Users Guide V7.0

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Users Guide V7.0:Introduction

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Geant4 Application for Emission Tomography: a simulation toolkit for PET and SPECT

OpenGATE Collaboration

http://www.opengatecollaboration.org

Authors

OpenGATE spokesperson: I. Buvat (CNRS-IMNC Paris)

OpenGATE technical coordinator: S. Jan (CEA-SHFJ Orsay)

Editorial board S. Glick (UMASS), S. Kerhoas (CEA Saclay), F. Mayet (CNRS-LPSC Grenoble)

The GATE mailing list

You are encouraged to participate in the dialog and post your suggestions, questions and answers to colleagues' questions on the gate-users mailing list, the GATE mailing list for users. You can subscribe to the gate-users mailing list, by registering at the GATE web site: http://www.opengatecollaboration.org

If you have a question, it is very likely that it has already been asked and answered, and is now stored in the archives. Please use the search engine to see if your question has already been answered before sending a mail to the GATE-users.

Forewords

Monte Carlo simulation is an essential tool in emission tomography to assist in the design of new medical imaging devices, assess new implementations of image reconstruction algorithms and/or scatter correction techniques, and optimise scan protocols. Although dedicated Monte Carlo codes have been developed for Positron Emission Tomography (PET) and for Single Photon Emission Computerized Tomography (SPECT), these tools suffer from a variety of drawbacks and limitations in terms of validation, accuracy, and/or support (Buvat). On the other hand, accurate and versatile simulation codes such as GEANT3 (G3), EGS4, MCNP, and GEANT4 have been written for high energy physics. They all include well-validated physics models, geometry modeling tools, and efficient visualization utilities. However these packages are quite complex and necessitate a steep learning curve.

GATE, the GEANT4 Application for Emission Tomography (MIC02, Siena02, ITBS02, GATE, encapsulates the GEANT4 libraries in order to achieve a modular, versatile, scripted simulation toolkit adapted to the field of nuclear medicine. In particular, GATE provides the capability for modeling time-dependent phenomena such as detector movements or source decay kinetics, thus allowing the simulation of time curves under realistic acquisition conditions.

GATE was developed within the OpenGATE Collaboration with the objective to provide the academic community with a free software, general-purpose, GEANT4-based simulation platform for emission tomography. The collaboration currently includes 21 laboratories fully dedicated to the task of improving, documenting, and testing GATE thoroughly against most of the imaging systems commercially available in PET and SPECT (Staelens, Lazaro).

Particular attention was paid to provide meaningful documentation with the simulation software package, including installation and user's guides, and a list of FAQs. This will hopefully make possible the long term support and continuity of GATE, which we intend to propose as a new standard for Monte Carlo simulation in nuclear medicine.

In name of the OpenGATE Collaboration

Christian MOREL CPPM CNRS/IN2P3, Marseille
Member institutes of the OpenGATE Collaboration (September 2009):

- CNRS Laboratory of Corpuscular Physics @ Clermont-Ferrand (LPC)
- CNRS Centre de Physique des Particules de Marseille (CPPM), Marseille
- CNRS Imaging and Modelling in Neurobiology and Cancerology lab @ Orsay (IMNC), Orsay
- CNRS IRES, Centre Pluridisciplinaire Hubert Curien (CPHC) @ Strasbourg
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- LATIM, U650 INSERM, Brest
- Service Hospitalier Frédéric Joliot (SHFJ), CEA-Orsay
- Healthgrid
- University of Ghent (ELIS)
- Sungkyunkwan University School of Medicine (DMN), Seoul
- Forschungszentrum-Juelich (IME)
- University of California (Crump Institute for Molecular Imaging), Los Angeles
- Memorial Sloan-Kettering Cancer Center (Department of Medical Physics), New York
- University of Athens (IASA)
- Delft University of Technology (IRI)
- John Hopkins University (Division of Medical Imaging Physics), Baltimore
- University of Santiago of Chile (USACH)
- Department of Atomic Physics, Sofia University, Sofia
- Walter Reed Army Medical Center, Washington DC
- University of Pennsylvania School of Medicine

Overview

GATE combines the advantages of the GEANT4 simulation toolkit well-validated physics models, sophisticated geometry description, and powerful visualization and 3D rendering tools with original features specific to emission tomography. It consists of several hundred C++ classes. Mechanisms used to manage time, geometry, and radioactive sources form a core layer of C++ classes close to the GEANT4 kernel [Figure 1]. An application layer allows for the implementation of user classes derived from the core layer classes, e.g. building specific geometrical volume shapes and/or specifying operations on these volumes like rotations or translations. Since the application layer implements all appropriate features, the use of GATE does not require C++ programming: a dedicated scripting mechanism - hereafter referred to as the macro language - that extends the native command interpreter of GEANT4 makes it possible to perform and to control Monte Carlo simulations of realistic setups.
One of the most innovative features of GATE is its capability to synchronize all time-dependent components in order to allow a coherent description of the acquisition process. As for the geometry definition, the elements of the geometry can be set into movement via scripting. All movements of the geometrical elements are kept synchronized with the evolution of the source activities. For this purpose, the acquisition is subdivided into a number of time-steps during which the elements of the geometry are considered to be at rest. Decay times are generated within these time-steps so that the number of events decreases exponentially from time-step to time-step, and decreases also inside each time-step according to the decay kinetics of each radioisotope. This allows for the modeling of time-dependent processes such as count rates, random coincidences, or detector dead-time on an event-by-event basis. Moreover, the GEANT4 interaction histories can be used to mimic realistic detector output. In GATE, detector electronic response is modeled as a linear processing chain designed by the user to reproduce e.g. the detector cross-talk, its energy resolution, or its trigger efficiency.

The first users guide was organized as follow: chapter 1 of this document guides you to get started with GATE. The macro language is detailed in Chapter 2. Visualisation tools are described in Chapter 3. Then, Chapter 4 illustrates how to define a geometry by using the macro language, Chapter 5 how to define a system, Chapter 6 how to attach sensitive detectors, and Chapter 7 how to set up the physics used for the simulation. Chapter 8 discusses the different radioactive source definitions. Chapter 9 introduces the digitizer which allows you to tune your simulation to the very experimental parameters of your setup. Chapter 10 draws the architecture of a simulation. Data output are described in Chapter 11. Finally, Chapter 12 gives the principal material definitions available in GATE. Chapter 13 illustrates the interactive, bathc, or cluster modes of running GATE.

Actually, the users guide is based on a wiki page and all chapters and informations are pointed on the following link: Users Guide V7.0.

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Users Guide V7.0: Getting started

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This paragraph is an overview of the main steps one must go through to perform a simulation using Gate for imaging applications (PET, SPECT and CT). It is presented in the form of a simple example and the user is encouraged to try out the example, in an interactive mode of Gate, while reading this section. A more detailed description of the different steps is given in the following sections of this user's guide. The use of Gate does not require any C++ programming, thanks to a dedicated scripting mechanism that extends the native command interpreter of Geant4. This interface allows the user to run Gate programs using command scripts only. The goal of this first section is to give a brief description of the user interface and to provide a basic understanding of the basic principles of Gate by going through the different steps of a simulation.

General simulation architecture for imaging applications

In each simulation, the user has to:

1. define the scanner geometry
2. define the phantom geometry
3. set up the physics processes
4. initialize the simulation
5. set up the detector model
6. define the source(s)
7. specify the data output format
8. start the acquisition

Steps 1) to 4) concern the initialization of the simulation.

Following the initialization, the geometry can no longer be changed. The following paragraph will illustrate these different steps.

General simulation architecture for dosimetry and radiotherapy
applications

1. define the beam geometry
2. define the phantom geometry
3. specify the output (actor concept for dose map etc...)
4. set up the physics processes
5. initialize the simulation
6. define the source(s)
7. start the simulation with the following command lines
8. `/gate/application/setTotalNumberOfPrimaries [particle_number]
9. `/gate/application/start

The user interface: a macro language

Gate, just as GEANT4, is a program in which the user interface is based on scripts. To perform actions, the user must either enter commands in interactive mode, or build up macro files containing an ordered collection of commands.

Each command performs a particular function, and may require one or more parameters. The Gate commands are organized following a tree structure, with respect to the function they represent. For example, all geometry-control commands start with geometry, and they will all be found under the `/geometry` branch of the tree structure.

When Gate is run, the `Idle>` prompt appears. At this stage the command interpreter is active; i.e. all the Gate commands entered will be interpreted and processed on-line. All functions in Gate can be accessed to using command lines. The geometry of the system, the description of the radioactive source(s), the physical interactions considered, etc., can be parameterized using command lines, which are translated to the Gate kernel by the command interpreter. In this way, the simulation is defined one step at a time, and the actual construction of the geometry and definition of the simulation can be seen on-line. If the effect is not as expected, the user can decide to re-adjust the desired parameter by re-entering the appropriate command on-line. Although entering commands step by step can be useful when the user is experimenting with the software or when he/she is not sure how to construct the geometry, there remains a need for storing the set of commands that led to a successful simulation.

Macros are ASCII files (with '.mac' extension) in which each line contains a command or a comment. Commands are GEANT4 or Gate scripted commands; comments start with the character `#`. Macros can be executed from within the command interpreter in Gate, or by passing it as a command-line parameter to Gate, or by calling it from another macro. A macro or set of macros must include all commands describing the different components of a simulation in the right order. Usually these components are visualization, definitions of volumes (geometry), systems, digitizer, physics, initialization, source, output and start. These steps are described in the next sections. A single simulation may be split into several macros, for instance one for the geometry, one for the physics, etc. Usually, there is a master macro which calls the more specific macros (see Users Guide V6.2:Architecture of the simulation). Splitting macros allows the user to re-use one or more of these macros in several other simulations, and/or to organize the set of all commands. Examples of complete macros can be found on the web site referenced above. To execute a macro (mymacro.mac in this example) from the Linux prompt, just type:

```
Gate mymacro.mac
```
To execute a macro from inside the Gate environment, type after the "Idle>" prompt:

```
Idle>/control/execute mymacro.mac
```

And finally, to execute a macro from inside another macro, simply write in the master macro:

```
/control/execute mymacro.mac
```

In the following paragraphs, the main steps to perform a simulation using Gate are explained. To test this example, the user can run Gate and can execute all the commands proposed in this section, line by line.

**First step: Defining a scanner geometry**

The first command lines entered at the Gate prompt usually concern the graphical interface. For on-line verification of the geometry being built, a visualization tool needs to be installed, using the following commands.

```
# VISUALIZATION
/vis/open OGLSX
/vis/viewer/reset
/vis/viewer/set/viewpointThetaPhi 60 60
/vis/viewer/zoom 1
/vis/viewer/set/style surface
/vis/drawVolume
/tracking/storeTrajectory 1
/vis/scene/endOfEventAction accumulate
/vis/viewer/update
```

The different visualization tools and associated commands are discussed in Users Guide V6.2:Visualization. The visualization being set, the user needs to define the geometry of the simulation based on volumes. All volumes are linked together following a tree structure where each branch represents a volume. Each volume is characterized by shape, size, position, and material composition. The default material assigned to a new volume is Air. The list of available materials is defined in the GateMaterials.db file. (See Users Guide V6.2:Materials). The location of the material database needs to be specified with the following command:

```
/gate/geometry/setMaterialDatabase MyMaterialDatabase.db
```

The base of the tree is represented by the world volume (fig 1.1) which sets the experimental framework of the simulation. All Gate commands related to the construction of the geometry are described in detail in Users Guide V6:Defining a geometry. The world volume is a box centered at the origin. It can be of any size and has to be large enough to include the entire simulation geometry. The tracking of any particle stops...
when it escapes from the world volume. The example given here simulates a system that fits into a box of 40 x 40 x 40 cm³. Thus, the world volume may be defined as follows:

```plaintext
# WORLD
/gate/world/geometry/setXLength 40. cm
/gate/world/geometry/setYLength 40. cm
/gate/world/geometry/setZLength 40. cm
```

The world contains one or more sub volumes referred to as daughter volumes.

```plaintext
/gate/world/daughters/name vol_name
```

The name `vol_name` of the first daughter of the world has a specific meaning and name. It specifies the type of scanner to be simulated. Users Guide V6.2:Defining a system gives the specifics of each type of scanner, also called system. In the current example, the system is a CylindricalPET system. This system assumes that the scanner is based on a cylindrical configuration (fig 1.2) of blocks, each block containing a set of crystals.

```plaintext
# SYSTEM
/gate/world/daughters/name cylindricalPET
/gate/world/daughters/insert cylinder
/gate/cylindricalPET/setMaterial Water
/gate/cylindricalPET/geometry/setRmax 100 mm
/gate/cylindricalPET/geometry/setRmin 86 mm
/gate/cylindricalPET/geometry/setHeight 18 mm
/gate/cylindricalPET/vis/forceWireframe
/vis/viewer/zoom 3
```

These seven command lines describe the global geometry of the scanner. The shape of the scanner is a cylinder filled with water with an external radius of 100 mm and an internal radius of 86 mm. The length of the cylinder is 18 mm. The last command line sets the visualization as wireframe.

You may see the following message when creating the geometry:

```plaintext
G4PhysicalVolumeModel::Validate() called.
Volume of the same name and copy number ("world_phys", co
WARNING: This does not necessarily guarantee it's the same
volume you originally specified in /vis/scene/add/.
```

This message is normal and you can safely ignore it.

At any time, the user can list all the possible commands. For example, the command line for listing the visualization commands is:

```plaintext
Idle> ls /gate/cylindricalPET/vis/
```

Let's assume that the scanner is made of 30 blocks (box1), each block containing textstyle 8 times 8 LSO crystals (box2).
The following command lines describe this scanner (see Users Guide V6.2:Defining a geometry to find a detailed explanation of these commands). First, the geometry of each block needs to be defined as the daughter of the system (here cylindricalPET system).

```
# FIRST LEVEL OF THE SYSTEM
/gate/cylindricalPET/daughters/name boxl
/gate/cylindricalPET/daughters/insert box
/gate/boxl/placement/setTranslation 91. 0 0 mm
/gate/boxl/geometry/setXLength 10. mm
/gate/boxl/geometry/setYLength 17.75 mm
/gate/boxl/geometry/setZLength 17.75 mm
/gate/boxl/setMaterial Water
/gate/boxl/vis/setColor yellow
/gate/boxl/vis/forceWireframe
```

Once the block is created (figure 1.3), the crystal can be defined as a daughter of the block (figure 1.4)

```
# CRYSTAL
/gate/box1/daughters/name box2
/gate/box1/daughters/insert box
/gate/box2/geometry/setXLength 10. mm
/gate/box2/geometry/setYLength 2. mm
/gate/box2/geometry/setZLength 2. mm
/gate/box2/setMaterial LSO
/gate/box2/vis/setColor red
/gate/box2/vis/forceWireframe
```

The zoom command line in the script allows the user to zoom the geometry and the panTo command translates the viewer window in 60 mm in horizontal and 40 mm in vertical directions (the default is the origin of the world (0,0)).

To obtain the complete matrix of crystals, the volume box2 needs to be repeated in the Y and Z directions (fig 1.5). To obtain the complete ring detector, the original block is repeated 30 times (fig 1.6).

```
# REPEAT CRYSTAL
/gate/box2/repeaters/insert cubicArray
/gate/box2/cubicArray/setRepeatNumberX 1
/gate/box2/cubicArray/setRepeatNumberY 8
/gate/box2/cubicArray/setRepeatNumberZ 8
/gate/box2/cubicArray/setRepeatVector 0. 2.25 2.25 mm
```

The geometry of this simple PET scanner has now been specified. The next step is to connect this geometry to the system in order to store data from particle interactions (called hits) within the volumes which represent detectors (sensitive detector or physical volume). Gate only stores hits for those volumes attached to a sensitive detector. Hits regarding interactions occurring in non-sensitive volumes are lost. A volume must belong to a system before it can be attached to a sensitive detector. Hits, occurring in a volume, cannot be scored in an output file if this volume is not connected to a sensitive detector.
system because this volume can not be attached to a sensitive detector. The concepts of system and sensitive detector are discussed in more detail in Users Guide V6.2:Defining a system and Users Guide V6.2:Attaching the sensitive detectors respectively.

The following commands are used to connect the volumes to the system.

```
# REPEAT RSECTOR
/gate/box1/repeaters/insert ring
/gate/box1/ring/setRepeatNumber 30

# ZOOM
/vis/viewer/zoom 0.25
/vis/viewer/panTo 0 0 mm

# ATTACH Volumes To a SYSTEM
/gate/systems/cylindricalPET/rsector/attach box1
/gate/systems/cylindricalPET/module/attach box2
```

The names rsector and module are dedicated names and correspond to the first and the second levels of the cylindricalPET system (see Users Guide V6.2:Defining a system).

In order to save the hits (see Users Guide V6.2:Digitizer and readout parameters) in the volumes corresponding to the crystals the appropriate command, in this example, is:

```
# Define a SENSITIVE Detector
/gate/box2/attachCrystalSD vglue 1cm
```

At this level of the macro file, the user can implement detector movement. One of the most distinctive features of Gate is the management of time-dependent phenomena, such as detector movements and source decay leading to a coherent description of the acquisition process. For simplicity, the simulation described in this tutorial does not take into account the motion of the detector or the phantom. Users Guide V6.2:Defining a geometry describes the movement of volumes in detail.

### Second step: Defining a phantom geometry

The volume to be scanned is built according to the same principle used to build the scanner. The external envelope of the phantom is a daughter of the world. The following command lines describe a cylinder with a radius of 10 mm and a length of 30 mm. The cylinder is filled with water and will be displayed in gray. This object represents the attenuation medium of the phantom.
To retrieve information about the Compton and the Rayleigh interactions within the phantom, a sensitive detector (*phantomSD*) is associated with the volume using the following command line:

```plaintext
# PHANTOM defined as SENSITIVE
/gate/my_phantom/attachPhantomSD
```

Two types of information will now be recorded for each hit in the hit collection:

- The number of scattering interactions generated in all physical volumes attached to the *phantomSD*.
- The name of the physical volume attached to the *phantomSD* in which the last interaction occurred.

These concepts are further discussed in Users Guide V6.2: Attaching the sensitive detectors.

**Third step: Setting-up the physics processes**

Once the volumes and corresponding sensitive detectors are described, the interaction processes of interest in the simulation have to be specified. Gate uses the GEANT4 models for physical processes. The user has to choose among these processes for each particle. Then, user can customize the simulation by setting the production thresholds, the cuts, the electromagnetic options...

Some typical physics lists are available in the directory *examples/PhysicsLists*:

- `egammaStandardPhys.mac` (physics list for photons, e- and e+ with standard processes and
recommended Geant4 "option3")
- egammaLowEPhys.mac (physics list for photons, e- and e+ with low energy processes)
- hadrontherapyStandardPhys.mac (physics list for hadrontherapy with standard processes and recommended Geant4 "option3")
- hadrontherapyLowEPhys.mac (physics list for hadrontherapy with low energy processes)

The details of the interactions processes, cuts and options available in Gate are described in Users Guide V6.2: Setting up the physics.

**Fourth step: Initialization**

When the 3 steps described before are completed, corresponding to the pre-initialization mode of GEANT4, the simulation should be initialized using:

```
# INITIALIZE
/gate/run/initialize
```

This initialization actually triggers the calculation of the cross section tables. After this step, the physics list cannot be modified any more and new volumes cannot be inserted into the geometry.

**Fifth step: Setting-up the digitizer**

The basic output of Gate is a *hit* collection in which data such as the position, the time and the energy of each hit are stored. The history of a particle is thus registered through all the *hits* generated along its track. The goal of the *digitizer* is to build physical observables from the *hits* and to model readout schemes and trigger logics. Several functions are grouped under the Gate *digitizer* object, which is composed of different modules that may be inserted into a linear signal processing sequence. As an example, the following command line inserts an *adder* to sum the hits generated per elementary volume (a single crystal defined as box2 in our example).

```
/gate/digitizer/Singles/insert adder
```

Another module can describe the readout scheme of the simulation. Except when one crystal is read out by one photo-detector, the readout segmentation can be different from the elementary geometrical structure of the detector. The readout geometry is an artificial geometry which is usually associated with a group of sensitive detectors. In this example, this group is box1.

```
/gate/digitizer/Singles/insert readout
/gate/digitizer/Singles/readout/setDepth 1
```

In this example, the readout module sums the energy deposited in all crystals within the block and determines the position of the crystal with the highest energy deposited ("winner takes all"). The setDepth command specifies at which geometry level (called "depth") the readout function is performed. In the
current example:

- base level (CylindricalPET) = depth 0
- 1st daughter (box1) of the system = depth 1
- next daughter (box2) of the system = depth 2
- and so on ....

In order to take into account the energy resolution of the detector and to collect singles within a pre-defined energy window only, other modules can be used:

```plaintext
# ENERGY BLURRING
/gate/digitizer/Singles/insert blurring
/gate/digitizer/Singles/blurring/setResolution 0.19
/gate/digitizer/Singles/blurring/setEnergyOfReference 511. keV

# ENERGY WINDOW
/gate/digitizer/Singles/insert thresholder
/gate/digitizer/Singles/thresholder/setThreshold 350. keV
/gate/digitizer/Singles/insert upholder
/gate/digitizer/Singles/upholder/setUphold 650. keV
```

Here, an energy resolution of 19% at 551 KeV is considered.

Furthermore, the energy window is set from 350 keV to 600 keV.

For PET simulations, the coincidence sorter is also implemented at the digitizer level.

```plaintext
# COINCIDENCESORTER
/gate/digitizer/Coincidences/setWindow 10. ns
```

Other digitizer modules are available in Gate and are described in Users Guide V6.2:Digitizer and readout parameters.

**Sixth step: Setting-up the source**

In Gate, a source is represented by a volume in which the particles (positron, gamma, ion, proton, ...) are emitted. The user can define the geometry of the source and its characteristics such as the direction of emission, the energy distribution, and the activity. The lifetime of unstable sources (radioactive ions) is usually obtained from the GEANT4 database, but it can also be set by the user.

A voxelized phantom or a patient dataset can also be used to define the source, in order to simulate realistic acquisitions. For a complete description of all functions to define the sources, see Users Guide V6.2:Activity, source, voxelized phantoms.

In the current example, the source is a 1 MBq line source. The line source is defined as a cylinder with a radius of 0.5 mm and a length of 50 mm. The source generates pairs of 511 keV gamma particles emitted 'back-to-back' (for a more realistic source model, the range of the positron and the non collinearity of the two gammas can also be taken into account).

```plaintext
# SOURCE
/gate/source/addSource twogamma
```
Seventh step: Defining data output

By default, the data output formats for all systems used by Gate are ASCII and ROOT as described in the following command lines:

```
# ASCII Output format
/gate/output/ascii/enable
/gate/output/ascii/setFileName test
/gate/output/ascii/setOutFileHitsFlag 0
/gate/output/ascii/setOutFileSinglesFlag 1
/gate/output/ascii/setOutFileCoincidencesFlag 1

# ROOT Output format
/gate/output/root/enable
/gate/output/root/setFileName test
/gate/output/root/setRootSinglesFlag 1
/gate/output/root/setRootCoincidencesFlag 1
```

Given this script, several ASCII files (.dat extension) and a ROOT file (test.root) will be created. Users Guide V6.2:Data output explains how to read the resulting files.

