

Serpent – a Continuous-energy Monte Carlo Reactor Physics Burnup Calculation Code

March 6, 2013

User's Manual

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Preface

This documentation is a User's Manual for the Serpent continuous-energy Monte Carlo reactor physics burnup calculation code.¹ Code development started at the VTT Technical Research Centre of Finland in 2004, under the working title “Probabilistic Scattering Game”, or PSG. This name is used in all publications dated before the pre-release of Serpent 1.0.0 in October 2008. The name was changed to due to the various ambiguities related to the acronym. The code is still under development and this manual covers only the main functionality available in March 2013.

The official Serpent website is found at <http://montecarlo.vtt.fi>. Support and minor updates in the source code are currently handled via the Serpent mailing list, in which all users are encouraged to join by sending e-mail to: Jaakko.Leppanen@vtt.fi. Any feedback is appreciated, including comments, bug reports, interesting results and ideas and suggestions for future development. A discussion forum for Serpent users is found at <http://ttuki.vtt.fi/serpent>.

For a quick start, experienced Monte Carlo code users are instructed to view the lattice input examples in Chapter 11 starting on page 133.

¹For referencing the code, use either the website: “<http://montecarlo.vtt.fi>” or this report: “J. Leppänen. *Serpent – a Continuous-energy Monte Carlo Reactor Physics Burnup Calculation Code*. VTT Technical Research Centre of Finland. (March 6, 2013)”

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Chapter 1

Installing and Running Serpent

1.1 Compiling Serpent

The Serpent code is written in standard ANSI-C language. The code is mainly developed in the Linux operating system, but it has also been compiled and tested in MAC OS X and some UNIX machines.¹ The Monte Carlo method is a computing-intensive calculation technique and raw computing power has a direct impact on the overall calculation time. It should be taken into account that the unionized energy grid format used in Serpent requires more computer memory compared to other continuous-energy Monte Carlo codes. One gigabyte of RAM should be sufficient for steady-state calculations, but a minimum of 3 Gb is recommended for burnup calculation.

The source code is compiled simply by running the GNU Make utility.² The Makefile provides for detailed instructions and various options for different platforms. Serpent uses the GD open source graphics library [1] for producing some graphical output. If this library is not installed in the system, the source code must be compiled with the “NO_GFX_MODE” option. The compilation should not result in any errors or warning messages and it should produce an executable named “sss”. Any problems in installation should be reported by e-mail to: Jaakko.Leppanen@vtt.fi.

Code updates are provided to registered users by distributing the updated source files by e-mail. New files replace old ones and the code must be re-compiled for the changes to take effect.

¹The main platforms in PSG/Serpent development have been a 2.6 GHz dual-core AMD Opteron PC with 5 Gb RAM running Fedora Core 4 and an iBook G4 with 1.2 GHz PowerPC processor and 768 Mb RAM running OS X v10.4.

²For a detailed description of Makefiles, see: <http://www.gnu.org/software/make>.

1.2 Running the Code

All interaction between the code and the user is handled through one or several input files and various output files, as described in the following chapters. The code is run from the command line interface. The general syntax is:

```
sss <inputfile> [<options>]
```

where <inputfile> is the name of the main input file
<options> are the options

The input file is a standard text file containing the input description. The input can also be divided into several files which are referred to in the main file.

The available options are:

-version	print version information and exit
-replay	run the simulation using random number seed from previous calculation
-plot	terminate run after geometry plot
-testgeom <N>	test the geometry using <N> randomly sampled neutron tracks
-checkvolumes <N>	calculate Monte Carlo estimates for material volumes by sampling <N> random points
-mpi <N>	run simulation in parallel mode (see Sec. 1.3)
-disperse	generate random particle or pebble distribution files for HTGR calculations

The replay option forces the code to use the same random number seed as in a previous run. Without this option, the seed is taken from system time and written in a separate seed file (named <inputfile>.seed) for later use. The seed can also be set manually in the input using the “set seed” option.³

The geometry test option can be used for debugging the geometry in addition to the geometry plotter. The code randomly samples neutron tracks across the geometry and checks that the cells are correctly defined. Some input errors can be spotted using this option.

The volume checking option can be used to verify that the volumes used in the calculation are correct. The code is able to calculate cell volumes for simple lattice geometries, but some more complicated geometry types require the values to be set by the user. The volumes

³The results of a Monte Carlo calculation depend on the sequence of pseudo random numbers used during the simulation. This sequence is fixed by the random number seed and the calculation can be repeated using the same seed. The capability to reproduce the same simulation is important, for example, for debugging purposes. Some codes, such as MCNP [2], use a fixed seed value, which results in the same results every time the code is run. The Serpent code uses by default a different seed for each run and hence the results are different as well. This behavior can be overridden by the replay command line option or by setting the seed manually in the input file.

are used for normalizing reaction rates for detectors and burnup calculation. The number of random points should be large (at least 1,000,000) for good statistical accuracy.

The random particle / pebble distribution generator works by prompting the user information on the volume type and dimensions, particle data and packing fractions. The code then generates a distribution inside the desired volume without overlapping any particles. The data is written in a file using format that can be directly read into the explicit HTGR geometry model (See Sec. 3.8.2 on page 40). The option is available from code version 1.1.5 on.

IMPORTANT NOTES ON RUNNING THE CODE:

1. The seed file is overwritten by a new value each time the code is run without the replay option and the old seed is lost.

SEE ALSO:

1. Dividing the input into several files (Sec.2.2.3 on page 16)
2. Setting the random number seed manually (Sec. 5.18 on page 73)
3. Geometry plotter (Sec. 3.9 on page 42)
4. Setting material volumes manually (Sec. 4.1.2 on page 49)

1.3 Parallel Calculation

Serpent uses the Message Passing Interface (MPI) [3] for parallel calculation. To activate this capability the code must be compiled with the “PARALLEL_MPI” option (see the Makefile for details) and the MPI libraries must be installed on the system.

There are two options for running the code in the parallel calculation mode. The first option is to use the standard MPI tools, such as mpirun:

```
[user@host mpitest]$ mpirun -np 10 sss input
```

This command executes the calculation in 10 hosts as defined in the parallel environment.

The second option is to use the built-in MPI runner and define the number of tasks in the command line:

```
[user@host mpitest]$ sss -mpi 10 input
```

In this calculation mode, the code attempts to run mpirun on its own. This may require small modifications in the source code or may not work at all in some systems. The file path for mpirun is defined by the “MPIRUN_PATH” precompiler variable in the “header.h” source file.

IMPORTANT NOTES ON PARALLEL CALCULATION:

1. Parallel calculation is available from version 1.0.3 on.
2. When multiple tasks are sharing the same memory space, the size of allocated memory is also multiplied. This should be taken into account when setting the memory size in the compilation.
3. The methodology is still under development. The calculation lacks error tolerance and load sharing and the mode should be used only in systems consisting of identical hosts. Most of the MPI routines were directly adopted from PSG and features exclusively available in Serpent (including burnup calculation) are not thoroughly tested.

SEE ALSO:

1. The MPI standard: <http://www-unix.mcs.anl.gov/mpi/>
2. The mpirun script:
<http://www-unix.mcs.anl.gov/mpi/www/www1/mpirun.html>

1.4 Nuclear Data

The Serpent code reads continuous-energy interaction data from ACE format cross section libraries. The current installation package contains libraries based on JEF-2.2, JEFF-3.1, ENDF/B-VI.8 and ENDF/B-VII evaluated data files. Since the data format is shared with MCNP, alternative data for various isotopes should be readily available to most users. There are also several ACE format data libraries based on different evaluations publicly available through the OECD/NEA Data Bank [4]. New libraries can be produced from raw ENDF format data using the NJOY nuclear data processing system [5].

1.4.1 Data Types

Three types of cross sections are available in the data files. Continuous-energy neutron cross sections (type 1) are used for the actual transport simulation. The data contains all necessary reaction cross sections, together with energy and angular distributions, fission neutron yields and delayed neutron parameters.

The second data type is the dosimetry cross section (type 2). Dosimetry cross sections exist for a large variety of materials and may include derived reaction modes not commonly encountered in transport calculation. The data may consist of one or several partial cross sections, but all energy and angular distributions are omitted. The data can be used with detectors but not in physical materials included in the transport calculation.

Thermal scattering cross sections (type 3) are used to replace the low-energy free-gas elastic scattering reactions for some important bound moderator nuclides, such as hydrogen in water or carbon in graphite. Thermal systems cannot be modelled using free-atom cross sections without introducing significant errors in the spectrum and the results.

1.4.2 Directory File

The cross section data is accessed by using a separate directory file, which differs from the “xsdir” file commonly used with ACE format data. A conversion between the two formats can be made by running the “xsdirconvert” utility script, included in the installation package:

```
[user@host xsdata]$ xsdirconvert.pl data.xsdir >> data.xsdata
```

The Serpent directory file contains the data necessary for the code for locating the cross section libraries and forming the material compositions. Each line in the directory file has the following format:

```
<alias> <zaid> <type> <ZA> <I> <AW> <T> <bin> <path>
```

where

<alias>	is the name identifying the nuclide in the input file
<zaid>	is the actual nuclide name in the data
<type>	is the type of the data
<ZA>	is the isotope identifier ($1000 \cdot Z + A$)
<I>	is the isomeric state number (0 = ground state)
<AW>	is the atomic weight
<T>	is the nuclide temperature (in K)
<bin>	is the binary format flag (0 = ASCII, 1 = binary)
<path>	is the data path for the library

EXAMPLES:

```
1001.06c 1001.06c 1 1001 0 1.00783 600.0 0 /xs/1001_06.ace
H-1.06c 1001.06c 1 1001 0 1.00783 600.0 0 /xs/1001_06.ace
8016.06c 8016.06c 1 8016 0 15.99492 600.0 0 /xs/8016_06.ace
O-16.06c 8016.06c 1 8016 0 15.99492 600.0 0 /xs/8016_06.ace
40000.06c 40000.06c 1 40000 0 91.21963 600.0 0 /xs/40000_06.ace
Zr-nat.06c 40000.06c 1 40000 0 91.21963 600.0 0 /xs/40000_06.ace
92235.09c 92235.09c 1 92235 0 235.04415 900.0 0 /xs/92235_09.ace
U-235.09c 92235.09c 1 92235 0 235.04415 900.0 0 /xs/92235_09.ace
92238.09c 92238.09c 1 92238 0 238.05078 900.0 0 /xs/92238_09.ace
U-238.09c 92238.09c 1 92238 0 238.05078 900.0 0 /xs/92238_09.ace
95342.09c 95342.09c 1 95242 1 242.05942 900.0 0 /xs/95342_09.ace
Am-242m.09c 95342.09c 1 95242 1 242.05942 900.0 0 /xs/95342_09.ace
lwtr.03t lwtr.03t 3 0 0 0.00000 0.0 0 /xs/tmccs1
Np-237.30y 93237.30y 2 93237 0 239.10201 0.0 0 /xs/l1ldos1
```

```
93237.30y 93237.30y 2 93237 0 239.10201 0.0 0 /xs/l11ldos1
```

The alias is the nuclide name used in the input file and it may or may not be the same as the actual isotope name. The `xsdircconvert` tool writes two entries for each nuclide, one using the original name and another one using the element symbol and the isotope number. The data types are: 1 = continuous-energy, 2 = ACE dosimetry. 3 = thermal scattering. The temperature entry is used with transport data only and the atomic mass with transport and dosimetry cross sections.

Isomeric states are identified from the state number⁴ (see Am-242m in the example). There is no standard convention on how to name these isotopes in the ACE format data, but the `xsdircconvert`-tool assumes that the mass number of isomeric state nuclides is increased above 300. If another convention is used, the state number must be set manually in the directory file. It is recommended that the isomeric state entries are always carefully checked after running `xsdircconvert`.

1.4.3 Radioactive Decay and Fission Yield Data

Radioactive decay and fission yield data is needed for running the Serpent code in the independent burnup calculation mode. It is recommended that the libraries are included in the coupled mode as well, since it enables the data to be reproduced in the output file, making it directly available to the coupled calculation.

The decay constants and fission product distributions are read from standardized ENDF format data files [6]. The format is directly accessible and the data requires no preprocessing. JEF-2.2, JEFF-3.1, ENDF/B-VI.8 and ENDF/B-VII data libraries are included in the installation package. More data can be downloaded from various Internet sources:

- OECD/NEA Data Bank: <http://www.nea.fr/html/dbdata/>
- Los Alamos T2 Nuclear Information Service: <http://t2.lanl.gov>
- US National Nuclear Data Center: <http://www.nndc.bnl.gov>
- US Radiation Safety Information Computational Center:
<http://www-rsicc.ornl.gov>
- IAEA Nuclear Data Centre: <http://www-nds.iaea.org>
- JAEA Nuclear Data Center: <http://wwwndc.tokai-sc.jaea.go.jp>

IMPORTANT NOTES ON INTERACTION DATA:

⁴The information on isomeric states is needed for burnup calculation only. All nuclides are treated similarly in the transport simulation.

1. The weight in the directory file is given as the *atomic weight*, not the atomic weight ratio as in MCNP xsdir files.
2. The temperature in the directory file is given in *Kelvin*, not in MeV as in the MCNP xsdir files.
3. Binary data is not supported in the current code version.
4. The data path in the directory file must refer to the absolute, not the relative location of the library file.
5. The code always uses the first matching entry in the directory file. The use of duplicate isotope names may lead to unexpected results.

SEE ALSO:

1. Setting up the file paths (Sec.5.4 on page 57)
2. Material definitions (Chapter 4 on page 47)

Chapter 2

Input

2.1 General

The Serpent code has no interactive user interface. All communication between the code and the user is handled through one or several input files and various output files discussed in Chapter 6. User-defined detectors are discussed as a separate item in Chapter 7 and burnup calculation in Chapter 8.

2.2 Input format

The format of the input file is unrestricted. The file consists of white-space (space, tab or newline) separated words, containing alphanumeric characters('a-z', 'A-Z', '0-9', '.', '-'). If special characters or white spaces need to be used within the word (file names, etc.), the entire string must be enclosed within quotation marks.

2.2.1 Input cards

The input file is divided into separate data blocks, denoted as cards. The file is processed one card at a time and there are no restrictions in what order the cards should be organized. The input cards are listed in Table 2.1 and detailed descriptions are provided in the following chapters. All input cards and special command words are case-insensitive. Each input card is delimited by the beginning of the next card. It is hence important that none of the parameter strings used within the card coincide with the card identifiers in Table 2.1.

Table 2.1: List of commands and input cards

Card	Description	Chapter / Section	Page
cell	cell definition	3.3	24
dep	irradiation history	8.3	110
det	detector definition	7.1	95
disp	implicit HTGR particle fuel model	3.8.1	39
ene	detector energy binning	7.1.2	99
include	read a new input file	2.2.3	16
lat	lattice definition	3.6.2	30
mat	material definition	4.1.2	48
mesh	reaction rate mesh plotter	10.1	131
nest	nest definition	3.5	27
particle	particle definition	3.8	39
pbed	explicit HTGR particle / pebble bed fuel model	3.8.2	40
pin	pin definition	3.4	27
plot	geometry plotter	3.9	42
set	misc. parameter definition	5.1	53
src	external source definition	9.2	126
surf	surface definition	3.2	20
therm	thermal scattering data definition	4.2	49
trans	universe transformation	3.6.1	29

2.2.2 Comment lines and sections

The Serpent code provides two types of comments for the input files. The percent-sign (%) or hash (#) are used to define a comment line. Anything from this character to the end of the line is omitted when the input file is read. The alternative is to use C-style comment sections beginning with “/*” and ending with “*/”. Everything within these delimiters is omitted, regardless of the number of newlines or special characters between them.

2.2.3 Dividing the input into several files

Complicated input descriptions can be simplified by dividing the cards into separate files. This capability may also be useful if different calculation cases share some partial data. Additional input files are recursively read from the main file using the `include`-command:

```
include "<filename>"
```

where `<filename>` is the file path for the input file

When this command is encountered, the program will first read the included file before

continuing with the main file. The number of nested input files is unrestricted. Since file names and paths often include non-alphanumeric characters, it is good practice to always enclose the strings within quotation marks.

2.2.4 Input errors

The Serpent code performs some error checking on the input file before proceeding with the calculation. These checks include:

- Checking that there are an even number of quotation marks.
- Checking the correct number of parameters for some input cards.
- Checking the type (string, integer, real) of some parameters.
- Checking that the values of some parameters are within a reasonable range.
- Checking that all cards that are referred to in other cards are defined.
- Checking that all referred files exist.
- Checking that the input contains sufficient data for running the simulation.
- Various checks related to specific input cards.

Failure in any of the checks results in an error message and the termination of the calculation.

Most common input errors are caused by missing parameters or mistyped command words. In the former case, the result is often an error message related to parameter type or number. The program does not recognize card names with typing errors, but rather processes the entire card as if was a set of parameters belonging to the previous card. Such errors may stop the calculation later on for entirely different reasons, or in the worst case, run the simulation with a set of parameters totally different from what the user intended. In case of any unexpected behavior, the typing of the card names should be the first thing to be checked.

IMPORTANT NOTES ON INPUT FORMAT:

1. The input file consists of white-space separated words containing alphanumeric characters. If special characters or white spaces need to be used (file names, etc.), the entire string must be enclosed within quotation marks.
2. Each card is delimited by the beginning of the next card and it is hence important that the card names are not used in for other purposes, for example as cell or material names. If the name of an input card is spelled incorrectly, the previous card is not delimited, which may result in a completely unexpected behavior.

3. Running the Serpent code should *never* result in crash or termination without an error message. In such case, please report the problem by e-mail to Jaakko.Leppanen@vtt.fi.

2.3 Units

Table 2.2 summaries the most essential units used in the code.

Table 2.2: Units used in the Serpent code.

Quantity	Unit	Notes
Distance	cm	
Area	cm ²	
Volume	cm ³	
Time	s	(depends on the case)
Energy	MeV	
Microscopic cross section	b	(barn = 10 ⁻²⁴ cm ²)
Macroscopic cross section	1/cm	
Mass	g	
Mass density	g/cm ³	
Atomic density	10 ²⁴ /cm ³	(= 1/barn × cm)
Power	W	
Power density	kW/g	
Neutron flux	1/cm ² s	
Reaction rate	1/cm ³ s	(reaction rate density)
Burnup	MWd/kgU	(per total initial heavy metal)
Burn time	days	

IMPORTANT NOTES ON UNITS:

1. Power, neutron flux, reaction rate and all related quantities depend on how the neutron source rate is normalized.

SEE ALSO:

1. Source rate normalization (Sec. 5.8 on page 61)

Chapter 3

Geometry

3.1 The Universe-based Geometry Model in Serpent

The Serpent code uses a universe-based geometry model for describing complicated structures, very similar to MCNP. This means that the geometry is divided into separate levels, which are all constructed independently and nested one inside the other. This approach allows the complexity of the geometry to be divided into smaller parts, which are much easier to handle. It also enables the use of regular geometry structures, such as square and hexagonal lattices, commonly encountered in reactor applications.

Perhaps the best example of a universe-based geometry construction is the reactor core. At the highest level, the geometry consists of fuel pins, in which the fuel pellets are surrounded by cladding and coolant. Each pin type is described independently in its own universe. The next level is the fuel assembly, in which the pin universes are arranged in a regular lattice. The assembly may also comprise flow channel walls, moderator channels or any support structures. In the next geometry level these assembly universes are arranged in another lattice to form the core layout, which can be surrounded by radial and axial reflectors and finally the reactor pressure vessel wall.

The basic building block of the geometry is the cell, which is a region of space determined using simple boundary surfaces. Each cell is filled with a homogeneous material composition, void or another universe.

3.2 Surface Definitions

Serpent provides for various elementary and derived surface types for geometry construction. A “derived” surface type refers here to a surface comprised of two or more elementary surfaces, such as a cube constructed of six planes. The input format does not make any dif-

ference between elementary and derived surfaces and the description below applies to both.

The syntax of the surface card is:

```
surf <id> <type> <param 1> <param 2> ...
```

where	<id>	is the surface identifier
	<type>	is the surface type (see Table 3.1)
	<param 1> <param 2> ...	are the surface parameters

The surface identifier is an arbitrarily chosen number identifying the surface in the cell definitions. Surface types and their use is described in the following subsections.

3.2.1 Surface types

The present code version contains 20 surface types, listed in Table 3.1. The number of parameters is fixed and depends on the type. Some surface types have parameters that are optional.

For the three types of planes, the x_0 , y_0 and z_0 coordinates refer to distances from the origin. For sphere, cube and the cylindrical surfaces these parameters define the coordinates of the surface center. Sphere, cube and cylinder radii are given by r . The square, hexagonal and cruciform cylinders also include an optional parameter r_0 , which defines the radius of rounded corners. If this parameter is omitted, it is assumed that the corners are sharp. The optional parameters z_1 and z_2 are bottom and top planes of truncated cylinder. The cylindrical surfaces are illustrated in Figure 3.1.

The cuboid is defined by the minimum and maximum coordinates in each direction.

The hexagonal prismatic surfaces are similar to the corresponding cylinders, with the difference that the enclosed space is limited by bottom and top planes at z_1 and z_2 .

The “pad” is a cylindrical surface type that was included in the code in order to model the neutron pad in the VENUS-2 reactor dosimetry benchmark [7]. The surface is defined as a sector between angles θ_1 and θ_2 cut out from a layer between cylinders of radii r_1 and r_2 .

The “cone” or “conz” surface type (see Fig. 3.2) is determined by the x_0 , y_0 and z_0 coordinates of the base, the base radius r and the height h . The height of the cone also determines the orientation: a positive value for a cone pointing in positive direction and a negative value for a cone pointing in the negative direction of the z-axis. Cones oriented in the x- and y-axes (“conx” and “cony”, respectively) are defined in a similar manner.

The “dode” and “octa” surface types (see Fig. 3.3) are determined by the x_0 and y_0 coordinates of the central axis and two distances r_1 and r_2 from the center. If the second value is omitted, the surface is a regular octa- or dodecagonal cylinder. The octagonal cylinder basically consists of two intersecting square and the dodecagonal surface of two intersecting

Table 3.1: Surface types in the Serpent code.

Type	Description	Parameters
inf	all space	-
px	plane perpendicular to x-axis	x_0
py	plane perpendicular to y-axis	y_0
pz	plane perpendicular to z-axis	z_0
sph	sphere	x_0, y_0, z_0, r
cylx	circular cylinder parallel to x-axis	y_0, z_0, r, x_1, x_2
cyly	circular cylinder parallel to y-axis	x_0, z_0, r, y_1, y_2
cylz or cyl	circular cylinder parallel to z-axis	x_0, y_0, r, z_1, z_2
sqc	square cylinder parallel to z-axis	x_0, y_0, r, r_0
cube	cube	x_0, y_0, z_0, r
cuboid	cuboid	$x_1, x_2, y_1, y_2, z_1, z_2$
hexxc	x-type hexagonal cylinder parallel to z-axis	x_0, y_0, r, r_0
hexyc	y-type hexagonal cylinder parallel to z-axis	x_0, y_0, r, r_0
hexxprism	x-type hexagonal prism parallel to z-axis	x_0, y_0, r, z_1, z_2
hexyprism	y-type hexagonal prism parallel to z-axis	x_0, y_0, r, z_1, z_2
cross	cruciform cylinder parallel to z-axis	x_0, y_0, r, d, r_0
pad	(see description below)	$x_0, y_0, r_1, r_2, \theta_1, \theta_2$
conx	cone oriented in the x-axis	x_0, y_0, z_0, r, h
cony	cone oriented in the y-axis	x_0, y_0, z_0, r, h
conz or cone	cone oriented in the z-axis	x_0, y_0, z_0, r, h
dode	dodecagonal cylinder parallel to z-axis	x_0, y_0, r_1, r_2
octa	octagonal cylinder parallel to z-axis	x_0, y_0, r_1, r_2
plane	general plane	A, B, C, D
quadratic	general quadratic surface	$A, B, C, D, E, F, G, H, J, K$

regular hexagons.

The general plane is defined by equation

$$Ax + By + Cz = D$$

This is a simplified case of the general quadratic surface, defined by

$$Ax^2 + By^2 + Cz^2 + Dxy + Eyz + Fzx + Gx + Hy + Jz + K = 0$$

3.2.2 Positive and negative surface sides

The surfaces are used for defining the geometry cells as will be described in the following section. For this purpose, each surface is associated with a positive side and a negative side. It is defined that a point is inside a surface if it is located on the negative side of the surface.

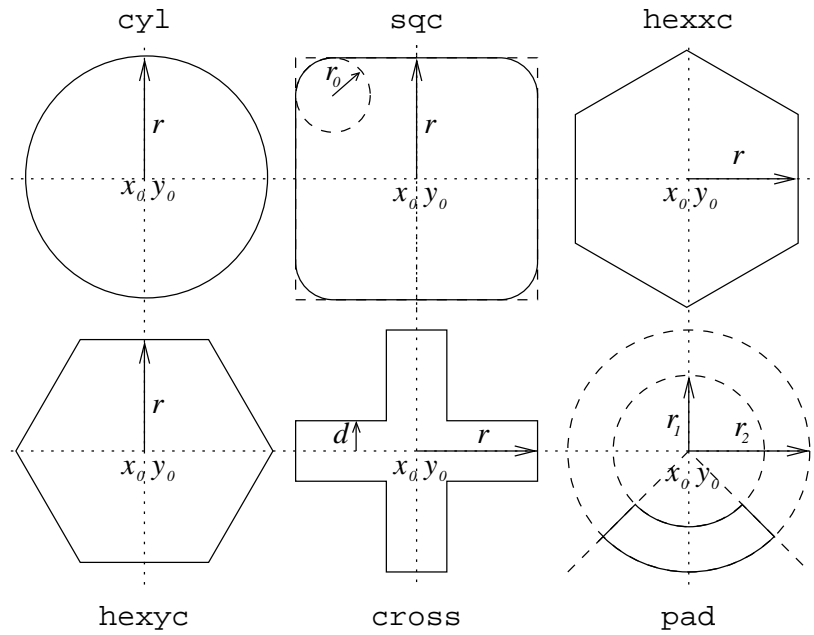


Figure 3.1: Basic cylinder types. The surfaces are infinite in the z -direction. The square cylinder illustrates the definition of rounded corners.

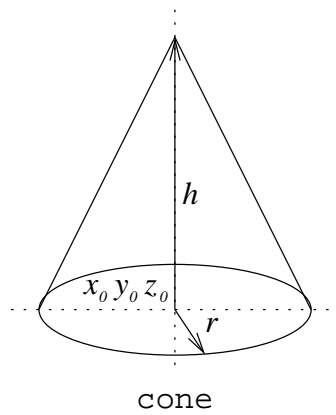


Figure 3.2: The cone surface.

For the three types of planes, the positive side is defined in the direction of the positive coordinate axis. The positive sides of the sphere, cube, cone and the cylindrical surfaces are defined outside the perimeter of the surface.

3.2.3 Surface examples

A few simple examples of surface definitions are given in the following.

