by a "/" and the keyword should be terminated by a "/".

Table 12.4: COMPIMB Keyword Description

Multiple grid block connections can be defined on one record for vertical wells by assigning different values to K1 and K2, for deviated and horizontal wells this may not be possible and therefore each grid block connection must be separately defines by setting K1 equal to K2.

See also the COMPDAT keyword in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

The following example defines the connections for two vertical oil wells using the COMPDAT keyword and then re-sets the imbibition saturation functions using the COMPIMP keyword.

```
WELL CONNECTION DATA
          --- LOCATION ---
                                                  WELL
                             OPEN
                                    SAT
                                           CONN
                                                          KΗ
                                                                SKTN
                                                                        D
                                                                               DTR
-- WFII
-- NAME
          ΙI
              JJ K1 K2
                             SHUT
                                           FACT
                                                  DIA
                                                          FACT
                                                                FACT
                                                                        FACT
                                                                              PFN
COMPDAT
                                                                               'Z' /
          1*
               1*
                   20 56
                             OPEN
                                           1*
                                                 0.708
                                                          1*
                                                                0.0
                                                                        1*
0P01
                                    1
                                                                               'Z' /
0P01
          1*
               1*
                   75 100
                                    2
                                           1*
                                                          1*
                                                                        1*
                             SHUT
                                                 0.708
                                                                0.0
                                                                        1*
                                                                               'Z' /
                                           1*
          35
                                                          1*
0P02
              96
                  75 100
                             OPEN
                                    1
                                                 0.708
                                                                0.0
-- ASSIGN IMBIBITION SATURATION TABLES TO CONNECTIONS
-- WELL
             ---LOCATION---
                               SAT
-- NAME
             II
                 JJ K1 K2
                               TAB
COMPTMR
                 1*
                     20
0P01
                         56
                               11
            1*
                 1*
0P01
                     75 100
                               12
                 1*
0P02
                     1*
                               11
```

Well OP01 has two sets of COMPIMB records to overwrite the imbibition saturation tables, one for connections (14, 13, 20) to (14, 13, 56) resetting the imbibition saturation table number from one to 11 and one for connections (14, 13, 75) to (14, 13, 100) that resets the imbibition table number from 2 to 12. Well OP02 has only one connection from cells (35, 96, 75) to cells (35, 96, 100), so all the default values for I, J, K1, and K2 can be used to set the imbibition table numbers from 2 to 11. Note in all cases the drainage saturation table retains the value as specified by the COMPDAT keyword, that is one, two and one.

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12.3.12 COMPLUMP - Assign Well Connections to Completions

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The COMPLUMP keyword assigns connections, as defined by the COMPDAT keyword in the SCHEDULE section, to completion intervals. This "lumping" of the connections to various completion intervals allows automatic workovers and economic criteria to be applied to the completions (that is a set of connections) as opposed to the connections. This allows for a more realistic approach for workovers operations.

For example, if the water cut criteria for working over a well was set to 95%, and the average grid block connection thickness was one meter, then once a well's water cut reached 95% the worst offending one meter connection would be shut-in. If the well's actual perforation interval was 10 meters and the 10 connections were lumped as one completion, then when the water cut limit of 95% is reach, the completion would be shut-in, that is all of the 10 connections within the completion would be shut-in.

As the keyword is used to lump connections into a completions, the simulator adds together the contribution from all connections in the completion and uses the total values to test the economic limits. Note that a connection can only belong to one completion. In addition, completions can be used instead of connections in the WELOPEN and WPIMULT keywords if the completions have been defined by COMPLUMP for a well.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	WELNAME		to eight characters in leng connection data are being		None		
		Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.					
2	I	A positive integer greater than or equal to zero and less than or equal to NX that defines the connection location in the I-direction. If set to zero or defaulted with I* then all connections in the I-direction that also satisfy J, KI and K2 criteria are assigned the ICOMP completion number.					
3	J	A positive integer greater than or equal to zero and less than or equal to NY that defines the connection location in the J-direction.					
			If set to zero or defaulted with 1* then all connections in the J-direction that also satisfy I, K1 and K2 criteria are assigned the ICOMP completion				
4	KI		r than or equal to one ar ER connection location in		0		
		If set to zero or defaulte well is used.	d with I* then the upper	most connection in the			
5	K2 A positive integer greater than or equal to K1 and less than or equal to NZ that defines the LOWER connection location in the K-direction.				0		
		If set to zero or defaulte well is used.	ed with 1* then the low	most connection in the			

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No.	Name		Description					
		Field	Field Metric Laboratory					
6	ICOMP	MXCONS as defined of	n integer greater than or equal to one and less than or equal to IXCONS as defined on the WELLDIMS keyword in the RUNSPEC ection, that defines the completion number of the currently defined set of connections.					
			If I, J, K1, K2 are all set to zero or defaulted to 1*, then all connections in the well have the same completion number of ICOMP.					

Revision: Rev-I

Notes:

- 1) The keyword is followed by any number of records.
- 2) Each record is terminated by a "/" and the keyword should be terminated by a "/".

Table 12.5: COMPLUMP Keyword Description

Multiple grid block connections can be defined on one record for vertical wells by assigning different values to K1 and K2, for deviated and horizontal wells this may not be possible and therefore each grid block connection must be separately defines by setting K1 equal to K2.

See also the COMPDAT keyword in the SCHEDULE section.

Example

The following example defines the connections for two vertical oil wells using the COMPDAT keyword and the re-allocation of the connections to completions intervals using the COMPLUMP keyword.

```
WELL CONNECTION DATA
- -
-- WELL
         --- LOCATION ---
                            OPEN
                                   SAT
                                          CONN
                                                 WELL
                                                         KH
                                                               SKIN
                                                                      D
                                                                             DIR
                                          FACT
                                                         FACT
-- NAMF
          II JJ K1 K2
                                                               FACT
                                                                      FACT
                                                                             PFN
                            SHUT
                                   TAB
                                                 DIA
COMPDAT
                                   1*
                                          1*
                                                        1*
              1*
                                                                             'Z' /
0P01
          1*
                   20 56
                            OPEN
                                                0.708
                                                               0.0
          1*
              1*
                                   1*
                                          1*
                                                        1*
                                                                      1*
                                                                             'Z' /
0P01
                  75 100
                            SHUT
                                                0.708
                                                               0.0
              96
                                                0.708
0P02
          35
                  75 100
                            OPEN
                                                               0.0
         ASSIGN WELL CONNECTIONS TO COMPLETIONS
-- WELL
         --- LOCATION ---
                            COMPL
          II
-- NAME
             JJ K1 K2
                            NO.
COMPLUMP
          1*
              1*
                   20 56
                                                              / COMPLETION NO. 01
0P01
                             1
          1*
              1*
0P01
                  75 100
                             2
                                                              / COMPLETION NO. 02
          1*
              1*
0P02
                   75 85
                             1
                                                              / COMPLETION NO. 01
                                                              / COMPLETION NO. 02
0P02
                   86 100
```

Here the well OP01 connections (14, 13, 20) to (14, 13, 56) are assigned to completion number one and connections (14, 13, 75) to (14, 13, 100) are assigned to completion number two. Well OP02 has only one set of connection data from cells (35, 96, 75) to cells (35, 96, 100), but they have split into two separate completion intervals, with connections (35, 96, 75) to (35, 96, 85) assigned to completion interval number one and (35, 96, 86) to (35, 96, 100) to completion number two.

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12.3.13 COMPORD - Define Well Connection Ordering

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The COMPORD keyword defines how the well connection data entered on the COMPDAT keyword in the SCHEDULE section are to be ordered for a well.

No.	Name			Default		
		Field	Metric	Laboratory		
I	WELNAME		A character string of up to eight characters in length that defines the well name for which the well connection data are being defined.			
		Note that the well n previously using the W otherwise an error may o				
2	COMPORD A character string that defines the method for ordering the well connections given on the COMPDAT keyword, and should be set to DEPTH, INPUT, or TRACK.					
		DEPTH: The column vertical depth from connections are a condered by the skeyword.				
		same sequence as the connections starting with the	on results in the connectic s entered via the COMPD should be declared in e connection nearest the e wellbore towards the bo	AT keyword. In this case the correct sequence, e well head and then		
		connections thro connections. If the (via the DIRECT	ption enables OPM Flough the grid to obtain the supplied COMPDAT indivariable being equal to e DEPTH option will be a	e correct order for the icates the well is vertical Z on the COMPDAT		

Notes:

- 1) The keyword is followed by any number of records.
- 2) Each record is terminated by a "/" and the keyword should be terminated by a "/".

Table 12.6: COMPORD Keyword Description

See also the COMPDAT keyword in the SCHEDULE section.

Note

If visual inspection of the well trajectories in the model indicate problematic or unrealistic well connections, the options on this keyword may be useful in correcting the issue.

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Example

The following example defines the connections for two vertical oil wells using the COMPDAT keyword and the COMPORD to defined the connection ordering for the wells.

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```
WELL CONNECTION DATA
- -
-- WELL
         --- LOCATION ---
                            OPEN
                                   SAT
                                         CONN
                                                 WELL
                                                        KH
                                                              SKIN
                                                                      D
                                                                            DIR
          II JJ K1 K2
                                                        FACT
                                                                      FACT
-- NAME
                            SHUT
                                   TAB
                                         FACT
                                                 DIA
                                                              FACT
                                                                            PEN
COMPDAT
                                   1*
                                                                            'Z' /
              1*
                                         1*
                                                        1*
0P01
                  20 56
                            OPEN
                                                0.708
                                                              0.0
                                                                     1*
          1* 1*
                                   1*
                                         1*
                                                        1*
                                                                            'Z' /
0P01
                  75 100
                            SHUT
                                                0.708
                                                              0.0
                                         1*
                                                        1*
0P02
          35 96 75 100
                            OPEN
                                                0.708
                                                              0.0
_ _
         DEFINE WELL CONNECTION ORDERING
-- WELL
        COMPL
-- NAME
         ORDER
COMPORD
         DEPTH
0P01
0P02
         DEPTH
```

The DEPTH option has been chosen because both wells are vertical. Also one could use the following format instead for the COMPORD:

```
--
-- DEFINE WELL CONNECTION ORDERING
--
-- WELL COMPL
-- NAME ORDER
COMPORD
* DEPTH /
```

as both wells should utilize the DEPTH option. This version would set all wells in the model to DEPTH connection ordering.

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12.3.14 COMPSEGS - Define Well Connections for Multi-Segment Wells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The COMSEGS keyword defines how a multi-segment well is connected to the reservoir by defining or modifying existing well connections. Note that well must have been previously define by the WELSPECS keyword in the SCHEDULE section and the well connections must have been previously defined via the COMPDAT keyword in the SCHEDULE section

The COMPSEGS keyword should be repeated for each multi-segment well in the model.

No.	Name		Default				
		Field	Metric	Laboratory			
1-1	WELNAME		to eight characters in leng egment well is being define		None		
		previously using the W	Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.				
1-2	1	Record terminated by a "	·/"		Not Applicable		
2-1	I		r than or equal to one an nection location in the I-di		None		
2-2	J		A positive integer greater than or equal to zero and less than or equal to NY that defines the connection location in the J-direction.				
2-3	К	A positive integer greate NZ that defines the conn	None				
2-4	IBRANCH	MXBRAN on WSEGDIM	A positive integer greater than or equal to one and less than or equal to MXBRAN on WSEGDIMS keyword in the RUNSPEC section that defines the branch number of the defined I, J and K connection.				
2.5	DEPTHI		DEPTH1 is a real positive value that defines the length of the tubing from the tubing head or wellhead at the surface to the <u>start</u> of the connection in the I, J, K cell.				
		feet	m	cm	None		
2-6	DEPTH2	DEPTH2 is a real positive the tubing head or wellhe the I, J, K cell.					
		feet	m	cm	None		
2-7	DIRECT	A one letter characte connections and should connections also determine	Z				
		The default value is for a to Z.	vertical connection, that	is DIRECT is defaulted			
		Currently this option is n	ot supported by OPM Flo	w.			

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No.	Name			Default			
		Field	Metric	Laboratory			
2-8	IEND	IEND is positive or negatione of the following:	IEND is positive or negative integer, that is not equal to zero that is set to one of the following:				
		*	-NX and +NX that is onnection location in the l				
		,	-NY and +NY that is in the John of the Joh				
			-NZ and +NZ that is onnection location in the k				
		that defines the end of value of DIRECT.					
		For example, if DIREC associated with the J-direct must be calculated to re 100 on the DIMENS ke record set to 50, then IEN					
		Currently this option is n	ot supported by OPM Flo	w.			
2-9	DEPTH3		e value that defines the dat aken as the mid-point of t f connections.				
		Currently this option is n	ot supported by OPM Flo	w.			
		feet	m	cm	None		
2-10	LENGTH	-	e value that defines the le used in thermal calculation	_			
		Currently this option is n	ot supported by OPM Flo	w.			
		feet	m	cm	None		
2-11	ISEG	A real positive values e coordinate in the x-dire used for display purposes					
		Currently this option is n					
		feet	m	cm	None		
2-12	1	Record terminated by a "	·" ₍₁₎ "		Not Applicable		

Revision: Rev-I

Notes:

- Each multi-segment wells must be defined by a separate COMPSEGS keyword that consists of two records, with entries 1-1 to 1-2 representing record one items and 2-1 to 2-12 representing record number two items in the "No." column in this table.
- 2) Record number two of the keyword, items 2-1 to 2-12 is followed by up to MXCONS records as declared on the WSEGDIMS keyword in the RUNSPEC section.
- Each of the records are terminated by a "l" and is explicitly shown in the above rows and the keyword should be terminated by a "l".

Table 12.7: COMPSEGS Keyword Description

The total number of wells and completions should be defined via the WELLSDIMS keyword and the number of multi-segment wells and completions should be declared on the WSEGDIMS keyword, both keywords are in the RUNSPEC section.

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See also the WELSPECS keyword to define wells, the COMPDAT keyword to define the well completions for both ordinary wells and multi-segment wells, and the COMPSEGS keyword to define a multi-segment segment completions. All the aforementioned keywords are described in the SCHEDULE section.

Revision: Rev-I

Example

The following example defines the completions for two oil producing segment oil wells (OP01 and OP02) using the COMPSEGS keywords.

```
_ _
         COMPLETION SEGMENT SPECIFICATION DATA
-- WELL
-- NAME
COMPSEGS
0P01
- -
         --LOCATION--
                       BRAN TUBING NODAL
                                               DIR LOC
                                                                    COMP
                                                                             ISEG
                                                            MTD
_ _
         ΙI
              JJ K1
                       NO
                              LENGTH DEPTH
                                               PEN I, J, K PERFS
                                                                    LENGTH
         10
                              2512.5
                                       2525.0
              10
                   1
                        1
         10
              10
                        1
                              2525.0
                                       2550.0
         10
              10
                   3
                        1
                              2550.0
                                       2575.0
              10
                              2575.0
                                       2600.0
         10
                   4
                        1
         10
              10
                   5
                              2600.0
                                       2625.0
         10
              10
                   6
                        1
                              2625.0
                                       2650.0
          9
              10
                         2
                              2637.5
                                       2837.5
          8
                   2
                        2
                              2837.5
              10
                                       3037.5
           7
              10
                   2
                         2
                              3037.5
                                       3237.5
                   2
           6
              10
                         2
                              3237.5
                                       3437.5
           5
                   2
                         2
              10
                              3437.5
                                       3637.5
         COMPLETION SEGMENT SPECIFICATION DATA
-- WELL
-- NAME
COMPSEGS
0P02
         --LOCATION--
- -
                       BRAN
                             TUBING NODAL
                                               DIR
                                                    LOC
                                                            MID
         II
              JJ K1
                       NO
                              LENGTH
                                       DEPTH
                                               PEN
                                                    I,J,K
                                                            PERFS
          1
               9
                        1
                              2662.5
                                       2862.5
           1
               8
                   3
                         1
                              2862.5
                                       3062.5
          1
               7
                                       3262.5
                   3
                        1
                              3062.5
           1
               6
                        1
                              3262.5
                                       3462.5
               5
                   3
                        1
                              3462.5
                                       3662.5
           2
              10
                         2
                              2712.5
                                       2912.5
           2
                         2
              10
                   5
                              2912.5
                                       3112.5
           4
                         2
              10
                   5
                              3112.5
                                       3312.5
           5
                         2
              10
                   5
                              3312.5
                                       3512.5
           6
              10
                   5
                        2
                              3512.5
                                      3712.5
           1
               9
                   6
                         3
                              2737.5
                                       2937.5
           1
               8
                         3
                              2937.5
                                       3137.5
                   6
           1
               7
                   6
                         3
                              3137.5
                                       3337.5
           1
               6
                   6
                         3
                              3337.5
                                       3537.5
                              3537.5
                                       3737.5
```

Note that the COMPDAT keyword in the SCHEDULE section must also be defines for these two wells.

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12.3.15 DATES - Advance Simulation by Reporting Date

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword advances the simulation to a given report date after which additional keywords may be entered to instruct OPM Flow to perform additional functions via the SCHEDULE section keywords, or further DATES data sets or keywords may be entered to advance the simulator to the next report date.

If the DATES keyword is to be used during the simulation, then the START keyword in the RUNSPEC section must be declared to set the start date for the run.

No.	Name	Description	Default
I	DAY	A positive integer that defines the day of the month for the data set, the value should be greater than or equal to one and less than or equal to 31.	None
2	MONTH	Character string for the month for the data set and should be one of the following 'JAN', 'FEB', 'MAR', 'APR', 'MAY', 'JUN', 'JUL' (or 'JLY'), 'AUG', 'SEP', 'OCT', 'NOV', or 'DEC'	None
3	YEAR	A positive four digit integer value representing the year for the data set, which must be specified fully by four digits, that is 1986.	None
4	TIME	A numeric character string that defines the time for the data set in the form of:	00:00:00
		HH;MM:SS:SSSS	
		The default value means in most cases this parameter can be defaulted. TIME is normally used when detailed DST matching is performed to enable the pressures and rates to be stated at specific dates and times.	

Notes:

- 1) The keyword is followed by a number of data sets (or rows) representing one DATE record per row.
- 2) Each record (or row) is terminated by "/" and the keyword is terminated by a "/".

Table 12.8: DATES Keyword Description

See also the TSTEP keyword in the SCHEDULE section.

Whenever possible it is a good idea to always set the start date to be at the beginning of the year, as like most simulators, OPM Flow reports are always stated at the number of days from the start date (and sometimes at a given date). If the start date is at the beginning of the year, then calculating the actual date is relatively straight forward and simple.

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Examples

Given a start date of January I, 2020 set via the START keyword in the RUNSPEC section, the following example advances the simulator from the start date of January I, 2020 to January I, 2021, using quarterly reporting time steps.

```
-- SCHEDULE SECTION
SCHEDULE
-- SCHEDULE SECTION - 2020-01-01
RPTSCHED
       'WELLS=2'
                 'WELSPECS'
                            'CPU=2'
                                    FIP=2'
DATES
       2 JAN
             2020 /
RPTSCHED
        'NOTHING'
DATES
       1 APR
              2020 /
       1 JLY
              2020 /
       1 OCT
              2020 /
-- SCHEDULE SECTION - 2021-01-01
RPTSCHED
       'WELLS=2' 'WELSPECS' 'CPU=2'
                                    FIP=2'
DATES
       1 JAN 2021 /
RPTSCHED
        'NOTHING'
DATES
       1 APR
              2021 /
       1 JLY
              2021 /
       1 OCT
              2021 /
```

The above example writes out a series of report at the start of the run and then advances the simulation one day to January 2, 2020 and switches off the reporting. The simulation then advances to April I, July I and October I, 2020 with no further changes to the run. After October I, 2020 reporting is switched on again to enable a report on January I, 2021, which is then subsequently switched off after the January I, 2021 report time step has been completed.

Note if one wishes to terminate the run at the end of year (as opposes to the beginning of the year and get a final report for the year, then the next example demonstrates the keyword sequence to enable this.

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```
-- SCHEDULE SECTION - 2021-01-01
RPTSCHED
         'WELLS=2' 'WELSPECS' 'CPU=2' FIP=2'
DATES
         2 JAN 2021 /
RPTSCHED
        'NOTHING'
DATES
         1 FEB
                 2021 /
                 2021 /
         1
           MAR
           APR
         1
                 2021 /
         1 MAY
                 2021 /
                 2021 /
         1
           JUN
           JLY
                 2021
         1
           AUG
                 2021
         1
                 2021 /
           SEP
           0CT
                 2021 /
         1 NOV
                 2021 /
                 2021 /
         1 DEC
-- FINAL REPORT AND RESTART AT YEAR END
RPTSCHED
                                 'CPU=2'
         'WELLS=2'
                    'WELSPECS'
                                            FIP=2'
RPTRST
         'BASIC=2'
DATES
                  2021 /
          31 DEC
```

In the above example monthly reporting time steps have been used instead of quarterly and report is requested after the December 1, 2021 time step and is therefore written out on December 31, 2021.

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12.3.16 DRSDT - Solution Gas (Rs) Maximum Rate of Increase Parameters

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

DRSDT defines the maximum rate at which the solution gas-oil ratio (Rs) can be increased in a grid cell. The keyword is similar in functionality to the DRSDTR keyword, that defines the maximum rate at which Rs can be increased in a grid cell by region. Both keywords should only be used if the OIL, GAS, and DISGAS keywords in the RUNSPEC section have been invoked to allow oil, gas and dissolved gas to be present in the model. The keyword only affects the behavior of an increasing Rs, for example when gas is being injected into an oil reservoir, and is subject to the availability of free gas and the ability of the undersaturated oil to adsorb this gas.

No.	Name	Description					
		Field	Metric	Laboratory	1		
I	DRSDTI	which the solution gas-oi	cive number that defines il ratio is allowed to incre as can dissolve into the a	ase in a grid cell, that is			
		dissolve into the unsatur	that Rs cannot increas rated oil in a grid cell. Al Rs to increase rapidly unt ock is fully saturated.	ternatively a very large			
		Note if the keyword is not present in the input deck then DRSDTI is assumed to be a very large number resulting in complete re-solution of the gas into the available undersaturated oil.					
		Mscf/stb/day	sm³/sm³/day	scc/scc/day	None		
2	DRSDT2	applied to either all grid gas: 1) ALL: means th	racter string that defines of blocks or just those grid the DRSDTI maximum rate a grid cell is applied to all	I blocks containing free	ALL		
		,	to increase in a grid cell is applied to grid blocks only containing				
		Note if the keyword is not present in the input deck then DRSDT2 is set to the default value of ALL.					

Table 12.9: DRSDT Keyword Description

Note this keyword can be used in history matching field performance to control the availability of the movable gas phase.

See also the VAPPAR keyword in the SOLUTION section and the DRSDTR, DRVDT and DRVDTR keywords in the SCHEDULE section that controls how vaporized oil is treated and the rate at which the dissolved phase ratio increases within a grid block.

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Examples

The first example prevents the solution gas-oil ratio from increasing and applies this to all grid cells.

Revision: Rev-I

```
-- SOLUTION GAS (RS) MAXIMUM RATE OF INCREASE FOR MODEL
-- DRSDT
-- MAX RS ALL/FREE
-- DRSDT1 DRSDT2
-- 0.000 ALL //
```

And the second example below applies 0.005 Mscf/stb/day as the maximum rate at which the solution gas-oil ratio is allowed to increase in a grid cell, and applies this to only cells containing free gas.

```
-- SOLUTION GAS (RS) MAXIMUM RATE OF INCREASE FOR MODEL
-- DRSDT
-- MAX RS ALL/FREE
-- DRSDT1 DRSDT2
-- 0.0005 FREE //
```

Again, the keyword parameters when applied are subject to the availability of free gas and the ability of the undersaturated oil to adsorb this gas.

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12.3.17 DRSDTR - Solution Gas (Rs) Maximum Rate of Increase Parameters by Region

F	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

DRSDTR defines the maximum rate at which the solution gas-oil ratio (Rs) can be increased in a grid cell for various regions in the model. The keyword is similar in functionality to the DRSDT keyword, that defines the maximum rate at which Rs can be increased in a grid cell for all cells in the model. The number of DRSDTR vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the DRSDTR records to different grid blocks in the model is done via the PVTNUM keyword in the REGION section. One data set consists of one record or line which is terminated by a "/".

DRSDTR should only be used if the OIL, GAS, and DISGAS keywords in the RUNSPEC section have been invoked to allow oil, gas and dissolved gas to be present in the model. The keyword only affects the behavior of an increasing Rs, for example when gas is being injected into an oil reservoir, and is subject to the availability of free gas and the ability of the undersaturated oil to adsorb this gas.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	DRSDTI	which the solution gas-oi	ive number that defines il ratio is allowed to incre as can dissolve into the a	ase in a grid cell, that is				
		dissolve into the unsaturality value of DRSDT1 allows	A value of zero means that Rs cannot increase and free gas cannot dissolve into the unsaturated oil in a grid cell. Alternatively a very large value of DRSDTI allows Rs to increase rapidly until there is no free gas or the oil within the grid block is fully saturated.					
		assumed to be a very la	Note if the keyword is not present in the input deck then DRSDTI is assumed to be a very large number resulting in complete re-solution of the gas into the available undersaturated oil.					
		Mscf/stb/day	sm³/sm³/day	scc/scc/day	None			
2	DRSDT2	applied to either all grid gas:	DRSDT2 is a defined character string that defines whether the DRSDT1 is applied to either all grid blocks or just those grid blocks containing free gas:					
		ALL: means the DRSDT1 maximum rate at which Rs is allowed to increase in a grid cell is applied to all grid blocks.						
		,	to increase in a grid cell is applied to grid blocks only containing					
		Note if the keyword is n to the default value of AL	ot present in the input de L.	ck then DRSDT2 is set				

Notes:

- 1) The keyword is followed by NTPVT records as declared on the TABDIMS keyword in the RUNSPEC section.
- 1) Each record is terminated by "/" and there is no "/" terminator for the keyword.

Table 12.10: DRSDTR Keyword Description

Note this keyword can be used in history matching field performance to control the availability of the movable gas phase.

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See also the VAPPAR keyword in the SOLUTION section and the DRSDT, DRVDT and DRVDTR keywords in the SCHEDULE section that controls how vaporized oil is treated and the rate at which the dissolved phase ratio increases within a grid block.

Revision: Rev-I

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness, as it is expected to be available in the next release of OPM Flow.

Examples

The first example prevents the solution gas-oil ratio from increasing and applies this to all regions for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to three.

```
-- SOLUTION GAS (RS) MAXIMUM RATE OF INCREASE BY REGION
-- DRSDTR
-- MAX RS ALL/FREE
-- DRSDT1 DRSDT2
-- 0.0000 ALL /
0.0000 ALL /
0.0000 ALL /
```

The second example below prevents the solution gas-oil ratio from increasing and applies this to all grid cells in PVTNUM region one. For PVTNUM regions one and two the keyword applies 0.005 Mscf/stb/day as the maximum rate at which the solution gas-oil ratio is allowed to increase in a grid cell, and applies this to only cells containing free gas.

```
-- SOLUTION GAS (RS) MAXIMUM RATE OF INCREASE BY REGION
-- DRSDTR
-- MAX RS ALL/FREE
-- DRSDT1 DRSDT2
-- 0.0000 ALL /
0.0005 FREE //
```

Again, the keyword parameters when applied are subject to the availability of free gas and the ability of the undersaturated oil to adsorb this gas.

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12.3.18 DRVDT - Solution OIL (Rv) MAXIMUM RATE OF INCREASE PARAMETERS

RUNSPEC GRID EDIT PROPS REGIONS SC	SOLUTION SUMMARY	SCHEDULE
------------------------------------	------------------	----------

Revision: Rev-I

Description

DRVDT defines the maximum rate at which the solution oil-gas ratio or condensate-gas ratio (Rv) can be increased in a grid cell. The keyword is similar in functionality to the DRVDTR keyword, that defines the maximum rate at which Rv can be increased in a grid cell by region. Both keywords should only be used if the OIL, GAS, and VAPOIL (condensate) keywords in the RUNSPEC section have been invoked to allow oil, gas and condensate to be present in the model. The keyword only affects the behavior of an increasing Rv, for example when gas is being injected into a gas condensate reservoir as part of as gas re-cycling scheme, and is subject to the availability of free oil (condensate) and the ability of the undersaturated gas to adsorb this condensate.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
I DRVE	DRVDTI	DRVDTI is a real posit which the solution oil-gas the maximum rate at undersaturated gas.	s ratio is allowed to incre	ase in a grid cell, that is		
		A value of zero means that Rv cannot increase and free oil cannot dissolve into the unsaturated gas in a grid cell. Alternatively a very large value of DRVDTI allows Rv to increase rapidly until there is no free oil or the gas within the grid block is fully saturated.				
		Note if the keyword is assumed to be a very lar the oil into the available u	rge number resulting in (
		stb/Mscf/day	sm³/sm³/day	scc/scc/day	None	

Table 12.11: DRVDT Keyword Description

Note this keyword can be used in history matching field performance to control the availability of the movable gas phase.

See also the VAPPARS keyword in the SOLUTION section and the DRVDTR, DRSDT and DRSDTR keywords in the SCHEDULE section that controls how vaporized oil is treated and the rate at which the dissolved phase ratio increases within a grid block.

Examples

The example prevents the solution oil-gas ratio from increasing.

```
- -
         SOLUTION OIL (RV) MAXIMUM RATE OF INCREASE FOR MODEL
DRVDT
         MAX RV
         DRVDT1
         0.000
```

Again, the keyword parameters when applied are subject to the availability of free oil and the ability of the undersaturated gas to adsorb this oil.

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12.3.19 DRVDTR - SOLUTION OIL (Rv) MAXIMUM RATE OF INCREASE PARAMETERS BY REGION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

DRVDTR defines the maximum rate at which the solution oil-gas ratio or condensate-gas ratio (Rv) can be increased in a grid cell for various regions in the model. The keyword is similar in functionality to the DRVDT keyword, that defines the maximum rate at which Rv can be increased in a grid cell for all cells in the model. The number of DRVDTR vector data sets is defined by the NTPVT parameter on the TABDIMS keyword in the RUNSPEC section and the allocation of the DRVDTR records to different grid blocks in the model is done via the PVTNUM keyword in the REGION section. One data set consists of one record or line which is terminated by a "/".

This keyword should only be used if the OIL, GAS, and VAPOIL (condensate) keywords in the RUNSPEC section have been invoked to allow oil, gas and condensate to be present in the model. The keyword only affects the behavior of an increasing Rv, for example when gas is being injected into a gas condensate reservoir as part of as gas re-cycling scheme, and is subject to the availability of free oil (condensate) and the ability of the undersaturated gas to adsorb this condensate.

No.	Name		Description				
		Field	Field Metric Laboratory				
I	DRVDTI	which the solution oil-ga	DRVDTI is a real positive number that defines the maximum rate at which the solution oil-gas ratio is allowed to increase in a grid cell, that is the maximum rate at which the oil can dissolve into the available undersaturated gas.				
		into the unsaturated gas DRVDTI allows Rv to in	A value of zero means that Rv cannot increase and free oil cannot dissolve into the unsaturated gas in a grid cell. Alternatively a very large value of DRVDTI allows Rv to increase rapidly until there is no free oil or the gas within the grid block is fully saturated.				
		assumed to be a very la	Note if the keyword is not present in the input deck then DRVDTI is assumed to be a very large number resulting in complete re-solution of the oil into the available undersaturated gas.				
		stb/Mscf/day	sm³/sm³/day	scc/scc/day	None		

Notes:

- 1) The keyword is followed by NTPVT records as declared on the TABDIMS keyword in the RUNSPEC section.
- 1) Each record is terminated by "/" and there is no "/" terminator for the keyword.

Table 12.12: DRVDTR Keyword Description

Note this keyword can be used in history matching field performance to control the availability of the movable gas phase.

See also the VAPPARS keyword in the SOLUTION section and DRVD, DRSDT, and DRSDTR keywords in the SCHEDULE section that controls how vaporized oil is treated and the rate at which the dissolved phase ratio increases within a grid block.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness, as it is expected to be available in the next release of OPM Flow.

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Examples

The first example prevents the solution oil-gas ratio from increasing and applies this to all regions for when NTPVT on the TABDIMS keyword in the RUNSPEC section is set to three.

Revision: Rev-I

```
-- SOLUTION OIL (RV) MAXIMUM RATE OF INCREASE PARAMETERS BY REGION
-- DRVDTR
-- MAX RV
-- DRVDT1
-- 0.000 /
0.000 /
0.000 /
```

The second example below prevents the solution oil-gas ratio from increasing and applies this to all grid cells in PVTNUM region one. For PVTNUM regions one and two the keyword applies 0.005 stb//Mscf/day as the maximum rate at which the solution oil-gas ratio is allowed to increase in a grid cell,

```
-- SOLUTION OIL (RV) MAXIMUM RATE OF INCREASE PARAMETERS BY REGION
-- DRVDTR
-- MAX RV
-- DRVDT1
-- 0.0000 /
0.0005 /
0.0005 /
```

Again, the keyword parameters when applied are subject to the availability of free oil and the ability of the undersaturated gas to adsorb this oil.

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12.3.20 ENDACTIO - END THE DEFINITION OF ACTION COMMANDS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

The ENDACTIO keyword defines the end of a series of conditions that invoke run time processing of the ACTION series of keywords, namely: ACTION, ACTIONG, ACTIONR, ACTIONS, ACTIONW and ACTIONX. Only the ACTIONX keyword is implemented in OPM Flow as this keyword implements the ACTION, ACTIONG, ACTIONR, ACTIONS, ACTIONW functionality with greater flexibility. See the ACTIONX keyword in the SCHEDULE section for a full description of the ACTION facility.

There is no data required for this keyword and there is no terminating "/" for this keyword.

Example

The example shows the use of the ACTIONX and ENDACTIO keywords to test if the field's gas production rate is less than 600 MMscf/d after 2020 and to open up additional wells if this occurs.

```
-- START OF ACTIONX FIELD PHASE-2 DEVELOPMENT DEFINITION
ACTIONX
    PHASE2
    GGPR 'FIELD' < 600E3 AND
    YEAR > 2020
-- WELL PRODUCTION STATUS
                   --LOCATION-- COMPLETION
   WELL
           WELL
                   I J K FIRST LAST
   NAME
            STAT
WELOPEN
GP10
            OPEN
GP11
            OPEN
-- END OF ACTIONX FIELD PHASE-2 DEVELOPMENT DEFINITION
ENDACTIO
```

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12.3.21 ENDBOX - Define the End of the BOX Defined Grid

This keyword marks the end of a previously defined BOX sub-grid as defined by a previously entered BOX keyword. The keyword resets the input grid to be the full grid as defined by the NX, NY, and NZ variables on the DIMENS keyword in the RUNSPEC section.

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See ENDBOX – Define the End of the BOX Defined Grid in the GRID section for a full description.

12.3.22 ENDFIN - END THE DEFINITION OF A LOCAL GRID REFINEMENT

ENDFIN defines the end of a Cartesian or radial local grid refinement ("LGR") definition and a LGR property definition data set.

See ENDFIN - End the Definition of a Local Grid Refinement in the GRID section for a full description.

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12.3.23 GCONINJE - Group Injection Targets and Constraints

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The GCONINJE keyword defines injection targets and constraints for groups, including the top most group in the group hierarchy known as the FIELD group. Wells are allocated to groups when the wells are specified by the WELSPECS keyword in the SCHEDULE section. Wells defined to be under group control will have their injection rates controlled by the group to which they belong, in addition to any well constraints defined for the wells.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	GRPNAME	group name for which the	p to eight characters in ne group target and constr is the top most group and or the whole field.	raints are being defined.	None
		keyword when there is	nierarchy should be defin more than one level of gr nder the FIELD group in th	oups, otherwise all the	
2	TYPE		ing that defines the type the following character stri	•	None
		I) GAS: for a gas inj			
		2) OIL: for a water i	·		
		3) WAT: for a water	injection well.		
3	TARGET	group, all the other phas will attempt to meet the	ing that sets the target in es will therefore act as co TARGET based on the ph rd. TARGET should be set	onstraints. The simulator wase rate stated in items	None
		I) NONE: the grou are still defined a	p has no target phase, bu nd active.	t if entered, constraints	
		2) FLD: this group the FIELD group.	is controlled from a high	er level group, including	
		for the phase de	on phase will be control I fined by the TYPE variable WAT then this would me vitem (4).	e. For example, if TYPE	
		4) RESV: the target defined by item (is set to the in situ re 5).	servoir volume rate as	
		by TYPE multiplie	is set to groups production and by the value on item (6 WAT then this would no plied by item (6).	6). For example, if TYPE	
		6) VREP: the target defined by item (is set to the groups voida 7).	ge replacement ratio as	

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No.	Name		Description		Default		
		Field	Metric	Laboratory			
4	RATE		nat defines the maximum he phase declared by the				
		Liquid stb/d	Liquid sm³/day	Liquid scc/hour			
		Gas Mscf/d	Gas sm³/day	Gas scc/hour	None		
5	RESV	•	A real positive value that defines the maximum reservoir volume injection rate target or constraint.				
		rtb/d	rm³/day	rcc/hour	None		
6	REIN		at defines the target or I phase defined by the TYF				
		For example, if TYPE is ed of the produced gas will I	qual to GAS and REINJ is be re-injected.	equal to 0.85, then 85%			
		dimensionless	dimensionless	dimensionless	None		
7	VREP	•	t defines the target or co	•			
			equal to WAT and VREP eservoir volume will be re				
		dimensionless	dimensionless	dimensionless	None		
8	8 GRPCNTL	higher level group contro	ing that determines if the old.		YES		
		the flow rates for					
		 NO: then this group is NOT subject to a higher level group's control and the flow rates for this group will only be control by the parameters for this group. 					
		This variable is ignored if	GRPNAME is equal to FI	ELD.			
		This option is currently n	ot supported by OPM Flo	w.			
9	GRPGUIDE	as a dimensionless number	t defines a group's injection er. A group requires a value ce a specified proportion	ue for GRPGUIDE only			
		This feature is not support.	orted by OPM Flow and s	hould be defaulted with			
		dimensionless	dimensionless	dimensionless	None		
10	GUIPHASE		ng that sets the guide ph GUIPHASE should be set		None		
		I) RATE: the guide p	hase is set to the surface	injection rate.			
		2) RESV: the guide p	hase is set to the in situ re	eservoir volume rate.			
			rate is calculated at the be up's net voidage rate.	ginning of each time step			
		This feature is not support.	orted by OPM Flow and s	hould be defaulted with			
- 11		Not used should be defau	ulted with 1*.				

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No.	Name		Default					
		Field	Metric	Laboratory				
12		Not used should be defau	Not used should be defaulted with 1*.					
13		Not used should be defau	Not used should be defaulted with 1*.					

Revision: Rev-I

Notes:

I) The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.13: GCONINJE Keyword Description

See also the GRUPTREE keyword to define the hierarchy of the groups below the FIELD level, the GCONPROD keyword to define a group's production targets and constraints, the WCONPROD keyword to define a production well's targets and constraints, and the WCONINJE keyword to define an injection well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines the injection targets and constraints for the field and two groups that are one level below the field group, since the GRUPTREE keyword has not been entered to define the group hierarchy.

```
GROUP INJECTION TARGETS AND CONSTRAINTS
- -
-- GRUP FLUID CNTL
                     SURF
                             RESV
                                    REINJ VOID GRUP GUIDE GUIDE GRUP GRUP
                                    FRAC FRAC CNTL RATE
-- NAMF
        TYPE MODE
                     RATE
                            RATE
                                                              DEF
                                                                   REINJ RESV
GCONINJE
                                   1*
               VREP
                      35E3
                             1*
                                          1*
                                                       1*
                                                              1*
                                                                   1*
FIELD
        WAT
                                                  NO
                                   1*
                                                 YES 1*
                                                              1*
                                                                         1*
                     1*
                             1*
                                          1.0
                                                                    1*
GRP01
        WAT
               VREP
GRP02
         WAT
                                           1.0
```

In this example, group GRP01 and GRP02 are injecting water via voidage replacement with a voidage replacement of one and are under the control on the field group, that imposes a 35,000 m³/day total water injection limit.

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12.3.24 GCONPROD - Group Production Targets and Constraints

RUN	NSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The GCONPROD keyword defines production targets and constraints for groups, including the top most group in the group hierarchy known as the FIELD group. Wells are allocated to groups when the wells are specified by the WELSPECS keyword in the SCHEDULE section. Wells defined to be under group control will have their production rates controlled by the group to which they belong, in addition to any well constraints defined for the wells.