For some scanner configurations, the events may be stored in a sinogram format or in List Mode Format (LMF). The sinogram output module stores the coincident events from a cylindrical scanner system in a set of 2D sinograms according to the parameters set by the user (number of radial bins and angular positions). One 2D sinogram is created for each pair of crystal-rings. The sinograms are stored either in raw format or ecat7 format. The List Mode Format is the format developed by the Crystal Clear Collaboration (LGPL licence). A library has been incorporated in Gate to read, write, and analyze the LMF format. A complete description of all available outputs is given in Users Guide V6.2:Data output.
Eighth step: Starting an acquisition

In the next and final step the acquisition is defined. The beginning and the end of the acquisition are defined as in a real life experiment. In addition, Gate needs a time slice parameter which defines time period during which the simulated system is assumed to be static. At the beginning of each time-slice, the geometry is updated according to the requested movements. During each time-slice, the geometry is kept static and the simulation of particle transport and data acquisition proceeds. Each slice corresponds to a Geant4 run.

If the sources of the simulation are not radioactive source or is they are not defined activity. User can fix the total number of events. In this case, the number of particles is splitted between slices in function of the time of each slice.

\[
\text{gate/application/setTotalNumberOfPrimaries \ [N]}
\]

User can also fix the same number of event per slice. In this case, each event is weighted by the ratio between the time slice and the total simulation time.

\[
\text{gate/application/setNumberOfPrimariesPerRun \ [N]}
\]

Regular time slice approach

This is the standard Gate approach for imaging applications (PET, SPECT and CT). User has to define the beginning and the end of the acquisition using the commands setTimeStart and setTimeStop. Each slice has the same duration. User has to define the slice duration (setTimeSlice).

\[
\begin{align*}
\text{gate/application/setTimeSlice} & \quad 1. \ s \\
\text{gate/application/setTimeStart} & \quad 0. \ s \\
\text{gate/application/setTimeStop} & \quad 1. \ s \\
\hline \\
\text{START the ACQUISITION} & \\
\text{gate/application/startDAQ}
\end{align*}
\]

The number of projections or runs of the simulation is thus defined by:

\[
N_{\text{run}} = \frac{\text{setTimeStop} - \text{setTimeStart}}{\text{setTimeSlice}}
\]
In the current example, there is no motion, the acquisition time equals 1 second and the number of projections equals one.

If you want to exit from the Gate program when the simulation time exceed the time duration, the last line of your program has to be `exit`.

**Slices with variable time**

In this approach, each slice has a specific duration. User has to defined the time of each slice. The first method is to use a file of time slices:

```
/gate/application/readTimeSlicesIn [File Name]
```

the second method is to add each slice with the command:

```
/gate/application/addSlice [value] [unit]
```

User has to define the beginning of the acquisition using the command `setTimeStart`. The end of acquisition is calculated by summing each time slice. The simulation is started with the commands:

```
/gate/application/start
```

or

```
/gate/application/startDAQ
```
NOTE: for dosimetry and radiotherapy applications, the general macro set-up should be quite different - A focus on the dedicated example folder is recommended.


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Users Guide V7.0: Defining a geometry

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- The world
- Creating a volume
- Repeating a volume
- Placing a volume
- Moving a volume
- Updating the geometry

The definition of a geometry is a key step in designing a simulation because it is through the geometry definition that the imaging device and object to be scanned are described. Particles are then tracked through the components of the geometry. This section explains how to define the different components of the geometry.

The world

Definition

The world is the only volume already defined in GATE when starting a macro. All volumes are defined as daughters or grand-daughters of the world. The world volume is a typical example of a GATE volume and has predefined properties. The world volume is a box centred at the origin. For any particle, tracking stops when it escapes from the world volume. The world volume can be of any size and has to be large enough to include all volumes involved in the simulation.

Use

The first volume that can be created must be the daughter of the world volume. Any volume must be included in the world volume. The geometry is built from the world volume.

Description and modification

The world volume has default parameters: shape, dimensions, material, visibility attributes and number of children. These parameters can be edited using the following GATE command:

```
/gate/world/describe
```

The output of this command is shown in figure 3.1. The parameters associated with the world volume can be modified to be adapted to a specific simulation configuration. Only the shape of the world volume, which is a box, cannot be changed. For instance, the X length can be changed from 50 cm to 2 m using:
The other commands needed to modify the *world* volume attributes will be given in the next sections.

### Creating a volume

#### Generality - Tree creation

When a volume is created with GATE, it automatically appears in the GATE tree (see Users Guide V6.2:Getting started). All commands applicable to the new volume are then available from this GATE tree. For instance, if the name of the created volume is *Volume_Name*, all commands applicable to this volume start with:

```
/gate/Volume_Name/
```

The tree includes the following commands:

- `setMaterial`: *To assign a material to the volume*
- `attachCrystalSD`: *To attach a crystal-SensitiveDetector to the volume*
- `attachPhantomSD`: *To attach a phantom-SensitiveDetector to the volume*
- `enable`: *To enable the volume*
- `disable`: *To disable the volume*
- `describe`: *To describe the volume*

The tree includes sub-trees that relate to different attributes of the volume *Volume_Name*. The available sub-trees are:

- `daughters`: *To insert a new 'daughter' in the volume*
- `geometry`: *To control the geometry of the volume*
- `vis`: *To control the display attributes of the volume*
- repeaters: To apply a new 'repeater' to the volume
- moves: To 'move' the volume
- placement: To control the placement of the volume

The commands available in each sub-tree will be described in #Building a volume, #Repeating a volume, #Placing a volume, #Moving a volume.

**Units**

Different units are predefined in GATE (see Table 3.1) and shall be referred to using the corresponding abbreviation. The list of units available in GATE can be edited using:

```plaintext
/units/list
```

inside the GATE environment (see Users Guide V6.2:Getting started).

**Axes**

Any position in the world is defined with respect to a three-axis system: X, Y and Z. These three axes can be seen in the display window using:

```plaintext
/vis/scene/add/axes
```
Figure 3.2: Three-axis system defined in GATE. The red, green and blue axes are the X, Y and Z axes respectively.

List of units available in GATE and corresponding abbreviations:

<table>
<thead>
<tr>
<th>LENGTH</th>
<th>SURFACE</th>
<th>VOLUME</th>
<th>ANGLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>parsec pc</td>
<td>kilometer2 km2</td>
<td>kilometer3 km3</td>
<td>radian rad</td>
</tr>
<tr>
<td>kilometer km</td>
<td>meter2 m2</td>
<td>kilometer3 km3</td>
<td>milliradian mrad</td>
</tr>
<tr>
<td>meter m</td>
<td>centimeter2 cm2</td>
<td>centimeter3 cm3</td>
<td>steradian sr</td>
</tr>
<tr>
<td>centimeter cm</td>
<td>millimeter2 mm2</td>
<td>millimeter3 mm3</td>
<td>degree deg</td>
</tr>
<tr>
<td>millimeter mm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>micrometer mum</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>nanometer nm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>angstrom Ang</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TIME</th>
<th>SPEED</th>
<th>ANGULAR SPEED</th>
<th>ENERGY</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Building a volume</th>
</tr>
</thead>
</table>

Any new volume must be created as the daughter of another volume (i.e., *World* volume or another volume previously created).

Three rules must be respected when creating a new volume:

- A volume which is located inside another must be its daughter
- A daughter must be fully included in its mother
- Volumes must not overlap
Errors in building the geometry yield wrong particle transportation, hence misleading results!

Creating a new volume

To create a new volume, the first step is to give it a name and a mother using:

```
/gate/mother_Volume_Name/daughters/name Volume_Name
```

This command prepares the creation of a new volume named `Volume_Name` which is the daughter of `mother_Volume_Name`.

Some names should not be used as they have precise meanings in gate. These names are the names of the GATE systems (see Users Guide V6.2:Defining a system) currently defined in GATE: `scanner`, `PETscanner`, `cylindricalPET`, `SPECTHead`, `ecat`, `CPET`, `OPET` and `OpticalSystem`.

The creation of a new volume is completed only when assigning a shape to the new volume. The tree

```
/gate/Volume_Name/
```

is then generated and all commands in the tree and the sub-trees are available for the new volume.

Different volume shapes are available, namely: `box`, `sphere`, `cylinder`, `cone`, `hexagon`, `general` or `extruded`, `trapezoid`, `wedge` and `elliptical tube`.

The command line for listing the available shapes is:

```
/gate/world/daughters/info
```

The command line for assigning a shape to a volume is:

```
/gate/daughter_Volume_Name/daughters/insert Volume_shape
```

where `Volume_shape` is the shape of the new volume.

`Volume_shape` must necessarily be one of the available names:

- `box` for a box - `sphere` for a sphere - `cylinder` for a cylinder - `ellipsoid` for an ellipsoid - `cone` for a cone - `eltub` for a tube with an elliptical base - `hexagon` for a hexagon - `polycone` for a polygon - `trap` for a general trapezoid - `trpd` for an extruded trapezoid and `wedge` for a wedge

The command line assigns the shape to the last volume that has been named.

The following command lists the daughters of a volume:

```
/gate/Volume_Name/daughters/list
```
Example

/gate/world/daughters/name Phantom
/gate/world/daughters/insert box

The new volume *Phantom* with a box shape is inserted in the *World* volume.

**Defining a size**

After creating a volume with a shape, its dimensions are the default dimensions associated with that shape. These default dimensions can be modified using the sub-tree `/geometry/`

The commands available in the sub-tree depend on the shape. The different commands for each type of shape are listed in table 3.2

These commands can be found in the directory

---

<table>
<thead>
<tr>
<th>BOX</th>
<th>TRPD</th>
</tr>
</thead>
<tbody>
<tr>
<td>setXLength: Set the length of the box along the X axis</td>
<td>setX1Length: Set half length along X of the plane at -dz position</td>
</tr>
<tr>
<td>setYLength: Set the length of the box along the Y axis</td>
<td>setY1Length: Set half length along Y of the plane at -dz position</td>
</tr>
<tr>
<td>setZLength: Set the length of the box along the Z axis</td>
<td>setX2Length: Set half length along X of the plane at +dz position</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>SPHERE</th>
<th>TRPD</th>
</tr>
</thead>
<tbody>
<tr>
<td>setRmin: Set the internal radius of the sphere (0 for full sphere)</td>
<td>setY2Length: Set half length along Y of the plane at +dz position</td>
</tr>
<tr>
<td>setRmax: Set the external radius of the sphere</td>
<td>setZLength: Set half length along Z of the trapezoid</td>
</tr>
<tr>
<td>setPhiStart: Set the start phi angle</td>
<td>setXBoxLength: Set half length along X of the extruded box</td>
</tr>
<tr>
<td>setDeltaPhi: Set the phi angular span (2PI for full sphere)</td>
<td>setYBoxLength: Set half length along Y of the extruded box</td>
</tr>
<tr>
<td>setThetaStart: Set the start theta angle</td>
<td>setZBoxLength: Set half length along Z of the extruded box</td>
</tr>
<tr>
<td>setDeltaTheta: Set the theta angular span (2PI for full sphere)</td>
<td>setXBoxPos: Set center position X of the box</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CYLINDER</th>
<th>TRPD</th>
</tr>
</thead>
<tbody>
<tr>
<td>setRmin: Set the internal radius of the cylinder (0 for full cylinder)</td>
<td>PARALLELEPIPED (... not yet implemented...)</td>
</tr>
<tr>
<td>setRmax: Set the external radius of the cylinder</td>
<td>setDx: Set Dx dimension of the parallelepiped</td>
</tr>
<tr>
<td>setHeight: Set the height of the cylinder</td>
<td>setDy: Set Dy dimension of the parallelepiped</td>
</tr>
<tr>
<td>Function</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>setPhiStart</td>
<td>Set the start phi angle</td>
</tr>
<tr>
<td>setDeltaPhi</td>
<td>Set the phi angular span (2PI for full cylinder)</td>
</tr>
<tr>
<td>setAlpha</td>
<td>Set Alpha angle</td>
</tr>
<tr>
<td>CONE</td>
<td></td>
</tr>
<tr>
<td>setRmin1</td>
<td>Set the internal radius of one side of the cone</td>
</tr>
<tr>
<td>setRmax1</td>
<td>Set the external radius of one side of the cone</td>
</tr>
<tr>
<td>setHeight</td>
<td>Set the height of the cone</td>
</tr>
<tr>
<td>setPhiStart</td>
<td>Set the start phi angle</td>
</tr>
<tr>
<td>setRmin2</td>
<td>Set the internal radius of one side of the cone</td>
</tr>
<tr>
<td>setRmax2</td>
<td>Set the external radius of one side of the cone</td>
</tr>
<tr>
<td>setHeight</td>
<td>Set the height of the cone</td>
</tr>
<tr>
<td>setPhiStart</td>
<td>Set the start phi angle</td>
</tr>
<tr>
<td>setDeltaPhi</td>
<td>Set the phi angular span (2PI for full cone)</td>
</tr>
<tr>
<td>POLYCONENOT YET IMPLEMENTED</td>
<td></td>
</tr>
<tr>
<td>setProfile</td>
<td>Set vectors of z, rInner, rOuter positions</td>
</tr>
<tr>
<td>HEXAGONE</td>
<td></td>
</tr>
<tr>
<td>setPhiStart</td>
<td>Set the start phi angle</td>
</tr>
<tr>
<td>setDeltaPhi</td>
<td>Set the phi angular span (2PI for full cone)</td>
</tr>
<tr>
<td>setRadius</td>
<td>Set the radius of the hexagon</td>
</tr>
<tr>
<td>ELLIPSOID</td>
<td></td>
</tr>
<tr>
<td>setXLength</td>
<td>Set the half axis length in the X direction</td>
</tr>
<tr>
<td>setYLength</td>
<td>Set the half axis length in the Y direction</td>
</tr>
<tr>
<td>setZLength</td>
<td>Set the half axis length in the Z direction</td>
</tr>
<tr>
<td>setXBottomCut</td>
<td>To cut the ellipsoide along the Z axis</td>
</tr>
<tr>
<td>setXTopCut</td>
<td>To cut the ellipsoide along the Z axis</td>
</tr>
<tr>
<td>setZBottomCut</td>
<td>To cut the ellipsoide along the Z axis</td>
</tr>
<tr>
<td>setZTopCut</td>
<td>To cut the ellipsoide along the Z axis</td>
</tr>
<tr>
<td>ELLIPTICAL TUBE</td>
<td></td>
</tr>
<tr>
<td>setXLength</td>
<td>Set the length of the semimajor axis</td>
</tr>
<tr>
<td>setYLength</td>
<td>Set the length of the semiminor axis</td>
</tr>
<tr>
<td>setHeight</td>
<td>Set the height of the tube</td>
</tr>
<tr>
<td>WEDGE</td>
<td></td>
</tr>
<tr>
<td>NarrowerXLength</td>
<td>Set the length of the shorter side of the wedge</td>
</tr>
<tr>
<td>XLength</td>
<td>Set the length of the wedge in the X direction</td>
</tr>
<tr>
<td>YLength</td>
<td>Set the length of the wedge in the Y direction</td>
</tr>
<tr>
<td>ZLength</td>
<td>Set the length of the wedge in the Z direction</td>
</tr>
</tbody>
</table>

For a box volume called *Phantom*, the X, Y and Z dimensions can be defined by:

```
/gate/Phantom/geometry/setXLength 20. cm
/gate/Phantom/geometry/setYLength 10. cm
/gate/Phantom/geometry/setZLength 5. cm
```

The dimensions of the *Phantom* volume are then 20 cm, 10 cm and 5 cm along the X, Y and Z axes respectively.

**Defining a material**
A material must be associated with each volume. The default material assigned to a new volume is Air. The list of available materials is defined in the GateMaterials.db file. (see Users Guide V7.0:Materials).

The following command fills the volume *Volume_Name* with a material called *Material*:

```bash
/gate/Volume_Name/setMaterial Material
```

**Example**

```bash
/gate/Phantom/setMaterial Water
```

The *Phantom* volume is filled with Water.

**Defining a color or an appearance**

To make the geometry easy to visualize, some display options can be set using the sub-tree /vis/

The commands available in this sub-tree are: setColor, setVisible, setDaughtersInvisible, setLineStyle, setLineWidth, forceSolid and forceWireframe (see Table 3.3)

<table>
<thead>
<tr>
<th>Command</th>
<th>Action</th>
<th>Argument</th>
</tr>
</thead>
<tbody>
<tr>
<td>setColor</td>
<td>Selects the color for the current volume</td>
<td>white, gray, black, red, green, blue, cyan, magenta and yellow</td>
</tr>
<tr>
<td>setVisible</td>
<td>Shows or hides the current volume</td>
<td></td>
</tr>
<tr>
<td>setDaughtersInvisible</td>
<td>Shows or hides the current volume daughters</td>
<td></td>
</tr>
<tr>
<td>setLineStyle</td>
<td>Sets the current volume line-style</td>
<td>dashed, dotted and unbroken</td>
</tr>
<tr>
<td>setLineWidth</td>
<td>Sets the current volume line-width</td>
<td></td>
</tr>
<tr>
<td>forceSolid</td>
<td>Forces solid display for the current volume</td>
<td></td>
</tr>
<tr>
<td>forceWireframe</td>
<td>Forces wireframe display for the current volume</td>
<td></td>
</tr>
</tbody>
</table>

These commands can be found in the tree /gate/Volume_Name/vis.

**Example**

```bash
/gate/Phantom/vis/setColor blue
/gate/Phantom/vis/forceWireframe
```

The *Phantom* volume will be displayed in blue and will be transparent.

**Enabling or disabling a volume**

A volume cannot be destroyed. The only possible action is to disable it: this makes the volume disappear.
from the display window but not from the geometry.

Only the *world* volume cannot be disabled.

To disable a volume *Volume_Name*, the command is:

```
/gate/Volume_Name/disable
```

The volume *Volume_Name* can be enabled again using:

```
/gate/Volume_Name/enable
```

- **Example**

```
/gate/Phantom/disable
```

The *Phantom* volume is disabled.

**Describing a volume**

The parameters associated with a volume *Volume_name* can be listed using:

```
/gate/Volume_Name/describe
```

- **Example**

```
/gate/Phantom/describe
```

The parameters associated with the *Phantom* volume are listed.

**Examples**

**How to build a NaI crystal**

```
/gate/mother_Volume_Name/daughters/name crystal
/gate/mother_Volume_Name/daughters/insert box
```

A volume named crystal is created as the daughter of a volume the shape of which is defined as a box.

```
/gate/crystal/geometry/setXLength 1. cm
/gate/crystal/geometry/setYLength 40. cm
/gate/crystal/geometry/setZLength 54. cm
```

The X, Y and Z dimensions of the volume crystal are set to 1 cm, 40 cm, and 54 cm respectively.
The new volume crystal is filled with NaI.

The new volume crystal is colored in yellow.

The previous command lists the parameters associated with the crystal volume.

The crystal volume is disabled

**How to build a "trpd" volume**

An alternative way of describing complicated geometries is to use a so-called "boolean" volume in order to describe one piece using a single volume instead of using a mother-children couple. This can make the description easier and more synthetic. The example below describes how the shape shown in Figure 3.3 can be defined using a trpd shape, based on a "boolean" volume consisting of a trapezoid "minus" a box:

The new volume called *trapeze_name*, which is...
the daughter of the *Volume_Name* volume, is described with 5+6 parameters. The first 5 parameters relate to the trapezoid, whereas the last 6 parameters describe the extruded volume using a box shape.

How to build a "Wedge" volume

Gate provides the class **Gate TrapCreator** to create and insert trapezoidal volumes into the geometry. To create a trapezoid, the user needs to specify eleven parameters (besides its name and material), which does not make it easy to use.

To model "slanted" crystals, a new class called **Gate WedgeCreator** (derived from **G4 Trap**) builds right angular wedges. As shown in Figure 3.4, a wedge is defined by only three parameters that are easily understood:

1. XLength: is the length of the wedge in the X direction.
2. Narrower XLength: is the length of the shorter side of the wedge in the X direction.
3. YLength: is the length in the Y direction.
4. ZLength: is the length in the Z direction.

For instance, the following macro lines insert a wedge crystal as a daughter of a module:

```
/gate/module/daughters/name wedge0
/gate/module/daughters/insert we
/gate/wedge0/geometry/setXLength 10 mm
/gate/wedge0/geometry/setNarrowerXLength 8.921 mm
/gate/wedge0/geometry/setYLength 2.1620 mm
/gate/wedge0/geometry/setZLength 2.1620 mm
/gate/wedge0/setMaterial LSO
/gate/wedge0/vis/setColor yellow
```

![Figure 3.4: When a wedge is inserted, it is oriented as shown in this figure](image)

**Repeating a volume**

To create X identical volumes, there is no need to create X different volumes. Only one volume must be created and then repeated. There are four different ways to repeat a volume: the linear repeater, the ring repeater, the cubic array repeater and the quadrant repeater.

To list the repeaters defined for the volume *Name_Volume*, use:

```
/gate/Name_Volume/repeaters/info
```

**Linear repeater**

The linear repeater is appropriate to repeat a volume along a direction (X, Y or Z axis). To use the linear repeater, first select this type of repeater using:
Then define the number of times \( N \) the volume \( Name\_Volume \) has to be repeated using:

\[/gate/Name\_Volume/linear/setRepeatNumber \ N\]

Finally, define the step and direction of the repetition using:

\[/gate/Name\_Volume/linear/setRepeatVector 0. 0. dZ. mm\]

A step of \( dZ \) mm along the \( Z \) direction is defined.

The "autoCenter" command allows the user to set the position of the repeated volumes:

\[/gate/Name\_Volume/linear/autoCenter true or false\]

The "true" option centers the group of repeated volumes around the position of the initial volume that has been repeated.

The "false" option centers the first copy around the position of the initial volume that has been repeated. The other copies are created by offset. The default option is true.

- Example

\[/gate/hole/repeaters/insert linear
\[/gate/hole/linear/setRepeatNumber 12
\[/gate/hole/linear/setRepeatVector 0. 4. 0. cm\]

The \( hole \) volume is repeated \( 12 \times \) every \( 4 \) cm along the \( Y \) axis. The application of this linear repeater is illustrated in figure 3.5.

**Ring repeater**

The ring repeater makes it possible to repeat a volume along a ring. It is useful to build a ring of detectors in PET.

To select the ring repeater, use:

\[/gate/Name\_Volume/repeaters/insert ring\]

To define the number of times \( N \) the volume \( Name\_Volume \) has to be repeated, use:

\[/gate/Name\_Volume/ring/setRepeatNumber \ N\]

Finally, the axis around which the volume \( Name\_Volume \) will be repeated must be defined by specifying
two points using:

```
/gate/Name_Volume/ring/setPoint1
```

```
/gate/Name_Volume/ring/setPoint2
```

The default rotation axis is the Z axis. Note that the default ring repetition goes counter clockwise.

These three commands are enough to repeat a volume along a ring over 360°. However, the repeat action can be further customized using one or more of the following commands. To set the rotation angle for the first copy, use:

```
/gate/Name_Volume/ring/setFirstAngle x deg
```

The default angle is 0 deg.

To set the rotation angle difference between the first and the last copy, use:

```
/gate/Name_Volume/ring/setAngularSpan x deg
```

The default angle is 360 deg.

The AngularSpan, the FirstAngle and the RepeatNumber allow one to define the rotation angle difference between two adjacent copies (AngularPitch).

\[
\frac{\text{Angular Span} - \text{First Angle}}{\text{Repeat Number} - 1} = \text{Angular Pitch}
\]

To set the number of objects in the periodic structure, hence the periodicity, use:

```
/gate/Name_Volume/ring/setModuloNumber M
```

When the volume auto-rotation option is enabled, the volume itself is rotated so that its axis remains tangential to the ring (see Figure 3.6). If this option is disabled, all repeated volumes keep the same orientation (see Figure 3.7). The commands for enabling or disabling the auto-rotation option are:
A volume can also be shifted along Z periodically. Each element of a sequence is shifted according to its position inside the sequence, defined as "j" below. In a sequence composed of $M_{\text{ModuloNumber}}$ elements, the shift values are defined as $Z_{\text{shift}_i} \equiv Z_{\text{shift}_j}$ where:

- $i$ is the position in the full ring
- $j = (i \% M_{\text{ModuloNumber}}) + 1$ is the position in a sequence, starting at 1.