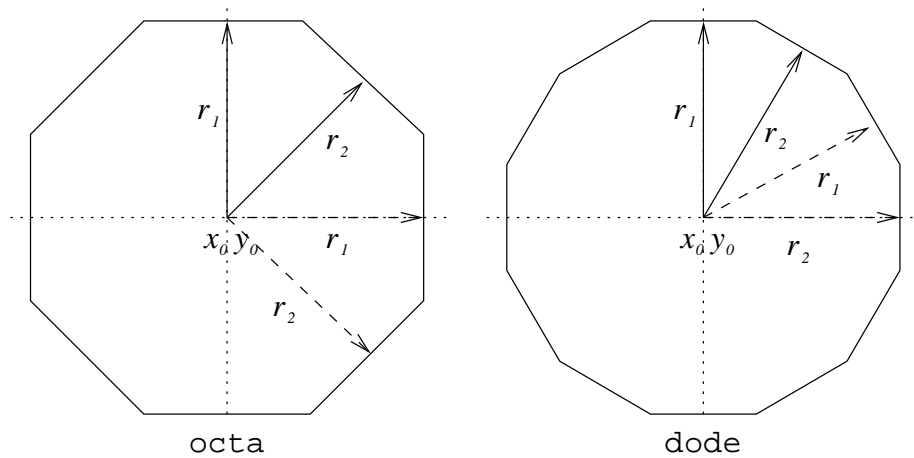


Figure 3.3: The octagonal and dodecagonal cylinder surfaces.

```
% --- plane perpendicular to x-axis, located at x = 4.0 cm:
surf 1 px      4.000

% --- sphere centered at (1.0, 0.0, 2.0), radius 5.0:
surf 2 sph     1.000  0.000  2.000  5.000

% --- cylinder centered at origin, radius 10.5 cm:
surf 3 cyl     0.000  0.000 10.500

% --- cube at origin with diameter 5.0 cm:
surf 4 cube    0.000  0.000  0.000  2.500

% --- square cylinder centered at origin, radius 10.0 cm,
%       rounded corners with radii 0.2 cm:
surf 5 sqc     0.000  0.000 10.000  0.200

% --- x-type hexagonal cylinder centered at (1.0, 0.0),
%       radius 2.0 cm:
surf 6 hexxc   1.000  0.000  2.000

% --- cruciform cylinder centered at origin, radius 20.0 cm,
%       half-thickness 5.0 cm:
surf 7 cross   0.000  0.000 20.000  5.000

% --- neutron pad used in the VENUS-2 benchmark:
surf 8 pad     0.000  0.000 11.250 54.750 59.073 65.073

% --- cone at origin, base diameter 2.0 cm, height 5.0 cm
surf 9 cone    0.000  0.000  0.000  1.000  5.000
```

IMPORTANT NOTES ON SURFACES:

1. In code versions earlier than 1.1.8 the cone surface type may only be used with the full delta-tracking calculation mode (threshold = 1).
2. Reflective and periodic boundary conditions may only be used in geometries where the outermost boundary is defined by a square or hexagonal cylinder or a cube.
3. The dodecagonal cylinder surface type is available from code version 1.1.4 on.
4. The octagonal cylinder and general plane and quadratic surface are available from code version 1.1.9 on.

SEE ALSO:

1. Delta-tracking options (Sec. 5.11 on page 66)
2. Boundary conditions (Sec. 5.7 on page 59)

3.3 Cell Definitions

The geometry description in the Serpent code consists of two- or three-dimensional regions, denoted as cells. Each cell is defined using a set of positive and negative surface numbers, which correspond to the surface identifiers defined in the surface cards. Unlike MCNP and other Monte Carlo codes, Serpent can only handle intersections of boundary surfaces. This means that *the neutron is inside the cell, if and only if it is on the same side of each boundary surface as given in the surface list* (see the examples below).

The lack of the union operator restricts the generality of the geometry description to some extent. This limitation is compensated for by providing a large collection of derived surface types, which in most cases can be used to replace the unions of the elementary surfaces. The advantage of this approach is that the geometry description remains relatively simple.¹

3.3.1 Cell types

The syntax of the cell card is:

¹ It is known that the use of derived surface types may slow down the neutron tracking routine in some cases when the conventional ray-tracing algorithm is used. Neutron transport in Serpent, however, is primarily based on the delta-tracking method which is not prone to such limitations. The use of derived surface types reduces the total number of surfaces, which may actually speed up the delta-tracking routine in complicated geometries.


```
cell <name> <u0> <mat> <surf 1> <surf 2> ...
```

where	<name>	is the cell name
	<u0>	is the universe number of the cell
	<mat>	is the cell material
	<surf 1> <surf 2> ...	are the boundary surfaces

The cell name is a text string that identifies the cell.² Each cell belongs to a universe, which is determined by the universe number (lattices and universes are thoroughly described in Section 3.6 on page 28). Cell material determines the name of the material that fills the cell (see Chapter 4 for material definitions). There are three exceptions:

1. If the cell is empty, the material name is set to “void”.
2. If the cell describes a region of space that is not part a of the geometry, the material name is set to “outside”.
3. If the cell is filled by another universe, the material name is replaced by command “fill” and the number of the filling universe.

The “outside” cells are required for filling the regions of space that are not a part of the actual geometry. When the neutron streams to such a region, the history is terminated or boundary conditions are applied.

The cell shape is determined by the list of boundary surfaces. Positive entries refer to positive (“outside”) surface sides and negative entries to negative (“inside”) surface sides. The cell is defined as the intersection of all surfaces in the list.

3.3.2 Cell examples

A few simple examples of cell definitions are given in the following.

```
% --- two half-planes separated by a plane in the z-axis at 5.0 cm:

surf 1 pz    5.000

cell 1 1  water  -1  % lower half-plane filled with "water"
cell 2 1  air    1   % upper half-plane filled with "air"

% --- solid uranium sphere ("Godiva") of radius 8.7407 cm:
```

²When the number of cells in the geometry is large, it is often easier to replace cell names with numerical constants. This is possible since the code treats cell numbers as any other text strings. This convention is followed in most example cases in this manual.

```

surf 1 sph  0.0  0.0  0.0  8.7407

cell 1 0  uranium  -1  % uranium inside sphere
cell 2 0  outside   1  % outside world

% --- tungsten-reflected plutonium sphere:

surf 1 sph  0.0  0.0  0.0  5.0419
surf 2 sph  0.0  0.0  0.0  9.7409

cell 1 0  plutonium  -1      % plutonium inside surface 1
cell 2 0  tungsten    1 -2    % tungsten between surfaces 1 and 2
cell 3 0  outside     2      % outside world

% --- a segment of LWR fuel rod in water:

surf 1 cyl  0.0  0.0  0.40
surf 2 cyl  0.0  0.0  0.45
surf 3 cyl  0.0  0.0  0.60
surf 4 pz  -50.0
surf 5 pz   50.0

cell 1 1  UO2          -1      4 -5  % UO2 fuel inside surface 1
cell 2 1  void          1 -2    4 -5  % gap between fuel and cladding
cell 3 1  clad          2 -3    4 -5  % cladding
cell 4 1  water         3       4 -5  % water outside cladding
cell 5 1  water        -4       % water below the segment
cell 6 1  water         5       % water above the segment

```

IMPORTANT NOTES ON CELLS:

1. Material names “void”, “outside” and “fill” are reserved for empty cells, cells not belonging to the geometry and cells filled by another universe, respectively.
2. Only the intersection operator is available for cell definitions. This means that a point is inside the cell if and only if it is inside (or outside if defined by a negative surface number) *all* the boundary surfaces in the list.

SEE ALSO:

1. Material definitions (Chapter 4 on page 47)
2. Boundary conditions (Sec. 5.7 on page 59)

3.4 Fuel pin definitions

Since Serpent is primarily a lattice physics code, the geometry has a simplified definition for fuel pins consisting of nested annular material layers. The syntax of the pin card is:

```
pin <id>
  <mat 1> <r1>
  <mat 2> <r2>
  ...
  <mat n>
```

where <id> is the pin identifier (universe number)
 <mat 1> <mat 2> ... are the materials
 <r1> <r2> ... are the outer radii of the material regions

The fuel pin is not an actual geometry object, but rather a macro that is used to define a pin universe. The material regions and their outer radii are given in ascending and the code constructs the cells using cylindrical surfaces. If the radius is negative, it is interpreted as layer thickness instead of absolute radius. The universe number is set by the pin identifier. Pin materials can also be other universes, which are defined using the fill command (See filled cells on page 28).

Pin definitions are most commonly used with lattices to define fuel assemblies. Examples are given in the following section.

IMPORTANT NOTES ON PIN DEFINITIONS:

1. The pin identifier is a universe number, which must not coincide with another universe.
2. The outermost material regions is given without a radius and it fills the rest of the universe.
3. Layer thickness are available from version 1.1.13 on.

SEE ALSO:

1. Filled cells (Sec. 3.6 on page 28)
2. Lattice examples (Sec. 3.6.3 on page 32)

3.5 Nests

Fuel pin and particle (see Sec. 3.8 on page 39) are special cases of the nest geometry type, defined as:

```

nest <id> <type>
<mat 1> <r1>
<mat 2> <r2>
...
<mat n>

```

where	<id>	is the nest identifier (universe number)
	<type>	is the surface type
	<mat 1> <mat 2> ...	are the materials
	<r1> <r2> ...	are the surface parameters

Nested objects consist of materials or sub-universes separated by similar surfaces. Nests can be defined using planar (`px`, `py`, `pz`), cylindrical (`cyl`, `sqc`, `hexxc`, `hexyc`), spherical (`sph`) or cubical (`cube`) surface types. In each case the parameters `<r1>`, `<r2>`, ... define the main dimension, all other parameters are set to zero.

3.6 Universes and Lattices

As mentioned above, a universe-based geometry allows the geometry to be divided into separate levels. Each universe is defined independently and must cover all space. Regions of space not belonging to the geometry must be defined using “outside” cells. The universes are defined by the cell universe numbers and the geometry is layered by replacing the material name with the fill command:

```
cell <name> <u0> fill <u1> <surf 1> <surf 2> ...
```

where	<name>	is the cell name
	<u0>	is the universe number of the cell
	<u1>	is the universe number of the filling universe
	<surf 1> <surf 2> ...	are the boundary surfaces

Each universe has its own origin, which can be shifted using the universe transformation command (see Sec. 3.6.1) The lowest level of the geometry belongs to universe 0, which must always exist.

3.6.1 Universe transformations and rotations

Each universe is by default centered at the origin, which may sometimes cause difficulties with filled cells. The origin can be shifted to another location using the universe transformation card:

```
trans <u> <x> <y> <z>
```

where <u> is the universe number
 <x> is the x coordinate of the new origin
 <y> is the y coordinate of the new origin
 <z> is the z coordinate of the new origin

Universe transformations are also convenient, for example, for positioning control rods in a reactor core. Universes filled in a lattice structure are automatically shifted to the appropriate position and transformations are not needed.

Universe rotations were implemented in Version 1.1.14. The syntax of the transformation card with rotations has two alternative formats:

```
trans <u> <x> <y> <z> <rx> <ry> <rz>
trans <u> <x> <y> <z> <a1> ... <a9>
```

where <u> is the universe number
 <x> is the x coordinate of the new origin
 <y> is the y coordinate of the new origin
 <z> is the z coordinate of the new origin
 <rx> is the rotation angle around x-axis
 <ry> is the rotation angle around y-axis
 <rz> is the rotation angle around z-axis
 <ai> are the coefficients of a rotation matrix

If three values are entered after the coordinates, they are interpreted as rotation angles around the three coordinate axes. If nine values are entered, they form the rotation matrix, which is used to multiply the position and direction vectors when the rotation is applied.

The coordinate translation always precedes the rotation.

3.6.2 Lattices

Lattices are special universes, filled with a regular structure of other universes. The Serpent code has eight lattice types: square lattice, two hexagonal lattices, the circular cluster array, three infinite 3D lattices filled with a single universe and the vertical stack.

Square and hexagonal lattices

The syntax of the lattice card for the square and hexagonal lattices is:

```
lat <u0> <type> <x0> <y0> <nx> <ny> <p>
```

where <u0> is the universe number of the lattice
 <type> is the lattice type (= 1, 2 or 3)
 <x0> is the x coordinate of the lattice origin
 <y0> is the y coordinate of the lattice origin
 <nx> is the number of lattice elements in the x-direction
 <ny> is the number of lattice elements in the y-direction
 <p> is the lattice pitch

The lattice card is followed by a list of universe numbers, which determines the layout. The lattice type numbers are:

1. Square lattice
2. X-type hexagonal lattice (unit cell is the x-type hexagonal cylinder, see Fig. 3.1)
3. Y-type hexagonal lattice (unit cell is the y-type hexagonal cylinder, see Fig. 3.1)

Each lattice defines a universe, which must be embedded inside a cell using the fill command. If the bounding cell is larger than the lattice, neutrons may stream to undefined lattice positions, which results in a geometry error. This can be avoided by increasing the lattice size by defining additional positions in the periphery (see examples below).

Circular cluster array

The circular cluster array (lattice type 4) is defined by:

```
lat <u0> <type> <x0> <y0> <nr>
```

where <u0> is the universe number of the lattice
 <type> is the lattice type (= 4)
 <x0> is the x coordinate of the lattice origin
 <y0> is the y coordinate of the lattice origin
 <nr> is the number of rings in the array

The lattice card is followed by a list of <nr> rings, which are defined by:

```
<n> <r> <theta> <u1> <u2> ... <un>
```

where <n> is the number of sectors in ring
 <r> is the central radius of the ring
 <theta> is the angle of rotation
 <u1> <u2> ... <un> are the universe numbers filling the sectors

The circular array is needed for constructing some cluster-type fuel assemblies, used in CANDU, MAGNOX, AGR and RBMK reactors. The array is also convenient for determining the fuel rod layout in some small research reactors, such as the TRIGA.

Infinite 3D lattices

The infinite 3D lattices are used to construct repeated structures of identical cells that fill the entire universe. This type of construction can be used, for example, for describing the microscopic fuel particles inside an HTGR fuel pebble or compact. The syntax is:

```
lat <u0> <type> <x0> <y0> <p>
```

where <u0> is the universe number of the lattice
 <type> is the lattice type (= 6, 7 or 8)
 <x0> is the x coordinate of the lattice origin
 <y0> is the y coordinate of the lattice origin
 <p> is the lattice pitch
 <u> is the filler universe

Lattice type 6 is a cubical lattice and types 7 and 8 x- and y-type hexagonal prismatic lattices, respectively.

Vertical stack

Universes can be vertically stacked, one on top of the other, using lattice type 9:

```
lat <u0> <type> <x0> <y0> <n1>
```

where <u0> is the universe number of the lattice
 <type> is the lattice type (= 9)
 <x0> is the x coordinate of the lattice origin
 <y0> is the y coordinate of the lattice origin
 <n1> is the number of axial layers

The lattice card is followed by a list of <n1> axial layers, which are defined by:

```
<z> <u>
```

where <z> is the axial position (lower boundary of the layer)
 <u> is the filler universe

The z-values must be given in ascending order. Space below the lowest z-value is not defined and the top layer fills the entire space above the highest value.

Cuboidal 3D lattice

The cuboidal lattice is a 3D structure composed of cuboids with different dimensions in the x-, y- and z-directions. The syntax is:

```
lat <u0> <type> <x0> <y0> <z0> <nx> <ny> <nz> <px> <py> <pz>
```

where	<u0>	is the universe number of the lattice
	<type>	is the lattice type (= 11)
	<x0>	is the x coordinate of the lattice origin
	<y0>	is the y coordinate of the lattice origin
	<z0>	is the z coordinate of the lattice origin
	<nx>	is the number of lattice elements in the x-direction
	<ny>	is the number of lattice elements in the y-direction
	<nz>	is the number of lattice elements in the z-direction
	<px>	is the lattice pitch in x-direction
	<py>	is the lattice pitch in y-direction
	<pz>	is the lattice pitch in z-direction

The lattice card is followed by a list of universes. This lattice type is available from version 1.1.17 on.

3.6.3 Universe and lattice examples

The universe and lattice definitions are best described using a few examples. The first example is a 17×17 PWR MOX fuel assembly containing three types of fuel pins and empty control rod guide tubes (see Figure 3.4 on page 45).

```
% --- MOX pin 1:
pin 1
MOX1    4.36250E-01
void     4.43750E-01
clad     4.75000E-01
water

% --- MOX pin 2:
pin 2
MOX2    4.36250E-01
void     4.43750E-01
clad     4.75000E-01
water

% --- MOX pin 3:
pin 3
MOX3    4.36250E-01
```



```

void      4.43750E-01
clad      4.75000E-01
water

% --- Empry guide tube:
pin 4
water     5.62500E-01
clad      6.12500E-01
water

% --- Pin lattice (pitch = 1.26 cm):

lat 10  1 0.0 0.0 17 17 1.26

1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 1 1
1 2 3 3 3 2 3 3 2 3 3 2 3 3 3 2 1
2 3 3 3 3 4 3 3 4 3 3 4 3 3 3 3 2
2 3 3 4 3 3 3 3 3 3 3 3 3 4 3 3 2
2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 2
2 2 4 3 3 4 3 3 4 3 3 4 3 3 4 2 2
2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 2
2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 2
2 2 4 3 3 4 3 3 4 3 3 4 3 3 4 2 2
2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 2
2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 2
2 2 4 3 3 4 3 3 4 3 3 4 3 3 4 2 2
2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 2
2 3 3 4 3 3 3 3 3 3 3 3 3 4 3 3 2
2 3 3 3 3 4 3 3 4 3 3 4 3 3 3 3 2
1 2 3 3 3 2 3 3 2 3 3 2 3 3 3 2 1
1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 1 1

```

The second example is a hexagonal VVER-440 lattice with 126 fuel pins and a central instrumentation tube. Empty lattice positions are filled with water (see Figure 3.5 on page 45).

```

% --- Fuel pin with central hole:

pin 1
void      0.08000
fuel      0.37800
void      0.38800
clad      0.45750
water

% --- Central instrumentation tube:

```

```

pin 2
water    0.44000
clad     0.51500
water

% --- Empty lattice position filled with water:

pin 3
water

% --- Pin lattice (x-type hexagonal, pitch = 1.23 cm):

lat 10  2  0.0 0.0 15 15 1.23

3 3 3 3 3 3 3 3 3 3 3 3 3 3 3
3 3 3 3 3 3 3 1 1 1 1 1 1 1 3
3 3 3 3 3 3 1 1 1 1 1 1 1 1 3
3 3 3 3 3 1 1 1 1 1 1 1 1 1 3
3 3 3 3 1 1 1 1 1 1 1 1 1 1 3
3 3 3 1 1 1 1 1 1 1 1 1 1 1 3
3 3 1 1 1 1 1 1 1 1 1 1 1 1 3
3 1 1 1 1 1 1 2 1 1 1 1 1 1 3
3 1 1 1 1 1 1 1 1 1 1 1 1 3 3
3 1 1 1 1 1 1 1 1 1 1 1 3 3 3
3 1 1 1 1 1 1 1 1 1 1 3 3 3 3
3 1 1 1 1 1 1 1 1 1 3 3 3 3 3
3 1 1 1 1 1 1 1 1 3 3 3 3 3 3
3 1 1 1 1 1 1 1 3 3 3 3 3 3 3
3 3 3 3 3 3 3 3 3 3 3 3 3 3 3

```

The third example is a CANDU cluster consisting of 37 pins in 4 rings. The third ring is rotated by 15 degrees (see Figure 3.6 on page 46).

```

% --- Fuel pin:

pin 1
fuel 0.6122
clad 0.6540
coolant

% --- Cluster:

lat 10  4  0.0 0.0 4
1  0.0000  0.0 1
6  1.4885  0.0 1 1 1 1 1 1
12 2.8755 15.0 1 1 1 1 1 1 1 1 1 1 1
18 4.3305  0.0 1 1 1 1 1 1 1 1 1 1 1 1 1 1

```

All three examples are illustrated using the geometry plotter in Section 3.9 on page 42. It should be noted that the plots contain cell structures not included in the above examples.

The following example demonstrates the use of the vertical stack:

```
% --- Uranium ball:

surf 1 sph 0.0 0.0 2.5 2.5

cell 1 1 uranium -1
cell 2 1 void     1

% --- Stack 5 balls:

lat 2 9 0.0 0.0 5

0.0 1
5.0 1
10.0 1
15.0 1
20.0 1
```

Notice that the origin of universe 1 is positioned at the bottom of each layer.

IMPORTANT NOTES ON UNIVERSES AND LATTICES:

1. Each universe is defined independently and must cover all space. Regions of space not belonging to the geometry must be defined using “outside” cells.
2. The lowest level of the geometry belongs to universe 0, which must always exist.
3. Each universe has its own origin, which can be shifted using the universe transformation command.
4. Cells in higher geometry levels can only be accessed through filled cells or lattices.
5. Each lattice defines a universe, which must be embedded inside a cell using the fill command. The lattice must fill the container cell completely to avoid neutron streaming to undefined lattice positions.
6. Hexagonal lattices are defined using a square matrix for the universe layout. To position the lattice cells correctly, a number of empty cells must be defined for each row. The definition is best described in the example in Sec. 3.6.3 on page 33.
7. Multi-level hexagonal structures (pin-assembly-core) are defined using both x- and y-type hexagonal lattices with different type at each level.

8. If the infinite lattice types are used in burnup calculation, material volumes must be set manually (see Sec. 4.1.2 on page 49).
9. The vertical stack lattice type is available from code version 1.1.8 on.

SEE ALSO:

1. Pin definitions (Sec. 3.4 on page 27)
2. Filled cells (Sec. 3.6 on page 28)

3.7 Repeated Boundary Conditions

What happens to neutrons that end up in a region defined as being outside the geometry is dictated by the boundary conditions. There are three options:

1. Black boundary - the neutron is killed
2. Reflective boundary - the neutron is reflected back into the geometry
3. Periodic boundary - the neutron is moved to the opposite side of the geometry

The condition is set by the “bc” parameter, described in Section 5.7 on page 59.

Reflective and periodic boundary conditions can be used to construct infinite and semi-infinite lattice structures. The way these boundary conditions are handled in Serpent is somewhat different from other Monte Carlo codes. Instead of stopping the neutron at the boundary surface, reflections and translations are handled by coordinate transformations. This limits the outermost boundary to a few specific surface types that can be used to define a square or hexagonal lattice structure. There are basically three options:

Infinite 2D geometry: The geometry has no dependence on the z-coordinate. The outer boundary is defined by a single square or hexagonal cylinder (“sqc”, “hexxc” or “hexyc”).

Radially infinite, axially finite 3D geometry: The outer boundary is defined by a square or hexagonal cylinder (“sqc”, “hexxc” or “hexyc”) and two axial planes (“pz”). The boundary condition takes effect in the radial direction only. The axial boundary conditions are black.

Infinite 3D geometry: The outer boundary is defined by a single cube, cuboid or hexagonal prism (“cube”, “cuboid”, “hexxprism” or “hexyprism”). The boundary condition takes effect in all directions.

The following examples illustrate the different geometry types. The details of the geometry are omitted for the sake of simplicity and replaced by a fill command.

An infinite 2D hexagonal geometry can be defined as:

```
surf 1 hexyc 0.0 0.0 7.350

% --- Cells:

cell 1 0 fill 10 -1
cell 99 0 outside 1

set bc 3
```

Note that the reflective boundary condition is unphysical in a hexagonal geometry. infinite 2D square geometry can be defined as:

```
surf 1 sqc -0.233 -0.233 7.68750

cell 1 0 fill 10 -1
cell 99 0 outside 1

set bc 2
```

In both cases the outer boundary is defined by a single surface.

If the geometry is finite in the axial dimension, the system becomes three-dimensional. A radially infinite square lattice can be defined as:

```
surf 1 sqc -0.233 -0.233 7.68750
surf 2 pz -200.0
surf 3 pz 200.0

cell 1 0 fill 10 -1 2 -3
cell 97 0 outside 1 2 -3
cell 98 0 outside -2
cell 99 0 outside 3

set bc 2
```

It is also possible to define the outside world as:

```
cell 97 0 outside 1
cell 98 0 outside -1 -2
cell 99 0 outside -1 3
```

but the code may run slower because the boundary condition will be handled also for some neutrons that end up outside the geometry.

As for the infinite 2D geometry, the boundary in an infinite 3D geometry must be defined by a single surface, such as a cube:

```
surf 1 cube 0.0 0.0 0.0 3.0

cell 1 0 fill 10 -1
cell 99 0 outside 1

set bc 2
```

or a hexagonal prism:

```
surf 1 hexxprism 0.0 0.0 1.880 0.0 4.93

cell 1 0 fill 10 -1
cell 99 0 outside 1

set bc 3
```

In both cases the boundary conditions are enforced in both radial and axial directions.

IMPORTANT NOTES ON REPEATED BOUNDARY CONDITIONS:

1. The outer boundary must be defined by a single surface in infinite 2D and 3D geometries. The allowed surface types for a 2D geometry are square and hexagonal cylinders. Infinite 3D geometries can be defined using a cube, cuboid or hexagonal prism.
2. Axially infinite, radially finite geometries are defined by a square or hexagonal cylinder and two axial planes. The way the outside world is defined may affect the running time.
3. The hexagonal cylinder and prismatic surfaces are physically reasonable only with periodic boundary conditions (reflective boundary conditions work if the geometry has a 30 degree symmetry). The use of reflective boundary conditions with these types was enabled in update 1.1.18. In earlier code versions the boundary condition is automatically changed into periodic.

SEE ALSO:

1. Surface types (Sec. 3.2.1 on page 20)
2. Defining the outside world (Sec. 3.3.1 on page 25)
3. Setting the boundary condition (Sec. 5.7 on page 59)

3.8 HTGR geometry types

The fuels in high-temperature gas-cooled reactors (HTGR) consist of microscopic TRISO particles dispersed in a graphite matrix. The multi-layer fuel particles can be defined similar to fuel pins (see Sec. 3.4 on page 27):

```
particle <id>
  <mat 1> <r1>
  <mat 2> <r2>
  ...
  <mat n>
```

where <id> is the particle identifier (universe number)
 <mat 1> <mat 2> ... are the materials
 <r1> <r2> ... are the outer radii of the material regions

The simplest approach is to describe the particle distribution as a regular lattice, using lattice types 6, 7 or 8 (See the infinite 3D lattices in Sec. 3.6.2). However, the regular arrangement fails to account for the random distribution of the particles and often leads to a distorted fuel-to-moderator ratio due to cell cut-off at the outer boundary. For this reason the Serpent code has two geometry models specifically designed for HTGR fuels.

3.8.1 Implicit particle fuel model

The implicit particle fuel model works by sampling new particles on the neutron flight path during the tracking process. The input syntax is:

```
disp <u0> <uf> <pfn> <r1> <u1> ... <pfn> <rn> <un>
```

where <u0> is the universe number of the dispersed medium
 <uf> is the universe filling the space between the particles
 <pfn> ... <pfn> are the packing fractions of the particle types
 <r1> ... <rn> are the radii of the particle types
 <u1> ... <un> are the universe numbers of the particle types

The number of particle types is not limited, but the sum of the packing fractions must be less than 1.0 (physical factors set the upper limit much lower, although this is not checked by the routine).

The implicit particle fuel model was revised in update 1.1.3. It should be noted that the model is not exact and there are statistically significant differences compared to the explicit model described below. The implicit model seems to work best for low packing fractions but no comprehensive validation has been carried out yet.

3.8.2 Explicit particle / pebble bed fuel model

A better choice for modeling HTGR geometries is the explicit particle fuel model, which reads the positions of the particles from a separate file. The same model can be used for setting up reactor-scale pebble-bed geometries. The input syntax is:

```
pbed <u0> <uf> "<inputfile>" [<options>]
```

where <u0> is the universe number of the dispersed medium
 <uf> is the universe filling the space between the particles / pebbles
 <inputfile> is the input file containing the particle / pebble coordinates
 <options> are the options

The particle / pebble distribution is handled explicitly, so there are no approximations done in the modeling. Each line in the input file describes the position of a single particle / pebble. The format is:

```
<x> <y> <z> <r> <u>
```

where <x> is the x coordinate of the particle / pebble
 <y> is the y coordinate of the particle / pebble
 <z> is the z coordinate of the particle / pebble
 <r> is the radius of the particle / pebble
 <u> is the universe number of the particle / pebble

The total number of entries is unlimited, although memory or running time may become a limiting factor if the number exceeds several million.

The options are used to activate the calculation of various particle / pebble-wise parameters. Currently the only available option is the power distribution, which is requested with option “pow”. The code writes the output in a separate file “<inputfile>.out”, where “<inputfile>” is the file where the distribution was read. The input data is included for convenience. The format of the output is:

```
<x> <y> <z> <r> <u> <P> <dP>
```

where <x> is the x coordinate of the particle / pebble
 <y> is the y coordinate of the particle / pebble
 <z> is the z coordinate of the particle / pebble
 <r> is the radius of the particle / pebble
 <u> is the universe number of the particle / pebble
 <P> is the power produced inside the particle / pebble
 <dP> is the associated relative statistical error

All results depend on source normalization (see Sec. 5.8 on page 61).

3.8.3 HTGR geometry examples

The following example shows how the particle distribution inside a single PBMR fuel pebble can be modeled using a regular 3D array and the two particle fuel models in the Serpent code.