No.	No. Name Description							
		Field	Metric	Laboratory				
ı	GRPNAME	group name for which the The group named FIELD targets and constraints for Note that the group has keyword when there is a	A character string of up to eight characters in length that defines the group name for which the group target and constraints are being defined. The group named FIELD is the top most group and should be used to set targets and constraints for the field. Note that the group hierarchy should be defined by the GRUPTREE keyword when there is more than one level of groups, otherwise all the groups will sit directly under the FIELD group in the group tree hierarchy.					
2	TARGET	A defined character string that sets the target production phase for the group, all the other phases will therefore act as constraints. The simulator will attempt to meet the TARGET based on the phase rate stated in items (3) to (6) on this keyword. TARGET should be set to one of the following character strings: 1) NONE: the group has no target phase, but if entered, constraints are still defined and active. 2) FLD: this group is controlled from a higher level group, including the FIELD group. 3) ORAT: the target is set to the surface oil production rate as defined by item (3).						
			4) WRAT: the target is set to the surface water production rate as defined by item (4).					
		5) GRAT: the targe defined by item (t is set to the surface 5).	gas production rate as				
			t is set to the surface s defined by item (6).	liquid (oil plus water)				
		7) RESV: the target defined by item (rget is set to the in situ reservoir volume rate as					
3	ORAT	A real positive value that target or constraint.	defines the maximum sur	face oil production rate				
		stb/d	sm³/day	scc/hour	None			
4	WRAT	A real positive value that rate target or constraint.	t defines the maximum su	rface water production				
		stb/d	sm³/day	scc/hour	None			
5	GAS	A real positive value the rate target or constraint	at defines the maximum	surface gas production				
		Mscf/d	sm³/day	scc/hour	None			

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No.	Name			Default				
		Field	Metric	Laboratory				
6	LRAT	A real positive value the water) production rate to	at defines the maximum arget or constraint.	surface liquid (oil plus				
		stb/d	sm³/day	scc/hour	None			
7	ACTION		ing that defines the acti re violated. ACTION shorts:		None			
		I) NONE: no action	NONE: no action is taken.					
		 CON: close the well. If connection worst offending closed. 						
		 +CON: close the the worst offend completions then in the worst offen 						
		4) WELL: close the v						
		5) RATE: control the This effectively constraint.						
		The corrective action tak constraint is violated.	es places at the end of the	e time step in which the				
8	GRPCNTL	A defined character str higher level group contro	ing that determines if th	nis group is subject to	None			
		,	up is subject to a higher l this group will be adjuste	•				
		 NO: then this group is NOT subject to a higher level group's control and the flow rates for this group will only be control by the parameters for this group. 						
		This feature is currently r	not supported by OPM Flo	ow.				
9	GRPGUIDE	A real positive value expressed as a dimensi GRPGUIDE only if it is higher level group's rate						
		This feature is not support i*.	orted by OPM Flow and si	hould be defaulted with				
		dimensionless	dimensionless	dimensionless	None			

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No.	Name		Description		Default			
		Field	Metric	Laboratory				
10	GUIPHASE		ing that sets the guide pl GUIPHASE should be set		*			
		I) ORAT: the guide	phase is set to the surface	e oil production rate.				
		2) WRAT: the guide	phase is set to the surfac	e water production rate.				
		e gas production rate.						
		 LRAT: the guide phase is set to the surface liquid (oil plus water) production rate. 						
	5) RESV: the guide phase is set to the in situ reservoir volume rate.							
		This feature is not support it.	orted by OPM Flow and s	should be defaulted with				
-11		Not used should be defa	I *					
12		Not used should be defa	*					
13		Not used should be defa	Not used should be defaulted with 1*.					
14	RESV	A real positive value production rate target o	that defines the maxing constraint.	mum reservoir volume				
		rtb/d	rm³/day	rcc/hour	None			
15		Not used should be defa	ulted with I*.		I *			
16		Not used should be defa	ulted with I*.		I *			
17		Not used should be defa	ulted with I*.		I *			
18		Not used should be defa	ulted with 1*.		I *			
19		Not used should be defa	ulted with 1*.		I *			
20		Not used should be defa	ulted with 1*.		I *			
21		Not used should be defa	ulted with 1*.		I *			

Revision: Rev-I

Notes:

Table 12.14: GCONPROD Keyword Description

See also the GRUPTREE keyword to define the hierarchy of the groups below the FIELD level, the GCONINJ keyword to define a group's injection targets and constraints, the WCONPROD keyword to define a production well's targets and constraints, and the WCONINEJ keyword to define an injection well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

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⁾ The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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Example

The following example defines the production targets and constraints for the field and two groups that are one level below the field group, since the GRUPTREE keyword has not been entered to define the group hierarchy.

```
GROUP PRODUCTION CONTROLS
_ _
-- GRUP
         CNTL
                OIL
                       WAT
                               GAS
                                      LIO
                                              CNTL
                                                    GRUP
                                                           GUIDE GUIDE
                                                                          CNTL
-- NAME
         MODE
                RATE
                       RATE
                               RATE
                                      RATE
                                              0PT
                                                    CNTL
                                                           RATE
                                                                  DEF
                                                                          WAT
GCONPROD
         ORAT
                40E3
                       60E3
                               300E3
                                      60E3
                                                     1*
FIELD
                                              1*
                                                     1*
                                                            1*
                                                                   1*
                                                                           1*
GRP01
         FLD
                25E3
                       1*
                                      1*
                               1*
                                              1*
                                                     1*
         FLD
                25E3
GRP02
```

All groups are controlled by oil rate targets or constraints, but only the field level has water, gas and liquid constraints to reflect the actual production facility constraints. The wells under group control will be produced based on oil potential of each of the wells under group control, such that the field oil production target of 40,000 stb/d is honored and subject to the other phase fluid constraints. In addition, GRP01 and GRP02 oil rate values of 25,000 stb/d are constraints as these two groups are subject to the FIELD level targets and constraints.

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12.3.25 GECON - GROUP ECONOMIC CRITERIA FOR PRODUCTION GROUPS

	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
--	---------	------	------	-------	---------	----------	---------	----------	--

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Description

The GECON keyword defines economic criteria for production groups, including the field level group FIELD, that have previously been defined by the GCONPROD keywords in the SCHEDULE section.

Note that wells are allocated to a group when they are specified by the WELSPECS keyword and wells can also have economic controls. Wells under group control are therefore subject to the economic criteria set via the GCONPROD and CECON keywords in the SCHEDULE section and the controls specified by the WECON keyword.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	GRPNAME	group name for which the The group named FIELD targets and constraints for Note that the group he keyword when there is it	A character string of up to eight characters in length that defines the group name for which the group target and constraints are being defined. The group named FIELD is the top most group and should be used to set targets and constraints for the field. Note that the group hierarchy should be defined by the GRUPTREE keyword when there is more than one level of groups, otherwise all the groups will sit directly under the FIELD group in the group tree hierarchy.				
2	ORAT	production rate, below stopping all the wells in WELSPECS keyword.	A value less than or equal to zero switches of this criteria.				
		stb/d	sm³/day	scc/hour	0.0		
3	GAS	A real positive value the production rate, below stopping all the wells in WELSPECS keyword. A value less than or equal					
		Mscf/d	sm³/day	scc/hour	0.0		
4	WCUT	A real positive value that cut, above which an econ Water cut is defined as: that are available if the veron (7). A value less than or equal dimensionless	0.0				
	GOR	A real positive value tha	t defines the maximum e	conomic surface gas-oil			
		ratio, above which an eco (7).	onomic action will take p	lace, as defined by item			
			I to zero switches of this				
		Mscf/stb	sm³/sm³	scc/scc	0.0		

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No.	Name		Description		Default
		Field	Metric	Laboratory	
6	WGR	A real positive value that gas ratio, above which as item (7).			
		A value less than or equa	criteria.		
		stb/Mscf	sm³/sm³	scc/scc	0.0
7	ACTION	A defined character str economic WCUT, GOR, set to one of the followin	None		
		I) NONE: no action			
		CON: close the v If connections ha offending complet			
		3) +CON: close the wort offending completions then the worst offending			
		4) WELL: shut or sto	op the well as per the AU	ΓO variable on the	
		WELSPECS keywo	ord.		
		The corrective action take constraint is violated.	es places at the end of the	e time step in which the	
8	END	A defined character string all the producing wells in stopped. END should be	NO		
		I) NO: no action is t	aken and the run continu	es.	
		2) YES: terminate the	e run at the next report ti	me step.t	
9		Not used			

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Notes:

The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.15: GECON Keyword Description

See also the WELSPECS keyword to define a wells shut-in or stop options, GCONPROD for group controls, and WECON for setting a well's economic criteria. All the aforementioned keywords are described in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Example

The following example defines the economic criteria for the field with a minimum oil rate of 2,000 m³/day and a maximum water cut of 95%.

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```
GROUP ECONOMIC CRITERIA FOR PRODUCTION GROUPS
_ _
-- GRUP
         OIL
               GAS
                              GOR
                                     WGR
                                            WORK
                                                    END
                                                          MAX
-- NAME
                                                          WELLS
        MIN
               MIN
                      MAX
                              MAX
                                     MAX
                                            OVER
                                                    RUN
GECON
         2E3
               1*
                      0.95
                                1*
                                      1*
                                             CON
                                                    'YES' 1*
FIELD
```

If the economic limits are violated then the run will stop at the next report time step.

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12.3.26 GEFAC - DEFINE GROUP EFFICIENCY

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Defines a group's efficiency or up-time as opposed to setting the efficient factors for individual wells.

Note that wells are allocated to a group when they are specified by the WELSPECS keyword and wells can also have efficient factors.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
Í	GRPNAME	group name for which							
		Note that the group hierarchy should be defined by the GRUPTREE keyword when there is more than one level of groups, otherwise all the groups will sit directly under the FIELD group in the group tree hierarchy.							
2	FACTOR	A real positive value that is less than or equal to one that defines the efficiency factor for the group. If a group's down time is 5% then FACTOR should be set to 0.95 $(1.0 - 0.05)$.							
		dimensionless	dimensionless	dimensionless	1.0				
3	NETOPTN	Not used							

Notes:

Table 12.16: GEFAC Keyword Description

See also the WELFAC keyword in the SCHEDULE section to define a well's' efficiency factor.

Example

```
-- GROUP EFFICIENCY FACTORS
-- GRUP EFF NETWK
-- NAME FACT OPTN
-- GEFAC
PLATFORM 0.950 //
SUBSEA1 0.860 //
```

In the above example the group PLATFORM has it's efficiency factor (up time) set to 0.95 and the subsea group SUBSEAI has an up time of 0.860.

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¹⁾ The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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12.3.27 GLIFTOPT - DEFINE GROUP GAS OPTIMIZATION LIMITS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

GLIFTOPT defines a group's gas rate limits for when gas lift optimization has been activated via the LIFTOPT keyword in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.28 GNETINJE - DEFINE GROUP INJECTION NETWORK CONFIGURATION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The GNETINJE keyword defines the configuration of a group injection network when the network option has been activated.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.29 GRUPNET - Define Group Standard Network Parameters

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The GRUPNET keyword defines the standard group network parameters used to model the flow and pressures behavior within the network. The group hierarchy is defined by the GRUPTREE keyword and wells are assigned to groups using the WELSPECS keyword, both keywords are in the SCHEDULE section.

Group pressure values are optionally entered for each group in the network together with a vertical lift performance ("VFP") table that the determines the pipeline pressure behavior from the LOWER group to the HIGHER given the current flowing conditions; the group relationship is defined by the GRUPTREE keyword. The VPP table is entered via the VFPPROD keyword for production pipelines and VFPINJ for injection pipelines. Although these keywords are the same as used for well modeling, they are also used for pipeline modelling as well; however, the manner in which they are generated by an external software is completely different.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
I	GRPNAME	group name for which	to eight characters in the network parameters the top most group a	are being defined. The	None				
2	PRES	A real value that defines the fixed pressure for this group when the group is a terminating group. If the group is not a terminating group then PRES should be defaulted with I* or set to a negative number.							
		psia	atma	*					
3	VFPTAB	A positive integer greater than or equal to zero that defines the VFPPROD or VFPINJ vertical lift performance table to be used for calculating the pipeline pressures connecting the LOWER and HIGHER group in the network. Note that: 1) The default value of zero implies that there is no pipeline connecting							
		the LOWER and HIGHER groups. 2) If PRES is set to a real positive number then VFPTAB should be set to zero as this implies that GRPNAME is a terminating group and therefore there is no pipeline connecting GRPNAME to a HIGHER group.							
		 If PRES and VFPTAB are defaulted with I* or zero, then GRPNAME is not part of the networf. 							
		 IF VFPTAB is set equal to 9999 then this implies that there is no pressure change between the LOWER and HIGHER group. 							
		If a non-zero value is entered then the vertical lift performance tables must be entered via the VFPPROD or VFPINJ keyword in the SCHEDULE section.							

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No.	Name		Description		Default				
		Field	Metric	Laboratory	1				
4	ALQ-PIPE	-	t defines the artificial lift PROD assigned to the gro		0.0				
		WELL are used with the	ormance table and the art ne pipeline fluid rates to OWER and HIFGHER gro	calculate the pipeline					
		Note that the units for A on the VFPPROD keywordepending how the VFPP external program.							
5	OPTIONI	A defined character string that defines if a group's production target should be achieved by adjusting the tubing pressure of the wells within the group or by the adjusting the well rates by their guide rate. OPTIONI should be set to one of the following character strings:							
		YES: the group pr pressure of the w same tubing head into a common manifold.							
		If a group is using this option and has a higher group with production targets or constraints, than this group should have have it's guide rate set via the GCONPROD keyword in the SCHEDULE section, to ensure that the well's within this group operate at the same tubing head pressure.							
		2) NO: the group production target is achieved by adjusting the guide rates of the wells within the group. This is the standard method in matching group targets and may result with the wells within the having different tubing head pressures.							
		Only groups containing wells can use OPTION1 equal to YES or NO, a group without wells should set OPTION1 to NO.							
		Numerical convergence controls and iteration limits for wells using OPTIONI set equal to YES are defined via the NETBALAN keyword in the SCHEDULE section.							

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No.	Name		Description		Default						
		Field	Metric	Laboratory							
6	OPTION2			lift gas flows through the o one of the following	NO						
		 NO: no gas lift gas is allowed to flow through the pipeline only produced reservoir gas is allowed to flow through the pipeline. 									
		2) FLO: both gas lift gas and produced reservoir gas are allowed to flow through the pipeline. Gas lift gas is calculated from summing the calculated gas lift values of all the subordinate wells. Here the summed well gas lift gas (ALQ-WELL), the pipeline gas lift gas (ALQ-PIPE), and the reservoir produced gas are added to the gas flow rate along the pipeline.									
		3) ALQ: both gas lift gas and produced reservoir gas are allowed to flow through the pipeline. Gas lift gas is calculated from summing the calculated gas lift values of all the subordinate wells. Here the summed ALQ-WELL gas lift gas is added to the reservoir produced gas flow rate along the pipeline. This means that ALQ-PIPE gas lift gas value declared on item (4) is ignored.									
		If either FLO or ALQ have been selected then artificial lift quantity for the pipeline (ALQ-PIPE) and the wells (ALQ-WELL) must be defined as gas lift gas on the VFPROD tables. A well's specific gas lift gas quantity is set via the ALQ-WELL variable on the WCONPROD keyword in the SCHEDULE section.									
7	OPTION3	be reset to an equivale	ent surface oil or gas	ALQ-PIPE variable should density flowing along the the following character	NONE						
		DENO: set ALQ- flowing along the	•	urface density of the oil							
		DENG: set ALQ- flowing along the		urface density of the gas							
		3) NONE: the group production target is achieved by adjusting the guide rates of the wells within the group. This is the standard method in matching group targets and may result with the wells within the having different tubing head pressures.									
		the VFPPROD tables mus	t be based on the same ed when a mixture of	en artificial lift quantity on density parameter. These oil or gas with different							

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The keyword is followed by any numbers records with each record terminated by a "/" and the keyword I) should be terminated by a "/".

Table 12.17: GRUPNET Keyword Description

See also the WELSPECS keyword to define wells, the VFPPROD and VFPINJ keywords that the define vertical lift performance table to be used for calculating the pipeline pressures connecting the LOWER and HIGHER group in the network. All the aforementioned keywords are described in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Example

The following example defines a network based on two groups

```
DEFINE GROUP STANDARD NETWORK PARAMETERS
-- GRUP
         CNTL
                VFP
                        PUMP
                               MANIFOLD
                                         INCLUDE
                                                    ALQ
-- NAME
         PRES
                TABLE POWER
                              GROUP
                                         LIFT GAS
                                                    DENS
GRUPNET
         1200.
                1*
PROD-A
                        1*
                              'YES'
                                         1*
                                                    1*
PROD-B
                1
```

The next example is more complex and is taken form the Norne model.

```
DEFINE GROUP STANDARD NETWORK PARAMETERS
- -
                VFP
-- GRUP
         CNTL
                        PUMP
                               MANIFOLD INCLUDE
                                                    ALQ
                TABLE
                       POWER
                               GROUP
-- NAME
         PRES
                                         LIFT GAS
                                                   DENS
GRUPNET
FIELD
         20.0
                5*
                5*
PROD
         20.0
                8
MANI-B2
        1*
                                NO
MANI-B1
        1*
                8
                        1*
                                NO
         1*
                       4*
MANI-K1
               9999
B1-DUMMY 1*
                       4*
               9999
                                NO
MANI-D1 1*
                8
                        1*
MANI-D2
                8
MANI-K2 1*
               9999
                        4*
D2-DUMMY 1*
               9999
                        4*
MANI-E1 1*
                        1*
                                NO
                                         2*
                9
MANI-E2 1*
                9
                        4*
```

Here the FIELD controlling pressure is set at 20 barsa and the same limit is used for group PROD which sits directly under the FIELD group (see Figure 12.1)

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12.3.30 GRUPRIG - Group Drilling and Workover Rig Specifications

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Description

Defines a groups drilling and workover specifications.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate.

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12.3.31 GRUPTREE - DEFINE GROUP TREE HIERARCHY

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

GRUPTREE defines the group hierarchy of groups that have been created by having wells assigned to them via the WELSPECS keyword in the SCHEDULE section, By default three group levels are defined that sets the wells as level three, reporting directly to defined groups at level two, and the level two groups reporting to the FIELD group at level one. If a different configuration is required then the GRUPTREE keyword should be used to define the group hierarchy by defining a lower level group that reports directly to a higher level group.

No.	Name	Description	Default
I	LOWER	A character string of up to eight characters in length that defines the group name which belongs to the HIGHER group. The group named FIELD is the top most group and should NOT be used as as a group name for the LOWER group name.	None
		Undefined group relationships are automatically assigned to the FIELD group.	
2	HIGHER	A character string of up to eight characters in length that defines the HIGHER group name that the LOWER group belongs to. The group named FIELD is the top most group and can be used as as the HIGHER group name.	None
		Undefined group relationships are automatically assigned to the FIELD group.	

Notes:

Table 12.18: GRUPTREE Keyword Description

A group hierarchy can have any number of levels but groups that have other groups as LOWER groups cannot also have wells for the HIGHER group. Thus, a group either contains wells or has LOWER groups

See also the GCONPROD and GCONINJE for defining group production and injection volumes, and the WELSPECS keywords to allocate wells to groups. All the aforementioned keywords are described in the SCHEDULE section.

Examples

The first example defines PLAT01 and PLAT03 reporting to the FIELD level (default if these records are omitted) and PLAT02 reporting to PLAT01.

```
-- DEFINE GROUP TREE HIERARCHY
-- LOWER HIGHER
-- GROUP GROUP
GRUPTREE
PLAT01 FIELD /
PLAT02 PLAT01 /
PLAT03 FIELD //
```

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The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

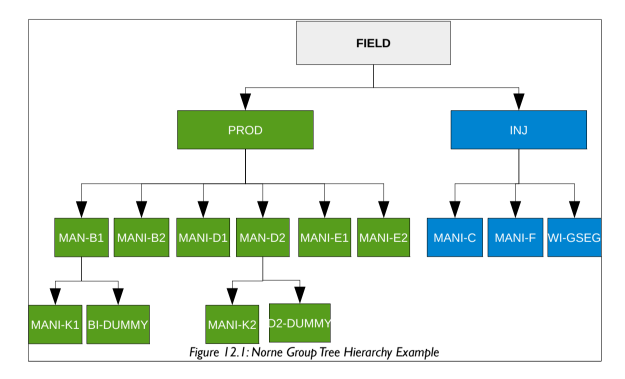
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The next example is more complex and is taken form the Norne model.

```
DEFINE GROUP TREE HIERARCHY
- -
          LOWER
                      HIGHER
          GROUP
                      GROUP
GRUPTREE
                     'FIELD'
         'INJE'
         'PROD'
                     'FIELD'
         'MANI-B2'
                     'PROD'
         'MANI-B1'
                     'PROD'
         'MANI-D1'
                     'PROD'
         'MANI-D2'
                     'PROD'
         'MANI-E1'
                     'PROD'
                     'PROD'
         'MANI-E2'
         'MANI-K1'
                     'MANI-B1'
         'MANI-K2'
                     'MANI-D2'
         'MANI-C'
                     'INJE'
         'MANI-F'
                     'INJE'
         'WI-GSEG'
                     'INJE'
         'B1-DUMMY' 'MANI-B1'
         'D2-DUMMY' 'MANI-D2'
```

The group hierarchy for this example is shown below.



Here groups PROD, INJ, MAN-B1, and MAN-D2 report to higher level groups and the other remaining groups all have individual wells allocated to them instead.

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12.3.32 GSATPROD - Define Group Satellite Production Rates

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

GSATPROD keyword defines a satellite group's oil, gas and water production rates in the model. Satellite groups are not connected to the reservoir model and therefore have no wells or subordinate groups associated with them, they are nevertheless connected to other higher level groups and higher level groups within a network model (if activated). They thus provide a means to "add-in" outside production to the model without modeling the "add-in" reservoir model.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.33 LIFTOPT - ACTIVATE GAS LIFT OPTIMIZATION

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The LIPTOPT keyword actives the gas lift optimization option.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.34 MULTFLT - MULTIPLY THE TRANSMISSIBILITY OF A DEFINED FAULT BY A CONSTANT

MULTFLT enables the transmissibilities across defined faults, as declared by the FAULTS keyword, to be modified. They keyword allows for the re-scaling of the existing fault transmissibilities calculated by OPM Flow, for example setting a fault to be completely sealing by setting the multiplier to zero..

Revision: Rev-I

See MULTFLT - Multiply the Transmissibility of a Defined Fault by a Constant in the GRID section for a full description.

12.3.35 MULTPV - MULTIPLY CELL PORE VOLUMES BY A CONSTANT

MULTPV multiples the pore volumes of a cell by a real positive constant for all the cells in the model via an array. An alternative to defining the complete array is to use the BOX keyword to define an area of the grid and then use the MULTPV keyword to set the multipliers just for the area defined by the BOX keyword (see the example).

See MULTPV - Multiply Cell Pore Volumes by a Constantin the GRID section for a full description.

12.3.36 MULTX - Multiply Cell Transmissibility in the +X Direction

MULTX multiples the transmissibility between two cell faces in the +X direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I, K) and (I+I, J, K).

See MULTX - Multiply Cell Transmissibility in the +X Direction in the GRID section for a full description.

12.3.37 MULTX- - MULTIPLY CELL TRANSMISSIBILITY IN THE -X DIRECTION

MULTX- multiples the transmissibility between two cell faces in the -X direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I-I, I, K) and (I, J, K).

See MULTX- - Multiply Cell Transmissibility in the -X Direction in the GRID section for a full description.

12.3.38 MULTY - Multiply Cell Transmissibility in the +Y Direction

MULTY multiples the transmissibility between two cell faces in the +Y direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I, K) and (I, J+I, K).

See MULTY - Multiply Cell Transmissibility in the +Y Direction in the GRID section for a full description.

12.3.39 MULTY- - MULTIPLY CELL TRANSMISSIBILITY IN THE -Y DIRECTION

MULTY- multiples the transmissibility between two cell faces in the -Y direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, I, K) between the cells (I, I-I, K) and (I, J, K).

See MULTY- - Multiply Cell Transmissibility in the -Y Direction in the GRID section for a full description.

	Α	В	С	D	Ε	F	G	Н	Κ	J	Κ	L	М	N	0	Р	Q	R	S	Т	U	V	W	Χ	Υ	Ζ
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12.3.40 MULTZ - MULTIPLY CELL TRANSMISSIBILITY IN THE +Z DIRECTION

MULTZ multiples the transmissibility between two cell faces in the +Z direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, J, K) between the cells (I, J, K) and (I, J, K+1).

Revision: Rev-I

See MULTZ - Multiply Cell Transmissibility in the +Z Direction in the GRID section for a full description.

12.3.41 MULTZ- - MULTIPLY CELL TRANSMISSIBILITY IN THE -Z DIRECTION

MULTZ- multiples the transmissibility between two cell faces in the -Z direction for all the cells in the model via an array, that is the keyword sets the transmissibility multiplier of block (I, J, K) between the cells (I, J, K-I) and (I, J, K).

See MULTZ- - Multiply Cell Transmissibility in the -Z Direction in the GRID section for a full description.

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12.3.42 NETBALAN - Network Balancing Parameters

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the network balancing parameters used to control how network balancing is perform on a network.

OPM Flow does not have this feature and hence this keyword is ignored by OPM Flow and the NETBALAN keyword has no effect on the simulation.

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12.3.43 NEXTSTEP - MAXIMUM NEXT TIME STEP SIZE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

This keyword defines the maximum time step size the simulator should take for the next time step. This keyword can be used to reset the time step for when known large changes to the model are taking place that may result in time step chops. For example, if the reporting time size is using monthly reporting steps via the DATES keyword in the SCHEDULE section, then if for example, a group of wells start production at a given date, then the NEXTSTEP keyword can be used to shorten the next step ito avoid a time step chop.

Time steps chops are computationally expensive as the simulator cannot solve the current time step at the given tolerance, and therefore has to reduce the time step size. For example, if the previous completed time step was at day 365 and the current time step ending at 396 days cannot be solved, then the simulator will reduce the current time step to perhaps end at day 370, if this still cannot solved then the time step will be be chopped back again to perhaps to less then one day. Using the NEXTSTEP keyword, the simulator is instructed to take a small time step in the anticipation that this will avoid time step chops and thus improve computational performance.

No.	Name	Description					
		Field	Metric	Laboratory			
1	NSTEPI	NSTEPI is a real positive value that defines the maximum length of the next time step.					
		days	days	hours	None		
2	NSTEP2	NSTEP2 is a character string that should be set to either NO or YES to state if the NSTEP1 should be applied to future reporting time steps. 1) NO: Means that NSTEP1 should not be applied to subsequent					
		reporting time steps. 2) YES: means that STEPI should be applied to subsequent reporting time steps.					
		The default value of NO	means that NSTEP1 will o	only be applied once.			

Table 12.19: NEXTSTEP Keyword Description

See also the DATES and TSTEP keyword in the RUNSPEC section that are used to advance the simulation through time.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness. See section 2.2 Running Flow on how to control time stepping for OPM Flow.

Examples

The first example shows the direct use of the NEXTSTEP keyword:,

```
-- NEXT ALL
-- STEP TIME
-- --- NEXTSTEP
1 'NO' /
```

Here the next step size is set to one day and should only be used once.

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The next example shows a more complete use of the keyword for when the field oil production has increased dramatically from 10,000 stb/d to 50,000 stb/d as indicated by the two GCONPROD keywords.

```
-- SCHEDULE SECTION - 2021-01-01
-- GROUP PRODUCTION CONTROLS
-- GRUP
           CNTL OIL
                          WAT
                                 GAS
                                         LIQ
                                                 CNTL
                                                       GRUP
                                                              GUIDE
                                                                     GUIDE
                                                                             CNTL
           MODE RATE
                                                       CNTL
-- NAME
                          RATE
                                 RATE
                                         RATE
                                                 OPT
                                                              RATE
                                                                     DEF
                                                                             WAT
GCONPROD
                                                                      1*
           'ORAT' 10E3
                                                 1*
                                                        1*
                                                               1*
                                                                              1*
'FIELD'
                          60E3
                                 300E3
                                         60E3
RPTSCHED
              'WELSPECS'
                             'CPU=2'
                                          'FTP=2'
'WELLS=2'
DATES
           2021 /
 2 JAN
RPTSCHED
'NOTHING'
DATES
1 FFB
           2021 /
1 MAR
           2021
-- GROUP PRODUCTION CONTROLS
-- GRUP
           CNTL
                  OIL
                                                 CNTI
                                                       GRUP
                                                              GUIDE
                                                                     GUTDE
                                                                             CNTI
                          WAT
                                 GAS
                                         LIO
-- NAME
           MODE
                 RATE
                          RATE
                                 RATE
                                         RATE
                                                 OPT
                                                       CNTL
                                                              RATE
                                                                     DEF
                                                                             WAT
GCONPROD
                                                        1*
                                                               1*
                                                                       1*
                                                                              1*
           'ORAT' 50E3
                                                 1*
'FIELD'
                          90E3
                                 300E3
                                         90F3
         NEXT
                 ALL
         STEP
                 TIME
- -
NEXTSTEP
         1
                'NO'
DATES
    APR
           2021
    MAY
           2021
 1
 1
    JUN
           2021
 1
    JLY
           2021
    AUG
           2021
 1
 1
    SEP
           2021
 1
    OCT
           2021
    NOV
           2021
 1
           2021
 1
    DEC
```

Given a start date of January 1, 2020 set via the START keyword in the RUNSPEC section, the above example shows the initial oil production of 10,000 stb/d starting in January 1, 2020. and continuing up to March 1, 2021. At the March 1, 2021 time step the field oil production rate is increased to 50,000 stb/d and the maximum next time step is set to one day. After the one day time step is completed (March 2, 2012), the simulator will progressively in increase the time step size until a maximum of 31 days is reached. The 31 day maximum is a result of requesting monthly time steps via the DATES keyword. The intent of using the NEXTSTEP keyword in this case is to prevent time step chops occurring due to the "shock" to the system caused by the large increase in oil production.

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12.3.44 NUPCOL - Define the Number of Newtonian Iterations Used to Update Well Targets

The NUPOL keyword defines the maximum number of Newtonian iterations within a time step that may be used to update the well production and injection targets, after which the well targets will be frozen until the time step calculations have converged and the time step is complete.

Revision: Rev-I

See NUPCOL – Define the Number of Newtonian Iterations Used to Update Well Targets in the RUNSPEC section for a full description.

See section 2.2 Running Flow on how to set the numerical control parameters for OPM Flow.

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12.3.45 PIMULTAB - Define Well Productivity Index versus Water Cut Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

PIMULTAB defines productivity index multiplier versus water cut tables that are used to scaled a well's connection factors based on connection's current producing water cut. The tables are used for modeling the productivity decline due to increasing water cut. Allocation of the tables to a well is via the WPITAB keyword in the SCHEDULE section.

No.	Name		Default					
		Field	Metric	Laboratory				
I	WCUT	maximum surface water	A real monotonically increasing positive columnar vector that defines the maximum surface water cut for the corresponding PIMULT vector. Water cut is defined as $f_{w} = \frac{q_{w}}{q_{w} + q_{o}} \; .$					
		dimensionless	dimensionless	dimensionless	None			
2	PIMULT	A real positive decreasir index multiplier used corresponding WCUT ve						
		dimensionless	dimensionless	dimensionless	None			

Notes:

- 1) The keyword is followed by NTPIMT tables as stated on the PIMTDIMS keyword in the RUNSPEC section.
- 2) Each table must contain a minimum of two rows and a maximum of NRPIMT rows as declared on the PIMTDIMS keyword in the RUNSPEC section.
- 3) Each table is terminated by "/" and there is no "/" terminator for the keyword.

Table 12.20: PIMULTAB Keyword Description

See also the WPITAB keyword that allocates the tables to the wells, and also the WPIMULT keyword that scales a well's productivity index by a constant value, both of which are in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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Examples

Given NTPIMT equals two and NRPIMT equals four on PIMTDIMS keyword in the RUNSPEC section, then:

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```
DEFINE WELL PRODUCTIVITY INDEX VERSUS WATER CUT TABLES
- -
         WCUT
                   MULT
- -
PIMULTAB
         0.0000
                  1.0000
         0.2500
                   0.9500
         0.5000
                  0.8500
         0.7500
                  0.7500
                   1.0000
         0.0000
         0.2500
                   0.9500
         0.5000
                   0.8500
         0.7500
                   0.7500
```

The next example is summarized from the Norne model with NTPIMT equals one and NRPIMT equals to 51 on the PIMTDIMS keyword in the RUNSPEC section.

```
DEFINE WELL PRODUCTIVITY INDEX VERSUS WATER CUT TABLES
_ _
         The following is the reviewed model in Aug-2006, low-high case
         a=0.25, b=0.1; PIMULT=(1-a)/exp(fw/b)+a
         MAX
                   PT
_ _
         WCUT
                   MULT
_ _
PIMULTAB
         0.000
                   1.0000
         0.025
                   0.8341
         0.050
                   0.7049
         0.075
                   0.6043
         0.100
                   0.5259
         0.125
                   0.4649
         0.150
                   0.4173
         0.175
                   0.3803
         0.200
                   0.3515
         0.225
                   0.3290
         0.250
                   0.3116
         0.275
                   0.2979
         0.300
                   0.2873
         0.325
                   0.2791
         0.350
                   0.2726
         0.375
                   0.2676
         0.400
                   0.2637
         0.425
                   0.2607
         0.450
                   0.2583
         0.475
                   0.2565
         0.500
                   0.2551
         0.525
                   0.2539
         0.550
                   0.2531
         0.575
                   0.2524
         0.600
                   0.2519
         0.625
                   0.2514
         0.650
                   0.2511
         0.675
                   0.2509
         0.700
                   0.2507
         0.725
                   0.2505
         0.750
                   0.2504
```

0.775

0.2503

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0.800	0.2503
0.825	0.2502
0.850	0.2502
0.875	0.2501
0.900	0.2501
0.925	0.2501
0.950	0.2501
0.975	0.2500
1.000	0.2500 /

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12.3.46 PLYADS - Define Polymer Rock Adsorption Tables

The PLYADS keyword defines the rock polymer adsorption tables for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section. Alternatively, the functions can be entered via the PLYADSS keyword in the PROPS section for when salt sensitivity is to be considered.

Revision: Rev-I

See PLYADS - Define Polymer Rock Adsorption Tables in the PROPS section for a full description.

12.3.47 PLYDHFLF - Define Polymer Thermal Degradation Half-Life Tables

The PLYDHFLF keyword defines the polymer thermal degradation half-life with respect to temperature functions for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

See PLYDHFLF - Define Polymer Thermal Degradation Half-Life Tables in the PROPS section for a full description.

12.3.48 PLYMAX - Define Polymer-Salt Viscosity Mixing Concentrations

The PLYMAX keyword defines maximum polymer and salt concentrations that are to be used in the mixing parameter calculation of the fluid component viscosities, for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

See X in the PROPS section for a full description.

12.3.49 PLYSHEAR - ACTIVATE AND DEFINE POLYMER SHEARING PARAMETERS

The PLYSHEAR keyword activates and the defines the polymer shear thinning-thickening option for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

See X in the PROPS section for a full description.

12.3.50 PLYSHLOG - ACTIVATE AND DEFINE THE POLYMER SHEARING LOGARITHMIC PARAMETERS

This keyword activates and defines the parameters for the logarithm-based polymer shear thinning/thickening option.

See PLYSHEAR – Activate and Define Polymer Shearing Parameters in the PROPS section for a full description.

12.3.51 PLYVISC - Define Polymer Viscosity Scaling Fact

PLYSVISC defines the polymer viscosity scaling factors used to determine the relationship of pure water viscosity with respect to increasing polymer saturation within a grid block. The polymer option must be activated by the POLYMER keyword in the RUNSPEC section in order to use this keyword.

See PLYVISC - Define Polymer Viscosity Scaling Factors in the PROPS section for a full description.

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12.3.52 PRORDER - DEFINE A GROUP PRODUCTION RULES SEQUENCE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Defines the order of group production rules to be implemented when a group's target is not satisfied.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.53 REFINE - START THE DEFINITION OF A LOCAL GRID REFINEMENT

The REFINE keyword defines the start of a Cartesian or radial local grid refinement ("LGR") definition that sets the properties of the selected LGR. The keyword is then followed by the property keywords associated with the section where the keyword is being invoked. For example, if the REFINE keyword is used in the GRID section then most of the keywords in that section can be used to set the grid properties for the LGR.

Revision: Rev-I

See REFINE – Start the Definition of a Local Grid Refinement in the GRID section for a full description.

12.3.54 RPTONLY - ACTIVATE THE REPORT TIME STEPS ONLY OPTION FOR THE SUMMARY FILE

This keyword activates the writing out of the SUMMARY file and RSM file data, if the RSM file option has been requested by the RUNSUM keyword in the SUMMARY section, at report time steps only. The default is for all time steps to be written out to the files. This keyword reduces the file size at the expense of lower resolution in the time domain. There is no data required for this keyword.

See RPTONLY - Activate the Report Time Steps Only Option for the SUMMARY File in the SUMMARY section for a full description.

12.3.55 RPTONLYO - DEACTIVATE THE REPORT TIME STEPS ONLY OPTION FOR THE **SUMMARY FILE**

This keyword deactivates the writing out of the SUMMARY file and RSM file data, if the RSM file option has been requested by the RUNSUM keyword in the SUMMARY section, at report time steps only, and switches on writing out all the time steps to the files. This option is the default behavior for when RPTONLY has not been activated. There is no data required for this keyword.

See RPTONLYO - Deactivate the Report Time Steps Only Option for the SUMMARY File in the SUMMARY section for a full description.

12.3.56 RPTRST - Define Data to be Written to the RESTART File

This keyword defines the data to be written to the RESTART file at each requested restart point.

See RPTRST – Define Data to be Written to the RESTART File in the SOLUTION section for a full description.

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12.3.57 RPTSCHED - Define SCHEDULE Section Reporting

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword defines the data in the SCHEDULE section that is to be printed to the output print file in human readable format. The keyword has two distinct forms, the first of which consists of the keyword followed by a series of integers on the next line indicating the data to be printed (see the first example). This is the original formal in the commercial simulator and was subsequently superseded by the second format. The second format consists of the keyword followed by a series of character strings that indicate the data to be printed. In most cases the character string is the keyword used to define the data in the OPM Flow input deck, for example WELSPECS to defined the basic well definitions. Its is anticipated that OPM Flow will eventually support the functionality of the second format only, the first format although recognized will be completely ignored.

For the purpose of controlling PRT output, this keyword is ignored by OPM Flow and has no effect on the simulation. However, the keyword can also be used to control the writing of restart files and those parts of this keyword are honored.

No.	Name	Description	Default
1	FIP	Print the fluid in-place report.	N/A
2	FIPRESV	Print the reservoir volumes in-place report.	N/A
3	RESTART	RESTART defines the frequency at which the restart data for restarting a run is written to the RESTART file. The parameter is assigned a value, OPTION, using the form RESTART = OPTION, where OPTION is an integer variable set to:	
		 OPTION = I then the restart files are written at every report time, but only the last one in the run is kept. This minimizes the restart file size but only the final results are stored, limiting the visualization in OPM ResInsight. 	
		2) OPTION = 2 then the phase inter-blocks are written to the restart files, in addition to the standard data.	
		3) OPTION = 3 then the fluid in-place and phase potentials are also written to the restart file.	
		4) OPTION = 6 then the restart files are written at every time step.	
		See the RPTRST keyword in the SOLUTION section for a more flexible way to write out restart files.	
4			
Notes	<u>s:</u>		

I) The keyword is terminated by "/".

Table 12.21: RPTSCHED Keyword Description

Development is current progressing on developing reports in a similar format to the commercial simulator and this section will be updated as additional reports are added to OPM Flow's functionality.

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Note

Unlike the other reporting keywords in the RUNSPEC, GRID, EDIT, PROPS and SOLUTION keywords, the requested reports on the this keyword in remain in effect until they are switch off by this keyword, that is the reports are written out every report time step until requested to stop.

Examples

The first example shows the original format of this keyword; although the keyword and format are recognized by OPM Flow, the format is ignored and is unlikely to be implemented in in the simulator.

```
-- DEFINE SCHEDULE SECTION REPORT OPTION (ORIGINAL FORMAT)
--
RPTSCHED
1 2*0 1 3*1 /
```

The next example shows the second format of the keyword which may be supported in a future release of OPM Flow.

```
-- SCHEDULE SECTION
SCHEDULE
-- SCHEDULE SECTION - 2000-01-01
RPTSCHED
      'WELLS=2' 'WELSPECS' 'CPU=2' 'FIP=2'
DATES
      1 JAN 2000 /
RPTSCHED
      'NOTHING'
DATES
      1 FEB
             2000 /
      1 MAR
             2000 /
      1 APR
             2000 /
      1 MAY
             2000 /
      1
         JUN
             2000
      1
         JLY
             2000
        AUG
      1
             2000
        SEP
             2000
      1
      1
        0CT
             2000
        NOV
             2000
      1
```

In the above example monthly reporting time steps have been used with a SCHEDULE section report on the January 1, 2000; after which all reports are switch off for the subsequent reporting time steps.

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12.3.58 SAVE - ACTIVATES OUTPUT OF A SAVE FILE FOR FAST RESTARTS

This keyword activates output of a SAVE file for fast restarts. There is no data required for this keyword.

See SAVE – Activates Output of a SAVE File for Fast Restarts in the RUNSPEC section for a full description.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.59 SCHEDULE - Define the Start of the SCHEDULE Section of Keywords

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The SCHEDULE activation keyword marks the end of the SUMMARY section and the start of the SCHEDULE section that defines the group and well definitions, operating and economic constraints, as well as how OPM Flow should advance through time. Numerical controls are also defined in this section and all parameters can be varied through time.

There is no data required for this keyword.

The above example marks the end of the SUMMARY section and the start of the SCHEDULE section in the OPM Flow data input file.

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12.3.60 SHRATE - ACTIVATE AND DEFINE THE POLYMER SHEARING LOGARITHMIC PARAMETERS

This keyword activates the logarithm-based polymer shear thinning/thickening option and defines the shear rate constant. This keyword can only be used in conjunction with the PLYSHLOG in the PROPS section

See SHRATE - Activate and Define the Polymer Shearing Logarithmic Parameters in the PROPS section for a full description.