To set a shift and the value of this shift, use:

```
/gate/Name_Volume/ring/setModuloNumber 1
/gate/Name_Volume/ring/setZShift1 Z mm
```

Up to 8 shifts and different shift values can be defined (setZShift1 to setZShift8).

Remark: This geometry description conforms to the document "List Mode Format Implementation: Scanner geometry description Version 4.1 M.Krieguer et al" and is fully described in the LMF output, in particular
in the ASCII header file entry:

\[ z \text{ shift sector } j \mod M_{\text{ModuloNumber}} : Z\text{shift}_j \text{ units} \]

Here \( j \) (starting here at 0) stands for the \( n^{th} \) object being shifted each \( M_{\text{ModuloNumber}} \) object. Each shift value introduced in the command line below corresponds to a new line in the .cch file.

The LMF version 22.10.03 supports a geometry with a cylindrical symmetry. As an example, a repeater starting at 0 degree and finishing at 90 degree (a quarter of ring) will not be supported by the LMF output.

- **Example 1**

```
/gate/hole/repeaters/insert ring
/gate/hole/ring/setRepeatNumber 10
/gate/hole/ring/setPoint1 0. 1. 0. mm
/gate/hole/ring/setPoint2 0. 0. 0. mm
```

The hole volume is repeated 10 times around the Y axis. The application of this ring repeater is illustrated in figure 3.8.

![Figure 3.6: Illustration of the application of the auto-rotation option](image-url)
Figure 3.7: Illustration of the application of the ring-repeater when the auto-rotation option is disabled
Figure 3.8: Illustration of the application of the ring repeater
Example 2

```
/gate/rsector/repeaters/insert ring
/gate/rsector/ring/setRepeatNumber 20
/gate/rsector/ring/setModuloNumber 2
/gate/rsector/ring/setZShift1 -3500 mum
/gate/rsector/ring/setZShift2 +3500 mum
/gate/rsector/ring/enableAutoRotation
```

The `rsector` volume is repeated 20 times along a ring. The sequence length is 2, with the first and the second volume shifted by -3500 µm and 3500 µm respectively. The `rsector` volume could also include several volumes itself, each of them being duplicated, which is illustrated in figure 3.9.

Cubic array repeater

The cubic array repeater is appropriate to repeat a volume along one, two or three axes. It is useful to build a collimator for SPECT simulations.

To select the cubic array repeater, use:

```
/gate/Name_Volume/repeaters/insert cubicArray
```
To define the number of times $N_x$, $N_y$ and $N_z$ the volume $Name\_Volume$ has to be repeated along the X, Y and Z axes respectively, use:

```
/gate/hole/cubicArray/setRepeatNumberX Nx
/gate/hole/cubicArray/setRepeatNumberY Ny
/gate/hole/cubicArray/setRepeatNumberZ Nz
```

To define the step of the repetition $X mm$, $Y mm$ and $Z mm$ along the X, Y and Z axes respectively, use:

```
/gate/hole/cubicArray/setRepeatVector X Y Z mm
```

The autocentering options are available for the cubic array repeater. If a volume is initially at a position $P$, the set of volumes after the repeater has been applied is centered on $P$ if autoCenter is true (default). If autoCenter is false, the first copy of the group is centered on $P$.

- Example

```
/gate/hole/repeaters/insert cubicArray
```
The hole volume is repeated 5 times each 5 cm along the Y axis and twice each 15 cm along the Z axis. The application of this cubic array repeater is illustrated in figure 3.10.

---

```
/gate/hole/cubicArray/setRepeatNumberX 1
/gate/hole/cubicArray/setRepeatNumberY 5
/gate/hole/cubicArray/setRepeatNumberZ 2
/gate/hole/cubicArray/setRepeatVector 0. 5. 15. cm
```

---

Figure 3.10: Illustration of the application of the cubic array repeater
Quadrant repeater

The quadrant repeater is appropriate to repeat a volume in a triangle-like pattern similar to that of a Derenzo resolution phantom.

To select the quadrant repeater, use:

```plaintext
/gate/Name_Volume/repeaters/insert quadrant
```

To define the number of repetition lines, use:

```plaintext
/gate/hole/quadrant/setLineNumber X
```

To define the orientation of the quadrant (the direction of line repetition), use:

```plaintext
/gate/hole/quadrant/setOrientation N deg
```

To define the distance between adjacent copies, use:
To define the maximum range of the repeater which is the maximum distance between a copy and the original volume, use:

```
/gate/hole/quadrant/setMaxRange xx cm
```

This command can be used to remove corner-copies that would fall outside your phantom

- **Example**

```
/gate/hole/repeaters/insert quadrant
/gate/hole/quadrant/setLineNumber 5
/gate/hole/quadrant/setOrientation 90 deg
/gate/hole/quadrant/setCopySpacing 6 cm
/gate/hole/quadrant/setMaxRange 30 cm
```

The *hole* volume is repeated in a triangle-like pattern. The application of this quadrant repeater is illustrated in figure 3.5.

![Illustration of the application of the cubic array repeater](image-url)
Remark: The repeaters that are applied to the Name_Volume volume can be listed using:

```
/gate/Name_Volume/repeaters/list
```

**Sphere repeater**

The sphere repeater makes it possible to repeat a volume along a spherical ring. It is useful to build rings of detectors for PET scanners having gantry of spherical shape (e.g. SIEMENS Ecat Accel, Hi-Rez, ....)

To select the sphere repeater, use:

```
/gate/Name_Volume/repeaters/insert sphere
```

Then, the radius R of the sphere can be set using:

```
/gate/Name_Volume /sphere/setRadius X cm
```

To define the number of times N1 and N2 the volume Name_Volume has to repeated in the transaxial plane and the axial plane respectively, use:

```
/gate/Name_Volume/sphere/setRepeatNumberWithTheta N1
/gate/Name_Volume/sphere/setRepeatNumberWithPhi N2
```

To set the rotation angle difference between two adjacent copies in the transaxial direction, use:

```
/gate/Name_Volume/sphere/setThetaAngle x deg
```

To set the rotation angle difference between two adjacent copies in the axial direction, use:
The replicates of the volume *Name_Volume* will be placed so that its axis remains tangential to the ring.

**Example 3.12**

```python
/gate/block/repeaters/insert sphere
/gate/block/sphere/setRadius 25. cm
/gate/block/sphere/setRepeatNumberWithTheta 10
/gate/block/sphere/setRepeatNumberWithPhi 3
/gate/block/setThetaAngle 36 deg
/gate/block/setThetaAngle 20 deg
```

The block volume is repeated 10 times along the transaxial plane, with a rotation angle between two neighbouring blocks of 36 deg, and is repeated 3 times in the axial direction with a rotation angle between...
two neighbouring blocks of 20 deg. The sphere defined here has a 25 cm radius.

**Generic repeater**

It is also possible to repeat a volume according to a list of transformations (rotation and translation). The following macros read the transformations into a simple text file:

```markdown
/gate/myvolume/repeaters/insert                genericRepeater
/gate/myvolume/genericRepeater/setPlacementsFilename  data/myvolume.placements
/gate/myvolume/genericRepeater/useRelativeTranslation 1
```

The text file "myvolume.placements" is composed as follows:

```markdown
##### List of placement (translation and rotation)
##### Column 1 is rotationAngle in degree
##### Columns 2,3,4 are rotation axis
##### Columns 5,6,7 are translation in mm
Rotation deg
Translation mm
0       0 1 0       0 0 10
10      0 1 0       0 0 10
15      0 1 0       0 0 20
```

- line with # are ignored
- first word must be Rotation or Translation followed with the unity (deg and mm here)
- Rotation are described with 4 columns, the first for the angle, three others for the rotation axis
- Translation are described with X Y Z.
- using "useRelativeTranslation 1" (default) allows to compose the transformation according to the initial volume translation. If set to 0, the transformation is set as is (in the coordinate system of the mother volume).

See example here

**Placing a volume**

The position of the volume in the geometry is defined using the sub-tree

```
/placement/
```

Three types of placement are available: translation, rotation and alignment.

**Translation**

To translate the *Name_Volume* volume along the X direction by x cm, the command is:

```
/gate>Name_Volume/placement/setTranslation x. 0. 0. cm
```
The position is always given with respect to the center of the mother volume.

To set the Phi angle (in XY plane) of the translation vector, use:

\[ /gate/Name\_Volume/placement/setPhiOfTranslation \ N \ deg \]

To set the Theta angle (with regard to the Z axis) of the translation vector, use:

\[ /gate/Name\_Volume/placement/setThetaOfTranslation \ N \ deg \]

To set the magnitude of the translation vector, use:

\[ /gate/Name\_Volume/placement/setMagOfTranslation \ xx \ cm \]

**Example**

\[ /gate/Phantom/placement/setTranslation \ 1. \ 0. \ 0. \ cm \]
\[ /gate/Phantom/placement/setMagOfTranslation \ 10. \ cm \]

The *Phantom* volume is placed at 10 cm, 0 cm and 0 cm from the center of the mother volume (here the *world* volume). The application of this translation placement is illustrated in figure 3.13.

**Rotation**

To rotate the *Name\_Volume* volume by *N* degrees around the *X* axis, the commands are:

\[ /gate/Name\_Volume/placement/setRotationAxis \ X \ 0 \ 0 \]
\[ /gate/Name\_Volume/placement/setRotationAngle \ N \ deg \]
\[ /gate/Name\_Volume/placement/setAxis \ 0 \ 1 \ 0 \]

The default rotation axis is the *Z* axis.

**Example**

\[ /gate/Phantom/placement/setRotationAxis \ 0 \ 1 \ 0 \]
\[ /gate/Phantom/placement/setRotationAngle \ 90 \ deg \]

The *Phantom* volume is rotated by 90 degrees around the *Y* axis. The application of this rotation placement is illustrated in figure 3.14.

**Alignment**

Using the alignment command, a volume having an axis of symmetry (cylinder, ellipsoid, cone and hexagon) can be aligned parallel to one of the three axes of the axis system.

To align the *Name\_Volume* volume along the *X* axis, use:
The rotation parameters of the Name_Volume volume are then set to +90 degree around the Y axis.

To align the Name_Volume volume along the Y axis, use:

```
/gate/Name_Volume/placement/alignToY
```

The rotation parameters of the Name_Volume volume are then set to -90 degree around the X axis.

To align the Name_Volume volume along the Z axis (default axis of rotation) use:

```
/gate/Name_Volume/placement/alignToZ
```

The rotation parameters of the Name_Volume volume are then set to 0 degree.

**Special example: Wedge volume and OPET scanner**

The wedge is always created as shown in figure 3.4, that is with the slanted plane oriented towards the positive X direction. If one needs to have it oriented differently, one could, for instance, rotate it:

```
/gate/wedge0/placement/setRotationAxis 0 1 0
/gate/wedge0/placement/setRotationAngle 180 deg
```

The center of a wedge in the Y and Z directions are simply

\[
setYLength = \frac{setZLength}{2}, \quad setZLength = \frac{2}{2}
\]

respectively. For the X direction, the center is located such that

\[
2\Delta = \frac{setXLength + setNarrowerXLength}{2}
\]

where Delta is the length of the wedge across the middle of the Y direction, as shown in Figure 3.15.
Wedge crystals are used to build the OPET scanner, in which the scanner ring geometry approximates a true circular ring.

By knowing the radius gantry R and the length of the longest crystal, it is possible to arrange a series of 8 crystals with varying the lengths as shown in Figure 3.16.

It is first necessary to create by-hand the first row of crystals. This is accomplished by first creating a module just big enough to contain one row of wedge crystals.

```
gate/rsector/daughters/name module
gate/rsector/daughters/insert box
/gate/module/geometry/setXLength 10 mm
/gate/module/geometry/setYLength 17.765 mm
/gate/module/geometry/setZLength 2.162 mm
/gate/module/setMaterial Air
```
Figure 3.14: Illustration of the rotation placement
Figure 3.14: Illustration of the rotation placement

Figure 3.16: A block approximating a true circular geometry
Then, a box that will contain the first wedge crystal is located inside the module:

```
/gate/module/daughters/name crystal0
/gate/module/daughters/insert box
/gate/crystal0/geometry/setXLength 10 mm
/gate/crystal0/geometry/setYLength 2.1620 mm
/gate/crystal0/geometry/setZLength 2.1620 mm
/gate/crystal0/placement/setTranslation 0. -7.8015 0. mm
/gate/crystal0/setMaterial Air
/gate/crystal0/vis/setColor black
/gate/crystal0/vis/setVisible false
```

Finally, the actual crystal is placed inside its box:

```
/gate/crystal0/daughters/name LSO0
/gate/crystal0/daughters/insert wedge
/gate/LSO0/geometry/setXLength 10 mm
/gate/LSO0/geometry/setNarrowerXLength 8.921 mm
/gate/LSO0/geometry/setYLength 2.1620 mm
/gate/LSO0/geometry/setZLength 2.1620 mm
/gate/LSO0/placement/setRotationAxis 0 1 0
/gate/LSO0/placement/setRotationAngle 180 deg
/gate/LSO0/placement/setTranslation 0.2698 0. 0. mm
```
It is necessary to locate each crystal in separate layers.

The last two steps are repeated for each crystal inside the module. Then the module is repeated along the Z axis and the block is repeated 6 times around the center of the scanner.

Figure 4.9 shows the final OPET scanner.

**Moving a volume**

The GEANT geometry architecture requires the geometry to be static during a simulation. However, the typical duration of a single event (e.g. ps for the particle transport, μs for scintillation, or ms for the response of the electronics) is very short when compared to most of the geometrical changes to be modeled (e.g. movements of the phantom or of the detector or bio-kinetics). Therefore, the elements of the geometry are considered to be at rest during each time-step. Between every time-step, the position and the orientation of a subset of daughter volumes can be changed to mimic a movement such as a rotation or a translation. These displacements are parametrized by their velocity. Hence, the amplitude of the volume displacement is deduced from the duration of the time-step multiplied by the velocity of the displacement.

Given the speed of the components of the geometry, it is the responsibility of the user to set the time step duration short enough in order to produce smooth changes.

A volume can be moved during a simulation using five types of motion: rotation, translation, orbiting, wobbling and eccentric rotation, as explained below.

**Translation**

To translate a *Name_Volume* volume during the simulation, the commands are:

```
/gate/Name_Volume/moves/insert translation
/gate/Name_Volume/translation/setSpeed x 0 0 cm/s
```

where *x* is the speed of translation and the translation is performed along the X axis. These commands can be useful to simulate table motion during a scan for instance.

- **Example**

```
/gate/Table/moves/insert translation
/gate/Table/translation/setSpeed 0 0 1 cm/s
```

The *Table* volume is translated along the Z axis with a speed of 1 cm per second.

**Rotation**

To rotate a *Name_Volume* volume around an axis during the simulation, with a speed of *N* degrees per second, the commands are:
Example

```
/gate/Phantom/moves/insert rotation
/gate/Phantom/rotation/setSpeed 1 deg/s
/gate/Phantom/rotation/setAxis 0 1 0
```

The *Phantom* volume rotates around the Y axis with a speed of 1 degree per second.

**Orbiting**

Rotating a volume around any axis during a simulation is possible using the orbiting motion. This motion is needed to model the camera head rotation in SPECT. To rotate the *Name_Volume* volume around the X axis with a speed of N degrees per second, the commands are:

```
/gate/SPECTHead/moves/insert orbiting
/gate/SPECTHead/orbiting/setSpeed N. deg/s
/gate/SPECTHead/orbiting/setPoint1 0 0 0 cm
/gate/SPECTHead/orbiting/setPoint2 1 0 0 cm
```

The last two commands define the rotation axis.

It is possible to enable or disable the volume auto-rotation option using:

```
/gate/Name_Volume/orbiting/enableAutoRotation
/gate/Name_Volume/orbiting/disableAutoRotation
```

Example

```
/gate/camera_head/moves/insert orbiting
/gate/camera_head/orbiting/setSpeed 1. deg/s
/gate/camera_head/orbiting/setPoint1 0 0 0 cm
/gate/camera_head/orbiting/setPoint2 0 0 1 cm
```

The *camera_head* volume is rotated around the Z axis during the simulation with a speed of 1 degree per second.

**Wobbling**

The wobbling motion enables an oscillating translation movement to the volume.

This motion is needed to mimic the behavior of certain PET scanners that wobble to increase the spatial sampling of the data during the acquisition.

The movement that is modeled is defined by $dM(t) = A \sin(2\pi f t + \phi)$ where $dM(t)$ is the translation vector at time $t$, $A$ is the maximum displacement vector, $f$ is the movement frequency, $\phi$ is the phase at
t=0, and t is the time.

To set the parameters of that equation, use:

```gate/Name_Volume/moves/insert osc-trans```

To set the amplitude vector of the oscillating translation:

```gate/Name_Volume/osc-trans/setAmplitude x. 0. 0. cm```

To set the frequency of the oscillating translation:

```gate/Name_Volume/osc-trans/setFrequency N Hz```

To set the period of the oscillating translation:

```gate/Name_Volume/osc-trans/setPeriod N s```

To set the phase at t=0 of the oscillating translation:

```gate/Name_Volume/osc-trans/setPhase N deg```

- Example

```gate/crystal/moves/insert osc-trans
/gate/crystal/osc-trans/setAmplitude 10. 0. 0. cm
/gate/crystal/osc-trans/setFrequency 50 Hz
/gate/crystal/osc-trans/setPeriod 1 s
/gate/crystal/osc-trans/setPhase 90 deg```

In this example, the movement that is modeled is defined by $dM(t) = 10\sin(100\pi t + 90)$

**Eccentric rotation**

The eccentric rotation motion enables an eccentric rotation movement of the volume. It is a particular case of the orbiting movement. To set the object in eccentric position (X-Y-Z) and rotate it around the OZ lab frame axis, use:

```gate/Name_Volume/moves/insert eccent-rot```

To set the shifts in the X-Y-Z directions:

```gate/Name_Volume/eccent-rot/setShiftXYZ x y z cm```
To set the orbiting angular speed:

```
/gate/Name_Volume/eccent-rot/setSpeed N deg/s
```

Remark: This particular move is closely related to the LMF definition since the move parameters (shifts in all 3 directions and angular speed) are propagated in the .cch header.

- Example

```
/gate/crystal/moves/insert eccent-rot
/gate/crystal/eccent-rot/setShiftXYZ 5. 0. 0. cm
/gate/crystal/eccent-rot/setSpeed 10 deg/s
```

The *crystal* volume is placed at 10 cm, 0 cm and 0 cm from the center of its mother volume and will rotate around the Z axis during the simulation with a speed of 10 degrees per second.

### Generic move

A volume can be move at given time value thanks to the following macros:

```
/gate/myvolume/moves/insert genericMove
/gate/myvolume/genericMove/setPlacementsFilename data/myvolume.placements
```

In the same idea than **GenericRepeater**, the placements file contains the transformations (rotation, translation) and the time value where this transformations is applied.

```
##### List of placement (translation and rotation) according to time
##### Column 1      is Time in s (second)
##### Column 2      is rotationAngle in degree
##### Columns 3,4,5 are rotation axis
##### Columns 6,7,8 are translation in mm

Time s   Rotation deg   Translation mm  
0        0            0 1 0 0  0 100
250.7    3            0 1 0 0  10 100
492.9    4            0 1 0 0  20 100
742.9    8            0 1 0 0  30 100
```

**WARNING**. The time values given here do not necessarily correspond to simulation's *run*. The real runs are defined with the time slices (see this section for example). At each new run, GATE look into the timeplacements list and choose the one that correspond to the starting time of the run. It leads that some placements can be not applied (if one run start before the placement time and the next run start after the next placement time). If run time is after the last placements time in the list, the last placements is applied.

See example here

### Generic repeater move
You can combine generic repeater and generic move to allow different repeated configurations according to time. This is for example useful to describe multi-leafs collimator from a single leaf which is repeated at different positions, and which move according to each beam.

```
/gate/myvolume/moves/insert genericRepeaterMove
/gate/myvolume/genericRepeaterMove/setPlacementsFilename data/myvolume.placements
/gate/myvolume/genericRepeaterMove/useRelativeTranslation 1
```

```text

# List of placement (translation and rotation)
# Column 1 is rotationAngle in degree
# Columns 2,3,4 are rotation axis
# Columns 5,6,7 are translation in mm

<table>
<thead>
<tr>
<th>Time s</th>
<th>NumberOfPlacements 3</th>
<th>Rotation degr</th>
<th>Translation mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>10 0 1 0 20 0 0</td>
<td>10 0 1 0 80 0 0</td>
<td>10 0 1 0 -60 0 0</td>
</tr>
<tr>
<td>1</td>
<td>20 0 1 0 20 10 0</td>
<td>20 0 1 0 80 10 0</td>
<td>20 0 1 0 -60 10 0</td>
</tr>
<tr>
<td>2</td>
<td>30 1 1 0 20 0 0</td>
<td>30 1 1 0 80 0 0</td>
<td>30 1 1 0 -60 0 0</td>
</tr>
<tr>
<td>4</td>
<td>40 0 1 1 20 0 40</td>
<td>40 0 1 1 80 0 40</td>
<td>40 0 1 1 -60 0 40</td>
</tr>
</tbody>
</table>
```

The 'NumberOfPlacements' is needed to indicate how many different repetition are performed at each motion.

## Updating the geometry

Updating the geometry is needed to take into account any change in the geometry. It also refreshes the display window. The geometry can be updated with the following command.

```
/gate/geometry/update
```

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Users Guide V7.0:Materials

From GATE collaborative documentation wiki

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The Gate material database

The primary method for defining the properties of the materials used in Gate is by a materials database. This file holds all the information required for Gate to assign the nuclear properties from the Geant4 data sets, and is easily modified by the user. The OpenGate collaboration supplies a fairly extensive listing of materials in this file as part of Gate. This chapter describes the details of how to modify this database.

As alluded to in the previous paragraph, there exists an alternate method for materials definitions. As discussed in previous chapters, Gate scripts are developed from Geant4 C++ data classes in order to simplify and standardize input for Geant4. As a result, materials definitions can be written and compiled in C++ directly using the Geant4 tools. Specifying materials in this manner is beyond the scope of this document. For those interested in direct access to Geant4’s materials should refer to the Geant4 User’s Guide: For Application Developers and the Geant4 User’s Guide: For Toolkit Developers for more detailed information.

The material database contains two Geant4 structures called elements and materials that are used to define the physical properties of the atoms, molecules, and compounds. In contrast with Geant4, Gate does not use isotopic abundances. This omission has little bearing on Gate applications because isotopic abundances are unimportant in low to mid energy photon and charged particle interactions. In fact, this distinction is only important for enriched or depleted materials interacting with neutrons or, high energy (>5 MeV) photons or charged particles.

It is possible to use several material database. If a material is defined in several database, Gate keeps the material in last database called (with a warning message). To call a database:

```
/gate/geometry/setMaterialDatabase MyMaterialDatabase.db
```

Elements

Elements are the building blocks of all the materials used in Gate simulations. Elements in Gate are defined as in a periodic table. Gate stores the elements name, symbol, atomic number, and molar mass. As stated above, isotopic abundances are not referenced or used. The supplied file GateMaterials.db contains the most commonly used elements and their molar masses as they are found in nature.

Some elements, particularly those that have an isotope with a large cross section for neutron absorption,
have isotopic abundances and thus molar masses that vary depending upon their source. One element that exhibits this behavior is boron. In practice this behavior is not important for Gate applications.