The definition of a fuel particle is very similar to the fuel pin:

```
% --- Definition of a coated fuel particle:

particle 1

fuel      0.0250
buffer    0.0340
PyC       0.0380
SiC       0.0415
PyC       0.0455
matrix
```

The first option is to describe the particle distribution as a regular cubical lattice:

```
% --- Option 1: regular 3D array:

lat  10 6 0.0 0.0 0.16341 1
```

The implicit particle fuel model is defined using a list of packing fractions and particle types:

```
% --- Filler universe composed of graphite:

surf 1 inf
cell 1 2 matrix -1

% --- Option 2: implicit particle fuel model:

disp 10 2 0.09043 4.55000E-02 1
```

The explicit particle fuel model reads particle coordinates from a separate input file (can be used for pebble distributions at reactor scale as well):

```
% --- Filler universe composed of graphite:

surf 1 inf
cell 1 2 matrix -1
% --- Option 3: explicit particle fuel model (read coordinates from file):

pbed 10 2 "particles.inp"
```

Finally the pebble description using one of the three options (all assigned with universe number 10):

```
% --- Pebble:

surf 10 sph 0.0 0.0 0.0 2.5
surf 20 sph 0.0 0.0 0.0 3.0
surf 30 cube 0.0 0.0 0.0 3.0

cell 10 0 fill 10 -10
cell 20 0 matrix 10 -20
cell 30 0 helium 20 -30
cell 40 0 outside 30
```

IMPORTANT NOTES ON HTGR GEOMETRY TYPES:

1. The implicit particle fuel model was revised in update 1.1.3. The model is not exact and should be used with caution. Test calculations show that the model works best for low packing fractions.
2. If the implicit particle fuel model is used in burnup calculation, material volumes must be set manually (see Sec. 4.1.2 on page 49).
3. Calculation of particle / pebble-wise power distributions is available from update 1.1.4 on.

SEE ALSO:

1. An earlier version of the implicit particle fuel model in Ref. [8]

3.9 Geometry plotter

The geometry plotter uses the GD open source graphics library [1] for producing png format output files for visualization. In order to use the plotter, the source code must be compiled with this library included (see the Makefile for detailed instructions). The syntax of the plotter command is:

```
plot <or> <nx> <ny> [<p> <min1> <max1> <min2> <max2>]
```

where

- <or> is the orientation of the plot plane (1, 2 or 3)
- <nx> is the width of the plot in pixels
- <ny> is the height of the plot in pixels
- <p> is the position on the axis perpendicular to the plot plane
- <min1> is the minimum value of the first coordinate
- <max1> is the maximum value of the first coordinate
- <min2> is the minimum value of the second coordinate
- <max2> is the maximum value of the second coordinate

The orientation of the plot plane is defined as:

1. yz-plot (perpendicular to the x-axis)
2. xz-plot (perpendicular to the y-axis)
3. xy-plot (perpendicular to the z-axis)

The plotted area is a rectangle defined by the orientation, the position on the perpendicular coordinate axis and the coordinates of the two corners. Zero position is assumed if the position parameter is omitted. If the corner coordinates are not given, the boundary values of the geometry are used.

Each plotter command produces an output file named “<input>_geom<n>.png”, where <input> is the name of the input file and <n> is the plot index. The resolution of the figure is defined by the width and height parameters. Each material is represented by a randomly selected color (void regions are in black, geometry errors bright green or red). Surfaces are drawn with black lines, which may overlap cell regions. It should be noted that the plotted surfaces may not necessarily represent the actual cell boundaries.

Example plots are shown in Figures 3.4–3.7. The lattices in the first three cases are described in the universe and lattice examples in Sec. 3.6.3. In each case the plotter command was:

```
plot 3 1000 1000
```

This generates a 1000 by 1000 pixel plot perpendicular to the z-axis, located at $z = 0$ and covering the entire geometry.

IMPORTANT NOTES ON GEOMETRY PLOTTER:

1. The geometry plotter uses the GD open source graphics library [1], which must be installed in the system.
2. The plotter produces png (portable network graphics) format output files.

3. The colors in the plot represent different materials. The color for each material is selected randomly (void regions are black, geometry errors bright green or red).
4. Surfaces are drawn with black lines, which may overlap cell regions. Plotted surfaces may not necessarily represent the actual cell boundaries.

SEE ALSO:

1. Compiling Serpent (Sec. 1.1 on page 8)
2. The GD open source graphics library: <http://www.libgd.org>

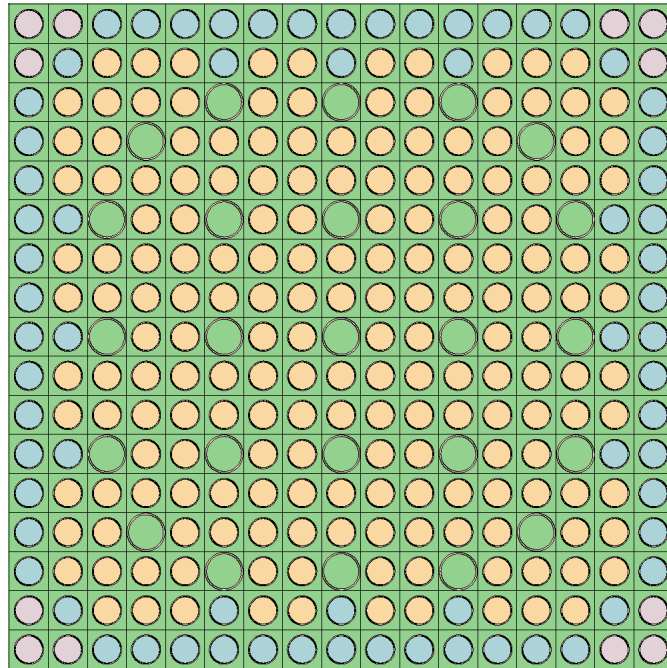


Figure 3.4: A 17×17 PWR MOX fuel assembly with 3 pin types.

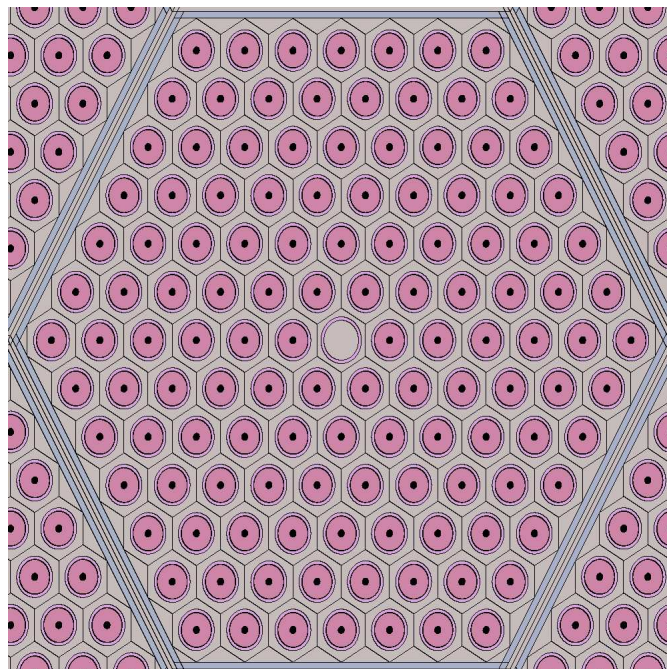


Figure 3.5: A hexagonal VVER-440 fuel assembly with 126 fuel pins and a central instrumentation tube in an infinite lattice. The proportions of the assembly are slightly distorted since the hexagonal assembly is fitted inside a square region.

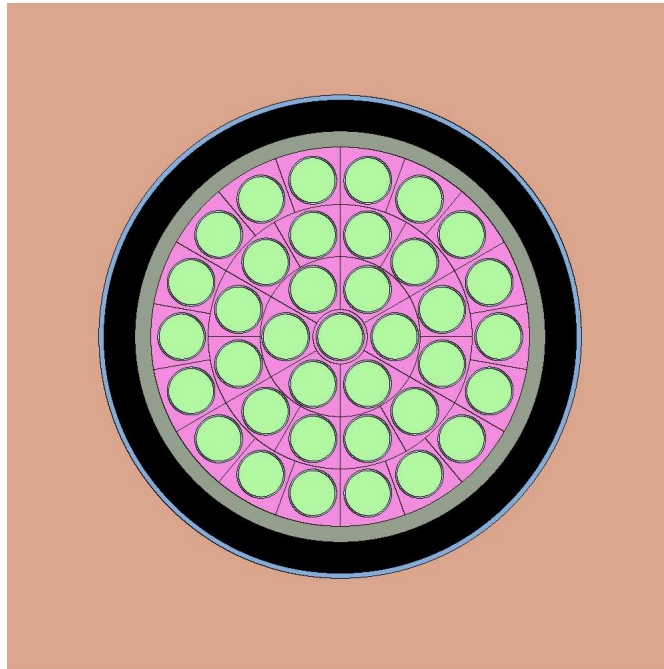


Figure 3.6: A CANDU cluster with 37 fuel pins in 4 rings. The third ring is rotated by 15 degrees.

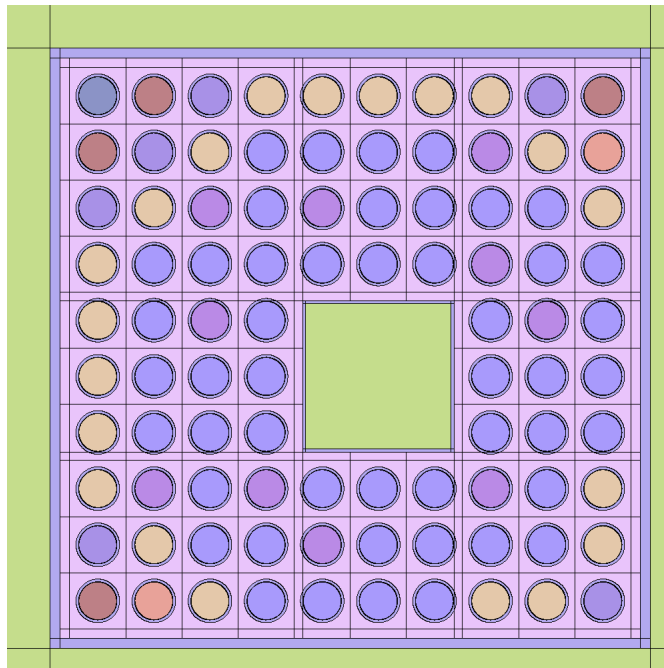


Figure 3.7: A 10×10 BWR fuel fuel assembly with 7 pin types and an asymmetrically positioned moderator channel.

Chapter 4

Materials

4.1 Material definitions

The geometry in Monte Carlo codes consists homogeneous material regions, which in Serpent are defined using cells and surfaces (see Chapter 3 for geometry definition).¹ Each material consists of a list of nuclides and each nuclide is associated with a cross section library, as defined in the directory file (see Sec. 1.4.2 on page 12).

Nuclide temperatures are fixed when the cross section data is generated and cannot be changed afterwards. It is important to use cross section libraries generated at the right temperature to correctly model the Doppler-broadening of resonance peaks. It is equally (or even more) important to use the appropriate bound-atom thermal scattering libraries for moderator nuclides.

Soluble absorbers can be defined by mixing two material compositions. This option is introduced in Sec. 5.14 on page 69. The concentration can be used for critical k_{eff} iteration. Serpent also has the option to use a built-in Doppler broadening routine to adjust nuclide temperatures before the calculation. This method is described in Sec. 4.3 on page 50.

4.1.1 Nuclides

Nuclide names may be arbitrary aliases defined in the directory file. The usual convention, also used by MCNP, is:

¹It is, in principle, possible to model continuously varying material compositions when the delta-tracking method is used for neutron transport. This option is considered for the future versions of the Serpent code.

`<Z><A>.<id>`

where `<Z>` is the element Z
`<A>` is the isotope mass number (three digits)
`<id>` is the library id

For example, “92235.09c” refers to ^{235}U . Natural element cross sections are denoted by mass number zero (“40000.06c” for natural zirconium). The library id usually refers to data evaluation or temperature (“60c” for ENDF/B-VI.0 based data, “09c” for data generated at 900K, and so on...).

There is no standard convention on how to name isomeric states. The `xmdirconvert`-utility used for producing Serpent directory files assumes a form in which the isotope mass number is simply increased above 300 (“95342.09c” for ^{242m}Am). In any case it is important to realize that the nuclide names are used for identification only and they do not contain any information used by the code in the calculation.

4.1.2 Material cards

The basic syntax of the material card is:

```
mat <name> <dens> [<options>]
<iso 1> <frac 1>
<iso 2> <frac 2>
...
```

where `<name>` is the material name
`<dens>` is the density (mass or atomic)
`<options>` are the options (depending on case)
`<iso 1> <iso 2> ...` are the names of the constituent nuclides
`<frac 1> <frac 2> ...` are the corresponding fractions (mass or atomic)

Material name is used to identify the material in cell cards (see Sec. 3.3.1 on page 25). The nuclide names correspond to the identifier determined in the directory file. These identifiers define the cross section data used in the calculation. Densities and fractions can be given as atomic or mass values. Positive entries refer to atomic densities (in units of $10^{24}/\text{cm}^3$) and atomic fractions, respectively, and negative entries to mass densities (in units of g/cm^3) and mass fractions. Isotopic compositions are normalized before the calculation and mixed entries are not allowed.

If the material density is set to zero or “sum”, the value is calculated from the isotopic composition. The isotope fractions must then be in absolute density units, not relative fractions.

Material volumes and masses are used for normalizing reaction rates, which is important, for example, in burnup calculation. The code calculates these automatically for simple pin structures, but the values must be entered manually for some more complicated geometries.

Material volume is set using option:

```
mat <name> <dens> vol <V>
...
```

where $\langle V \rangle$ is the total material volume in cm^3

and material mass (alternatively):

```
mat <name> <dens> mass <M>
...
```

where $\langle M \rangle$ is the total material mass in g

Colors for the geometry plotter (see Sec. 3.9 on page 42) can be set using:

```
mat <name> <dens> rgb <R> <G> <B>
...
```

where $\langle R \rangle$ is the value for red channel (between 0 and 255)

$\langle G \rangle$ is the value for green channel (between 0 and 255)

$\langle B \rangle$ is the value for blue channel (between 0 and 255)

if the colors are not set, random values are used in the plots.

Other options are used to set up depletion zones in burnup calculation and to determine thermal scattering libraries for moderator materials and temperatures for Doppler broadening. Material definitions in burnup calculation is a separate topic in Section 8.2 on page 109. Thermal scattering and Doppler broadening are discussed below.

4.2 Thermal scattering libraries

Thermal scattering cross sections are used to replace the low-energy free-gas elastic scattering reactions for some important bound moderator nuclides, such as hydrogen in water or carbon in graphite. Thermal systems cannot be modelled using free-atom cross sections without introducing significant errors in the spectrum and results. Thermal scattering data is defined using:

```
therm <thname> <lib>
```

where $\langle \text{thname} \rangle$ is the name of the data library

$\langle \text{lib} \rangle$ is the library identifier as defined in the directory file

The library identifier is the actual name of the library in the directory file. The library name is used to associate the data with a material, in which case the material card has an additional entry:

```

mat <name> <dens> moder <thname> <ZA>
<iso 1> <frac 1>
<iso 2> <frac 2>
...

```

where	<name>	is the material name
	<dens>	is the density (mass or atomic)
	<thname>	is the name of the thermal scattering data library
	<ZA>	is the nuclide ZA of the moderator nuclide
	<iso 1> <iso 2> ...	are the names of the constituent nuclides
	<frac 1> <frac 2> ...	are the corresponding fractions (mass or atomic)

The nuclide ZA identifies the moderator nuclide (in the form of: 1000*Z + A). The “moder” entry can be used several times to define thermal scattering libraries for multiple nuclides, such as hydrogen and deuterium in heavy water (see the example in Sec. 4.4).

4.3 Doppler broadening

The Doppler broadening routine is initiated by adding a “tmp” entry in the material card:

```

mat <name> <dens> tmp <T>
<iso 1> <frac 1>
<iso 2> <frac 2>
...

```

where	<name>	is the material name
	<dens>	is the density (mass or atomic)
	<T>	is the Doppler temperature in Kelvin
	<iso 1> <iso 2> ...	are the names of the constituent nuclides
	<frac 1> <frac 2> ...	are the corresponding fractions (mass or atomic)

The broadening is performed after the data is read from the ACE format libraries and there is slight increase in the overall calculation time, depending on the number of nuclides. If the the same nuclide is broadened to several temperatures in different materials, there is also an increase in memory usage. The routine works only if the given temperature is above the original one. The cross section libraries provided with the Serpent code are generated between 300K intervals and it is recommended that the closest temperature below the broadened value is used as the basis.

4.4 Material examples

A few simple examples of material definitions are given in the following.

```

% --- Fuel consisting of enriched UO2 and burnable absorber.
%      Atomic densities given in units of 1/(barn*cm):

mat UO2Gd    sum          % Atomic density from composition
92234.09c    4.2940E-06    % Atomic density of U-234
92235.09c    5.6226E-04    % Atomic density of U-235
92238.09c    2.0549E-02    % Atomic density of U-238
64154.09c    4.6173E-05    % Atomic density of Gd-154
64155.09c    2.9711E-04    % Atomic density of Gd-155
64156.09c    4.1355E-04    % Atomic density of Gd-156
64157.09c    3.1518E-04    % Atomic density of Gd-157
64158.09c    4.9786E-04    % Atomic density of Gd-158
64160.09c    4.3764E-04    % Atomic density of Gd-160
8016.09c     4.5243E-02    % Atomic density of O-16

% --- Zircaloy cladding:

mat clad     -6.55000      % Mass density given in g/cm3
40000.06c    -0.98135     % Mass fraction of natural zirconium
24000.50c    -0.00100     % Mass fraction of natural chromium
26000.55c    -0.00135     % Mass fraction of natural iron
28000.42c    -0.00055     % Mass fraction of natural nickel
50000.42c    -0.01450     % Mass fraction of natural tin
8016.06c     -0.00125     % Mass fraction of O-16

% --- Boronized light water with thermal scattering data:

mat water    -0.7207      moder lwtr 1001
1001.06c     -1.1180E-1
8016.06c     -8.8755E-1
5010.06c     -1.1890E-4
5011.06c     -5.3110E-4

therm lwtr lwtr.04t

% --- Heavy water with thermal scattering data (two libraries):

mat D2O      -0.812120    moder lwtr1 1001 moder hwtr1 1002
8016.06c     -7.99449E-1
1002.06c     -1.99768E-1
1001.06c     -7.83774E-4

therm lwtr1 lwtr.04t
therm hwtr1 hwtr.04t

% --- Doppler broadening from 900K to 1000K:

mat fuel     -10.45700    tmp 1000

```

92235.09c	-0.03173
92238.09c	-0.84977
8016.09c	-0.11850

IMPORTANT NOTES ON MATERIALS:

1. Nuclide names are used for identification only. All information used in the calculation is read from the directory file and the cross section data.
2. Positive entries in material density and isotopic composition refer to atomic densities and atomic fractions, respectively, and negative entries to mass densities and mass fractions. Typical input errors in material compositions are related to confusing the two definitions.
3. Isotopic compositions can be given as density values, rather than fractions, since the compositions are normalized before the calculation.
4. The mass fraction of oxygen in UO_2 fuel is ~ 0.1185 . Natural boron consists of 20% ^{10}B and 80% ^{11}B (atomic fractions).
5. Thermal scattering data must be used for moderator materials (water, graphite, etc.) when modelling thermal systems. The use of free-atom cross sections will introduce significant errors in the results.
6. Doppler broadening is available from code version 1.1.0 on, and completed in version 1.1.2. The broadened temperature must be above the original nuclide temperature.

SEE ALSO:

1. Directory files and the “xsdirconvert” utility (Sec. 1.4.2 on page 12)
2. Soluble absorber definitios (Sec. 5.14 on page 69)

Chapter 5

Options

5.1 General

Serpent has various calculation parameters determined using the “set” command:

```
set <param> <value 1> <value 2> ...
```

where <param> is the parameter name
 <value 1> <value 2> ... are the parameter values

The available options are listed in Table 5.1 and described in more detail in the following sections. Parameters used for burnup calculation are described in Section 8.4 of Chapter 8. Table 8.2 on page 111 summarizes the options in the burnup calculation mode.

5.2 Neutron Population and Criticality Cycles

The default calculation mode in Serpent is the k -eigenvalue criticality source method, in which the simulation is run in cycles and the source distribution of each cycle is formed by the fission reaction distribution of the previous cycle. External source simulation is discussed as a separate topic in Chapter 9. The parameters for criticality source calculation are set using:

```
set pop <npop> <cycles> <skip> [<keff0> <int>]
```

where <npop> is the number of source neutrons per cycle
 <cycles> is the number of active cycles run
 <skip> is the number of inactive cycles run
 <keff0> is the initial guess for k_{eff}
 <int> is the collection interval

The number of source neutrons per cycle is fixed. Since the number of generated source points usually differs from this value, the source size is increased ($k_{\text{eff}} < 1$) or decreased ($k_{\text{eff}} > 1$) to match the given source size.

Inactive cycles are cycles that are run in order to allow the initial fission source distribution to converge before starting to collect the results. In lattice calculations the convergence is typically reached well within the first 20 cycles. Source convergence in full-core calculations, however, may take much longer.

The initial source points are randomly selected inside the fissile cells in the geometry and no source input is needed from the user. The simulation requires an initial guess for k_{eff} , which by default is set to unity. This is usually sufficient, but if the system is far from criticality, the simulation may die out during the first few cycles. This problem may be overcome by setting the initial k_{eff} guess to a more appropriate value.

If all fissile material is located in a small region compared to the geometry dimensions, initial source sampling may fail. The default source can be overridden by defining an external source, as described in Section 9.2 of Chapter 9. If the “nps” parameter is not set, the user-defined source is used as the initial guess only, and the simulation proceeds in power iterations.

The statistical accuracy of the results depends on the total number of active neutron histories run, which is determined by the population size per cycle and the total number of active cycles. Appropriate values for a typical lattice calculation are 500 active cycles of 5000 source neutrons. If more precision is required or the geometry is larger, it is suggested that the population size, rather than the number of cycles is increased.

The collection interval defines the number of generations run for each batch of results. By default this value is set to one, and increasing the number has essentially the same impact as running more neutrons per generation and fewer generations.¹

Serpent uses two buffers to store data for new source points and neutrons produced in multiplying scattering reactions and certain special calculation modes. In some cases these buffers may be overrun, which terminates the simulation. To overcome such problems, the buffer size may be increased by setting:

```
set nbuf <f>
```

where <f> is the buffer factor (criticality mode) or total size (external source mode)

In criticality source mode the buffer size is population size multiplied by the given factor (set to 2.5 by default). In external source mode neutron histories are run one at a time and the value of nbuf sets the absolute size of the buffer (set to 1000 by default).

IMPORTANT NOTES ON NEUTRON POPULATION AND CRITICALITY CYCLES:

¹The difference is that the correlations between adjacent batches could be weaker, which may have some impact in the statistics in large geometries (the effects have not yet been studied).

1. It is important that a sufficient number of cycles is discarded to allow the initial fission source to converge before starting to collect the results. This number depends on the size and the complexity of the geometry. Fission source convergence is a complicated research topic, subject to both theoretical and practical considerations [9–14]. It should be noted, however, that problems with source convergence are practically never encountered in lattice calculations where the neutron migration distance is long compared to the dimensions of the geometry.
2. The k -eigenvalue criticality source calculation yields physically consistent results only in the special case that $k_{\text{eff}} = 1$. When the system is far from criticality, the importance of fission neutrons is either over- ($k_{\text{eff}} < 1$) or underestimated ($k_{\text{eff}} > 1$). The result is that the neutron population becomes biased in energy and space (and time), which may affect the final results as well. The problem originates from the basic methodology and the fact that a physically sub- or super-critical chain reaction is simulated as a steady-state system. Deterministic lattice transport codes use neutron leakage models to overcome this problem, but the methodology for Monte Carlo calculation is not well established.
3. All source neutrons are born in fission. Other neutron-multiplying (n,xn) reactions are treated as scattering within the criticality cycle.
4. External source definitions are available from version 1.1.11 on.
5. Buffer size and collect interval are options available from version 1.1.13 on.

SEE ALSO:

1. Simulating the neutron chain reaction and the k -eigenvalue criticality source calculation mode in Ref. [15] (Sec. 5.5 on page 112).
2. Discussion on neutron leakage models in Monte Carlo calculation in Ref. [15] (Sec. 9.5 on page 171).
3. External source simulation (Chapter 9).

5.3 Energy grid reconstruction

The continuous-energy reaction cross sections in Serpent are reconstructed using a single unionized energy grid for all nuclides. The reason for this approach is the major speed-up in calculation, achieved by minimizing the number of grid search iterations.² The default

²Each nuclide in the continuous-energy ACE format data is associated with its own energy grid. The calculation of material total cross sections, for example, is carried out by summing over all the constituent nuclides. This requires an iterative energy grid search to be performed for each nuclide, which may take a significant fraction of the overall CPU time, especially since the procedure has to be repeated each time the neutron enters a new material. The advantage of using the same grid for all nuclides is that the grid search has to be performed only once, each time the neutron scatters to a new energy.

method for grid reconstruction is that all grid points of all nuclides in the ACE format data are included in the master grid. The disadvantage of this method is that computer memory is wasted for storing a large number of redundant data points. The available memory is usually not a problem in fresh fuel calculations, but the introduction of actinide and fission product isotopes in burnup calculation may result in an excessively large master grid. Serpent has a method for avoiding this problem by combining adjacent grid points. The reconstruction parameters are given by:

```
set egrid <tol> [<Emin> <Emax>]
```

where <tol> is the fractional reconstruction tolerance
<Emin> is the minimum energy in the grid (MeV)
<Emax> is the maximum energy in the grid (MeV)

The fractional reconstruction tolerance is the minimum relative difference between two grid points, below which the points are combined. All points below the lower limit and above the upper limit are discarded.

The default value for the fractional reconstruction tolerance is zero in the transport calculation mode and $5 \cdot 10^{-5}$ in the burnup calculation mode. Test calculations have shown that the results are not significantly affected until the tolerance is raised above 10^{-3} . There is no absolute guarantee, however, that the results are valid in all imaginable cases when the grid size is significantly reduced. It is therefore suggested that the grid reduction is not used unless necessary because of insufficient computer memory.

The lower limit of the energy grid is by default set to 10^{-9} MeV and the upper limit to 15 MeV. Very few neutrons are born or scattered to higher or lower energies in fission reactor applications.

If a reduction in memory size is necessary, an alternative to grid thinning is the double indexing method, in which the data is stored in the original ACE format and the unionized grid used only for accessing the nuclide-wise grids. This method is activated by:

```
set dix <mode>
```

where <mode> is 1 if the method is used and 0 if not.

The double indexing method reduces the memory usage, but may lead to an increase in processing time, which may become significant in burnup calculation. Double indexing is turned off by default.

IMPORTANT NOTES ON ENERGY GRID:

1. Grid reduction inevitably leads to some loss of data. There is no guarantee that this reduction does not affect the results.
2. Test calculations have shown that the reduction in grid size does *not* significantly affect the overall calculation time.

SEE ALSO:

1. Cross section data in the PSG code in Ref. [15] (Sec. 8.2 on page 143). NOTE: Some of the described methods are outdated.
2. A more recent description of the unionized energy grid formats in Serpent is found in Ref. [16].

5.4 Library File Paths

The Serpent code uses a single directory file for determining the cross sections used in the transport simulation. The directory file can be generated from an MCNP xsdir file using the “xsdirconvert” utility (see Sec.1.4 on page 11). The cross section library directory file path is set using:

```
set acelib "<file>"
```

where `<file>` is the file path for the ACE directory file

A default directory path can be set by defining environment variable `SERPENT_DATA`. The code looks for cross section directory files in this path if not found at the absolute location.

IMPORTANT NOTES ON DATA FILES AND FILE PATHS:

1. The cross section library directory file is a text file generated by the “xsdirconvert” utility.
2. The environment variable feature is available from code version 1.1.8 on.