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12.3.61 SKIPREST - ACTIVATE SKIPPING OF RESTART SCHEDULE DATA

RUNSPEC GRID EDIT PROPS REGIONS SC	SOLUTION SUMMARY	SCHEDULE
------------------------------------	------------------	----------

Revision: Rev-I

Description

This keyword turns on skipping of keywords up to the start of the restart point, as defined on the RESTART keyword in the RUNSPEC section. The RESTART keyword defines the parameters to restart the simulation from a previous run that has written a RESTART file out to disk. Activating the SKIPREST keyword causes the simulator to only read in data it requires for restarting the run up to the RESTART point (RSNUM on the RESTART keyword in the RUNSPEC section). Note that certain keywords always need to be present in a restart run in the SCHEDULE section as the data is not stored on the RESTART file, for example the VFP tables (VFPPROD and VFPINJ keywords). The SKIPREST keyword automatically processes the input deck and reads the required data.

There is no data required for this keyword.

Note that SKIPREST is not necessary for OPM Flow to restart, the simulator will restart gracefully at the chosen step even without it, and the keyword itself is ignored. It is however advisable to include it if compatibility with other simulators is important.

Examples

The example below defines a restart from the previously run NOR-OPM-A01 case at time step number 40.

```
-- SOLUTION SECTION
SOLUTION
      FLEXIBLE RESTART FROM PREVIOUS SIMULATION RUN
                     RESTART
                            RESTART
      FILE
      NAME
                     NUMBER
                            TYPE
                                  FORMAT
RESTART
                                  1*
      'NOR-OPM-A01'
                     40
                            1*
```

Then in the SCHEDULE section the SKIPREST keyword is used to correctly read in the schedule data up to the RESTART point.

```
-- SCHEDULE SECTION
-- SCHEDULE SECTION
-- SCHEDULE
-- ACTIVATE SKIPREST OPTION TO AVOID MODIFYING SCHEDULE SECTION
-- SKIPREST
```

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12.3.62 SUMTHIN - Define SUMMARY DATA Reporting Time Steps

This keyword defines a time interval for writing out the SUMMARY data to the SUMMARY file and the RSM file, if the RUNSUM keyword has been has also been activated in the SUMMARY section. Only the data for the first time step in the time interval is written out and the other time steps are skipped until the next time interval. This enable the size of the SUMMARY files to be reduced depending on the size of the time interval. However, the keyword will produce irregular time steps reports of the SUMMARY data.

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See SUMTHIN - Define SUMMARY DATA Reporting Time Steps in the SUMMARY section for a full description.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.63 TSTEP - Advance Simulation by Reporting Time

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword advances the simulation to a given report time after which additional keywords may be entered to instruct OPM Flow to perform additional functions via the SCHEDULE section keywords, or further TSTEP data sets or keywords may be entered to advance the simulator to the next report time.

No.	Name		Default						
		Field	Metric Labora						
	TSTEP		A vector of real positive numbers that define the length of the time intervals to subsequent report steps						
1		days	days	hours	None				

Notes:

Table 12.22:TSTEP Keyword Description

See also the DATES keyword in the SCHEDULE section. Note that since OPM Flow uses the standard Gregorian calendar, then leap years are accounted for in the DATES keyword. Thus, it is more accurate to use the DATES keyword to progress the simulator through time if one is matching actual production data.

Whenever possible it is a good idea to always set the start date to be at the beginning of the year, as like most simulators, OPM Flow reports are always stated at the number of days from the start date (and sometimes at a given date). If the start date is at the beginning of the year, then calculating the actual date is relatively straight forward and simple.

Examples

The fist example shows how to advance the simulation via the reporting time steps from the given start date of January 1, 2022 set via the START keyword in the RUNSPEC section, to the next year, without any actions or reporting taking place.

```
-- SCHEDULE SECTION
SCHEDULE
-- SCHEDULE SECTION - 2022-01-01
     ADVANCE SIMULATION BY REPORTING TIME
     JAN FEB MAR APR MAY JUN JLY AUG SEP
                                  OCT NOV
                                         DEC
TSTEP
     31
        28
           31
              30
                 31
                      30
                         31 31 30
                                   31
                                      30
                                         31
```

The keyword is followed by a vector of numbers separated by a space and the keyword is terminated by a "/".

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The second example is similar to the previous example but with quarterly reporting time steps used instead

Again, if the simulated production targets are actual production data or the results are going to be used in economic evaluations then the DATES keyword may be more useful in advancing the simulation via the reporting time steps, as the exact dates will be honoured.

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12.3.64 TUNING - Numerical Tuning Control

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Defines the parameters used for controlling the commercial simulator's numerical convergence parameters. The keyword is similar to the TUNINGDP keyword in the SCHEDULE section that is optimized for high throughput runs. The keyword is mostly ignored by OPM Flow; however, the simulator can be instructed to read the first record of the TUNING keyword if the appropriate command line parameter has been activated (see section 2.2 Running Flow).

No.	Name		Description		Default
		Field	Metric	Laboratory	
1-1	TSINIT	TSINT is a real positive v time step.	alue that defines the maxi	mum length of the next	
		Note that whenever the default value of one, unless	keyword is used TSINIT is ss explicitly over written.	s always set back to the	
		days	days	hours	1.0
1-2	TSMAXZ	TSMAXZ is a real position	ve value that defines the TSINIT.	maximum length of the	
		days	days	hours	365.0
1-3	TSMINZ	TSMINZ is a real positive time steps.	ve value that defines the	minimum length of all	
		days	days	hours	0.1
1-4	TSMCHP	TSMCHP is a real posit chopped time steps.	tive values that sets the	minimum length of all	
		days	days	hours	0.15
I-5	TSFMAX		e value that specifies the r sed by, subject to the m		
		TSFMAX is set to the de	rent time step has conv efault value, then the next ovided it is less than TSM.	time step will be 3.0 x	
		dimensionless	dimensionless	dimensionless	3.0
I-6	TSFMIN		e value that specifies the ased by, subject to the m		
		TSFMAX is set to the de	ent time step has not co efault value, then the next num of 0.3 days and TSMIN	time step will be 0.3 x	
		dimensionless	dimensionless	dimensionless	0.3
1-7	TSFCNV		lue that specifies the dec number of target iteratio		
		dimensionless	dimensionless	dimensionless	0.1

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No.	Name	Description					
		Field	Metric	Laboratory			
1-8	TFDIFF	TFDIFFA is a real positi	ve value that sets the tim	e step growth factor of			
		TFDIFF is set to the def	For example, if the chopped current convergent time step is 10 days and TFDIFF is set to the default value, then the time step will be increased to 1.25×10 days, that is the minimum of 11.25 days and TSMAXZ.				
		dimensionless	dimensionless	dimensionless	1.25		
1-9	THRURPT	THRURPT is a real posi ratio over a time step.	tive value that specifies th	e maximum throughput			
		dimensionless	dimensionless	dimensionless	1.0×10^{20}		
1-10	TMAXWC		ble precision value that deent; for example, when a v				
		days	days	hours	None		
1-11	1	Record terminated by a	up,		Not Applicable		
2-1	TRGTTE	TRGTTE is a real positiv	e value that sets the time	truncation error target.			
		dimensionless	dimensionless	dimensionless	0.1		
2-2	TRGCNV	TRGCNV a real positive error.	TRGCNV a real positive value that defines the non-linear convergence error.				
		dimensionless	dimensionless	dimensionless	0.001		
2-3	TRGMBE	TRGMBE is a real positive error.	TRGMBE is a real positive value that specifies then target material balance error.				
		dimensionless	dimensionless	dimensionless	1.0 x 10 ⁻⁷		
2-4	TRGLCV	TRGLCV is a real positive error target.	re value that specifies the I	inear convergence			
		dimensionless	dimensionless	dimensionless	0.00001		
2-5	XXXTTE	XXXTTE is a real positive error.	ve value that sets the maxi	mum time truncation			
		dimensionless	dimensionless	dimensionless	10.0		
2-6	XXXCNV	XXXCNV is a real post convergence error.	itive value that defines th	ne maximum non-linear			
		dimensionless	dimensionless	dimensionless	0.01		
2-7	XXXMBE	XXXMBE is a real positive value that specifies the maximum mass balance error, that is the tolerated mass balance error relative to total mass present.					
		dimensionless	dimensionless	dimensionless	1.0 × 10 ⁻⁶		
2-8	XXXLCV	XXXLCV is a real p convergence error.	ositive values that sets	the maximum linear			
		dimensionless	dimensionless	dimensionless	0.001		

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No.	Name	Name Description						
		Field	Metric	Laboratory				
2-9	XXXWFL	XXXWFL is a real post convergence error.	sitive values that fixes th	ne maximum well flow				
		dimensionless	dimensionless	dimensionless	0.001			
2-10	TRGFIP		TRGFIP is a real positive value that stipulates the target fluid in-place error in Local Grid Refinements.					
		dimensionless	dimensionless	dimensionless	0.025			
2-11	TRGSFT	TRGSFT is a real positive when the Surfactant Mod	e values that defines the t lel is active in the run.	target surfactant change				
		dimensionless	dimensionless	dimensionless	None			
2-12	THIONX		al value used to set the to on for when the Brine Mo					
		dimensionless	dimensionless	dimensionless	0.01			
2-13	TRWGHT	tracer updates within the						
		dimensionless	dimensionless	dimensionless	I			
2-14	1	Record terminated by a '	Record terminated by a "/"					
3-1	NEWTMX		e integer greater or equ number of Newtonian iter					
		dimensionless	dimensionless	dimensionless	12			
3-2	NEWTMN		nteger that is less or equa nber of Newtonian iteration					
		dimensionless	dimensionless	dimensionless	I			
3-3	LITMAX		eger greater or equal to LI ar iterations within a New					
		dimensionless	dimensionless	dimensionless	25			
3-4	LITMIN		ger less or equal to LIMM ar iterations within a New					
		dimensionless	dimensionless	dimensionless	l I			
3-5	MXWSIT		MXWSIT is a positive integer that defines the maximum number of iterations within a well flow calculation.					
		dimensionless	dimensionless	dimensionless	8			
3-6	MXWPIT	iterations for solving the	leger that stipulates the m bottom-hole pressure for thin a well flow calculation	wells under tubing				
		dimensionless	dimensionless	dimensionless	8			

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No.	Name		Description		Default
		Field	Metric	Laboratory	
3-7	DDPLIM	naximum pressure			
		psia	barsa	atma	1.0 x 10 ⁻⁶
3-8	DDSLIM	DDSLIM a real positive va		um saturation change at	
		dimensionless	dimensionless	dimensionless	1.0 x 10 ⁻⁶
3-9	TRGDPR	TRGDP is a real positive within a time step.	value that defines the tar	get pressure change	
		psia	barsa	atma	1.0 x 10 ⁻⁶
3-10	XXXDPR	XXXDPR is a real positive pressure change within a		e maximum tolerable	
		psia	barsa	atma	1.0 × 10 ⁻⁶
3-11	MNWRFP	MNWRFP is a positive in that defines the minimum the bisection algorithm for via the POLYMER keywork.			
		dimensionless	dimensionless	dimensionless	4
3-12	1	Record terminated by a "	,,,	ı	Not Applicable

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Notes:

- 1) The keyword consists of three records, with items 1-1 to 1-11 representing record one items, 2-1 to 2-14 record two and 3-1 to 3-12 representing record number three. Each of the records are terminated by a "/" and is explicitly shown in the above rows.
- 2) There is no keyword terminating "/".

Table 12.23:TUNING Keyword Description

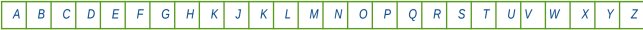
Note that for record number two (items 2-1 to 2-13) the maximum values should always be greater than the associated target value; for example, XXXCNV should be greater than TRGCNV. Also note that the TUNING keyword is stored on the restart files (see *RPTRST – Define Data to be Written to the RESTART File*) enabling the parameters to be utilized in a restart run without re-specifying the keyword.

Example

```
-- DEFAULT TUNING PARAMETERS
-- TUNING
1.0 365.0 0.1 0.15 3 0.3 0.1 1.25 1E20 1* /
```

The above example explicitly sets the default parameters for OPM Flow for when the appropriate command line parameter has been activated (see section 2.2 Running Flow) to instruct the simulator to read the first record of the TUNING keyword. Alternatively one could just use:

```
TUNING
/
/
```



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12.3.65 TUNINGDP - NUMERICAL TUNING CONTROL FOR HIGH THROUGHPUT CASES

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Defines the parameters used for controlling the commercial simulator's numerical convergence parameters. The keyword is similar to the TUNING keyword in the SCHEDULE section, but the defaults on this keyword are optimized for high throughput runs.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.66 UDQ - Declare User Define Quantities ("UDQ")

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword starts the definition of a UDQ section that stipulates the variables and operations used to access the User Defined Quantities features in OPM Flow. UDQ variables can be constants, SUMMARY variables, as defined in the SUMMARY section, or a formula using various mathematical functions together with constants and SUMMARY variables. Available operation commands include the ASSIGN, DEFINE UNITS and UPDATE that are sub-keywords to the UDQ section keyword. An UDQ definition section is terminated by a "/" on a single line.

Although this keyword is read by OPM Flow, the ACTION and UDQ computational logic and calculations have not been fully implemented and therefore this keyword should not be used as it may result in OPM Flow aborting.

No.	Name	Description	Default
UDQ		Define the start of UDQ Definition Section. This is then followed on a new line by any number of UDQ records that define the various operations to be performed using the ASSIGN, DEFINE UNITS and UPDATE sub-keywords for the OPERATOR.	
1	OPERATOR	OPERATOR is a character sting that that defines the type of operations to perform, and should be one of the following:	
		 ASSIGN: This option assigns a value to a variable and sets the UPDATE status to OFF. 	
		 DEFINE: In this case a mathematical formula is defined and assigned to a variable. The variable is initialized with the formula and the UPDATE status is set to ON. 	
		3) UNITS: This option sets the reporting units for a defined variable and has no effect on the calculations. The variable must already have been defined prior to using this option.	
		 UPDATE: Stipulates when the defined variable should be re- calculated. 	

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No.	Name	Description	Default
2	VARIABLE	VARIABLE is a character string of length eight that stipulates the name of the user defined variable that will processed by the OPERATOR command. The first two characters of VARIABLE must be set based on the type of of variable being defined, that is:	
		 CU: For variables that are associated with connections, for example SUMMARY variable COFR (Connection Oil Flow Rate). 	
		 FU: For variables that are associated with field data, for example SUMMARY variable FOPR (Field Oil Production Rate). 	
		3) GU: For variables that are associated with groups, for example SUMMARY variable GLPR (Group Liquid Production Rate).	
		4) RU: For variables that are associated with regions, for example SUMMARY variable RPR (Region Pressure).	
		 SU: For variables that are associated with multi-segment wells, for example SUMMARY variable SOFR (Segment Oil Flow Rate). 	
		 WU: For variables that are associated with wells, for example SUMMARY variable WWCT (Well Water Cut). 	
		 AU: For variables that are associated with aquifers, for example SUMMARY variable AAQP (Analytical Aquifer Pressure). 	
		8) BU: For variables that are associated with blocks, for example SUMMARY variable BPR (Block oil phase Pressure).	
3	EXPRESSION	The data type for EXPRESSION is based on the OPERATOR option above, namely if OPERATOR is set to:	
		ASSIGN: Then EXPRESSION should be a numerical value.	
		 DEFINE: In this case a mathematical formula is defined and assigned to a variable. The variable is initialized with the formula and the UPDATE status is set to ON. 	
		 UNITS: Then EXPRESSION should be a character string enclosed in quotes it it contains blanks, with a maximum length of eight characters, that declares the units for VARIABLE that will be used for reporting. 	
		4) UPDATE: In this case EXPRESSION can have the value ON to evaluate VARIABLE at all time steps, OFF to not evaluate VARIABLE, or NEXT to evaluate VARIABLE at the next time step.	
	1	Termination of a UDQ record. Note that multiple numbers of records can be entered within a UDQ section with each record terminated by a "/".	
/		Define the end of UDQ Definition Section	
Notes	<u>. </u>		
I)	The keyword is t	terminated by a "/".	

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Table 12.24: UDQ Keyword Description

Currently, the simulator only supports well variable names (WU type) variables and simple mathematical formula consisting of opening and closing brackets (()), and the plus, minus, multiply and divide operators, as illustrated in the examples below.

See also the UDADIMS, UDQDIMS and UDQPARAM keywords in the RUNSPEC section to define the dimensions for the UDQ keyword and associated variables.

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Examples

The fist example shows how to define some constant field variables used for calculating facilities corrected condensate and Liquefied Petroleum Gas⁶³ ("LPG") yields in a wet gas model:

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```
-- DEFINE START OF USER DEFINED QUANTITY SECTION
UDQ
-- OPERATOR VARIABLE EXPRESSION
ASSIGN
            FUNGLYLD 1.100000
                                       / Condensate Yield (stb/Mscf)
ASSIGN
            FUNGLSHK
                     0.000000
                                       / Condensate Shrinkage Factor set to Zero
                                       / LPG Sep Gas Yield (stb/Mscf)
ASSIGN
            FULPGYLD
                     0.065775
                                                     Shrinkage Factor
            FUI PGSHK
                     0.080410
                                       / IPG
ASSTGN
ASSIGN
            FUFACSHK
                     0.000935
                                       / Facilities
                                                     Shrinkage Factor
ASSIGN
            FUFULSHK
                     0.052924
                                       / Fuel
                                                     Utilization
ASSTGN
            FUDFI TA
                      1F-10
                                       / Value to avoid diving by zero errors
  DEFINE END OF USER DEFINED QUANTITY SECTION
```

The next example is a continuation of this example by showing how one can calculate the adjusted field condensate and LPG rates. Note both examples could be merged into a single UDQ definition but have been stated separately for ease of reference.

```
-- DEFINE START OF USER DEFINED QUANTITY SECTION
UDQ
-- OPERATOR VARIABLE EXPRESSION
            FU FNGLR
                     FGPR *(FOGR * FUNGLYLD) / Calculate Condensate Rate Field
DEFINE
UPDATE
            FU FNGLR
                     ON
UNITS
            FU_FNGLR
                      STBD
DEFINE
            FU FLPGR
                      FU FWGPR * FULPGYLD
                                                 Calculate LPG Rate Field
UPDATE
            FU_FLPGR
                      ON
            FU_FLPGR
UNITS
                      STBD
  DEFINE END OF USER DEFINED QUANTITY SECTION
```

In the above the DEFINE operator is use to define the equations to calculate the corrected condensate (FU_FNGLR) and LPG rates (FU_FLPGR) with the UPDATE operator set to ON so that the rates are calculate at every time step, and finally, the UNITS operator is used to set the units of of the calculated rates.

The final example show the use of the UDADIMS and UDQDIMS keywords in the RUNSPEC section, followed by the keywords in the SCHEDULE section that define a UDQ definition that users the DEFINE operator to calculate adjusted well rates based on an expression. The final set of keywords show how the UDQ defined variables are employed on the WCONPROD keyword to control the production constraints for several wells.

⁶³ Liquefied Petroleum Gas or LPG consists mainly of propane, propylene, butane, and butylene in various mixtures. It is produced as a by-product of natural gas processing and petroleum refining. The components of LPG are gases at standard conditions.

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```
RUNSPEC SECTION KEYWORDS
         USER DEFINED ARGUMENT DIMENSIONS
- -
         NO.
                 NOT
                         TOTAL
                         UDQ
_ _
         ARGS
                 USED
UDADIMS
                  1*
         10
                         10
_ _
         USER DEFINED ARGUMENT DIMENSIONS FACILITY
         MAX
                 MAX
                         MAX
                                MAX
                                     MAX
                                               MAX
                                                     MAX
                                                           MAX
                                                                  MAX MAX
                                                                              RAND
                         CONNS FIELD GROUP REGS SEGTM WELL
- -
         FUNCS
                 ITEMS
                                                                  AQUF BLCKS
                                                                              0PT
UDODIMS
         50
                 25
                                 50
                                        50
                                                           0
                                                                  0
                                                                              N /
                             SCHEDULE SECTION KEYWORDS
-- DEFINE START OF USER DEFINED QUANTITY SECTION
UDQ
-- OPERATOR VARIABLE EXPRESSION
            WUOPRL (WOPR OPLO1 - 150) * 0.90 / OIL & LIQ CAPACITIES
DEFINE
            WULPRL (WLPR OPLO1 - 200) * 0.90 / at GEFAC = 0.8995
DEETNE
            WUOPRU (WOPR OPU01 - 250) * 0.80 /
DEFINE
            WULPRU (WLPR 0PU01 - 300) * 0.80 /
DEFINE
            WUOPRL SM3/DAY
UNITS
                                              / DEFINE REPORTING UNITS
UNITS
            WULPRL SM3/DAY
                                              / FOR UDQ VARIABLES
            WUOPRU SM3/DAY
UNITS
            WULPRU SM3/DAY
/ DEFINE END OF USER DEFINED QUANTITY SECTION
- -
         WELL PRODUCTION WELL CONTROLS
- -
-- WELL
        OPEN/
               CNTL
                       OIL
                              WAT
                                      GAS
                                            LIO
                                                   RES
                                                          BHP
                                                                 THP
                                                                       VFP
                                                                              VFP
-- NAME
         SHUT
                MODE
                       RATE
                              RATE
                                      RATE
                                            RATE
                                                   RATE
                                                          PRES PRES TABLE ALFO
WCONPROD
                       1*
                                      1*
0P01
         SHUT
                GRUP
                              1*
                                            1*
                                                   1*
                                                          200.0
                       1*
                              1*
                                      1*
                                            1*
                                                   1*
0P02
         SHUT
                GRUP
                                                          200.0
DATES
         1 FEB
                  2020 /
         WELL PRODUCTION WELL CONTROLS
-- WELL
        OPEN/
                CNTL
                       OIL
                              WAT
                                      GAS
                                            LIO
                                                   RES
                                                          BHP
                                                                 THP
                                                                       VFP
-- NAME
         SHUT
                MODE
                       RATE
                              RATE
                                      RATE RATE
                                                   RATE
                                                          PRES
                                                                PRES TABLE ALFQ
WCONPROD
                GRUP
                       WUOPRL 1*
                                      1*
                                            WULPRL 1*
0P01
         OPEN
                                                          60.0
                       WUOPRL 1*
                                      1*
                                            WULPRL 1*
0P02
         OPEN
                GRUP
                                                          00.0
DATES
         1 MAR
                  2020
         1
            APR
                  2020
         1
            MAY
                  2020
         1
            JUN
                  2020
            JLY
                  2020
         1
            AUG
                  2020
            SEP
                  2020
         1
```

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12.3.67 VAPPARS - OIL VAPORIZATION PARAMETERS

VAPPARS defines the rate of oil vaporization in the presence of undersaturated gas and the rate at which the remaining oil gets "heavier" via the reduction in the solution gas-oil ratio ("Rs"). This keyword should only be used if the OIL, GAS, DISGAS and VAPOIL keywords in the RUNSPEC section have been invoked to allow oil, gas, dissolved gas and vaporized oil to be present in the model.

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In OPM Flow, VAPPARS can only be set once, and not changed multiple times in the SCHEDULE section.

See VAPPARS - Oil Vaporization Parameters in the SOLUTION section for a full description.

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12.3.68 VFPINJ - Define Injection Vertical Flow Performance Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The VFPINJ keyword defines injection Vertical Flow Performance ("VFP") tables that are used to determine the outflow or downstream pressure based on the inlet or upstream pressure and the phases being injected into the system. For a well this means the table relates the flowing bottom-hole pressure ("BHP") to the well's tubing head pressure ("THP") based on the oil, gas or water injection rates. The table is also used to describe the pressure relationship when the network option is being used, although the Network option is not currently implemented in OPM Flow. In this case the table describes the pipeline pressure behavior from the HIGHER group (inlet node) to the LOWER group (outlet node) given the current flowing conditions (the group relationship is defined by the GRUPTREE keyword in SCHEDULE section).

The data for this keyword is generated by an external program and is normally included into the input deck using the INCLUDE keyword as described in section 4 GLOBAL SECTION KEYWORDS, as the data can be quite voluminous.

Each VFPINJ table must be entered with a separate VFPINJ keyword that consists of four records, with I-I, I-2 representing record one items (I) and (2) and 2-I, 2-2 representing record number two items (I) and (2) and so on in the "No." column in Table 12.25. Each record is terminated by a "/". The fourth record must be repeated to give BHP data as a function of FLO for all THP values.

No.	Name		Description		Default
		Field	Metric	Laboratory	
1-1	VFPTAB	A positive integer greater than zero and less than or equal to the MXVFPTAB variable as defined on the VFPIDIMS keyword in the RUNSPEC section, that defines the vertical flow performance table number.			None
1-2	VFPREF	A real positive value that defines the reference depth used to generate this VFPINJ table data set. OPM Flow automatically corrects any difference between VFPREF and the BHPREF on the WELSPECS and WPAVDEP keywords in the SCHEDULE section, using the current hydrostatic head.			None
1-3	FLO	set to one of the followin 1) OIL: for injecting 2) GAS: for injecting	A defined character string that defines the injection phases, and should be set to one of the following character strings: 1) OIL: for injecting phase being oil. 2) GAS: for injecting phase being gas. 3) WAT: for injecting phase being water.		
1-4	VFPTYPE	A defined character string	g that should be defaulted	or set equal to THP.	THP
1-5	VFPUNITS	Units used for the BHP-DATA on this keyword. This variable is ignored by OPM Flow and should be defaulted with I*. FIELD METRIC LAB		*	
1-6	VFPVALUE				1.
1-6	VFFVALUE		g that should be defaulted y OPM Flow and should b	•	ВНР
1-7	1	Record terminated by a "	·","		Not Applicable

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No.	Name		Description		Default		
		Field	Metric	Laboratory			
2-1	FLO-DATA		ically increasing vector that ase declared by the FLO v				
			nust greater than two and the VFPIDIMS keyword in				
		Liquid: stb	Liquid: sm³	Liquid: scc			
		Gas: Mscf	Gas: sm³	Gas: scc	None		
2-2	1	Record terminated by a "	Record terminated by a "/"				
3-I THP-DATA		A real positive monotoni values of the tubing head	cally increasing vector that pressure values.	at defines the numerical			
		The number of entries m MXMTHP as defined on t					
		psia	barsa	atma	None		
3-2	1	Record terminated by a "	·//"		Not Applicable		
4-1	NTHP	values entered via the THTHP-DATA is equal to 1	of an integer value that d IP-DATA records on this k 000, 2000, 3000 and 3500 to third entry, that is THP	keyword. For example, if and NTHP is equal to			
	BHP-DATA		NTHP is then followed by a real vector of BHP values for each FLO injection rate for the corresponding index value (NTHP) and is then terminated with a"/"				
			consists of both NTHP combinations of (NTHeen entered.				
		psia	barsa	atma	None		
4-2	1	Each Index (NTHP, BHP-I	DATA) data set is terminat	ted by a "/"	Not Applicable		

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Notes:

- Each VFPINJ table must be entered with a separate VFINJ keyword that consists of four records, with items I-I to I-7 representing record one items and 2-I to 2-2 representing record number two items, etc., in the "No." column in this table.
- 2) Each of the records are terminated by a "/" and is explicitly shown in the above rows.
- 3) There is no keyword terminating "/".

Table 12.25:VFPINI Keyword Description

See also the WELSPECS keyword to define wells and the WCONINJE keyword that is used to allocate the VFPINJ tables to specific wells. Note that one VFPINJ table can be allocated to one or more wells, provided the wells in question have a similar trajectory and similar flow characteristics, for example vertical water injection wells injecting into the same reservoir.

The VFPROD keyword is used to enter VFP tables for production wells or to describe the pressure relationship when the network option is being used. In this case the table describes the pipeline pressure behavior from the LOWER group (inlet node) to the HIGHER group (outlet node) given the current production conditions.

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All the aforementioned keywords are described in the SCHEDULE section.

Examples

The following example shows the VFPINJ table for a water injection well and is taken from the Norne OPM Flow model.

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```
VFPINJ
```

```
-- Table Datum Depth Rate Type
-- ----
           _____
                           'WAT' /
              2718.07
      12
-- 'WAT' units - SM3/DAY
  500.0 1263.2 2026.3 2789.5 3552.6
 4315.8 5078.9 5842.1 6605.3 7368.4
8131.6 8894.7 9657.9 10421.1 11184.2
11947.4 12710.5 13473.7 14236.8 15000.0 /
-- 'THP' units - BARSA
         63.24 105.46 147.68 189.90
  21.01
 232.12 274.35 316.57 358.79 401.01 /
1 254.51 253.95 252.27 249.83 246.69
   242.88 238.42 233.32 227.59 221.22
   214.23 206.62 198.38 189.53 180.06
   169.97 159.26 147.95 136.00
                                    123.46
  297.02 296.49 294.82 292.39
                                    289.26
   285.47 281.01 275.92 270.20
                                    263.84
   256.87 249.28 241.05 232.22
                                    222.76
   212.70 202.01 190.71 178.79
                                   166.27
 594.67 594.29 592.70 590.34 587.29
   583.57 579.16 574.17 568.55
                                   562.25
   555.40 547.92 539.79 531.09
511.82 501.25 490.13 478.34
                                    521.74
                                   466.01
10 637.19 636.83 635.26 632.91
                                    629.86
   626.16 621.76 616.78 611.17
                                    604.89
   598.05 590.59 582.47 573.79 564.45
554.56 544.01 532.91 521.14 508.83
```

The example shows the first two and the last two records of the fourth kind, as the data is too voluminous to be included.

Note

The VFPTAB variable defines the table number of the VFPINJ data set; if more then one VFPINJ keyword is entered with the same VFPTAB number then the VFPINJ data set will be overwritten by the last VFPINJ keyword with the same VFPTAB number.

The same comment is also applicable to the VFPPROD keyword.

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12.3.69 VFPPROD - Define Production Vertical Flow Performance Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The VFPROD keyword defines production Vertical Flow Performance ("VFP") tables that are used to determine the outflow or downstream pressure based on the inlet or upstream pressure and the phases flowing through the system. For a well this means the table relates the flowing bottom-hole pressure ("BHP") to the well's tubing head pressure ("THP") based on the oil, gas and water rates (and any artificial lift quantities like gas lift gas), or phases ratios, flowing up the wellbore. The table is also used to describe the pressure relationship when the network option is being used, although the Network option is not currently implemented in OPM Flow. In this case the table describes the pipeline pressure behavior from the LOWER group (inlet node) to the HIGHER group (outlet node) given the current flowing conditions (the group relationship is defined by the GRUPTREE keyword in SCHEDULE section).

Each VFPPROD table must be entered with a separate VFPPROD keyword that consists of seven records, with I-I, I-2 representing record one items (I) and (2) and 2-I, 2-2 representing record number two items (I) and (2) in the "No." column in Table 12.26. Each record is terminated by a "/". The seventh record must be repeated to give BHP data as a function of FLO for all THP values.

No.	Name	Description				
		Field	Metric	Laboratory		
1-1	VFPTAB	MXVFPTAB variable as	A positive integer greater than zero and less than or equal to the MXVFPTAB variable as defined on the VFPPDIMS keyword in the RUNSPEC section, that defines the vertical lift performance table number.			
1-2	VFPREF	A real positive value that defines the reference depth used to generate this VFPPROD table data set. OPM Flow automatically corrects any difference between VFPREF and the BHPREF on the WELSPECS and WPAVDEP keywords in the SCHEDULE section, using the current hydrostatic head.			None	
1-3	FLO	set to one of the following p 1) GAS: for flowing p 2) OIL: for flowing p	A defined character string that defines the flowing phases, and should be set to one of the following character strings: 1) GAS: for flowing phase being the gas rate. 2) OIL: for flowing phase being the oil rate. 3) LIQ: for flowing phase being the liquid (oil plus water) rate.			
1-4	WFR	should be set to one of the should be used if 2) WCT: for the was should be used if 3) WGR: for the was	ing that defines the flow he following character stricter fraction being the war FLOW is set to OIL or LI ter fraction being the war FLOW is set to OIL or LI ter fraction being the war FLOW is set to GAS.	ings: $\frac{q_w}{q_o} \text{and} Q'$ $\text{ter cut} \frac{q_w}{q_o + q_w} \text{and} Q$	None	

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No.	Name		Description		Default
		Field	Metric	Laboratory	
1-5	GFR	A defined character string be set to one of the follo	g that defines the flowing wing character strings:	gas fraction and should	None
		I) GOR: for the gas	fraction being the gas-oil	ratio $\frac{q_w}{q_o}$ and should	
		be used if FLOW	is set to OIL or LIQ'	10	
		2) GLR: for the gas f	fraction being the gas-liqui	d ratio $\frac{q_g}{q_o + q_w}$ and	
		should be used if	FLOW is set to OIL or LI	Q	
		3) OGR: for the gas	fraction being the oil-gas	ratio $\frac{q_o}{q_g}$ and should	
		be used if FLOW	is set to GAS.		
I-6	VFPTYPE	A defined character strin	g that should be defaulted	or set equal to THP.	THP
1-7	ALQ	A defined character strin be set to one of the follo	g that defines the artificial wing character strings:	lift quantity and should	 *
		I) GRAT: for the ar rate.	tificial lift quantity being	the gas lift gas injection	
		IGLR: for the art gas-liquid ratio.	ificial lift quantity being t	he gas lift gas, injection	
		3) TGLR: for the ar total gas-liquid ra	tificial lift quantity being t tio.	the gas lift gas, injection	
		4) COMP: for the a for a compressor.	rtificial lift quantity being	the compressor power,	
		5) PUMP: for the a pump.	artificial lift quantity being	the pump rating for a	
		6) DENO: for oil sur	face density.		
		7) DENG: for gas su	•		
		The default value of I* is ALQ variable is not enter	"" or undefined that cored.	vers the case when the	
1-8	VFPUNITS	Units used for the BHP-D	OATA on this keyword.		
		This variable is ignored b	y OPM Flow and should b	e defaulted with 1*.	
		FIELD	METRIC	LAB	 *
1-9	VFPVALUE	A defined character strir	ng that should be defaulted	or set equal to BHP.	
		This variable is ignored b	y OPM Flow and should b	e defaulted with 1*.	BHP
1-10	1	Record terminated by a '	<i>(</i>)"		Not Applicable
2-1	FLO-DATA		ically increasing vector that se declared by the FLOW		
			nust greater than two an the VFPPDIMS keyword in		
		Liquid: stb	Liquid: sm³	Liquid: scc	
		Gas: Mscf	Gas: sm ³	Gas: scc	None

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No.	Name		Description		Default	
		Field	Metric	Laboratory		
2-2	1	Record terminated by a "	<i>J</i> ,,		Not Applicable	
3-1	THP-DATA	A real positive monotoni values of the tubing head	cally increasing vector that pressure values.	at defines the numerical		
			nust greater than two an the VFPPDIMS keyword in			
		psia	barsa	atma	None	
3-2	1	Record terminated by a "	/ ''		Not Applicable	
4-1	WFR-DATA		cally increasing vector tha er fraction declared by the			
			nust greater than two an on the VFPPDIMS keyw			
		WOR: dimensionless	dimensionless	dimensionless		
		WCT: dimensionless	dimensionless	dimensionless		
		WGR: stb/Mscf	dimensionless	dimensionless	None	
4-2	1	Record terminated by a "	<i>)</i> "		Not Applicable	
5-1	GFR-DATA	A real positive monotoni values of the flowing gas f				
		The number of entries must greater than two and less than or equal to MXMGFR as defined on the VFPPDIMS keyword in the RUNSPEC section.				
		GOR: Mscf/stb	dimensionless	dimensionless		
		GLR: Mscf/stb	dimensionless	dimensionless		
		OGR: stb/Mscf	dimensionless	dimensionless	None	
5-2	1	Record terminated by a "	/ ,,		Not Applicable	
6-1	ALQ-DATA		cally increasing vector tha quantity declared by the A			
			nust greater than two and the VFPPDIMS keywood			
		GRAT: Mscf/day	sm³/day	scc/hour		
		IGLR: Mscf/stb	dimensionless	dimensionless		
		TGLR: Mscf/day	dimensionless	dimensionless		
		DENO: lb/ft ³	kg/m³	gm/cc		
		DENG: lb/ft	kg/m³	gm/cc	None	
6-2	1	Record terminated by a "	<i>)</i> "		Not Applicable	

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No.	Name		Description		Default		
		Field	Metric	Laboratory			
7-1	NTHP	index of THP, WFR, GFI keyword. The first index, NTHP, is values entered via the TH THP-DATA is equal to 10	s of a series of integer R, ALQ entered via the an integer value that de IP-DATA records on this RO, 200, 300 and 350 and d entry, that is THP equal	fines the index of THP ceyword. For example, if NTHP is equal to three	None		
	NWFR	water fraction values e keyword. For example, if	R, is an integer value that centered via the WFR-D WFR-DATA is equal to 0 two then NWFR refers t	ATA records on this .00, 0.25, 0.50 and 0.75	None		
	NGFR	fraction values entered vexample, if GFR-DATA is	The third index, NGFR, is an integer value that defines the index of the gas fraction values entered via the GFR-DATA records on this keyword. For example, if GFR-DATA is equal to 100.0, 200.0, 500.0 and 750.0 and NGFR is equal to three then NGFR refers to third entry, that is GFR equal to 500.0.				
	NALQ	The fourth and final inde the index of artificial li keyword. For example, if NALQ is equal to one th to 50.	None				
	BHP-DATA	The fourth index is then followed by the BHP values. BHP-DATA is a real vector of BHP values for each FLO production rate for the corresponding index value (NTHP, NWFR, NGFR, NALQ) and is then terminated with a"/"					
			consists of the four indice ations of (NTHP, NWFR, een entered.				
		psia	barsa	atma	None		
7-2	1	Each Index (NTHP, NWF terminated by a "/"	R, NGFR, NALQ. BHP-DA	TA) data set is	Not Applicable		

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Notes:

- Each VFPPROD table must be entered with a separate VFPPROD keyword that consists of seven records, with entries 1-1 to 1-10 representing record one items and 2-1 to 2-2 representing record number two items, etc., in the "No." column in this table.
- 2) Each of the records are terminated by a "/" and is explicitly shown in the above rows.
- 3) There is no keyword terminating "/".

Table 12.26:VFPPROD Keyword Description

The data for this keyword is generated by an external program and is normally included into the input deck using the INCLUDE keyword as described in section 4 GLOBAL SECTION KEYWORDS, as the data can be quite voluminous.

See also the WELSPECS keyword to define wells and the WCONPROD keyword that is used to allocate the VFPPROD tables to specific wells. Note that one VFPPROD table can be allocated to one or more wells, provided the wells in question have a similar trajectory and similar flow characteristics, for example vertical oil wells producing from the same reservoir, or different reservoirs with similar PVT properties.

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he the pressure relationship

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The VFPINJ keyword is used to enter VFP tables for injection wells or to describe the pressure relationship when the network option is being used. In this case the table describes the pipeline pressure behavior from the HIGHER group (inlet node) to the LOWER group (outlet node) given the current injection conditions.

All the aforementioned keywords are described in the SCHEDULE section.

Examples

The following example shows the VFPROD table for a production gas well and is taken from the Norne OPM Flow model. Here WFR has been set to water-gas ratio and GFR has been set to the oil-gas ratio, and the ALQ value is defaulted.

```
VFPPROD
```

```
-- Table
          Datum Depth
                      Rate Type
                                 WFR Type
                                            GFR Type
                      -----
                                 ------
                                  'WGR'
                       'GAS'
                                            'OGR' /
           2623.39
-- 'GAS' units - SM3/DAY
 50000.0 100000.0 200000.0 400000.0 800000.0
1200000.0 1600000.0 1999999.9 3000000.0 3999999.8
5000000.5 /
-- 'THP' units - BARSA
 10.00
        20.00 40.00
                       80.00 120.00
150.00 200.00 250.00 /
-- 'WGR' units - SM3/SM3
       1e-9 1e-6
    0
                        1e-5 0.0001
 0.001
         0.01
                 0.1 /
-- 'OGR' units - SM3/SM3
  1e-7
        1e-6 1e-5 0.0001 0.001
  0.01 /
-- 'ALQ' units -
     0 /
           11.93
   1 1 1
                   12.22 13.35
                                  17.24
                                        27.93
            39.83
                    52.06 64.38
                                  95.20 125.89
           156.52
  1 2 1
            11.93
                    12.22 13.35
                                  17.24
                                          27.94
            39.84
                    52.07 64.39
                                  95.21 125.91
           156.55
   8 5 1 483.75 511.15 614.09 1044.78 2757.56
           5592.55 9528.36 14567.24 32005.79 56375.24
            87684
   8 6 1 487.68 516.24 624.74 1075.40 2860.16
           5803.92 9880.58 15093.76 33119.59 58297.57
```

The example shows the first two and the last two records of type seven, as the data is too voluminous to be included.

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The next example below shows an example oil producing well VFPPROD, again taken from Norne OPM Flow model. Here WFR has been set to water cut and GFR has been set to the gas-oil ratio, and the ALQ value is defaulted.