Materials

In Gate, materials are defined as combinations of elements, and are an important parameter that Gate uses for all of the particle interactions that take place during a simulation. These combinations of elements require defining four additional parameters. These are the material's name, density, constituent element(s), and their individual abundances.

The composition of elements within a material can be defined in two different ways. If the material is a chemical compound then its relative amounts of elements are specified by the number of atoms in the chemical formula of the compound. For example, methane $\text{CH}_4$ would be defined as having one carbon atom and four hydrogen atoms. If the material is better described as a mixture, such as 304-stainless steel, then the relative combinations of the elements are given by mass fraction. In the case of 304-stainless steel, the various mass fractions are given as 0.695 Iron, 0.190 Chromium, 0.095 Nickel, and 0.020 Manganese. Note that the mass fractions from the elements must all sum to one.

Densities of materials often vary greatly between different sources and must be carefully selected for the specific application in mind. Units of density must also be defined. These are typically given in g/cm$^3$ but can be given in more convenient units for extreme cases. For example, a vacuum's density may be expressed in units of mg/cm$^3$.

Modifying the Gate material database

New element

Defining a new element is a simple and straightforward process. Simply open the GateMaterials.db file with the text editor of your choice. At the top of the file is the header named [Elements] and somewhere in the middle of the file is another header named [Materials]. All element definitions required by the application must be included between these two headers. The format for entering an element is given by the elements name, symbol, atomic number, and molar mass. Below is an example:

```
Element Example GateMaterials.db:

[Elements]
Hydrogen:   S= H   ; Z=  1. ; A=   1.01  g/mole
Helium:     S= He  ; Z=  2. ; A=   4.003 g/mole
Lithium:    S= Li  ; Z=  3. ; A=   6.941 g/mole
Beryllium:  S= Be  ; Z=  4. ; A=   9.012 g/mole
Boron:      S= B   ; Z=  5. ; A=  10.811 g/mole
Carbon:     S= C   ; Z=  6. ; A=  12.01  g/mole
```

In this example the name of the element is given first and is followed by a colon. Next, the standard symbol for the element is given by $S=$symbolic name followed by a semi-colon. The atomic number and molar mass follow the symbolic name given by $Z=$atomic number with a semi-colon and by $A=$molar mass units for the molar mass and its units.

New material
Materials are defined in a similar manner to elements but contain some additional parameters to account for their density and composition. Defining density is straightforward and performed the same way for all materials. However, material compositions require different definitions depending upon their form. These compositional forms are pure substances, chemical compounds, and mixtures of elements.

To add or modify a material in the material database, the GateMaterials.db file should be open using a text editor. The new entry should be inserted below the header named Materials. All material definitions required by the application must be included below this second header. Material definitions require several lines. The first line specifies their name, density, number of components, and an optional parameter describing the materials state (solid, liquid, or gas). The second and following lines specify the individual components and their relative abundances that make up this material.

The compositional forms of materials that Gate uses are pure substances, chemical compounds, mixtures of elements, and mixtures of materials. Gate defines each of these cases slightly differently and each will be dealt with separately below. In every case, the elements being used in a material definition must be previously defined as elements.

**Elements as materials**

Substances made of a pure element are the easiest materials to define. On the first line, enter the name of the material (the name of the material can be the same as that of the element), its density, its number of constituents (which is one in this case), and optionally its state (solid, liquid, or gas). The default state is gaseous. On the second line enter the element that it is composed of and the number of atoms of that element (in the case of an element as a material this number is 1). For example:

```plaintext
Elements as materials example GateMaterials.db:

[Materials]
Vacuum: d=0.000001 mg/cm3 ; n=1
   +el: name=Hydrogen ; n=1
Aluminium: d=1.350 g/cm3 ; n=1 ; state=solid
   +el: name=auto ; n=1
Uranium: d=18.90 g/cm3 ; n=1 ; state=solid
   +el: name=auto ; n=1
```

On the first line the density (with units) is defined by \(d=\text{material density units}\) and is separated by a semi-colon from the number of constituents in the material defined by \(n=\text{number of elements}\). If the optional material form parameter is used it is also separated by a semi-colon. The available forms are gas, liquid, and solid. On the second line the individual elements and their abundances are defined by \(+el: \text{name=of the element } ; n=\text{number of atoms}\). If the name of the element and the material are the same, the element name can be defined by \(+el: \text{name=auto}\) command.

**Compounds as materials**

Chemical compounds are defined based upon the elements they are made of and their chemical formula. The first line is identical to the first line of a pure substance except that the number of constituent elements is now greater than one. On the second and subsequent lines, the individual elements and their abundances are defined by \(+el: \text{name=of the element } ; n=\text{number of atoms}\).

For example:
## Compounds as materials example GateMaterials.db:

```
[Materials]
NaI: d=3.67 g/cm3; n=2; state=solid
  +el: name=Sodium ; n=1
  +el: name=Iodine ; n=1
PWO: d=8.28 g/cm3; n=3 ; state=Solid
  +el: name=Lead; n=1
  +el: name=Tungsten; n=1
  +el: name=Oxygen; n=4
```

## Mixtures as materials

Mixture of elements are defined by indicating the mass fraction of the elements that make up the mixture. The first line of this definition is identical to the first line of the definition of a chemical compound. On the second and subsequent lines, the individual elements and their mass fractions are defined by `+el: name=name of element; f=mass fraction`.

In the case of material mixtures, the sum of the mass fractions should be one. For example:

```
[Materials]
Lung:  d=0.26 g/cm3 ; n=9
  +el: name=Hydrogen  ; f=0.103
  +el: name=Carbon    ; f=0.105
  +el: name=Nitrogen  ; f=0.031
  +el: name=Oxygen    ; f=0.749
  +el: name=Hydrogen  ; f=0.002
  +el: name=Sodium    ; f=0.002
  +el: name=Phosphor  ; f=0.003
  +el: name=Sulfur    ; f=0.003
  +el: name=Chlorine  ; f=0.003
  +el: name=Potassium ; f=0.002
SS304:  d=7.92 g/cm3 ; n=4 ; state=solid
  +el: name=Iron      ; f=0.695
  +el: name=Chromium  ; f=0.190
  +el: name=Nickel    ; f=0.095
  +el: name=Manganese ; f=0.020
```

## Mixtures of materials as materials

Another way material can be defined is as mixtures of other materials and elements. As an example:

```
[Materials]
Aerogel:  d=0.200 g/cm3 ; n=3
  +mat: name=SiO2     ; f=0.625
  +mat: name=Water    ; f=0.374
  +el: name=Carbon    ; f=0.001
```

In this example, the material, Aerogel, is defined to be made up of two materials, silicon dioxide and water, and one element, carbon. Mass fractions of the silicon dioxide, water, and carbon are given to specify the atom densities of the material when related to the density of the Aerogel. When specifying materials rather
than elements the +mat: name=identifier must be used.

**Ionization potential**

The ionization potential is the energy required to remove an electron to an atom or a molecule. By default, the ionization potential is calculated thanks to the Bragg’s additivity rule. It is possible to define the ionization potential of each material defined in gate. For example:

```
/gate/geometry/setIonisationPotential Water 75 eV
```


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Users Guide V7.0: Setting up the physics

From GATE collaborative documentation wiki

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New physics list mechanism

WARNING: big change from Gate V7.0

Up to now, physic lists were set in GATE according to macro files containing list of command to add physical processes. From GATE Version 7.0, we highly recommend to switch to the "physic list builder" mechanism. The physic-list builders are provided by the Geant4 community. To use a builder, use the
"addPhyslicsList" command as follow:

```
//gate/physics/addPhyslicsList QGSP_BERT_EMV
```

The name of the builder is one of the following for the electromagnetic physic-lists:

- emstandard
- emstandard_opt1
- emstandard_opt2
- emstandard_opt3
- emlivermore
- emlivermore_polar
- empenelope

And for the hadronic physics list builders:

- QGSP
- QGSP_EMV
- QGSC
- QGSC_EMV
- QGSP_EFLOW
- QGSP_BERT
- QGSP_BERT_EMV
- QGSP_BERT_HP
- QGSP_BERT_TRV
- QGSP_BIC
- QGSP_BIC_HP
- QGSP_NEQ
- QGSP_EMV_NQE
- QGSP_BERT_NQE
- QGSP_INCLXX
- FTFP_BERT
- FTFP
- FTFP_EMV
- ...


The production cuts special cuts can still be used as exemplified below:

```
//gate/physics/Gamma/SetCutInRegion world 10 mm
//gate/physics/Electron/SetCutInRegion world 10 mm
//gate/physics/Positron/SetCutInRegion world 10 mm
//gate/physics/Gamma/SetCutInRegion patient 1 mm
//gate/physics/Electron/SetCutInRegion patient 1 mm
//gate/physics/Positron/SetCutInRegion patient 1 mm
```
Physics list selection

Processes have three possible states: "Available", "Enabled" and "Initialized". In a simulation, GATE uses initialized processes. By default, all processes are only "Available". To initialize processes you want to use, you have to choose them in the list of available processes and put them in the list of "Enabled" processes with the `addProcess/removeProcess` commands. Then, you have to define models, data sets or energy range if it is necessary. Enabled processes are initialized when the command `/gate/run/initialize` is called. After initialization, physics list cannot be modified. Processes must be initialized before the source definition.

To obtain the list of processes available, enabled or initialized:

```plaintext
/gate/physics/processList [State] [Particle]
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>State</th>
<th>Available, Enabled, Initialized</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter type</td>
<td>s</td>
<td>True</td>
</tr>
<tr>
<td>Default value</td>
<td>Available</td>
<td></td>
</tr>
<tr>
<td>Candidates</td>
<td>Available, Enabled, Initialized</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Particle</th>
<th>All</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter type</td>
<td>s</td>
<td>True</td>
</tr>
<tr>
<td>Default value</td>
<td>All</td>
<td></td>
</tr>
</tbody>
</table>

User can print the list of initialized processes for each particles and the list of enabled processes with all informations (particles, models, data sets and energy range):

```plaintext
/gate/physics/print [File name]
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>File Name</th>
<th>False</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter type</td>
<td>s</td>
<td></td>
</tr>
</tbody>
</table>

Particles

To designate a particle in a macro command for physics, you have to use the Geant4 name of the particle. The command `/particle/list` gives the list of particles available in GATE. Note that there are 5 particles defined for nuclei: "deuteron", "triton", "alpha", "He3" and a generic particle for all ions called "GenericIon". For more information about particles in Geant4, see Geant4 user support[1].

User can also use a particle group name. For example, "Charged" designates all charged particles or “EM” designates gamma, e+ and e-. Here, the list of particle groups available in GATE:

- Default : defaults particles defined for a process (see the "processList" command before for more
informations)
- EM : e+, e-, gamma
- Charged : all charged particles

< ! > Particle name in non-physics macro command could be slightly different in particular for ions!

**Adding a process**

To add a process to the list of "Enabled" processes for a particle or a group of particles:

```
//gate/physics/addProcess [Process] [Particle]
```

Parameter : Process  
Parameter type : s  
Omittable : False

Parameter : Particle  
Parameter type : s  
Omittable : True  
Default value : Default

**Removing a process**

To remove a process of the "Enabled" processes list:

```
//gate/physics/removeProcess [Process] [Particle]
```

Parameter : Process  
Parameter type : s  
Omittable : False

Parameter : Particle  
Parameter type : s  
Omittable : True  
Default value : Default

**Models and Data sets**

Some processes have several models or several data sets available. To known if you can choose a model or a data set for a process, use the commands:

```
//gate/physics/processes/[Process Name]/modelList [Particle]
//gate/physics/processes/[Process Name]/dataSetList [Particle]
```

Parameter : Particle  
Parameter type : s  
Omittable : True  
Default value : Default

To select or unselect a model or a data set, use the command:

```
//gate/physics/processes/[Process Name]/setModel [Model Name] [Particle]
//gate/physics/processes/[Process Name]/unSetModel [Model Name] [Particle]
```
Models can be selected for an energy range or for specific materials or elements. To do this, use commands:

```
/gate/physics/processes/[Process Name]/[Model Name]/setEmax [Value] [Unit] [Particle] [Option]
/gate/physics/processes/[Process Name]/[Model Name]/setEmin [Value] [Unit] [Particle] [Option]
```

The parameter "Option" allow to define material or element for this model (see chapter on detector construction for more informations on materials and elements). For example:

```
/gate/physics/processes/IonInelastic/G4BinaryLightIonReaction/setEmin 100 keV
/gate/physics/processes/IonInelastic/G4BinaryLightIonReaction/setEmin 200 keV GenericIon Water
```

A command allows to clear energy ranges defined for a model:

```
/gate/physics/[Process Name]/[Model Name]/clearEnergyRange [Particle]
```

---

**Physics processes**

**Electromagnetic processes**

Electromagnetic processes are used to simulate the electromagnetic interaction of particles with matter. The mean free path of a process, $\lambda$, also called the interaction length, can be given in terms of the total cross section:

$$\lambda(E) = \left( \sum_i \left[ n_i \cdot \sigma(Z_i, E) \right] \right)^{-1}$$

where $\sigma(Z_i, E)$ is the cross section of the process for atom $i$ composing the material. Cross-sections per atom and mean free path values are tabulated during initialization.

In Geant4, three models are available for electromagnetic processes:

- Standard processes are effective between 1 keV and 100 TeV
- Low energy processes are effective between 250 eV and 100 GeV (there is also the LivermorePolarizedModel for polarized gamma)
- Penelope processes are effective between 250 eV and 1 GeV

Models and cross-sections are based on the theoretical calculations and on exploitation of evaluated data. For the standard processes based on data, models and cross-sections rely on parameterizations of these data. Because atomic shell structure is more important in most cases at low energies, the low energy processes make direct use shell cross section data. The data used for the determination of cross-sections and for sampling of the final state are extracted from a set of freely distributed evaluated data libraries:

- EPDL97 (Evaluated Photons Data Library)
- EEDL (Evaluated Electrons Data Library)
- EADL (Evaluated Atomic Data Library)
- stopping power data
- binding energy values based on data of Scofield.

The Penelope models have been specifically developed for Monte Carlo simulation and great care was given to the low energy description (i.e. atomic effects, etc.). These processes are the Geant4 implementation of the physics models developed for the PENELOPE code (PENetration and Energy LOss of Positrons and Electrons), version 2001.

> For the low energy processes, the download of G4EMLOW data files is required.

> Positron have no low energy process.

> Since Geant4 9.3, users can mixed electromagnetic models.

**Photoelectric effect**

The photoelectric effect is the absorption of a photon by an atomic electron with the ejection of this electron from the atom. The energy of the outgoing electron is:
\[ E = h\nu - L \]

where \( L \) is the binding energy of the electron. Since a free electron cannot absorb a photon and also conserve momentum, the photoelectric effect always occurs on bound electrons while the nucleus absorbs the recoil momentum. The cross-section calculation is complex due to the combinaison of the electron Dirac wave functions. It is simulated by using a parameterized photon absorption cross section to determine the mean free path, atomic shell data to determine the energy of the ejected electron, and the K-shell angular distribution to sample the direction of the electron.

The cross-section depends on the atomic number \( Z \) of the material. The photoelectric process is favored by high \( Z \) materials. In the current implementation the relaxation of the atom is not simulated, but instead is counted as a local energy deposit. For low energy process, the deexcitation of the atom is simulated.

To add photoelectric effect in GATE:

```
/gate/physics/addProcess PhotoElectric gamma
```

Note that the two following macro are equivalent because the default particle for the photoelectric process is the gamma only:

```
/gate/physics/addProcess PhotoElectric
/gate/physics/addProcess PhotoElectric Default
/gate/physics/addProcess PhotoElectric gamma
```

Then, choose the model:

```
/gate/physics/processes/PhotoElectric/setModel StandardModel
```

or

```
/gate/physics/processes/PhotoElectric/setModel LivermoreModel
```

or

```
/gate/physics/processes/PhotoElectric/setModel LivermorePolarizedModel
```

or

```
/gate/physics/processes/PhotoElectric/setModel PenelopeModel
```

### Compton scattering

Compton process describes the photon scattering by free electrons. Although electrons are bound in matter, the electron can be considered as essentially free for photons of energy much greater than the binding energy.
of the electron.

In the simulation, an empirical cross section formula is used, which reproduces the cross section data down to 10 keV. The final state is generated following the Klein-Nishina formula. For low energy incident photons, the simulation of the Compton scattering process is performed according to the same procedure used for the standard Compton scattering simulation, with the addition that Hubbel’s atomic form factor (or scattering function) is taken into account. The angular and energy distribution of the incoherently scattered photon is then given by the product of the Klein-Nishina formula and the scattering function.

To add Compton scattering in GATE:

```
gate/physics/addProcess Compton gamma
```

Note that the two following macro are equivalent because the default particle for the Compton process is only the gamma:

```
gate/physics/addProcess Compton
```
```
gate/physics/addProcess Compton Default
```
```
gate/physics/addProcess Compton gamma
```

Then, choose the model:

```
gate/physics/processes/Compton/setModel StandardModel
```

or

```
gate/physics/processes/Compton/setModel LivermoreModel
```

or

```
gate/physics/processes/Compton/setModel LivermorePolarizedModel
```

or

```
gate/physics/processes/Compton/setModel PenelopeModel
```

Rayleigh scattering

Thomson and Rayleigh scattering are linked to Compton scattering. Thomson scattering is similar to Compton scattering in the classical limit i.e. the scattering of photons by free electrons. For Rayleigh scattering, all the electrons of the atom contribute in a coherent way (this process is also called coherent scattering).
For these processes, no energy is transferred to the target. The direction of the photon is the only modified parameter. Atoms are not excited or ionized. At high energies, the cross-sections of Thomson and Rayleigh scattering are very small and are neglected. For these reasons, the Rayleigh process is defined only for low energy and Penelope models:

```plaintext
/gate/physics/addProcess RayleighScattering gamma
/gate/physics/processes/RayleighScattering/setModel LivermoreModel
/gate/physics/processes/RayleighScattering/setModel LivermorePolarizedModel
/gate/physics/processes/RayleighScattering/setModel PenelopeModel
```

**Pair production**

The process of pair production describes the transformation of a photon into an electron-positron pair. In order to conserve momentum, this can only occur in the presence of a third body, usually a nucleus. Moreover, the minimum energy required to create a pair is equal to the sum of the electron mass and positron mass (1.022 MeV).

To add pair production process in GATE:

```plaintext
/gate/physics/addProcess GammaConversion
```

Then, choose the model:

```plaintext
/gate/physics/processes/GammaConversion/setModel StandardModel
```

or

```plaintext
/gate/physics/processes/GammaConversion/setModel LivermoreModel
```

or

```plaintext
/gate/physics/processes/GammaConversion/setModel LivermorePolarizedModel
```

or

```plaintext
/gate/physics/processes/GammaConversion/setModel PenelopeModel
```

**Ionization**

A charged particle passing through matter loses energy due to inelastic collision with atomic electrons of the material. Lost energy is transferred to the atom causing ionization or excitation. The ionization energy loss
is calculated using the Bethe-Bloch formula. The particle energy loss $E$ is divided into continuous energy loss and production of secondary electrons. The production threshold is defined as the minimum energy $E_{cut}$ above which secondary particles will be produced and tracked. When $E < E_{cut}$, $E$ is included into the continuous energy loss and when $E > E_{cut}$, secondary electrons are produced. Energy loss due to excitation is included into continuous energy loss.

The mean excitation potential $I$ is the main parameter of the Bethe-Bloch formula. This quantity can be defined by user for each material (see chapter about geometry and materials).

There are three processes to handle ionization:

- for electron and positron
  
  ```
  /gate/physics/addProcess ElectronIonisation e+
  /gate/physics/addProcess ElectronIonisation e-
  ```

- for hadrons and ions
  
  ```
  /gate/physics/addProcess HadronIonisation [Particle Name]
  ```

- for ions
  
  ```
  /gate/physics/addProcess IonIonisation [Particle Name]
  ```

The electron ionization process has three models available:

```
/gate/physics/processes/ElectronIonisation/setModel StandardModel e+
/gate/physics/processes/ElectronIonisation/setModel StandardModel e-
/gate/physics/processes/ElectronIonisation/setModel LivermoreModel e-
/gate/physics/processes/ElectronIonisation/setModel PenelopeModel e+
/gate/physics/processes/ElectronIonisation/setModel PenelopeModel e-
```

The new Geant4 model selection has not yet implemented for hadron ionization process in GATE V6.2. So users have to the old process/model selection.

The low energy model have a specific process for ionization of hadrons, ions, muons and taus:

```
/gate/physics/addProcess LowEnergyHadronIonisation
```

For the energy range between 1 keV and 2 MeV, this process have several models for the parametrization of electronic and nuclear stopping power:

```
/gate/physics/processes/LowEnergyHadronIonisation/setModel Elec_Ziegler1977p proton
```

Electronic stopping power models:

- Elec_ICRU_R49p (default)
Elec_ICRU_R49He
Elec_Ziegler1977p
Elec_Ziegler1977He
Elec_Ziegler1985p
Elec_SRIM2000p

Nuclear stopping power models:

- Nuclear_ICRU_R49 (default)
- Nuclear_Ziegler1977
- Nuclear_Ziegler1985

It is possible to enable/disable nuclear stopping power for all hadrons and ions ionization processes:

```xml
/gate/physics/processes/[Process name]/setNuclearStoppingOn
/gate/physics/processes/[Process name]/setNuclearStoppingOff
```

Example:

```xml
/gate/physics/processes/HadronIonisation/setNuclearStoppingOff
```

**Bremsstrahlung**

Bremsstrahlung process is the production of an electromagnetic radiation by a charged particle accelerated in the field of another charged particle, such as a nucleus. The cross-section of bremsstrahlung is inversely proportional to the mass squared. Thus this process is most important for electron and positron than other charge particles. As for ionization process, above a given threshold energy the energy loss is simulated by the explicit production of photons. Below the threshold, emission of soft photons is treated as a continuous energy loss. The bremsstrahlung energy spectrum is continuous.

```xml
/gate/physics/addProcess Bremsstrahlung e+
/gate/physics/addProcess Bremsstrahlung e-
```

The electron ionization process has three models available:

```xml
/gate/physics/processes/Bremsstrahlung/setModel StandardModel e+
/gate/physics/processes/Bremsstrahlung/setModel StandardModel e-
/gate/physics/processes/Bremsstrahlung/setModel LivermoreModel e-
/gate/physics/processes/Bremsstrahlung/setModel PenelopeModel e+
/gate/physics/processes/Bremsstrahlung/setModel PenelopeModel e-
```

**Positron and electron annihilation**

In Geant4, the process which simulated the in-flight annihilation of a positron with an atomic electron is attached to positron. As is usually done in shower programs, it is assumed here that the atomic electron is
initially free and at rest. Also, annihilation processes producing one, or three or more, photons are ignored because these processes are negligible compared to the annihilation into two photons.

```
/gate/physics/addProcess G4PositronAnnihilation e+
```

Then, choose the model:

```
/gate/physics/processes/G4PositronAnnihilation/setModel StandardModel
```

or

```
/gate/physics/processes/G4PositronAnnihilation/setModel PenelopeModel
```

An improvement of the positron-electron annihilation has been developed to take into account the \( \gamma\gamma \) non-colinearity. The mean value of the angle distribution is \( \approx 0.5^0 \) (do not need model selection).

```
/gate/physics/addProcess PositronAnnihilation e+
```

**Single and multiple Scattering**

In addition to inelastic collisions with atomic electrons, charged particles passing through matter also suffer repeated elastic Coulomb scatterings from nuclei. Elastic cross section is huge when particle energy decreases, so multiple scattering approach (MSC) should be introduced in order to have acceptable CPU performance of the simulation. The MSC model used in GEANT4 belongs to the class of condensed algorithm in which the global effects of the collisions are simulated at the end of a track segment (step). The global effects generally computed in these codes are the net displacement, energy loss, and change of direction of the charged particle. The model is based on Lewis' MSC theory and uses model functions to determine the angular and spatial distributions after a step. The functions have been chosen in such a way as to give the same moments of the (angular and spatial) distributions as the Lewis theory. Two processes are available for multiple scattering. These processes are similar but they allow to define some sets of options for group of particles (electron/positron and hadron). The new Geant4 model selection has not yet implemented for multiple scattering process in GATE V6.2. So users have to the old process/model selection. The old MultipleScattering is now deprecated.

```
/gate/physics/addProcess eMultipleScattering e-
/gate/physics/addProcess eMultipleScattering e+
```

```
/gate/physics/addProcess hMultipleScattering proton
/gate/physics/addProcess hMultipleScattering alpha
/gate/physics/addProcess hMultipleScattering GenericIon
```
Single elastic scattering process is an alternative to the multiple scattering process. The advantage of the single scattering process is in possibility of usage of theory based cross sections, in contrary to the Geant4 multiple scattering model, which uses a number of phenomenological approximations on top of Lewis theory. Because each of elastic collisions are simulated the simulation CPU time of charged particles significantly increasing in comparison with the multiple scattering approach.

```
/gate/physics/addProcess SingleScattering ...
```

**Muon electromagnetic processes**

Muons have their own electromagnetic processes:

- **Ionization**
  ```
  /gate/physics/addProcess MuIonisation mu+
  /gate/physics/addProcess MuIonisation mu-
  ```

- **Bremsstrahlung**
  ```
  /gate/physics/addProcess MuBremsstrahlung mu+
  /gate/physics/addProcess MuBremsstrahlung mu-
  ```

- **Direct production of (e+, e-) pairs by mu+ and mu-**
  ```
  /gate/physics/addProcess MuPairProduction mu+
  /gate/physics/addProcess MuPairProduction mu-
  ```

For ionization, the low energy model can handle muons.