SEE ALSO:

1. Setting up the directory file (Sec.1.4 on page 11)
2. Setting up file paths for burnup calculation (Sec.8.4 on page 111)

5.5 Unresolved resonance data

The use of unresolved resonance probability tables can be switched on and off using:

```
set ures <use> [<mode>] [dilu] [<iso 1> <iso 2> ...]
```

where <use> is the option (1 = use data, 0 = omit data)
 <mode> is the calculation mode
 <dilu> is the infinite dilution cut-off
 <iso n> are the nuclides for which the data is used or omitted

Since the probability table sampling has to be carried out during tracking, the transport cycle tends to slow down quite significantly.³ There are three options to treat the probability table data:

1. Sample all cross sections at once, each time the neutron scatters to a new energy, adjust material totals and majorant.
2. Sample cross sections when the neutron enters a new material. Use a pre-calculated majorant cross section corresponding to the maximum probability table values.
3. Sample cross sections when the neutron enters a new material. Switch to surface tracking when neutron is in the unresolved range.

The overall calculation time using the different options depends on the case, in particular the flux spectrum and the number of nuclides with probability table data. Mode 1 is used by default. It should also be noted that options 2 and 3 work by sampling the cross sections for physical materials. If a material is used for detector calculation only, the probability tables may not be appropriately sampled. This is not a problem for method 1.

Since the overall impact of using unresolved resonance cross sections is a self-shielding effect, the calculation routine can be optimized by omitting the probability table sampling for nuclides with low concentration. This limit is given by the infinite dilution cut-off, which is set to zero by default.

If the options are followed by a list of nuclide names (94239.09c, etc.), the probability table treatment is used or omitted only for the listed nuclides. If no list is given, the options cover all nuclides with probability table data.

In order for the methodology to work, the probability table data must be available in the ACE format cross section libraries. This data is not included, for example, in the default libraries provided with installation package 1.1.0. The methodology is available from update 1.1.4 on and is still under development. The mode and infinite dilution cut-off options were added in update 1.1.5. The treatment is currently switched off by default.

³Serpent pre-calculates certain material-wise total cross sections to avoid having to sum over the constituent nuclides during the transport cycle. This pre-calculation cannot be combined with probability table sampling, which has to be carried out on-the-fly.

5.6 Doppler-Broadening Rejection Correction (DBRC)

There is a physical flaw in the ENDF reaction laws of the ACE format data, that ignores the impact of thermal motion on angular distributions of elastic scattering near resonances. The phenomenon becomes important in heavy nuclides ($A > 200$) with scattering resonances at low energy (< 0.2 keV), and ignoring it may result in a noticeable under-prediction of resonance absorption and over-prediction of k_{eff} . To correct this flaw, Serpent can apply a Doppler-broadening rejection correction (DBRC) in the thermal free-gas model:

```
set dbrc <Emin> <Emax> [<iso 1> <iso 2> ...]
```

where <Emin> is the minimum energy for DBRC (MeV)
 <Emax> is the maximum energy for DBRC (MeV)
 <iso n> are the zero-Kelvin cross section data of the nuclides involved

The method uses zero-Kelvin elastic scattering cross sections in the rejection sampling loop and the provided data tells the code which nuclides should use the correction. If the correction is used with U-238, for example, the entry is the nuclide name for U-238 generated at 0K ("92238.00t"). It is usually sufficient to use DBRC for the primary heavy nuclide only. The energy range should cover the low resonance peaks. Typical range for U-238 is from 0.4 to 210 eV. The method is not used by default.

IMPORTANT NOTES ON DBRC:

1. The correction increases resonance absorption, which may reduce k_{eff} by few hundred pcm.
2. DBRC is not widely used by other Monte Carlo codes, so switching the correction on may increase differences to any reference results.
3. The method is available from update 1.1.14 on.
4. Zero-Kelvin cross section data is not available in the cross section libraries distributed with the current base versions.

SEE ALSO:

1. Theory behind DBRC is discussed in reference [17].

5.7 Boundary conditions

Boundary conditions determine the fate of neutrons escaping outside the defined geometry. The boundary conditions are set using:

```
set bc <c>
```

where <c> is the boundary condition

The Serpent code has three available boundary condition options: 1 - black, 2 - reflective and 3 - periodic. Default is the black boundary, which means that all neutrons streaming into outside cells are killed. Reflective and periodic boundary conditions can be used for setting up infinite lattices. When the neutron encounters a reflective boundary, it is diverted back into the geometry. In the case of a periodic boundary, the neutron is moved to the opposite surface.

Different boundary conditions can be applied in x-, y- and z- surfaces of square cylinder, cube and cuboidal boundary. The syntax is then:

```
set bc <cx> <cy> <cz>
```

where <cx> is the boundary condition in the x-direction
<cy> is the boundary condition in the y-direction
<cz> is the boundary condition in the z-direction

All three entries must be given, even if the geometry is two-dimensional. This capability is available in code version 1.1.17 on.

Symmetries in finite geometries can be taken into account using the universe symmetry option:

```
set usym <uni> <sym> <x> <y>
```

where <uni> is the universe number
<sym> is symmetry type
<x> is the x-coordinate of symmetry origin
<y> is the y-coordinate of symmetry origin

Present version of Serpent allows only quadrant symmetries (<sym> = 4) in universe 0.

IMPORTANT NOTES ON BOUNDARY CONDITIONS:

1. The reflective and periodic boundary conditions can only be used in geometries where the outer boundary surface is either a square or a hexagonal cylinder or a cube.
2. Even though the reflective and the periodic boundary conditions produce the same results in many cases, it should be noted that they are not equivalent when the geometry is asymmetric. This is the case, for example, for BWR assemblies surrounded by an asymmetric moderator channel. Infinite BWR lattices must always be defined using reflective, instead of periodic boundary conditions.
3. If black boundary conditions are used, the outer geometry boundary must be non-reentrant or leakage will produce unphysical results.

4. The universe symmetry option is available from version 1.1.14 on.

SEE ALSO:

1. Outside cells (Sec. 3.3.1 on page 24)

5.8 Source rate normalization

The integral reaction rate estimates given by a Monte Carlo simulation are more or less arbitrarily normalized, unless fixed by a given constant. The Serpent code provides for seven options for source rate normalization.

Normalization to fission neutron generation rate is set using:

```
set genrate <N>
```

where $\langle N \rangle$ is the number of fission neutrons emitted per second

The neutron generation rate includes only prompt and delayed neutrons emitted in fission. All (n,xn) reactions are treated as neutron-multiplying scattering within the criticality cycle. Normalization to source rate is set using:

```
set srcrate <N>
```

where $\langle N \rangle$ is the number of neutrons emitted per second

Normalization to source rate is recommended to be used only in external source calculation mode, in which case the total source rate refers to the rate at which neutrons are emitted from the user-defined source. In criticality source mode, the normalization is done for the number of neutron histories started per generation. Normalization to total fission rate is set using:

```
set fisstrate <N>
```

where $\langle N \rangle$ is the number of fission reactions per second

Normalization to total absorption rate is set using:

```
set abstrate <N>
```

where $\langle N \rangle$ is the number of neutrons absorbed per second

Absorption includes all reactions in which the incident neutron is lost, i.e. all capture reactions and fission. The default normalization is absorption rate set to unity. Normalization to total loss rate is set using:

```
set lossrate <N>
```

where <N> is the number of fission neutrons lost per second

Loss rate includes absorption rate and leakage. Normalization to total flux is set using:

```
set flux <flx>
```

where <flx> is the total neutron flux

Normalization to total heating power is set using:

```
set power <P>
```

where <P> is the total heating power (W)

The total heating power includes all heat generated in the system. If the geometry is two-dimensional, the value is the linear power in W/cm. The source rate normalization can be changed during burnup calculation by re-defining the value between burnup intervals. The first value is used during the first interval, the second during the second interval and so on.

It should be noted that the heating power is *not* the same thing as the total fission power (recoverable fission energy production rate), mainly because a significant fraction of heat is produced in (n,γ) reactions. The direct calculation of this heating power is difficult and Serpent uses an approximation based on the total fission rate and empirical heating values directly proportional to fission energy. The heating value for U-235 fission is 202.27 MeV and the values for other nuclides are scaled according to the ratios of fission Q-values. The U-235 heating value can also be set manually using:

```
set U235H <H>
```

where <H> is the heating value for U-235 (MeV)

Heating values for individual actinides can be overridden using:

```
set fissh <ZAI1> <H1> <ZAI2> <H2> ...
```

where <ZAI n > is the actinide ZAI
<H n > is the heating value

Power density, instead of power can be used for source normalization by setting:

```
set powdens <pde>
```

where <pde> is the average power density (kW/g)

The value is the total heating power divided by the total initial mass of fissile isotopes. This mass is calculated automatically by the code. If the calculation is not possible, the value must be set manually (see Sec. 8.4.2 on page 112).

IMPORTANT NOTES ON SOURCE RATE NORMALIZATION:

1. The source rate normalization affects only integral reaction rates encountered, for example, in detector calculation. Homogenized group constants are unaffected since the normalization cancels out.
2. The default normalization is unit loss rate. It should be noted that the value generally differs from source and generation rates due to neutron-producing reactions.
3. If the geometry is two-dimensional, the values are divided by unit length. The total heating power (W), for example, becomes the linear power (W/cm).
4. Power density is given in units of kW/g, not W/g used in several other codes. The typical order of magnitude for this parameter in LWR calculations is $20\text{E-}3 \dots 50\text{E-}3$.

SEE ALSO:

1. Definition of irradiation history (Sec. 8.3 on page 110)
2. Discussion on source normalization in Ref. [15] (Sec. 9.4 on page 169).
3. Additional options for source rate normalization in burnup calculation (Sec. 8.4.2 on page 112).

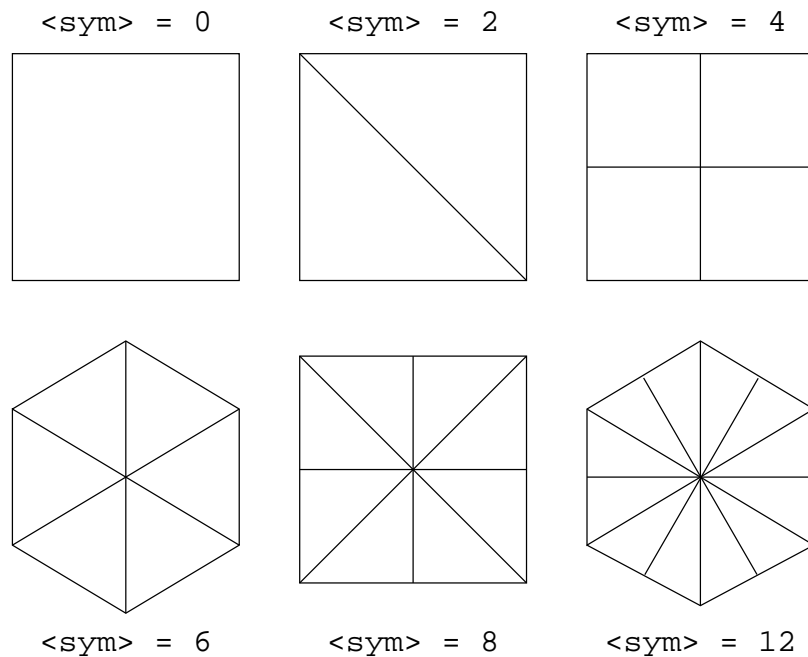


Figure 5.1: Symmetry options.

5.9 Group constant generation

The universes in which the group constants are calculated can be set by:

```
set gcu <u1> <u2> ...
```

where <un> are the universe numbers

The homogenization is carried out in the given universes and all higher universes accessed from lattices and filled cells. The results are printed in the output file (see Sec. 6.1 on page 77) using a different run index for each universe in the list. The default is <u1> = 0, i.e. a single universe that covers the entire geometry. It should be noted that the universe options affect only some of the output parameters, mainly the homogenized group constants. The methodology was included in code version 1.1.5 and is still under development.

The statistical error in assembly discontinuity factors can be reduced by taking advantage of the symmetry of the geometry. The symmetry option is set by:

```
set sym <sym>
```

where <sym> is the symmetry option

The available symmetries are illustrated in Figure 5.1. Options 2, 4 and 8 are used with square lattice geometries and options 6 and 12 with hexagonal geometries. Default option is 0 (no symmetry).

All group constants are generated using the same few-energy group structure. The default structure consists of two energy groups: fast group above 0.625 eV and thermal group below that. This can be overridden by setting the group boundaries manually:

```
set nfg <ne> [<E1> <E2> ...]
```

where <ne> is the number of energy groups
<E1> <E2> ... are the group boundaries (in MeV)

The boundaries are listed in ascending order without the upper and lower limits and the number must be consistent with the number of given values (<ne> - 1 values for <ne> groups).

When it comes to multiplying scattering reactions, such as (n,2n), (n,3n) or (n, 2n α), there is some ambiguity in the way group-to-group scattering matrices and removal cross sections are defined and used in nodal diffusion codes.

The first option is to reflect only the scattering rate, i.e. to disregard the number of neutrons produced in each reaction. In this case, the sum of each matrix column equals the group-wise

total scattering cross section:

$$\sum_{h=1}^G \Sigma_{s,g \rightarrow h} = \Sigma_{s,g} = \Sigma_{\text{ela},g} + \Sigma_{\text{inl},g} + \Sigma_{2n,g} + \Sigma_{3n,g} + \dots = \Sigma_{\text{tot},g} - \Sigma_{\text{capt},g} - \Sigma_{\text{fiss},g} .$$

The second option is to include neutron production in the cross sections, so that the product of group flux and the corresponding group-transfer cross section yields the rate at which neutrons enter group h from scattering reactions in group g , taking into account the multiplication in (n,xn) reactions. The summation to total scattering cross section no longer holds.

Serpent versions from 1.1.15 on calculate both matrixes (see Sec. 6.1.23). The definition of the scattering matrix also affects the removal cross section:

$$\Sigma_{\text{rem},g} = \Sigma_{\text{tot},g} - \Sigma_{s,g \rightarrow g} ,$$

and the whether production of secondary neutrons is included or not is selected by:

```
set remxs <opt>
```

where <opt> is the scattering matrix option (0 = include only scattering rate,
1 = include also production)

The option is available from version 1.1.15 on. The methods used in previous versions correspond to option 0. Option 1 is currently used as the default.

IMPORTANT NOTES ON GROUP CONSTANT GENERATION:

1. The methodology has been thoroughly tested only in cases where group constants are homogenized over the entire geometry. The calculation may produce incorrect results for diffusion coefficients and assembly discontinuity factors if the homogenization is restricted to a higher universe.
2. The list of universes given after the `gcu` option is exclusive. If a collision point is located in several universes in the list, only the highest universe is scored.
3. The use of the symmetry option will lead to erroneous results if the geometry is not truly symmetric.
4. The listed energy values cover only the group boundaries between the minimum and maximum energy of the cross section data. The absolute boundary values are defined in the reconstruction of the master energy grid.
5. The energy groups are indexed in increasing lethargy (decreasing energy) (1 = highest group, <ne> = lowest group).

SEE ALSO:

1. Setting the master energy grid (Sec. 5.3 on page 55)
2. Group constant output (Sec. 6.1 on page 77)

5.10 Full-core power distributions

Serpent can calculate assembly or pin-wise power distributions in full-core simulations. This option is set by:

```
set cpd <depth> [<nz> <zmin> <zmax>]
```

where <depth> is the number of levels included
<nz> is the number axial bins
<zmin> is the lower axial boundary
<zmax> is the upper axial boundary

The level argument determines whether the calculation is carried out at assembly only (1) or both assembly and pin-levels (2). The axial variables determine the number and location of bins in the z-direction.

The code calculates integral fission power inside nested lattice structures (core and assembly lattices). The output data is printed in a separate file named “<input>_core<n>.png”, where <input> is the name of the input file and <n> is the burnup step.

IMPORTANT NOTES ON FULL-CORE POWER DISTRIBUTIONS:

1. This is an experimental feature, available from version 1.1.8 on. The routine has not been thoroughly tested. The results may not be considered reliable, especially when used in combination with the the track-length estimator option.
2. When used in full 3D mode with axial binning, the routine produces very large output files.

SEE ALSO:

1. The track-length estimator option (Sec. 5.18 on page 74)

5.11 Delta-tracking options

The Woodcock delta-tracking tracking method used by Serpent loses its efficiency in the presence of localized heavy absorbers, such as control rods or burnable absorber pins. To overcome this problem, the code switches to the conventional surface-to-surface ray-tracing

when the probability of sampling a physical collision falls below a user-defined threshold. The value is set by:

```
set dt <thresh>
```

where `<thresh>` is the delta-tracking threshold value

This parameter determines the probability limit below which the delta-tracking method is used (0 = never, 1 = always).⁴

Finding the optimal value for the threshold parameter can only be accomplished by trial and error. The default value is 0.9, which seems to work well for most cases.

Serpent also has a built-in optimization routine that tries to find the best value for the cut-off criterion. From version 1.1.9 on the optimization handles each material separately, which has shown to improve the efficiency at least in some complicated HTGR full-core geometries. The optimization is switched on by giving a negative threshold value. This value also serves as the initial guess, so `<thresh> = -0.9` is the recommended choice for optimization.

The use of delta-tracking can be blocked in given materials by setting:

```
set blockdt <mat 1> <mat 2> ...
```

where `<mat 1> <mat 2> ...` are the materials where delta-tracking will not be used

The tracking routine in serpent selects between surface and delta-tracking, based on the cut-off criterion described above. Some geometries may run faster, however, if surface tracking is always used in very large material regions comprised of simple cells. A good example of such region is the outer reflector in a full-core geometry. It should be noted, however, that this option may also impair the efficiency if not properly used.

IMPORTANT NOTES ON DELTA-TRACKING:

1. The cut-off value is set to 0.9 by default in code version 1.1.1 and later. Earlier versions use the optimization by default. The optimization routine was changed in update 1.1.9 to handle each material separately.
2. The optimization has not been thoroughly tested and the methodology is not guaranteed to result in the optimal threshold value in terms of CPU time.
3. The code should always yield consistent results with and without delta-tracking. If any discrepancies are observed, please report them by e-mail to Jaakko.Leppanen@vtt.fi
4. The block option is available from version 1.1.8 on.

⁴The delta-tracking method is essentially a rejection probability sampling technique, and the threshold parameter determines the highest rejection probability at which the method is used. If the probability is higher than the threshold value, the code switches to the conventional ray-tracing method.

SEE ALSO:

1. Description of the basic Woodcock delta-tracking method in Ref. [15] (Sec. 5.3.3 on page 100).
2. Description of the extended delta-tracking method used in PSG in Ref. [15] (Sec. 8.3.1 on page 149). NOTE: The described methods are partially outdated.
3. A more recent description of the method is found in Ref. [18].

5.12 Cross section data plotter

Serpent has the option to plot all cross sections in a matlab m-file format. The cross section data plotter is activated using:

```
set xsplot [<ne> <Emin> <Emax> ]
```

where <ne> is the number of energy points in plot
 <Emin> is the lower limit of the energy grid (MeV)
 <Emax> is the upper limit of the energy grid (MeV)

The energy grid used for the plot is uniform with respect to the lethargy variable. The plotter produces a file “<input>_xs<n>.png”, where <input> is the name of the input file and <n> is the burnup step. The file contains the energy grid vector, isotopic reaction cross sections, material total cross sections and fission nubar.

5.13 Fission source entropy

The fission source entropy for convergence studies is calculated by default and the total entropy is divided in x-, y- and z-components. The entropy mesh is set by:

```
set entr [<nx> <ny> <nz> <x0> <x1> <y0> <y1> <z0> <z1>]
```

where <nx> is the number of x bins
 <ny> is the number of y bins
 <nz> is the number of z bins
 <x0> is the minimum x-coordinate in mesh
 <x1> is the maximum x-coordinate in mesh
 <y0> is the minimum y-coordinate in mesh
 <y1> is the maximum y-coordinate in mesh
 <z0> is the minimum z-coordinate in mesh
 <z1> is the maximum z-coordinate in mesh

The source entropies are written in the history output file as function of criticality cycle.

SEE ALSO:

1. History output (Sec. 6.2 on page 94)
2. Discussion on fission source entropy in Ref. [14].

5.14 Soluble absorber

Materials with soluble absorber, most commonly boron in light water, can be defined by mixing two material compositions. This is considerably simpler than explicitly listing the associated isotopic fractions. The soluble absorber is defined using:

```
set abs <solu> <conc> <mat 1> <mat 2> ...
```

where	<code><solu></code>	is the soluble absorber material name
	<code><conc></code>	is the absorber concentration
	<code><mat 1> <mat 2> ...</code>	are the materials where the absorber is dissolved

The code mixes material “<abs>” into materials “<mat 1>”, “<mat 2>” ... in concentration defined by “<conc>”. Positive concentrations refer to atomic fractions and negative concentrations to mass fractions. A simple example is given in the VVER-440 calculation case in Sec. 11.1.1 on page 134.

The absorber concentration can be changed during burnup calculation by re-defining the value between burnup intervals. The first value is used during the first interval, the second during the second interval and so on.

IMPORTANT NOTES ON SOLUBLE ABSORBER:

1. If soluble absorber is used with multiple materials, all must share the same isotopic composition.
2. Only the total absorption channel of the absorber material is used and fission, scattering and all the other reaction modes are discarded. This is a good approximation if the concentration is low and the material is high-absorbing. The maximum concentration for natural boron in water is around few-thousand ppm per weight. If the concentration is higher, it is better to determine the isotopic composition explicitly.
3. The methodology is available from code version 1.0.2 on.

SEE ALSO:

1. Definition of irradiation history (Sec. 8.3 on page 110)

5.15 Iteration

k_{eff} can be iterated to a desired value by allowing the variation in some geometry, material or physics variable. Iteration is defined by:

```
set iter <mode> <keff> [<spec> <ne>]
```

where <mode> is the iteration mode
 <keff> is the target k_{eff}
 <opt> is the leakage spectrum mode (B_1 -iteration only)
 <ne> is number of energy bins in the spectrum (B_1 -iteration only)

The iteration modes are:

- Iteration of soluble absorber concentration, <mode> = “abs”
- α -eigenvalue calculation, <mode> = “alpha”
- Iteration of albedo boundary condition, <mode> = “albedo”
- B_1 iteration, <mode> = “B1”

The soluble absorber iteration works by varying the concentration of soluble absorber (see Sec. 5.14) to yield the desired k_{eff} .

The α -eigenvalue mode is a standard transport technique that allows time-absorption or -multiplication to balance neutron source and loss rates. The “cross section” for the reaction is equal to the α -eigenvalue divided by neutron speed. The calculation is basically equivalent with a time-dependent simulation.

The albedo boundary condition iteration is an attempt to simulate the effects of neutron leakage in an infinite-lattice geometry. The method works by sampling leakage ($k > 1$) or multiplication reactions ($k < 1$) each time a neutron crosses a repeated or periodic boundary. It should be noted that this method is highly experimental, and does not have physical foundation similar to deterministic leakage models.

The second experimental leakage model is the B_1 iteration. The method works similar to the α -eigenvalue simulation: leakage absorption or multiplication reactions are introduced to balance neutron source and loss rates. The cross section for the reaction is equal to the B_1 -factor multiplied by the leakage spectrum, given by the energy-dependence of the volume-integrated diffusion coefficient. Since the diffusion coefficient cannot be defined as a continuous-energy parameter, the code calculates a fine-group spectrum using an estimate based on the diffusion area and the removal cross section (<spec> = 1) or the P1-approximation (<spec> = 2). The energy variable is divided into <ne> equal lethargy-width bins (default = 500).

IMPORTANT NOTES ON ITERATION:

1. When iteration is used in burnup calculation mode, the procedure is repeated independently for each burnup step.
2. Soluble absorber must be defined in the absorber iteration mode.
3. The α -eigenvalue calculation, albedo iteration and B_1 mode are available from update 1.1.5 on.
4. The albedo- and B_1 -iteration modes are experimental, rather than standard Monte Carlo techniques. The theory is not on a particularly solid foundation and the results are generally not satisfactory when compared to deterministic calculations.
5. The B_1 iteration mode must not be confused with the fundamental mode calculation, discussed in Section 5.16.
6. The α -eigenvalue simulation is a widely-used method, but the implementation in Serpent has not been validated. The mode does not account for delayed neutron emission.
7. Some of the k_{eff} estimates are different from a standard calculation, depending on the iteration mode used.

SEE ALSO:

1. Definition of soluble absorber (Sec. 5.14 on page 69)
2. Ref. [19] and Sec. 5.5.2 in Ref. [15] for discussion on the α -eigenvalue method.
3. Diffusion coefficients in output (Sec. 6.1.27 and 6.1.29).
4. Discussion on neutron leakage models in Monte Carlo calculation in Sec. 9.5 in Ref. [15]

5.16 Fundamental mode calculation

The fundamental mode calculation can be considered as an intermediate solution to the criticality spectrum problem, until the development of a valid Monte Carlo based leakage correction. The calculation consists of two stages. First, the Monte Carlo simulation is run to produce homogenized micro-group cross sections for B_1 equations. The solution of these equations yields the criticality spectrum, which is used to re-homogenize the cross sections.

The syntax is:

```
set fum <egrid>
```

where `<egrid>` is the micro-group structure used for the calculation

The energy grid determines the micro-group structure used to form the B_1 equations and it is set up using the “ene” parameter (see Sec. 7.1.2). The method produces a separate set of

output parameters (see Sec. 6.1.30) and does not affect the values of other group constants. The energy group boundaries in the few-group structure must match the boundaries in the micro-group structure.

IMPORTANT NOTES ON FUNDAMENTAL MODE CALCULATION:

1. The fundamental mode calculation is available from version 1.1.14 on. The spectrum correction affects only a set of separately produced few-group constants. Extending the correction to burnup calculation is under development.
2. The group boundaries in the few-group structure must match the boundaries in the micro-group structure.
3. Relative statistical errors are not included in the results.
4. The fundamental mode calculation must not be confused with the experimental B_1 iteration, discussed in Section 5.15.

SEE ALSO:

1. Definition of energy grids (Sec. 7.1.2 on page 99).
2. Output parameters for fundamental mode calculation (Sec. 6.1.30 on page 92).
3. Definition of the few-group structure (Sec. 5.9 on page 64).

5.17 Equilibrium xenon calculation

Serpent can iterate the concentration of fission product poison Xe-135 to an equilibrium value in transport or burnup calculation. The equilibrium xenon calculation is set by:

```
set xenon <mode> [<mat 1> <mat 2> ...]
```

where <mode> is the calculation mode (0 = off, 1 = on)
<mat 1> <mat 2> ... are the materials involved in the calculation

The mode option is followed by a list of materials for which the calculation is turned on or off. If no list is given, the option affects all fissile materials. Each material is handled separately.

The calculation is based on the production rates of Xe-135 and its precursors (I-135, Te-135, Sb-135 and Sn-135), the absorption rate of Xe-135 and the radioactive decay of the isotopes. The production and absorption rates are normalized to source rate. The decay and fission yield data are read from external libraries, similar to a burnup calculation.

IMPORTANT NOTES ON EQUILIBRIUM XENON CALCULATION:

1. The equilibrium concentration depends on source rate normalization.
2. When used in the burnup calculation mode, the concentration of Xe-135 is handled separate from the other nuclides. The equilibrium concentration is copied in the depletion output.
3. The capability was included in code version 1.1.9 and currently it may not work with unresolved resonance probability table sampling, soluble absorbers or k-eff iteration.

SEE ALSO:

1. Source rate normalization (Sec. 5.8 on page 61)
2. Setting the decay and fission yield library file paths (Sec. 8.4 on page 111)

5.18 Miscellaneous parameters

A title string for the calculation can be set using

```
set title "<ttl>"
```

where <ttl> is the title string

This text string is reproduced in the output files together with date and time and version information.