```
VFPPROD
                        Rate Type
                                    WFR Type
          Datum Depth
-- Table
                                               GFR Type
                                                         TAB Type
                         'LIQ'
                                      'WCT'
     37
              2641.02
                                                  'GOR'
-- Prosper files are corrected from RKB to MSL depth. lmarr
                                               GFR Type
-- Table
          Datum Depth
                        Rate Type
                                    WFR Type
                                                         TAB Type
                         'LIQ'
                                      'WCT'
                                               'GOR'
     37
              2617.02
-- 'LIQ' units - SM3/DAY
         500.0 1000.0 1500.0 2000.0
 200.0
        3000.0 3500.0 4000.0 4500.0
5500.0 6000.0 6500.0 7000.0
 2500.0
 5000.0
 7500.0 8000.0 10000.0 14000.0 /
-- 'THP' units - BARSA
        51.01 61.01
 21.01
                        81.01 101.01
        141.01 161.01 181.01
 121.01
                                201.01 /
-- 'WCT' units - FRACTION
                   0.2
         0.1
                           0.3
                                   0.4
   0.5
                   0.7
           0.6
                           0.8
                                    1 /
-- 'GOR' units - SM3/SM3
          100
                           200
                                   500
    90
                   150
   1000
          2000 /
-- 'ALO' units -
     0 /
 1 1 1 1 160.82 136.70 119.79 115.86 117.38
            121.16 126.08 131.56 137.48 143.74
            150.29 157.07 164.02 171.07
                                           178.13
            185.11 192.09
                            220.38 280.86
   1 2 1
            155.63 129.40
                            112.32 108.64
                                            110.44
                    120.15
            114.74
                            126.09
                                   132.47
            146.02 153.41 160.67 167.91
                                           175.13
            182.34 189.55 218.81 281.02
10 10 6 1 439.30 437.95 437.53 437.79 438.39
            439.26
                    440.36
                            441.67
                                   443.19
                                           444.92
                            451.32 453.85
            446.85
                    448.99
                                           456.58
            459.51 462.64
                            477.11 515.47
10 10 7 1
            439.30
                    437.95
                            437.53 437.79
                                            438.39
                            441.67
            439.26
                    440.36
                                   443.19
                                            444.92
                    448.99
                            451.32
                                   453.85
                                            456.58
            446.85
            459.51 462.64
                            477.11 515.47
```

The example shows the first two and the last two records of type seven, as the data is too voluminous to be included.

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12.3.70 WCONHIST - Define Well Historical Production Rates and Pressures

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WCONHIST keyword defines production rates and pressures for wells that have been declared history matching wells by the use of this keyword. History matching wells are handled differently then ordinary wells that use the WCONPROD keyword for controlling their production targets and constraints. However, the wells still need to be defined like ordinary production wells using the WELSPECS keyword in the SCHEDULE section.

Note that although wells can be allocated to a group when they are specified by the WELSPECS keyword, history matching wells cannot operate under group control. Field and group reporting is still consistent for all wells allocated to a group, but history matching wells cannot be under group control.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I WELNAME			to eight characters in leng Is observed production r		None		
			ame (WELNAME) must ELSPECS keyword in tho occur.				
2	STATUS		ng that declares the stat he following character str		OPEN		
		 OPEN: the well is open to flow and will attempt to produce the required production volumes. 					
		fluids to surface; may occur within depending on a other connectior prevented by se	"stopped" at the surface a however, if there any open the wellbore and betwee connection's potential was. Inter-connection flootting the XFLOW varial In this case the well's be described below.	n connections then flow on the open connections ith respect to all the w (cross flow) can be ble on the WELSPECS			
		SHUT: the well is shut at the surface and downhole, this results in no flow at the surface and no cross flow downhole.					
		well's production is to be	nould always be set either e set to zero. Just setting rell is open to flow with a	a well's production rate			

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No.	Name	Description				
		Field	Metric	Laboratory		
3	TARGET	for the well, all the other reporting only. The simul the phase rate stated i	g that sets the observed of phases are calculated uncator will attempt to meet in items (4) to (6) and to one of the following cha	onstrained and used for the TARGET based on (10) on this keyword.	None	
		ORAT: the target defined by item (4)	et is set to the surface 4).	oil production rate as		
		WRAT: the targe defined by item (!	t is set to the surface w	ater production rate as		
		3) GRAT: the targe defined by item (6	t is set to the surface (5).	gas production rate as		
		4) LRAT: the target	t is set to the surface nd is calculated by the sim			
			is set to the in situ reser simulator using items (4),			
	6) BHP: the target r by item (10).	ate is set to the bottom-l	nole pressure as defined			
	keyword in the SCHEDI	rol mode may be reset of ULE section, from the tire changing the control mo	ne the WHISTCNTL is			
4	ORAT	A real positive value that target or constraint.	defines the observed sur	face oil production rate		
		stb/d	sm³/day	scc/hour		
		0.0	0.0	0.0	Defined	
5	WRAT	A real positive value that defines the observed surface water production rate target or constraint.				
		stb/d	sm³/day	scc/hour		
		0.0	0.0	0.0	Defined	
6	GRAT	A real positive value that target or constraint	defines the observed surf	ace gas production rate		
		Mscf/d	sm³/day	scc/hour		
		0.0	0.0	0.0	Defined	
7	VFPTAB		er than or equal to zero to be used for calculating the		None	
			ntered then the vertical vVFPPROD keyword in t via this item.			
	If this value is then rese	implies no vertical lift pe t to be greater than zero l's tubing head pressure. S eclared table number.	then the table will be			

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No.	Name		Description		Default	
		Field	Metric	Laboratory		
8	ALQ-WELL		t defines the artificial life PROD assigned to the w		None	
		VFPTAB vertical lift performance table and the artificial lift quantity ALQ-WELL are used with the well fluid rates to calculate the well's tubing head pressures values from the bottom-hole pressure.				
		Note that the units for variable on the VFPPROE		dent on the associated		
9	THP	This parameter is only us	A real positive value that defines the observed tubing head pressure. This parameter is only used for comparing the actual tubing head pressure given here with those calculated by the simulator, that is history marching wells can only controlled by either the surface injection rate or their bottom-hole pressure.			
		psia	barsa	atma		
		0.0	0.0	0.0	Defined	
10	ВНР	A real positive value that	defines the observed bo	ttom-hole pressure.		
		psia	barsa	atma		
		0.0	0.0	0.0	Defined	
11		Not Used				
		1				

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Notes:

Table 12.27:WCONHIST Keyword Description

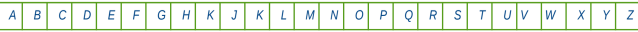
See also the WHISTCNTL that can be used to reset the TARGET phase, the GCONPROD and GCONINJ keywords to define a group's production and injection targets and constraints, and the WCONPROD keyword to define a production well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Note

One can use TARGET set to RESV in the initial history matching runs to get a "reasonable" pressure match, this ensures that the total reservoir withdrawals are correct, although the individual phase withdrawals will not match. Once a reasonable pressure match is achieved for the reservoir then one can reset TARGET to the sales phase, OIL or GAS, and continue with the matching of all the phases.

In oil reservoirs some engineers prefer to use LIQ rather than OIL as the TARGET phase, although one should consider as the water phase has no commercial value, the measurement accuracy is significantly less than the oil sales phase.

History matching wells are converted to ordinary wells by restating a well's control mode using either the WCONPROD or WELTARG keywords in the SCHEDULE section.



¹⁾ The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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Examples

The following example below shows the observed gas rates for the OP01 oil producer for the first quarter of 2000.

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```
-- 01 JAN 2000 START OF SCHEDULE SECTION
         WELL HISTORICAL PRODUCTION CONTROLS
-- WELL
         OPEN/
                 CNTL
                        OIL
                                WAT
                                       GAS
                                               VFP
                                                      VFP
                                                             THP
                                                                   RHP
                 MODE
                        RATE
                                RATE
                                       RATE
                                                      ALFQ
                                                             PRES
                                                                   PRFS
-- NAMF
         SHUT
                                               TABLE
WCONHIST
0P01
         OPEN
                 ORAT 15.5E3 100.0 1550
                                               10
                                                       1*
                                                             900.0 1*
DATES
01 FEB 2000 /
         WELL HISTORICAL PRODUCTION CONTROLS
- -
-- WELL
         OPEN/
                 CNTL
                        OIL
                                WAT
                                       GAS
                                               VFP
                                                      VFP
                                                             THP
                                                                   BHP
-- NAME
         SHUT
                 MODE
                        RATE
                                RATE
                                       RATE
                                               TABLE
                                                      ALFQ
                                                             PRES
                                                                   PRES
WCONHIST
0P01
         OPEN
                 ORAT 15.2E3 150.0 1520
                                               1*
                                                       1*
                                                             875.0 3250.0 /
DATES
01 MAR 2000 /
         WELL HISTORICAL PRODUCTION CONTROLS
- -
         OPEN/
                 CNTL
                        OIL
                                WAT
                                       GAS
                                               VFP
                                                      VFP
                                                             THP
                                                                   BHP
-- WFII
                 MODE
                                                      ALFQ
-- NAME
         SHUT
                        RATE
                                RATE
                                       RATE
                                               TABLE
                                                             PRES
                                                                   PRES
WCONHIST
                               200.0
                                               1 *
0P01
         OPEN
                 ORAT 15.0E3
                                      1500
                                                       1*
                                                             850.0 1*
```

From January I, 2000 well OP01 is open and is on oil rate control, and produces 15,500 stb/d oil, with the observed rates of 100 stb/d of water and 15.5 MMscf/d of gas. The well uses VFPPROD vertical lift table number 10 so that OPM Flow can calculate the tubing head pressures based on the fluids produced and the calculated pressures in the simulator.

The next example illustrates how to convert OP01 from a history match well to a normal production well at the start for the forecast run at August 1, 2017 using the WELTARG keyword.

```
DATES
01 AUG 2017 /
/
--
-- WELL PRODUCTION AND INJECTION TARGETS
--
-- WELL WELL TARGET
-- NAME TARG VALUE
WELTARG
OP01 THP 1* /
```

Here by defaulting the bottom-hole pressure via I* OPM Flow automatically applies the last bottom-hole pressure from the previous time step as the "constraining phase" together with the last historical rates as constraints. This ensures a smooth transition between history and prediction without having to resort to unreasonable changes to the model. This option is currently not implemented in OPM Flow but is expected to be incorporated in a future release.

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12.3.71 WCONINJ - Well Injection Targets and Constraints

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The WCONINJ is a legacy keyword that is no longer used in the commercial simulator and is not supported by OPM Flow. Instead well injection targets and constraints should be defined using the WCONINJE keyword in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.72 WCONINJE - Well Injection Targets and Constraints

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WCONINJE keyword defines injection targets and constraints for wells that have previously been defined by the WELSPECS keyword in the SCHEDULE section. Note that wells can be allocated to a group when they are specified by the WELSPECS keyword. Wells defined to be under group control will have their injection rates controlled by the group to which they belong, in addition to any well constraints defined for the wells using this keyword.

No.	Name	Description					
		Field	Metric	Laboratory			
I	WELNAME	IAME A character string of up to eight characters in length that defines the well name for which the well injection targets and constraints data are being defined.					
			ame (WELNAME) mus ELSPECS keyword in th occur.				
2	TYPE		ng that defines the type he following character str		None		
		I) GAS: for a gas inje	ection well.				
		2) OIL: for an oil inje					
		3) WAT: for a water	injection well.				
3	STATUS	A defined character stri should be set to one of t	OPEN				
		OPEN: the well is required injection	open for injection and wolumes.	vill attempt to inject the			
		fluids; however, if within the wellbo on a connection connections. Into by setting the XFI	"stopped" at the surface there any open connection re and between the open n's potential with resper-connection flow (cross LOW variable on the WEL vell's behavior will be similar	ons then flow may occur a connections depending ect to all the other flow) can be prevented LSPECS keyword to NO.			
		,	shut at the surface and one cross flow downhole.	downhole, this results in			
			s initially SHUT, but may l limit is violated. This o M Flow.				
	Note a well's STATUS should always be set either STOP or SHUT if well's production is to be set to zero. Just setting a well's injection rate zero means that the well is open for injection with a zero rate, this cause numerical issues especially for wells under THP control.						

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No.	Name	Description					
		Field	Metric	Laboratory			
4	TARGET		ng that sets the target injection be set to one of the follo		None		
		for the given we example, if TYPE	on phase will be control ell type as defined by t has been set to WAT th ection rate as defined by it	he TYPE variable. For en this would mean the			
		volume fluid rate variable. For exar	on phase will be control for the given well type mple, if TYPE has been set servoir volume injection	as defined by the TYPE to GAS then this would			
		3) BHP: the target r by item (7).	ate is set to the bottom-	hole pressure as defined			
		by item (8). If performance table	rate is set to the tubing he this option is selected es must be entered via th on and allocated to the w	then the vertical lift e VFPINJ keyword in the			
		group's target a	 GRUP: the well is under group control and injects its share of the group's target as set using the GCONINJ keyword in the SCHEDULE. Section. 				
5	RATE	A real positive value that defines the maximum surface injection rate target or constraint.					
		Liquid stb/d	Liquid sm³/day	Liquid scc/hour			
		Gas Mscf/d	Gas sm³/day	Gas scc/hour	None		
6	RESV	RESV A real positive value that defines the maximum reservoir volume injerate target or constraint.					
		rtb/d	rm³/day	rcc/hour	None		
7	ВНР	A real positive value the target or constraint.	at defines the maximum	bottom-hole pressure			
		Note the default value of basically means unlimited injection or no constraint and should therefore be avoided as the BHP will result in unrealistic well potentials as well as optimistic injection forecasts for the well.					
		psia	barsa	atma			
		10,0000	6,895	6,803	Defined		
8	THP	A real positive value the target or constraint.	nat defines the maximun	n tubing head pressure	None		
		psia	barsa	atma			
9	VFPTAB	A positive integer greater than or equal to zero that defines the vertical lift performance tables to be used for calculating the tubing head pressure for the well.					
		If a non-zero value is e must be entered via the allocated to the well via t					
			implies no vertical lift pe t be set to THP and in a ro.				

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No.	Name		Default		
		Field	Metric	Laboratory	
10		Not Used			
Ш		Not Used			
12		Not Used			
13		Not Used			
14		Not Used			
15		Not Used			

Revision: Rev-I

Notes:

1) The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.28:WCONINJE Keyword Description

See also the GCONPROD the GCONINJ keywords to define a group's production and injection targets and constraints, and the WCONPROD keyword to define a production well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines the injection targets and constraints for one gas injection well and one water injection well as follows:

```
WELL INJECTION CONTROLS
        FLUID OPEN/
                       CNTL
                                     RESV
                                             BHP
                                                   THP
                                                         VFP
-- WELL
                              SURF
-- NAME
         TYPE
                SHUT
                        MODE
                              RATE
                                     RATE
                                             PRSES PRES
                                                         TABLE
WCONINJE
                                     1*
                                                    1*
         GAS
                OPEN
                        GRUP
                              50E3
                                                          1*
GT01
WI01
         WAT
                OPEN
                        RATE
                                             5000.
                                                          1*
                              25E3
```

Well GI01 is a gas injection well directly under group control constrained by a maximum surface gas injection rate of 50 MMscf/d and well WI01 is an open water injection well with a surface water injection rate target of 25,000 stb/d, subject to a maximum bottom-hole pressure constraint 5,000 psia.

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12.3.73 WCONINJH - Well Historical Observed Injection Rates and Pressures

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WCONINJH keyword defines injection rates and pressures for wells that have been declared history matching wells by the use of this keyword. History matching wells are handled differently then ordinary wells that use the WCONINJE keyword for controlling their injection targets and constraints. However, the wells still need to be defined like ordinary injection wells using the WELSPECS keyword in the SCHEDULE section.

Note that although wells can be allocated to a group when they are specified by the WELSPECS keyword, history matching wells cannot operate under group control. Field and group reporting is still consistent for all wells allocated to a group, but history matching wells cannot be under group control.

No. Name		Description				
		Field	Metric	Laboratory		
I	WELNAME	A character string of up to eight characters in length that defines the well name for which the wells observed injection rates and pressures are being defined. Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.				
2	TYPE		njection well.		None	
3	STATUS	should be set to one of t 1) OPEN: the well is observed injectio 2) STOP: the well is however, if there the wellbore and connection's pote Inter-connection XFLOW variable the well's behavious below. 3) SHUT: the well is no injection and r Note a well's STATUS she well's injection is to be s	ng that declares the statiche following character stricts open for injection and we not volumes. "stopped" at the surface any open connections the between the open connectial with respect to all flow (cross flow) can be pon the WELSPECS keywor will be similar to the shot at the surface and concross flow downhole. Included always be set either et to zero. Just setting a wopen to flow with a zero.	ings: will attempt to inject the and will not inject fluids; in flow may occur within ections depending on a the other connections. Drevented by setting the ord to NO. In this case SHUT option described downhole, this results in a STOP or SHUT if the well's inject rate to zero	OPEN	

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No.	Name		Description		Default		
		Field	_				
4	RATE	A real positive value that	defines the observed surf	ace injection rate.			
		Liquid stb/d	Liquid sm³/day	Liquid scc/hour			
		Gas Mscf/d	Gas sm³/day	Gas scc/hour	0.0		
5	ВНР	A real positive value that	defines the observed bott	com-hole pressure.			
		psia	barsa	atma	0.0		
6	THP	A real positive value that	defines the observed tubi	ng head pressure.			
		This parameter is only used for comparing the actual tubing head pressure given here with those calculated by the simulator, that is history marching wells can only controlled by either the surface injection rate or their bottom-hole pressure.					
		psia	barsa	atma	None		
7	VFPTAB	A positive integer greater than or equal to zero that defines the vertical lift performance tables to be used for calculating the tubing head pressure for the well.					
		If a non-zero value is entered then the vertical lift performance tables must be entered via the VFPINJ keyword in the SCHEDULE section and allocated to the well via this item.					
		The default value of zero implies no vertical lift performance table initially. If this value is then reset to be greater than zero then the table will be used to calculate the well's tubing head pressure. Subsequently, the default is to use the previously declared table number.					
8		Not used and should be	defaulted with 1*.				
9		Not used and should be d	defaulted with 1*.				
10		Not used and should be o	defaulted with 1*.				
-11		Not used and should be d	defaulted with 1*.				
12	TARGET	A defined character string that sets the target injection control mode for the well. TARGET should be set to one of the following character strings:					
		rate for the given example, if TYPE	on well will be controlled well type as defined by has been set to WAT the ction rate as defined by it	the TYPE variable. For en this would mean the			
		BHP: the injection pressure as define	n well will be controlled by item (5).	ed by the bottom-hole			

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Notes:

The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.29:WCONINJH Keyword Description

This keyword should be repeated at various time steps to fully describe the historical injection performance of the wells. For example, as most production and injection data is reconciled on a monthly basis, then monthly time steps covering the injection history of the wells should be used with WCONINJH keyword entered on a monthly basis.

History matching well are converted to ordinary wells by restating a well's control mode using either the WCONINJE or WELTARG keywords in the SCHEDULE section.

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Example

The following example below shows the observed gas rates for the GI01 gas injector for the first quarter of 2000.

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```
-- 01 JAN 2000 START OF SCHEDULE SECTION
         WELL HISTORICAL INJECTION CONTROLS
-- WELL FLUID OPEN/
                       SURF
                               RESV
                                      BHP
                                            THP
                                                  VFP
                                                          NOT
                                                                CNTL
-- NAME TYPE
                SHUT
                       RATE
                               RATE
                                      PRSES PRES
                                                         USED MODE
                                                  TABLE
WCONINJH
                                                          4*
                OPEN 15.5E3 1*
                                      1*
                                                                1* /
GI01
         GAS
                                            5462
                                                   12
DATES
01 FEB 2000 /
         WELL HISTORICAL INJECTION CONTROLS
- -
-- WELL FLUID OPEN/
                       SURF
                               RESV
                                      BHP
                                            THP
                                                  VFP
                                                          NOT
                                                                CNTL
-- NAME TYPE
                SHUT
                       RATE
                               RATE
                                      PRSES PRES
                                                  TABLE
                                                         USED
                                                               MODE
WCONINJH
                                                          4*
                                      1*
                                                                1*
GI01
         GAS
                OPEN 15.9E3 1*
                                            5468
                                                   1*
DATES
01 MAR 2000 /
- -
         WELL HISTORICAL INJECTION CONTROLS
                                      BHP
        FLUID OPEN/
                       SURF
                               RFSV
                                            THP
                                                  VFP
                                                         NOT
                                                                CNTI
-- WELL
-- NAME
         TYPE
                SHUT
                       RATE
                               RATE
                                      PRSES PRES
                                                  TABLE
                                                         USED
                                                               MODE
WCONINJH
                                      1*
                                                                1* /
GI01
         GAS
                OPEN 17.2E3 1*
                                            5489
                                                   1*
```

Well Gl01is declared as a gas injection well under gas rate control as TARGET variable is defaulted to rate control by using 1* (the last entry on the record). In addition, the well users vertical lift table VFPINJ number 12 (as shown at January 1, 2000) to calculate the tubing head pressures for the well. Note that it is not necessary to declare the VFPINJ table number if it remains the same for subsequent time steps and thus the default 1* is used to indicate the last entry should be used.

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12.3.74 WCONPROD - Define Well Production Targets and Constraints

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WCONPROD keyword defines production targets and constraints for wells that have previously been defined by the WELSPECS keyword in the SCHEDULE section. Note that wells can be allocated to a group when they are specified by the WELSPECS keyword. Wells defined to be under group control will have their production rates controlled by the group to which they belong, in addition to any well constraints defined for the wells using this keyword.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	WELNAME		to eight characters in leng production targets and co		None			
			ame (WELNAME) must ELSPECS keyword in tho occur.					
2	STATUS		ng that declares the state he following character stri		OPEN			
		 OPEN: the well is open to flow and will attempt to produce the required production volumes. 						
		2) STOP: the well is "stopped" at the surface and will not produce any fluids to surface; however, if there any open connections then flow may occur within the wellbore and between the open connections depending on a connection's potential with respect to all the other connections. Inter-connection flow (cross flow) can be prevented by setting the XFLOW variable on the WELSPECS keyword to NO. In this case the well's behavior will be similar to the SHUT option described below.						
		,	shut at the surface and offace and no cross flow do	-				
		 AUTO: the well is initially SHUT, but may be opened automatically if an economic limit is violated. This option is currently not supported by OPM Flow. 						
		well's production is to be to zero means that the	nould always be set either e set to zero. Just setting well is open to flow wit pecially for wells under Th	a well's production rate th a zero rate, this will				

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No.	Name		Description		Default	
		Field	Metric	Laboratory		
3	TARGET	well, all the other phase will attempt to meet the	ng that sets the target pr s will therefore act as co TARGET based on the ph ord. TARGET should be set	nstraints. The simulator hase rate stated in items	None	
		ORAT: the target defined by item (et is set to the surface 4).	oil production rate as		
	WRAT: the target defined by item ()	et is set to the surface w 5).	ater production rate as			
	GRAT: the target defined by item (et is set to the surface 6).	gas production rate as			
		t is set to the surface as defined by item (7).	liquid (oil plus water)			
	5) RESV: the target defined by item (is set to the in situ re	servoir volume rate as			
	6) BHP: the target r by item (9).	rate is set to the bottom-	nole pressure as defined			
	by item (10). If performance tabl	rate is set to the tubing h this option is selected es must be entered via th section and allocated to th	then the vertical lift e VFPPROD keyword in			
			s under group control and the GCON on.			
4	ORAT	A real positive value that target or constraint.	defines the maximum sur	face oil production rate		
		stb/d	sm³/day	scc/hour	None	
5	WRAT	A real positive value that defines the maximum surface water production rate target or constraint.				
		stb/d	sm³/day	scc/hour	None	
6	GRAT		A real positive value that defines the maximum surface gas production rate target or constraint			
		Mscf/d	sm³/day	scc/hour	None	
7	LRAT	A real positive value that defines the maximum surface liquid (oil plus water) production rate target or constraint.				
		stb/d	sm³/day	scc/hour	None	
8	RESV	A real positive value production rate target of	that defines the maxin r constraint.	num reservoir volume		
		rtb/d	rm³/day	rcc/hour	None	

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No.	Name		Description		Default	
		Field	Metric	Laboratory		
9	ВНР	A real positive value the target or constraint.	nat defines the minimum	bottom-hole pressure		
			f one atmosphere should well potentials as well as			
		psia	barsa	atma		
		14.70	1.01325.	1.0	Defined	
10	THP	A real positive value that or constraint.	defines the minimum tubi	ng head pressure target		
		TARGET has been set to	of zero should be avoided o THP, as this will result in nce a well must flow ag acilities.	n optimistic production		
		psia	barsa	atma		
		0.0	0.0	0.0	Defined	
Ш	VFPTAB	A positive integer greate lift performance tables to for the well.	0			
			ntered then the vertical VFPPROD keyword in t via this item.			
		this case TARGET cannot	The default value of zero implies no vertical lift performance tables and in this case TARGET cannot be set to THP and in addition item (10) should be defaulted or set to zero.			
12	ALQ-WELL		t defines the artificial lift PROD assigned to the we		0.0	
		VFPTAB vertical lift perfo WELL are used with the pressures values from the				
		Note that the units fo variable on the VFPPROD	r ALQ-WELL is depend keyword.	ent on the associated		
13		Not Used				
14		Not Used				
15		Not Used				
16		Not Used				
17		Not Used				
18		Not Used				
19		Not Used				
20		Not Used				

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Notes:

1) The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.30:WCONPROD Keyword Description

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See also the GCONPROD and GCONINJ keywords to define a group's production and injection targets and constraints, and the WCONINJE keyword to define an injection well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines the production targets and constraints for five wells as follows:

```
WELL PRODUCTION WELL CONTROLS
- -
                 CNTL OIL
                              WAT
                                     GAS
                                           LIO
                                                   RES
                                                           RHP
                                                                         VFP
                                                                                 VFP
-- WELL
         OPEN/
                                                                  THP
                 MODE
                       RATE
-- NAME
         SHUT
                              RATE
                                     RATE
                                           RATE
                                                   RATE
                                                           PRES
                                                                  PRES
                                                                         TABLE
                                                                                ALF0
WCONPROD
                              1*
                                            1*
                                                   1 *
         OPEN
                 GRUP
                       5E3
                                                           500.0
0P01
                              1*
                                      1*
                                             1*
                                                   1*
0P02
         OPEN
                 GRUP 10E3
                                                           200.0
                                                                  500.0
                                                                                 0.0
                              1*
                                            1*
                                                   1*
                                      1*
0P03
         OPEN
                 GRUP 15E3
                                                           200.0
                                                                  500.0
                                                                                10.0
                              1*
                                      1*
                                            1*
                                                   1*
0P04
         OPEN
                 ORAT 20E3
                                                           500.0
0P05
          SHUT
                 GRUP 20E3
                                                           500.0
```

Well OP01 is open and is on group control, subject to a maximum oil rate constraint of 5,000 stb/d and a minimum bottom-hole pressure of 500 psia. OP02 is also open and on group control but it's maximum oil rate constraint has been set 10,000 stb/d, and is subject to a minimum bottom-hole pressure limit of 200 psia and a minimum tubing head pressure limit of 500 psia using VFPPROD vertical lift table number two. Well OP03 is very similar to OP02, but with a 15,000 stb/d maximum oil constraint and using VFPPROD vertical lift table number three with an artificial lift parameter of 10. The next three wells are not on group control, for example, well OP04 is open and has an oil rate target of 20,000 stb/d, subject to a minimum bottom-hole pressure of 500 psia. Finally, well OP05 is shut and will not be brought back on production despite being put under group control, as the well has been declared shut.

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12.3.75 WDFACCOR - GAS FLOW DEPENDENT SKIN FACTOR

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Description

Defines the parameters to calculate a gas well's connection D-factor based on a correlation for the coefficient of inertial resistance, usually known as β , in Forchheimer's flow equation 64.65.66 and 67.

This keyword is ignored by OPM Flow and has no effect on the simulation.

⁶⁴ Dake, L.P. Fundamentals of Reservoir Engineering, Amsterdam, The Netherlands, Elsevier Science BV (1978) Chapter 8.6, pages 252-257.

⁶⁵ Geertsma, J., 1974. Estimating the Coefficient of Inertial Resistance in Fluid Flow Through Porous Media. Soc.Pet.Eng.J., October: 445-450.

⁶⁶ Gewers, C.W.W. and Nichol, L.R., 1969. Gas Turbulence Factor in a Microvugular Carbonate. J.Can.Pet.Tech., April.

⁶⁷ Wong, S.W., 1970. Effects of Liquid Saturation on Turbulence Factors for Gas Liquid Systems. J.Can.Pet.Tech., October

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12.3.76 WDRILTIM - Define Drilling Parameters for Automatic Drilling of New Wells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

WDRILTIM defines the automatic drilling parameters used to describe the numbers of days taken to to drill a well, the drilling status of the well, and status of other wells when drilling an automatically drilled well.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.77 WECON - WELL ECONOMIC CRITERIA FOR PRODUCTION WELLS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WECON keyword defines economic criteria for production wells that have previously been defined by the WELSPECS and WCONPROD keywords in the SCHEDULE section.

Note that wells can be allocated to a group when they are specified by the WELSPECS keyword and groups can also have economic controls. Wells under group control are therefore subject to the economic criteria set via the GCONPROD and GECON keywords in the SCHEDULE section and the controls specified by the WECON keyword. Note that GECON is not supported by OPM Flow in the current release.

No.	Name		Description		Default
		Field	Metric	Laboratory	
I	WELNAME		to eight characters in leng economic criteria data is l	-	None
			name (WELNAME) must ELSPECS keyword in the occur.		
2	ORAT		nat defines the minimum which an economic act		
		STATUS variable	le set to AUTO on the Co ection, then one of	os in the well with the OMPDAT keyword in the these connections (or	
		STATUS variab	le set to AUTO on the C e shut or stopped as requ	s in the well with the COMPDAT keyword, then uested by item (9) of the	
			orted by OPM Flow as ST d is currently not suppo shut or stopped.		
		A value less than or equa	ll to zero switches off this	criterion.	
		stb/d	sm³/day	scc/hour	0.0

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No.	Name		Description		Default
		Field	Metric	Laboratory	
3	GAS	-	nat defines the minimum which an economic act	- 1	
		STATUS variable	ection, then one of	OMPDAT keyword in the	
		STATUS variab	no remaining connection le set to AUTO on the C e shut or stopped as requ word.	OMPDAT keyword, then	
			orted by OPM Flow as ST d is currently not suppo shut or stopped.	-	
		A value less than or equa	I to zero switches off this	criterion.	
		Mscf/d	sm³/day	scc/hour	0.0
4 WCUT			t defines the maximum e		
			$f_{w}=rac{q_{w}}{q_{w}+q_{o}}$, a vater cut limit is exceede		
		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	l to zero switches off this	criterion.	
		dimensionless	dimensionless	dimensionless	0.0
5	GOR		t defines the maximum e onomic action will take p		
		A value less than or equa	I to zero switches off this	criterion.	
		Note that this feature is	currently not supported in	OPM Flow	
		Mscf/stb	sm³/sm³	scc/scc	0.0
6	WGR		t defines the maximum e n economic action will ta		
		A value less than or equa	I to zero switches off this	criterion.	
		Note that this feature is	currently not supported ir	OPM Flow.	
		stb/Mscf	sm³/sm³	scc/scc	0.0

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No.	Name		Description		Default	
		Field	Metric	Laboratory		
7	ACTION		, or WGR limits are vio	ction to be taken if the slated. ACTION should be	None	
		I) NONE: no actio	n is taken.			
				ection. If connections have worst offending completion		
		connections have		nection and all below it. If ompletions then the worst libe closed.		
		4) WELL: shut or st	top the well as per the A	UTO variable on the		
	WELSPECS keyword.					
	The corrective action takes places at the end of the time step in which the constraint is violated.					
		Only ACTION equal to	CON is currently suppo	orted by OPM Flow.		
8	END		A defined character string that defines if the simulation should terminate if the well is shut or stopped. END should be set to one of the following character strings:			
		I) NO: no action is	taken and the run conti	nues.		
		2) YES: terminate the	he run at the next repor	t time step.		
		Only the default value o	f NO is supported in OF	PM Flow.		
9		Not used				
10		Not used				
11		Not used				
12		Not used				
13		Not used				
14		Not used				
15		Not used				
16		Not used				

Revision: Rev-I

Notes:

Table 12.31:WECON Keyword Description

See also the WELSPECS keyword to define a wells shut-in or stop options, GCONPROD for group controls, and GECON for setting a group's economic criteria. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines one oil well and one gas well using the WELSPECS keyword, together with their economic criteria.

The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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```
-- WELL SPECIFICATION DATA
-- WELL
             GROUP
                          LOCATION
                                     BHP
                                            PHASE
                                                    DRAIN
                                                            INFLOW
                                                                     SHUT
                                                                            CROSS
                                                                                    PRESS
-- NAME
             NAME
                          Ι
                                J
                                     DEPTH
                                            FLUID
                                                    AREA
                                                            EQUA.
                                                                     ΙN
                                                                            FLOW
                                                                                    TABLE
WELSPECS
GP01
           PLATFORM
                          14
                               13
                                     1*
                                              GAS
                                                    1*
                                                            GPP
                                                                     SHUT
                                                                             NO
                                     1*
                                                    1*
                                                                                    1*
0P01
           PLATFORM
                          28
                               96
                                              OIL
                                                            STD
                                                                     SHUT
                                                                             NO
/
          WELL ECONOMIC CRITERIA FOR PRODUCTION WELLS
-- WELL
          MIN
                  MIN
                         MAX
                                 MAX
                                         MAX
                                                 CNTL
                                                          END
-- NAME
          ORAT
                  GRAT
                         WCUT
                                 GOR
                                         WGR
                                                 MODE
                                                          RUN
WECON
          1*
                         1*
                                 1*
                                         1*
GP01
                  5.0E3
                                                'WELL'
                                                         'NO'
                                         1*
                 1*
                                                'WELL'
                                                         'YES'
0P01
          500
                         0.95
                                 15E3
```

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Well GP01 has a minimum economic gas rate of 5 MMscf/d and will shut-in if the gas rate falls below this rate, but the simulation will continue even if this occurs. Well OP02 as a minimum economic oil rate of 500 stb/d, a maximum water cut limit of 95%, and a maximum GOR of 15 MMscf/d, if any any of these limits are violated the well will be shut-in and the run terminated at the next reporting time step.

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12.3.78 WEFAC - Define Well Efficiency

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

Defines a well's efficiency or up-time.

This keyword is ignored by OPM Flow and has no effect on the simulation.

See the GEFAC keyword in the SCHEDULE section that is implemented in OPM Flow and can be used to set a group's efficiency.

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12.3.79 WELCNTL - Modify Well Control and Targets

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WELCNTL keyword modifies a wells' target control and value, both rates and pressures, for previously defined wells without having to define all the variables on the well control keywords: WCONPROD, WCONHIST, WCONDINJE, or WCONDINJH keywords. Variables not changed by the WELCNTL keyword remain the same as those previously entered via the well control keywords or previously entered WELCNTL keywords. Note that the well must still be initially be fully defined using the WCONPROD or WCONINJE keywords. All the aforementioned keywords are described in the SCHEDULE section.

OPM Flow does not recognize this keyword and the keyword may cause the program to terminate, but it is documented here for completeness.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
1	WELNAME	A character string of up to eight characters in length that defines the well name for which the well production rates and pressures data are being redefined.				
		Note that the well n previously using the WE keywords in the SCHEDU	ELSPECS and WCONPRO	OD (or WELCONINJE)		
2	TARGET	A defined character striit the value of the item is so	_	be changed for the well	None	
		I) ORAT: reset the item (3).	surface oil production	rate value as defined by		
		2) WRAT: reset the item (3).	surface water production	rate value as defined by		
		3) GRAT: reset the item (3).	surface gas production	rate value as defined by		
		4) LRAT: reset the star defined by (3).	urface liquid (oil plus wate	er) production rate value		
		5) RESV: reset he in	situ reservoir volume rat	e value as defined by (3).		
		6) BHP: reset the bo	ttom-hole pressure value	as defined by item (3).		
		7) THP: reset the tuby item (3).	ıbing head pressure value	e for the well as defined		
		8) VFP: reset the ver (3).	rtical lift performance tab	ole number as defined by		
		9) LIFT: reset the performance table		r use with vertical lift		
		10) GUID: reset the control.	guide rate value for well	s operating under group		
		Note TARGET redefines value on item (4). For ex defined by the previous TARGET equal to LRAT the given value. That is in previously requested of SCHEDULE section to check the section of the section in the section is the section of the section is a section of the section of the section is a section of the section is a section of the section of	ample, if a well is operatiusly entered WCONPR with a value, sets the TAI set well will be targetial ratel. Use the WELT	ing on ORAT control, as OD keyword, entering RGET to liquid rate with ng anliquid rate not the FARG keyword in the		

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No.	Name	Description				
		Field	Metric	Laboratory		
3	VALUE	A real positive value the TARGET	at defines the value of the	he variable declared by		
	Liquid	stb/d	sm³/day	scc/hour		
	Gas	Mscf/d	sm³/day	scc/hour		
	Res Vol	rb/d	rm³/day	rcc/hour		
	Pressure	psia	barsa	atma		
	VFP	dimensionless	dimensionless	dimensionless		
	LIFT	same as	same as	same as		
		VFPPROD or VFPINJ	VFPPROD or VFPINJ	VFPPROD or VFPINJ	None	

Revision: Rev-I

Notes:

The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.32:WELCNTL Keyword Description

If a well is currently a history matching well, then WELCNTL can be used to change the well to a standard well.

See also the WELTARG keyword, in the SCHEDULE section that can be used to reset a well's target and constraints of both rates and pressures.

Example

The following example below shows the oil rates for the OP01 oil producer at the start of the schedule section (January I, 2000).

```
-- 01 JAN 2000 START OF SCHEDULE SECTION
        WELL PRODUCTION WELL CONTROLS
                                                                       VFP
-- WELL OPEN/
               CNTL
                       OIL
                              WAT
                                     GAS
                                           LIO
                                                  RES
                                                         BHP
                                                                 THP
                                                                              VFP
                MODE
                                     RATE RATE
-- NAME SHUT
                       RATE
                              RATE
                                                  RATE
                                                         PRES
                                                                PRES TABLE
                                                                             ALFQ
WCONPROD
                ORAT
                              1*
                                     1*
                                           1*
                                                  1*
                                                                              1* /
0P01
         OPEN
                       3000
                                                         750.0 500.
DATES
01 FEB 2000 /
         WELL CONTROL MODE AND OPERATING TARGET
   WELL WELL
                TARGET
   NAME CNTL
                VALUE
- -
WELCNTL
                5000
0P01
         LRAT
```

From January I, 2000 to February I, 2000 well OP01 is open and is on oil rate control and has a target oil rate of 3,000 stb/d and uses VFPPROD vertical lift table number 9 with a minimum tubing head pressure constraint of 500 psia. After February I, 2000 the well is changed to liquid control with a target rate of 5,000 stb/d of liquid and all the other parameters remain unchanged.

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12.3.80 WELOPEN - Define Well and Well Connections Flowing Status

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WELOPEN keyword defines the status of wells and well connection and is used to open and shut previously defined well and well connections without having to re-specify all the data on the well control keywords: WCONPROD, WCONHIST, WCONINJE, or WCONINJH keywords. Note that the well must still be initially be fully defined using the WCONPROD or WCONINJE keywords. All the aforementioned keywords are described in the SCHEDULE section

No.	Name	Description							
		Field	Metric	Laboratory					
I	WELNAME	A character string of up name for which the well a			None				
		Note that the well n previously using the W otherwise an error may o	ELSPECS keyword in th						
2	STATUS	A character string of leading connections' operational following character string	status, STATUS should		OPEN				
		I) OPEN: the connec	ctions are open to flow.						
		2) SHUT: the connec	ctions are closed to flow (shut-in).					
			ection are initially close economic limit is violated						
3	I	An integer less than or e in the I-direction.	qual to NX that defines t	the connection location	I *				
4	J	An integer less than or e in the J-direction.	equal to NY that defines t	the connection location	I *				
5	К	An integer less than or e in the K-direction.	equal to NZ that defines t	the connection location	 *				
6	KI	An integer less than or of location in the K-direction		the UPPER connection	 *				
		If connections have been keyword, then K1 refer connection (layer) value.							
7	K2	An integer less than or e location in the K-directio		he LOWER connection	 *				
		If connections have been lumped into completions via the COMPLUMP keyword, then K2 refers to the completion number instead of the connection (layer) value.							

Notes:

Table 12.33:WELOPEN Keyword Description

If variables I, J K, K I and K2 are all set to a negative number or defaulted with I* then STATUS is applied to the well and the well connection remain unchanged.

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The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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If variables I, J K, K I and K2 are all set to zero or a positive value then STATUS is applied to the defined connections and the well status remains unchanged. The defined connections are those with the I, J, K variables the specified location and a completion number in the range specified by K I and K2.