**Electromagnetic options**

< ! > This part is recommended for advanced users only!

< ! > Valid only from Geant4 version 9.2 - Gate V6.0 & V6.0.p01

< ! > In Geant4 version 9.2, options are available only for standard processes - Gate V6.0 & V6.0.p01

< ! > In case of using Gate V6.2, users must compile with Geant4 version 9.5 patch 01 - This Gate release is NOT compatible with previous Geant4 version

Options are available for steering the standard electromagnetic processes. Some options modify all electromagnetic processes initialized in the simulation (Global options). Some options have to be defined for each processes.

**Global options:**
The following options manage the DEDX, mean free path and cross sections tables. User can defined the table range (default 0.1 keV - 100 TeV):

```
/gate/physics/setEMin 0.1 keV
/gate/physics/setEMax 10 GeV
```

the number of bins of the DEDX table (default = 84):

```
/gate/physics/setDEDXBinning 500
```

and the number of bins of the mean free path table (default = 84):

```
/gate/physics/setLambdaBinning 500
```

Using cubic spline interpolation of DEDX and cross section tables, better interpolation was found to increase stability when varying transport parameters, such as cuts, of energy deposition of hadrons. Cubic spline interpolation is enabled by default. If the option was disable, the old linear interpolation is used:

```
/gate/physics/setSplineFlag true
/gate/physics/setSplineFlag false
```

**Step function**

Continuous energy loss imposes a limit on the step size because of the energy dependence of the cross sections. It is generally assumed in MC programs that the particle cross sections are approximately constant along a step, i.e. the step size should be small enough that the change in cross section, from the beginning of the step to the end, is also small. In principle one must use very small steps in order to insure an accurate simulation, however the computing time increases as the step size decreases. A good compromise is to limit the step size by not allowing the stopping range of the particle to decrease by more than a value \( [\text{Ratio}] \) during the step \( (\text{Ratio} = \Delta \text{range}/\text{range}) \). This condition works well for particles with kinetic energies > 1 MeV, but for lower energies it gives very short step sizes.

To cure this problem a lower limit on the step size was introduced \( (\text{Final range}) \). The step size limit varies smoothly with decreasing energy from the value \( \text{Ratio} \) to the lowest possible value range cut \( \text{Final range} \). By default for electron, \( \text{Ratio} = 0.2 \) and \( \text{Final range} = 0.1 \) mm.

```
/gate/physics/processes/[Process name]/SetStepFunction [Particle] [Ratio] [Final range] [Unit]
```

Parameter : [Particle]
Parameter type : s
Omittable : False

Parameter : [Ratio] (step/range)
Parameter type : d
Omittable : False

Parameter : [Final range]
Parameter type : d
Omittable : False
Parameter : [Unit]  
Parameter type : s  
Omittable : False

Example:

```
/gate/physics/processes/ElectronIonisation/setStepFunction e+ 0.01 1 mm
```

**Linear loss limit**

This cut is another approach to limit the step size. In a step, the energy loss by a particle with a kinetic energy $E$ cannot exceed a value $E_{cut}$ such as $E_{cut}/E < [Ratio]$. By default for the ionization of an electron, the limit is 0.01.

```
/gate/physics/processes/IonIonisation/setLinearLossLimit [Particle] [Ratio]
```

Parameter : Particle or Group of particles  
Parameter type : s  
Omittable : False

Parameter : Limit  
Parameter type : d  
Omittable : False

Example:

```
/gate/physics/processes/HadronIonisation/setLinearLossLimit proton 0.0001
```

**Geometrical step limit type**

This option allows to choose the transport algorithm for the multiple scattering process near to boundary. The two options proposed are *safety* and *distanceToBoundary*. For instance, the *distanceToBoundary* algorithm limit the step size near to geometrical boundaries: only single scattering is applied very close to the boundaries.

```
/gate/physics/processes/MultipleScattering/setGeometricalStepLimiterType [Particle] [Limit type]
```

Parameter : Particle or Group of particles  
Parameter type : s  
Omittable : False

Parameter : Limit type  
Parameter type : s  
Omittable : False

Example:

```
/gate/physics/processes/MultipleScattering/setGeometricalStepLimiterType proton distanceToBoundary
```
Hadronic processes

Hadronic processes described the interactions between incident hadrons/ions and the target nuclei. We also include decays of hadron and nuclei in this part. There is no strict frontier between nucleon-nucleon collision processes but we can distinguish four main process types in function of the energy and the impact parameter. At low energies, collisions lead to incomplete fusion in central collisions and to elastic scattering or inelastic scattering in peripheral collisions. At higher energies, there is fragmentation into several lighter fragments or nucleons for central collisions and into participant and the spectator regions for peripheral collisions. In Geant4, fusion, inelastic scattering and fragmentation processed are included in the inelastic process type.

Inelastic processes had three main steps:

- cascade: the incident particle interacts strongly with the target and produces secondary particles
- preequilibrium (thermalization process): the excited target nucleus switches into equilibrated state by emitting excitons and light nuclei.
- de-excitation: the equilibrated nuclear residues evaporates into nucleons/light nuclei or breaks up into several fragments.

Each hadronic process may have one or more data sets associated with it. The term "data set" is meant, in a broad sense, to be an object that encapsulates methods and data for calculating total cross sections for a given process. The methods and data may take many forms, from a simple equation using a few hard-wired numbers to a sophisticated parameterization using large data tables. For the evaluation of cross sections, the list has a LIFO (Last In First Out) priority, meaning that data sets added later to the list will have priority over those added earlier to the list.

The final state is produced using models coupled to processes. In Geant4, any model can be run together with any other model and the ranges of applicability for the different models can be steered. This way, highly specialized models (valid only for one material and particle, and applicable only in a very restricted energy range) can be used in the same application, together with more general code, in a coherent fashion. Each model has an intrinsic range of applicability, and the model chosen for a simulation depends very much on the use-case. Three types of hadronic models have been implemented: parametrization driven models, data driven models, and theory driven models.

Most of hadronic processes need an explicit choice of models while a lots of processes have a default data set.

**Main Geant4 models**

**LHEP**

LHEP is the model used by default. It is based on code GHEISHA developed since 1978 by H. Fesefeldt to simulate hadron-nucleus interactions. The low energy part is valid from few hundred MeV to 20 GeV. The model is based on the principle of the intranuclear cascade and only the first hadron-nucleus interaction is simulated in detail. Other interactions in the nucleus are simulated by generating additional hadrons, simply treated as secondary particles can themselves generate their own intranuclear cascade. LHEP is a fully parameterized model but the physical meaning of the large number of adjustable parameters is sometimes unclear. Its main assets are the broad energy range covered, good reproduction of average values of distributions and computation times.

Example: G4LCapture, G4LENeutronInelastic, G4LFission, G4LCapture, G4LEProtonInelastic,
G4LEPionMinusInelastic, G4LEPionPlusInelastic...

*Bertini cascade*

The intranuclear cascade model of Bertini has been developed by H.W. Bertini in 1963. This code includes the intranuclear cascade model of Bertini, a pre-equilibrium model, a simple model of nucleus explosion, a model of fission and evaporation model.

*Binary cascade*

The Geant4 Binary Cascade is an intranuclear cascade propagating primary and secondary particles in a nucleus. The energy range and type of projectile covered are the same as Bertini model. From a theoretical point of view, this model is much more evolved taking into account a large number of resonances and fully modeled in three dimensions. Interactions are between a primary or secondary particle and an individual nucleon of the nucleus, leading to the name Binary Cascade. Cross section data are used to select collisions. Where available, experimental cross sections are used by the simulation. Propagating of particles is the nuclear field is done by numerically solving the equation of motion. The cascade terminates when the average and maximum energy of secondaries is below threshold. The remaining fragments are treated by precompound and de-excitation models.

*QMD*

*Pre compound*

**Elastic scattering**

In elastic scattering, the projectile and the target particles do not changed during the collision and no other particles are produced. Two processes are available for hadrons and ions. The first one is the old elastic process.

```
/gate/physics/addProcess HadronElastic Default
/gate/physics/processes/HadronElastic/setModel G4LElastic Default
```

or

```
/gate/physics/addProcess HadronElastic
/gate/physics/processes/HadronElastic/setModel G4LElastic
```

or

```
/gate/physics/addProcess HadronElastic proton
/gate/physics/processes/HadronElastic/setModel G4LElastic proton
/gate/physics/addProcess HadronElastic alpha
/gate/physics/processes/HadronElastic/setModel G4LElastic alpha
```

This process has a default dataset (G4HadronElasticDataSet) and 6 models.
The alternative process is an improvement of the first process. It is supposed to be a good mix between the models of the first process:

```
/gate/physics/addProcess HadronElastic
/gate/physics/processes/HadronElastic/setModel G4HadronElastic
/gate/physics/processes/HadronElastic/setDataSet G4HadronElasticDataSet
```

For this process, there is only one model and a main dataset (G4HadronElasticDataSet). An other dataset for low energy neutrons is also available (G4NeutronHPElasticData).

**Inelastic process for proton**

This process manage inelastic interaction of proton with matter. For example, the selection of two models with energy range for proton inelastic process (the only default particle is the proton):

```
/gate/physics/addProcess ProtonInelastic
/gate/physics/processes/ProtonInelastic/setModel G4BinaryCascade
/gate/physics/processes/ProtonInelastic/G4BinaryCascade/setEmin 170 MeV
/gate/physics/processes/ProtonInelastic/G4BinaryCascade/setEmax 500 GeV
/gate/physics/processes/ProtonInelastic/setModel PreCompound
/gate/physics/processes/ProtonInelastic/PreCompound/setEmin 0 MeV
/gate/physics/processes/ProtonInelastic/PreCompound/setEmax 170 MeV
```

This process has a default dataset (G4HadronInelasticDataSet) and an alternative dataset (G4ProtonInelasticCrossSection). There are 6 models available:

- G4LEProtonInelastic
- G4BertiniCascade
- G4BinaryCascade
- PreCompound
- G4QMDReaction

**Inelastic process for ion**

This process manage inelastic interaction of ions with matter. This process is valid for GenericIon, alpha, deuteron and triton. For example, a complete selection of models and data set for ions:

```
/gate/physics/addProcess IonInelastic Default
/gate/physics/processes/IonInelastic/setModel G4BinaryLightIonReaction Default
/gate/physics/processes/IonInelastic/setModel G4DEuteronInelastic deuteron
/gate/physics/processes/IonInelastic/setModel G4LETritonInelastic triton
/gate/physics/processes/IonInelastic/setModel G4LEAlphaInelastic alpha
/gate/physics/processes/IonInelastic/G4BinaryLightIonReaction/setEmin 80 MeV deuteron
/gate/physics/processes/IonInelastic/G4BinaryLightIonReaction/setEmax 20 GeV deuteron
/gate/physics/processes/IonInelastic/G4BinaryLightIonReaction/setEmin 80 MeV triton
```

---

The IonInelastic process includes the G4IonInelasticProcess for GenericIon, the G4DeuteronInelasticProcess for deuteron, the G4TritonInelasticProcess for triton and the G4AlphaInelasticProcess for alpha. The G4QMDReaction model and the G4BinaryLightIonReaction model are available for all ions. For GenericIon, one additional model (G4BinaryLightIonReaction) and 5 datasets are available:

- G4TripathiCrossSection
- G4IonsKoxCrossSection
- G4IonsShenCrossSection
- G4IonsSihverCrossSection
- G4TripathiLightCrossSection

Alpha, deuteron and triton have a default data set (G4HadronInelasticDataSet) and a alternative dataset (G4TripathiLightCrossSection). There are also specific models for each particle: G4LEDeuteronInelastic, G4LETritonInelastic, G4LEAlphaInelastic.

### Pions

The inelastic interaction of pi+ and pi- with matter is handled by PionPlusInelastic and PionMinusInelastic processes. These processes have two specific models (G4LEPionMinusInelastic - G4LEPionPlusInelastic) and three common models:

- Bertini Cascade
- Binary Cascade
- Leading Particle Bias

The default dataset is G4HadronInelasticDataSet. There is an alternative dataset: G4PiNuclearCrossSection.

### Neutrons

The interactions of neutrons at low energies are split into four parts. We consider radiative capture, elastic scattering, fission, and inelastic scattering as separate processes. Each processes have standard models and datasets like others particles. In additions, some "high precision" models and datasets are provided for low energy interactions. The high precision neutron models depend on an evaluated neutron data library.
(G4NDL) for cross sections, angular distributions and final state information. G4NDL data comes largely from the ENDF/B-VI library.

< ! > For the low energy processes, the download of G4NDL data files is required.

Radiative Capture

The G4LCapture model generates the final state for neutron capture. The G4NeutronHPCapture model generates the final state for neutron capture using the high precision neutron model. The G4NeutronHPorLCapture model generates the final state for neutron capture using the high precision neutron model when sufficient high precision data is available for the selected element or isotope. When there is insufficient data, the neutron is captured using the less precise Low Energy Parameterized model.

The G4HadronCaptureDataSet is the default dataset for this process. The alternative high precision dataset is G4NeutronHPCaptureData.

```
/gate/physics/addProcess NeutronCapture
/gate/physics/processes/NeutronCapture/setModel G4LCapture
```

Inelastic scattering

The G4NeutronInelasticProcess is similar than proton inelastic and ion inelastic processes. In addition to the standard models (G4LENeutronInelastic, G4BertiniCascade, G4BinaryCascade, PreCompound, LeadingParticleBias), two models using the high precision data are available. The G4NeutronHPIInelastic model generates the final state for inelastic neutron scattering. The G4NeutronHPorLEInelastic model generates the final state for inelastic neutron scattering using the high precision neutron model when sufficient high precision data is available for the selected element or isotope. When there is insufficient data, the neutron is scattered inelastically using the less precise Low Energy Parameterized model (G4LENeutronInelastic).

The G4HadronInelasticDataSet is the default dataset for this process. An alternative dataset is the G4NeutronInelasticCrossSection. The high precision dataset is G4NeutronHPIInelasticData.

```
/gate/physics/addProcess NeutronInelastic
/gate/physics/processes/NeutronInelastic/setModel PreCompound
```

Fission

The G4LFission model generates the final state for fission. The G4NeutronHPFission model generates the final state for neutron-induced fission using the high precision neutron model. The G4NeutronHPorLFission model generates the final state for neutron-induced fission using the high precision neutron model when sufficient high precision data is available for the selected element or isotope. When there is insufficient data, neutron-induced fission is performed using the less precise Low Energy Parameterized model.

The G4HadronFissionDataSet is the default dataset for this process. The alternative high precision dataset is G4NeutronHPFissionData.

```
/gate/physics/addProcess Fission
```
Particle decay

Particle decay is the spontaneous process of one elementary particle transforming into other elementary particles. If the particles created are not stable, the decay process can continue. The majority of decays in Geant4 are implemented using the G4PhaseSpaceDecayChannel class. It simulates phase space decays with isotropic angular distributions in the center-of-mass system.

Radioactive decay

Radioactive decay is the process in which an unstable atomic nucleus spontaneously loses energy by emitting ionizing particles and radiation. In Geant4, the decay of radioactive nuclei by α, β+, and β emission and by electron capture are taken into account. The simulation model is empirical and data-driven, and uses the Evaluated Nuclear Structure Data File (ENSDF).

< ! > The download of radioactive decay data files is required.

Optical physics processes

For detailed information, see Users Guide V6.2:Generating and tracking optical photons.

Bulk Absorption

This process kills the optical photon. It requires the Material properties filled by the user with the Absorption length (average distance traveled by a photon before being absorbed by the medium).

Rayleigh Scattering

This process depends on the particle’s polarization. A photon which is not assigned a polarization at production may not be Rayleigh scattered. The photon is scattered in a new direction that is required to be perpendicular to the photon’s new polarization in such a way that the final direction, initial and final polarizations are all in one plane. The process requires the Material properties filled with Rayleigh scattering length (average distance traveled by a photon before it is Rayleigh scattered in the medium).
N.B: For Water ONLY, when scattering lengths are not specified but the user, the Geant4 code calculates them following the Einstein-Smoluchowski formula.

Mie Scattering

Mie Scattering is an analytical solution of Maxwell’s equations for scattering of optical photons by spherical particles. It is significant only when the radius of the scattering object is of order of the wave length. The analytical expressions for Mie Scattering are very complicated since they are a series sum of Bessel functions. One common approximation made is call Henyey-Greenstein (HG). The implementation in GATE (Geant4) follows the HG approximation and the treatment of polarization and momentum are similar to that of Rayleigh scattering.

The process requires Material properties to be filled by the user with Mie scattering length data (MIEHG). In practice, the user not only needs to provide the attenuation length of Mie scattering, but also needs to provide the constant parameters of the approximation: MIEHG_FORWARD, MIEHG_BACKWARD, and MIEHG_FORWARD_RATIO.

Processes at Boundaries

The optical boundary process design relies on the concept of surfaces: physical properties of the surface itself (stored in Materials.xml) and characteristics of the surface specifying the two ordered pairs of physical volumes touching at the surface (Surface.xml).

When the surface concept is not needed, and a perfectly smooth surface exists between two dielectric materials, the only relevant property is the index of refraction, a quantity stored with the material.

Wavelength Shifting or Fluorescence

Fluorescence is the result of a three-stage process that occurs in certain molecules called fluorophores or fluorescent dyes. A fluorescent probe is a fluorophore designed to respond to a specific stimulus or to localize within a specific region of a biological specimen. The process responsible for the fluorescence involves the creation of an excited electronic singlet state by optical absorption and subsequent emission of an optical photon of lower energy than the excitation photon.
A magnetic field can be defined. It will be attached and thus active in the whole world volume. It is currently not possible to confine the field to another volume. The following command can be used to activate and define the magnetic field properties:

```
/gate/geometry/setMagField Bx By Bz Unit
```

Here is the help for the command:

```
Command /gate/geometry/setMagField
Guidance :
Define magnetic field.

Parameter : Bx
Parameter type : d
Omittable : False

Parameter : By
Parameter type : d
Omittable : False

Parameter : Bz
Parameter type : d
Omittable : False

Parameter : Unit
Parameter type : s
Omittable : True
Default value : tesla
Candidates : T kG G tesla kilogauss gauss
```


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Users Guide V7.0: Cut and Variance Reduction Technics

From GATE collaborative documentation wiki

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    - Splitting
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Production threshold, step limiter and special cuts

Production threshold

To avoid infrared divergence, charged particles processes (ionization and bremsstrahlung) require a threshold below which no secondary will be generated. Because of this requirement, gammas, electrons and positrons require production thresholds which the user should define. This threshold should be defined as a distance, or range cut-off, which is internally converted to an energy for individual materials. Production thresholds are defined for a geometrical region. In GATE, each volume is considered as a geometrical region. If no cut is defined, the region inherited the threshold of the parent volume. The default cut value of the world is set to 1.0 mm.

```
/gate/physics/Gamma/SetCutInRegion  [Volume Name]  [Cut value]  [Unit]
/gate/physics/Electron/SetCutInRegion  [Volume Name]  [Cut value]  [Unit]
/gate/physics/Positron/SetCutInRegion  [Volume Name]  [Cut value]  [Unit]
```

For example:

```
/gate/physics/Gamma/SetCutInRegion  world  1.0 mm
```

The list of production threshold in range and in energy for each volume can be display with the command:

```
/gate/physics/displayCuts
```

User should use this command after the initialization.
X-Ray cuts and auger electrons in photo-electric process

Secondary electron and photon production in low energy photo-electric process (livermore and penelope models) could be customized.

The auger electron could be activated in livermore and penelope models:

```
/gate/physics/processes/PhotoElectric/setAugerElectron true
```

In livermore model, an energy threshold could be defined from which secondary particles are produced. The threshold is defined for the whole world of the simulation.

```
/gate/physics/processes/PhotoElectric/setDeltaRayCut [value] [unit]
/gate/physics/processes/PhotoElectric/setXRayCut [value] [unit]
```

Step limiter

< ! > This part is recommended for advanced users only!

The step limiter can be considered as a process with fixed step size. It allows to limit the maximum size of step. As for production threshold, the step limiter is defined for a geometrical region and the region can inherit the step limiter of the parent volume. User have to define the step size in the region:

```
/gate/physics/SetMaxStepSizeInRegion [Volume Name] [Step size] [Unit]
```

For example:

```
/gate/physics/SetMaxStepSizeInRegion world 1.0 mm
```

Then, the step limiter process has to be add to particles:

```
/gate/physics/ActivateStepLimiter proton
```

Special cuts

< ! > This part is recommended for advanced users only!

The user can define four cuts to limit the tracking of a particle:

- the maximum total track length
- the maximum total time of flight
- the minimum kinetic energy
- the minimum remaining range

While step limiter is affected to a step, special cuts are affected to a track. When a particle is stopped, the
energy is deposited locally. As for production threshold, the special cuts are defined for a geometrical region and the region can inherit the special cuts of the parent volume.

User have to define the values of the special cuts in the region and activate the cuts for particles.

```
gate/physics/SetMaxToFInRegion world 5 s
gate/physics/SetMinKineticEnergyInRegion world 1 keV
gate/physics/SetMaxTrackLengthInRegion world 0.01 mm
gate/physics/SetMinRemainingRangeInRegion world 0.02 mm
gate/physics/ActivateSpecialCuts proton
```

The user does not have to define all the cuts. The `ActivateSpecialCuts` command is effective for all the special cuts that are defined.

**Variance reduction**

Two standard reduction variance techniques are available in GATE: splitting and russian roulette. The weight of secondary particles is recalculated in function of the number of secondaries generated. User can also defined filters to increase the efficiency of these techniques.