The Monte Carlo simulation uses a sequence of random numbers, generated from an initial seed value. This seed is by default taken from system time. The calculation can be reproduced using the “replay” command line option, which forces the code to use the same seed as in the previous run. The seed value can also be set manually using:

```
set seed <val>
```

where <val> is the seed value (a large integer)

Temperatures used in the free-gas model for elastic scattering are read from the ACE format data. The free-gas temperatures in cells can be overridden by defining a list of cell temperatures:

```
set ctmp <cell 1> <T1> <cell 2> <T2> ...
```

where <cell n> are the cell names
<Tn> are the temperatures

User-defined variables can be set up for labeling different runs. The syntax is:

```
set var <name> <value>
```

where <name> is the variable name
<value> is the value

The variable name and value are printed in the main output (see Sec. 6.1 on page 77). The type (numeric or string) is identified from the value.

The use of track-length flux estimate can be forced in place of the collision estimator using:

```
set tle <n>
```

where <n> is the option (1 = use tle, 0 = use cfe)

If the track-length estimator is used, delta-tracking is switched off.

By default, Serpent uses various pre-calculated summation cross sections for each material to speed-up the transport simulation. This increases the overall memory demand per material, which may become a limiting factor in burnup calculation. To reduce the demand, the calculation can be switched off using:

```
set sumxs <use>
```

where <use> is the option (1 = use pre-calculated cross sections, 0 = calculate cross sections on-the-fly)

It should be noted that switching off the option results in an increase in the overall calculation time. The option is available from version 1.1.13 on.

The emission of delayed neutrons can be switched on and off using:

```
set delnu <use>
```

where <use> is the option (1 = emission on, 0 = emission off)

This option was added in version 1.1.16. Delayed neutron emission is on by default in criticality source problems and off in external source problems.

Calculation of fission product poison cross section (production of I-135, Xe-135, Pm-149 and Sm-149 and absorption of Xe-135 and Sm-149) can be switched on and off by setting:

```
set poi "<opt>"
```

where <opt> is the option (0 = off, 1 = on)

Calculation is off by default. Switching the mode on requires setting the file path for fission yield data (see Sec. 8.4.1). This feature is available from version 1.1.17 on.

SEE ALSO:

1. Running the code in replay mode (Sec. 1.2 on page 9)
2. Main output file (Sec. 6.1 on page 77)

Table 5.1: List of parameters and options.

Option		Description	Section	Page
pop	(3-4)	population size and number of cycles	5.2	53
nbuf	(1)	source buffer	5.2	53
egrid	(1-3)	energy grid reconstruction	5.3	55
dix	(1)	double indexing of energy grids	5.3	56
acelib	(1)	file path for xs library directory file	5.4	57
ures	(1-N)	probability table treatment for ures data	5.5	58
dbrc	(3-N)	DBRC correction for scattering kernel	5.6	59
bc	(1)	boundary conditions	5.7	59
usym	(2-4)	universe symmetry	5.7	60
genrate	(1)	source normalisation to generation rate	5.8	61
srcrate	(1)	source normalisation to source rate	5.8	61
fissrate	(1)	source normalisation to fission rate	5.8	61
absrate	(1)	source normalisation to absorption rate	5.8	61
lossrate	(1)	source normalisation to loss rate	5.8	62
flux	(1)	source normalisation to total flux	5.8	62
power	(1)	source normalisation to total heating power	5.8	62
powdens	(1)	source normalisation to power density	5.8	62
U235H	(1)	heating value for U-235 fission	5.8	62
fissh	(1-N)	fission heating values for individual actinides	5.8	62
gcu	(1)	universe for homogenization	5.9	64
sym	(1)	symmetry option	5.9	64
nfg	(1-N)	few-group structure for homogenization	5.9	64
remxs	(1)	scattering matrix used with removal cross section	5.9	65
cpd	(1)	full-core power distributions	5.10	66
dt	(1)	delta-tracking threshold	5.11	67
blockdt	(1)	delta-tracking block	5.11	67
xsplot	(1-4)	cross section data plot file	5.12	68
entr	(1-9)	parameters for source entropy calculation	5.13	68
abs	(3-N)	soluble absorber	5.14	69
iter	(2)	k_{eff} iterations	5.15	70
fum	(2)	fundamental mode calculation	5.16	71
xenon	(1-N)	equilibrium Xe-135 calculation	5.17	72
title	(1)	case title	5.18	73
seed	(1)	random number seed value	5.18	73
ctmp	(1)	override cell temperatures	5.18	73
var	(1)	user-defined variable	5.18	74
tle	(1)	track-length estimate of neutron flux	5.18	74
sumxs	(1)	use pre-calculated summation cross sections	5.18	74
delnu	(1)	switch delayed neutron emission on and off	5.18	74
poi	(1)	fission product poison cross sections	5.18	74

Chapter 6

Output

6.1 Main output file

The main output file contains all results calculated by default during the transport cycle. User-defined detectors produce another file, described in Section 7.2 on Page 105. Inventory data in burnup calculation is discussed in Section 8.5 on Page 116.

The file is named “<input>_res.m”, where “<input>” is the name of the input file. The data is written in matlab m-file format to simplify the simultaneous post-processing of several files. Each parameter is read to a variable (scalar or vector) and a run index “idx” is assigned to each file. Each time a new file is read, the index is first increased by, 1 so that the new data is placed on the next line in the result matrix. The following Octave example illustrates the procedure:¹

```
octave:1> idx = 0
idx = 0
octave:2> run1_res;
octave:3> FISSXS
FISSXS =

    0.0160550  0.0005253  0.0033174  0.0006745  0.0863956  0.0005001

octave:4> run2_res;
octave:5> FISSXS
FISSXS =

    0.0160550  0.0005253  0.0033174  0.0006745  0.0863956  0.0005001
    0.0158454  0.0005277  0.0032059  0.0006817  0.0833996  0.0005078
```

¹GNU Octave is a Matlab-compatible open-source language for numerical computations.

```

octave:6> run3_res;
octave:7> FISSXS
FISSXS =

    0.0160550  0.0005253  0.0033174  0.0006745  0.0863956  0.0005001
    0.0158454  0.0005277  0.0032059  0.0006817  0.0833996  0.0005078
    0.0119486  0.0005737  0.0031909  0.0005741  0.0694696  0.0005300

```

Three input files: “run1_res.m”, “run2_res.m” and “run3_res.m” are read and the data from each file is placed on a different row in the variables. Variable “FISSXS” is the homogenized fission cross section, calculated in this case using a two-energy group structure. The first two columns are the total (one-group) value and the associated relative statistical error, respectively. The following four columns contain the same data for the two energy groups in ascending order.

Output data in burnup calculation is written in a single file. The run index is updated for each burnup step. The variables in the main output file are listed in the following.

6.1.1 Version, title and date

Parameter	Values	Description
VERSION	1	Code version used in calculation
TITLE	1	Case title
DATE	1	Date and time at the beginning

6.1.2 Run parameters

Parameter	Values	Description
POP	1	Number of source neutrons per cycle
CYCLES	1	Number of active cycles
SKIP	1	Number of inactive cycles
SRC_NORM_MODE	1	Fission source normalization mode (1 = preserve size, 2 = preserve weight)
SEED	1	Random number seed
MPI_TASKS	1	Number of MPI tasks in parallel calculation
MPI_MODE	1	Results collection in MPI mode
DEBUG	1	Debug mode flag (1 = yes, 0 = no)
CPU_TYPE	1	CPU type
CPU_MHZ	1	CPU MHz
AVAIL_MEM	1	Available memory in Mb

NOTES:

1. In parallel calculation mode, the number of source neutrons per cycle is the number for each parallel task.
2. CPU type and MHz are read from /proc/cpuinfo and available memory from /proc/meminfo.

6.1.3 File paths

Parameter	Values	Description
XS_DATA_FILE_PATH	1	Cross section directory file path
DECAY_DATA_FILE_PATH	1	Decay data file path
NFY_DATA_FILE_PATH	1	Fission yield data file path

NOTES:

1. Only the first given xs directory file path is printed

6.1.4 Delta-tracking parameters

Parameter	Values	Description
DT_THRESH	1	Probability threshold for using delta-tracking
DT_FRAC	1	Fraction of path lengths sampled using delta-tracking
DT_EFF	1	Efficiency of DT rejection algorithm
MIN_MACROXS	1	Minimum macroscopic cross section for sampling the collision distance

6.1.5 Run statistics

Parameter	Values	Description
TOT_CPU_TIME	1	Total CPU time
RUNNING_TIME	1	Cumulative total running time (wall-clock)
CPU_USAGE	1	CPU usage (ratio of CPU time to wall-clock time)
INIT_TIME	1	Total initialization time before transport or burnup cycle
TRANSPORT_CYCLE_TIME	1	Cumulative transport cycle running time
BURNUP_CYCLE_TIME	1	Cumulative time used for solving the depletion equations
PROCESS_TIME	1	Cumulative time used for data processing between transport cycles
CYCLE_IDX	1	Current cycle index
SOURCE_NEUTRONS	1	Number of simulated source neutrons
MEAN_POP_SIZE	1	Mean population size
MEMSIZE	1	Size of allocated memory block in megabytes
SIMULATION_COMPLETED	1	Flag to indicate that all neutron histories are run (1 = yes, 0 = no)

NOTES:

1. The total RUNNING_TIME is the sum of INIT_TIME, PROCES_TIME, TRANSPORT_CYCLE_TIME and BURNUP_CYCLE_TIME.

6.1.6 Energy grid parameters

Parameter	Values	Description
ERG_EMIN	1	Minimum energy in unionized grid (MeV)
ERG_EMAX	1	Maximum energy in unionized grid (MeV)
ERG_TOL	1	Fractional grid reconstruction tolerance
ERG_NE	1	Number of grid points
ERG_NE_INI	1	Number of grid points before thinning
ERG_NE_IMP	1	Number of important grid points
ERG_DIX	1	Double indexed energy grids (1 = yes, 0 = no)
USE_DBRC	1	Doppler-broadening rejection correction (1 = yes, 0 = no)

6.1.7 Unresolved resonance data

Parameter	Values	Description
URES_MODE	1	Probability table sampling mode
URES_DILU_CUT	1	Infinite dilution cut-off
URES_EMIN	1	Minimum energy for unresolved resonance probability table data (MeV)
URES_EMAX	1	Maximum energy for unresolved resonance probability table data (MeV)
URES_AVAIL	1	Number of isotopes with ures data available
URES_USED	1	Number of isotopes with ures data used

6.1.8 Nuclides and reaction channels

Parameter	Values	Description
TOT_ISOTOPES	1	Total number of isotopes
TOT_TRANSPORT_ISOTOPES	1	Total number of isotopes with cross section data
TOT_DECAY_ISOTOPES	1	Total number of isotopes without cross section data
TOT_REA_CHANNELS	1	Total number of reaction channel
TOT_TRANSMU_REA	1	Total number of transmutation reactions

NOTES:

1. TOT_REA_CHANNELS includes neutron reactions only, no decay.

6.1.9 Reaction mode counters

Parameter	Values	Description
COLLISIONS	1	Total number of collisions
FISSION_FRACTION	1	Fraction of fission reactions
CAPTURE_FRACTION	1	Fraction of capture reactions
ELASTIC_FRACTION	1	Fraction of elastic scattering reactions
INELASTIC_FRACTION	1	Fraction of inelastic scattering reactions
ALPHA_FRACTION	1	Fraction of time-absorption or -multiplication reactions in α -eigenvalue calculation mode
BOUND_SCATTERING_FRACTION	1	Fraction of bound atom scattering reactions
NXN_FRACTION	1	Fraction of (n,xn) reactions
UNKNOWN_FRACTION	1	Fraction of unsampled reactions
VIRTUAL_FRACTION	1	Fraction of virtual collisions
FREGAS_FRACTION	1	Fraction of free-gas elastic scattering reactions
TOTAL_ELASTIC_FRACTION	1	Fraction of free and bound atom elastic scattering reactions
FISSILE_FISSION_FRACTION	1	Fraction of fission reactions in fissile isotopes
LEAKAGE_REACTIONS	1	Number of leakage reactions
REA_SAMPLING_EFF	1	Reaction mode sampling efficiency

NOTES:

1. Leakage in B1 and albedo iteration modes is counted in LEAKAGE_REACTIONS

6.1.10 Slowing-down and thermalization

Parameter	Values	Description
COL_SLOW	2	Average number of collisions before thermalization
COL_THERM	2	Average number of collisions after thermalization
COL_TOT	2	Average total number of collisions
SLOW_TIME	2	Average slowing-down time
THERM_TIME	2	Average thermal life time
SLOW_DIST	2	Average slowing-down distance
THERM_DIST	2	Average thermal migration distance
THERM_FRAC	2	Average fraction of neutrons reaching thermalization

6.1.11 Parameters for burnup calculation

Parameter	Values	Description
BURN_MODE	1	Burnup mode (1 = TTA, 2 = CRAM)
BURN_STEP	1	Burnup step index
BURN_TOT_STEPS	1	Total number of burnup steps
BURNUP	1	Burnup at current step (in MWd/kgU)
BURN_DAYS	1	Number of burn days at current step
ENERGY_OUTPUT	1	Total cumulative energy output (in J)
DEP_TTA_CUTOFF	1	TTA cut-off value
DEP_STABILITY_CUTOFF	1	Stability cut-off value
DEP_FP_YIELD_CUTOFF	1	Fission product yield cut-off value
DEP_XS_FRAC_CUTOFF	1	Depletion fraction cut-off value
DEP_XS_ENERGY_CUTOFF	1	Depletion reaction energy cut-off value
BURN_MATERIALS	1	Number of depleted materials
FP_NUCLIDES_INCLUDED	1	Total number of fission product nuclides included in the calculation
FP_NUCLIDES_AVAILABLE	1	Total number of fission products available before yield cut-off
TOT_ACTIVITY	1	Total activity at current step
TOT_DECAY_HEAT	1	Total decay heat at current step (in W)
TOT_SF_RATE	1	Total spontaneous fission rate
ACTINIDE_ACTIVITY	1	Actinide activity at current step
ACTINIDE_DECAY_HEAT	1	Actinide decay heat at current step (in W)
FISSION_PRODUCT_ACTIVITY	1	Fission product activity at current step
FISSION_PRODUCT_DECAY_HEAT	1	Fission product decay heat at current step (in W)
DH_N_PREC	1	Number of decay heat precursor groups
DH_PREC_BOUNDS	$J_d + 1$	Decay heat precursor group boundaries
DH_PREC_LAMBDA	J_d	Decay heat group-wise decay constants
DH_PREC_HEAT	J_d	Decay heat group-wise heat production (in W)

NOTES:

1. Precursor-group wise decay heat production is available from version 1.1.17 on. The option for setting the group boundaries is described in Sec. 8.4.8.

6.1.12 Fission source entropies

Parameter	Values	Description
ENTROPY_X	2	X-component of fission source entropy
ENTROPY_Y	2	Y-component of fission source entropy
ENTROPY_Z	2	Z-component of fission source entropy
ENTROPY_TOT	2	Total fission source entropy

6.1.13 Fission source center

Parameter	Values	Description
SOURCE_X0	2	X-coordinate of fission source center
SOURCE_Y0	2	Y-coordinate of fission source center
SOURCE_Z0	2	Z-coordinate of fission source center

6.1.14 Soluble absorber

Parameter	Values	Description
SOLU_ABS_AFRAC	1	Atomic fraction of soluble absorber
SOLU_ABS_MFRAC	1	Mass fraction of soluble absorber

NOTES:

1. The values are printed only if soluble absorber defined (see Sec. 5.14 on page 69).

6.1.15 Iteration

Parameter	Values	Description
ITER_MODE	1	Iteration mode
ITER_KEFF	1	Target k_{eff} for iteration
ITER_VAR	2	Iteration variable
B1_MODE	1	Method used for calculating leakage spectrum in B1 iteration mode
B1_NE	1	Number of equal lethargy-width bins in the leakage spectrum
B1_ERG	B1_NE	Energy bin limits for the leakage spectrum
B1_SPECTR	B1_NE	Cycle-averaged leakage spectrum

NOTES:

1. The values are printed only if iteration is in use (see Sec. 5.15 on page 70).

6.1.16 Equilibrium Xe-135 calculation

Parameter	Values	Description
XE135_EQUIL_CONC	2	Equilibrium Xe-135 concentration
I135_EQUIL_CONC	2	Equilibrium I-135 concentration

NOTES:

1. The values are printed only if xenon iteration is in use (see Sec. 5.17 on page 72).
2. The concentrations are averaged over all regions involved in the iteration

6.1.17 Criticality eigenvalues

Parameter	Values	Description
ANA_KEFF	2	Analog estimate of k_{eff}
IMP_KEFF	2	Implicit estimate of k_{eff}
COL_KEFF	2	Collision estimate of k_{eff}
ABS_KEFF	2	Absorption estimate of k_{eff}
ABS_KINF	2	Absorption estimate of k_{∞}
ABS_GC_KEFF	2	Absorption estimate of k_{eff} in group constant generation universe
ABS_GC_KINF	2	Absorption estimate of k_{∞} in group constant generation universe
EXT_K	10	External source multiplication factor in 5 generations
IMPL_ALPHA_EIG	2	Implicit estimate of α -eigenvalue
FIXED_ALPHA_EIG	2	Fixed or iterated value in α -eigenvalue calculation
GEOM_ALBEDO	2	Fixed or iterated value for albedo

NOTES:

1. The absorption estimate of k_{eff} is currently used as the implicit estimate.
2. External source multiplication factor is not printed in criticality source mode.

6.1.18 Normalization

Parameter	Values	Description
TOT_POWER	2	Total power
TOT_GENRATE	2	Total neutron generation rate
TOT_FISSRATE	2	Total fission rate
TOT_ABSRATE	2	Total absorption rate
TOT_LEAKRATE	2	Total leakage rate
TOT_LOSSRATE	2	Total loss rate
TOT_SRCRATE	2	Total source rate
TOT_FLUX	2	Total flux
TOT_RR	2	Total reaction rate
TOT_SOLU_ABSRATE	2	Total absorption rate in soluble absorber
TOT_XE135_ABSRATE	2	Total absorption rate in Xe-135
TOT_FMASS	1	Total fissile mass
TOT_POWDENS	2	Total power density
BURN_POWER	2	Power in burnable materials
BURN_GENRATE	2	Neutron generation rate in burnable materials
BURN_FISSRATE	2	Fission rate in burnable materials
BURN_ABSRATE	2	Absorption rate in burnable materials
BURN_FLUX	2	Flux in burnable materials
BURN_FMASS	1	Fissile mass in burnable materials
BURN_POWDENS	2	Power density in burnable materials
BURN_VOLUME	1	Total combined volume of all burnable materials

NOTES:

1. Normalization is set by the user (see Sec. 5.8 on page 61).
2. By default, the loss rate is normalized to unity.
3. Total power density is printed only if total fissile mass is calculated or given by the user.
4. Parameters in burnable materials are printed only in burnup calculation mode.
5. Total (external) source rate is not printed in criticality source mode.
6. Xe-135 absorption rate is printed only in equilibrium xenon mode.
7. All flux values are divided by volume

6.1.19 Point-kinetic parameters

Parameter	Values	Description
ANA_PROMPT_LIFETIME	2	Analog estimate of prompt neutron lifetime
IMPL_PROMPT_LIFETIME	2	Implicit estimate of prompt neutron lifetime
ANA_REPROD_TIME	2	Analog estimate of neutron reproduction time
IMPL_REPROD_TIME	2	Implicit estimate of neutron reproduction time
DELAYED_EMTIME	2	Mean delayed neutron emission time

NOTES:

1. The neutron reproduction time is also commonly known as the “neutron generation time”.
2. The analog estimates and delayed neutron emission time are calculated for the entire geometry. The implicit estimates are calculated in the universe set by the user.

6.1.20 Six-factor formula

Parameter	Values	Description
SIX_FF_ETA	2	Average number of neutrons emitted per thermal neutron absorbed in fuel
SIX_FF_F	2	Thermal utilization factor
SIX_FF_P	2	Resonance escape probability
SIX_FF_EPSILON	2	Fast fission factor
SIX_FF_LF	2	Fast non-leakage probability
SIX_FF_LT	2	Thermal non-leakage probability
SIX_FF_KINF	2	Six-factor k_{∞} (four-factor k_{eff})
SIX_FF_KEFF	2	Six-factor k_{eff}

NOTES:

1. The parameters are calculated using simple analog estimates and intended mainly for the demonstration of basic reactor physics phenomena.

6.1.21 Delayed neutron parameters

Parameter	Values	Description
USE_DELNU	1	Delayed neutron emission (0 = off, 1 = on)
PRECURSOR_GROUPS	1	Number of delayed neutron precursor groups
BETA_EFF	$2J_d + 2$	Effective delayed neutron fraction
BETA_ZERO	$2J_d + 2$	Physical delayed neutron fraction
DECAY_CONSTANT	$2J_d + 2$	Precursor group-wise decay constants

NOTES:

1. The number of precursor groups J_d depends on data. The usual number is 6 or 8. The first two entries refer to the total value and the associated relative statistical error.
2. Since different precursor group structures cannot be combined, the number of groups is fixed to the value used in the first actinide in the input. Delayed neutron emission is entirely omitted for nuclides using a different group structure.

6.1.22 Parameters for group constant generation

Parameter	Values	Description
GC_UNI	1	Universe for group constant generation
GC_SYM	1	Symmetry option
GC_NE	1	Number of energy groups
GC_BOUNDS	$G + 1$	Group boundaries

6.1.23 Few-group cross sections

Parameter	Values	Description
FLUX	$2G + 2$	Integral flux
LEAK	$2G + 2$	Leakage rate
TOTXS	$2G + 2$	Total cross section
FISXS	$2G + 2$	Fission cross section
CAPTXS	$2G + 2$	Capture cross section
ABSXS	$2G + 2$	Absorption cross section
RABSXS	$2G + 2$	Reduced absorption cross section
ELAXS	$2G + 2$	Elastic scattering cross section
INELAXS	$2G + 2$	Inelastic scattering cross section
SCATTXS	$2G + 2$	Total scattering cross section
SCATTPRODXS	$2G + 2$	Total scattering production cross section
N2NXS	$2G + 2$	(n,2n) cross section
REMXS	$2G + 2$	Group-removal cross section
NUBAR	$2G + 2$	Average number of emitted fission neutrons
NSF	$2G + 2$	Fission neutron production cross section ($\nu \Sigma_{fissg}$)
RECIPVEL	$2G + 2$	Inverse mean neutron speed
FISSE	$2G + 2$	Average fission heating value (in MeV)

MAJOR FLAW IN CALCULATION METHODS:

Earlier code versions, including base version 1.1.0, contain a serious flaw in group constant calculation. The collision flux estimator yields zero values in void regions, resulting in a systematic over-prediction of the homogenized values. The problem was fixed in code update 1.1.3.

NOTES:

1. The first two entries are the total (one-group) value and the associated relative statistical error. The remaining $2G$ entries are few-group values.
2. The normalization of group-flux does not work in the pre-release version 1.0.0 of the Serpent code (corrected in version 1.0.1). The one-group value should be equal to variable TOT_FLUX (see Sec. 6.1.18).
3. All cross sections are macroscopic.
4. Capture cross section includes all (n,0n) reactions.
5. Absorption cross section includes capture and fission.
6. Elastic scattering includes thermal bound-atom reactions.
7. Group-removal cross section includes absorption and scattering out of the energy group. Option to include neutron multiplication was added in version 1.1.15 (see Sec. 5.9).
8. Reduced absorption cross section is defined as absorption minus production in (n,xn) reactions.

6.1.24 Fission product poison cross sections

Parameter	Values	Description
I135PRODXS	$2G + 2$	Production cross section for I-135
XE135PRODXS	$2G + 2$	Production cross section for Xe-135
PM149PRODXS	$2G + 2$	Production cross section for Pm-149
SM149PRODXS	$2G + 2$	Production cross section for Sm-149
XE135ABSXS	$2G + 2$	Absorption cross section of Xe-135
SM149ABSXS	$2G + 2$	Absorption cross section of Sm-149

NOTES:

1. The option to switch on the calculation of fission product poison cross sections is described in Sec. 5.18

2. All values are microscopic cross sections
3. Available from version 1.1.17 on

6.1.25 Fission spectra

Parameter	Values	Description
CHI	$2G$	Energy spectrum of all fission neutrons
CHIP	$2G$	Energy spectrum of prompt fission neutrons
CHID	$2G$	Energy spectrum of delayed fission neutrons

6.1.26 Group-transfer probabilities and cross sections

Parameter	Values	Description
GTRANSFP	$2G^2$	Group-transfer probability matrix
GTRANSFXS	$2G^2$	Group-transfer cross section matrix
GPRODP	$2G^2$	Group-production probability matrix
GPRODXS	$2G^2$	Group-production cross section matrix

NOTES:

1. The matrices are given in vector format:

$$P_{1 \rightarrow 1} \ P_{2 \rightarrow 1} \ ... \ P_{G \rightarrow 1} \ P_{1 \rightarrow 2} \ P_{2 \rightarrow 2} \ ... \ P_{G \rightarrow 2} \ ...$$

Each probability and cross section is followed by the associated relative statistical error. Index for reaction $j \rightarrow i$ is given by:

$$n = 2(i - 1)G + 2j - 1$$

2. The production matrixes include neutron multiplication in (n,xn) reactions.

6.1.27 Diffusion parameters

Parameter	Values	Description
DIFFAREA	$2G + 2$	Diffusion area
DIFFCOEF	$2G + 2$	Diffusion coefficient
TRANSPXS	$2G + 2$	Transport cross section
MUBAR	$2G + 2$	Average scattering angle
MAT_BUCKLING	$2G + 2$	Material buckling
LEAK_DIFFCOEF	$2G + 2$	Diffusion coefficient from leakage mode

NOTES:

1. The first two entries are the total (one-group) value and the associated relative statistical error. The remaining $2G$ entries are few-group values.
2. The values are based on the analog estimate of group-wise diffusion area. The results usually differ from the P_1 -values below.
3. Leakage diffusion coefficient is defined as buckling divided by leakage, which can be physical or from a leakage model. The theoretical basis is very questionable.

6.1.28 P_n scattering cross sections

Parameter	Values	Description
SCATT0	$2G + 2$	P_0 scattering cross section
SCATT1	$2G + 2$	P_1 scattering cross section
SCATT2	$2G + 2$	P_2 scattering cross section
SCATT3	$2G + 2$	P_3 scattering cross section
SCATT4	$2G + 2$	P_4 scattering cross section
SCATT5	$2G + 2$	P_5 scattering cross section

NOTES:

1. The first two entries are the total (one-group) value and the associated relative statistical error. The remaining $2G$ entries are few-group values.

6.1.29 P_1 diffusion parameters

Parameter	Values	Description
P1_TRANSPXS	$2G + 2$	Transport cross section
P1_DIFFCOEF	$2G + 2$	Diffusion coefficient
P1_MUBAR	$2G + 2$	Average scattering angle

NOTES:

1. The first two entries are the total (one-group) value and the associated relative statistical error. The remaining $2G$ entries are few-group values.
2. The values are based on the P_1 approximation. The results usually differ from values calculated using the analog estimate of diffusion area (see above).

6.1.30 B_1 fundamental mode calculation

Parameter	Values	Description
B1_KINF	1	Iterated multiplication factor
B1_BUCKLING	1	Iterated buckling
B1_FLUX	$2G + 2$	B_1 integral flux
B1_TOTXSXS	$2G + 2$	B_1 total cross section
B1_NSF	$2G + 2$	B_1 fission neutron production cross section
B1_FISSXS	$2G + 2$	B_1 fission cross section
B1_CHI	$2G$	B_1 fission spectrum
B1_ABSXS	$2G + 2$	B_1 absorption cross section
B1_RABSXS	$2G + 2$	B_1 reduced absorption cross section
B1_REMXS	$2G + 2$	B_1 removal cross section
B1_DIFFCOEF	$2G + 2$	B_1 diffusion coefficient
B1_SCATTXS	$4G^2$	B_1 scattering matrix
B1_SCATTPRODXS	$4G^2$	B_1 scattering production matrix
B1_I135PRODXS	$2G + 2$	Production cross section for I-135
B1_XE135PRODXS	$2G + 2$	Production cross section for Xe-135
B1_PM149PRODXS	$2G + 2$	Production cross section for Pm-149
B1_SM149PRODXS	$2G + 2$	Production cross section for Sm-149
B1_XE135ABSXS	$2G + 2$	Absorption cross section of Xe-135
B1_SM149ABSXS	$2G + 2$	Absorption cross section of Sm-149

NOTES:

1. B_1 fundamental mode calculation is performed after the transport cycle using homogenized multi-group cross sections (see Sec. 5.16).
2. The definition of scattering matrix was changed in version 1.1.15 (see Sec. 5.9).
3. Reduced absorption cross section is defined as absorption minus production in (n,xn) reactions.
4. Scattering production matrix includes neutron multiplication in (n,xn) reactions.
5. The option to switch on the calculation of fission product poison cross sections is described in Sec. 5.18
6. The capability is available from version 1.1.14 on. Some parameters were added in versions 1.1.15 and 1.1.17.