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See also the COMPDAT keyword to define a well's connections, the COMPLUMP keyword to group well connections into well completions, the WCONPROD and WCONDINJE keywords to define a well's production and injections targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Examples

The following example defines three vertical oil wells using the WELSPECS keyword and their associated connection data.

```
-- WELL SPECIFICATION DATA
                                      BHP
                                                                       SHUT
                          LOCATION
                                              PHASE
                                                                              CROSS
                                                                                      PRESS
-- WELL
             GROUP
                                                     DRAIN
                                                              INFLOW
-- NAME
             NAME
                           Т
                                 J
                                      DEPTH
                                              FLUID
                                                     AREA
                                                              EQUA.
                                                                       ΤN
                                                                              FLOW
                                                                                      TABLE
WELSPECS
                                      1*
                                                                                      1*
                                                      1 *
           PLATFORM
                                               OTI
                                                                       OPEN
                                                                               NO
0P01
                          14
                                13
                                                              STD
0P02
           PLATFORM
                          28
                                96
                                      1*
                                               OIL
                                                      1*
                                                              STD
                                                                       OPEN
                                                                               NO
                                                                                      1*
                                                                                      1*
                                      1*
                                                     1*
0P03
           PLATFORM
                         128
                                56
                                               OIL
                                                              STD
                                                                       OPEN
                                                                               NO
- -
          WELL PRODUCTION WELL CONTROLS
          OPEN/
                  CNTL
                          OIL
                                  WAT
                                                                 BHP
                                                                                VFP
                                                                                        VFP
-- WELL
                                          GAS
                                                 LIQ
                                                         RES
                                                                         THP
-- NAME
          SHUT
                  MODE
                          RATE
                                  RATE
                                          RATE
                                                 RATE
                                                         RATE
                                                                 PRES
                                                                         PRES
                                                                                TABLE
                                                                                        ALFQ
WCONPROD
          SHUT
                  GRUP
                          1*
                                  1*
                                          1*
                                                 1*
                                                         1*
                                                                 200.0
/
- -
          WELL CONNECTION DATA
-- WELL
              LOCATION ---
                               OPEN
                                       SAT
                                              CONN
                                                      WELL
                                                              KH
                                                                     SKIN
                                                                             D
                                                                                    DTR
-- NAME
           II
                JJ
                    K1
                         K2
                               SHUT
                                       TAB
                                              FACT
                                                      DIA
                                                              FACT
                                                                    FACT
                                                                             FACT
                                                                                   PEN
COMPDAT
                                                                                   'Z' /
           1*
               1*
                                       1*
                                              1*
                                                     0.708
                                                             1*
                                                                    0.0
                                                                            1*
0P01
                     1
                         10
                               SHUT
           1*
                                                                                   'Z' /
               1*
                                       1*
                                                            1*
                                                                           1*
                                              1*
0P01
                    15
                         30
                               SHUT
                                                    0.708
                                                                   0.0
                                                                                   'Z' /
           1*
                                       1*
                                              1*
                                                            1*
                                                                            1*
0P01
               1*
                    35
                                                    0.708
                                                                   0.0
                         90
                               SHUT
                                                                                   'Z' /
                                                                            1*
0P02
                               SHUT
                                                     0.708
                                                                    0.0
                     1
                         10
0P03
           1*
                1*
                                       1*
                                              1*
                                                                            1 *
                    35
                         90
                               SHUT
                                                    0.708
                                                                    0.0
          DEFINE WELL AND WELL CONNECTIONS FLOWING STATUS
    WELL WELL
                  --LOCATION--
                                  COMPLETION
    NAME STAT
                    Ι
                              Κ
                                  FIRST LAST
                         J
WELOPEN
          OPEN
0P01
0P01
          OPEN
                    0
                         0
                              0
                                    35
                                           90
0P02
          OPEN
          OPEN
                                      2
                                            5
0P02
                    0
                         0
                               0
0P03
          OPEN
0P03
          OPEN
                    0
                         0
                               0
                                      0
                                            0
```

In this example the first record of each well WELOPEN keyword changes the well status from shut (as per the WCONPROD keyword) to open. Then for well OP01 well connections 35 to 90 are opened for flow, connections two to five for well OP02 and all the connections for well OP03.

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The next example shows the use of the COMPLUMP keyword to group the well connections into well completions for wells OP01 and OP03, and then use the WELOPEN keyword to open the well and the well connections.

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```
- -
         ASSIGN WELL CONNECTIONS TO COMPLETIONS
- -
         --- LOCATION ---
-- WELL
                            COMPL
-- NAME
          ΙI
              JJ K1 K2
                             NO.
COMPLUMP
                                                               / COMPLETION NO. 01
0P01
                       10
                                                               / COMPLETION NO. 02
0P01
           0
               0 15
                       30
                              2
0P01
           0
               0
                   35
                       90
                              3
                                                               / COMPLETION NO. 03
0P03
           0
                0
                   35
                       90
                              3
                                                               / COMPLETION NO. 03
- -
         DEFINE WELL AND WELL CONNECTIONS FLOWING STATUS
    WELL WELL
                 --LOCATION--
                                COMPLETION
    NAME STAT
                   Ι
                       J
                            Κ
                               FIRST LAST
WELOPEN
         OPEN
0P01
0P01
         OPEN
                   0
                       0
                            0
                                   3
                                          3
0P02
         OPEN
0P02
         OPEN
                             0
         OPEN
0P03
0P03
         OPEN
                                          3
```

Again, the first record of each well WELOPEN keyword changes the well status from shut (as per the WCONPROD keyword) to open. Then for well OP01 well completion number three is opened (connections 35 to 90), connections two to five for well OP02 and completion number three (all the connections) for well OP03.

Note the completion number for well OP03 was named completion number three, but it could have been named number one as well. The reason why it was named number three instead of one was because it was assumed (for the example) that layers 35 to 90 represent a particular reservoir, and therefore allowing for the tracking of completions for individual reservoirs., as shown in the example.

This example shows how one can open all the wells and well completions for a given reservoir.

```
DEFINE WELL AND WELL CONNECTIONS FLOWING STATUS
_ _
    WELL WELL
                --LOCATION--
                               COMPLETION
    NAME STAT
                  I J K FIRST LAST
WELOPEN
         OPEN
1 * 1
         OPEN
                                  3
                  0
                       0
                            0
                                        3
0P02
         SHUT
                                  0
                                         0
0P02
         OPEN
                                        5
```

In this case well OP01 and OP03 are opened via completion number three, and well OP02 is opened on well connection (or layer) number three – which we do not want. Hence, all the connection for OP02 are shut, and then connections two to five are opened instead for well OP02.

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12.3.81 WELPI - Define Well Productivity and Injectivity Indices

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

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Description

The WELPI keyword is used to define a well's productivity or injectivity index and values enter on this keyword for a given well will override any previously calculated values and values previously entered using this keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.82 WELSEGS - Define Multi-Segment Wells and Their Segment Structure

1.10.10.10.10.10.10.10.10.10.10.10.10.10	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

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Description

The WELSEGS keyword defines a well to be a multi-segment well and defines the well's segment structure. Note that the well must have been previously define by the WELSPECS keyword in the SCHEDULE section and that the WELSEGS keyword should be repeated for each multi-segment well in the model.

No.	Name		Description		Default				
		Field	Metric	Laboratory					
1-1	WELNAME		to eight characters in leng		None				
			name (WELNAME) must ELSPECS keyword in th occur.						
1-2	BHPREF	reporting the bottom ho	be entered that defines t le pressure for the multi-s e nodal point of the top se	egment well. Ideally this					
		example the well config	ed multiple times for the guration changing througl ata the first time the keyw	n time, then it is only					
		feet	m	cm	None				
1-3	TUBDZ		TUBDZ is a real positive value that defines the length of the tubing from the tubing head or wellhead at the surface to the nodal point of the top segment (BHPREF). Tubing pressures from BHREF up to the tubing length of TUBDZ to the						
		surface are not calculate taken into account by the	HREF up to the tubing lend by the multi-segment we VFP tables allocated to words in the SCHEDULE so	vell option as these are vell and entered via the					
			or defaulted then the tul he top segment, that is BH						
		feet	m	cm	0				
I-4	WBORVOL	volume for the top segm	ositive value that defines nent, that is from the tubi point of the top segment (ng head or wellhead at					
		The default value of 1.0 x	c 10 ⁻⁵ results in minimal we	ellbore storage.					
		ft ³	m³	cm³	1.0E-5				
1-5	TUBOPT	TUBOPT is a character string that defines the type of length and depth data entered for DEPTH1 and DEPTH2 on the second record and should be set to one of the following:							
		I) INC: Increment	tal values, that is the length	of each segment.					
		2) ABS: Absolute v	values, that si the depth of	each segment.					
		There is no default value explicitly defined.	bove options must be						

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No. Name Description **Default** Field Metric Laboratory 1-6 **PRESOPT** PRESOPT is a character string that defines the pressure drop calculation **HFA** used for each well segment and should be set to one of the following: HFA: Sets the pressure calculation to include the hydrostatic, friction and acceleration terms. HF-: Sets the pressure calculation to include the hydrostatic and friction terms only. H--: Sets the pressure calculation to include the hydrostatic pressure drop term only. The default value for PRESOPT of HFA sets the pressure calculation to include the hydrostatic, friction and acceleration terms. 1-7 **FLOWOPT** HO FLOWOPT is a character string that defines the type of multi-phase calculation used for each well segment and should be set to one of the following: HO: Sets the multi-phase calculation to the homogeneous model, that is all phases flow at the same velocity. DF-: Sets the multi-phase calculation to the Drift Flux Model. OPM Flow only supports the default value of HO. 1-8 **XCORD** A real positive values equal to or greater than zero that defines the coordinate in the x-direction of the nodal point of the top segment that is used for display purposes only. Currently this option is not supported by OPM Flow. feet cm None 1-9 **YCORD** A real positive value equal to or greater than zero that defines the coordinate in the y-direction of the nodal point of the top segment that is used for display purposes only. Currently this option is not supported by OPM Flow. None cm 1-10 **XAREA** XAREA is real positive value equal to or greater than zero that defines the cross sectional area of the pipe wall used in thermal conductivity calculations for when the temperature calculation is activated by the TEMP keyword in the RUNSPEC section. Currently this option is not supported by OPM Flow. ft^2 m^2 cm^2 None 1-11 **VHEATCAP** VHEATCAP is real positive value equal to or greater than zero that defines the volumetric heat capacity of the pipe wall used in thermal conductivity calculations for when the temperature calculation is activated by the TEMP keyword in the RUNSPEC section. Currently this option is not supported by OPM Flow. J/cm/hr/K Btu/ft/day/°R kJ/m/day/K None

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2-8

EIPSILON

No. Name Description **Default** Field Metric Laboratory 1-12 **THCON** THCON is real positive value equal to or greater than zero that defines the thermal conductivity of the pipe wall used in thermal conductivity calculations for when the temperature calculation is activated by the TEMP keyword in the RUNSPEC section. Currently this option is not supported by OPM Flow. Btu/ft/day/°R kJ/m/day/K I/cm/hr/K None 1-13 Record terminated by a "/" Not **Applicable** 2-1 ISEG I A positive integer greater than or equal to two and less than or equal to None MXSEGS on WSEGDIMS keyword in the RUNSPEC section that defines the start of a segment 2-2 ISEG2 A positive integer greater than or equal to two and less than or equal to None ISEG1 on this record and MXSEGS on the WSEGDIMS keyword in the RUNSPEC section that defines the end of a segment. 2-3 **IBRANCH** A positive integer greater than or equal to one and less than or equal to None MXBRAN on WSEGDIMS keyword in the RUNSPEC section that defines the branch number of a segment. All segments on the main stem must have IBRANCH set to one and lateral branches should have values between two and MXSEGS on the WSEGDIMS keyword in the RUNSPEC section. ISEG3 2-4 A positive integer greater than or equal to two and less than or equal to None MXSEGS on the WSEGDIMS keyword in the RUNSPEC section that defines the outlet segment. 2-5 **DEPTHI** DEPTH I is a real positive value that: 1) If TUBOPT is set to INC then DEPTH1 is the total length of the tubing for this segment. If TUBOPT is set to ABS then DEPTH is the length of the tubing from the tubing head or wellhead at the surface to the last segment in the range. feet cm None 2-6 DEPTH2 DEPTH2 is a real positive value that: If TUBOPT is set to INC then DEPTH2 is the total incremental depth change of the tubing for this segment. If TUBOPT is set to ABS then DEPTH defines the depth of the tubing at the last nodal point of this segment. in this range. None 2-7 ID A real positive value that defines the tubing internal diameter of the segment for the well.

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None

None

cm

	Α	В	С	D	Ε	F	G	Н	Κ	J	Κ	L	М	N	0	Р	Q	R	S	Т	U	V	W	Χ	Υ	Ζ
--	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

segment for the well.

A real positive value that defines the tubing absolute roughness of the

m

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No. Name Description **Default** Field Metric Laboratory 2-9 **XAREASEG** XAREA is real positive value equal to or greater than zero that defines the cross sectional area of the pipe wall used in thermal conductivity calculations for when the temperature calculation is activated by the TEMP keyword in the RUNSPEC section. Currently this option is not supported by OPM Flow. cm²None 2-10 **VOLSEG** VOLSEG is a real positive value that defines the effective segment volume for the this segment. Currently this option is not supported by OPM Flow. cm³None 2-11 **XCORDS** A real positive values equal to or greater than zero that defines the coordinate in the x-direction of the nodal point of this segment that is used for display purposes only. Currently this option is not supported by OPM Flow. None cm 12-2 **YCORDS** A real positive value equal to or greater than zero that defines the coordinate in the y-direction of the nodal point of this top segment that is used for display purposes only. Currently this option is not supported by OPM Flow. None 2-13 **XAREAS** XAREAS is real positive value equal to or greater than zero that defines the cross sectional area of the pipe wall for this segment, that is used in thermal conductivity calculations for when the temperature calculation is activated by the TEMP keyword in the RUNSPEC section. Currently this option is not supported by OPM Flow. cm^{2} None 2-14 **VHEATSEG** VHEATCAP is real positive value equal to or greater than zero that defines the volumetric heat capacity of the pipe wall for this segment, that is used in thermal conductivity calculations for when the temperature calculation is activated by the TEMP keyword in the RUNSPEC section. Currently this option is not supported by OPM Flow. Btu/ft/day/°R kJ/m/day/K J/cm/hr/K None 2.15 **THCSEG** THCON is real positive value equal to or greater than zero that defines the thermal conductivity of the pipe wall for this segment, that is used in thermal conductivity calculations for when the temperature calculation is activated by the TEMP keyword in the RUNSPEC section. Currently this option is not supported by OPM Flow. Btu/ft/day/°R kJ/m/day/K J/cm/hr/K None 2-16 Record terminated by a "/" Not **Applicable**

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No.	Name		Description		Default
		Field	Metric	Laboratory	

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Notes:

- Each multi-segment wells must be defined by a separate WELSEGS keyword that consists of two records, with entries 1-1 to 1-12 representing record one items and 2-1 to 2-15 representing record number two items in the "No." column in this table.
- 2) Record number two of the keyword, items 2-1 to 2-15, is followed by up to MXCONS records as declared on the WSEGDIMS keyword in the RUNSPEC section.
- B) Each of the records are terminated by a "/" and is explicitly shown in the above rows and the keyword should be terminated by a "/".

Table 12.34:WELSEGS Keyword Description

The total number of wells should be defined via the WELLSDIMS keyword and the number of multi-segment wells should be declared on the WSEGDIMS keyword, both keywords are in the RUNSPEC section.

See also the WELSPECS keyword to define wells, the COMPDAT keyword to define the well completions for both ordinary wells and multi-segment wells, and the COMPSEGS keyword to define a multi-segment segment completions. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines one producing well segment oil well (OP01) using the WELSPECS, WELSEGS COMPDAT and COMPSEGS keywords, and one water injection well (WI01) using the WELSPECS and COMPDAT keywords.

WELL	SPE	CIFI	CATIO	N DATA									
										OPEN SHUT	CROSS FLOW	PVT TABLE	
PLAT													/
WELL	CON	NECT	ION D	ATA									
II	LOCA JJ	TION K1	K2	OPEN SHUT	SAT TAB	CONN FACT						DIR T PEN	
10 10	10 10	1 2	1 2	OPEN OPEN	1* 1*	200. 200.							/
10 10	10 10	3 4	3 4	OPEN OPEN	1* 1*	200. 200.	0.	4					/
10 10	10 10	5 6	5 6	OPEN OPEN	1* 1*	200. 200.							/
9	10 10	2	2	OPEN OPEN	1* 1*	200.							/
7 6	10 10	2	2	OPEN OPEN	1* 1*	200. 200.	0.	4					, /
5	10	2	2	OPEN	1*	200.							/
10 10	9 8 7	3	3	OPEN OPEN	1*	200.	0.	4					//
10 10 10	6 5	3	3 3	OPEN OPEN	1* 1*	200. 200.	0.	4					/
	GROUNAME PLAT PLAT WELL II 10 10 10 10 10 10 10 10 10 10 10 10 10	GROUP NAME PLATFORM PLATFORM WELL CON LOCA II JJ 10 10 10 10 10 10 10 10 10 10 10 10 10 1	GROUP LOONAME PLATFORM 1 PLATFORM WELL CONNECT LOCATION II JJ K1 10 10 1 10 10 2 10 10 3 10 10 4 10 10 5 10 10 6 9 10 2 8 10 2 7 10 2 6 10 2 5 10 2 10 9 3 10 8 3 10 7 3 10 6 3	GROUP NAME I PLATFORM 10 1 PLATFORM 1 1 WELL CONNECTION D LOCATION II JJ K1 K2 10 10 1 1 10 10 2 2 10 10 3 3 10 10 4 4 10 10 5 5 10 10 6 6 9 10 2 2 8 10 2 2 7 10 2 2 6 10 2 2 7 10 2 2 6 10 2 2 7 10 2 2 6 10 2 2 7 10 2 2 6 10 2 2 7 10 2 2 6 10 2 2 7 10 2 2 6 10 2 2 7 10 2 3 10 9 3 3 10 8 3 10 7 3 3 10 6 3	NAME I J DEPORT PLATFORM 10 10 1* PLATFORM 1 1 1 1* WELL CONNECTION DATA LOCATION OPEN II JJ K1 K2 SHUT 10 10 1 1 OPEN 10 10 2 2 OPEN 10 10 3 3 OPEN 10 10 4 4 OPEN 10 10 5 5 OPEN 10 10 6 6 OPEN 9 10 2 2 OPEN 10 10 6 6 OPEN 9 10 2 2 OPEN 7 10 2 2 OPEN 7 10 2 2 OPEN 6 10 2 2 OPEN 7 10 2 2 OPEN 6 10 2 2 OPEN 5 10 2 2 OPEN 10 9 3 3 OPEN 10 8 3 3 OPEN 10 7 3 3 OPEN 10 7 3 3 OPEN 10 6 3 3 OPEN	GROUP	GROUP	GROUP	GROUP LOCATION BHP PHASE DRAIN INFL EQUAL PLATFORM 10 10 1* OIL PLATFORM 1 1 1 1* WATER WELL CONNECTION DATA LOCATION OPEN SAT CONN WELL II JJ K1 K2 SHUT TAB FACT DIA 10 10 1 1 0 OPEN 1* 200. 0.5 10 10 2 2 OPEN 1* 200. 0.4 10 10 4 4 OPEN 1* 200. 0.4 10 10 5 5 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 9 10 2 2 OPEN 1* 200. 0.4 10 10 2 2 OPEN 1* 200. 0.4 10 10 5 5 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 7 3 3 OPEN 1* 200. 0.4 10 9 3 3 OPEN 1* 200. 0.4 10 7 3 3 OPEN 1* 200. 0.4	GROUP LOCATION BHP PHASE DRAIN INFLOW RAME I J DEPTH FLUID AREA EQUANS PLATFORM 10 10 1* OIL WATER WELL CONNECTION DATA LOCATION OPEN SAT CONN WELL KH II JJ K1 K2 SHUT TAB FACT DIA FACT 10 10 1 1 1 OPEN 1* 200. 0.5 10 10 2 2 OPEN 1* 200. 0.4 10 10 3 3 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 9 10 2 2 OPEN 1* 200. 0.4 10 10 5 5 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 7 10 2 2 OPEN 1* 200. 0.4 10 10 2 3 OPEN 1* 200. 0.4 10 10 3 3 OPEN 1* 200. 0.4 10 7 10 2 2 OPEN 1* 200. 0.4 10 7 3 3 OPEN 1* 200. 0.4	GROUP LOCATION BHP PHASE DRAIN INFLOW OPEN NAME I J DEPTH FLUID AREA EQUANS SHUT PLATFORM 10 10 1* OIL PLATFORM 1 1 1 1* WATER WELL CONNECTION DATA LOCATION OPEN SAT CONN WELL KH SKI II JJ K1 K2 SHUT TAB FACT DIA FACT FACT 10 10 1 1 1 OPEN 1* 200. 0.5 10 10 2 2 OPEN 1* 200. 0.5 10 10 3 3 OPEN 1* 200. 0.4 10 10 4 4 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 2 2 OPEN 1* 200. 0.4 10 10 2 2 OPEN 1* 200. 0.4 10 10 5 5 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 7 3 3 OPEN 1* 200. 0.4 10 9 3 3 OPEN 1* 200. 0.4 10 7 3 3 OPEN 1* 200. 0.4 10 7 3 3 OPEN 1* 200. 0.4 10 7 3 3 OPEN 1* 200. 0.4	GROUP LOCATION BHP PHASE DRAIN INFLOW OPEN CROSS SAME I J J DEPTH FLUID AREA EQUANS SHUT FLOW PLATFORM 10 10 1* OIL WATER WELL CONNECTION DATA LOCATION OPEN SAT CONN WELL KH SKIN DII JJ K1 K2 SHUT TAB FACT DIA FACT FACT FACT FACT 10 10 10 1 1 OPEN 1* 200. 0.5 10 10 2 2 OPEN 1* 200. 0.5 10 10 3 3 OPEN 1* 200. 0.4 10 10 6 6 OPEN 1* 200. 0.4 10 10 6 0 OPEN 1* 200. 0.4 10 10 0 OPEN 1* 200. 0.4 10 10 0 OPEN 1* 200. 0.4 10 10 0 OPEN 1* 200. 0.4 10 OPEN	CROUP CROSS PVT AREA CQUANS SHUT FLOW TABLE

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0P01	9	10 5	5	OPEN 1	* 200.	0.4				/
0P01	8	10 5		OPEN 1	* 200.	0.4				/
0P01	7	10 5	5	OPEN 1	* 200.	0.4				/
0P01	6	10 5	5	OPEN 1	* 200.	0.4				/
0P01	5	10 5		OPEN 1						/
	_			-						
0P01	10	9 6	6	OPEN 1	* 200.	0.4				/
0P01	10	8 6		OPEN 1						/
0P01	10	7 6		OPEN 1						,
0P01	10	6 6		OPEN 1						,
0P01	10	5 6		OPEN 1						,
OFUI	10	5 0	0	OFLN I	200.	0.4				,
WI01	1	1 7	9	OPEN 1	* 200.	0.5				/
/		1 /	9	OI LIN I	200.	0.5				,
	WELL	SEGMEN	IT SDEC	IFICATION	DATA					
	WLLL	SEGNEN	II SFLC	IFICATION	DATA					
WELL	NODA	ı	LEN	WELL	DEPH	PRESS	FLOW			
NAME	DEPT		TUBIN		OPTN	CALC	MODEL			
WELSEGS	DEPTI	П	LODIN	G VOLM	OPTN	CALC	MODEL			
	2512	-	2512	- 1 OF F	ABC	115	ш			,
0P01	2512	.5	2512.	5 1.0E-5	ABS	HFA	НО			/
	CEC	CEC	DDAN	CEC	TUDTNO	NODAL	TUDE	TUDE	VCEC	\/OI
	SEG	SEG	BRAN	SEG	TUBING	NODAL	TUBE	TUBE	XSEC	V0L
	ISTR	IEND	NO	NO	LENGTH	DEPTH	ID	ROUGH	AREA	SEG
	2	2	1	1	2537.5	2534.5	0.3	0.00010		/
	3	3	1	2	2562.5	2560.5	0.3	0.00010		/
	4	4	1	3	2587.5	2593.5	0.3	0.00010		/
	5	5	1	4	2612.5	2614.5	0.3	0.00010		/
	6	6	1	5	2637.5	2635.5	0.3	0.00010		/
	7	7	2	2	2737.5	2538.5	0.2	0.00010		/
	8	8	2	7	2937.5	2537.5	0.2	0.00010		/
	9	9	2	8	3137.5	2539.5	0.2	0.00010		/
	10	10	2	9	3337.5	2535.5	0.2	0.00010		/
	11	11	2	10	3537.5	2536.5	0.2	0.00010		/
	12	12	3	3	2762.5	2563.5	0.2	0.00010		/
	13	13	3	12	2962.5	2562.5	0.1	0.00010		/
	14	14	3	13	3162.5	2562.5	0.1	0.00010		/
	15	15	3	14	3362.5	2564.5	0.1	0.00010		/
	16	16	3	15	3562.5	2562.5	0.1	0.00010		/
	17	17	4	5	2812.5	2613.5	0.2	0.00010		/
	18	18	4	17	3012.5	2612.5	0.1	0.00010		/
	19	19	4	18	3212.5	2612.5	0.1	0.00010		/
	20	20	4	19	3412.5	2612.5	0.1	0.00010		,
	21	21	4	20	3612.5	2613.5	0.1	0.00010		
			•		3322.3	_0_0.0	0.2	0.00020		,
	22	22	5	6	2837.5	2634.5	0.2	0.00010		/
	23	23	5	22	3037.5	2637.5	0.2	0.00010		/
	24	24	5	23	3237.5	2638.5	0.2	0.00010		,
	25	25	5	24	3437.5	2639.5	0.1	0.00010		,
	26	26	5	25	3637.5	2639.5	0.1	0.00010		,
/	_0	_0	Ü		0007.10	_000.0	V. ±	0.00010		,
, 										
	COMP	IETTON	SEGMEN	T SPECIFI	CATTON DA	ΤΔ				
	COMP	CC 1 TON	JEUNEN	i Si LCIPI	CALLON DA	NIΛ				
WELL										
WELL NAME										
COMPSEGS										
0P01										/
0P01										/

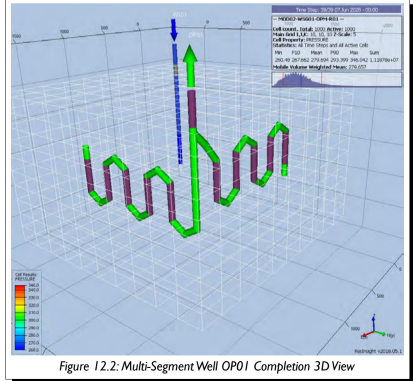
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	L0 II 10 10 10 10 10	JJ 10 10 10 10 10	ON K1 1 2 3 4 5	BRAN NO 1 1 1 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0	DIR PEN	LOC I,J,K	MID PERFS	COMP LENGTH	ISEG NO.	/ / / / / /
	9 8 7 6 5	10 10 10 10 10	2 2 2 2 2	2 2 2 2 2 2	2637.5 2837.5 3037.5 3237.5 3437.5	2650.0 2837.5 3037.5 3237.5 3437.5 3637.5						/ / / / /
	10 10 10 10 10	9 8 7 6 5	3 3 3 3	3 3 3 3	2662.5 2862.5 3062.5 3262.5 3462.5	2862.5 3062.5 3262.5 3462.5 3662.5						///////////////////////////////////////
	9 8 7 6 5	10 10 10 10 10	5 5 5 5	4 4 4 4	2712.5 2912.5 3112.5 3312.5 3512.5	2912.5 3112.5 3312.5 3512.5 3712.5						/ / /
,	10 10 10 10 10	9 8 7 6 5	6 6 6 6	5 5 5 5	2737.5 2937.5 3137.5 3337.5 3537.5	2937.5 3137.5 3337.5 3537.5 3737.5						/ / / /

Note the use of both the COMPDAT and COMPSEGS keywords to fully define a multi-segment well's completion.

Finally Figure 12.2 depicts the resulting well configuration for both wells.



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12.3.83 WELSPECL - Define Well Specifications for Local Grid Refinements

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WELSPECL keyword defines the general well specification data for all well types and must be used for all wells contained within a Local Grid Refinement ("LGR") instead of the WELSPECS keyword. WELSPECL must declare wells first before any other LGR well specification keywords are used in the input file. The keyword declares the name of well, the group the well belongs to, the LGR the well is incorporated into, the wellhead location and other key parameters.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

No.	Name			Default	
		Field	Metric	Laboratory	
I	WELNAME		to eight characters in leng connection data is being d		None
2	GRPNAME		o to eight characters in ne well is assigned to. The		None
		keyword when there is a	ierarchy should be defin more than one level of gr der the FIELD group in th	oups, otherwise all the	
3	LGRNAME		o to eight characters in inement for which the we		None
4	I	A positive integer greate NX on the CARFIN I wellhead location for a horizontal well in the I-di For radial LGRs this para	None		
5	J	A positive integer greate NY on the CARFIN keyw location for a vertical or the J-direction within the For radial LGRs this para	hat defines the wellhead for a horizontal well in	None	
6	BHPREF	A real value that defines hole pressure for the we of the perforations as SCHEDULE section.	be set to the midpoint DATL keyword in the	Mid-point of shallowest	
		If defaulted by I* or set mid-point of shallowest of will be used.		connection defined by the COMPDAT	
		feet	m	cm	keyword

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No.	Name		Description		Default		
		Field	Metric	Laboratory			
7	TYPE	A defined character that be set to one of the follo	for the well, and should	None			
		I) GAS: for a gas we	II.				
		2) OIL: for an oil we	II.				
		3) WAT: for a water	injection well.				
		4) LIQ: for an oil we for the well.	ell when the liquid produ	ctivity index is required			
		or injectivity index and the when a group's production keyword in the SCHEDU	the phase used to calculathe type of well, or a we on constraints, as defined LE section, have been viol I well, then excessive gas .	II's connection, to close d on the GCONPROD ated. For example, if the			
8	DRADIUS	A real value that define calculate a well's product	s the well drainage radiu	s for the well used to			
		A default of zero results in the pressure equivalent radius of the grid blocks containing the well connections are used.					
		feet	m	cm	0.0		
9	INFLOW	A defined character string that defines the inflow equation to be used for the well in calculating the well's flow rates. INFLOW should be set to one of the following character strings:					
		STD: the standard inflow equation will be used. This is normally used for wells that are primary oil or water wells.					
		2) NO: an alias for STD.					
		3) R-G: the Russell Goodrich ⁶⁸ pressure square inflow equation will used. This option can be used for dry gas wells.					
		4) YES: an alias for R	-G.				
			dry gas pseudo pressure sed for dry gas wells.	inflow equation will be			
		6) GPP: the generalized gas pseudo pressure inflow equation used with wet gas wells, that is condensate gas wells. This inflow equation is based on the formulation of Whitson et. al. 69					
		For oil and water wells the INFLOW should be set to STD, why for dry gas wells INFLOW can be set to either R-G or P-P; however, the P-P option is preferred for dry gas wells due to the more rigorous treatment of gas flow. For wet gas wells, that is gas condensate wells, INFLOW should be set to GPP.					
		Only INFLOW equal to Flow.	STD and NO are current	ly implemented in OPM			

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⁶⁹ Whitson, C. H. and Fevang, Ø. "Generalised Pseudopressure Well Treatment in Reservoir Simulation," Presented at the IBC Technical Services Conference on Optimisation of Gas Condensate Fields, Aberdeen, UK (June 26-27, 1997).

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Russell, D.G., Goodrich, J.H., Perry, G.E and Bruskotter, J.F "Methods of Predicting Gas Well Performance", Transactions of the ASME, Journal of Petroleum Technology (1966) 99-108.

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No.	Name	Description						
		Field	Metric	Laboratory				
10	AUTO	A defined character strin the economic WCUT, GC cease production. AUTO strings: 1) STOP: the well is fluids to surface; may occur within depending on a connections. Inte setting the XFLO will be similar to the setting the string t	SHUT					
		no flow at the sur	shut at the surface and face and no cross flow do es places at the end of the					
		constraint is violated.	es places at the end of the	s carrie scep in which the				
11	XFLOW	within the wellbore, and s 1) YES: to allow cros 2) NO: to disallow	s flow in the wellbore thr	ough well connections. ellbore, even if the flow	YES			
			issues can occur if this varesolve the issue; howevocess in this case.	-				
12	PVTNUM		than or equal to zero the lbore fluid properties that Irface volume rates.		0			
		The default value of zer deepest connection in the	ro sets PVTNUM to be e well.	the PVT table of the				
13	DENOPT		g that sets the type of de nydrostatic head, and shou is:		SEG			
		this cases the connections and the more accurate the well connections.	density is calculated be the volumes flowing from te calculation if the fluid	culation to segmented. In tween neighboring well in the connections. This is properties flowing from ensity calculation itself is last time step.				
		density calculation given reservoir ar	n. Here the density is cor	alculation to the average isidered uniform across a inflow rates of each phase				
		The default option of I* implemented in OPM Flo	invokes the SEG option w.	and is the only option				

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No.	Name	Description								
		Field Metric Laboratory								
14	FIPNUM	An integer value defines the FIPNUM region used to determine the reservoir conditions in calculating the well's reservoir volumes.								
		If set to a negative integer value then the FIPNUM region of the deepest connection in the well will be used.								
		If set to zero, the default will be used.	value, then the average	properties for the field						
		If set to an integer value this value will be used.	greater than zero, then th	ne FIPNUM indicated by						
15		Not used.								
16		Not used.								
17		Not used.								
18		Not used.								

Revision: Rev-I

Notes:

- 1) The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".
- 2) Note that the total number of individual wells declared by the WELSPECS cannot exceed the valued entered via the MXWELS variable on the WELLDIMS keyword in the RUNSPEC section.

Table 12.35:WELSPECL Keyword Description

See also the COMPDATL keyword to define a well's connections in a LGR, the WCONPROD and WCONDINJE keywords to define a well's production and injections targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines three wells using the WELSPECL keyword

```
-- WELL SPECIFICATION DATA FOR LGR WELLS
                                      BHP
                                                                             CROSS PVT
-- WELL
         GROUP
                   I GR
                           LOCATION
                                              PHASE DRATN
                                                             TNFI OW
                                                                      SHUT
                   NAME
                                      DEPTH
                                              FLUID
                                                                                   TABLE
-- NAME
         NAME
                                                      AREA
                                                              EQUA.
                                                                      IN
                                                                             FLOW
WELSPECL
         PLATFORM LGR01
                           14
                                      1*
                                              GAS
                                                      1*
                                                              P-P
                                                                     SHUT
                                                                             NO
                                                                                     1* /
GT01
                                 13
                                                                                     1* /
1* /
GP01
          PLATFORM LGR01
                                      1*
                                              GAS
                                                      1*
                                                              GPP
                                                                     SHUT
                                                                             NO
                           64
                                 80
                                      1*
                                                      1*
0P01
         PLATFORM LGR02
                           24
                                 10
                                              OIL
                                                              STD
                                                                     SHUT
                                                                             NO
```

Here, well Gl01 and GP01 are in the same LGR named LGR01 and OP01 is in a separate LGR named LGR02. Gl01 is a dry gas injection well that users the dry gas pseudo inflow equation, GP01 is a gas condensate well that users the generalized gas pseudo pressure inflow equation, and finally, OP01 is an oil well that users the standard inflow equation. All wells: will be shut if they are required to cease production, all wells disallow cross flow, and the hydrostatic head calculation is defaulted to the segment option for all wells.

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12.3.84 WELSPECS - Define Well Specifications

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WELSPECS keyword defines the general well specification data for all well types, and must be used for all wells before any other well specification keywords are used in the input file. The keyword declares the name of well, the wellhead location and other key parameters.

No.	Name		Description	Default					
		Field	Metric	Laboratory	-				
I	WELNAME		to eight characters in leng connection data is being d		None				
2	GRPNAME		o to eight characters in ne well is assigned to. The		None				
		keyword when there is	Note that the group hierarchy should be defined by the GRUPTREE keyword when there is more than one level of groups, otherwise all the groups will sit directly under the FIELD group in the group tree hierarchy.						
3	1	NX that defines the wel	A positive integer greater than or equal to zero and less than or equal to NX that defines the wellhead location for a vertical or deviated well, or the heel for a horizontal well in the I-direction.						
4	J	NY that defines the wel	A positive integer greater than or equal to zero and less than or equal to NY that defines the wellhead location for a vertical or deviated well, or the heel for a horizontal well in the J-direction						
5	BHPREF	A real value that defines hole pressure for the we of the perforations as SCHEDULE section.	Mid-point of shallowest						
			If defaulted by I* or set to a value less than or equal to zero, then the mid-point of shallowest connection defined by the COMPDAT keyword will be used.						
		feet	m	cm	COMPDAT keyword				
6	TYPE	A defined character that be set to one of the follo	defines the "main" phase wing character strings:	for the well, and should	None				
		I) GAS: for a gas we	II.						
		2) OIL: for an oil we							
		3) WAT: for a water injection well.							
		4) LIQ: for an oil was for the well.							
		This parameter defines or injectivity index and the when a group's production keyword in the SCHEDU well is declared as an oin will be subject to closure							

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No.	Name		Description		Default		
		Field	Metric	Laboratory			
7	DRADIUS	A real value that define calculate a well's product	s the well drainage radiu	s for the well used to			
		A default of zero result blocks containing the well	s in the pressure equivall connections are used.	lent radius of the grid			
		feet	m	cm	0.0		
8	INFLOW		g that defines the inflow of well's flow rates. INFLOV r strings:	•	STD		
			d inflow equation will be t are primary oil or water				
		2) NO: an alias for S					
			Goodrich ⁷⁰ pressure squ can be used for dry gas w				
		4) YES: an alias for R-G.					
		P-P: the general dry gas pseudo pressure inflow equation will be used. Normally used for dry gas wells.					
		6) GPP: the generalized gas pseudo pressure inflow equation used with wet gas wells, that is condensate gas wells. This inflow equation is based on the formulation of Whitson et. al. ⁷¹					
		For oil and water wells a gas wells INFLOW can option is preferred for d of gas flow. For wet gas should be set to GPP.	P-P; however, the P-P ore rigorous treatment				
		Only INFLOW equal to Flow.	STD and NO are current	y implemented in OPM			
9	AUTO	the economic WCUT, GO	g that defines the automa DR, or WGR limits are vic should be set to one of	lated and the well is to	SHUT		
		fluids to surface; may occur within depending on a co connections. Inte setting the XFLO	"stopped" at the surface however, if there any ope the wellbore and betweennection's potential with r-connection flow (cross f W variable to NO. In this the SHUT option describe	n connections then flow en the open connections respect to all the other low) can be prevented by a case the well's behavior			
			s shut at the surface and face and no cross flow do				
		The corrective action tak constraint is violated.	es places at the end of the	e time step in which the			

Revision: Rev-I

Whitson, C. H. and Fevang, Ø. "Generalised Pseudopressure Well Treatment in Reservoir Simulation," Presented at the IBC Technical Services Conference on Optimisation of Gas Condensate Fields, Aberdeen, UK (June 26-27, 1997).

Α	В	С	D	Ε	F	G	Н	Κ	J	Κ	L	М	N	0	Р	Q	R	S	Τ	U	V	W	Χ	Υ	Ζ
Date	e: No	veml	ber 6	, 201	8						Tab	le of (Conte	ents								Pa	ge 70	09 of	792

⁷⁰ Russell, D.G., Goodrich, J.H., Perry, G.E and Bruskotter, J.F "Methods of Predicting Gas Well Performance", Transactions of the ASME, Journal of Petroleum Technology (1966) 99-108.

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No.	Name		Description		Default
		Field	Metric	Laboratory	
10	XFLOW	A defined character stri within the wellbore, and	ing that defines the if cr should be set to either:	oss flow should occur	YES
		I) YES: to allow cros	ss flow in the wellbore thr	ough well connections.	
			cross flow within the w well connections would all	rellbore, even if the flow ow such flow to occur.	
			issues can occur if this varesolve the issue; howevocess in this case.	· ·	
11	used to calculate the wellbore fluid properties that define the relationship between reservoir and surface volume rates.				
		The default value of ze deepest connection in the	ro sets PVTNUM to be e well.	the PVT table of the	
12	DENOPT		g that sets the type of de nydrostatic head, and shou ss:		SEG
		this cases the connections and the more accurathe well connections	density is calculated be the volumes flowing from		
		density calculation given reservoir ar		alculation to the average isidered uniform across a inflow rates of each phase	
		The default option of I* implemented in OPM Flo	invokes the SEG option w.	and is the only option	
13	FIPNUM		es the FIPNUM region unalculating the well's reserve		0
		If set to a negative integer connection in the well with	er value then the FIPNUN ill be used.	1 region of the deepest	
		If set to zero, the defaulwill be used.	t value, then the average	properties for the field	
		If set to an integer value this value will be used.	greater than zero, then th	ne FIPNUM indicated by	
14		Not used.			
15		Not used.			
16		Not used.			
17		Not used.			

Revision: Rev-I

Notes:

- The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".
- Note that the total number of individual wells declared by the WELSPECS cannot exceed the valued entered 2) via the MXWELS variable on the WELLDIMS keyword in the RUNSPEC section.

Table 12.36:WELSPECS Keyword Description

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See also the COMPDAT keyword to define a well's connections, the WCONPROD and WCONDINJE keywords to define a well's production and injections targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Revision: Rev-I

Example

The following example defines three wells using the WELSPECS keyword

```
WELL SPECIFICATION DATA
- -
         GROUP
                   LOCATION
                              RHP
                                     PHASE
                                            DRAIN
                                                    INFLOW
                                                            OPEN
                                                                  CROSS
                                                                         PVT
-- WELL
                                     FLUID
                                            AREA
                                                                          TABLE
-- NAME
         NAME
                     Т Л
                              DEPTH
                                                    EQUANS
                                                            SHUT
                                                                  FLOW
WELSPECS
         PLATFORM
                               1*
                                             1*
                                                     P-P
                                                            SHUT
                                                                           1*
                    14
                         13
                                      GAS
GI01
                                                                   NO
                               1*
                                             1*
                                                                           1*
GP01
         PLATFORM
                    64
                         80
                                      GAS
                                                     GPP
                                                            SHUT
                                                                   NO
                               1*
                                             1*
0P01
         PLATFORM
                    24
                        110
                                      OIL
                                                     STD
                                                            SHUT
                                                                   NO
```

Here, well Gl01 is a dry gas injection well that users the dry gas pseudo inflow equation, GP01 is a gas condensate well that users the generalized gas pseudo pressure inflow equation, and finally, OP01 is an oil well that users the standard inflow equation. All wells: will be shut if they are required to cease production, all wells disallow cross flow, and the hydrostatic head calculation is defaulted to the segment option for all wells.

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12.3.85 WELTARG - Modify Well Targets and Constraints Values

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WELTARG keyword modifies the target and constraints values of both rates and pressures for previously defined wells without having to define all the variables on the well control keywords: WCONPROD, WCONHIST, WCONINJE, or WCONINJH keywords. Variables not changed by the WELTARG keyword remain the same as those previously entered via the well control keywords or previously entered WELTARG keywords. Note that the well must still be initially be fully defined using the WCONPROD or WCONINJE keywords.

All the aforementioned keywords are described in the SCHEDULE section.

No.	Name	Description							
		Field	Metric	Laboratory					
I	WELNAME		to eight characters in leng I production rates and pi		None				
		previously using the W	Note that the well name (WELNAME) must have been declared previously using the WELSPECS and WCONPROD (or WCONINJE) keywords in the SCHEDULE section, otherwise an error may occur.						
2	TARGET	A defined character strir the value of the item is se	ng that sets the item to bet by item (3).	e changed for the well	None				
		I) ORAT: reset the item (3).	surface oil production r	ate value as defined by					
		 WRAT: reset the surface water production rate value as defined by item (3). GRAT: reset the surface gas production rate value as defined by item (3). LRAT: reset the surface liquid (oil plus water) production rate value as defined by (3). 							
		5) RESV: reset he in	situ reservoir volume rate	e value as defined by (3).					
		6) BHP: reset the bo	ttom-hole pressure value	as defined by item (3).					
		7) THP: reset the tubing head pressure value for the well as defined by item (3).		for the well as defined					
		8) VFP: reset the ver (3).	rtical lift performance tab	e number as defined by					
		9) LIFT: reset the performance table	artificial lift quantity for es.	use with vertical lift					
		10) GUID: reset the control.	guide rate value for wells	operating under group					
		how a well is controlled control, as defined by the entering TARGET equal constraint but the well st	es the variable to be chard. For example, if a well the previously entered V to LRAT with a valuill remains on ORAT cont LE section to change the	is operating on ORAT VCONPROD keyword, i.e, changes the liquid rol. Use the WELCNTL					

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No.	Name	Description					
		Field	Metric	Laboratory			
3 VALUE		A real positive value the TARGET	A real positive value that defines the value of the variable declared by TARGET				
	Liquid	stb/d	sm³/day	scc/hour			
	Gas	Mscf/d	sm³/day	scc/hour			
	Res Vol	rb/d	rm³/day	rcc/hour			
	Pressure	psia	barsa	atma			
	VFP	dimensionless	dimensionless	dimensionless			
	LIFT	same as	same as	same as			
		VFPPROD or VFPINJ	VFPPROD or VFPINJ	VFPPROD or VFPINJ	None		

Revision: Rev-I

Notes:

1) The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.37:WELTARG Keyword Description

If a well is currently a history matching well, then WELTARG should only be used to change a wells bottom-hole pressure limit, vertical flow performance table number or the artificial lift quantity.

See also the WELCNTL keyword, in the SCHEDULE section that can be used to reset the control mode, as well as a well's target and constraints of both rates and pressures.

Example

The following example below shows the oil rates for the OP01 oil producer at the start of the schedule section (January I, 2000).