< ! > User have to verify that all the tools he used in his simulation take into account particle weight!

**Splitting and russian roulette**

**Splitting**

In this technique, the final state of the process is generated N times and the weight of each secondary is 1/N.

```
gate/physics/processes/Bremsstrahlung/activateSplitting [Particle] [N]
```

Example: to split 100 times the electron bremsstrahlung photon (not that we specify that the e- is the particle which do the bremsstrahlung, but the split is applied on the generated photon):

```
gate/physics/processes/Bremsstrahlung/activateSplitting e- 100
```

**Russian roulette**

In this technique, Russian roulette is played on secondary particles. The survival probability is 1/N and the weight of each secondary is N.
/gate/physics/processes/Bremsstrahlung/activateRussianRoulette [Particle] [N]

Parameter : [Particle]
Parameter type : s
Omittable       : False

Parameter : [N]
Parameter type : i
Omittable       : False

Example: to keep 2% of electron bremsstrahlung photon (2/100 = 1/50):

/gate/physics/processes/Bremsstrahlung/activateRussianRoulette e- 50

Selective splitting and russian roulette

To increase the efficiency of the splitting and the russian roulette technique, user can add selections criteria on the incident (primary) or secondary particles. The selection is done with filters. The filters for splitting and russian roulette are the same as for Actors. For example, to split bremsstrahlung photons with a vector direction inside a cone of 20 degrees around the x axis:

/gate/physics/processes/Bremsstrahlung/addFilter angleFilter secondaries
/gate/physics/processes/Bremsstrahlung/secondaries/angleFilter/setAngle 20
/gate/physics/processes/Bremsstrahlung/secondaries/angleFilter/setDirection 1 0 0

There are several filters types: filters on particle, particle ID, energy, direction, volume... See the chapter on Actor for a description of all filters.


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Users Guide V7.0:Source

From GATE collaborative documentation wiki

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The GPS (General Particle Source)

To introduce a source into a GATE simulation, the user has to define the type of source (voxelized, linacBeam, phaseSpace, PencilBeam, TPSBeam or GPS) ans its feature (angle, energy, and position). Many activity distributions are available in GATE. At each new event, the source manager decides randomly which source decays, and generates for it one or more primary particles.

Creating a source

Different type of sources can be defined in the same GATE simulation. Each source is independent. The command to create a source is given below:

```
//gate/source/NAME
```

where "NAME" defines the name of the source.

Adding a source

The next step is to add the source. For this, the users has to type the following GATE command:
In this example, a source "NAME" is added. Once a source has been added, a series of properties must be assigned to it. These properties are: activity, type of particle(s), energy distribution, energy value or bounds, angular emission, spatial distribution and location, and half-life. The commands required to assign these properties are described in the following paragraphs.

**Defining the type of source**

Many options are available to suit your specific applications:
- Using keywords (e.g. Fluor18, backtoback, linacBeam, phaseSpace, fastI124) to help you to define one or several properties.

**Ion source**

The ion source type can simulate any ion by defining its atomic number (Z), atomic weight (A), ionic charge in units of energy (Q), and its excitation energy in keV (E). It incorporates both the radioactive decay and the atomic de-excitation. This is the most "realistic" way of simulating a radio-nuclide; however, it is also the slowest.

To use the ion source:

```bash
/gate/source/NAME/gps/particle ion
/gate/source/NAME/gps/ion 8 15 0 0
/gate/source/NAME/setForcedUnstableFlag true
/gate/source/NAME/useDefaultHalfLife
```

In the above example, an ion source of oxygen-15 has been defined with Z=8, A=15, Q=0, E=0. If is too slow, other options are available as described below.

**Simple particles**

You can choose from a long list of simple particles ($e^-$, $e^+$, gamma, etc) to use in your simulations (use the help command to obtain the full list). For example:

```bash
/gate/source/NAME/gps/particle gamma
```

defines a photon source and

```bash
/gate/source/NAME/gps/particle e+
```
defines a positron source. If you choose to use a particle, you will have to define more properties. As an example, the correct use of a positron source simulating fluorine-18 is:

```plaintext
/gate/source/NAME/gps/particle e+
/gate/source/NAME/gps/energytype Fluor18
/gate/source/NAME/setForcedUnstableFlag true
/gate/source/NAME/setForcedHalfLife 6586 s
```

In the above example, more properties of fluorine-18 have been added by using the helper keyword Fluor18 (see below): half-life and energy distribution. Note that the branching ratios are not respected with this kind of simulation because no decay is simulated, i.e. emission is 100% positron. You may want to lower the activity by an appropriate factor to take this better point into account.

**Helper keywords**

These keywords are defined to help you to define the particle properties. The first three keywords define an positron energy distribution resulting from common beta emitters, and the last one defines a special two-particle source.

For instance, **Fluor18** defines the positron energy spectrum of fluorine-18. Note that this keyword define only the energy spectrum, you still have to specify the particle $e^+$ and the half-life.

**Back-to-back** This keyword is implemented for PET simulations where two annihilation photons are generated at 180 degrees. This type of source is faster to simulate than the ion source or the positron source and allows for selecting emission angle. To use the back-to-back source type:

```plaintext
/gate/source/NAME/setType backtoback
```

Note that there are no radioactive decays simulated when using the back-to-back type and that you still have to define the particle (gamma), energy type (Mono) and energy-value (0.511 MeV).

**FastI124**

FastI124 is a special source implementing a simplified decay scheme of the non-pure beta emitter iodine-124 in which positrons are emitted but not neutrinos, there is no nuclear recoil, gammas are emitted if their emission probability is > 1%; and no atomic de-excitation occurs (no x-rays, Auger electrons). These simplifications allow for an increase in speed with respect to the ion source while retaining important features of iodine-124, i.e. gammas may be emitted concurrently with positrons to possibly create "dirty" coincidences. Since decay is simulated, branching ratios are respected hence no activity compensation is necessary.

To use the fastI124 source:

```plaintext
/gate/source/NAME/setType fastI124
```

The source takes care of particle definitions (gamma, positron) and energy distribution so that there is no need to specify a particle or mention its energy.

**Defining the activity**
To define the activity of the given source, the user defines the amount of activity and its unit using the following command:

```
/gate/source/NAME/setActivity 5. becquerel
```

In this example, the total activity of the source referred to as "NAME" is set to 5 Bq. The activity can be defined in Curie (Ci) as well as in Becquerel (Bq).

### Defining the energy

#### Energy distribution

If the source does not take care of the type of energy distribution (e.g. fastI124), then it has to be explicitly defined. This can be achieved either by using a pre-defined spectrum (see helper keywords above) or by using built-in distributions.

Candidates for built-in energy distributions are: mono-energetic "Mono", linear "Lin", powerlaw "Pow", exponential "Exp", Gaussian "Gauss", bremsstrahlung "Brem", black-body "Bbody", cosmic diffuse gamma ray "Cdg", user-defined histogram "User", arbitrary point-wise spectrum "Arb", and user-defined energy per nucleon histogram "Epn". Capitalisation is important: only strings given exactly as above will be recognized.

In the following example, all particles have the same energy:

```
/gate/source/NAME/gps/energytype Mono
```

#### Energy value

You may have to specify the energy value (or bounds) depending on the type of energy distribution you have selected. For example, for monoenergetic distributions (like back-to-back sources), you specify the energy value with:

```
/gate/source/NAME/gps/monoenergy 511. keV
```

In the case of ions, the kinetic energy must be 0 since the ions are at rest:

```
/gate/source/NAME/gps/monoenergy 0. ev
```

Any type of energy unit within the International System of Units (SI) can be used: eV, GeV, MeV, keV...

### Examples

1. ion source for fluorine-18
2. ion source for fluorine-18
Defining the angular distribution of the emission

An emission angle distribution can be defined with the angular span using:

```
gate/source/NAME/gps/angtype iso
gate/source/NAME/gps/mintheta 90. deg
gate/source/NAME/gps/maxtheta 90. deg
gate/source/NAME/gps/minphi 0. deg
gate/source/NAME/gps/maxphi 360. deg
```

In this case, all particles have the same polar angle (theta) of 90 degrees. They are all emitted along directions orthogonal to the z-axis. The particles are emitted with an azimuthal angle (phi) between 0 and 360 degrees, along all possible directions.

By default, a full span of 0-180 degrees for the polar angle and 0-360 degrees for the azimuthal angle are defined. The emission span can be reduced for back-to-back sources to speed up the simulation.
Defining the shape of the source

The last step is to define its geometry. The following command defines the type of source distribution:

```
/gate/source/NAME/gps/type Volume
```

In the above description, a volumic source distribution has been chosen. Other types of source distribution can be used: *Point*, *Beam*, *Plane*, or *Surface*. The default value is *Point*.

For a *Plane* source, the source shape type can be *Circle*, *Annulus*, *Ellipsoid*, *Square*, or *Rectangle*. For both *Surface* and *Volume* sources, this can be *Sphere*, *Ellipsoid*, *Cylinder*, or *Para*. The default source is a *Point* source and so *Shape* is not set to any of the above types. Each shape has its own parameters:

```
/gate/source/NAME/gps/shape Cylinder
/gate/source/NAME/gps/radius 1. cm
/gate/source/NAME/gps/halfz 1. mm
```

In the previous commands, the source is a cylinder with a radius of 1 cm and a length of 2 mm. Very often, the half-length is given rather than the full length.

- To define a circle, the radius (*radius*) should be set. (In reality it is not really a circle but a disk).
- To define an annulus, the inner (*radius0*) and outer radii (*radius*) should be given.
- To define an ellipse, square, or rectangle, the half-lengths along *x* (*halfx*) and *y* (*halfy*) have to be given.
- To define a sphere, only the radius (*radius*) only has to be specified.
- To define an ellipsoid, its half-lengths in *x* (*halfx*), *y* (*halfy*), and *z* (*halfz*) have to be given.
- To define a cylinder with its axis along the *z*-axis, only the radius (*radius*) and the *z* half-length (*halfz*) have to be specified.
- To define parallelepipeds, the *x* (*halfx*), *y* (*halfy*), and *z* (*halfz*) half-lengths, and the angles alpha (*paralp*), theta (*parthe*), and phi (*parphi*) have to be given.

Define the placement of the source

The position of the source distribution can be defined using:

```
/gate/source/NAME/gps/centre 1. 0. 0. cm
```

In that example, the centre of the source distribution is 1 cm off-centered along the x-axis.

Movement of a source

Attach to a volume

The source can be attached to a volume:

```
/gate/source/[Source name]/attachTo [Volume Name]
```
If the volume moves during the simulation, the source moves along with the volume. Note that when attaching a source to a volume, the source's placement becomes relative to the volume.

**Confining a source**

*Note: this is the old way of moving a source. It is very inefficient. Please consider using the "Attach to a volume" method instead.*

To define sources in movement, the source distribution have to be confined in a Geant4 volume. This volume will be animated using the usual GATE command as described in Chapter 4 of this manual.

The command:

```
/gate/source/NAME/gps/confine NAME_P
```

specifies that the emission must be confined to a volume of the Geant4 geometry. In this case, the emission distribution is the intersection of the General Particle Source (GPS) and the Geant4 volume. The Geant4 volume must be specified by its physical volume name: GATEnname + '_P'.

One should note that the confinement slows down the simulation, the confinement volume must have an intersection with the GPS shape, and the confinement volume must not be too large as compared to the GPS shape.

A complete example of a moving source can be found in the SPECT benchmark or in the macro hereafter:

```
# Define the shape/dimensions of the moving source
/gate/MovingSource/geometry/setRmax 5. cm
/gate/MovingSource/geometry/setRmin 0. cm
/gate/MovingSource/geometry/setHeight 20. cm
/gate/MovingSource/moves/insert translation
/gate/MovingSource/translation/setSpeed 0 0 0.04 cm/s

# Define the shape/dimensions of the large sourcecontainer that should contain the full trajectory of the moving source
/gate/source/SourceContainer/gps/type Volume
/gate/source/SourceContainer/gps/shape Cylinder
/gate/source/SourceContainer/gps/radius 4. cm
/gate/source/SourceContainer/gps/halfz 30. cm
# Define the placement of the SourceContainer
/gate/source/SourceContainer/gps/centre 0. 0. 0. cm
# Define the source as a gamma source
/gate/source/SourceContainer/gps/particle gamma
# Define the gamma energy
/gate/source/SourceContainer/gps/energy 140. keV
# Set the activity of the source
/gate/source/SourceContainer/setActivity 5000. Bq
# Define a confinement and confine the large container to the MovingSource at a position defined by the time and the translation speed
/gate/source/SourceContainer/gps/confine MovingSource_P
```
Example: two gammas

The following example gives a script to insert a point source of back-to-back type.

```
# A new source with an arbitrary name #(``twogamma) is created
/gate/source/addSource twogamma
# The total activity of the source is set
/gate/source/twogamma/setActivity 0.0000001 Ci
# The source emits pairs of particles back-to-back
/gate/source/twogamma/setType backtoback
# The particles emitted by the source are gammas
/gate/source/twogamma/gps/particle gamma
# The gammas have an energy of 511 keV
/gate/source/twogamma/gps/energytype Mono
/gate/source/twogamma/gps/monoenergy 0.511 MeV
# The source is a full sphere with radius 0.1 mm,
# located at the centre of the FOV
/gate/source/twogamma/gps/type Volume
/gate/source/twogamma/gps/shape Sphere
/gate/source/twogamma/gps/radius 0.1 mm
/gate/source/twogamma/gps/centre 0. 0. 0. cm
# The angular distribution of emission angles is isotropic
/gate/source/twogamma/gps/angtype iso
# The parameters below mean that the source emits
# at all angles along the z axis
/gate/source/twogamma/gps/mintheta 0. deg
/gate/source/twogamma/gps/maxtheta 180. deg
# Uncomment the parameters below if you want the source
# to emit in an XY (transverse) plane
/gate/source/twogamma/gps/mintheta 90. deg
/gate/source/twogamma/gps/maxtheta 90. deg
# The parameters below mean that the source emits
# at all angles in the transverse (XY) directions
/gate/source/twogamma/gps/minphi 0. deg
/gate/source/twogamma/gps/maxphi 360. deg
```

Defining a cold source

To define a cold (i.e. with no activity) volume in a phantom, a dedicated command is available.

The command:

```
/gate/source/NAME/gps/Forbid Volume_Name
```

The following example explains how to use this option. First you must define a volume that defines the cold region:

```
/gate/world/daughters/name cold_area
/gate/world/daughters/insert cylinder
/gate/cold_area/vis/forceWireframe
/gate/cold_area/vis/setColor green
/gate/cold_area/geometry/setRmax 3.0 cm
/gate/cold_area/geometry/setHeight 1. cm
```

Then you describe your source with the Forbid command:
It is important to remember that the `/gate/run/initialize` command must have been executed prior to using the `Forbid` command because phantom geometries are not available until after they are initialized.

### Visualizing a source

To check that sources are at the right location in the geometry, you can use the following command

```
/gate/source/\[Source name\]/visualize
```

along with a real time viewer (e.g. OpenGL). To visualize a source, Gate will randomly pick a certain number of points within the source and display them on the screen, along with the geometry. The full syntax is:

```
/gate/source/\[Source name\]/visualize count color size
```

where `name` is the name of the source, `count` is the number of random points to pick up (must be > 0 and <= 10000), `color` is the color to assign to those points (valid colors are: white, gray, grey, black, red, green, blue, cyan, magenta, yellow), and `size` is the screen size (in pixels) of each point (must be > 0 and <= 20).

Depending on the size and shape of the source, more or fewer points may be necessary.

- **Example:**

```
/gate/source/backgroundSource/visualize 2000 yellow 3
/gate/source/hotRegion/visualize 5000 red 2
```

### Intensity

If several sources have been added and no activity is defined, user can use intensity to define the source priorities. A high intensity correspond to a high priority. For each event, the source is randomly selected taking into account the intensity of each sources.
Pencil Beam source

The simulation source can be a pencil beam. This source allows for characterizing a beam of particles having energy and optical properties. This beam can be used for instance in order to characterize a clinical beam interacting in a passive beam line or to characterize a specific spot from an active beam scanning delivery system.

```
/gate/source/addSource [Source name] PencilBeam
```

One can select the type of particle used for the pencil beam (proton, e-, etc.):

```
/gate/source/ [Source name] /setParticleType [particle_type]
```

Alternatively, one can define a specific type of ion, by defining the particle type as "GenericIon" and then specifying the particle parameters of the ion to be generated: Z: AtomicNumber, A: AtomicMass, Q: Charge of Ion (in unit of e), E: Excitation energy (in keV). As an example, the definition of a C12 ion beam is given:

```
/gate/source/ [Source name] /setParticleType GenericIon
/gate/source/PBS/setIonProperties 6 12 6 0
```

The energy spectrum of the source is Gaussian and is defined by a mean energy and standard deviation:

```
/gate/source/ [Source name] /setEnergy [mean_energy] [Unit]
/gate/source/ [Source name] /setSigmaEnergy [energy_standard_deviation] [Unit]
```

The source position can be set as follows:

```
/gate/source/ [Source name] /setPosition [Pos_X Pos_Y Pos_Z] [Unit]
```

The pencil beam shape is Gaussian. The spot size can is defined by the standard deviation of the normal probability density function in x and y directions. The beam default direction being +z.

```
/gate/source/ [Source name] /setSigmaX [spot_size_X] [Unit]
/gate/source/ [Source name] /setSigmaY [spot_size_Y] [Unit]
```

The beam is also characterized by its divergences: Theta in the XoZ plan and Phi in the YoZ plan. The beam divergence is defined by the standard deviation of the normal probability density function:

```
/gate/source/ [Source name] /setSigmaTheta [divergence_Theta] [Unit]
/gate/source/ [Source name] /setSigmaPhi [divergence_Phi] [Unit]
```
The correlation between spot size and divergence (in the two plans) is characterized by the beam emittance. The beam emittance is defined by the standard deviation of the normal probability density function. The Emittance of the beam has to be lower (or equal) than the ellipse phase space area: \[\text{Emittance}_{X,\Theta} \leq \pi \times \text{divergence}_{\Theta} \times \text{spot}_X\] and \[\text{Emittance}_{Y,\Phi} \leq \pi \times \text{divergence}_{\Phi} \times \text{spot}_Y\].

Please note that for emittance, the unit cannot be selected and has to be "mm*mrad".

\[
\text{/gate/source/ [Source name] /setEllipseXThetaEmittance [Emittance}_{X,\Theta} \ mm*mrad}
\]
\[
\text{/gate/source/ [Source name] /setEllipseYPhiEmittance [Emittance}_{Y,\Phi} \ mm*mrad}
\]

When defining the beam parameters, one can define the beam convergence or divergence in the two plans (XoZ and YoZ), by setting the "RotationNorm" either to "positive" for a convergent beam or to "negative" for a divergent beam.

\[
\text{/gate/source/ [Source name] /setEllipseXThetaRotationNorm [negative or positive]}
\]
\[
\text{/gate/source/ [Source name] /setEllipseYPhiRotationNorm [negative or positive]}
\]

Users can also define the direction of the beam, which is by default +z (0 0 1), by rotating the beam along the x, y and z axis. For instance, to rotate the beam direction around the x-axis by 90°.

\[
\text{/gate/source/ [Source name] /setRotationAxis 1 0 0}
\]
\[
\text{/gate/source/ [Source name] /setRotationAngle 90 \ deg}
\]

The number of particles simulated is defined using the conventional command:

\[
\text{/gate/application/setTotalNumberOfPrimaries [number_of_particles]}
\]

Example

In the following example, we defined a 180 MeV proton beam, with 1 MeV energy spread. The beam is asymmetrical and convergent. The direction is -Y.

\[
\text{/gate/source/addSource PBS PencilBeam}
\]
\[
\text{/gate/source/PBS/setParticleType proton}
\]
\[
\text{/gate/source/PBS/setEnergy 188.0 \ MeV}
\]
\[
\text{/gate/source/PBS/setSigmaEnergy 1.0 \ MeV}
\]
\[
\text{/gate/source/PBS/setPosition 0 0 0 \ mm}
\]
\[
\text{/gate/source/PBS/setSigmaX 2 \ mm}
\]
\[
\text{/gate/source/PBS/setSigmaY 4 \ mm}
\]
\[
\text{/gate/source/PBS/setSigmaTheta 3.3 \ mrad}
\]
\[
\text{/gate/source/PBS/setSigmaPhi 3.8 \ mrad}
\]
\[
\text{/gate/source/PBS/setEllipseXThetaEmittance/PBS/setEllipseXThetaEmittance 15 \ mm*mrad}
\]
\[
\text{/gate/source/PBS/setEllipseXThetaRotationNorm negative}
\]
\[
\text{/gate/source/PBS/setEllipseYPhiEmittance 20 \ mm*mrad}
\]
\[
\text{/gate/source/PBS/setEllipseYPhiRotationNorm negative}
\]
\[
\text{/gate/source/PBS/setRotationAxis 1 0 0}
\]
\[
\text{/gate/source/PBS/setRotationAngle 90 \ deg}
\]
\[
\text{/gate/application/setTotalNumberOfPrimaries 10}
\]

**TPS Pencil Beam source**
The source of the simulation can be a stack of pencil beams. This source has been designed in order to allow the simulation of real treatment plans for active beam scanning delivery techniques.

```
/gate/source/addSource [Source name] TPSPencilBeam
```

One can select the type of particle used for the pencil beam (proton, e-, etc.):

```
/gate/source/ [Source name] /setParticleType [particle_type]
```

A treatment plan is made of one or multiple fields, each field being described by a gantry angle and a collection of pencil beams having different energies, directions, weights etc. user has to select the "plan description file" of the simulation:

```
/gate/source/ [Source name] /setPlan [plan_description_file]
```

It is possible to simulate all fields simultaneously or only some of them, by setting the "NotAllowedField" option. This option allows for ignoring specific fields:

```
/gate/source/ [Source name] /setNotAllowedFieldID [field_ID_1]
/gate/source/ [Source name] /setNotAllowedFieldID [field_ID_2]
/gate/source/ [Source name] /setNotAllowedFieldID [field_ID_3]
```

etc.

In the "plan description file", each single spot is characterized by its position at treatment isocenter and also by its weight (intensity). By default, the weight corresponds to the number of Monitor Units (MU) delivered and is converted into a number of particles according to the proton stopping power in air. Warning! The conversion of MU to number of particles has been developped for protons only ! However, it is possible to define the particle weight as a number of particles (and not as MU), by setting the following command to true:

```
/gate/source/ [Source name] /setSpotIntensityAsNbProtons true
```

As each spot has an intensity in the treatment plan, it is possible to simulate either each spot with the same probability and setting the dose scoring weight equal to the intensity, or simulate each spot with a probability related to the intensity and setting the dose scoring weight to 1. The second option is strongly advised (for efficiency) and used by default. It is possible to select the first or second option by setting the "FlatGenerationFlag" to true or false, respectively.

```
/gate/source/ [Source name] /setFlatGenerationFlag false
```

The physical properties of each single pencil beam delivered are computed using the "source description file". This file consists in a set of polynomial equations allowing to define the physical and optical properties of each single pencil beam with energy. Pencil beam properties are those described in the previous section "Pencil Beam source".
Irradiation systems can be tuned with either a convergent beam or a divergent beam at the delivery system exit. By default, the system is defined as divergent. User can define either a convergent or a divergent beam by setting the following command either to "true" or "false", respectively:

```
/setBeamConvergence [true or false]
```

The number of particles simulated is defined using the conventional command

```
/setTotalNumberOfPrimaries 10
```

### Example

The following example shows how to simulate a proton treatment plan based on the 2 following input files: "MyPlanDescriptionFile.txt" and "MySourceDescriptionFile.txt". The beam is considered convergent and the spot intensities are defined as number of protons.

```
/addSource PBS TPSPencilBeam
/setParticleType proton
/setPlan MyPlanDescriptionFile.txt
/setNotAllowedFieldID 1
/setFlatGenerationFlag false
/setSourceDescriptionFile MySourceDescriptionFile.txt
/setSpotIntensityAsNbProtons true
/setBeamConvergence true
/setTotalNumberOfPrimaries 10
```

**About the "source_description_file"** It contains the source to isocenter distance, and scanning magnets distance to isocenter in x- and y-directions. These parameters allow for computing the position and direction of each single pencil beam at the nozzle exit. It contains also 8 polynomial equations: 2 describing the energy properties (mean energy in MeV and energy spread in % of the mean energy) and 6 describing the optical properties of the beam (spot size in mm, beam divergence in rad and beam emittance in mm.rad, in x- and y-directions at nozzle exit). Polynomials are functions of the system energy, which is read in the "plan description file" for each pencil beam. For each polynomial, one has to give the polynomial order and then the polynomial parameters. For instance, for a second order polynomial \( ax^2 + bx + c \), one has to give the polynomial order: 2, followed by the a, b and c parameters in this order. Please note that their is no choice about the units used for the different polynomials!! Please have a look to example "example_Radiotherapy/example5" in the source code.