6.1.31 Assembly discontinuity factors

Parameter	Values	Description
ADFS	$2GN_V$	Surface discontinuity factors
ADFC	$2GN_V$	Corner discontinuity factors

NOTES:

1. The assembly discontinuity factors are calculated only for square and hexagonal cylinder boundaries. The ADF surface is the outermost surface in the universe where the group constants are calculated.
2. The number of vertices N_V is 4 for square boundary and 6 for hexagonal boundary.
3. The index for vertice (corner) n and group g is given by:

$$i = 2(n - 1)G + 2g - 1$$

4. The methodology is tested only group constant generation is extended over the entire geometry.
5. For square assemblies the numbering of vertices is: 1 - West, 2 - North, 3 - East, 4 - South and for the corners: 1 - South-West, 2 - South-East, 3 - North-East, 4 - North-West. Also note that geometry plots are inverted in the north-south direction.

6.1.32 Power distributions in lattices

Parameter	Values	Description
LAT<nl>	3	Lattice type and size
POWDISTR<nl>	$2N_L$	Power distribution in lattice
FG_POWDISTR<nl>	$2N_L(2G + 1)$	Power distribution in lattice divided into energy groups
PEAKF<nl>	4	Peaking factor in lattice

NOTES:

1. Lattice parameters are calculated for each lattice, regardless of the content. Variable names include the lattice number “<nl>”.
2. For square and hexagonal lattices the type and number of rows and columns is given. For cluster-type lattices the entries are type, number of rings and total number of elements.
3. The values in the power distribution are given as a single vector. The order is determined by the universe map in the lattice definition.

4. Peaking factor gives the position and the peak value in the lattice.
5. Energy-group wise power distribution is calculated from version 1.1.17 on.

6.2 History output

Detectors

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Table 7.1: Detector parameters.

Param.	Description	Comments
dr	Reaction multiplier	Determines the response function
dv	Detector volume	Used for normalization
dc	Detector cell	Defines the cell where the reactions are scored
du	Detector universe	Defines the universe where the reactions are scored
dm	Detector material	Defines the material where the reactions are scored
dl	Detector lattice	Defines the lattice where the reactions are scored
de	Detector energy grid	Defines the energy bins for the response function
dx	Detector mesh	Defines the x-mesh where the reactions are scored
dy	Detector mesh	Defines the y-mesh where the reactions are scored
dz	Detector mesh	Defines the z-mesh where the reactions are scored
dt	Detector type	Special detector types
ds	Surface current detector	Defines surface for current detector

IMPORTANT NOTES ON THE COLLISION FLUX ESTIMATOR:

1. The Serpent code uses the collision estimate of neutron flux, simply because the track-length estimate is not available when delta-tracking is used for neutron transport. The two estimates are equally well-suited for typical reactor lattice calculations, in which the neutron source is distributed over the entire geometry. The efficiency of the collision estimator becomes poor, however, if reaction rates are calculated inside small or optically thin volumes located in regions of low collision density. This is why the code is not the best choice for dosimetry calculations (see Ref. [20]). On the other hand, the use of the collision estimate requires less computational effort, especially for mesh detectors, which is directly reflected in the overall calculation time.

7.1.1 Setting the Response Function

The detector response function determines the type of the calculation. In the simplest case, $f = 1$, and (7.1) is reduced to the neutron flux integrated over space and energy. If a reaction cross section is used, the result is the corresponding reaction rate. It should be noted that the absolute value of the integral depends on source normalization (see Sec. 5.8).

The detector response function is defined by the “dr” entry:

```
det <name> dr <mt> <mat>
```

where <name> is the detector name
 <mt> is the response function number
 <mat> is the material name (or “void” for void material)

Table 7.2: Detector response functions. For a complete list of ENDF reaction MT's, see Ref. [6].

	MT	Reaction mode
Material total reactions	0	None
	-1	Total
	-2	Total capture
	-3	Total elastic
	-5	Total (n,2n)
	-6	Total fission
	-7	Total fission neutron production
	-8	Total fission energy deposition
	-9	Majorant
ENDF Reaction modes	1	Total
	2	Elastic scattering
	16	(n,2n)
	17	(n,3n)
	18	Total fission
	19	First-chance fission
	20	Second-chance fission
	51	Inelastic scattering to 1st excited state
	52	Inelastic scattering to 2nd excited state
	...	
	90	Inelastic scattering to 40th excited state
	91	Continuum inelastic scattering
	102	(n, γ)
	103	(n,p)
	104	(n,d)
	105	(n,t)
	106	(n, ^3He)
	107	(n, α)

If multiple responses are defined for a detector, an equal number of bins are created for the results. The response functions are listed in Table 7.2. Negative entries define total reaction rates related to materials. The total cross section (mt = -1), for example, is calculated from:

$$R = \frac{1}{V} \int_V \int_{E_{i+1}}^{E_i} \sum_j \left[\Sigma_{\text{tot},j}(\mathbf{r}, E) \phi(\mathbf{r}, E) \right] d^3r dE, \quad (7.2)$$

where the summation is carried over all nuclides in the material. If the material entry is set to void, the material at each collision point is used in the calculation. This allows the integration of reaction rates in volumes extending over several material regions.

Positive response numbers are related to isotopic, rather than material total reaction rates,

and they correspond to the reaction MT's used in ENDF format data. The list in Table 7.2 is not complete and a more detailed description is found in Ref. [6]. The detector material for an isotopic response function must consist of a single nuclide.

Detector values can be multiplied or divided by other values by setting the detector type to 2 or 3, respectively. The type is then followed by the name of the multiplier or divider detector. The total number of values must be equal for both detectors or the divider / multiplier detector single-valued.

EXAMPLES:

```
% Total flux in material "fuel":

det 1 dm fuel

% Detector materials:

mat U235 1.0 92235.09c 1.0
mat U238 1.0 92238.09c 1.0

% Calculate microscopic fission and capture cross sections of
% U-235 and U-238 by dividing the reaction rate by total flux:

det 2 dm fuel dr 18 U235 dt 3 1
det 3 dm fuel dr 102 U235 dt 3 1
det 4 dm fuel dr 18 U238 dt 3 1
det 5 dm fuel dr 102 U238 dt 3 1
```

IMPORTANT NOTES ON DETECTOR RESPONSE FUNCTIONS:

1. If multiple response functions are defined for a detector, an equal number of bins are created for the results.
2. Dosimetry cross sections (type 2 or 'y') can be used with detectors and with detectors only.
3. The ENDF reaction MT numbers are universal and related to isotopic cross sections. These reactions may not be used with materials consisting of more than one nuclide. The result is multiplied by the material atomic density and microscopic reaction rates can be calculated by setting the density to unity.
4. Some high-energy reaction modes, such as (n,3n), are excluded from the transport simulation. These modes are not available in the detector calculation either. All reaction modes are included for dosimetry cross sections.
5. The negative MT numbers are specific to Serpent and not universally defined. The reaction rates are calculated by summing over all nuclides in the material. MCNP also

uses some code-specific negative reaction MT's, but the interpretations are slightly different.

6. The fission energy deposition function defined by $mt = -8$ yields the total energy absorbed in the system (in J). This is not equivalent with the fission Q-value (see source normalization in Sec. 5.8).
7. The mt 's 0, -9 and -10 are not material-specific and the entry must be set to void.
8. If the "dr" entry is omitted entirely, the result is the total flux integrated over space and energy.

SEE ALSO:

1. Dosimetry cross sections (Sec. 1.4.1 on page 11)
2. Source rate normalization (Sec. 5.8 on page 61)

7.1.2 Setting the Energy Domain

The energy boundaries $[E_{i+1} E_i]$ of the integration (7.1) are set by a user-defined energy grid, linked to the detector by the "dt" entry:

```
det <name> de <ene>
```

where <name> is the detector name
<ene> is the grid name

The same energy grid definition is also used with B_1 fundamental mode calculation (See Sec. 5.16).

The number of energy bins is defined by the grid size. There are four types of energy grids

1. arbitrarily defined
2. equal energy-width bins
3. equal lethargy-width bins
4. predefined energy group structure

The grid definition has three entry formats:

```

ene <name> 1 <E1> <E2> ... <En>
ene <name> <type> <N> <Emin> <Emax>
ene <name> 4 <struct>

```

where <name>	is the grid name
<type>	is the grid type
<E1> <E2> ... <En>	are the bin boundaries in type 1 grid
<N>	is the number of bins in type 2 and 3 grids
<Emin>	is the minimum energy in type 2 and 3 grids
<Emax>	is the maximum energy in type 2 and 3 grids
<struct>	is the name of a predefined structure

The predefined energy grid names and descriptions are listed in Table 7.3. Bin boundaries are not listed here, but the values are easily readable in Serpent source file “egroups.c”.

The detector energy grid is often used for calculating spectral quantities. There are three special detector types for spectral calculations, determined by the “dt” detector type entry:

1. Cumulative spectrum (“dt -1”)
2. Division by energy width (“dt -2”)
3. Division by lethargy width (“dt -3”)

In the default mode, the bin values are independent and undivided.

EXAMPLES:

```

% Flux per lethargy using energy grid 1:

det 1 de 1 dt -3

% Differential capture, fission and production spectra:

det 2 de 1 dt -2 dr -2 void
det 3 de 1 dt -2 dr -6 void
det 4 de 1 dt -2 dr -7 void

% Integral capture, fission and production spectra:

det 5 de 1 dt -1 dr -2 void
det 6 de 1 dt -1 dr -6 void
det 7 de 1 dt -1 dr -7 void

```

Table 7.3: Predefined energy grid types.

Grid name	Description
nj2	csewg 239 group structure
nj3	lanl 30 group structure
nj4	anl 27 group structure
nj5	rrd 50 group structure
nj8	laser-thermos 35 group structure
nj9	epri-cpm 69 group structure
nj11	lanl 70 group structure
nj14	eurlib 100-group structure
nj16	vitamin-e 174-group structure
nj17	vitamin-j 175-group structure
nj18	xmas 172-group structure
nj19	ecco 33-group structure
nj20	ecco 1968-group structure
nj21	tripoli 315-group structure
nj22	xmas lwpc 172-group structure
nj23	vit-j lwpc 175-group structure
wms69	WIMS 69-group structure (equivalent with nj9)
wms172	WIMS 172-group structure
cas70	CASMO 70-group structure
cas40	CASMO 40-group structure
cas25	CASMO 25-group structure
cas23	CASMO 23-group structure
cas18	CASMO 18-group structure
cas16	CASMO 16-group structure
cas14	CASMO 14-group structure
cas12	CASMO 12-group structure
cas9	CASMO 9-group structure
cas8	CASMO 8-group structure
cas7	CASMO 7-group structure
cas4	CASMO 4-group structure
cas3	CASMO 3-group structure
cas2	CASMO 2-group structure
mupo43	MUPO 43-group structure
scale44	SCALE 44-group structure
scale238	SCALE 238-group structure

7.1.3 Setting the Spatial Domain

There are five options for setting the spatial domain of the integration:

1. By defining the cell where the reaction rates are scored using the “dc” parameter.
2. By defining the universe where the reaction rates are scored using the “du” parameter.
3. By defining the material where the reaction rates are scored using the “dm” parameter.
4. By defining the lattice where the reaction rates are scored using the “dl” parameter.
5. By setting up a one-, two- or three-dimensional mesh using the “dx”, “dy” and “dz” parameters.

All these options can be used without restrictions in various combinations. It should be noted, however, that some combinations may result in physically impossible configurations and produce zero results.

Detector cells, materials and universes

Detector cell, material and universe parameters all work on the same principle: the collision is scored if it occurs inside the cell, material or universe, respectively. A separate bin is created for each entry and the combination of different types creates a combination of bins. The syntax is:

```
det <name> dc <cell> dm <mat> du <univ>
```

where <name> is the detector name
 <cell> is the detector cell
 <mat> is the detector material
 <univ> is the detector universe

Detector cells can be either physical or super-imposed on the geometry. Super-imposed cells are not used for defining material regions. They must contain void material and the universe number must be set to a negative value. Universes containing super-imposed cells can be created for defining complicated geometry regions. These universes are not bound by the restrictions of physical universes discussed in Section 3.6. Leakage rate can be calculated by scoring collisions in outside cells.

Fuel pin definitions are geometry macros that are converted into ordinary geometry objects constructed using cells and surfaces. The cells in fuel pins are named using convention:

```
nst<np>c<nr>
```

where <np> is the pin (universe) number
 <nr> is the ring index starting from the innermost region (= 1)

Burnable materials in fuel pins are renamed and divided into a user-defined number of annular depletion zones (see Sec. 8.2 on page 109). The naming convention is:

```
<mat>p<np>r<nr>
```

where <mat> is the original material name
 <np> is the pin (universe) number
 <nr> is the ring index starting from the innermost region (= 1)

EXAMPLES:

```
% Simple cell, material and universe detectors:
```

```
det 1 dc 1          % Score collisions in cell 1
det 2 dm fuel       % Score collisions in material "fuel"
det 3 du 2          % Score collisions in universe 2
```

```
% Combined detectors:
```

```
det 4 dc 1 dc 2     % Two bins: collisions in cells 1 and 2
det 5 du 1 dm H2O   % Collisions in material "H2O" in universe 2
```

```
% Super-imposed cells:
```

```
cell 10 -1 void -1
cell 11 -1 void 1 -2
```

```
det 6 dc 10         % Collisions in super-imposed cell 10
det 7 du -1         % Collisions in super-imposed universe -1
```

Lattice detectors

The input format for the lattice detector is:

```
det <name> dl <lat>
```

where <name> is the detector name
 <lat> is the detector lattice number

A bin is created for each lattice position. The results can be combined with cell, material and universe bins. For example, the flux distribution in material “clad” in a fuel pin lattice “10” can be calculated using:

```
det 1
dm clad           % Score in material "clad"
dl 10            % Lattice bins in lat 10
```

Mesh detectors

The mesh detector creates a super-imposed uniform square mesh over the geometry. The mesh structure is given separately in x-, y- and z-directions and the input format for the x-type is:

```
det <name> dx <xmin> <xmax> <nx>
```

where <name> is the detector name
 <xmin> is the minimum x-coordinate of the mesh
 <xmax> is the maximum x-coordinate of the mesh
 <nx> is the number of mesh bins in the x-direction

EXAMPLES:

```
% One-dimensional mesh (axial power distribution in fuel pin):
```

```
det 1
du 1 % Score in universe (pin) 1
dm fuel % Score in material "fuel"
dz 0.0 120.0 50 % 50 axial bins between z = 0 and z = 120 cm
```

```
% Two-dimensional mesh (total fission rate distribution):
```

```
det 2
dr -6 void % Multiply by total fission rate
dx -225.0 225.0 30 % 30 bins in x-direction
dy -225.0 225.0 30 % 30 bins in y-direction
```

```
% Three-dimensional mesh (thermal flux distribution):
```

```
ene 1 1 1E-11 0.625E-6 % Detector energy grid (single bin)
```

```
det 3
de 1 % Use energy grid 1
dx -225.0 225.0 30 % 30 bins in x-direction
dy -225.0 225.0 30 % 30 bins in y-direction
dz 0.0 400.0 10 % 10 bins in z-direction
```

7.1.4 Surface Current Detectors

Serpent 1.1.17 and later versions have the capability to calculate neutron current through surfaces. The syntax for the current detector is:


```
det <name> ds <surf> <dir>
```

where <name> is the detector name
 <surf> is the surface name
 <dir> is the direction vector (-1 = inward, 0 = net, 1 = outward)

The surface associated with the detector is assumed to be located relative to the origin of universe zero, and it may or may not be a part of the geometry definition. The direction vector determines which surface crossings are included in the result. Inward current has positive and outward current negative value, respectively. Net current is calculated as the sum of the two.

The surface current detector was added mainly for the calculation of reflector group constants, and the first implementation had some limitations with respect to boundary conditions (see note below). Reflector geometries typically involve the use of partial boundary conditions (see Sec. 5.7 on page 59), available from code version 1.1.17 on.

IMPORTANT NOTES ON THE SURFACE CURRENT DETECTOR:

1. The surface current detector in version 1.1.17 cannot cope with some of the coordinate transformations performed when repeated boundary conditions are applied, which limits its use to geometries with black boundary conditions, or reflected or periodic boundary condition perpendicular to the detector surface. This limitation was lifted in update 1.1.18, and the most recent implementation should work in all geometry types.

7.2 Detector output

The output from all detectors is printed in matlab m-file format in a single file named “<input>_det<n>.m”, where “<input>” is the name of the input file and “<n>” is the burnup step.

The results for each detector are written in a 13-column table, one bin value per row. The variable is named “DET<name>.m”, where “<name>” is the detector name. The values in each column are:

1. Value index (total number in “DET<name>_VALS”)
2. Energy bin index (total number in “DET<name>_EBINS”)
3. Universe bin index (total number in “DET<name>_UBINS”)
4. Cell bin index (total number in “DET<name>_CBINS”)
5. Material bin index (total number in “DET<name>_MBINS”)

6. Lattice bin index (total number in “DET<name>_LBINS”)
7. Reaction bin index (total number in “DET<name>_RBINS”)
8. Z-mesh bin index (total number in “DET<name>_ZBINS”)
9. Y-mesh bin index (total number in “DET<name>_YBINS”)
10. X-mesh bin index (total number in “DET<name>_XBINS”)
11. Mean value
12. Relative statistical error
13. Total number of scores

Detector volume is given in variable “DET<name>_VOL”. All results have been divided by this number.

If an energy bin structure is defined, the corresponding bin boundaries are written in variable “DET<name>E”. The variable has three columns:

1. Lower energy boundary of bin
2. Upper energy boundary of bin
3. Mean energy of bin

The number of rows is equal to the number of energy bins.

If x-, y- or z-bins are defined, the corresponding bin boundaries are written in variables “DET<name>X”, “DET<name>Y”, “DET<name>Z”, respectively. The variables have three columns:

1. Coordinate of the lower bin boundary
2. Coordinate of the upper bin boundary
3. Coordinate of bin center

The number of rows is equal to the number of x-, y- or z-bins.

IMPORTANT NOTES ON DETECTOR OUTPUT:

1. Some variables are missing and the names are in lower-case in the pre-release version 1.0.0 of the Serpent code (corrected in version 1.0.1).
2. Detector volume is printed in version 1.1.13 on.

7.3 Detectors in Burnup Calculation

There are a few things that need to be considered when using detectors in the burnup calculation mode. First, the output is printed in a different file for each burnup step (see previous section). The file names are separated by the step index, which is set to zero for the initial composition. Second, when burning materials inside pin and particle structures (see Sec.3.4 and 3.8), the materials are renamed according to pin / particle index and region number if the material is divided into multiple depletion zones (see Sec. 8.2). The original material names no longer exist and the new names must be used instead with the “dm” parameter.

Chapter 8

Burnup calculation

8.1 General

Serpent can be run both as a stand-alone burnup calculation code and as a part of a coupled system. In the first case, the code uses an internal calculation routine for solving the set of Bateman equations describing the changes in the material compositions caused by neutron-induced reactions and radioactive decay. In the second case, the code is used as the neutronics solver in an externally coupled system.

The additional input for burnup calculation consists of identifying the depleted materials (Sec. 8.2) and setting up the irradiation history (Sec. 8.3). There are also some additional parameters for determining file paths and options used by the calculation routines (Sec. 8.4). A few simple examples are given in Sec. 8.7 and complete input listings in Sec. 11.2.

It should be noted that burnup calculations are more sensitive to small changes in the geometry, materials and calculation parameters compared to a steady state simulation. The length of burnup steps and predictor-corrector calculation (see Sec. 8.3 and Sec. 8.4) may have a significant impact on the accumulation of certain isotopes, and especially the depletion of burnable absorbers. In thermal systems, the build-up rate of plutonium is strongly dependent on moderator conditions, such as density and the $S(\alpha, \beta)$ scattering laws (see Sec. 4.2 on Page 49). As low as a 30K difference in moderator temperature may result in over 1% discrepancy in Pu-239 concentration at high burnup.¹ Differences originating from the evaluated nuclear data should always be taken into account, especially for older libraries, such as JEF-2.2 and ENDF/B-VI.

¹It should be noted that the thermal scattering data provided with the installation package is generated at slightly different temperatures for different libraries.

8.2 Depleted materials

Depleted materials are identified by an additional “burn” entry in the material card:

```
mat <name> <dens> burn <nr>
<iso 1> <frac 1>
<iso 2> <frac 2>
...
```

where	<name>	is the material name
	<dens>	is the density (mass or atomic)
	<nr>	is the number of annular regions in depleted fuel pins
	<iso 1> <iso 2> ...	are the names of the constituent nuclides
	<frac 1> <frac 2> ...	are the corresponding fractions (mass or atomic)

If the irradiation history is not set up, the “burn” entry activates the coupled calculation mode and one-group transmutation cross sections, radioactive decay constants and fission yields are written in a separate output file (see Sec. 8.6) without running the depletion calculation.

The code treats depleted materials in fuel pins different from materials in ordinary cells. Each pin type is treated separately and further divided into <nr> annular depletion zones of equal volume. The division is important for accounting for the rim-effects caused by spatial self-shielding. The code automatically renames the depleted pin materials using convention:

```
<mat>p<np>r<nr>
```

where	<mat>	is the original material name
	<np>	is the pin (universe) number
	<nr>	is the ring index starting from the innermost region (= 1)

Depleted materials in ordinary cells are not renamed or divided into sub-regions.

IMPORTANT NOTES ON DEPLETED MATERIALS:

1. Each fuel pin type containing a depleted material is treated separately and divided into a user-given number of annular depletion zones.
2. The separation of material regions is based on pin type, not lattice position. If similar pins in different positions need to be treated as different materials, a new (identical) pin type must be assigned for each position (See examples in Sec. 8.7.1 on page 117).
3. Fuel pins containing burnable absorber should always be divided into ~10 rings in order to account for the rim-effects caused by spatial self-shielding.
4. The current code version can only handle burnup calculation in cylindrical or spherical material regions, such as fuel pins or HTGR micro particles.

SEE ALSO:

1. Material cards (Sec. 4.1.2 on page 48)

8.3 Irradiation history

The irradiation history in the independent burnup calculation mode consists of one or several burnup intervals, defined by the “dep” card:

```
dep <stype>
  <step 1>
  <step 2>
  ...
```

where <stype> is the step type
 <step 1> <step 2> ... are the burnup steps

The step types are listed in Table 8.1

Table 8.1: Burnup step types.

<stype>	Step values
bustep	depletion step, burnup intervals given in MWd/kgU
butot	depletion step, cumulative burnup given in MWd/kgU
daystep	depletion step, time intervals given in days
daytot	depletion step, cumulative time given days
decstep	decay step, time intervals given in days
dectot	decay step, cumulative time given in days

Source rate normalization and soluble absorber concentration can be changed between intervals by re-defining the values. The first value is used during the first burnup interval, the second during the second interval and so on. Examples are given in Sec. 8.7.2 on page 121.

The last two options omit the transport cycle and handle only radioactive decay, which makes the calculation run significantly faster. This mode is intended to be used for calculating activities and inventories after the irradiation is completed. Downtime between cycles is better handled by setting the power to zero.

IMPORTANT NOTES ON IRRADIATION HISTORY:

1. If source rate normalization or soluble absorber concentration are changed between burnup intervals, it is important that the number of definitions is equal to the number of intervals.

2. The structure of the “dep” card is different in the early code versions (before 1.0.2).
3. The soluble absorber definition is available from version 1.0.2 on.
4. The decay mode is available from code version 1.1.10 on.

SEE ALSO:

1. Source rate normalization options (Sec. 5.8 on page 61)
2. Soluble absorber (Sec. 5.14 on page 69)

8.4 Options for Burnup Calculation

The calculation parameters in the burnup mode are summarized in Table 8.2.

Table 8.2: List of parameters and options in burnup calculation mode.

Option		Description	Section	Page
declib	(1)	file path for radioactive decay data	8.4.1	112
nfylib	(1)	file path for fission yield data	8.4.1	112
sfylib	(1)	file path for spontaneous fission yield data	8.4.1	112
bunorm	(1)	normalization mode in burnup calculation	8.4.2	112
fmass	(1)	total fissile mass	8.4.2	112
bumode	(1)	solution method for Bateman equations	8.4.3	113
pcc	(1)	flag for predictor-corrector calculation	8.4.3	113
xscal	(1)	transmutation cross sections generation	8.4.4	113
fpcut	(1)	fission product yield cut-off	8.4.5	114
axs	(2)	actinide mass chains included in calculation	8.4.5	114
stabcut	(1)	stability cut-off	8.4.5	114
ttacut	(1)	TTA chain cut-off	8.4.5	114
xsfcut	(1)	XS fraction cut-off	8.4.5	114
xsecut	(1)	XS threshold energy cut-off	8.4.5	114
inventory	(1-N)	nuclide list for burnup calculation output	8.4.6	115
printm	(1)	flag for printing material compositions	8.4.7	115
dhprec	(1)	precursor-group wise decay heat production	8.4.8	115

8.4.1 Library File Paths

In addition to the continuous-energy cross section libraries, burnup calculation requires radioactive decay data and neutron-induced and spontaneous fission product yields. These files are read in the raw ENDF format. The decay data library file path is set using:

```
set declib "<file>"
```

where `<file>` is the file path for the ENDF format decay data library
the neutron-induced fission yield library using:

```
set nfylib "<file>"
```

where `<file>` is the file path for the ENDF format fission yield library
and the spontaneous fission yield library:

```
set sfylib "<file>"
```

where `<file>` is the file path for the ENDF format fission yield library

The spontaneous fission yield library is optional. If the file path is not set, the code uses neutron-induced yields for spontaneous fission. The present code version does not model spontaneous fission.

A default directory path can be set by defining environment variable `SERPENT_DATA`. The code looks for data files in this path if not found at the absolute location.

8.4.2 Normalization

The normalization of fission source is described in Sec. 5.8 on page 61. In some burnup calculation problems, the geometry may contain fissile materials that are not depleted, which may also affect the source normalization. Serpent offers three options, set using:

```
set bunorm <mode>
```

where `<mode>` is the normalization mode

Mode 1 is the default treatment which normalizes the given reaction rate or power to all materials. Mode 2 includes only burnable materials and mode 3 only non-burnable materials. The option is available from update 1.1.5 on and earlier code versions use all materials in the normalization.

The code automatically calculates the total fissile mass in the system, which is needed for normalizing the reaction rates. If the calculation fails, the value can be set manually using:

```
set fmass <m>
```

where `<m>` is the total fissile mass in the system (in grams)

8.4.3 Solution of Depletion Equations

The Serpent code has three options and two methods for solving the Bateman equations describing the changes in the isotopic compositions caused by neutron-induced reactions and radioactive decay. The calculation mode is set using:

```
set bumode <mode>
```

where <mode> is the method used for depletion calculation

The first method (<mode> = 1) is Transmutation Trajectory Analysis (TTA), based on the analytical solution of linearized transmutation chains. The second method (<mode> = 2), used by default, is an advanced matrix exponential solution based on the Chebyshev Rational Approximation Method (CRAM). The third option (<mode> = 3) is the variation TTA method, in which cyclic transmutation chains are handled by inducing small variations in the coefficients instead of solving the extended TTA equations.

Predictor-corrector calculation is activated using:

```
set pcc <corr>
```

where <corr> is the flag for running the corrector step (0 = no, 1 = yes)

The method is used by default and results in a more accurate estimation of isotopic changes during each burnup step. The drawback is that the transport cycle is repeated, which increases the overall calculation time.