```
-- 01 JAN 2000 START OF SCHEDULE SECTION
        WELL PRODUCTION WELL CONTROLS
                                                                      VFP
-- WELL OPEN/
               CNTL
                       OIL
                              WAT
                                     GAS
                                           LIO
                                                  RES
                                                         BHP
                                                                THP
                                                                             VFP
               MODE
                                    RATE RATE
-- NAME SHUT
                       RATE
                              RATE
                                                  RATE
                                                         PRES
                                                                PRES TABLE
                                                                            ALFQ
WCONPROD
               ORAT
                              1*
                                     1*
                                           1*
                                                  1*
                                                                             1* /
0P01
        OPEN
                       3000
                                                         750.0 500.
DATES
01 FEB 2000 /
        WELL PRODUCTION AND INJECTION TARGETS
   WELL WELL
                TARGET
   NAME TARG
                VALUE
- -
WELTARG
                2000
0P01
         ORAT
```

From January I, 2000 to February I, 2000 well OP0I is open and is on oil rate control and has a target oil rate of 3,000 stb/d, and uses VFPPROD vertical lift table number 9 with a minimum tubing head pressure constraint of 500 psia. After February I, 2000 the well's oil rate is reduced to 2,000 stb/d and all the other parameters remain unchanged.

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12.3.86 WGASPROD - Define Sale Gas Well Production Targets

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

WGASPROD keyword declares wells to be Sales Gas producers and sets the incremental gas rate for a well and the maximum number of increments that this rate can be increased. Wells must have been previously been defined via the WELSPECS and WCONPROD keywords in the SCHEDULE section and are subject to any targets or constraints on WCONPROD keyword.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.87 WGRUPCON - Define Well Guides for Group Control

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WGRUPCON keyword defines a well's production or injection guide rate for when a well is under group control. The guide rate is used to determine a well's production target under group control in order to satisfy a group's targets and constraints, including any higher level related groups as well as the FIELD group.

Wells must have been previously defined and allocated to a group by the WELSPECS keyword in the SCHEDULE section. Wells defined to be under group control will have their production rates controlled by the group to which they belong, in addition to any well constraints defined for the wells via the WCONPROD and WCONINJE keywords in the SCHEDULE section.

No.	Name		Default			
		Field	Metric	Laboratory		
I	WELNAME	A character string of up name for which the well defined.	None			
		Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.				
2	STATUS	A defined character string that declares the status of the well to be under group control or not under group control. STATUS should be set to one of the following character strings:				
		I) YES: the well is u will be influenced level groups as we				
		NO: the well is NOT under group control and its production behavior will only be influenced by its own targets and constraints.				
		specified otherwise by the	of YES puts all wells unden the STATUS variable, or the CONINJE keywords in the	ne TARGET variable on		
3 GUIDERAT	GUIDERAT	A dimensionless real nun production (or injection)	nber that determines the target rate.	well's share of it's group		
		If GUIDERAT is a positive number then the guide rate for the well is fixed until modified by this keyword at a subsequent time. If TARGET variable on this keyword is not equal to the group's controlling phase, then the GUIDERAT is converted into the groups' controlling phase and is updated every time step.				
		If GUIDERAT is less that based on the well's po calculated every time ste				
		dimensionless	dimensionless	dimensionless	-1.0	

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No.	Name	Description				
		Field	Metric	Laboratory		
4	TARGET	A defined character string that sets the well's guide rate phase that the GUIDERAT value should be applied to. TARGET should be set to one of the following character strings:				
		OIL: the well's guide rate applies to the surface oil production rate.				
		WAT: the well's guide rate applies to the surface water production rate.				
		GAS: the well's guide rate applies to the surface gas production rate.				
		4) LRAT: the well's guide rate applies to the surface liquid (oil plus water) production rate.				
		5) RES: the well's guide rate applies to the in situ reservoir volume rate.				
		6) RAT: the well's guide rate applies to the injection phase. This should only be used if the well has been declared an injection via the WCONINJE keyword in the SCHEDULE section.				
5	SCALE A real value that is used to multiple the GUIDERAT or the calculated well potentials to determine the final GUIDERAT for the well.					
		dimensionless	dimensionless	dimensionless	1.0	

Revision: Rev-I

Notes:

Table 12.38:WGRUPCON Keyword Description

See also the GCONPROD the GCONINJ keywords to define a group's production and injection targets and constraints, and the WCONPROD and WCONINJE keyword to define a well's production and injection characteristics.. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines the guides rates all oil and gas producers and the gas injectors as follows:

```
DEFINE WELL GUIDES FOR GROUP CONTROL
-- WELL
        GRUP GUIDE GUIDE
                            SCALE
-- NAME
        CNTL RATE
                     PHASE
                            FACT
WGRUPCON
'GI*'
         YES
              0
                     RAT
                            1.0
'GP*'
        YES
            0
                     GAS
                             1.0
'0P*'
                     OIL
                             1.0
```

Both the gas producers ('GP*') and injectors ('GI'*) are under group control with their guide rates based on their potentials. The gas injector wells are controlled based on their reservoir potential volumes and the producers on their potential gas rates. In comparison, all the oil wells controlled by their oil rates.

The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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12.3.88 WHISTCTL - Define Well Historical Target Phase

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WHISCTL keyword changes the target control for wells declared as history match wells via the WCONHIST keyword in the SCHEDULE section. The target phase is set on the WCONHIST keyword and WHISCTL overrides this value for all subsequent entries on the WCONHIST keyword.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
I	TARGET	A defined character strin for the well, all the other reporting only. The simulathe phase rate stated in keyword.	phases are calculated uncator will attempt to meet	onstrained and used for the TARGET based on	None	
		TARGET should be set to	one of the following cha	racter strings:		
			t is set to the surface b) on the WCONHIST key			
		 2) WRAT: the target is set to the surface water production rate as defined by item (5) on the WCONHIST keyword. 3) GRAT: the target is set to the surface gas production rate as defined by item (6) on the WCONHIST keyword. 				
			is set to the surface nd is calculated by the sin ST keyword.			
		,	is set to the in situ reser simulator using items (word.			
		,	ate is set to the bottom-line WCONHIST keyword.	nole pressure as defined		
		7) NONE: revert b WCONHIST key	oack to the TARGET word.	control mode on the		
		The TARGET control moc control mode on the W from the time the WHI control model on all subs	CONHIST keyword in the STCNTL is invoked, thus	ne SCHEDULE section, savoiding changing the		
2	END	A defined character string the well has switch to Bh one of the following character	HP control by the simulate		NO	
		I) NO: no action is t	aken and the run continu	es.		
		2) YES: terminate the	e run at the next report ti	me step.		
		Wells set to BHP control ignored. Only END equal				

The keyword is terminated by a "/".

Table 12.39:WHISTCTL Keyword Description

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keywords in the SCHEDULE section.

History matching wells are handled differently then ordinary wells that use the WCONPROD keyword for controlling their production targets and constraints. However, the wells still need to be defined like ordinary production wells using the WELSPECS keyword in the SCHEDULE section. History matching well are converted to ordinary wells by restating a well's control mode using either the WCONPROD or WELTARG

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See also the WCONHIST and WCONINJH keywords that are used to define the historical production and injection data, All the aforementioned keywords are described in the SCHEDULE section.

Example

The example below shows the observed gas rates for the OP01 oil producer for the first quarter of 2000.

```
01 JAN 2000 START OF SCHEDULE SECTION
   ______
         DEFINE WELL HISTORICAL TARGET PHASE
- -
         CNTL
                 BHP
         MODE
                 ST<sub>0</sub>P
WHISTCTL
         RESV
                 NO
         WELL HISTORICAL PRODUCTION CONTROLS
- -
-- WELL
         OPEN/
                CNTL
                        OIL
                               WAT
                                      GAS
                                              VFP
                                                     VFP
                                                           THP
                                                                  RHP
-- NAME
         SHUT
                MODE
                        RATE
                               RATE
                                      RATE
                                              TABLE
                                                     ALFQ
                                                           PRES
                                                                 PRFS
WCONHIST
                                                      1*
         OPFN
                ORAT 15.5E3 100.0 1550
                                                           900.0 1*
0P01
                                              10
DATES
01 FEB 2000 /
- -
         WELL HISTORICAL PRODUCTION CONTROLS
- -
         OPEN/
                CNTL
                                              VFP
                                                     VFP
-- WELL
                        OIL
                               WAT
                                      GAS
                                                           THP
                                                                  BHP
-- NAME
                MODE
                        RATE
                               RATE
                                      RATE
                                                                 PRES
         SHUT
                                              TABLE
                                                     ALFQ
                                                           PRES
WCONHIST
                                              1*
                ORAT 15.2E3 150.0
                                                      1*
                                                           875.0 3250.0 /
0P01
         OPEN
                                     1520
DATES
01 MAR 2000 /
         WELL HISTORICAL PRODUCTION CONTROLS
- -
         OPEN/
                CNTL
                                              VED
                                                     VFP
                                                           THP
                                                                  RHP
-- WELL
                        OTI
                               WAT
                                      GAS
                                      RATE
-- NAME
         SHUT
                MODE
                        RATE
                               RATE
                                              TABLE
                                                     ALFQ
                                                           PRES
                                                                 PRES
WCONHIST
                              200.0 1500
                                              1*
                                                      1*
                                                           850.0 1*
         OPEN
                ORAT 15.0E3
0P01
```

From January I, 2000 the WCONHIST keyword defines well OP01, which is open and is on oil rate control, to produce 15,500 stb/d oil, with the observed rates of 100 stb/d of water and 15.5 MMscf/d of gas. However the WHISCTL keyword resets the target control to reservoir voidage from January I, 2000 and onward. This is useful in initial history matching runs to get a "reasonable" pressure match, by ensuring that the total reservoir withdrawals are correct, although the individual phase withdrawals will not match. Once a reasonable pressure match is achieved for the reservoir then one can reset TARGET to the sales phase, OIL or GAS, and continue with the matching of all the phases.

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12.3.89 WINJMULT - Define Well Pressure Dependent Injectivity Multipliers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WINJMULT keyword defines pressure dependent injectivity multipliers for injection wells and can be used to approximate the increase or decrease in a well's injectivity due to hydraulic fracturing in water injection wells.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.90 WINJTEMP - Define Injection Fluid Thermal Properties

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

WINJTEMP defines the injection fluid thermal properties for when the thermal option has been activated by the THERMAL keywords in the RUNSPEC. Only water and gas injection is supported.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	WELNAME		to eight characters in lengell injection fluid therma		None		
			ame (WELNAME) mus ELSPECS keyword in thoccur.				
2	STEAMQAL		itive value greater than o defines the steam quality				
		This parameter should be defaulted using I* as STEAMQUAL is not used by OPM FLOW, as only water and gas injection is supported.					
		This data is used by the commercial simulator's THERMAL option and is not supported by OPM Flow's THERMAL option.					
		dimensionless	dimensionless	dimensionless	I *		
3	TEMP	TEMP is a real positive v fluid for the defined well.	alue that defines the tem	perature of the injected			
		°F	°C	°C	None		
4	PRES	PRES is a real positive va	lue that defines the press	ure of the injected fluid			
		psia	barsa	atma	None		
5	ENTHALPY	ENTHALPY is a real pos		e specific enthalpy of the			
		This is data is used by the not supported by OPM F	e commercial simulator's ⁻ flow's THERMAL option.	THERMAL option and is			
		Btu/lbs-M	kJ/kg-M	J/gm-M	None		

Notes:

Table 12.40:WINJTEMP Keyword Description

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¹⁾ The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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Example

The following example shows the WINJTEMP keyword for when OPM Flow's temperature option has been activated by the THERMAL keyword in the RUNSPEC section.

Revision: Rev-I

```
INJECTION FLUID THERMAL PROPERTIES
- -
-- WELL
         STEAM
                INJ
                        INJ
                               SPEC
-- NAME
         QUAL
                 TEMP
                        PRES
                               ENTH
WINJTEMP
         1*
WI01
                 68.0
                        220.0
         1*
                                 1*
WI02
                 70.0
                        230.0
```

Here the water injection fluid's temperature and pressure, in field units, for two water injections well are defined. Notice that both the steam quality and the specific enthalpy of the injected fluid for the defined wells are defaulted (or skipped), as OPM Flow's THERMAL option does not support this data.

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12.3.91 WLIFT - Define Well Re-Tubing, THP and Lift Switching Workover Operations

F	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The WLIFT defines the automatic workovers parameters for changing out wellbore tubing, changing the THP limit (for example switching from the high stage pressure separator to the low stage pressure separator), or changing the artificial lift parameters, for wells.

OPM Flow does not have this feature and hence this keyword is ignored by OPM Flow and the WLIFT keyword has no effect on the simulation.

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12.3.92 WLIMTOL - Define Constraint Tolerance

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

WLIMTOL keyword defines the tolerance to be used for various constraints applied to connections, completions (if connections have been lumped via the COMPLUMP keyword in the SCHEDULE section), wells, and groups, including the field group.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

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12.3.93 WORKLIM - DEFINE WELL WORKOVER TIME

	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
--	---------	------	------	-------	---------	----------	---------	----------	--

Revision: Rev-I

Description

WORKLIM sets the numbers of days taken to complete a workover.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.94 WORKTHP - Define Well Workover Options for THP Killed Wells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WORKTHP keyword defines workover options for when a well dies, that is unable to produce at the current operating conditions, when under tubing head pressure ("THP") control. For example, if a well is producing to the high pressure separator and therefore has a high THP constraint, then the WORKTHP keyword can be used to switch the well to the lower pressure separator via re-setting the THP constraint.

This keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.95 WPAVE - Define Well Block Average Pressure Calculation Parameters

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
	_						

Revision: Rev-I

Description

The WPAVE keyword defines the method and parameters for calculating a well's block average pressures for all wells in the model. The resulting average pressure can be written out to the summary file in order to compared with field observed data.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	WPAVEI		ue that defines the weight ounding blocks used to ca pressures.		0.5			
		then the average pressur A value of zero indicates	n or equal to zero and les es are calculate based on s only the surrounding bl e of one indicates only the	the connection factors. ocks should be used in				
		•	If WPAVEI is equal to zero, then average pressure is calculate based or the pore volumes of the blocks.					
2	WPAVE2	equal to one, that define	ue greater than or equal t es the weighting factor b ures and the pore volu	etween the connection	1.0			
			ie, then the average pressi n factor calculated pressur					
		If WPAVE2 is equal to ze only using the pore volun	ro, then average pressures nes calculated pressures.	s are calculate based on				
3	WPAVE3	calculation is performed depth on the WELSPEC	A defined character string that determines how the hydrostatic head calculation is performed in correcting the pressures to the BHP reference depth on the WELSPECS or WPAVEDEP keywords in the RUNSPEC section. WPAVE3 should be set to one of the following character strings:					
		,	static head is calculated user at the well connections	,				
			ic head is calculated using with well connections					
		3) NONE: no hydros	static correction is applied	I to the pressures.				
4	WPAVE4		ng that determines which WPAVE4 should be set t		OPEN			
		used in the cal	connections and associate culations. This option n connections are opened	nay result in pressure				
		associated grid b discontinuities iss	ly defined open and clocks are used in the casue mentioned above caling all the well connectun.	lculations. The pressure n be avoided with this				

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No.	Name		Default		
		Field	Metric	Laboratory	
Notes:					
1)	The keyword sho	ould be terminated by a "/"			

Revision: Rev-I

Table 12.41:WPAVE Keyword Description

See also the WELSPECS keyword that defines a well and a well's bottom-hole pressure reference depth, the WPAVEDEP keyword that also defines a well's bottom-hole pressure reference depth, and the COMPDAT keyword to define a well's connections. All the aforementioned keywords are described in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.

Example

The following example defines the default well block average pressure calculation parameters

```
DEFINE WELL BLOCK AVERAGE PRESSURE CALCULATION PARAMETERS

INNER PORV WELL OPEN
OUTER CONN RES ALL

WPAVE

0.5 1.0 WELL ALL

And the next example shows the parameters used in the Norne model.
```

```
DEFINE WELL BLOCK AVERAGE PRESSURE CALCULATION PARAMETERS

INNER PORV WELL OPEN
OUTER CONN RES ALL

WPAVE

1* 0.0 WELL ALL
```

Here only pore volume weighting is used instead of connection weighting.

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12.3.96 WPAVEDEP - Define Well Reference Depth for Pressure Calculations

1.10.10.10.10.10.10.10.10.10.10.10.10.10	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
------------------------------------------	---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The WPAVEDEP keyword defines the reference depth to be used to calculate and report grid block average bottom hole pressures for a well. This keyword can be used to override the values entered or defaulted on the WELPSPECS keyword in the SCHEDULE section. The simulator corrects the grid block calculated pressures to a well's reference depth using the hydrostatic well of the producing fluids.

No.	Name		Description		Default				
		Field Metric Laboratory							
I	WELNAME	A character string of up name for which the well a Note that the well n previously using the Wotherwise an error may contact the string of the well of the well and the well of the well and the well an	None						
2	BHPREF	hole pressure for the we	A real value that defines the reference depth for reporting the bottom hole pressure for the well. Ideally this value should be set to the midpoint of the perforations as defined by the COMPDAT keyword in the SCHEDULE section.						
		If defaulted by I* or set to a value less than or equal to zero, then the mid-point of shallowest connection defined by the COMPDAT keyword will be used.							
		feet	m	cm	keyword				

Notes:

Table 12.42:WPAVDEP Keyword Description

See also the WELSPECS keyword that defines a well, the COMPDAT keyword to define a well's connections, and the WPAVE for defining how the average bottom hole pressure should be calculated. All the aforementioned keywords are described in the SCHEDULE section.

Note

The keyword is normally used to reset a well's bottom hole pressure depth to match the pressure gauge depth for when observed pressure is available, for example when conducting a history match for a well test, or when attempting to match static bottom hole surveys conducted on a well.

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The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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Example

The following example illustrates how to set the bottom hole reference depth for wells completed in different reservoirs that have different datum depths. Here it is assumed that all wells in a reservoir A have RES-A as part of their well name, and similarly for reservoirs B and C.

Revision: Rev-I

```
-- WELL SPECIFICATION DATA
-- WELL
             GROUP
                        LOCATION
                                   BHP
                                           PHASE
                                                  DRAIN
                                                          INFLOW
                                                                   SHUT
                                                                         CROSS
                                                                                 PRESS
-- NAME
             NAME
                         Ι
                               J
                                   DEPTH
                                           FLUID
                                                  AREA
                                                          EQUA.
                                                                   IN
                                                                         FLOW
                                                                                 TABLE
WELSPECS
                                                                                 1*
          PLATFORM
                              13
                                            OIL
                                                  1*
                                                                   OPEN
                                                                          NO
RES-A0P1
                                                  1*
                                                                                 1*
                                   1*
RES-A0P2
          PLATFORM
                        17
                              16
                                            OIL
                                                          STD
                                                                   OPEN
                                                                          NO
                                            OIL
                              19
                                   1*
                                                  1*
                                                                                 1*
RES-A0P3
          PLATFORM
                        21
                                                          STD
                                                                   OPEN
                                                                          NO
                                                                                 1*
RES-B0P4
                              96
                                   1*
                                                  1*
                                                                   OPEN
          PLATFORM
                        28
                                            OIL
                                                          STD
                                                                          NO
                                   1*
                                                  1*
                                                                                 1*
                                                                   OPEN
RES-B0P5
          PLATFORM
                              89
                        34
                                            OIL
                                                          STD
                                                                          NO
                                   1*
                                                  1*
                                                                                 1*
RES-COP6
          PLATFORM
                       128
                              52
                                            OTI
                                                          STD
                                                                   OPEN
                                                                          NO
                                   1*
                                                  1*
                                                                                 1*
RES-COP7
          PLATFORM
                       134
                              56
                                            OIL
                                                          STD
                                                                   OPEN
                                                                          NO
                                                                                 1*
          PLATFORM
                                   1*
                                                  1*
                                                                          NO
RES-COP8
                       138
                              50
                                            OIL
                                                          STD
                                                                   OPEN
RES-COP9
          PLATFORM
                       120
                              52
                                   1*
                                            OIL
                                                  1*
                                                          STD
                                                                   OPEN
                                                                          NO
         DEFINE WELL REFERENCE DEPTH FOR PRESSURE CALCULATIONS
         REF
-- WELL
  NAME
         DEPTH
WPAVEDEP
'RES-A*' 3100.0
'RES-B*' 3300.0
'RES-C*' 5909.0
```

In the example the all wells dedicated to RES-A will have their bottom hole reference depth set to 3,000 ft. TVDSS, RES-B wells to 3,300 ft. TVDSS and well RES-C wells to 5909 ft. TVDSS.

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12.3.97 WPIMULT - Define Well Connection Multipliers

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WPIMULT keyword defines a well connection multiplier factor that scales the existing well connection values. The resulting effect is scale the well' productivity at the reporting time step the keyword is entered.

No.	Name		Description		Default	
		Field	Metric	Laboratory		
I	WELNAME	A character string of up name for which the well a			None	
		Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.				
2	PIMULT	A real positive value that will be used to scale the well connection factors defined by I, J, K, K I and K2 below.				
3	1	An integer less than or equal to NX that defines the connection location in the I-direction.				
4	J	An integer less than or equal to NY that defines the connection location in the J-direction.				
5	К	An integer less than or equal to NZ that defines the connection location in the K-direction.				
6	KI	An integer less than or ellocation in the K-direction	•	the UPPER connection	*	
		If connections have been keyword, then K1 refer connection (layer) value.	n lumped into completion rs to the completion r			
7	K2	An integer less than or e location in the K-directio		the LOWER connection	*	
		If connections have been keyword, then K2 refer connection (layer) value.				

Notes:

Table 12.43:WPIMULT Keyword Description

If variables I, J K, KI and K2 are all defaulted with zero or I* then PIMULT is applied to all the well connections in the well. If variables I, J K, KI and K2 are set to zero (meaning any or all values), or a positive value then PIMULT is applied to the defined connections. The defined connections are those with the I, I, K variables in the specified location and a completion number in the range specified by K1 and K2.

Note that PIMULT variable is applied at the time the WPIMULT keyword is entered and is cumulative if there are intervening time steps between consecutive WPIMULT keywords.

See also the PIMULTAB keyword that defines productivity index multiplier versus water cut tables that are used to scaled a well's connection factors based on a wells connection current producing water cut. The keyword is documented in the SCHEDULE section.

The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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Revision: Rev-I

Examples

The following example defines three vertical oil wells using the WELSPECS keyword and their associated connection data.

```
-- WELL SPECIFICATION DATA
-- WELL
             GROUP
                         LOCATION
                                    BHP
                                            PHASE
                                                    DRAIN
                                                            INFLOW
                                                                     SHUT
                                                                            CROSS
                                                                                    PRESS
-- NAME
             NAME
                          Ι
                                J
                                     DEPTH
                                            FLUID
                                                    AREA
                                                            EQUA.
                                                                     ΙN
                                                                            FLOW
                                                                                    TABLE
WELSPECS
                                                    1*
                                                                                    1*
0P01
           PLATFORM
                         14
                               13
                                     1*
                                              OIL
                                                            STD
                                                                     OPEN
                                                                             NO
                                    1*
                                                                                   1*
                                                    1*
0P02
           PLATFORM
                         28
                               96
                                             OIL
                                                            STD
                                                                     OPFN
                                                                             NO
                                                                                        /
                                                                                    1*
0P03
           PLATFORM
                        128
                               56
                                     1*
                                             OIL
                                                    1*
                                                            STD
                                                                     OPEN
                                                                             NO
          WELL PRODUCTION WELL CONTROLS
                                                                BHP
                                                                              VFP
                                                                                      VFP
-- WELL
          OPEN/
                 CNTL
                         OTI
                                 WAT
                                         GAS
                                                        RES
                                                                       THP
                                                LIQ
-- NAME
          SHUT
                 MODE
                         RATE
                                 RATE
                                         RATE
                                                RATE
                                                        RATE
                                                                PRES
                                                                       PRES
                                                                              TABLE
                                                                                      ALF0
WCONPROD
                         1*
                                         1*
                                                1*
          SHUT
                 OPEN
                                 1*
                                                        1*
                                                                200.0
- -
          WELL CONNECTION DATA
-- WELL
          --- LOCATION ---
                              OPEN
                                      SAT
                                            CONN
                                                    WELL
                                                            KΗ
                                                                   SKTN
                                                                           D
                                                                                  DIR
           II
-- NAME
              JJ K1
                        K2
                              SHUT
                                      TAB
                                            FACT
                                                    DIA
                                                            FACT
                                                                   FACT
                                                                           FACT
                                                                                  PEN
COMPDAT
           1*
                              OPEN
                                      1*
                                                   0.708
                                                            1*
                                                                           1*
                                                                                  'Z'
0P01
               1 *
                        10
                                            1 *
                                                                   0.0
                     1
                                                                                  'Z' /
           1*
                                      1*
0P01
               1*
                    15
                        30
                              OPEN
                                            1*
                                                   0.708
                                                            1*
                                                                   0.0
                                                                           1*
           1*
               1*
                                     1*
                                            1*
                                                                          1*
                                                            1*
                    35
0P01
                              OPEN
                                                   0.708
                        90
                                                                   0.0
                                                                                  'Z' /
           1*
                                      1*
                                            1*
                                                            1*
                                                                           1*
0P02
               1*
                              OPEN
                                                   0.708
                                                                   0.0
                     1
                        10
0P03
                        90
                              OPEN
                                                   0.708
                                                                   0.0
          DEFINE WELL CONNECTION MULTIPLIERS
- -
                  --LOCATION--
                                 COMPLETION
-- WELL
          PΙ
-- NAME
          MULT
                    Ι
                        J
                              K
                                 FIRST LAST
WPIMULT
0P01
          1.250
                    1*
                        1*
                              1*
                                   1*
                                          1*
                        1*
                                   1*
                              1*
0P02
                    1*
          0.750
                                          10
0P03
                        1*
                              1*
          1.100
                                   35
                                          50
```

In this example the WPIMULT scales the well productivity of well OP01 by 1.25, scales the well connection factor in layer 10 only by 0.75 for well OP02, and for OP03, scales the top most 15 well connections by 1.100.

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12.3.98 WPITAB - Assign Well Productivity Index versus Water Cut Tables

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WPITAB keyword assigns the well productivity index multiplier versus water cut tables, that are used to scaled a well's connection factors based on the connection's current producing water cut, to a well. The tables are defined via the PIMULTAB keyword in the SCHEDULE section.

No.	Name		Description					
		Field	Field Metric Laboratory					
I	WELNAME	A character string of up name for which the well	None					
		Note that the well n previously using the W otherwise an error may o						
2	NTABLE	An integer positive value that defines the corresponding PIMULTAB table to be allocated to the well.						
		A value less than or ed allocated to the well	qual to zero means that	no PIMULTAB table is				

Notes:

Table 12.44:WPITAB Keyword Description

See also the PIMULTAB keyword that defines productivity index multiplier versus water cut tables and also the WPIMULT keyword that scales a well's productivity index by a constant value, both of which are in the SCHEDULE section.

This keyword is ignored by OPM Flow and has no effect on the simulation.

Example

Given NTPIMT equals two on PIMTDIMS keyword in the RUNSPEC section, then:

```
-- ASSIGN WELL PRODUCTIVITY INDEX VS WATER CUT TABLE
-- WELL PI
-- NAME TABLE
WPITAB
OP01 1 /
OP02 1 /
OP03 2 /
```

Assigns PIMULTAB table one to wells OP01 and OP02 and table two to OP03.

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¹⁾ The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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12.3.99 WPOLYMER - Define Water Injection Well Polymer and Salt

CONCENTRATIONS

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
---------	------	------	-------	---------	----------	---------	----------

Revision: Rev-I

Description

The WPOLYMER keyword defines a water injection well's polymer and salt injection stream concentrations that are to be used for when the polymer and salt options have been activated by the POLYMER and BRINE keywords in the RUNSPEC section.

Note that if the BRINE option has not be activated by the BRINE keyword in the RUNSPEC section, then the salt concentrations in the third column are ignored.

Currently the BRINE option is not implemented in OPM Flow and therefore both the SALTCON and GRPSALT variables on this keyword are ignored.

No.	Name		Description		Default		
		Field	Metric	Laboratory			
I	WELNAME		A character string of up to eight characters in length that defines the well name for which the well connection data is being defined.				
		Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.					
2	POLCON	A real positive value that injection stream.	t defines the polymer con	centration of the well's			
		lb/stb	kg/sm³	gm/scc	None		
3	SALTCON	A real positive value the injection stream.	at defines the salt conc	entration of the well's			
		This variable is ignored by OPM Flow and has no effect on the simulation but is documented here for completeness.					
		lb/stb	kg/sm³	gm/scc	None		
4	GRPPOL	A character string of up to eight characters in length that defines the group name for which the group's produced polymer concentration should be used instead of the well's POLCON value stated on this keyword.					
5	GRPSALT A character string of up to eight characters in length that defines the group name for which the group's produced salt concentration should be used instead of the well's SALTCON value stated on this keyword.						
		This variable is ignored but is documented here f	y OPM Flow and has no or completeness.	effect on the simulation			

Notes:

Table 12.45:WPOLYMER Keyword Description

Water injection wells that are not declared via this keyword have their concentrations defaulted to zero.

See also the GCONPROD and GCONINJE keywords to define a group's production and injection targets and constraints, and the WCONINJE keyword to define an injection well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

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I) The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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Example

The following example defines the polymer and salt injection stream concentrations for three water injection wells for when the polymer option has been activated by the POLYMER keyword in the RUNSPEC section.

Revision: Rev-I

```
DEFINE WATER INJECTION WELL POLYMER AND SALT CONCENTRATIONS
- -
-- WELL
        POLYMER
                   SALT
                               POLYMER
                                          SALT
                   SALTCON
                                          GROUP
-- NAME POLCON
                              GROUP
                    -----
WPOLYMER
WI01
        0.2500
        1*
                              GRPINJ1
WI02
                   1*
WI03
         0.2500
                               GRPINJ1
```

The polymer concentration for well WI01 is set to 0.25 and the stated polymer concentration for well WI02 will be ignored, as both WI02 and WI03 will re-inject the produced polymer from the GRPINJI group.

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12.3.100 WRFT - ACTIVATE WELL RFT REPORTING TO THE RFT FILE

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

This keyword activates reporting of a well's depth pressure profile to the RFT file for the requested wells at the time the keyword is activated. Data written out by OPM Flow is used to match the field measured data collected from a Repeat Formation Tester ("RFT") tool.

The RFT tool is an open hole device which is an updated version of the Formation Interval Tester (FIT), both of which are run on wire line. Both tools take multiple pressure readings (at various depths) thus enabling a pressure depth profile to be obtained from the formation, and, in addition, they can also take fluid samples from the formation. The latest tool available from Schulmberger is the Modular Formation Dynamics Tester (MDT), which, as its name suggests, is a modular tool that can be assembled in different configurations depending on what are the objectives for running the tool. Note other vendors have similar wire line logging tools with alternative names for the tools. Throughout this section the term RFT applies to all tools that measure a pressure profile versus depth (RFT/FIT/MDT etc.).

No.	Name	Description						
		Field Metric Laboratory						
I	WELNAME	for each item, that defin be written to the RFT declared previously usin	A columnar vector of character strings of up to eight characters in length for each item, that defines the well name for which the RFT data should be written to the RFT file. Note that the WELNAME must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.					
		If the WELNAME is left blank then the data is written out for all wells at the time they are first opened to flow.						
		If the WELNAME is given, then the RFT data for the well at the time step the keyword is invoked is written out.						

Notes:

- The keyword is followed by any number of records. I)
- Each record is terminated by a "/" and the keyword should be terminated by a "/".

Table 12.46:WRFT Keyword Description

See also the WRFTPLT keyword in the SCHEDULE section that has more flexible reporting options.

Examples

The first example activates RFT reporting for all wells at the time a well is first opened to flow:

```
ACTIVATE WELL RFT REPORTING TO THE RFT FILE
-- WELL
-- NAME
WRFT
```

Ideally, this version of the keyword should be place at the beginning of the SCHEDULE section to obtain the data for the wells in the run before they are opened up through time.

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The next example shows how to use the keyword to request the output for several wells at different reporting time steps.

```
-- 01 JAN 2000 START OF SCHEDULE SECTION
DATES
15 JAN 2000 /
         WELL HISTORICAL PRODUCTION CONTROLS
- -
                                                      VFP
-- WELL
        OPEN/
                CNTL
                        OIL
                               WAT
                                       GAS
                                              VFP
                                                            THP
                                                                  BHP
                                              TABLE ALFQ
-- NAME
        SHUT
                MODE
                        RATE
                               RATE
                                       RATE
                                                            PRES PRES
WCONHIST
0P01
         OPEN
                 ORAT 15.5E3 0.0
                                       1550
                                              10
                                                       1*
                                                            900.0 1*
0P02
         SHUT
- -
         ACTIVATE WELL RFT REPORTING TO THE RFT FILE
-- WELL
-- NAME
WRFT
0P01
0P02
DATES
01 FEB 2000 /
         WELL HISTORICAL PRODUCTION CONTROLS
- -
-- WELL
        OPEN/
                CNTL
                               WAT
                                       GAS
                                              VFP
                                                      VFP
                                                            THP
                                                                  BHP
                        OTI
-- NAME
         SHUT
                MODE
                        RATE
                               RATE
                                       RATE
                                              TABLE ALFQ
                                                            PRES
                                                                  PRES
WCONHIST
         OPEN
                ORAT 15.5E3 0.0
                                       1550
                                                       1*
                                                            900.0 1*
0P01
                                              10
0P02
         SHUT
/
_ _
         ACTIVATE WELL RFT REPORTING TO THE RFT FILE
-- WELL
-- NAME
WRFT
0P01
0P02
DATES
01 MAR 2000 /
_ _
--
         WELL HISTORICAL PRODUCTION CONTROLS
                CNTL
                                              VFP
                                                      VFP
                                                            THP
                                                                  BHP
-- WELL
        OPEN/
                        OIL
                               WAT
                                       GAS
-- NAME
         SHUT
                MODE
                        RATE
                               RATE
                                      RATE
                                              TABLE ALFQ
                                                            PRES PRES
WCONHIST
         OPEN
                 ORAT 15.5E3
                               0.0
                                       1550
                                                       1*
                                                            900.0 1*
0P01
                                              10
0P02
         OPEN
                ORAT 10.5E3
                               0.0
                                       1000
                                              10
                                                       1*
                                                            900.0 1*
In this example, both well's have their RFT written out on February I and March I 2000.
```

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12.3.101 WRFTPLT - ACTIVATE WELL RFT AND PLT REPORTING TO THE RFT FILE

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY	ISPEC	GRID EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
--------------------------------------------------	-------	-----------	-------	---------	----------	---------	----------

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Description

This keyword activates reporting of a well's depth pressure and fluid rates profile to the RFT file for the requested wells at the time the keyword is activated. Data written out by the simulator is used to match the field measured data collected from both the Repeat Formation Tester ("RFT") tool and various Production Logging Tools ("PLT").

See the WRFT keyword in the SCHEDULE section for a brief description of the RFT data set. This keyword also actives the writing out of each well connection's fluid rates, connection factors and KH data, etc., as the PLT data. The PLT data is used to compare with measured data from wire line production logging tools.

No.	Name			Default	
		Field	Metric	Laboratory	
I	WELNAME	for each item, that define be written to the RFT	aracter strings of up to ei es the well name for whi file. Note that the WELI g the WELSPECS keywo or may occur.	ch the RFT data should NAME must have been	None
		If the WELNAME is left the time they are first op	blank then the data is wri ened to flow.	tten out for all wells at	
		If the WELNAME is given the keyword is invoked is	n, then the RFT data for the written out.	ne well at the time step	
2	RFT		ng that sets the RFT data he following character stri		NO
		I) NO: do not not w	vrite RFT data for the wel		
		2) YES: write out the	e RFT data at the current	reporting time step.	
		,	the RFT data at the curr t <u>reporting</u> time steps.	ent reporting time step	
		4) TIMESTEP: write step and all subse	out the RFT data at the quent time steps.	current reporting time	
		for the well if it i	the RFT data at the curr is opened, otherwise writ red well is opened.		
3	PLT		ng that sets the PLT data he following character stri		NO
		I) NO: do not not w	vrite PLT data for the well		
		2) YES: write out the	e PLT data at the current r	reporting time step.	
		,	the PLT data at the curret reporting time steps.	ent reporting time step	
		4) TIMESTEP: write step and all subse	out the PLT data at the quent time steps.	current reporting time	
4		Not Used.			

Notes:

I) The keyword is followed by any number of records terminated by a "/" and the keyword should be terminated by a "/".

Table 12.47:WRFTPLT Keyword Description

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See also the WRFT keyword in the SCHEDULE section that has less flexible reporting options.

Examples

0P02

The first example activates RFT output at the current reporting time step for all the wells that are opened to flow, otherwise the RFT data is written out the first time a well is opened.

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```
WELL RFT, PLT AND SEGMENT DATA
-- WELL
         RFT
                PLT
                       SEGMENT
-- NAME
          DATA DATA DATA
WRFTPLT
          FOPN
The next example writes out the RFT and PLT data for two wells at the current reporting time step.
- -
          WELL RFT, PLT AND SEGMENT DATA
          RFT
                PLT
                       SEGMENT
-- WELL
-- NAME
          DATA DATA DATA
WRFTPLT
0P01
          YES
                YES
```

The final example is shown below:

YES

YES

```
-- WELL RFT, PLT AND SEGMENT DATA
-- WELL RFT PLT SEGMENT
-- NAME DATA DATA DATA
WRFTPLT
OP01 REPT NO /
OP02 NO YES /
```

In this case the RFT data for well OP01 is written out at the current reporting time step and all subsequent reporting time steps. For well OP02, no RFT is written out but the PLT data is written out for the current report time step only.