**About the "plan_description_file"** It contains many informations about the plan, but not all of them are taken into account for the simulation, as for instance the number of fractions. These additional informations may be used in further releases. Please have a look to example "example_Radiotherapy/example5" and "Gate/examples/example_Radiotherapy/example5/data/PlanDescriptionToGATE.txt" file. Warning, the unused fields of the plan description file cannot be removed. The main parameters of the file are the number of fields, gantry angle for each field, energy of each layer from each field, number of spot in each layer, spot description (position in x- and y- direction at isocenter and intensity) for each spot from each layer.

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Users Guide V7.0: Voxelized Source and Phantom

From GATE collaborative documentation wiki

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Voxelized Source and Phantom

Introduction

Voxelized source and phantom's purpose is the use of digital phantoms or patient data as inhomogeneous anthropomorphic attenuation maps or emission sources so as to simulate and thus reproduce realistic acquisitions.

From its first public release, GATE is able to read digital phantom or clinic data stored in various image file formats so as to allow users to read attenuation maps or emission data from these voxelized phantoms and sources.

To read in voxelized geometry, GATE requires a text file for the description of materials (AttenuationRange.dat for instance) and a 3D image stored in one of the following formats: ASCII, Interfile, Analyze, MetaImage and DICOM (coming soon). The text file must provide a number of subdivisions, define intervals associated to each subdivision and attach them with a correlated material name.

Example of AttenuationRange.dat file:

```
# Number of subdivisions
3
# Define the intervals and attach a correlated material
0 10 Air
```
To read in voxelized sources, GATE requires a text file for the description of activity levels (ActivityRange.dat for instance) and a 3D image stored in one of the following formats: ASCII and Interfile. The text file must provide a number of subdivisions, define intervals associated to each subdivision and attach them with a correlated activity (in Bq). The possibility to extend image file formats for voxelized sources is still under development.

Example of ActivityRange.dat file:

```
# Number of subdivisions
6
# Define the intervals and attach a correlated activity (in Bq)
200 210 1.
211 220 3.
221 230 5.
231 240 10.
241 250 20.
251 255 40.
```

### Voxelized phantoms

#### Description of voxelized geometry

To import digital phantom or patient data to voxelized geometry, GATE needs a convenient method to describe multiple copies of a volume inside a mother volume. Obviously, this description can lead to problem of memory consumption and hence affect the efficiency of tracking algorithm (computation of the distance to the next boundary, search for the neighbouring voxels, particles stopped at each boundary etc.)

To save both memory consumption and computation time when simulating voxelized phantoms, new parameterization methods derived from Geant4 together with their corresponding navigation algorithms are often developed.

#### Regular parameterization method

Since Geant4.9.1 a new navigation algorithm, dubbed regular navigation, can be used for the tracking of particles in voxelized volumes. The regular navigation algorithm performs fast direct neighbouring voxel identification without a large memory overhead. This is the major source of acceleration of the implemented regular navigation algorithm. In addition, boundaries between voxels which share the same material can be ignored. Using this option, the geometry only limits tracking at the boundary between voxels with different materials, providing a significant reduction of the number of propagation steps. The regular navigator uses a new algorithm that performs this search only for the neighbours of the actual voxel. It therefore highly reduces the time spent on this search, as much as the number of voxels is large. It also includes a new method called ComputeStepSkippingEqualMaterials; when a boundary is encountered, the navigator searches for the next voxel it should enter and check if its material is the same as the actual one. If this is the case, this method is directly called again without passing through the navigator manager which loads the new properties of the next voxel, etc. Therefore the fewer the materials, the faster the simulation. In conclusion, the time saved using the regular navigator is directly dependent on the number of voxels and the
number of different materials contained in the voxelized phantom. The better acceleration factors were obtained while simulating PET acquisitions (3 different materials: air, water, bone) with finely sampled phantom definitions. This factor could be around 3 in those cases. However in any case, even with a lot of different materials, this navigator will always be faster than using older navigators such as parameterizedBoxMatrix or compressedMatrix. That is the reason why these navigators still be progressively deprecated.

Additionally, as the SkipEqualMaterials method can lead to fewer G4steps, one may want to force the stepping process at each boundary. In that case, the method can be inactivated using the following command:

```
/gate/world/anyname/setSkipEqualMaterials 0
```

Also the particle tracking will inevitably be less effective.

**Nested parameterization method**

Another method of creating parametrized volumes in Geant4, using nested parametrization and the corresponding navigation algorithm has been available in GATE since version 6.1. Based on parametrized approach, this method allows GATE storing a single voxel representation in memory and dynamically changing its location and composition at run-time during the navigation. The main advantage of this method is high efficiency in memory space. While reusing the same mechanism as parameterized volume, Nested representation also splits the 3D volume along the three principal directions, allowing logarithmic finding of neighbouring voxels. Nested approach supposes geometry has three-dimensional regular reputation of same shape and size of volumes without gap between volumes and material of such volumes are changing according to the position. Instead of direct three-dimensional parameterized volume, one can use replicas for the first and second axes sequentially, and then use one-dimensional parameterization along the third axis. This approach requires much less memory access and consumption for geometry optimization and gives much faster navigation for ultra-large number of voxels. Using Nested representation, images are split into sub-volumes of homogeneous composition, which are parallelepipeds, either of the voxel size or larger. The main drawback is that all the particles are forced to stop at the boundaries of all parallelepipeds, generating a supplementary step and additional time cost, even if the two neighboring parallelepipeds share the same content. Such artificial steps occur very often as human organs are far from being parallelepipedic.

**Regionalized parameterization method**

Recently, some GATE developers have proposed a new method for efficient particle transportation in voxelized geometry for Monte Carlo simulations, especially for calculating dose distribution in CT images for radiation therapy. The proposed approach, based on an implicit volume representation named segmented volume, coupled with an adapted segmentation procedure and a distance map, allows them to minimize the number of boundary crossings, which slows down simulation. Before being implemented within GATE, the method was developed using the GEANT4 toolkit and compared to four other methods: one box per voxel, parameterized volumes, octree-based volumes, and nested parameterized volumes. For each representation, they compared dose distribution, time, and memory consumption. The proposed method allows them to decrease computational time by up to a factor of 15, while keeping memory consumption low, and without any modification of the transportation engine. Speeding up is related to the geometry complexity and the number of different materials used. They obtained an optimal number of steps with removal of all
unnecessary steps between adjacent voxels sharing a similar material. However, the cost of each step is increased. When the number of steps cannot be decreased enough, due for example, to the large number of material boundaries, such a method is not considered suitable. Thus, optimizing the representation of an image in memory potentially increases computing efficiency.

**Fictitious interaction**

**Important note:** so far, this method is available in GATE v7.0 version using Interfile reader only.

For detailed information, please refer to Users_Guide_V6.2:Voxelized_Source_and_Phantom#Speeding-up_tracking:_Fictitious_interaction

**Description of voxelized phantoms**

Regular, Nested and Regionalized parametrization methods together with their corresponding navigation algorithms are available in GATE V7.0 for voxelized phantoms. Note that so far their current implementations do not support voxel visualization attributes on a per material basis.

To create a parameterized phantom object using any of the three above-mentioned methods, one can use the corresponding command lines as follows:

```
/gate/world/daughters/name anyname
/gate/world/daughters/insert ImageRegularParametrisedVolume
```

Or

```
/gate/world/daughters/name anyname
/gate/world/daughters/insert ImageNestedParametrisedVolume
```

Or

```
/gate/world/daughters/name anyname
/gate/world/daughters/insert ImageRegionalizedVolume
```

All these three methods support 3D images stored in various image file formats, which is automatically defined from their extension:

- ASCII
- Interfile format: header .h33 + raw image .i33
- Analyze format: header .hdr + raw image .img
- MetaImage format: header. mhd + raw image .raw
- DICOM format (dcm) is expected for the next GATE version (7.1)

**Conversion into material definitions**

Whatever the navigation algorithm selected, conversion of image grayscales into material definitions is performed as follows:
When already defined in GateMaterials.db file, appropriate material properties are assigned to each voxel encoded value using either a range translator (to be used with images providing label values) or a units to materials conversion descriptor (to be used with images providing label or HU values).

**Range translator**

Tables read by the range translator have a prescribed format.

The first line defines the number of material subdivisions (i.e. the number of subsequent lines). The following lines describe the intervals (i.e. range) of encoded values (bits or segmented values) associated to each subdivision (i.e material), followed by a material name. A particular material will be assigned to each voxel whose value falls within the material range.

One can keep specifying visibility boolean (true or false) and color attribute values (red, green, blue components and transparency coefficient) within the range translator as he did when using a previous GATE version. However, as previously mentioned, so far current implementations of any of the three new parametrization methods do not support voxel visualization attributes on a per material basis, hence preventing the voxelized phantom from being displayed.

Example of a range translation table (AttenuationRange.dat, for instance)

```
4
0 Air false 0.0 0.0 0.0 0.2
4 4 Water true 1.0 0.0 1.0 0.2
5 5 Water true 0.0 1.0 0.0 0.2
15 SpineBone true 0.0 0.0 1.0 0.2
```

In this example, the number of material subdivisions is 4. Material Air is assigned to pixels with value 0, Water to pixels with value 4 and 5, and SpineBone to pixels with value between 6 and 15.

**Units to materials conversion descriptor**

Units to materials conversion descriptor is a simple text file with three columns: Label or HU_start, Label or HU_end and material_name. It allows to associate a material to each label or HU voxel encoded value in the image. This text file can be written by hand or generated with the automated method, especially for HU values (see For RT applications only: Automated HU stoichiometric calibration section)

Example of a units to materials conversion descriptor (AttenuationRange.dat, for instance)

```
5
1 Air
4 Water
6 Bone
16 SpineBone
```

In this example, the material Air is assigned to pixels with value 0, Water to pixels with value between 1 and 3 and 4 and 5, and SpineBone to pixels with value between 6 and 15.

**Example of voxelized phantom description macro**
Example of Interfile format header

```
INTERFILE :=
!GENERAL IMAGE DATA :=
type of data := Tomographic
total number of images := 16
study date := 1997:11:28
study time := 00:00:00
imagedata byte order := LITTLEENDIAN
number of images/energy window := 16
process status := Reconstructed
matrix size [1] := 32
number format := unsigned integer
number of bytes per pixel := 2
scaling factor (mm/pixel) [1] := +8.8
scaling factor (mm/pixel) [2] := +8.8
number of projections := 16
extent of rotation :=
time per projection (sec) := 0
study duration (sec) := 0
maximum pixel count := +2.000000e+02
patient orientation := head_in patient
rotation := supine
!GENERAL DATA :=
data offset in bytes := 0
name of data file := brain_phantom.i33
```
Using such an image reader, digital phantom or patient data can be read in as voxelized attenuation geometries. Additionally, when a sensitive detector (phantomSD) is associated to this phantom, the system can retrieve information about the Compton and Rayleigh interactions within this volume.

Figure 7.1: attenuation map from digital Hoffman phantom (left: data - right: after translation to material definition).

For RT applications only: Automated HU stoichiometric calibration

To generate a correspondence between HU (voxel values) and material, you may use the following commands:

```
/gate/HounsfieldMaterialGenerator/SetMaterialTable                  data/Schneider2000MaterialsTable.txt
/gate/HounsfieldMaterialGenerator/SetDensityTable                   data/Schneider2000DensitiesTable.txt
/gate/HounsfieldMaterialGenerator/SetDensityTolerance               0.1 g/cm3
/gate/HounsfieldMaterialGenerator/SetOutputMaterialDatabaseFilename data/patient-HUmaterials.db
/gate/HounsfieldMaterialGenerator/SetOutputHUMaterialFilename       data/patient-HU2mat.txt
/gate/HounsfieldMaterialGenerator/Generate
```

In this case, you need to provide:

- "Schneider2000MaterialsTable.txt" calibration text file allowing to split the HU range into several materials (see [Schneider2000]).
- "Schneider2000DensitiesTable.txt" calibration text file to indicate the relation between HU and mass density (g/cm3). It is normally given by calibration of your CT scanner.
- the parameter "DensityTolerance" allows the user to define the density tolerance. Even if it is possible to generate a new Geant4 material (atomic composition and density) for each different HU, it would lead to too many different materials, with a long initialization time. So we define a single material for a range of HU belonging to the same material range (in the first calibration Table) and with densities differing for less than the tolerance value.
- the files "patient-HUmaterials.db" and "patient-HU2mat.txt" are generated and can be used with setMaterialDatabase and SetHUtoMaterialFile macros.
Examples are available here (http://wiki.opengatecollaboration.org/?title=GateRT)

## Voxelized sources

So far, there is only one parametrization method within GATE allowing to read in voxelized sources. It requires the user to provide 3D image stored in Interfile format. Its extension to other image file formats is expected for the next GATE release (7.1).

### Conversion into activity values

Each voxel of the grayscale image is converted into activity value using a range translation table that is the same kind as the voxelized phantom one.

An example of a range translation table from voxel encoded values to activities (ActivityRange.dat in the example) is shown below:

```
3
4 4 1
5 5 3
14 15 5
```

where you specify the number of subdivisions or intervals (3 in this example), followed by the intervals definition and the correlated activity attached to each interval. If the number in the ASCII file, for a given voxel, is for instance between 14 and 15, then the activity for that voxel is set to 5 Bq. The resulting voxelized source has thus a discretized number of activity values (preliminar segmentation).

### Example of voxelized source description macro

Example of voxelized source description macro reading in an InterFile image as source distributions is detailed below:

```plaintext
# DECLARATION OF THE FACT THAT A VOXELIZED SOURCE WILL BE USED
/gate/source/addSource hof_brain voxel
# DECLARATION THAT THE VOXELIZED SOURCE WILL BE ENTERED USING INTERFILE DATA
/gate/source/voxel/reader/insert interfile
# INSERT THE TRANSLATOR THAT WILL CONVERT THE IMAGE FROM DIGITAL VALUES TO ACTIVITY VALUES (HERE THE NUMBERS AND ARE THEN CONVERTED TO PREDEFINED ACTIVITIES
/gate/source/hof_brain/interfileReader/translator/insert range
/gate/source/hof_brain/interfileReader/rangeTranslator/readTable ActivityRange.dat
/gate/source/hof_brain/interfileReader/rangeTranslator/describe 1
# THE FOLLOWING LINE ALLOWS YOU TO INSERT THE INTERFILE DATAFILE
/gate/source/hof_brain/interfileReader/readFile hof_brain_phantom.h33
/gate/source/hof_brain/interfileReader/verbose 1
# THE DEFAULT POSITION OF THE VOXELIZED SOURCE IS IN THE 1ST QUARTER. SO THE VOXELIZED SOURCE HAS TO BE SHIFTED OVER HALF ITS DIMENSION IN THE NEGATIVE DIRECTION ON EACH AXIS
/gate/source/hof_brain/setPosition -128. -128. 0. mm
```

Using this InterFile reader any digital phantom or patient data, stored in InterFile format, can be read in as emission distribution. Afterwards, activity levels can be used to determine the number of primary particles for each voxel.

An example of the Hoffman brain phantom, where the gray scales have been translated to activity distributions is shown in Figure 7.2.

![Figure 7.2: emission map from digital Hoffman phantom (left: data - right: translated activity values).](image)

### Dose collection

To collect absorbed dose deposited in the phantom, attach the phantom sensitive detector with the new attachVoxelPhantomSD command (and not attachPhantomSD) and add a dose output module:

```plaintext
/gate/anyname/attachVoxelPhantomSD
/gate/anyname/addOutput outputModuleName
```

The output module responds to the following commands:

```plaintext
/gate/output/outputModuleName/saveUncertainty [true|false]
/gate/output/outputModuleName/setFileName anyFileName
```
The output file is a binary file (number format is 4 bytes float) containing the absorbed dose in cGy. It has the same dimensions as the phantom. The output module optionally writes a second binary file containing the uncertainty on absorbed dose expressed as a fraction between 0 and 1. The uncertainty file also has the same dimensions as the phantom and its creation is controlled by the `saveUncertainty` command. The file name is the same as the absorbed dose file with a capital **U** appended. By default, the output file name is `doseMatrix.bin` and the uncertainty file is not created.

**Example**

```plaintext
# Create a simple phantom called CCD
/gate/world/daughters/name CCD
/gate/world/daughters/insert parameterizedBoxMatrix

# Read the file : a 300x300x1 array
/gate/CCD/geometry/insertReader image
/gate/CCD/imageReader/insertTranslator tabulated
/gate/CCD/imageReader/tabulatedTranslator/readTable ccdTable.dat
/gate/CCD/imageReader/readFile ccd300Phantom.dat

# Place the phantom and rotate it so that it is in the XZ plane
/gate/CCD/placement/setTranslation 0 -82.269 0 mm
/gate/CCD/placement/setRotationAxis 1 0 0
/gate/CCD/placement/setRotationAngle 90 deg

# Attach the phantom SD and the output module
/gate/CCD/attachVoxelPhantomSD
/gate/CCD/addOutput doseOutput
/gate/output/doseOutput/saveUncertainty true
/gate/output/doseOutput/setFileName ccdDose.bin
```

**Comments**

Depending on the phantom dimensions, the use of a parameterizedBoxMatrix may increase memory usage by up to a factor of 2 and increase CPU time by 5-50.

If you plan to collect only dose in the phantom, it is suggested that you disable other types of output, for example:

```plaintext
/gate/output/ascii/disable
```

**Dose calculations**

The relative uncertainty on dose is calculated on a per voxel basis. Let \( \{d_i\} \) \( i = 1, \ldots, N \) be the sequence of energy deposits in a given voxel, we can calculate the following quantities:

Mean energy deposit:

\[
\bar{d} = E(d) = \frac{1}{N} \sum_{i=1}^{N} d_i
\]

Sample variance:
$s^2 = E(d^2) - E(d)^2$

$$s^2 = \frac{1}{N^2} \left[ N \sum d_i^2 - (\sum d_i)^2 \right]$$

Population variance estimator:

$$s^2 = \frac{N}{N - 1} s_0^2$$

Standard deviation:

$$s = s \sqrt{\frac{N}{N - 1}}$$

Standard error of the mean:

$$\hat{d} = \frac{s}{N} = \frac{s}{\sqrt{N - 1}}$$

**Real-time motion management for voxelized source and phantom**

- Generate N frames for the phantom corresponding to the time acquisition desired for example 50 frames for 5s so each frame is for .1 s;
- I assume the 50 frames are NCAT_frame_i33, NCAT_frame_2.i33, NCAT_frame_3.i33, NCAT_frame_4.i33 $...$ NCAT_frame_N.i33.

```plaintext
/gate/world/daughters/name Ncat
/gate/world/daughters/insert compressedMatrix
/gate/Ncat/geometry/insertReader interfile
/gate/RTPhantom/insert RTVPhantom
/gate/RTVPhantom/AttachTo Ncat
/gate/RTVPhantom/setBaseFileName NCAT
/gate/RTVPhantom/setStatusFile NCAT_header.h33
/gate/RTVPhantom/setNumberOfFrames 50
/gate/RTVPhantom/setTimePerFrame 0.1 s
/gate/Ncat/interfileReader/insertTranslator range
/gate/Ncat/interfileReader/rangeTranslator/readTable range.dat
/gate/Ncat/interfileReader/rangeTranslator/describe 1
/gate/Ncat/attachPhantomSD
/gate/Ncat/placement/setTranslation 0.0.0.0 mm
/gate/Ncat/interfileReader/describe 1
```

The header NCAT_header.h33 looks like:

```plaintext
!matrix size [1] := 128
```

number format := unsigned integer
scaling factor (mm/pixel) [1] := +3.125000e+00
scaling factor (mm/pixel) [2] := +3.125000e+00
number of slices := 128
slice thickness (pixels) := +3.125000e+00

For the activity source:

```plaintext
# VOLUME SOURCE
/gate/source/addSource voxel voxel
/gate/source/voxel/reader/insert interfile
/gate/RTVPhantom/AttachToSource voxel
/gate/source/voxel/interfileReader/translator/insert range
/gate/source/voxel/interfileReader/rangeTranslator/readTable activityRange_test.dat
/gate/source/voxel/interfileReader/rangeTranslator/describe 1
/gate/source/voxel/setType backtoback
/gate/source/voxel/gps/particle gamma
/gate/source/voxel/setForcedUnstableFlag true
/gate/source/voxel/gps/mono energy Mono
/gate/source/voxel/gps/monoenergy 0.511 MeV
/gate/source/voxel/setPosition -200. -200. -200 mm
/gate/source/voxel/gps/confine NULL
/gate/source/voxel/gps/angtype iso
/gate/source/voxel/dump 1
```

The activityRange_test.dat is a text file looking like this:

```
12
46  46  0
61  61  0
123 123 0
215 215 0
246 246 0
261 261 0
352 352 352
369 369 0
770 770 0
831 831 0
2300 2300 0
950 950 950
```

The TimeActivity_Tables.dat is a text file looking like this:

```
2
950 lesion.dat
352 liver.dat
```
Where the value 950 is the key corresponding in the attenuation map to the lesion and 352 is the key corresponding in the attenuation map to the liver.\ Note that lesion.dat is a text file which contains the time activity table curve for the lesion, as explain here:

```
19
0.0 0
0.01 89.5
0.02 158.8
0.03 199.3
0.04 228.1
0.05 250.4
0.06 268.7
0.08 297.4
0.1 319.7
0.15 360.3
0.2 389.1
0.3 429.6
0.4 458.4
0.5 480.7
0.6 498.9
0.7 514.3
0.8 527.7
0.9 539.5
1 550.0
```

Where first column is the time in second and the second one is the activity in Bq at time t.