8.4.4 Calculation of Transmutation Cross Sections

There are two options for calculating the isotopic one-group transmutation cross sections:

```
set xscal <mode>
```

where <mode> is the method used for cross section calculation

In the default method (<mode> = 2), the code calculates these parameters using a high-resolution flux spectrum recorded during the transport calculation. This procedure results in a reduction of calculation time by a factor of 3-4 compared to the direct calculation of the cross sections during the transport cycle (<mode> = 1). The drawback is that the method is an approximation and that the information on statistical accuracy is lost.²

²The flux spectrum is calculated using the main energy-grid structure. The resolution is high and the only approximation is that the continuous-energy cross sections are assumed constant between two grid points. It is therefore assumed that the difference to the direct calculation are negligible, although the methodology still requires some thorough validation. The two methods are automatically compared by setting <mode> = 3.

8.4.5 Cut-offs

Burnup calculation uses various cut-offs for reducing the computational effort.

Fission product yield cut-off determines which fission products are included in the calculation. The selection is based on the cumulative yield of each fp mass chain:

```
set fpcut <lim>
```

where <lim> is the limit for fission product yield cut-off

By default, the range of actinide mass chains included in the calculation extends from $A_{\min} - 1$ to $A_{\max} + 7$, where A_{\min} and A_{\max} are the minimum and maximum actinide mass numbers in the initial composition. This range can be set manually by:

```
set axs <Amin> <Amax>
```

where <Amin> is the lightest actinide mass chain included in the calculation

<Amax> is the heaviest actinide mass chain included in the calculation

Stability cut-off:

```
set stabcut <lim>
```

where <lim> is the limit for stability cut-off

TTA chain cut-off:

```
set ttacut <lim>
```

where <lim> is the limit for TTA chain cut-off

Cross section fraction cut-off:

```
set xsfcut <lim>
```

where <lim> is the limit for cross section fraction chain cut-off

Threshold energy cut-off:

```
set xsecut <lim>
```

where <lim> is the energy boundary

8.4.6 Nuclide Inventory

The standard output in the independent calculation mode consists of material compositions, transmutation cross sections, activities and decay heating values. The isotopes, elements,

etc. included in the output are set by the inventory option:

```
set inventory <id1> <id2> ...
```

where <idn> are the identifiers.

The list consists of numerical values that identify the nuclides ($1000 \cdot Z + 10 \cdot A + I$) or elements (Z). Isotope and elemental names and symbols (“Pu-239”, “Gd155”, “PM148M”, “Cs”, “plutonium”, etc.) are also accepted. Elemental values are calculated by summing over the isotopes. Table 8.3 lists additional options that can be used in the inventory list to sum over several elements.

Table 8.3: Special entries in the inventory list. The list entry may consist of name or ID.

ID	Name	Description
201	act	Actinides ($Z > 89$)
202	fp	Fission products
204	dp	Decay products below thorium in the natural actinide decay series
208	ng	Noble gases (in the fission product range, helium and radon excluded)

8.4.7 Additional Output

The code has an option for writing the compositions of depleted materials in a separate output file after each step:

```
set printm <mode>
```

where <mode> is the flag for printing material compositions (0 = no, 1 = yes)

The code produces for each step a file named “<input>.bumat<n>”, where <input> is the name of the input file and <n> is the burnup step. The material compositions can be used in another Serpent calculation or converted to MCNP format for validation purposes.

8.4.8 Decay heat production in multiple precursor groups

Decay heat production can be divided into multiple precursor groups based on the nuclide decay constant. The syntax for the option is:

```
set dhprec [ <l0> <l1> ... ]
```

where <ln> are the group boundaries in ascending order

Default values are used if the option is not given. The output is printed in the main output file (see Sec. 6.1.11). The feature is available from code version 1.1.17 on.

IMPORTANT NOTES ON BURNUP CALCULATION PARAMETERS:

1. Decay and fission yield libraries are raw ENDF data files in ASCII format.
2. Symbolical names can be used in the inventory list from version 1.1.3 on. Elemental and special identifiers are available from version 1.1.10 on. If the list is empty, only material total values are printed.
3. The code looks for the daughter nuclide cross section data libraries in the ACE directory file. It is important that the directory file contains as many nuclides as possible.
4. Mode 2 (matrix exponential solution) is available and used by default from version 1.1.0 on.
5. It is important to use the predictor-corrector step in cases involving burnable absorbers.
6. The environment variable feature is available from code version 1.1.8 on.

SEE ALSO:

1. Setting up the cross section library file path (Sec. 5.4 on page 57).
2. Description of the CRAM method in Ref. [21].

8.5 Output in independent mode

The burnup calculation output in the independent calculation mode is written in Matlab m-file format in file “<input>_dep.m”, where <input> is the name of the input file. The variables are summarized in Table 8.4. The number of burnup steps is N and the number of inventory nuclides I . The material-wise parameters are printed for each depleted material.

IMPORTANT NOTES ON OUTPUT:

1. If the predictor-corrector method is used, the material compositions are given at the beginning of each step. The transmutation cross sections are not equivalent with the corrected values used for solving the depletion equations.
2. The variable names are slightly different in the pre-release version 1.0.0 of the Serpent code (corrected in version 1.0.1).
3. The “lost” in the output file refers to data that is lost to undefined nuclides.

SEE ALSO:

1. Setting up burnup inventory list (Sec. 8.4.6 on page 115).

Table 8.4: Variables in the Matlab m-format burnup calculation output file.

Variable	Size	Contents
BU	(1, N)	Cumulative burnup in MWd/kgU
DAYS	(1, N)	Cumulative burn time in days
i<ZAI>	1	Table index for nuclide "<ZAI>"
iTOT	1	Table index for total values
iLOST	1	Table index for lost data
ZAI	($I + 2$, 1)	Nuclide ZAI's
NAMES	($I + 2$, 8)	Nuclide names (character strings)
MAT_<mname>_VOLUME	(1, N)	Volume of material "<mname>"
MAT_<mname>_FLUX	(1, N)	Volume-integrated flux in material "<mname>"
MAT_<mname>_ADENS	($I + 2$, N)	Atomic densities in material "<mname>"
MAT_<mname>_MDENS	($I + 2$, N)	Mass densities in material "<mname>"
MAT_<mname>_A	($I + 2$, N)	Activities in material "<mname>"
MAT_<mname>_H	($I + 2$, N)	Decay heat in material "<mname>"
MAT_<mname>_FISSXS	($I + 2$, N)	(n,f) cross sections in material "<mname>"
MAT_<mname>_CAPTXS	($I + 2$, N)	(n, γ) cross sections in material "<mname>"
MAT_<mname>_N2NXS	($I + 2$, N)	(n,2n) cross sections in material "<mname>"
TOT_VOLUME	1	Total volume of depleted materials
TOT_ADENS	($I + 2$, N)	Total averaged atomic densities
TOT_MASS	($I + 2$, N)	Total mass
TOT_A	($I + 2$, N)	Total activities
TOT_H	($I + 2$, N)	Total decay heat

8.6 Output in coupled mode

8.7 Burnup calculation examples

8.7.1 Material and lattice examples

A simple assembly burnup calculation consisting of two pin types:

```
% --- Fuel pin:

pin 1
UO2      0.4025
clad     0.4750
water

% --- Gd-pin:
```

```

pin 3
UO2Gd    0.4025
clad      0.4750
water

% --- Guide tube:

pin 4
water     0.5730
tube      0.6130
water

% --- Pin lattice:

lat 110   1   0.0 0.0 17 17   1.265

1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 3 4 1 1 4 1 1 4 3 1 1 1
1 1 1 4 1 1 1 1 3 1 1 1 1 4 1 1
1 1 3 1 1 1 1 1 1 1 1 1 1 3 1 1
1 1 4 1 1 4 1 1 4 1 1 4 1 1 4 1
1 1 1 1 1 1 3 1 1 1 3 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 4 3 1 4 1 1 4 1 1 4 1 3 4 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 3 1 1 1 3 1 1 1 1 1
1 1 4 1 1 4 1 1 4 1 1 4 1 1 4 1
1 1 3 1 1 1 1 1 1 1 1 1 1 3 1 1
1 1 1 4 1 1 1 1 3 1 1 1 1 4 1 1
1 1 1 1 3 4 1 1 4 1 1 4 3 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

% --- Fuel in normal pins, no division into rings:

mat UO2      6.7402E-02  burn 1
92234.09c    9.1361E-06
92235.09c    9.3472E-04
92238.09c    2.1523E-02
8016.09c     4.4935E-02

% --- Fuel in Gd pins, division into 10 rings:

mat UO2Gd    6.8366E-02  burn 10
92234.09c    4.2940E-06
92235.09c    5.6226E-04

```

```
92238.09c  2.0549E-02
64154.09c  4.6173E-05
64155.09c  2.9711E-04
64156.09c  4.1355E-04
64157.09c  3.1518E-04
64158.09c  4.9786E-04
64160.09c  4.3764E-04
8016.09c   4.5243E-02
```

Similar case, but each lattice position treated as a separate depletion zone, taking into account the 1/12 symmetry of the pin layout:

```
% --- Fuel pins:
```

```
pin 10
UO2      0.4025
clad     0.4750
water
```

```
pin 11
UO2      0.4025
clad     0.4750
water
```

```
...
```

```
(identical definition of pins 12-45 omitted for simplicity)
```

```
...
```

```
% --- Gd-pins:
```

```
pin 50
UO2Gd    0.4025
clad     0.4750
water
```

```
pin 51
UO2Gd    0.4025
clad     0.4750
water
```

```
pin 52
UO2Gd    0.4025
clad     0.4750
water
```

```
% --- Guide tube:
```

```
pin 90
water    0.5730
tube     0.6130
water
```

```
% --- Pin lattice:
```

```
lat 110  1  0.0 0.0 17 17  1.265
```

```
45 44 43 42 41 40 39 38 37 38 39 40 41 42 43 44 45
44 36 35 34 33 32 31 30 29 30 31 32 33 34 35 36 44
43 35 28 27 52 90 26 25 90 25 26 90 52 27 28 35 43
42 34 27 90 24 23 22 21 51 21 22 23 24 90 27 34 42
41 33 52 24 20 19 18 17 16 17 18 19 20 24 52 33 41
40 32 90 23 19 90 15 14 90 14 15 90 19 23 90 32 40
39 31 26 22 18 15 50 13 12 13 50 15 18 22 26 31 39
38 30 25 21 17 14 13 11 10 11 13 14 17 21 25 30 38
37 29 90 51 16 90 12 10 90 10 12 90 16 51 90 29 37
38 30 25 21 17 14 13 11 10 11 13 14 17 21 25 30 38
39 31 26 22 18 15 50 13 12 13 50 15 18 22 26 31 39
40 32 90 23 19 90 15 14 90 14 15 90 19 23 90 32 40
41 33 52 24 20 19 18 17 16 17 18 19 20 24 52 33 41
42 34 27 90 24 23 22 21 51 21 22 23 24 90 27 34 42
43 35 28 27 52 90 26 25 90 25 26 90 52 27 28 35 43
44 36 35 34 33 32 31 30 29 30 31 32 33 34 35 36 44
45 44 43 42 41 40 39 38 37 38 39 40 41 42 43 44 45
```

```
% --- Fuel in normal pins, no division into rings:
```

```
mat UO2      6.7402E-02  burn 1
92234.09c    9.1361E-06
92235.09c    9.3472E-04
92238.09c    2.1523E-02
8016.09c     4.4935E-02
```

```
% --- Fuel in Gd pins, division into 10 rings:
```

```
mat UO2Gd    6.8366E-02  burn 10
92234.09c    4.2940E-06
92235.09c    5.6226E-04
92238.09c    2.0549E-02
64154.09c    4.6173E-05
64155.09c    2.9711E-04
64156.09c    4.1355E-04
64157.09c    3.1518E-04
64158.09c    4.9786E-04
```



```
64160.09c  4.3764E-04
8016.09c   4.5243E-02
```

8.7.2 Irradiation history examples

Irradiation at constant power density, cumulative burnup steps:

```
set powdens 40.0E-3
```

```
dep butot
```

```
0.10000
0.50000
1.00000
1.50000
2.00000
2.50000
3.00000
3.50000
4.00000
4.50000
5.00000
5.50000
6.00000
6.50000
7.00000
7.50000
8.00000
8.50000
9.00000
9.50000
10.00000
10.50000
11.00000
11.50000
12.00000
12.50000
13.00000
13.50000
14.00000
14.50000
15.00000
20.00000
25.00000
30.00000
35.00000
```

40.00000

Similar case with step size given and history divided into 3 irradiation intervals with cooling period. Nuclide inventory is traced for 1000 years after the fuel is removed from the reactor:

```
% --- Cycle 1: 650 ppm boron, final burnup 13.5 MWd/kgU
```

```
set powdens 40.0E-3
```

```
set abs boron -650E-6 water
```

```
dep bustep
```

```
0.10000
```

```
0.40000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
0.50000
```

```
% --- Downtime for 80 days:
```

```
set powdens 0.0
```

```
set abs boron -650E-6 water
```

```
dep daystep 80
```

```
% --- Cycle 2: 300 ppm boron, final burnup 25.0 MWd/kgU
```

```
set powdens 40.0E-3
set abs boron -300E-6 water
```

```
dep bustep
```

```
0.50000
0.50000
0.50000
5.00000
5.00000
```

```
% --- Downtime for 80 days:
```

```
set powdens 0.0
set abs boron -300E-6 water
```

```
dep daystep 80
```

```
% --- Cycle 3: no boron, final burnup 40.0 MWd/kgU
```

```
set powdens 40.0E-3
set abs boron 0.0 water
```

```
dep bustep
```

```
5.00000
5.00000
5.00000
```

```
% --- Decay after fuel is removed from the reactor
```

```
dep decstep
```

```
365    % 1. year
365    % 2. year
365    % 3. year
365    % 4. year
365    % 5. year
365    % 6. year
365    % 7. year
365    % 8. year
365    % 9. year
365    % 10. year
3650   % 20. year
3650   % 30. year
```

3650	% 40. year
3650	% 50. year
3650	% 60. year
3650	% 70. year
3650	% 80. year
3650	% 90. year
3650	% 100. year
36500	% 200. year
36500	% 300. year
36500	% 400. year
36500	% 500. year
36500	% 600. year
36500	% 700. year
36500	% 800. year
36500	% 900. year
36500	% 1000. year

Chapter 9

External Source Mode

9.1 General

External source simulation mode, available from version 1.1.11 on, can be used to replace the k -eigenvalue criticality source method in sub-critical and non-multiplying systems. Instead of performing power iterations on the fission source, all source neutrons are started from a user-defined distribution. The calculation mode is activated by replacing the “pop” input parameter (see Sec. 5.2 on page 53) with:

```
set nps <Nsrc> [ <Nbatch> ]
```

where <Nsrc> is the total number of source neutrons run
<Nbatch> is the number of batches run

By default, the simulation is run by dividing the source size into 200 batches. Apart from the source definition, described in the following section, the external source simulation works very similar to the criticality source method. All features, including detectors and burnup calculation are available.

IMPORTANT NOTES ON EXTERNAL SOURCE SIMULATION:

1. The calculation mode is available from version 1.1.11 on, and still very much under development.
2. External source simulations can only be run in non-multiplying or sub-critical systems. Geometries with $k_{\text{eff}} \geq 1$ produce infinite multiplication and the simulation diverges.

9.2 Source definition

The external source simulation requires one or several source definitions. A user-defined source can also be used as the initial guess for criticality source calculations (see Sec. 5.2). The syntax for the source definition is:

```
src <name> <param 1> <param 2> ...
```

where <name> is the source name
<param 1> <param 2> ... are the source parameter sets

The parameters are listed in Table 9.1 and they can be combined in different ways as described in the following subsections. If multiple sources are used, the relative importances are determined by the weights, set to unity by default.

Table 9.1: Detector parameters.

Param.	Description	Comments
sw	Source weight	Determines the relative importance of the source
sc	Source cell	Defines the cell where the neutrons are started
sm	Source material	Defines the material where the neutrons are started
sp	Source point	Defines the coordinates of a point source
sx, sy, sz	Source boundaries	Defines the boundaries of the source distribution
sd	Source direction	Defines the source direction vector
se	Source energy	Multiple uses
sb	Source energy bins	Defines a bin-wise energy spectrum
sr	Source reaction	Defines the source reaction
ss	Source surface	Defines a surface source

9.2.1 Setting the Spatial Distribution

If spatial distribution is not defined, neutrons are started uniformly all over the geometry. The sampling volume can be limited by setting the boundaries in x-, y- and z-directions using:

```
src <name> sx <x0> <x1> sy <y0> <y1> sz <z0> <z1>
```

where <name> is the source name
 <x0> is the minimum boundary in x-direction
 <x1> is the maximum boundary in x-direction
 <y0> is the minimum boundary in y-direction
 <y1> is the maximum boundary in y-direction
 <z0> is the minimum boundary in z-direction
 <z1> is the maximum boundary in z-direction

The source can be defined by a single cell using:

```
src <name> sc <cell>
```

where <name> is the source name
 <cell> is the cell where the neutrons are started

or to a single material using:

```
src <name> sm <mat>
```

where <name> is the source name
 <mat> is the material where the neutrons are started

The cell and material definitions can be used in combination with the boundaries set by “sx”, “sy” and “sz”.

An alternative to a volume source is the point source, defined as:

```
src <name> sp <x> <y> <z>
```

where <name> is the source name
 <x> is x-coordinate of the point source
 <y> is y-coordinate of the point source
 <z> is z-coordinate of the point source

Surface sources can be defined as:

```
src <name> ss <surf>
```

where <name> is the source name
 <surf> is the source surface

The surface is defined using the “surf” card (see Sec. 3.2 on page 19). Positive and negative entries refer to neutrons being emitted in the direction of positive and negative surface normal, respectively. The feature is available from version 1.1.15 on, and the allowed surface types include sphere (“sph”) and cylinder (“cyl”).

9.2.2 Setting the Directional Distribution

By default, all source neutrons in point and volume sources are emitted isotropically. To define a mono-directional source, the direction vector can be set by the “sd” parameter:

```
src <name> sd <u> <v> <w>
```

where <name> is the source name
 <u> is direction cosine in the x-direction
 <v> is direction cosine in the y-direction
 <w> is direction cosine in the z-direction

Directional distributions will be added in future code versions.

9.2.3 Setting the Energy Distribution

A mono-energetic source is defined by setting the “se” parameter:

```
src <name> se <E>
```

where <name> is the source name
 <E> is neutron energy

By default, the emission energy is set to 1 MeV.

Another option is to take the energy distribution from a nuclear reaction using the “sr” option:

```
src <name> sr <iso> <mt>
```

where <iso> is the nuclide identifier
 <mt> is the reaction mt

The reaction can be any scattering or fission reaction for which the distribution data exists in the ACE format data (notice that this is not the case for elastic scattering and inelastic level scattering). If source energy is defined using the “se” option, the value is used as the energy of the incoming neutron when the emission energy is sampled. If the value is not set, the minimum value allowed by the distribution is used.

The third option is to define discrete energy bins as:

```
src <name> sb <nb> <E0> <w0> <E1> <w1> ... <En> <wn>
```

where <nb> is the number of source energy bins
 <Ei> are the energy bin boundaries
 <wi> are the bin weights

The code samples the energy bin according to the probability calculated from the bin weights,

and the energy uniformly between the bin boundaries. The energy entries correspond to the upper boundaries of each bin, and the weight of the first bin must be set to zero. The feature is available from version 1.1.15 on.

9.2.4 Source files

Source distribution can be read from a file using:

```
src <name> sf <file> <type>
```

where <name> is the source name
<file> is the source file
<type> is the file type (must be 1)

The source file contains coordinates, direction cosines, energy, weight and time for every source neutron, one entry per line. This feature was added in version 1.1.17, and the format of the source file may change in later updates.

9.3 Source Examples

Source definition using default parameters – isotropic, mono-energetic 1 MeV source, uniformly distributed over the geometry:

```
src 1
```

Setting the spatial and directional distribution:

```
% Uniform source in a cuboid:
```

```
src 2 sx -1.0 1.0 sy -1.0 1.0 sz -1.0 1.0
```

```
% Source in cell:
```

```
src 3 sc 1
```

```
% Source in material, bounded in axial direction:
```

```
src 4 sm fuel sz -10.0 10.0
```

```
% Point source in origin, directed in the positive x-axis:
```

```
src 5 sp 0.0 0.0 0.0 sd 1.0 0.0 0.0
```

Setting the energy distribution:

```
% Three point sources with different energy and importance

src 6 sw 0.5 sp 0.0 0.0 0.0 se 1.0
src 7 sw 0.3 sp 1.0 0.0 0.0 se 2.0
src 8 sw 0.2 sp 0.0 1.0 0.0 se 3.0

% U-235 fission source in material fuel:

src 9 sc fuel sr 92235.03c 18

% U-238 fission source induced by 14 MeV neutrons:

src 10 sr 92238.03c 18 se 14.0

% Histogram energy distribution defined using 5 bins:

src 6 sb 5
1E-11 0.0 % Energy below 1E-11 MeV (weight must be zero)
1E-6   0.5 % Between 1E-11 and 1E-6 MeV, weight 0.5
1E-3   1.0 % Between 1E-6 and 1E-3 MeV, weight 1.0
1.0    2.0 % Between 1E-3 and 1.0 MeV, weight 2.0
20.0   1.0 % Between 1.0 and 20.0 MeV, weight 1.0
```

Chapter 10

Reaction rate mesh plotter

10.1 Mesh input

Serpent has a built-in capability to visualize the neutronics in thermal systems by plotting the fission power and thermal flux distributions in a single png graphics file. The parameters for a reaction rate mesh plotter are defined as:

```
mesh <or> <nx> <ny> [ <sym> <x0> <x1> <y0> <y1> <z0> <z1> ]
```

where

<or>	is the orientation of the plot plane (1, 2 or 3)
<nx>	is the width of the plot in pixels
<ny>	is the height of the plot in pixels
<sym>	is the symmetry option (0, 2, 4 or 8)
<x0>	is the minimum value of the x-coordinate
<x1>	is the maximum value of the x-coordinate
<y0>	is the minimum value of the y-coordinate
<y1>	is the maximum value of the y-coordinate
<z0>	is the minimum value of the z-coordinate
<z1>	is the maximum value of the z-coordinate

The code calculates reaction rates in an <nx>by <ny> mesh, and projects the data according to the orientation of the plot plane, defined as:

1. yz-plot (perpendicular to the x-axis)
2. xz-plot (perpendicular to the y-axis)
3. xy-plot (perpendicular to the z-axis)

If the optional coordinate boundaries are not given, the code uses the boundaries of the defined geometry.

The symmetry option can be used to attain better statistics. The symmetry types are illustrated in Fig. 5.1 on page 63, and only options 0, 2, 4 and 8 are allowed with mesh plots. The option is set to zero by default (no symmetry).

10.2 Mesh output

Output is written in a png format file “<input>_mesh<n>.png”, where <input> is the name of the input file and <n> is the plot index. Burnup mode produces new plots for each depletion step. The files are named “<input>_mesh<n>_bstep<m>.png”, where <m> is the step index.

The colour scheme consists of “hot” shades of red and yellow, representing relative fission power, and “cold” shades of blue, representing relative thermal flux (flux below 0.625 eV). The normalization is fixed after the first burnup step, so changes in flux and power level can be observed in the color schemes. Examples of reaction rate mesh plots can be found at the Serpent website: <http://montecarlo.vtt.fi/development.htm>.

IMPORTANT NOTES ON REACTION RATE MESH PLOTTER:

1. The mesh plots are subject to random noise, and the figures become smoother along with better statistics.
2. The geometry plotter uses the GD open source graphics library [1], which must be installed in the system.
3. The plotter produces png (portable network graphics) format output files.

SEE ALSO:

1. Compiling Serpent (Sec. 1.1 on page 8)
2. The GD open source graphics library: <http://www.libgd.org>
3. Mesh plot gallery at Serpent website:
<http://montecarlo.vtt.fi/development.htm>.

Chapter 11

Complete Input Examples

11.1 Quick start

For an experienced Monte Carlo code user the easiest way to get started with Serpent is to look at the lattice input examples in the following subsections. Installation and running the code is described in Chapter 1 and a general description of the input syntax is given in Chapter 2. The input cards used in the example cases include:

- Fuel pin definitions (Sec. 3.4 on page 27)
- Lattice definitions (Sec. 3.6 on page 28)
- Surface definitions (Sec. 3.2 on page 19)
- Cell definitions (Sec. 3.3 on page 24)
- Material definitions (Sec. 4.1.2 on page 48, see also Sec. 4.1.1)
- Thermal scattering libraries (Sec. 4.2 on page 49)
- Soluble absorber (Sec. 5.14 on page 69)
- File paths (Sec. 5.4 on page 57)
- Neutron population and criticality cycles (Sec. 5.2 on page 53)
- Boundary conditions (Sec. 5.7 on page 59)
- Parameters for group constant generation (Sec. 5.9 on page 64)
- Detectors (Chapter. 7 on page 95)

The examples describe the three main lattice types: square and hexagonal lattices and the circular cluster array. All geometries are two-dimensional and infinite in the axial direction.

The VVER-440 example in Sec. 11.1.1 demonstrates the use of soluble absorber and the calculation of various spectral quantities using detectors. The BWR case in Sec. 11.1.2 demonstrates the calculation of fast neutron flux ($E > 1$ MeV) in cladding and flow channel walls.

A more complicated mixed UOX/MOX lattice example is given in Sec. 11.1.4. The homogenization is carried over the central MOX assembly, but the use of a simple infinite MOX lattice would result in a distorted flux spectrum near the boundary between the two fuel types.

The input format is free and unrestricted. The only limitation is that command words must be separated by one or more white space characters. Due to the universe-based approach, similarities to MCNP input files are easy to see. To differentiate from the other examples, the mixed lattice case in Sec. 11.1.4 is prepared following a “SCALE-style” formulation.

More example cases are available at the Serpent website: <http://montecarlo.vtt.fi>.