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12.3.102 WSEGITER - Define Multi-Segment Wells Iteration Parameters

RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDULE

Description

The WSEGSITER keyword defines the multi-segment well solution iteration sequence and solution controls.

OPM Flow users a different numerical scheme which makes this keyword redundant; hence, OPM Flow ignores this keyword. It is documented here for completeness. See section 2.2 Running Flow on how to set the numerical control parameters for OPM Flow.

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12.3.103 WSEGSICD - Define Multi-Segment Well Spiral ICD Connections

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

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Description

The WSEGSICD keyword defines a multi-segment well segment to be a spiral Inflow Control Device. ("ICD") as part of a completion for a multi-segment well. Note that the well must have been previously define by the WELSPECS and WELSEGS keywords in the SCHEDULE section and that the data for the keyword should be repeated for each multi-segment completion that contains a spiral ICD.

An ICD is a well completion component usually installed along the producing section of a well to minimize the unwanted water and gas breakthrough in an oil well, or early water production in a gas well, due to an uneven flow profile over the completed interval. Permeability variations over the producing interval cause the high permeability zones to produce higher quantities of fluids than the lower permeability zones and this uneven producing fluid profile may result in bypassed hydrocarbons. Secondly, for horizontal wells, the pressure loss from the "toe" to the "heel" of the well again results in an uneven fluid profile over the producing interval. In order to rectify this ICDs can be installed so that the well fluids have to flow through an ICD before entering the tubing; thus, creating an additional "designed" pressure loss. A spiral ICD is a type of frictional ICD that adds an additional pressure loss by directing the fluid along a "channel" (spiral/helix) before entering the tubing. The channel flow path is designed in such a manner as to create the desired pressure loss for a given ICD. By placing various ICD's over the production interval one can design a completion that results in a more uniform producing fluid profile throughout the length of the producing interval.

No.	Name		Description		Default			
		Field	Metric	Laboratory				
I	WELNAME	A character string of up to eight characters in length that defines the well name for which a multi-segment well is being defined.						
		previously using the W	Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.					
2	ISEGI		A positive integer greater than or equal to two and less than or equal to MXSEGS on WSEGDIMS keyword in the RUNSPEC section that defines the start of a segment					
3	ISEG2	A positive integer greater than or equal to two and less than or equal to ISEGI on this record and MXSEGS on the WSEGDIMS keyword in the RUNSPEC section that defines the end of a segment.						
4	ICDSTREN	A real positive value greater than zero that defines an empirical constant for the strength of the given ICD as determined from measurements using the calibrated fluid.						
		psia(rft³/day)²	barsa/(rm³/day)²	atma/(rcc/day) ²	None			

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No.	Name		Description		Default				
		Field	Metric	Laboratory					
5	ICDLEN	NSCAFAC to calculate a	A real value defines the length of the ICD used in conjunction with NSCAFAC to calculate a scaling factor to be applied to the reservoir flow to adjust the flow through each ICD, that is:						
		length of the ICI section, that is the	 If NSCAFAC equals zero: then the scale factor is equal to the length of the ICD (ICDLEN) divided by the length of the tubing section, that is the parent of the ICDs, then this allows for the case when the ICD segment may represent a number of ICDs in parallel. 						
		4) If NSCAFAC equal absolute value of	uals one: then the scale ICDLEN.	factor is equal to the					
			als two: then the scale fact ed by the total length of						
			ets which of the above to l, then option I) is used nen ICDLEN is negative.						
		feet	m	cm					
		39.37	12.00	1,2000	Defined				
6	CALDEN	CALDEN is a real positive of the calibrating fluid at	re value greater than zero surface conditions.	that defines the density					
		lb/ft³	kg/m³	gm/cc					
		62.416	1000.25	1.00025	Defined				
7	CALVISC		itive value greater than g fluid at surface condition						
		cP	cP	cP	0.45				
8	EMLCRT	water" in liquid fraction (e value greater than zero t used to determine whethe mulation equation should	r the "water-in-oil" or					
		dimensionless	dimensionless	dimensionless	0.5				
9	EMLTRANS	EMLTRANS is a real positive value greater than zero that defines the width of the transition zone around EMLCRT and is used to ensure that the calculated viscosity forms a continuous function of water in liquid fraction. Within this region, the emulsion viscosity is a linear interpolation between the "water-in-oil" and "oil-in-water" viscosity values either side of the region.							
		dimensionless	dimensionless	dimensionless	0.05				
10	EMLMAX		EMLMAX is a real positive value greater than zero that defines the maximum emulsion viscosity to continuous phase viscosity (oil or water)						
		dimensionless	dimensionless	dimensionless	5.0				
		1	I		I				

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No.	Name		Description		Default
		Field	Metric	Laboratory	
11	NSCAFAC	NSCALFAC is a positive zero, that sets the metho- should be set to one of the	d to be used when applyir		
		length of the ICE section, that is the	als zero: then the scale O (ICDLEN) divided by the parent of the ICDs, then segment may represent	he length of the tubing this allows for the case	
		If NSCAFAC equals absolute value of limits.	uals one: then the scale	factor is equal to the	
			als two: then the scale fact ed by the total length of		
		NSCALFAC explicitly se NSCALFAC is defaulted positive and option 2) wh	, then option 1) is used		
		dimensionless	dimensionless	dimensionless	None
12	CALRATE	A real positive value that the ICD was calibrated.	defines the maximum sur	face flow rate for which	
		scf/d	sm³/day	scc/hour	None
13	STATUS	A character string of leng			OPEN
		I) OPEN: the ICD co	onnection is are open to f	low.	
		2) SHUT: the ICD co	onnections is closed to flo	w (shut-in)	

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The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.48:WSEGSICD Keyword Description

The total number of wells should be defined via the WELLSDIMS keyword and the number of multi-segment wells should be declared on the WSEGDIMS keyword, both keywords are in the RUNSPEC section. In addition, the WELSPECS keyword should be used to define wells, the COMPDAT keyword to define the well completions for both ordinary wells and multi-segment wells, and the COMPSEGS keyword to define a multi-segment segment completions. Finally, the WSEGSICD keyword can then be use to define ICD connections for the well. All the aforementioned keywords are described in the SCHEDULE section.

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Example

The following example defines one producing well segment oil well (OP01) using the WELSPECS, WELSEGS COMPDAT and COMPSEGS keywords, followed by the WSEGSICD keyword to define the spiral inflow control devices for the well.

		WELL	SPEC1	[FICA	ΓΙΟN	DATA								
NAME NAME I J DEPTH FLUID AREA EQUINS SHUT FLOW TABLE		00011			FTON	DUD			DATN T	IELOU O	DEN OF	2000 5		
Decision														
OPO		IVALLE			J	DEI II	' '	LUID A	NLA L	S CNIVO?	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	-OW 17	ADEL	
WELL CONNECTION DATA WELL LOCATION OPEN SAT CONN WELL KH SKIN D DIR NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT PEN NAME II JJ KI K2 SHUT TAB FACT DIA FACT PEN NAME DIA FACT DIA FACT PEN NAME DEPH II JUBING NOLL TUBE TUBE XSEC VOL SEG SEG BRAN SEG TUBING NOLL TUBE TUBE XSEC VOL SEG SEG BRAN SEG TUBING NOLL TUBE TUBE XSEC VOL STATE TEND NOLL LEN WELL SEGNET SEG		PLAT	FORM	10	10	1*		OIL					/	
WELL CONVECTION DATA WALL	/													
WELL S LOCATION OPEN SAT CONN WELL KH SKIN D D DIR FACT NAME II JJ KI K2 SHUT TAB FACT DIA FACT FACT FACT PEN SHUT TAB FACT DIA FACT PEN SHUT TAB														
NAME ON THE COMPOND TO THE COMPON		WELL	CONNE	ECTIO	N DAT	Ά								
NAME ON THE COMPOND TO THE COMPON			LOCATI	ron.	0	DEN	CAT	CONN	MELL	IZH	CIZTN	D	DTD	
COMPDAT OPO1 10 10 10 1 1 0 PEN 1* 200 0.5														
OPO1			33 F	_		1101	IAL	J IACI	DIA	IACI	IACI	IACI	I LIV	
OPO1		10	10	1	1 0	PEN	1*	200.	0.5				/	
OPD1													/	
OPO1	0P01	10	10		6 0	PEN	1*	200.	0.4				/	
OPO1	0P01	10	10			PEN	1*	200.	0.4				/	
OPO1		10											/	
OPO1	0P01	10	10	6	6 0	PEN	1*	200.	0.4				/	
OPO1	0004	0	10	_		DEN	4 *	200	0 4				,	
OPO1														
OP01														
OP01														
OP01													,	
OP01														
OP01	0P01	10	9			PEN	1*	200.	0.4				/	
OP01		10				PEN			0.4				/	
OP01														
OP01 9 10 5 5 OPEN 1* 200. 0.4 OP01 8 10 5 5 OPEN 1* 200. 0.4 OP01 7 10 5 5 OPEN 1* 200. 0.4 OP01 6 10 5 5 OPEN 1* 200. 0.4 OP01 5 10 5 5 OPEN 1* 200. 0.4 OP01 15 10 5 5 OPEN 1* 200. 0.4 OP01 10 9 6 6 OPEN 1* 200. 0.4 OP01 10 8 6 6 OPEN 1* 200. 0.4 OP01 10 7 6 6 OPEN 1* 200. 0.4 OP01 10 7 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4 OP01 10 5 6 6 OPEN 1* 200. 0.4													/	
OP01	0001	10	5	3	3 0	PEN	1^	200.	0.4				/	
OP01	0P01	a	10	5	5 0	DEN	1*	200	0.4				/	
OP01														
OP01														
OPO1 10 9 6 6 OPEN 1* 200. 0.4 OPO1 10 8 6 6 OPEN 1* 200. 0.4 OPO1 10 7 6 6 OPEN 1* 200. 0.4 OPO1 10 6 6 6 OPEN 1* 200. 0.4 OPO1 10 5 6 6 OPEN 1* 200. 0.4	0P01	6	10			PEN	1*	200.	0.4				/	
OPO1 10 8 6 6 OPEN 1* 200. 0.4 OPO1 10 7 6 6 OPEN 1* 200. 0.4 OPO1 10 6 6 6 OPEN 1* 200. 0.4 OPO1 10 5 6 6 OPEN 1* 200. 0.4 // OPO1 10 5 6 6 OPEN 1* 200. 0.4 //	0P01	5	10	5	5 0	PEN	1*	200.	0.4				/	
OPO1 10 8 6 6 OPEN 1* 200. 0.4 OPO1 10 7 6 6 OPEN 1* 200. 0.4 OPO1 10 6 6 6 OPEN 1* 200. 0.4 OPO1 10 5 6 6 OPEN 1* 200. 0.4 // OPO1 10 5 6 6 OPEN 1* 200. 0.4 //			_											
OP01 10 7 6 6 OPEN 1* 200. 0.4 / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / /														
OP01 10 6 6 0 PEN 1* 200. 0.4 / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / / /													/	
OPO1 10 5 6 6 OPEN 1* 200. 0.4 //													/	
/													,	
WELL NODAL LEN WELL DEPH PRESS FLOW NAME DEPTH TUBING VOLM OPTN CALC MODEL WELSEGS OP01 2512.5 2512.5 1.0E-5 ABS HFA HO / SEG SEG BRAN SEG TUBING NODAL TUBE TUBE XSEC VOL ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG			Ŭ		, ,		_	2001	0				,	
WELL NODAL LEN WELL DEPH PRESS FLOW NAME DEPTH TUBING VOLM OPTN CALC MODEL WELSEGS OP01 2512.5 2512.5 1.0E-5 ABS HFA HO / SEG SEG BRAN SEG TUBING NODAL TUBE TUBE XSEC VOL ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG														
NAME DEPTH TUBING VOLM OPTN CALC MODEL WELSEGS OP01 2512.5 2512.5 1.0E-5 ABS HFA HO / SEG SEG BRAN SEG TUBING NODAL TUBE TUBE XSEC VOL ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG		WELL	SEGME	ENT S	PECIF	ICATIO	ON E	DATA						
NAME DEPTH TUBING VOLM OPTN CALC MODEL WELSEGS OP01 2512.5 2512.5 1.0E-5 ABS HFA HO / SEG SEG BRAN SEG TUBING NODAL TUBE TUBE XSEC VOL ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG														
WELSEGS 0P01														
OP01 2512.5 2512.5 1.0E-5 ABS HFA HO / SEG SEG BRAN SEG TUBING NODAL TUBE TUBE XSEC VOL ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG		DEPT	Н	TU	SING	VULM		OPIN	CALC	MODEL				
SEG SEG BRAN SEG TUBING NODAL TUBE TUBE XSEC VOL ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG		2512	5	25	12 5	1 05	- 5	ΔRS	НΕΔ	но			,	
SEG SEG BRAN SEG TUBING NODAL TUBE TUBE XSEC VOL ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG		2312	. 5	23.	12.5	1.02	J	ADO	111 🗸	110			/	
ISTR IEND NO NO LENGTH DEPTH ID ROUGH AREA SEG		SEG	SEG	BR	AΝ	SEG		TUBING	NODAL	TUBE	TUBE	XSE	C VOL	
2 2 1 1 2537.5 2534.5 0.3 0.00010 /														
									2534.5					
3 3 1 2 2562.5 2560.5 0.3 0.00010 /		3	3	1		2		2562.5	2560.5	0.3	0.0001	-0	/	

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	4 5 6		4 5 6	1 1 1	3 4 5	2587.5 2612.5 2637.5	2614	4.5	0.3 0.3 0.3	3 0	.00010 .00010 .00010	
	7		7	2	2	2737.5			0.2		.00010	
	8		8	2	7	2937.5			0.2		.00010	
	9		9	2	8	3137.5			0.2		.00010	
	10 11		10 11	2 2	9 1 0	3337.5 3537.5			0.2		.00010 .00010	
	12		12	3	3	2762.5	2563	2 5	0.2)	.00010	
	13		13	3	12	2962.5			0.1		.00010	
	14		14	3	13	3162.5			0.1		00010	
	15		15	3	14	3362.5			0.1		.00010	
	16		16	3	15	3562.5	2562	2.5	0.1	L 0.	.00010	
	17		17	4	5	2812.5			0.2		.00010	
	18		18	4	17	3012.5			0.1		.00010	
	19 20		19 20	4 4	18 19	3212.5 3412.5			$0.1 \\ 0.1$		00010	
	20		21	4	20	3412.5			0.1		.00010 .00010	
						5012.5	2010	3.3	0.1			
	22		22	5	6	2837.5			0.2		.00010	
	23		23	5	22	3037.5			0.2		.00010	
	24		24	5 5	23	3237.5 3437.5			0.2		.00010	
	25 26		25 26	5 5	24 25	3637.5			$0.1 \\ 0.1$.00010 .00010	
/	20		20	J	20	000710	200	0.0	0.1		.00010	
 WEL	LL	PLET	ION S	SEGMENT	SPECIFI	CATION D	АТА					
NAN COMPSE OP01	LL ME	PLET	ION S	SEGMENT	SPECIFI	CATION D	ATA					
NAN COMPSE	LL ME EGS						ATA DIR	LOC		MID	COMP	ISEG
NAN COMPSE OP01	LL ME EGS		ON K1	BRAN NO	TUBING LENGTH	NODAL DEPTH		LOC I,J		MID PERFS	COMP LENGTH	ISEG H NO.
NAN COMPSE OP01	LL ME EGS LO II 10	CATI JJ 10	ON K1 1	BRAN NO 1	TUBING LENGTH 2512.5	NODAL	DIR					
NAN COMPSE OP01	LL ME EGS LO II 10 10	CATI JJ 10 10	ON K1 1 2	BRAN NO 1 1	TUBING LENGTH 2512.5 2525.0	NODAL DEPTH 2525.0 2550.0	DIR					
NAN COMPSE OP01	LO II 10 10	CATI JJ 10 10	ON K1 1 2 3	BRAN NO 1 1	TUBING LENGTH 2512.5 2525.0 2550.0	NODAL DEPTH 2525.0 2550.0 2575.0	DIR					
NAN COMPSE OP01	LO II 10 10 10	CATI JJ 10 10 10	ON K1 1 2 3 4	BRAN NO 1 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0	DIR					
NAN COMPSE OP01	LO II 10 10	CATI JJ 10 10	ON K1 1 2 3	BRAN NO 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0	NODAL DEPTH 2525.0 2550.0 2575.0	DIR					
NAN COMPSE OP01	LO II 10 10 10 10	CATI JJ 10 10 10 10 10	ON K1 1 2 3 4 5 6	BRAN NO 1 1 1 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10	CATI JJ 10 10 10 10	ON K1 1 2 3 4 5 6	BRAN NO 1 1 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 7	CATI JJ 10 10 10 10 10 10	XON K1 1 2 3 4 5 6	BRAN NO 1 1 1 1 1 2 2	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5	DIR					
NAN COMPSE OP01	L0 II 10 10 10 10 10 9 8 7 6	CATI JJ 10 10 10 10 10 10 10	XON K1 1 2 3 4 5 6	BRAN NO 1 1 1 1 1 2 2 2 2	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5 3237.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 7	CATI JJ 10 10 10 10 10 10	XON K1 1 2 3 4 5 6	BRAN NO 1 1 1 1 1 2 2	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5	DIR					
NAN COMPSE OP01	L0 II 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 10 10 10 10 10	50N K1 1 2 3 4 5 6 2 2 2 2 2	BRAN NO 1 1 1 1 1 1 1 1 2 2 2 2 2 2 2 3	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5 3237.5 3437.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 10 10 10 10 10	50N K1 1 2 3 4 5 6 2 2 2 2 2	BRAN NO 1 1 1 1 1 1 1 1 1 2 2 2 2 2 2 2 3 3 3	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5 3237.5 3437.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 10 7 8 7	50N K1 1 2 3 4 5 6 2 2 2 2 2 2 3 3	BRAN NO 1 1 1 1 1 1 1 1 1 2 2 2 2 2 2 2 3 3 3 3	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5 3237.5 3437.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 10 10 10 10 10	50N K1 1 2 3 4 5 6 2 2 2 2 2	BRAN NO 1 1 1 1 1 1 1 1 1 2 2 2 2 2 2 2 3 3 3	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5 3237.5 3437.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 10 5 6 5	ON K1 1 2 3 4 5 6 2 2 2 2 2 2 3 3 3 3 3	BRAN NO 1 1 1 1 1 1 1 1 2 2 2 2 2 2 3 3 3 3 3 3	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5 3237.5 3437.5 2662.5 2862.5 3062.5 3262.5 3462.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5 3632.5 3662.5 3662.5	DIR					
NAN COMPSE OP01	L0 II 10 10 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 7 6	50N K1 1 2 3 4 5 6 2 2 2 2 2 2 3 3 3 3	BRAN NO 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 2837.5 3037.5 3237.5 3437.5 2662.5 2862.5 3062.5 3262.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5 3637.5	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 10 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 10 10 10 10 10	50N K1 1 2 3 4 5 6 2 2 2 2 2 2 3 3 3 3 5 5 5 5 5	BRAN NO 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 3037.5 3237.5 3437.5 2662.5 3062.5 3262.5 3462.5 2712.5 2912.5 3112.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5 3662.5 3662.5 3662.5 3662.5 3112.5 3312.5	DIR					
NAN COMPSE OP01	LO II 10 10 10 10 10 10 10 10 10 10 10 10 10	CATI JJ 10 10 10 10 10 10 10 10 10 10 10 10 10	50N K1 1 2 3 4 5 6 2 2 2 2 2 2 3 3 3 3 3 5 5 5	BRAN NO 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	TUBING LENGTH 2512.5 2525.0 2550.0 2575.0 2600.0 2625.0 2637.5 3037.5 3237.5 3437.5 2662.5 3062.5 3262.5 3462.5 2712.5 2912.5	NODAL DEPTH 2525.0 2550.0 2575.0 2600.0 2625.0 2650.0 2837.5 3037.5 3237.5 3437.5 3637.5 362.5 3662.5 3662.5 3662.5 3662.5 3112.5	DIR					

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10 10 10 10 10	9 6 8 6 7 6 6 6 5 6	5 5	2937. 3137. 3337.	5 31 5 33 5 35	37.5 37.5 37.5						/ / / /
MULTI	-SEGM	ENT WELL	ICD S	EGMEN	T SPEC	IFICAT	ION DAT	A			
SEG ISTR	SEG IEND	ICD STRNEN	ICD LEN	CAL DEN	CAL VISC	EML CRIT	EML TRANS	EML MAX	SCAL FAC	CAL RATE	OPEN CLOSE
7	10	0.00025	1*	1.0	0.45	0.50	0.05	5.0	2	1*	OPEN /
12	15	0.00025	1*	1.0	0.45	0.50	0.05	5.0	2	1*	OPEN /
17	20	0.00025	1*	1.0	0.45	0.50	0.05	5.0	2	1*	OPEN /
22 23 24 25	22 23 24 25	0.00025 0.00025 0.00025 0.00050	1* 1* 1* 10.0	1.0 1.0 1.0	0.45 0.45 0.45 0.45	0.50 0.50 0.50 0.50	0.05 0.05 0.05 0.05	5.0 5.0 5.0 5.0	2 2 2 2	1* 1* 1* 1*	OPEN / OPEN / OPEN /
	10 10 10 10 MULTI SEG ISTR 7 12 17 22 23 24	10 8 6 10 7 6 10 6 6 10 5 6 MULTI-SEGM SEG SEG ISTR IEND 7 10 12 15 17 20 22 22 23 23 24 24	10 8 6 5 10 7 6 5 10 6 6 5 10 5 6 5 MULTI-SEGMENT WELL SEG SEG ICD ISTR IEND STRNEN 7 10 0.00025 12 15 0.00025 17 20 0.00025 22 22 0.00025 23 23 0.00025 24 24 0.00025	10 8 6 5 2937. 10 7 6 5 3137. 10 6 6 5 3337. 10 5 6 5 3537. MULTI-SEGMENT WELL ICD S SEG SEG ICD ICD ISTR IEND STRNEN LEN 7 10 0.00025 1* 17 20 0.00025 1* 17 20 0.00025 1* 22 22 0.00025 1* 23 23 0.00025 1* 24 24 0.00025 1*	10 8 6 5 2937.5 31 10 7 6 5 3137.5 33 10 6 6 5 3337.5 35 10 5 6 5 3537.5 37 MULTI-SEGMENT WELL ICD SEGMEN SEG SEG ICD ICD CAL ISTR IEND STRNEN LEN DEN 7 10 0.00025 1* 1.0 17 20 0.00025 1* 1.0 17 20 0.00025 1* 1.0 22 22 0.00025 1* 1.0 23 23 0.00025 1* 1.0 24 24 0.00025 1* 1.0	10 8 6 5 2937.5 3137.5 10 7 6 5 3137.5 3337.5 10 6 6 5 3337.5 3537.5 10 5 6 5 3537.5 3737.5 MULTI-SEGMENT WELL ICD SEGMENT SPEC SEG SEG ICD ICD CAL CAL ISTR IEND STRNEN LEN DEN VISC 7 10 0.00025 1* 1.0 0.45 12 15 0.00025 1* 1.0 0.45 17 20 0.00025 1* 1.0 0.45 22 22 0.00025 1* 1.0 0.45 23 23 0.00025 1* 1.0 0.45 24 24 0.00025 1* 1.0 0.45	10 8 6 5 2937.5 3137.5 10 7 6 5 3137.5 3337.5 10 6 6 5 3337.5 3537.5 10 5 6 5 3537.5 3737.5 MULTI-SEGMENT WELL ICD SEGMENT SPECIFICAT SEG SEG ICD ICD CAL CAL EML ISTR IEND STRNEN LEN DEN VISC CRIT 7 10 0.00025 1* 1.0 0.45 0.50 12 15 0.00025 1* 1.0 0.45 0.50 17 20 0.00025 1* 1.0 0.45 0.50 22 22 0.00025 1* 1.0 0.45 0.50 23 23 0.00025 1* 1.0 0.45 0.50 24 24 0.00025 1* 1.0 0.45 0.50	10 8 6 5 2937.5 3137.5 10 7 6 5 3137.5 3337.5 10 6 6 5 3337.5 3537.5 10 5 6 5 3537.5 3737.5 MULTI-SEGMENT WELL ICD SEGMENT SPECIFICATION DAT SEG SEG ICD ICD CAL CAL EML EML ISTR IEND STRNEN LEN DEN VISC CRIT TRANS 7 10 0.00025 1* 1.0 0.45 0.50 0.05 12 15 0.00025 1* 1.0 0.45 0.50 0.05 17 20 0.00025 1* 1.0 0.45 0.50 0.05 22 22 0.00025 1* 1.0 0.45 0.50 0.05 23 23 0.00025 1* 1.0 0.45 0.50 0.05 24 24 0.00025 1* 1.0 0.45 0.50 0.05	10 8 6 5 2937.5 3137.5 10 7 6 5 3137.5 3337.5 10 6 6 5 3337.5 3537.5 10 5 6 5 3537.5 3737.5 MULTI-SEGMENT WELL ICD SEGMENT SPECIFICATION DATA SEG SEG ICD ICD CAL CAL EML EML EML EML ISTR IEND STRNEN LEN DEN VISC CRIT TRANS MAX 7 10 0.00025 1* 1.0 0.45 0.50 0.05 5.0 12 15 0.00025 1* 1.0 0.45 0.50 0.05 5.0 17 20 0.00025 1* 1.0 0.45 0.50 0.05 5.0 17 22 22 0.00025 1* 1.0 0.45 0.50 0.05 5.0 18 23 23 0.00025 1* 1.0 0.45 0.50 0.05 5.0 18 24 24 0.00025 1* 1.0 0.45 0.50 0.05 5.0 18 24 24 0.00025 1* 1.0 0.45 0.50 0.05 5.0	10 8 6 5 2937.5 3137.5 10 7 6 5 3137.5 3337.5 10 6 6 5 3337.5 3537.5 10 5 6 5 3537.5 3737.5 MULTI-SEGMENT WELL ICD SEGMENT SPECIFICATION DATA SEG SEG ICD ICD CAL CAL EML EML EML SCAL TRANS MAX FAC 1STR IEND STRNEN LEN DEN VISC CRIT TRANS MAX FAC 7 10 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 12 15 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 17 20 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 22 22 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 23 23 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 24 24 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2	10 8 6 5 2937.5 3137.5 10 7 6 5 3137.5 3337.5 10 6 6 5 3337.5 3537.5 10 5 6 5 3537.5 3737.5 MULTI-SEGMENT WELL ICD SEGMENT SPECIFICATION DATA SEG SEG ICD ICD CAL CAL EML EML EML SCAL CAL ISTR IEND STRNEN LEN DEN VISC CRIT TRANS MAX FAC RATE 7 10 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 1* 17 20 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 1* 18 22 22 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 1* 22 23 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 1* 23 23 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 1* 24 24 0.00025 1* 1.0 0.45 0.50 0.05 5.0 2 1*

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Branch number two users one ICD for segments seven to ten, branch number three again users only one ICD for segments I2 to I5 and similarly branch number three users one ICD. The fifth and final branch has a total of four ICDs with the last interval having a I0 foot length. Since NSCAFAC equals two for the ICDs, then the scale factor is equal to the length to ICDLEN, divided by the total length of the completions which supply the ICD. Where ICDLEN is defaulted, the default value of 39.37 ft will be used for field units and I2 m for SI units.

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12.3.104 WSOLVENT - Define Gas Injection Well Solvent Fraction

	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
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Revision: Rev-I

Description

WSOLVENT defines a gas injection well's solvent faction in the injection stream that is to be used when the solvent option been activated by the SOLVENT keyword in the RUNSPEC section.

No.	Name		Description		Default					
		Field	Field Metric Laboratory							
1	WELNAME		A character string of up to eight characters in length that defines the well name of a gas injection well for which the solvent fraction data is being defined.							
		previously using the W	Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.							
4	SOLFRA	A real positive value greater than or equal to zero and less than or equal to one that defines the fraction of solvent in the gas well's injection stream.								
		fraction	fraction fraction fraction							

Notes:

Table 12.49:WSOLVENT Keyword Description

Gas injection wells that are not declared via this keyword have their solvent fractions set to zero.

See also the GCONINJE keyword to define a group's injection targets and constraints, and the WCONINJE keyword to define an injection well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines the solvent fractions for three gas injection wells for when the solvent option has been activated by the SOLVENT keyword in the RUNSPEC section.

```
-- DEFINE GAS INJECTION WELL SOLVENT FRACTION
-- WELL SOLVENT
-- NAME FRACTION
-- WSOLVENT
GI01 0.0000 //
GI02 0.5000 //
GI03 0.5000 //
```

The solvent fraction for the GI01 gas injector is set to zero and both GI02 and GI03 gas injectors have solvent fraction values of 0.5 for their injection streams.

¹⁾ The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

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12.3.105 WTEMP - Define An Injection Well's Fluid Temperature

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WTEMP keyword defines the temperature of the injection fluid being injected by an injection well.

This keyword can only be used if OPM Flow's thermal option has been activated by the THERMAL keyword in the RUNSPEC section. Note this is different to the commercial simulator that users the TEMP keyword in the RUNSPEC section to activate the "black-oil" thermal model.

No.	Name		Description						
		Field							
I	WELNAME	A character string of up name for an injection we data is being defined.	None						
		Note that the well n previously using the W otherwise an error may o							
2	TEMP	A real positive value greatinjected fluid.							
		°F	°C	°C	None				

Notes:

- Injection wells that are not declared via this keyword have their injection fluid temperatures set to zero degrees in the run's units.
- 2) The keyword is followed by any numbers records with each record terminated by a "/" and the keyword should be terminated by a "/".

Table 12.50:WTEMP Keyword Description

See also the GCONINJE keyword to define a group's injection targets and constraints, and the WCONINJE keyword to define an injection well's targets and constraints. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines the injected fluid temperatures for three water injection wells for when the thermal option has been activated by the THERMAL keyword in the RUNSPEC section.

```
-- DEFINE INJECTION WELL FLUID TEMPERATURE
-- WELL FLUID
-- NAME TEMP.
-- WTEMP
WI01 39.00 /
WI02 37.00 /
WI03 39.00 /
```

Here wells WI01 and WI03 inject water with a water temperature of 39 $^{\circ}F$ and well WI02's injection water temperature is 37 $^{\circ}F$.

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12.3.106 WTEST - Well Testing Criteria for Re-Opening Closed Wells

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE

Revision: Rev-I

Description

The WTEST keyword outlined the testing procedures to be be applied to wells that are closed for various reason to see if the wells are capable flowing under the current operating conditions. The keyword can be applied to single wells or groups of wells.

No.	Name	Description					
		Field	Metric	Laboratory			
1	WELNAME	A character string of up to eight characters in length that defines the well name for which the well connection data is being defined.					
		Note that the well name (WELNAME) must have been declared previously using the WELSPECS keyword in the SCHEDULE section, otherwise an error may occur.					
2	TIME	A real value greater than to zero that defines the period before another test is performed, for example if TIME is set equal to 365.25 (days), the test is performed every year.					
		days days hours					
3	TEST	A character string of up to five characters that defines the reason the well was closed. If a well was closed for one of the criteria then the well is tested to see if it can be put back on production. The characters that can be used to define TEST are as follows:					
		 P: meaning the well was closed due to a bottom-hole or tubing head pressure limit, or other physical limit then the well is tested to see if it can flow, if it can then it is put back on production, otherwise it remains closed. 					
		 E: meaning the well was closed due to a well or a well connection economic constraint then the well is tested to see if it can flow, if it can then it is put back on production, otherwise it remains closed. 					
		 G: meaning the constraint then the it is put back on p 					
		4) D: not used by OPM Flow.					
		5) C: not used by OPM Flow.					
		The default value is an empty string "" that switches of testing.					
		Note that only the E of	ption is currently suppor	ted in OPM Flow.			
4	NTIME	A positive integer greater than or equal to zero that define the number of times a well can be tested.					
		The default value of zero means an infinite number of times.					
5	START	START A real positive value that defines the start up time used to prorate the rate at which the well is put back on production. If START is large compared to the tilme step size, then the well is brought on gradually, if it is less then the well is opened faster.					
		The default value of 0.0 r	neans the well is opened i	mmediately.			
		days	days	hours	0.0		

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No.	Name		Default		
		Field	Metric	Laboratory	

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Notes:

The keyword is followed by any number of records and each record is terminated by a "/" and the keyword should be terminated by a "/".

Table 12.51:WTEST Keyword Description

See also the WELSPECS keyword to define a wells shut-in or stop options, WECON for setting a well's economic criteria, GCONPROD and GCONINJE for group controls, and GECON for setting a group's economic criteria. All the aforementioned keywords are described in the SCHEDULE section.

Example

The following example defines test criteria for all gas wells ("GP*") and three oil wells (OP01, OP02, and OP03).

```
- -
          WELL TESTING CRITERIA FOR RE-OPENING CLOSED WELLS
-- WELL
          TST
                 TST
                         NO.
                                 STRT
-- NAME
         INTV
                 TYPE
                         TSTS
                                 TIME
WTEST
'GP*'
          365.25 P
                                 0.0
0P01
          30.0
                 PFG
                         0
                                 0.0
0P02
          30.0
                 PEG
                         0
                                 0.0
0P03
          30.0
                 PEG
                         0
                                 0.0
```

All the gas wells are test annually if they have been shut-in due to a bottom-hole or tubing head pressure limit, are tested five times after they have been closed, and are opened up immediately. The oil wells are tested every 30 days if they have been closes due bottom-hole or tubing head pressure limit, a well economic limit or a group economic limit. All the oil wells are tested an infinite amount of times and are opened up immediately. **Note that only the E option is currently supported in OPM Flow**

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12.3.107 WTRACER - Define An Injection Well's Tracer Concentration

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Description

The WELTRACER keyword defines the tracer concentration of the injection fluid being injected by an injection well. This keyword should only be used if the tracer option has been invoked by the TRACER keyword in the RUNSPEC section.

OPM Flow has not implemented tracer modeling and therefore this keyword is ignored by OPM Flow and has no effect on the simulation.

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12.3.108 ZIPPY2 - ACTIVATE AUTOMATIC TIME STEP CONTROL

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Description

The ZIPPY2 keyword actives the commercial simulator's alternative automatic time step selection algorithm that assumes no <u>prior</u> knowledge of the problem, as opposed to the standard time step algorithm that is controlled via the TUNNING keyword in the SCHEDULE section, combined with posterior knowledge gained from previous time steps.

This keyword is ignored by OPM Flow and has no effect on the simulation.

See section 2.2 Running Flow on how to control time stepping for OPM Flow.

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13 KEYWORD INDEX - ALPHABETIC LISTING

A

	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter A	Status
ACTDIMS – ACTION Keyword Dimensions	
ACTION – Define Action Conditions and Command Processing (Field)	
ACTIONG – Define Action Conditions and Command Processing (Groups)	
ACTIONR – Define Action Conditions and Command Processing (Regions)	
ACTIONS – Define Action Conditions and Command Processing (Well Segments)	
ACTIONW – Define Action Conditions and Command Processing (Wells)	
ACTIONX – Define Action Conditions and Command Processing	
ACTNUM – Set the Status of a Grid Block To Active or Inactive	
ADD – Add a Constant to a Specified Array	
ADDREG – Add a Constant to an Array based on a Region Number	
ADSALNOD – Salt Concentration Based on SATNUM Array	
AITS - Intelligent Time Stepping Activation	
ALL – Export Standard Summary Variable Vectors to File	
API – Activate API Tracking	
APIGROUP – Define API Tracking Number of Grouped Oil PVT Tables	
AQUANCON – Define Analytical Connections to the Grid	
AQUCON – Define Numerical Aquifer Connections to the Grid	
AQUCT – Define Carter-Tracy Analytical Aquifers	
AQUDIMS – Aquifer Dimensions	
AQUFETP – Fetkovich Aquifer Definition	
AQUNUM – Numerical Aquifer Assignment	
AQUTAB – Define Carter-Tracy Aquifer Influence Functions	
ASPKDAM – Define Asphaltene Permeability Damage	
ASPPW2D – Define Asphaltene Two Parameters Precipitation Data	
ASPREWG -Define Asphaltene as Percentage Weight	
ASPWETF – Define Asphaltene Wettability Factor Data	

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B

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter B	Status
BLACKOIL – Activate Black Oil Phases	
BOX - Define a Range of Grid Blocks to Enter Property Data	

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C

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Alphabetic Listing of Keywords Starting with the Letter C	Status
CARFIN – Define a Cartesian Local Grid Refinement	
CECON – Define Well Connections Economic Limit Criteria	
CIRCLE – Completion of Radial Grid Circle Activation	
COALNUM – Define the Coal Region Numbers	
COMPDAT – Define Well Connections to the Grid	
COMPDATL – Define Well Connections to a LGR Grid	
COMPIMB – Assign Imbibition Saturation Tables to Well Connections	
COMPLUMP – Assign Well Connections to Completions	
COMPORD - Define Well Connection Ordering	
COMPSEGS – Define Well Connections for Multi-Segment Wells	
COORD – Define a Set of Coordinates Lines for a Reservoir Grid	
COORDSYS – Define Coordinate Grid Options	
COPY – Copy Array Data to Another Array	
Error: Reference source not found	
CPR – Activate Constrained Pressure Residual ("CPR") Linear Solver	

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D

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Alphabetic Listing of Keywords Starting with the Letter D	Status
DATE - Activate the DATE Option for the SUMMARY File	
DATES – Advance Simulation by Reporting Date	
DATUM – Define the Datum Depth for the Model	
DATUMR – Define Datum Depths for the FIPNUM Regions	
DEBUG – Define the Debug Data to be Printed to File	
DEADOIL – Activate the Dead Oil Phase (No Free or Dissolved Gas)	
DENSITY – Define the Surface Oil, Water Gas Densities for the Fluids	
DEPTH - Edits the Depth at the Center of Each Cell	
DIMENS – Define the Dimension of the Model	
DISGAS – Activate the Dissolved Gas Phase in the Model	
DR - Define the Size of Grid Blocks in the R Direction for All Cells	
DRSDT – Solution Gas (Rs) Maximum Rate of Increase Parameters	
DRSDTR – Solution Gas (Rs) Maximum Rate of Increase Parameters by Region	
DRV - Define the Size of Grid Blocks in the R Direction via a Vector	
DRVDT – Solution Oil (Rv) Maximum Rate of Increase Parameters	
DRVDTR – Solution Oil (Rv) Maximum Rate of Increase Parameters by Region	
DTHETA - Define the Size of Grid Blocks in the THETA Direction for All Cells	
DTHETAV - Sets the Size of Grid Blocks in THETA Direction via a Vector	
DUMPFLUX – Activate Writing Out of a Flux File	
DX - Define the Size of Grid Blocks in the X Direction for All Cells	
DXV - Define the Size of Grid Blocks in the X Direction via a Vector	
DY - Define the Size of Grid Blocks in the Y Direction for All Cells	
DYV - Define the Size of Grid Blocks in the Y Direction via a Vector	
DZ - Define the Size of Grid Blocks in the Z Direction for All Cells	
DZV - Define the Size of Grid Blocks in the Z Direction via a Vector	

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Alphabetic Listing of Keywords Starting with the Letter E	Status
ECHO – Activate Echoing of User Input Files to the Print File	
EDIT - Define the Start of the EDIT Section of Keywords	
EDITNNC – Scale Non-Neighbor Connections Between Cells Manually	
EDITNNCR – Reset Non-Neighbor Connections Between Cells Manually	
EHYSTR – Define Hysteresis Model and Parameters	
END – Define the End of the Input File	
ENDACTIO – End the Definition of ACTION Commands	
ENDBOX – Define the End of the BOX Defined Grid	
ENDFIN – End the Definition of a Local Grid Refinement	
ENDINC – Define the End of an Include File	
ENDNUM – Define the End-Point Scaling Depth Region Numbers	
ENDSCALE – Activate Relative Permeability End-Point Scaling Option	
ENDSKIP – Deactivates Skipping of Keywords and Input Data	
ENKRVD – Define Relative Permeability End-Points versus Depth Functions	
ENKRVDX ENKRVD – Define Relative Permeability End-Points versus Depth Functions	
ENKRVDX- ENKRVD – Define Relative Permeability End-Points versus Depth Functions	
ENKRVDY ENKRVD – Define Relative Permeability End-Points versus Depth Functions	
ENKRVDY- ENKRVD – Define Relative Permeability End-Points versus Depth Functions	
ENKRVDZ ENKRVD – Define Relative Permeability End-Points versus Depth Functions	
ENKRVDZ- ENKRVD – Define Relative Permeability End-Points versus Depth Functions	
ENPTVD – Define Relative Permeability Saturation End-Points versus Depth	
ENPTVDX ENPTVD – Define Relative Permeability Saturation End-Points versus Depth	
ENPTVDX- ENPTVD – Define Relative Permeability Saturation End-Points versus Depth	
ENPTVDY ENPTVD – Define Relative Permeability Saturation End-Points versus Depth	
ENPTVDY- ENPTVD – Define Relative Permeability Saturation End-Points versus Depth	
ENPTVDZ ENPTVD – Define Relative Permeability Saturation End-Points versus Depth	
ENPTVDZ- ENPTVD – Define Relative Permeability Saturation End-Points versus Depth	
EQLDIMS – Define the Equilibration Data Dimensions	