11.1.1 VVER-440 lattice calculation

```
% --- VVER-440 Assembly -----

set title "VVER-440"

% --- Fuel pin with central hole:

pin 1
void    0.08000
fuel    0.37800
void    0.38800
clad    0.45750
water

% --- Central tube:

pin 2
water   0.44000
clad    0.51500
water

% --- Empty lattice position:

pin 3
water
```

```

% --- Lattice (type = 2, pin pitch = 1.23 cm):

lat 10  2  0.0 0.0 15 15 1.23
3 3 3 3 3 3 3 3 3 3 3 3 3 3 3
 3 3 3 3 3 3 3 1 1 1 1 1 1 1 3
    3 3 3 3 3 3 1 1 1 1 1 1 1 1 3
      3 3 3 3 3 1 1 1 1 1 1 1 1 1 3
        3 3 3 3 1 1 1 1 1 1 1 1 1 1 3
          3 3 3 3 1 1 1 1 1 1 1 1 1 1 3
            3 3 3 1 1 1 1 1 1 1 1 1 1 1 3
              3 3 1 1 1 1 1 1 1 1 1 1 1 1 3
                3 1 1 1 1 1 1 2 1 1 1 1 1 1 3
                  3 1 1 1 1 1 1 1 1 1 1 1 1 1 3 3
                    3 1 1 1 1 1 1 1 1 1 1 1 1 3 3 3
                      3 1 1 1 1 1 1 1 1 1 1 3 3 3 3
                        3 1 1 1 1 1 1 1 1 1 3 3 3 3 3
                          3 1 1 1 1 1 1 1 3 3 3 3 3 3
                            3 3 3 3 3 3 3 3 3 3 3 3 3

% --- Surfaces (assembly pitch = 14.7 cm):

surf 1  hexyc    0.0  0.0  7.100  % Shroud tube inner radius
surf 2  hexyc    0.0  0.0  7.250  % Shroud tube outer radius
surf 3  hexyc    0.0  0.0  7.350  % Outer boundary

% --- Cells:

cell  1  0  fill 10  -1          % Pin lattice
cell  4  0  tube    1    -2      % Shroud tube
cell  5  0  water   2    -3      % Water in channel
cell 99  0  outside        3      % Outside world

% --- UO2 fuel enriched to 3.6 wt-% U-235:

mat fuel  -10.45700
92235.09c  -0.03173
92238.09c  -0.84977
 8016.09c  -0.11850

% --- Zr-Nb cladding and shroud tube:

mat clad  -6.55000
40000.06c  -0.99000
41093.06c  -0.01000

mat tube  -6.58000
40000.06c  -0.97500
41093.06c  -0.02500

```

```
% --- Water:

mat water    -0.7207  moder lwtr 1001
  1001.06c    2.0
  8016.06c    1.0

% --- Thermal scattering data for light water:

therm lwtr lwj3.11t

% --- Natural boron (used as soluble absorber):

mat boron     1.0
  5010.06c     0.2
  5011.06c     0.8

% --- 650 ppm soluble absorber in water:

set abs boron -650E-6 water

% --- Cross section library file path:

set acelib "/xs/sss_jeff31.xsdata"

% --- Periodic boundary condition:

set bc 3

% --- Group constant generation:

% universe = 0 (homogenization over all space)
% symmetry = 12
% 2-group structure (group boundary at 0.625 eV)

set gcu 0
set sym 12
set nfg 2 0.625E-6

% --- Neutron population and criticality cycles:

set pop 2000 500 20

% --- Geometry and mesh plots:

plot 3 500 500
mesh 3 500 500
```



```
% --- Detector energy grid (uniform lethargy):

ene 1 3 1000 1E-9 12.0

% --- Flux per lethargy:

det 1 de 1 dt -3

% --- Differential capture, fission and production spectra:

det 2 de 1 dt -2 dr -2 void
det 3 de 1 dt -2 dr -6 void
det 4 de 1 dt -2 dr -7 void

% --- Integral capture, fission and production spectra:

det 5 de 1 dt -1 dr -2 void
det 6 de 1 dt -1 dr -6 void
det 7 de 1 dt -1 dr -7 void

% -----
```

11.1.2 BWR lattice calculation

```
% --- Asymmetric BWR assembly with Gd-pins -----

set title "BWR+Gd"

% --- Fuel Pin definitions:

pin 1
fuel1 4.33500E-01
void 4.42000E-01
clad 5.02500E-01
cool

pin 2
fuel2 4.33500E-01
void 4.42000E-01
clad 5.02500E-01
cool

pin 3
fuel3 4.33500E-01
void 4.42000E-01
clad 5.02500E-01
```

```

cool

pin 4
fuel4  4.33500E-01
void    4.42000E-01
clad    5.02500E-01
cool

pin 5
fuel5  4.33500E-01
void    4.42000E-01
clad    5.02500E-01
cool

pin 6
fuel6  4.33500E-01
void    4.42000E-01
clad    5.02500E-01
cool

pin 7
fuel7  4.33500E-01
void    4.42000E-01
clad    5.02500E-01
cool

% --- Empty lattice position:

pin 9
cool

% --- Lattice (type = 1, pin pitch = 1.295):

lat 10  1  0.0 0.0 12 12 1.295
9 9 9 9 9 9 9 9 9 9 9 9
9 1 2 3 5 5 5 5 5 3 2 9
9 2 3 5 6 6 6 6 7 5 4 9
9 3 5 7 6 7 6 6 6 6 5 9
9 5 6 6 6 6 6 6 7 6 6 9
9 5 6 7 6 9 9 9 6 7 6 9
9 5 6 6 6 9 9 9 6 6 6 9
9 5 6 6 6 9 9 9 6 6 6 9
9 5 7 6 7 6 6 6 7 6 5 9
9 3 5 6 6 7 6 6 6 6 5 9
9 2 4 5 6 6 6 6 5 5 3 9
9 9 9 9 9 9 9 9 9 9 9 9

% --- Outer channel (assembly pitch = 15.375):

```

```
surf 1  sqc    0.0    0.0    6.70000
surf 2  sqc    0.0    0.0    6.93000
surf 3  sqc   -0.233  -0.233  7.68750

% --- Channel inside assembly:

surf 4  sqc    0.6475  0.6475  1.6742
surf 5  sqc    0.6475  0.6475  1.7445

% --- Cell definitions:

cell  1  0  moder    -4      % Water inside moderator channel
cell  2  0  box       4 -5    % Moderator channel walls
cell  3  0  fill 10   -1  5    % Pin lattice
cell  4  0  box       1 -2    % Channel box wall
cell  5  0  moder     2 -3    % Water outside channel box
cell 99  0  outside   3      % Outside world

% --- Fuel materials:

mat fuel1  -10.424
  92235.09c  -0.015867
  92238.09c  -0.86563
  8016.09c   -0.1185

mat fuel2  -10.424
  92235.09c  -0.018512
  92238.09c  -0.86299
  8016.09c   -0.1185

mat fuel3  -10.424
  92235.09c  -0.022919
  92238.09c  -0.85858
  8016.09c   -0.1185

mat fuel4  -10.424
  92235.09c  -0.026445
  92238.09c  -0.85505
  8016.09c   -0.1185

mat fuel5  -10.424
  92235.09c  -0.029971
  92238.09c  -0.85153
  8016.09c   -0.1185

mat fuel6  -10.424
  92235.09c  -0.032615
```

```
92238.09c    -0.84888
8016.09c     -0.1185

% --- Fuel with Gd:

mat fuel7    -10.291
92235.09c    -3.13109E-02
92238.09c    -8.14929E-01
64152.09c    -6.70544E-05
64154.09c    -7.13344E-04
64155.09c    -5.06012E-03
64156.09c    -7.08860E-03
64157.09c    -5.43718E-03
64158.09c    -8.64341E-03
64160.09c    -7.69426E-03
8016.09c     -1.19056E-01

% --- Cladding and channel box wall:

mat clad     -6.55
40000.06c    -0.98135
24000.06c    -0.00100
26000.06c    -0.00135
28000.06c    -0.00055
50000.06c    -0.01450
8016.06c     -0.00125

mat box      -6.55
40000.06c    -0.98135
24000.06c    -0.00100
26000.06c    -0.00135
28000.06c    -0.00055
50000.06c    -0.01450
8016.06c     -0.00125

% --- Coolant (40% void fraction):

mat cool     -0.443760  moder lwtr 1001
1001.06c     0.66667
8016.06c     0.33333

% --- Moderator:

mat moder    -0.739605  moder lwtr 1001
1001.06c     0.666667
8016.06c     0.333333

% --- Thermal scattering data for light water:
```

```
therm lwtr lwj3.11t

% --- Cross section data library file path:

set acelib "/xs/sss_jeff31.xsdata"

% --- Reflective boundary condition:

set bc 2

% --- group constant generation:

% universe = 0 (homogenization over all space)
% symmetry = 4
% 4-group structure (3 group boundaries)

set gcu 0
set sym 4
set nfg 4 0.625E-6 5.5E-3 0.821

% --- Neutron population and criticality cycles:

set pop 2000 500 20

% --- Geometry and mesh plots:

plot 3 500 500
mesh 3 500 500

% --- Total power for normalization:

set power 1.96329E+04

% --- Detector energy grid (1 bin, E > 1.0 MeV):

ene 1 1 1.0 20

% --- Average fast flux in cladding:

det 1
de 1          % Use energy grid 1
dm clad       % Score in material "clad"
dv 16.3361    % Volume for normalization

% --- Pin-wise fast flux in cladding:

det 2
```

```

de 1          % Use energy grid 1
dm clad       % Score in material "clad"
dl 10         % Lattice bins in lat 10
dv 0.17952    % Volume for normalization

% --- Fast flux in inner moderator channel wall:

det 3
de 1          % Use energy grid 1
dc 2          % Score in cell 2
dv 0.96134    % Volume for normalization

% --- Fast flux in outer channel wall:

det 4
de 1          % Use energy grid 1
dc 4          % Score in cell 4
dv 12.5396    % Volume for normalization

% -----

```

11.1.3 CANDU lattice calculation

```

% --- CANDU cluster -----

set title "CANDU"

% --- Fuel pin:

pin 1
fuel 0.6122
clad 0.6540
cool

% --- Lattice (type = 4, 4 rings, 3rd ring rotated 15 deg.):

lat 10 4 0.0 0.0 4
1 0.0000 0.0 1
6 1.4885 0.0 1 1 1 1 1 1
12 2.8755 15.0 1 1 1 1 1 1 1 1 1 1 1
18 4.3305 0.0 1 1 1 1 1 1 1 1 1 1 1 1 1 1

% --- Surfaces (core pitch = 18.191 cm):

surf 1 cyl 0.0 0.0 5.16890 % Pressure tube inner wall
surf 2 cyl 0.0 0.0 5.60320 % Pressure tube outer wall

```

```

surf 3   cyl 0.0 0.0 6.44780 % Calandria tube inner wall
surf 4   cyl 0.0 0.0 6.58750 % Calandria tube outer wall
surf 5   sqc 0.0 0.0 9.09570 % Outer boundary

% --- Cells:

cell 1   0   fill 10   -1      % Pin lattice
cell 2   0   tube      1   -2   % Pressure tube
cell 3   0   void      2   -3   % Void between tubes
cell 4   0   caltube   3   -4   % Calandria tube
cell 5   0   moder     4   -5   % Moderator channel
cell 6   0   outside   5       % Outside world

% --- Fuel (UO2, natural uranium, 0.7% U-235):

mat fuel      -10.4375010
  8016.09c     -1.18473E+1
  92235.09c    -6.27118E-1
  92238.09c    -8.75256E+1

% --- Cladding:

mat clad      -6.44
  25055.06c    -1.60000E-1
  28000.06c    -6.00000E-2
  24000.06c    -1.10000E-1
  40000.06c    -9.97100E+1
   5010.06c    -5.7409e-05
   5011.06c    -2.5259E-04

% --- Pressure tube:

mat tube      -6.57
  40000.06c    -9.75000E+1
   5010.06c    -3.8889E-05
   5011.06c    -1.7111E-04

% --- Calandria tube:

mat caltube   -6.44
  25055.06c    -1.60000E-1
  28000.06c    -6.00000E-2
  24000.06c    -1.10000E-1
  40000.06c    -9.97100E+1
   5010.06c    -5.7409e-05
   5011.06c    -2.5259E-04

% --- Coolant water:

```

```
mat cool      -0.812120    moder lwtr 1001 moder hwtr 1002
  8016.06c    -7.99449E-1
  1002.06c    -1.99768E-1
  1001.06c    -7.83774E-4
```

```
% --- Moderator water:
```

```
mat moder     -1.082885    moder lwtr 1001 moder hwtr 1002
  8016.06c    -7.98895E-1
  1002.06c    -2.01016E-1
  1001.06c    -8.96000E-5
```

```
% --- Thermal scattering data for light and heavy water:
```

```
therm lwtr lwj3.11t
therm hwtr hwj3.11t
```

```
% --- Cross section data library file path:
```

```
set acelib "/xs/sss_jeff31.xsdata"
```

```
% --- Periodic boundary condition:
```

```
set bc 3
```

```
% --- group constant generation:
```

```
% universe = 0 (homogenization over all space)
% symmetry = 2
% 4-group structure (3 group boundaries)
```

```
set gcu 0
set sym 2
set nfg 4 0.625E-6 5.5E-3 0.821
```

```
% --- Neutron population and criticality cycles:
```

```
set pop 2000 500 20
```

```
% --- Geometry and mesh plots:
```

```
plot 3 500 500
mesh 3 500 500
```

```
% -----
```


11.1.4 Mixed UOX/MOX PWR lattice calculation

```
% --- PWR MOX/UOX lattice (SCALE-style input formulation) ----

% --- Problem title:

set title "MOX assembly in UOX lattice"

% --- Cross section library file path:

set acelib "/xs/sss_jeff31.xsdata"

% -----

% --- Material definitions ("comp block"):

% --- UOX fuel, initial enrichment 3.25%, burnup 25 MWd/kgU:

mat UO2      6.585000E-02
92235.09c    3.0000E-04
92236.09c    8.0000E-05
92238.09c    2.0000E-02
93237.09c    7.1000E-06
94238.09c    1.7000E-06
94239.09c    1.2000E-04
94240.09c    3.8000E-05
94241.09c    2.1000E-05
94242.09c    5.3000E-06
95241.09c    4.2000E-07
54131.09c    1.4000E-05
54135.09c    8.0000E-09
63153.09c    2.8000E-06
62149.09c    9.0000E-08
45103.09c    1.8000E-05
60143.09c    2.5000E-05
55133.09c    3.5000E-05
64155.09c    8.4000E-10
43099.09c    3.2000E-05
42095.09c    3.2000E-05
61147.09c    6.4000E-06
62150.09c    7.5000E-06
62151.09c    4.1000E-07
62152.09c    3.2000E-06
8016.09c     4.5100E-02

% --- Low Pu-content (2.9%) MOX fuel:
```

```
mat MOX1      6.702700E-02
92234.09c     4.3391E-07
92235.09c     4.9682E-05
92236.09c     8.6782E-07
92238.09c     2.1644E-02
94238.09c     5.4861E-06
94239.09c     4.3144E-04
94240.09c     1.3387E-04
94241.09c     4.8185E-05
94242.09c     1.8859E-05
95241.09c     9.1090E-06
8016.09c      4.4685E-02
```

% --- Medium Pu-content (4.4%) MOX fuel:

```
mat MOX2      6.702100E-02
92234.09c     4.2718E-07
92235.09c     4.8271E-05
92236.09c     8.5435E-07
92238.09c     2.1309E-02
94238.09c     8.1476E-06
94239.09c     6.5555E-04
94240.09c     2.0151E-04
94241.09c     7.4065E-05
94242.09c     2.7751E-05
95241.09c     1.4626E-05
8016.09c      4.4681E-02
```

% --- High Pu-content (5.6%) MOX fuel:

```
mat MOX3      6.701800E-02
92234.09c     4.2175E-07
92235.09c     4.9766E-05
92236.09c     8.4350E-07
92238.09c     2.1037E-02
94238.09c     1.0815E-05
94239.09c     8.3501E-04
94240.09c     2.5798E-04
94241.09c     9.4430E-05
94242.09c     3.6112E-05
95241.09c     1.7374E-05
8016.09c      4.4678E-02
```

% --- Zircaloy in cladding and guide tube:

```
mat can       4.004642E-02
40000.06c     3.9550E-02
26000.06c     1.3830E-04
```

```
24000.06c      7.0720E-05
8016.06c       2.8740E-04

% --- Water with 550 ppm boron:

mat water      7.088200E-02  moder lwtr 1001
1001.06c      4.7240E-02
8016.06c      2.3620E-02
5010.06c      4.3210E-06
5011.06c      1.7390E-05

% --- Thermal scattering data for light water:

therm lwtr lwj3.11t

% -----

% --- Parameters ("param block"):

% --- Periodic boundary condition:

set bc 3

% --- Group constant generation:

% universe = 200 (homogenization over MOX assembly)
% symmetry = 8
% 2-group structure (group boundary at 0.625 eV)

set gcu 200
set sym 8
set nfg 2 0.625E-6

% --- Neutron population and criticality cycles:

set pop 2000 500 20

% -----

% --- Geometry ("geom block"):

% --- UOX Pin ("unit 1"):

pin 1
UO2 0.41260
can 0.47400
water
```

```
% --- Guide tube ("unit 2"):
```

```
pin 2
water 0.57100
can 0.61300
water
```

```
% --- MOX Pins ("units 3-5"):
```

```
pin 3
MOX1 0.41260
can 0.47400
water
```

```
pin 4
MOX2 0.41260
can 0.47400
water
```

```
pin 5
MOX3 0.41260
can 0.47400
water
```

```
% --- UOX-assembly ("unit 100"):
```

```
surf 1000 sqc 0.0 0.0 10.727
```

```
cell 100 100 fill 110 -1000
cell 101 100 water 1000
```

```
% --- MOX-assembly ("unit 200"):
```

```
surf 2000 sqc 0.0 0.0 10.727
```

```
cell 200 200 fill 210 -2000
cell 201 200 water 2000
```

```
% --- Core lattice ("global unit 0"):
```

```
surf 3000 sqc 0.0 0.0 21.612
```

```
cell 300 0 fill 300 -3000
cell 301 0 outside 3000
```

```
% -----
```

```
% --- Lattices ("array block"):
```

```
% --- UOX pin lattice:
```

```
lat 110 1 0.0 0.0 17 17 1.262
```

```
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 2 1 1 2 1 1 2 1 1 1 1
1 1 1 2 1 1 1 1 1 1 1 1 1 2 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 2 1 1 1 1 1 1 1 1 1 2 1 1
1 1 1 1 1 2 1 1 2 1 1 2 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
```

```
% --- MOX pin lattice:
```

```
lat 210 1 0.0 0.0 17 17 1.262
```

```
3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3
3 4 4 4 4 4 4 4 4 4 4 4 4 4 4 3
3 4 4 4 4 2 4 4 2 4 4 2 4 4 4 3
3 4 4 2 4 5 5 5 5 5 5 5 4 2 4 3
3 4 4 4 5 5 5 5 5 5 5 5 5 4 4 3
3 4 2 5 5 2 5 5 2 5 5 2 5 5 2 4 3
3 4 4 5 5 5 5 5 5 5 5 5 5 4 4 3
3 4 4 5 5 5 5 5 5 5 5 5 5 4 4 3
3 4 2 5 5 2 5 5 2 5 5 2 5 5 2 4 3
3 4 4 5 5 5 5 5 5 5 5 5 5 4 4 3
3 4 4 5 5 5 5 5 5 5 5 5 5 4 4 3
3 4 2 5 5 2 5 5 2 5 5 2 5 5 2 4 3
3 4 4 4 5 5 5 5 5 5 5 5 5 4 4 3
3 4 4 2 4 5 5 5 5 5 5 5 4 2 4 3
3 4 4 4 4 2 4 4 2 4 4 2 4 4 4 3
3 4 4 4 4 4 4 4 4 4 4 4 4 4 4 3
3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3
```

```
% --- Core lattice:
```

```
lat 300 1 0.0 0.0 3 3 21.612
```

```

100 100 100
100 200 100
100 100 100

% -----

% --- Plotters ("plot block"):

% --- Geometry and mesh plots:

plot 3 500 500
mesh 3 500 500

% -----

```

11.2 Burnup calculation examples

11.2.1 Pin-cell burnup calculation

```

% --- Pin-cell burnup calculation -----

set title "Pin-cell burnup calculation"

% --- Pin definition:

pin 1
fuel 0.412
clad 0.475
water

% --- Geometry:

surf 1 sqc 0.0 0.0 0.665

cell 1 0 fill 1 -1
cell 2 0 outside 1

% --- Fuel (composition given in atomic densities):

mat fuel -10.045 burn 1
92234.09c 6.15169E+18
92235.09c 6.89220E+20
92236.09c 3.16265E+18
92238.09c 2.17103E+22

```

```
6012.09c    9.13357E+18
7014.09c    1.04072E+19
8016.09c    4.48178E+22

% --- Zircalloy cladding:

mat clad    -6.560
40000.06c   -0.9791
50000.06c   -0.0159
26000.06c   -0.0050

% --- Water (composition given in atomic densities):

mat water   -0.7569    moder lwtr 1001
1001.06c    5.06153E+22
8016.06c    2.53076E+22
5010.06c    2.75612E+18
5011.06c    1.11890E+19

% --- Thermal scattering data for light water:

therm lwtr lwj3.11t

% --- Cross section library file path:

set acelib  "/xs/sss_jeff31.xsdata"

% --- Periodic boundary condition:

set bc 3

% --- Group constant generation:

% universe = 0 (homogenization over all space)
% symmetry = 12
% 2-group structure (group boundary at 0.625 eV)

set gcu 0
set sym 12
set nfg 2 0.625E-6

% --- Neutron population and criticality cycles:

set pop 2000 500 20

% --- Geometry and mesh plots:

plot 3 500 500
```

```
mesh 3 500 500

% --- Decay and fission yield libraries:

set declib "/xs/JEFF311RDD"
set nfylib "/xs/JEFF31NIFY"

% --- Reduce energy grid size:

set egrid 5E-5 1E-9 15.0

% --- Cut-offs:

set fpcut 1E-9
set stabcut 1E-12
set ttacut 1E-18
set xsfcut 1E-6

% --- Options for burnup calculation:

set bumode 1 % TTA method
set pcc 1 % Predictor-corrector calculation on
set xscal 2 % Cross sections from spectrum
set printm 0 % No material compositions

% --- Depletion steps:

% Power density 40 kW/kgU
% Depletion steps given in units of total burnup

set powdens 40.0E-3

dep butot

0.1
0.5
1
5
10
15
20
25
30
35
40

% --- Isotope list for inventory calculation:
```



```
set inventory
922340
922350
922360
922380
932370
942380
942390
942400
942410
942420
952410
952430
420990
430990
441010
451030
471090
551330
621470
621490
621500
621510
621520
601430
601450
631530
641550
```

```
% -----
```

11.2.2 PWR assembly burnup calculation

```
set title "PWR Burnup Calculation Based on NEA Benchmark"

% --- Fuel pins:

pin 10
UO2      0.4025
clad     0.4750
water

pin 11
UO2      0.4025
clad     0.4750
```

water

pin 12

UO2 0.4025

clad 0.4750

water

pin 13

UO2 0.4025

clad 0.4750

water

pin 14

UO2 0.4025

clad 0.4750

water

pin 15

UO2 0.4025

clad 0.4750

water

pin 16

UO2 0.4025

clad 0.4750

water

pin 17

UO2 0.4025

clad 0.4750

water

pin 18

UO2 0.4025

clad 0.4750

water

pin 19

UO2 0.4025

clad 0.4750

water

pin 20

UO2 0.4025

clad 0.4750

water

pin 21

UO2 0.4025
clad 0.4750
water

pin 22
UO2 0.4025
clad 0.4750
water

pin 23
UO2 0.4025
clad 0.4750
water

pin 24
UO2 0.4025
clad 0.4750
water

pin 25
UO2 0.4025
clad 0.4750
water

pin 26
UO2 0.4025
clad 0.4750
water

pin 27
UO2 0.4025
clad 0.4750
water

pin 28
UO2 0.4025
clad 0.4750
water

pin 29
UO2 0.4025
clad 0.4750
water

pin 30
UO2 0.4025
clad 0.4750
water

pin 31
UO2 0.4025
clad 0.4750
water

pin 32
UO2 0.4025
clad 0.4750
water

pin 33
UO2 0.4025
clad 0.4750
water

pin 34
UO2 0.4025
clad 0.4750
water

pin 35
UO2 0.4025
clad 0.4750
water

pin 36
UO2 0.4025
clad 0.4750
water

pin 37
UO2 0.4025
clad 0.4750
water

pin 38
UO2 0.4025
clad 0.4750
water

pin 39
UO2 0.4025
clad 0.4750
water

pin 40
UO2 0.4025

clad 0.4750
water

pin 41
UO2 0.4025
clad 0.4750
water

pin 42
UO2 0.4025
clad 0.4750
water

pin 43
UO2 0.4025
clad 0.4750
water

pin 44
UO2 0.4025
clad 0.4750
water

pin 45
UO2 0.4025
clad 0.4750
water

% --- Gd-pins:

pin 50
UO2Gd 0.4025
clad 0.4750
water

pin 51
UO2Gd 0.4025
clad 0.4750
water

pin 52
UO2Gd 0.4025
clad 0.4750
water

% --- Guide tube:

pin 90

```

water    0.5730
tube     0.6130
water

```

```
% --- Pin lattice:
```

```
lat 110  1  0.0 0.0 17 17  1.265
```

```

45 44 43 42 41 40 39 38 37 38 39 40 41 42 43 44 45
44 36 35 34 33 32 31 30 29 30 31 32 33 34 35 36 44
43 35 28 27 52 90 26 25 90 25 26 90 52 27 28 35 43
42 34 27 90 24 23 22 21 51 21 22 23 24 90 27 34 42
41 33 52 24 20 19 18 17 16 17 18 19 20 24 52 33 41
40 32 90 23 19 90 15 14 90 14 15 90 19 23 90 32 40
39 31 26 22 18 15 50 13 12 13 50 15 18 22 26 31 39
38 30 25 21 17 14 13 11 10 11 13 14 17 21 25 30 38
37 29 90 51 16 90 12 10 90 10 12 90 16 51 90 29 37
38 30 25 21 17 14 13 11 10 11 13 14 17 21 25 30 38
39 31 26 22 18 15 50 13 12 13 50 15 18 22 26 31 39
40 32 90 23 19 90 15 14 90 14 15 90 19 23 90 32 40
41 33 52 24 20 19 18 17 16 17 18 19 20 24 52 33 41
42 34 27 90 24 23 22 21 51 21 22 23 24 90 27 34 42
43 35 28 27 52 90 26 25 90 25 26 90 52 27 28 35 43
44 36 35 34 33 32 31 30 29 30 31 32 33 34 35 36 44
45 44 43 42 41 40 39 38 37 38 39 40 41 42 43 44 45

```

```
% --- assembly data:
```

```

surf  1000  sqc  0.0 0.0 10.752
surf  1001  sqc  0.0 0.0 10.806

```

```

cell 110  0  fill 110  -1000
cell 111  0  water      1000 -1001
cell 112  0  outside    1001

```

```
% --- Materials:
```

```

mat UO2      6.7402E-02  burn 1
92234.09c    9.1361E-06
92235.09c    9.3472E-04
92238.09c    2.1523E-02
8016.09c     4.4935E-02

mat UO2Gd    6.8366E-02  burn 10
92234.09c    4.2940E-06
92235.09c    5.6226E-04
92238.09c    2.0549E-02
64154.09c    4.6173E-05

```

```
64155.09c 2.9711E-04
64156.09c 4.1355E-04
64157.09c 3.1518E-04
64158.09c 4.9786E-04
64160.09c 4.3764E-04
8016.09c 4.5243E-02

mat clad 3.8510E-02
26000.06c 1.3225E-04
24000.06c 6.7643E-05
40000.06c 3.8310E-02

mat tube 4.3206E-02
26000.06c 1.4838E-04
24000.06c 7.5891E-05
40000.06c 4.2982E-02

mat water 7.2216E-02 moder lwtr 1001
1001.06c 4.8132E-02
8016.06c 2.4066E-02
5010.06c 3.6487E-06
5011.06c 1.4686E-05

therm lwtr lwj3.11t

% --- Cross section library file path:

set acelib "/xs/sss_jeff31.xsdata"

% --- Periodic boundary condition:

set bc 3

% --- Neutron population and criticality cycles:

set pop 5000 500 20

% --- Geometry and mesh plots:

plot 3 500 500
mesh 3 500 500

% --- Decay and fission yield libraries:

set declib "/xs/JEFF311RDD"
set nfylib "/xs/JEFF31NFY"

% --- Reduce energy grid size:
```

```
set egrid 5E-5 1E-9 15.0

% --- Cut-offs:

set fpcut 1E-6
set stabcut 1E-12

% --- Options for burnup calculation:

set bumode 2 % CRAM method
set pcc 1 % Predictor-corrector calculation on
set xscal 2 % Cross sections from spectrum

% --- Irradiation cycle:

set powdens 38.6E-3

dep butot

0.10000
0.50000
1.00000
1.50000
2.00000
2.50000
3.00000
3.50000
4.00000
4.50000
5.00000
5.50000
6.00000
6.50000
7.00000
7.50000
8.00000
8.50000
9.00000
9.50000
10.00000
10.50000
11.00000
11.50000
12.00000
12.50000
13.00000
13.50000
```


14.00000
14.50000
15.00000
17.50000
20.00000
22.50000
25.00000
27.50000
30.00000
32.50000
35.00000
37.50000
40.00000

% --- Nuclide inventory:

set inventory

922340
922350
922360
922370
922380
922390
932360
932370
932380
932390
942360
942380
942390
942400
942410
942420
942430
952410
952420
952430
952440
952421
962420
962430
962440
962450
962460
962470
962480
962490

972490
972500
982490
982500
982510
982520
360830
451030
451050
471090
531350
541310
541350
551330
551340
551350
551370
561400
571400
601430
601450
611470
611480
611490
611481
621470
621490
621500
621510
621520
631530
631540
631550
631560
641520
641540
641550
641560
641570
641600

‰ -----

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