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RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHEDU

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Alphabetic Listing of Keywords Starting with the Letter E	Status
EQLNUM – Define the Equilibration Region Numbers	
EQLOPTS – Activates the Equilibration Options	
Error: Reference source not found	
Error: Reference source not found	
EQUIL – Define the Equilibration Initialization Data	
EXCEL - Activate the EXCEL Option for the SUMMARY File	
EXTRAPMS – Activate Extrapolation Warning Messages	

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F

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Alphabetic Listing of Keywords Starting with the Letter F	Status
FAULTDIM – Define the Number of Fault Segments	
FAULTS – Define Faults in the Grid Geometry	
FIELD – Activates the Oil Field System of Units for the Model	
FILLEPS – Activate Saturation End-Point Export to the INIT File	
FIPNUM – Define the Fluid In-Place Region Numbers	
FIPOWG – Activate Oil, Gas, and Water FIP Zone Reporting	
FLUXNUM – Define the Flux Regions	
FMTIN – Activates The Format Input File Option	
FMTOUT – Activates The Format Output File Option	
FWSET - Export Well Status Vectors for the Field to File	
FULLIMP – Activates Fully Implicit Solution Option	

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G

	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
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Revision: Rev-I

Alphabetic Listing of Keywords Starting with the Letter G	Status
GAS – Activate the Gas Phase in the Model	
GASDENT – Define Gas Density Temperature Coefficients	
GASVISCT – Define Gas Viscosity versus Temperature Functions	
GCONINJE – Group Injection Targets and Constraints	
GCONPROD – Group Production Targets and Constraints	
GDFILE – Load a Grid File	
GDORIENT - Define Grid Orientation Parameters	
GECON – Group Economic Criteria for Production Groups	
GEFAC – Define Group Efficiency	
GLIFTOPT – Define Group Gas Optimization Limits	
GMWSET - Export Well Status Vectors by Group to File	
GNETINJE – Define Group Injection Network Configuration	
GRAVITY- Define the Surface Oil, Water Gas Gravities for the Fluids	
GRID - Define the Start of the GRID Section of Keywords	
GRIDFILE – Set the Grid File Output Options	
GRIDOPTS - Grid Processing Options	
GRIDUNIT – Define the Grid Units	
GRUPNET – Define Group Standard Network Parameters	
GRUPRIG – Group Drilling and Workover Rig Specifications	
GRUPTREE – Define Group Tree Hierarchy	
GSATPROD – Define Group Satellite Production Rates	

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RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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HEATCR – Define Reservoir Rock Heat Capacity for All Cells			
HEATCRT – Define Reservoir Rock Heat Capacity Temperature Dependence for All Cells			

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Alphabetic Listing of Keywords Starting with the Letter I	Status
IMBNUM – Define the Imbibition Saturation Table Region Numbers	
IMBNUMX IMBNUM – Define the Imbibition Saturation Table Region Numbers	
IMBNUMY IMBNUM – Define the Imbibition Saturation Table Region Numbers	
IMBNUMZ IMBNUM – Define the Imbibition Saturation Table Region Numbers	
IMBNUMX- IMBNUM – Define the Imbibition Saturation Table Region Numbers	
IMBNUMY- IMBNUM – Define the Imbibition Saturation Table Region Numbers	
IMBNUMZ- IMBNUM – Define the Imbibition Saturation Table Region Numbers	
IMKRVD – Imbibition Relative Permeability End-Points versus Depth Functions	
IMKRVDX IMKRVD – Imbibition Relative Permeability End-Points versus Depth Functions	
IMKRVDX- IMKRVD – Imbibition Relative Permeability End-Points versus Depth Functions	
IMKRVDY IMKRVD – Imbibition Relative Permeability End-Points versus Depth Functions	
IMKRVDY- IMKRVD – Imbibition Relative Permeability End-Points versus Depth Functions	
IMKRVDX IMKRVD – Imbibition Relative Permeability End-Points versus Depth Functions	
IMKRVDZ- IMKRVD – Imbibition Relative Permeability End-Points versus Depth Functions	
IMPES – Activates Implicit Pressure Explicit Saturation Solution Option	
IMPTVD – Imbibition Relative Permeability Saturation End-Points versus Depth	
IMPTVDX IMPTVD – Imbibition Relative Permeability Saturation End-Points versus Depth	
IMPTVDX- IMPTVD – Imbibition Relative Permeability Saturation End-Points versus Depth	
IMPTVDY IMPTVD – Imbibition Relative Permeability Saturation End-Points versus Depth	
IMPTVDY- IMPTVD – Imbibition Relative Permeability Saturation End-Points versus Depth	
IMPTVDZ IMPTVD – Imbibition Relative Permeability Saturation End-Points versus Depth	
IMPTVDZ- IMPTVD – Imbibition Relative Permeability Saturation End-Points versus Depth	
INCLUDE – Load Another Data File at the Current Position	
INIT – Activates the INIT File Option	
IPCG – End-Point Scaling of Grid Cell Gas Capillary Pressure (Imbibition)	
IPCW – End-Point Scaling of Grid Cell Water Capillary Pressure (Imbibition)	
INRAD – Define the Inner Radius of a Radial Grid	
ISGCR – End-Point Scaling of Grid Cell Critical Gas Saturation (Imbibition)	

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Alphabetic Listing of Keywords Starting with the Letter I	Status
ISGL – End-Point Scaling of Grid Cell Connate Gas Saturation (Imbibition)	
ISGU – End-Point Scaling of Grid Cell Maximum Gas Saturation (Imbibition)	
ISOGCR – End-Point Scaling of Grid Cell Critical Oil Saturation to Gas (Imbibition)	
ISOLNUM – Define the Independent Reservoir Regions	
ISOWCR – End-Point Scaling of Grid Cell Critical Oil Saturation to Water (Imbib.)	
ISWCR – End-Point Scaling of Grid Cell Critical Water Saturation (Imbibition)	
ISWL – End-Point Scaling of Grid Cell Connate Water Saturation (Imbibition)	
ISWU – End-Point Scaling of Grid Cell Maximum Water Saturation (Imbibition)	

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J

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords	Starting with the Letter J	Status
JFUNC - Activates the Leverett J-function Option		

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K

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Alphabetic Listing of Keywords Starting with the Letter K	Status
KRG – End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)	
KRGX KRG – End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)	
KRGX- KRG – End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)	
KRGY KRG – End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)	
KRGY- KRG – End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)	
KRGZ KRG – End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)	
KRGZ- KRG – End-Point Scaling of Grid Cell Krg(Sgu) (Drainage)	
KRGR – End-Point Scaling of Grid Cell Krgr(1-Sogcr) (Drainage)	
KRGRX KRGR – End-Point Scaling of Grid Cell Krgr(1-Sogcr) (Drainage)	
KRGRX- KRGR – End-Point Scaling of Grid Cell Krgr(1-Sogcr) (Drainage)	
KRGRY KRGR – End-Point Scaling of Grid Cell Krgr(1-Sogcr) (Drainage)	
KRGRY- KRGR – End-Point Scaling of Grid Cell Krgr(1-Sogcr) (Drainage)	
KRGRZ KRGR – End-Point Scaling of Grid Cell Krgr(1-Sogcr) (Drainage)	
KRGRZ- KRGR – End-Point Scaling of Grid Cell Krgr(1-Sogcr) (Drainage)	
KRO – End-Point Scaling of Grid Cell Kro(Swl) (Drainage)	
KROX KRO – End-Point Scaling of Grid Cell Kro(Swl) (Drainage)	
KROX- KRO – End-Point Scaling of Grid Cell Kro(Swl) (Drainage)	
KROY KRO – End-Point Scaling of Grid Cell Kro(Swl) (Drainage)	
KROY- KRO – End-Point Scaling of Grid Cell Kro(Swl) (Drainage)	
KROZ KRO – End-Point Scaling of Grid Cell Kro(Swl) (Drainage)	
KROZ- KRO – End-Point Scaling of Grid Cell Kro(Swl) (Drainage)	
KRORG – End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)	
KRORGX KRORG – End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)	
KRORGX- KRORG – End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)	
KRORGY KRORG – End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)	
KRORGY- KRORG – End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)	
KRORGZ KRORG – End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)	
KRORGZ- KRORG – End-Point Scaling of Grid Cell Kro(Sgcr) (Drainage)	

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K

RUNSPEC GRI	ID EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter K	Status
KRORW – End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)	
KRORWX KRORW – End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)	
KRORWX- KRORW – End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)	
KRORWY KRORW – End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)	
KRORWY- KRORW – End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)	
KRORWZ KRORW – End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)	
KRORWZ- KRORW – End-Point Scaling of Grid Cell Kro(Swcr) (Drainage)	
KRW – End-Point Scaling of Grid Cell Krw(Sw =1.0) (Drainage)	
KRWX KRW – End-Point Scaling of Grid Cell Krw(Sw =1.0) (Drainage)	
KRWX- KRW – End-Point Scaling of Grid Cell Krw(Sw =1.0) (Drainage)	
KRWY KRW – End-Point Scaling of Grid Cell Krw(Sw =1.0) (Drainage)	
KRWY- KRW – End-Point Scaling of Grid Cell Krw(Sw =1.0) (Drainage)	
KRWZ KRW – End-Point Scaling of Grid Cell Krw(Sw =1.0) (Drainage)	
KRWZ- KRW – End-Point Scaling of Grid Cell Krw(Sw =1.0) (Drainage)	
KRWR – End-Point Scaling of Grid Cell KRWR(Sw =1.0) (Drainage)	
KRWRX KRWR – End-Point Scaling of Grid Cell KRWR(Sw =1.0) (Drainage)	
KRWRX- KRWR – End-Point Scaling of Grid Cell KRWR(Sw =1.0) (Drainage)	
KRWRY KRWR – End-Point Scaling of Grid Cell KRWR(Sw =1.0) (Drainage)	
KRWRY- KRWR – End-Point Scaling of Grid Cell KRWR(Sw =1.0) (Drainage)	
KRWRZ KRWR – End-Point Scaling of Grid Cell KRWR(Sw =1.0) (Drainage)	
KRWRZ- KRWR – End-Point Scaling of Grid Cell KRWR(Sw =1.0) (Drainage)	

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L

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Alphabetic Listing of Keywords Starting with the Letter L	Status
LAB - Activates the Laboratory System of Units for the Model	
LGR – Define Local Grid Refinement Parameters	
LICENSES – Define Required Licenses for Run	
LIFTOPT – Activate Gas Lift Optimization	
LIVEOIL – Activate the Live Oil Phase (Oil with Free and Dissolved Gas)	

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M

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Alphabetic Listing of Keywords Starting with the Letter M	Status
MAPAXES- Define the Map Origin Input Data	
MAPUNITS – Define the Map Axes Units	
Error: Reference source not found	
MEMORY – Define Allocated Memory	
MESSAGES – Define Message Print Limits and Stop Limits	
METRIC – Activates the Metric System of Units for the Model	
MINPV – Set a Minimum Grid Block Pore Volume Threshold for All Cells	
MINPVV – Set a Minimum Grid Block Pore Volume Threshold for Individual Cells	
Error: Reference source not found	
MISC – Define Solvent Miscibility-Immiscibility Transform Functions	
MISCIBLE – Define Miscibility Todd-Longstaff Parameters	
MISNUM – Define the Miscibility Region Numbers	
MONITOR – Activates Output of the Monitoring Data and File	
MSFN – Miscible Normalized Relative Permeability Tables	
MSGFILE – Active or Deactivate Message File Output	
MULTFLT – Multiply the Transmissibility of a Defined Fault by a Constant	
Error: Reference source not found	
Error: Reference source not found	
MULTNUM – Define the Multiple Transmissibility Regions	
MULTPV – Multiply Cell Pore Volumes by a Constant	
Error: Reference source not found	
Error: Reference source not found	
MULTX - Multiply Cell Transmissibility in the +X Direction	
MULTX Multiply Cell Transmissibility in the -X Direction	
MULTY - Multiply Cell Transmissibility in the +Y Direction	
MULTY Multiply Cell Transmissibility in the -Y Direction	
MULTZ - Multiply Cell Transmissibility in the +Z Direction	
MULTZ Multiply Cell Transmissibility in the -Z Direction	

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N

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Alphabetic Listing of Keywords Starting with the Letter N	Status
NETBALAN – Network Balancing Parameters	
NEWTRAN – Activate Irregular Corner-Point Grid Transmissibilities	
NEXTSTEP – Maximum Next Time Step Size	
NNC – Define Non-Neighbor Connections Between Cells Manually	
NOCASC – Activate Linear Solver Tracer Algorithm	
NOECHO – Deactivate Echoing of User Input Files to the Print File	
NOGGF – Deactivate Output of Grid Geometry File	
NOINSPEC – Deactivate Output of the INIT Index File	
NOMONITO – Deactivates Output of the Monitoring Data and File	
NONNC – Deactivates Non-Neighbor Connections	
NORSSPEC – Deactivate Output of the RESTART Index File	
NOSIM – Activates the No Simulation Mode for Data File Checking	
NOWARN – Deactivate Warning Messages	
NSTACK – Define the Stack Length for the Iterative Linear Solver	
NTG – Define the Net-to-Gross Ratio for All the Cells	
NUMRES – Define the Number of Reservoir Grids	
NUPCOL – Define the Number of Newtonian Iterations Used to Update Well Targets	

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RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter O	Status
OIL – Activate the Oil Phase in the Model	
OILDENT – Define Oil Density Temperature Coefficients	
OILVISCT – Define Oil Viscosity versus Temperature Functions	
OLDTRAN – Activate Cartesian Regular Grid Transmissibilities	
OPERATE – Define Mathematical Operations on Arrays	
OPERATER – Define Mathematical Operations on Arrays by Region	
OPERNUM – Define Regions for Mathematical Operations on Arrays	
OPTIONS – Activate Various Program Options	
OUTRAD - Define the Outer Radius of a Radial Grid	

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P

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Alphabetic Listing of Keywords Starting with the Letter P	Status
PARALLEL – Define Run Configuration	
PATHS – Define Filename Directory Path Aliases	
PBUB – Define the Initial Equilibration Bubble-Point Pressure for All Grid Blocks	
PBVD – Equilibration Bubble-Point versus Depth Tables	
PCG – End-Point Scaling of Grid Cell Maximum Gas Capillary Pressure (Drainage)	
PCW – End-Point Scaling of Grid Cell Water Capillary Pressure (Drainage)	
PDEW – Define the Initial Equilibration Dew-Point Pressure for All Grid Blocks	
PDVD – Define Equilibration Dew-Point versus Depth Tables	
PERMR – Define the Permeability for Each Cell in the R Direction	
PERMTHT – Define the Permeability for Each Cell in the THETA Direction	
PERMX - Define the Permeability in the X Direction for All the Cells	
PERMXX – Define the Permeability Tensor in the XX Direction for All the Cells	
PERMXY – Define the Permeability Tensor in the XY Direction for All the Cells	
PERMY - Define the Permeability in the Y Direction for All the Cells	
PERMYY – Define the Permeability Tensor in the YY Direction for All the Cells	
PERMYZ – Define the Permeability Tensor in the YZ Direction for All the Cells	
PERMZ - Define the Permeability in the Z Direction for All the Cells	
PERMZX – Define the Permeability Tensor in the ZX Direction for All the Cells	
PERMZZ – Define the Permeability Tensor in the ZZ Direction for All the Cells	
PIMTDIMS – Define Well Productivity Scaling Table Dimensions	
PIMULTAB – Define Well Productivity Index versus Water Cut Tables	
PINCH – Define Pinch-Out Layer Options	
PINCHNUM – Define Pinch-Out Regions for the PINCHREG Keyword	
PINCHREG - Define Pinch-Out Region Options	
PLMIXPAR – Define the Polymer Todd-Longstaff Mixing Parameters	
PLYADS - Define Polymer Rock Adsorption Tables	
PLYADSS - Define Polymer Rock Adsorption with Salt Dependence Tables	
PLYDHFLF - Define Polymer Thermal Degradation Half-Life Tables	

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RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter P	Status
PLYMAX - Define Polymer-Salt Viscosity Mixing Concentrations	
PLMIXNUM – Define the Polymer Region Numbers	
PLYROCK - Define Polymer-Rock Properties	1
PLYSHEAR – Activate and Define Polymer Shearing Parameters	
PLYSHLOG - Activate and Define the Polymer Shearing Logarithmic Parameters	
PLYVISC – Define Polymer Viscosity Scaling Factors	
PMISC – Define Miscibility versus Pressure Tables	
POLYMER – Activate the Polymer Phase in the Model	
PORO - Define the Porosity Values for All the Cells	
PORV - Define the Pore Volumes for All the Cells	
PPCWMAX – Define SWATINIT Calculated Capillary Pressure Constraints	
PRESSURE – Define the Initial Equilibration Pressures for All Grid Blocks	
PROPS - Define the Start of the PROPS Section of Keywords	
PRORDER – Define a Group Production Rules Sequence	
PVCDO - Oil PVT Properties for Dead Oil (Constant Compressibility)	
PVDG - Oil PVT Properties for Dead Oil (Constant Compressibility)	
PVDO – Oil PVT Properties for Dead Oil	
PVDS - Solvent PVT Properties for the Solvent Model	
PVTG - Gas PVT Properties for Wet Gas	
PVTNUM – Define the PVT Regions	
PVTO - Oil PVT Properties for Live Oil	
PVTW - Define Water Fluid Properties for Various Regions	

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Q

Revision: Rev-I

Alphabetic Listing of Keywords Starting with the Letter Q	Status
There Are No Keywords Beginning with the Letter Q	

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R

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Revision: Rev-I

Alphabetic Listing of Keywords Starting with the Letter R	Status
RADFIN – Define a Radial Local Grid Refinement with One Column	
RADFIN4 – Define a Radial Local Grid Refinement with Four Columns	
RADIAL – Radial Grid Activation Option	
REFINE – Start the Definition of a Local Grid Refinement	
REGDIMS – Define the Maximum Number of Regions for a Region Array	
REGIONS - Define the Start of the REGIONS Section of Keywords	
RESTART – Restart Run From an Existing Restart File	
RKTRMDIR - Activate ROCKTAB Keyword Directional Transmissibility Multipliers	
ROCK - Define the Rock Compressibility for Various Regions	
ROCKCOMP – Activate Rock Compaction	
ROCKNUM – Define Rock Compaction Table Region Numbers	
ROCKOPTS – Define Rock Compaction and Compressibility Options	
ROCKTAB – Rock Compaction Tables	
RPTGRID – Define GRID Section Reporting	
RPTONLY - Activate the Report Time Steps Only Option for the SUMMARY File	
RPTONLYO - Deactivate the Report Time Steps Only Option for the SUMMARY File	
RPTPROPS – Define PROPS Section Reporting	
RPTREGS – Define REGIONS Section Reporting	
RPTRST – Define Data to be Written to the RESTART File	
RPTRUNSP – Activates RUNSPEC Reporting	
RPTSCHED – Define SCHEDULE Section Reporting	
RPTSMRY - Activate or Deactivate Summary List Report	
RPTSOL – Define SOLUTION Section Reporting	
RS – Define the Initial Equilibration GOR (Rs) for All Grid Blocks	
RSCONST – Define Constant GOR for Dead Oil PVT Fluids	
RSCONSTT – Define Constant GOR for Dead Oil PVT Fluids	
RSVD – Equilibration Dissolved Gas-Oil Ratio (Rs) versus Depth Tables	
RTEMP - Define the Initial Reservoir Temperature for the Model	

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Alphabetic Listing of Keywords Starting with the Letter R	Status
RTEMPA - Define the Initial Reservoir Temperature for the Model	
RTEMPVD - Define the Initial Reservoir Temperature versus Depth Tables	
RUNSPEC -Define the Start of the RUNSPEC Section of Keywords	
RUNSUM – Activate RSM File Output of the SUMMARY Data	
RV – Define the Initial Equilibration CGR (Rv) for All Grid Blocks	
RVVD – Equilibration Vaporized Oil-Gas Ratio (Rv) versus Depth Tables	

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RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter S	Status
SALNODE – Salt Concentration Based PVTNUM Array	
SATNUM – Define the Saturation Table Region Numbers	
SATOPTS – Activate Relative Permeability Assignment Options	
SAVE – Activates Output of a SAVE File for Fast Restarts	
SCALECRS – Set End-Point Scaling Option	
SCHEDULE - Define the Start of the SCHEDULE Section of Keywords	
SDENSITY – Define the Miscible or Solvent Surface Gas Density	
SEPARATE – Activate the Separate RSM File Output Option	
SGAS – Define the Initial Equilibration Gas Saturation for All Grid Blocks	
SGCR – End-Point Scaling Grid Cell Critical Gas Saturations	
SGCRX SGCR – End-Point Scaling Grid Cell Critical Gas Saturations	
SGCRX- SGCR – End-Point Scaling Grid Cell Critical Gas Saturations	
SGCRY SGCR – End-Point Scaling Grid Cell Critical Gas Saturations	
SGCRY- SGCR – End-Point Scaling Grid Cell Critical Gas Saturations	
SGCRZ SGCR – End-Point Scaling Grid Cell Critical Gas Saturations	
SGCRZ- SGCR – End-Point Scaling Grid Cell Critical Gas Saturations	
SGCWMIS – Miscible Critical Gas versus Water Saturation Functions	
SGFN – Gas Saturation Tables (Format Type 2)	
SGL – End-Point Scaling Grid Cell Connate Gas Saturations	
SGLX SGL – End-Point Scaling Grid Cell Connate Gas Saturations	
SGLX- SGL – End-Point Scaling Grid Cell Connate Gas Saturations	
SGLY SGL – End-Point Scaling Grid Cell Connate Gas Saturations	
SGLY- SGL – End-Point Scaling Grid Cell Connate Gas Saturations	
SGLZ SGL – End-Point Scaling Grid Cell Connate Gas Saturations	
SGLZ- SGL – End-Point Scaling Grid Cell Connate Gas Saturations	
SGOF – Gas-Oil Saturation Tables versus Gas (Format Type 1)	
SGU – End-Point Scaling Grid Cell Gas Saturation	
SGUX SGU – End-Point Scaling Grid Cell Gas Saturation	

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RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter S	Status
SGUX- SGU – End-Point Scaling Grid Cell Gas Saturation	
SGUY SGU – End-Point Scaling Grid Cell Gas Saturation	
SGUY- SGU – End-Point Scaling Grid Cell Gas Saturation	
SGUZ SGU – End-Point Scaling Grid Cell Gas Saturation	
SGUZ- SGU – End-Point Scaling Grid Cell Gas Saturation	
SGWFN – Gas-Water Saturation Tables (Format Type 2)	
SHRATE - Activate and Define the Polymer Shearing Logarithmic Parameters	
SKIP – Activate Skipping of All Keywords and Input Date	
SKIP100 – Activate Skipping of "Black-Oil" Keywords and Input Date	
SKIP300 – Activate Skipping of "Compositional" Keywords and Input Date	
SKIPREST – Activate Skipping of Restart Schedule Data	
SLGOF – Gas-Oil Saturation Tables versus Gas (Format Type 1)	
SMRYDIMS – Define Maximum Number of Summary Vectors to be Written	
SOF2 – Oil Saturation Tables with Respect to Gas or Water (Format Type 2)	
SOF3 – Oil Saturation Tables with Respect to Gas and Water (Format Type 2)	
SOGCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Gas	
SOGCRX SOGCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Gas	
SOGCRX- SOGCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Gas	
SOGCRY SOGCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Gas	
SOGCRY- SOGCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Gas	
SOGCRZ SOGCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Gas	
SOGCRZ- SOGCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Gas	
SOIL – Define the Initial Equilibration Oil Saturation for All Grid Blocks	
SOLUTION - Define the Start of the SOLUTION Section of Keywords	
SOLVENT – Activate the SOLVENT Phase in the Model	
SORWMIS – Miscible Residual Oil versus Water Saturation Functions	
SOWCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Water	
SOWCRX SOWCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Water	

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RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter S	Status
SOWCRX- SOWCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Water	
SOWCRY SOWCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Water	
SOWCRY- SOWCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Water	
SOWCRZ SOWCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Water	
SOWCRZ- SOWCR – End-Point Scaling Grid Cell Critical Oil Saturation with Respect to Water	
SPECGRID- Define the Dimensions of a Corner-Point Grid	
SPECHEAT – Define the Specific Heat of Oil, Water and Gas	
SPECROCK – Define the Specific Heat of the Reservoir Rock	
SPOLY – Define the Initial Equilibration Polymer Concentration for All Grid Blocks	
SSFN – Solvent and Gas Relative Permeability Tables	
SSOL – Define the Initial Equilibration Solvent Saturation for All Grid Blocks	
START – Simulation Start Date	
STONE1 – Activate Stone's First Three Phase Oil Relative Permeability Model	
STONE1EX – Define Stone's First Three Phase Oil Relative Permeability Parameter	
STONE2 – Activate Stone's Second Three Phase Oil Relative Permeability Model	
SUMMARY - Define the Start of the SUMMARY Section of Keywords	
SUMTHIN – Define SUMMARY DATA Reporting Time Steps	
SWAT – Define the Initial Equilibration Water Saturation for All Grid Blocks	
SWATINIT – Define the Initial Water Saturation Array for Capillary Pressure Scaling	
SWCR – End-Point Scaling Grid Cell Critical Water Saturation	
SWCRX SWCR – End-Point Scaling Grid Cell Critical Water Saturation	
SWCRX- SWCR – End-Point Scaling Grid Cell Critical Water Saturation	
SWCRY SWCR – End-Point Scaling Grid Cell Critical Water Saturation	
SWCRY- SWCR – End-Point Scaling Grid Cell Critical Water Saturation	
SWCRZ SWCR – End-Point Scaling Grid Cell Critical Water Saturation	
SWCRZ- SWCR – End-Point Scaling Grid Cell Critical Water Saturation	
SWFN – Water Saturation Tables (Format Type 2)	
SWL – End-Point Scaling Grid Cell Connate Water Saturation	

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RUNSPEC GRID EDIT PROPS REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter S	Status
SWLX SWL – End-Point Scaling Grid Cell Connate Water Saturation	
SWLX- SWL – End-Point Scaling Grid Cell Connate Water Saturation	
SWLY SWL – End-Point Scaling Grid Cell Connate Water Saturation	
SWLY- SWL – End-Point Scaling Grid Cell Connate Water Saturation	
SWLZ SWL – End-Point Scaling Grid Cell Connate Water Saturation	
SWLZ- SWL – End-Point Scaling Grid Cell Connate Water Saturation	
SWOF – Water-Oil Saturation Tables (Format Type 1)	
SWU – End-Point Scaling Grid Cell Gas Saturation	
SWUX SWU – End-Point Scaling Grid Cell Gas Saturation	
SWUX- SWU – End-Point Scaling Grid Cell Gas Saturation	
SWUY SWU – End-Point Scaling Grid Cell Gas Saturation	
SWUY- SWU – End-Point Scaling Grid Cell Gas Saturation	
SWUZ SWU – End-Point Scaling Grid Cell Gas Saturation	
SWUZ- SWU – End-Point Scaling Grid Cell Gas Saturation	

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Alphabetic Listing of Keywords Starting with the Letter T	Status
TABDIMS – Define the Number of Tables and the Table Dimensions	
TEMP – Activate the Temperature Modeling Option	
TEMPI – Define the Initial Temperature Values for All Cells	
TEMPVD - Define the Initial Reservoir Temperature versus Depth Tables	
THCGAS – Define Gas Phase Thermal Conductivity for All Cells	
THCOIL – Define Oil Phase Thermal Conductivity for All Cells	
THCONR – Define Rock and Fluid Thermal Conductivity for All Cells	
THCONSF – Define Gas Saturation Dependent Thermal Conductivity Scaling Factor for All Cells	
THCROCK – Define Reservoir Rock Thermal Conductivity for All Cells	
THCSOLID – Define Solid Phase Thermal Conductivity for All Cells	
THCWATER – Define Water Phase Thermal Conductivity for All Cells	
THERMAL – Activate the Thermal Modeling Option	
THERMEXI – Define Component Thermal Expansion Coefficients	
THPRES - Define Equilibration Region Threshold Pressures	
TITLE – Define the Title for the Input Deck	
TLMIXPAR – Define the Miscible Todd-Longstaff Mixing Parameters	
TNUM – Define Passive Tracer Concentration Regions	
TOLCRIT – Define The Critical Saturation Tolerance	
TOPS - Define the Depth at the Center of the Top Face for Each Cell	
TRACER – Define Passive Tracer Variables	
TRACERS – Activate Tracer Options and Set Tracer Array Dimensions	
TRANX - Define the Transmissibility in the X Direction for All the Cells	
TRANY - Define the Transmissibility in the Y Direction for All the Cells	
TRANZ - Define the Transmissibility in the Z Direction for All the Cells	
TREF - Define Component Fluid Densities Reference Temperatures	
TREFS – Define Component Fluid Densities Reference Temperature at Surface	
TSTEP – Advance Simulation by Reporting Time	
TUNING - Numerical Tuning Control	

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Alphabetic Listing of Keywords Starting with the Letter T	Status
TUNINGDP – Numerical Tuning Control for High Throughput Cases	
TVDP – Define the Initial Equilibration Tracer Saturation versus Depth Functions	

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	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE	
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Alphabetic Listing of Keywords Starting with the Letter U	Status
UDADIMS – Define the Dimensions of the User Defined Arguments	
UDQ - Declare User Define Quantities ("UDQ")	
UDQDIMS – Define the Dimensions of the User Defined UDQ Feature	
UDQPARAM – Define Parameters for the User Defined Quantity Feature	
UNIFIN – Activates The Unified Input File Option	
UNIFOUT – Activates The Unified Output File Option	

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Regions Section Somman Solution	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter V	Status
VAPOIL – Activate the Vaporize Oil in Wet Gas Phase in the Model	
VAPPARS – Oil Vaporization Parameters	
VFPIDIMS – Injection Vertical Flow Performance Table Dimensions	
VFPINJ – Define Injection Vertical Flow Performance Tables	
VFPPDIMS – Production Vertical Flow Performance Table Dimensions	
VFPPROD – Define Production Vertical Flow Performance Tables	
VISCREF - Define Viscosity-Temperature Reference Conditions	

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RUNSPEC GR	GRID EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter W	Status
WARN – Activate Warning Messages	
WATDENT – Define Water Density Temperature Coefficients	
WATER – Activate the Water Phase in the Model	
WATVISCT – Define Water Viscosity versus Temperature Functions	
WCONHIST – Define Well Historical Production Rates and Pressures	
WCONINJ – Well Injection Targets and Constraints	
WCONINJE – Well Injection Targets and Constraints	
WCONINJH – Well Historical Observed Injection Rates and Pressures	
WCONPROD – Define Well Production Targets and Constraints	
WDFACCOR – Gas Flow Dependent Skin Factor	
WDRILTIM – Define Drilling Parameters for Automatic Drilling of New Wells	
WECON – Well Economic Criteria for Production Wells	
WEFAC – Define Well Efficiency	
WELCNTL – Modify Well Control and Targets	
WELLDIMS – Define the Wells and Group Dimensions	
WELOPEN – Define Well and Well Connections Flowing Status	
WELPI – Define Well Productivity and Injectivity Indices	
WELSEGS – Define Multi-Segment Wells and Their Segment Structure	
WELSPECL – Define Well Specifications for Local Grid Refinements	
WELSPECS – Define Well Specifications	
WELTARG – Modify Well Targets and Constraints Values	
WGASPROD – Define Sale Gas Well Production Targets	
WGRUPCON – Define Well Guides for Group Control	
WHISTCTL - Define Well Historical Target Phase	
WINJMULT – Define Well Pressure Dependent Injectivity Multipliers	
WINJTEMP – Define Injection Fluid Thermal Properties	
WLIFT – Define Well Re-Tubing, THP and Lift Switching Workover Operations	
WLIMTOL – Define Constraint Tolerance	

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Alphabetic Listing of Keywords Starting with the Letter W	Status
WORKLIM – Define Well Workover Time	
WORKTHP – Define Well Workover Options for THP Killed Wells	
WPAVE – Define Well Block Average Pressure Calculation Parameters	
WPAVEDEP – Define Well Reference Depth for Pressure Calculations	
WPIMULT – Define Well Connection Multipliers	
WPITAB - Assign Well Productivity Index versus Water Cut Tables	
WPOLYMER - Define Water Injection Well Polymer and Salt Concentrations	
WRFT – Activate Well RFT Reporting to the RFT File	
WRFTPLT – Activate Well RFT and PLT Reporting to the RFT File	
WSEGDIMS – Define Multi-Segment Well Dimensions	
WSEGITER – Define Multi-Segment Wells Iteration Parameters	
WSEGSICD – Define Multi-Segment Well Spiral ICD Connections	
WSOLVENT - Define Gas Injection Well Solvent Fraction	
WTEMP – Define An Injection Well's Fluid Temperature	
WTEST – Well Testing Criteria for Re-Opening Closed Wells	
WTRACER - Define An Injection Well's Tracer Concentration	

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Alphabetic Listing of Keywords Starting with the Letter X	Status
There Are Keywords Beginning with the Letter X	

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RUNSPEC GRID EDIT PROPS REGIONS SOLUTION SUMMARY SCHED	RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDULE
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Alphabetic Listing of Keywords Starting with the Letter Y	Status
There Are Keywords Beginning with the Letter Y	

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Z

RUNSPEC	GRID	EDIT	PROPS	REGIONS	SOLUTION	SUMMARY	SCHEDUL	
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Alphabetic Listing of Keywords Starting with the Letter Z				
ZCORN – Define the Depth of Each Corner-Point of a Grid Block				
ZIPPY2 – Activate Automatic Time Step Control				

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14 OPM FLOW RELEASE HISTORY

14.1 RELEASE 2018-10

The Open Porous Media project is please to announce that version 2018.10 of the OPM suite of simulation software has been released. Binary packages for Red-Hat Enterprise Linux 6 and 7 and Ubuntu 16.04 and 18.04 are available for download at the OPM website download page https://opm-project.org/?page_id=36. Installation instructions can also be found at the download page as well as in section 2 INSTALLING AND RUNNING FLOW of this manual.

In addition to the usual list of fixes and improvements two significant improvements include:

- I) OPM Flow can now write restart files which can be used to restart runs using the commercial simulator.
- 2) Performance has been improved significantly, on selected field models, where OPM Flow is significantly faster than the commercial simulator.

Due to problems with the Zoltan package we have unfortunately been forced to disable MPI for the Ubuntu 18.04 package. We are looking into this, and if possible we will make updated packages at a later stage.

In terms of new features OPM Flow now includes an option for thermal modeling. The energy "black-oil" implementation in OPM Flow is a mixture of the commercial simulators "black-oil" and the commercial simulators "compositional thermal" keywords, as well as some OPM Flow specific keywords. The temperature option (TEMP keyword) and the thermal option (THERMAL keyword) are two separate modeling facilities in the commercial simulator. OPM Flow's thermal implementation is based on solving the energy equation fully coupled with the "black-oil" equations so the results are not directly equivalent to commercial simulator's "black-oil" TEMP or compositional THERMAL formulations. See the 5.2.57 THERMAL-Activate the Thermal Modeling Option keyword in the RUNSPEC section outlining the available keywords.

Apart form the new thermal keywords summarized in section 5.2.57 THERMAL – Activate the Thermal Modeling Option, the following new keywords have been incorporated in this release and are active:

- 1) 6.3.36 GDFILE Load a Grid File.
- 2) 9.3.21 PLMIXNUM Define the Polymer Region Numbers.
- 3) 8.3.113 TOLCRIT Define The Critical Saturation Tolerance.
- 4) 12.3.103 WSEGSICD Define Multi-Segment Well Spiral ICD Connections.

Joakim Hove

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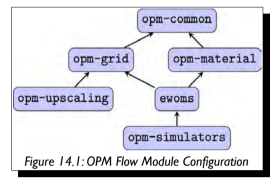
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14.2 Release 2018-04

The Open Porous Media project is please to announce that version 2018.04 of the OPM suite of simulation software has been release. Installation instructions can found on OPM website download page https://opm-project.org/?page_id=36 and in section 2 INSTALLING AND RUNNING FLOW of this manual. Packages for Ubuntu 16.04 and Red-Hat Enterprise Linux 6 and 7 is prepared. New in this release is that also the Red-Hat packages are shipped with mpi support.

In addition to various minor bug-fixes and a reorganization of the code, the release contains new functionality for the OPM Flow simulator like DRSDT support, experimental CPR preconditioning and enhanced output capability. Note that number of modules are reduced to simplify the build process and the code maintenance. The new module organization is shown opposite.

Since the 2017.10 release the opm-core module has been removed, and the modules opm-parser and opm-output have been folded into opm-common.



In terms of new features OPM Flow now includes analytical aquifers using the Carter-Tracy analytical aquifer and the ability model multi-segment wells. The following new keywords have been incorporated in this release and are active:

- 1) 6.3.4 AQUANCON Define Analytical Connections to the Grid.
- 2) 6.3.6 AQUCT Define Carter-Tracy Analytical Aquifers.
- 3) 5.2.4 AQUDIMS Aquifer Dimensions.
- 4) 8.3.10 AQUTAB Define Carter-Tracy Aquifer Influence Functions.
- 5) 12.3.14 COMPSEGS Define Well Connections for Multi-Segment Wells.
- 6) 12.3.16 DRSDT Solution Gas (Rs) Maximum Rate of Increase Parameters.
- 7) 12.3.18 DRVDT Solution Oil (Rv) Maximum Rate of Increase Parameters.
- 8) 12.3.82 WELSEGS Define Multi-Segment Wells and Their Segment Structure.
- 9) 5.2.70 WSEGDIMS Define Multi-Segment Well Dimensions.

Tor Harald Sandve

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14.3 Release 2017-10 Update 1

We have created an update for the 2017.10 release. It fixes a few bugs in Flow that could affect simulation results slightly and also lead to very bad performance when running Flow in parallel using MPI. The effect was most pronounced when running with more than 4 MPI processes.

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Binary packages for Ubuntu 16.04 and Red Hat Enterprise Linux 6 and 7 have been updated, and for most users your systems will ask to install the updated version or do it automatically.

For those who compile OPM from source, the release branches on GitHub have been updated and tagged with

release/2017.10/update1

The master branch of course includes the same fixes.

Atgeirr Flø Rasmussen

14.4 RELEASE 2017-10

On behalf of the OPM project, I'm happy to announce that version 2017.10 has been released. Packages for Ubuntu 16.04 and Red-Hat Enterprise Linux 6 and 7 have been prepared or should be available soon.

As usual, this release contains a multitude of new features and improvements. Most notable are probably that the Flow simulator is now considerably faster than in the 2017.04 release, Flow now supports the solvent and polymer black-oil extensions and that there now is freely available documentation for the file format that is used to specify the input.

Finally, as the release manager, I'd take the opportunity and thank everyone involved in making the release process for 2017.10 go as smoothly as it did.

Andreas Lauser

14.5 RELEASE 2017-04

The Open Porous Media project is glad to announce that version 2017.04 of the OPM suite of simulation software has been released! Installation instructions can found on our download page.

Certainly the most significant change of this release is the introduction of the "flow_ebos" simulator. Compared to the previous "flow" simulators, "flow_ebos" uses a different approach to linearize the nonlinear system of partial differential equations and as a result exhibits significantly better performance. The new simulator is intended to eventually fully replace the current family of "flow" simulators (i.e., "flow", "flow_mpi", "flow_solvent", "flow_polymer", etc.) and should already provide a proper superset of the capabilities of the "flow" simulator of previous OPM releases. For this reason, the name "flow" has been made an alias for "flow_ebos" in OPM 2017.04. If, for some reason, the previous "flow" simulator must be used, it is still shipped under the name "flow_legacy", but we strongly encourage you to send us bug reports if you encounter any case that can be simulated using "flow_legacy" but not using "flow_ebos".

Besides the introduction of "flow_ebos", plenty unit tests have been added, a plethora of bugs has been fixed, well handling has been considerably improved and now supports e.g. top-hole pressure controls and vertical flow performance tables, ECL output and restart capabilities have been made much more comprehensive, and all grid related functionality of opm-core has been moved to the opm-grid module in preparation of the former module's eventual retirement. In addition, a Docker container has been uploaded to Docker Hub to

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ease deployment for people who are into container technologies.

Last but not least, I would like to thank everyone who contributed to making the many changes of this release happen so smoothly.

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Modules involved in the release (maintainers are given in parenthesis):

- opm-data (Alf Birger Rustad)
- opm-common (Atgeirr Rasmussen, Bård Skaflestad, Arne Morten Kvarving, Joakim Hove, Robert Klöfkorn, Tor Harald Sandve, and Andreas Lauser)
- opm-parser (Joakim Hove)
- opm-output (Joakim Hove)
- opm-grid (Atgeirr Rasmussen, Robert Klöfkorn, and Bård Skaflestad)
- opm-material (Andreas Lauser, Robert Klöfkorn, and Tor Harald Sandve)
- opm-core (Atgeirr Rasmussen, Robert Klöfkorn, and Bård Skaflestad)
- ewoms (Andreas Lauser, Robert Klöfkorn, and Tor Harald Sandve)
- opm-simulators (Atgeirr Rasmussen, Robert Klöfkorn, Tor Harald Sandve, and Andreas Lauser)
- opm-upscaling (Arne Morten Kvarving, Atgeirr Rasmussen, and Bård Skaflestad)

Andreas Lauser

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End of Document

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