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Users Guide V7.0: How to run Gate

From GATE collaborative documentation wiki

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- Interactive mode
- Running GATE in Qt mode
- Running parameterized macros
- How to launch DigiGate
- How to separate the phantom and detector tracking - Phase space approach
- Batch mode

Interactive mode

To start Gate in interactive mode, simply type:

```
$ Gate
```

and the following output (or something similar) will appear on the screen:

```
1  **********************************************
2  Geant4 version $Name:  
3                                 (3-October-2003)
4              Copyright : Geant4 Collaboration
5 **********************************************
6 Time set to (s) 0
7 Visualization Manager instantiating...
8 Visualization Manager initialising...
9 Registering graphics systems...
10 You have successfully chosen to use the following graphics systems.
11 Current available graphics systems are:
12   DAWNFILE (DAWNFILE)
13   VRML1FILE (VRML1FILE)
14   VRML2FILE (VRML2FILE)
15   OpenGLES2file (OGL2ES)
16   OpenGLstoredx (OGLSX)
17 /control/saveHistory
18 /run/verbose 0
19 /event/verbose 1
20 /tracking/verbose 1
21 /gate/timing/setTime 0. s
22 Time set to (s) 0
23 /gate/application/setTimeSlice 1. s
24 /control/Reset
25 /control/setEventVerbosity 0
26 /control/setRunVerbosity 0
27 /control/setTrackingVerbosity 0
28 /control/setEventVerbosity 0
29 /control/setRunVerbosity 0
30 /control/setTrackingVerbosity 0
31 /gate/Reinitialize
2 Idle>
```

This output will vary depending on your Gate installation, that is what software was installed and how it was installed. Notice that the numbers on the left do not appear in the actual output. They are shown here just for didactic purposes.

Lines 1-5 indicates the version of the Geant4 software in your installation. Lines 6-9 are initialization messages from Gate. If you installed Gate with visualization functions, then you should see messages like lines those appearing in lines 10-15. Then, Gate runs the file `prerunGate.mac` located in the `petsim` directory. It then outputs the command lines 16-22 found in that file. Finally, and if everything went right, then Gate outputs the interpreter command prompt (line 23). This means Gate is ready to read commands entered by the user.

If you are not yet familiar with Gate commands, you can get help by typing `ls`:

```
Idle> ls
Command directory path : /
Sub-directories :
   /control/ UI control commands.
   /units/ Available units.
   /persistency/ Control commands for Persistency package
   /geometry/ Geometry control commands.
   /tracking/ TrackingManager and SteppingManager control commands.
   /event/ EventManager control commands.
   /run/ Run control commands.
   /random/ Random number status control commands.
   /particle/ Particle control commands.
   /process/ Process Table control commands.
   /gate/ Gate detector control.
   /hits/ Sensitive detectors and Hits
   /digi/ DigitizerModule
   /vis/ Visualization commands.
 Commands :
19 Idle>
```

When the `Sub-directories` names (lines 4-17) end with a `/` (slash), it means you can go deeper in that sub-directory. For instance, let's say you want to find out more about how to run macros:

```
Idle> ls /control
Command directory path : /control/
Guidance :
UI control commands.
Sub-directories :
Commands : 
   execute * Execute a macro file.
   loop * Execute a macro file more than once.
   foreach * Execute a macro file more than once.
```

supressAbortion * Suppress the program abortion caused by G4Exception.
verbose * Applied command will also be shown on screen.
saveHistory * Store command history to a file.
stopSavingHistory * Stop saving history file.
alias * Set an alias.
unalias * Remove an alias.
listAlias * List aliases.
shell * Execute a (Unix) SHELL command.
manual * Display all of sub-directories and commands.
createHTML * Generate HTML files for all of sub-directories and commands.
maximizeStoredHistory * Set maximum number of stored UI commands.

At the end of the Sub-directories names means that it is the last level for that subdirectory. In line 7, it is explained that the command /control/execute executes a macro file. This command basically reads the macro file and executes the lines as they appear in the file. Suppose, you have a file named myScanner.mac that contains all the necessary commands to run a particular simulation. Then type:

```
Idle> /control/execute myScanner.mac
```

to run the macro file. The macro file myScanner.mac can contain additional /control/execute commands to run other macro files and so on. Gate will read and execute those files in the order in which they appear. Notice that /control/execute does not start a simulation (data acquisition), it simply reads the commands and executes them. The command that starts the actual simulation is /gate/application/startDAQ, which is usually the last command found in your macro files.

Depending on the level of verbosity that you have specified in your macro, you will see more or less messages about the simulation. If your simulation contains visualization commands, you will see an OpenGL window appear with a beautiful picture of your scanner.

At the end of your simulation, the command line interpreter prompt will appear again. To exit the interpreter, type:

```
Idle> exit
Graphics systems deleted.
Visualization Manager deleting...
```

You can display the help typing this command:

```
$ Gate -h
```

and the following output (or something similar) will appear on the screen:

```
Printing help!!!
Usage: Gate [OPTION]... MACRO_FILE
Mandatory arguments to long options are mandatory for short options too.
-h, --help             print the help
-a, --param            set alias. format is '{alias1,value1} {alias2,value2} ...'
-d                    use the DigiMode
-q                    use the Qt visualization mode
```

Running GATE in Qt mode

First you need to compile Geant4 with the variable 'GEANT4_USE_QT' setting to 'ON'. You can visualize the position of your system using the Qt mode. First you need to type the following command to your console:

```
$ Gate --qt
```

A window will display:
After you launch your macro GATE in the label 'Session:', you could visualize your system in the viewer windows:

In order to use Qt you have to write this line in your GATE macro:
IMPORTANT!!! Qt visualization mode is a visualization after the simulation. In fact you could zoom, translate, etc... only at the end of the simulation.

**Running parameterized macros**

It is very common for users to run several simulations that differ in only a few parameters. For instance, a user might have designed a small animal PET scanner and would like to estimate its performance for five different crystal materials and three energy windows. In that case, the user does not need to write a complete set of macros for each simulation scenario. Instead, he can write parameterized macros. The actual values of the parameters are specified on the command line when running Gate or they can be defined with the interpreter.

For instance, suppose we want to parameterize the lower and upper level energy discriminators and the length of coincidence window. Then, we may have the following `Acquisition.mac` macro file:

```plaintext
# DIGITIZER
/gate/digitizer/Singles/insert adder
/gate/digitizer/Singles/insert readout
/gate/digitizer/Singles/readout/setDepth 1
/gate/digitizer/Singles/insert blurring
/gate/digitizer/Singles/blurring/setResolution 0.26
/gate/digitizer/Singles/insert thresholder
/gate/digitizer/Singles/thresholder/setThreshold {lld} keV
/gate/digitizer/Singles/insert upholder
/gate/digitizer/Singles/upholder/setUphold {uld} keV

# COINCI SORTER
/gate/digitizer/Coincidences/setWindow {CoincWindow} ns
```

Lines 8, 10, and 11 define aliases for the lower level discriminator, the upper level discriminator, and the length of the coincidence window, respectively. An alias is always specified between `{}` (curled brackets) and it can consist of any set of characters.

To pass actual values to the macro file, we run Gate, for instance, as follows:

```bash
$ Gate -a [CoincWindow,10] [lld,350] [uld,650]
```

It is worth emphasizing the following points about aliases:

- The order of the aliases at the command line does not matter.
- Aliases are case sensitive, so `[lld,350]` is not the same as `[Lld,350]`.
- All aliases in your macro file(s) must be defined when you run Gate. If some are undefined the simulation will fail.

**How to launch DigiGate**

GATE offers an operating mode dedicated to digitizer optimization, known as DigiGate (see Users Guide V6.2: Digitizer and readout parameters). DigiGate works by re-reading a previously generated ROOT hit-file.

The use of DigiGate consists of two steps.

- In the first step, the simulation runs according to `MacroTest.mac`. This macro file should save the Hits data in the root output file with the name `gate.root` (which is the default name).
- In the second step, the digitizer modifications are made in `MacroTest.mac` (like a new module for the energy resolution, or a different dead-time...), and then the analysis is repeated by using the `gate.root` file as an input file for the program DigiGate. This is achieved by launching Gate with a `--d` option.

```bash
Gate < MacroTest.mac
-> a root output file is produced with Hits information.
-> the digitizer of MacroTest.mac is changed along with the name of the root output file.
```

```bash
Gate --d < MacroTest.mac
-> a new root output file is produced which incorporates the changes due to a different digitizer without having to repeat the particle generation and its propagation.
```

**How to separate the phantom and detector tracking - Phase space approach**

To speed-up the simulation, it is possible to split and separate the particle tracking. This is a phase space approach with the possibility to store the phantom tracking particle history in a root file and to use it as an input file for the detector tracking.

**Using Gate in the tracker mode: phantom tracking**

Basically, as illustrated in the folder example_TrackerDetector, 3 major command lines are available to use the phantom tracker mode:

```plaintext
# Selection of the tracking mode
/gate/stepping/SetMode Tracker
```

The **Option1** variable can be chosen from the following list:

- StopOnPhantomBoundary (Default): the particles are tracked until the last hit before the phantom boundary;
- StopAfterPhantomBoundary: the particles are tracked until the first hit after phantom boundary;
- KillTrackAndSecondaries: StopOnPhantomBoundary + no secondary production.

The **Option2** variable may be chosen from:

- KeepAll (Default): all particles (primary and secondary) are stored
- KeepOnlyPrimaries: only source particles are stored
- KeepOnlyPhotons: only photons are stored
- KeepOnlyElectrons: only electrons are stored

Two additional command line options are also available:

```
/gate/stepping/SetEnergyThreshold aThreshold keV
```

With this option, only particles reaching the phantom boundary with an energy greater than a threshold in energy of **aThreshold** (expressed in keV) will be stored.

```
/gate/stepping/SetTextOutput status
```

with a status flag set as On or Off. This command will print Tracks information in the PostStepInfo.txt file.

Finally the tracker mode acquisition will generate N root files named OutPutRoot_TrackerData_number.root. The base output name, which is OutPutRoot in the case of this example, is chosen by the user with the usual output command line to set the file name:

```
/output/root/setFileName OutPutRoot
```

### Using Gate in the detector mode: detector tracking

During the tracker mode acquisition, N files are generated with the following name architecture:

- OutPutRoot_TrackerData.root
- OutPutRoot_TrackerData_1.root
- OutPutRoot_TrackerData_2.root
  ...
- OutPutRoot_TrackerData_(N-1).root

To use the Detector Mode, the user must select the mode and specify that N TrackerData files were generated during the tracker mode. All this can be done using the 2 following command lines:

```
/gate/stepping/SetMode Detector
/gate/stepping/SetNumberOfTrackerDataFiles N
```

### New commands in detector mode

In Detector Mode, we need to tell GATE that N TrackerData files were generated during tracker mode and we should use these command lines:

```
/gate/stepping/SetMode Detector
/gate/stepping/SetNumberOfTrackerDataFiles N
```

### Batch mode

It is possible to run a Gate simulation in **batch** mode, i.e. the mode in which you do not need to enter the interpreter and run the `/control/execute` and `exit` commands every time.

If you want to run a simulations in **batch** mode, you can do so by typing the alias values before the file name of the macro you want to run. (In some previous versions this was accomplished by redirecting the standard input of Gate with the `<` symbol and the name of the file you wanted to run.) For example,

```
$ Gate -a [CoincWindow,10] [lld,350] [uld,650] myScanner.mac
```

In order to return to command prompt, the last line in **myScanner.mac** file must be

```exit```

This is very important, especially when you are running a series of simulations in sequence. If Gate does not find the exit command, it will return to the user interface prompt and the rest of the simulations will not run.
It is recommended to redirect the terminal output of the simulation (listing of physics processes, sources, run time, etc.) by writing it to a text file instead of printing it in the terminal. This allows one to store the terminal output of each simulation for later viewing. For example,

```
$ Gate -a [CoincWindow,10] [lld,350] [ulld,650] myScanner.mac > terminal_output.txt
```

The above command creates a file named terminal_output.txt and does not print to the terminal window.

When running multiple simulations simultaneously from the command line in batch mode, it is often desirable to have the process run in the background. This can be accomplished by inserting an ampersand "&" symbol at the end of the command. For example,

```
$ Gate -a [CoincWindow,10] [lld,350] [ulld,650] myScanner.mac > terminal_output.txt &
```

It is recommended (although not compulsory) to avoid running visualization commands in batch mode.

Users Guide V7.0: Visualization

From GATE collaborative documentation wiki

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- Important Hints
- Command Lines
  - Visualization with OpenGL
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  - Visualization with DAWN
  - Visualization with VRML
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Introduction

The visualization options in GATE provide the same functionalities as provided in GEANT4. Most options in GEANT4 to visualize detector geometry, particle trajectories, tracking steps, etc. are also available in GATE. The graphics systems that can be selected in GATE are: DAWNFILE, VRMLFILE (versions 1 and 2) and OpenGL in stored and immediate mode, with OpenGL required as an external library. Most of the libraries are freely available.

Important Hints

When loading digital images using OpenGL, the OpenGL Immediate-X viewer is recommended instead of the frequently used Stored-X viewer.

Using DAWN and VRMLVIEW, complicated geometries, like a huge number of crystals in a cylindrical PET system, may take very long to get rendered. To decrease the file size and to speed up the visualization, the following option may be used:

```
/gate/crystal/vis/setVisible 0
```

Using that option, the individual crystals are not rendered but they are shown as a wireframe instead.

Command Lines

Basic commands provided by the GEANT4-package can be used to set basic visualization options, as shown below.

Visualization with OpenGL

```
# VIEWER # Opening of the OpenGL Stored-X viewer which is the standard viewer
# /vis/open OGLSX
# or opening of an OpenGL Immediate-X viewer used for digital images
# /vis/open OGLIX
# Let Geant4 choose the best
/vis/open OGL
# define the zoom factor
/vis/viewer/zoom 1.5
# Set the viewing angle
/vis/viewer/viewpointThetaPhi 5 60
# Set the drawing style
/vis/viewer/set/style surface
# Tell the viewer to draw the volumes
/vis/drawVolume
# The trajectories for each run should be drawn together
# don't store trajectories = 0; store trajectories = 1
/tracking/storeTrajectory 1
# Requests viewer to refresh hits, tracks, etc., at end of event.
# Or to accumulate drawings. Detector remains or is redrawn.
/vis/scene/endOfEventAction accumulate
```

The following commands implement additional options relevant within GATE:

```
# draw object in Wireframe Mode
/gate/block/vis/forceWireframe
```

Instead of block as in the example here, objects like crystal, source, scanner can be assigned specific visualization properties.

The "autoUpdate" capability has been removed since version 6.0.0 and manual geometry update has to be used instead. This can be done using the following command at any moment in the GATE command prompt environment:

```
/gate/geometry/rebuild
```

Otherwise the complete geometry building is done when /gate/run/initialize command is given.

**Visualization of Images**

Since 7.0, GATE can show images with OpenGL but only in Immediate mode and the volume of the image must be in WireFrame mode. This functionality works with X11 and Qt.

**Immediate mode**

```
/vis/open OGLI
```
**Stored mode**

```bash
/vis/open OGLS
```
Visualization with DAWN

Instead of real-time visualization based on OpenGL, storing images in a file (mostly eps) for further processing might be useful. DAWN offers such options.

The package can be downloaded from the Internet and installed following the instruction given at http://geant4.kek.jp/tanaka/src/dawn_3_85e.taz

To use DAWN and DAWNFILE in your macro, a specific open command should be used, in replacement of the opening of OpenGL.
Specific environment variables have to be set in your shell script to have access to DAWN inside GATE. For instance, in a C-shell:

```bash
if ( Xn == Xy ) then
    setenv G4VIS_BUILD_DAWN_DRIVER 1
    echo "On this machine the G4VIS_BUILD_DAWN_DRIVER= G4VIS_BUILD_DAWN_DRIVER"
endif
```

and also

```bash
if ( Xn == Xy ) then
    setenv G4VIS_USE_DAWN 1
    echo "On this machine the G4VIS_USE_DAWN= G4VIS_USE_DAWN"
endif
```

### Visualization with VRML

Sometimes, it may be helpful to check a geometry setup by interactively manipulating the visualized scene. These features are offered by the option VRML2FILE in connection with an appropriate viewer like vrmlview. Such a viewer can be freely downloaded from: http://www.sim.no/products/VRMLview/

A specific environment variable has to be set first:

```bash
setenv G4VRMLFILE_VIEWER vrmlview
```

For using this option in Gate, the following line has to be added to the macro instead of the corresponding OpenGL opening:

```bash
/vis/open VRML2FILE
```

Again, the environment variables have to be properly set (here C-shell example):

```bash
if [ Xn = Xy ] ;
    then G4VIS_BUILD_VRML_DRIVER=1
    export G4VIS_BUILD_VRML_DRIVER
    echo "On this machine the G4VIS_BUILD_VRML_DRIVER=$G4VIS_BUILD_VRML_DRIVER"
fi
```

```bash
if [ Xn = Xy ] ;
    then G4VIS_USE_VRML=1
    export G4VIS_USE_VRML
    echo "On this machine the G4VIS_USE_VRML=$G4VIS_USE_VRML"
fi
```

During processing in GATE, a file is written with the extension wrl.

### Axes

Any position in the world is defined with respect to a three-axis system: X, Y and Z. These three axes can be seen in the display window using:

```bash
/vis/scene/add/axes
```

Users Guide V7.0:Bibliography

From GATE collaborative documentation wiki

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Users Guide V7.0: Architecture of the simulation

From GATE collaborative documentation wiki

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Main rules

GATE simulations are based on the execution of scripted commands (Users Guide V6.2: Getting started) gathered in macros.

A simulation is generally divided into 7 steps (Fig. 9.1) as follows:

1. Verbosity and Visualization (Users Guide V6.2: Visualization)
3. Digitizer (Users Guide V6.2: Digitizer and readout parameters)
4. Physics (Users Guide V6.2: Setting up the physics)
5. Sources (Users Guide V6.2: Activity, source, voxelized phantoms)
6. Outputs (Users Guide V6.2: Data output)
7. Experiment (Users Guide V6.2: Digitizer and readout parameters)

The first 4 steps correspond to the PreInit mode of GEANT4 whereas the last 3 occur after the initialization of the simulation. The first 4 steps are validated by the GEANT4 command:

```
/gate/run/initialize
```

Once this phase is completed, the sources can be inserted in the setup and the simulation can be run.


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Verbosity and Visualization

Verbosity

For each simulation module, one can set a verbosity level between 0 and 2. The higher the verbosity level, the higher the level of information returned by GATE. By default, the verbosity level is set to 0, but if one wants to follow in detail each step of the simulation, it can be set to higher values. As an example, to get the computation time of the simulation written by Gate on the screen at the end of a simulation like this:

```
User simulation time (sec) := 13.9
Real simulation time (sec) := 15.92
System simulation time (sec) := 0.03
```

the verbosity level of the output should be set to 2:

```
/gate/output/verbose 2
```

Visualization

There are several tools available for visualization OpenGL, VRML, DAWN. They can be activated as a function of the visualization options selected at the GEANT4 configuration step. The on line visualization is a useful tool when developing new geometries. It allows one to visually check the scanner geometry (positions, physical volume overlaps, etc). Once the geometry is checked and one wants to run a complete simulation, it is recommended to disable the on line visualization in order not to overload the CPU.

Geometry

The world

The first volume to be created is the world. Each new volume will be inserted in this one, with a given name, and will be a daughter of the world. The world dimensions must be large enough to include the scanner and the phantom.

The system

Next, the system type must be chosen, as a function of the scanner to be modelled: scanner, PETscanner, cylindricalPET, ecat, CPET or SPECTHead. Each system has a defined number of levels with a specific hierarchical organization (tree geometry) and it is linked with a specific data output format. Two output formats are also available, independent of the selected system (ROOT and ASCII). Once the scanner is built, all the scanner elements should be attached to the system.

The phantom

A phantom has to be defined, using materials included in the materials database. The phantom can be an analytical or voxelized volume. Each voxel of the volume can be made of a specific material with its own
density.

Finally, the sensitive volumes \textit{crystalSD} and \textit{phantomSD} have to be attached. Only the interactions (\textit{hits}) occurring in a volume attached to a sensitive detector will be stored by GATE for the digitization.

\section*{Digitizer}

The \textit{digitizer} pre-processes the \textit{hits} by sorting, regrouping and adding them to create \textit{singles}. The \textit{singles} are time-stamped and stored in the events history. In PET, the \textit{coincidences} are then sorted out as a function of the coincidence window width. The detection parameters (\textit{temporal resolution, crystal blurring, dead time, threshold, uphold}...) are set at the \textit{digitizer} level.

\section*{Physics}

The physics part of the simulation defines the simulated physical processes by:

\begin{itemize}
  \item selecting the appropriate interactions library (\textit{Standard} or \textit{Low Energy package}),
  \item enabling or disabling the physical processes (\textit{Photoelectric effect, Compton effect, Rayleigh, gamma conversion}...),
  \item setting cutoffs in energy or range.
\end{itemize}

\section*{Sources}

A source is defined by:

\begin{itemize}
  \item its nature (\textit{particle / ion}),
  \item its activity (\textit{initial activity, half live}...),
  \item its geometry (\textit{shape or voxelized}),
  \item its emission angle,
  \item its movement if necessary.
\end{itemize}

The source activity can be confined in a specific volume (\textit{e.g} the phantom volume).

\section*{Data outputs}

The data output formats are of two types in GATE:

\begin{itemize}
  \item Standard outputs: ASCII, Root
  \item System dependent outputs: LMF, sinogram, ecat7, Interfile
\end{itemize}

\subsection*{Standard outputs}

ASCII outputs are spread out into 3 files (\textit{Hits, Singles, Coincidences}). The Root output is composed of one NTuple (\textit{Gate}) and three TTrees (\textit{Hits, Singles, Coincidences}) in which the interaction type, position and time information are stored. By default these two types of output are enabled. Each one of these outputs can be enabled or disabled according to the kind of information one is interested in.

\subsection*{Specific outputs}
In addition to the two standards outputs, GATE provides system dependent outputs. The LMF output is linked to the cylindricalPET system. The Sinogram and the ecat7 outputs are related to the ecat system, while the Interfile output relates to the SPECTHead system.

All these outputs are characterized by several parameters which have to be correctly set up as explained in Users Guide V6.2:Data output.

Experiments

The last step of the simulation consists in setting up the experiment. In this part, the duration of the simulated acquisition is set by defining the beginning and the end of the simulation. The overall acquisition time can be subdivided in several time slices of fixed duration. This feature is very useful in GATE, since the geometry is updated only between two time slices. This provides the possibility to take into account the movements of the sources or the detectors by subdividing a run in time slices during the geometry remains unchanged.

Random generator

As a Monte Carlo tool, GATE needs a random generator. The CLHEP libraries provide various ones. Three different random engines are currently available in GATE, the Ranlux64, the James Random and the Mersenne Twister. The default one is the Mersenne Twister, but this can be changed easily using:

```
/gate/random/setEngineName aName
```

where aName can be: Ranlux64, JamesRandom, or MersenneTwister.

NB Several users have reported artifacts in PET data when using the Ranlux64 generator. These users have said that the artifacts are not present in data generated with the Mersenne Twister generator.

The choice of the generator seed is also extremely important. There are 3 ways to make that choice:

- The 'default' option. In this case the default CLHEP internal seed is taken. This seed is always the same.
- The 'auto' option. In this case, a new seed is automatically generated each time GATE is run.

To randomly generate the seed, the time in millisecond since January 1, 1970 and the process ID of the GATE instance (i.e. the system ID of the running GATE process) are used. So each time GATE is run, a new seed is used.

- The 'manual' option. In this case, the user can manually set the seed. The seed is an unsigned integer value and it is recommended to be included in the interval [0,900000000].

The commands associated to the choice of the seed with the 3 different options are the following:

```
/gate/random/setEngineSeed default
/gate/random/setEngineSeed auto
/gate/random/setEngineSeed 123456789
```
It is also possible to control directly the initialization of the engine by selecting the file containing the seeds with the command:

```
/gate/random/resetEngineFrom fileName
```

Finally, the level of verbosity of the random engine can be chosen. It consists into printing the random engine status, depending on the type of generator used. The command associated to the verbosity is:

```
/gate/random/verbose 1
```

Values from 0 to 2 are allowed, higher values will be interpreted as 2. A value of 0 means no printing at all, a value of 1 results in one printing at the beginning of the acquisition, and a value of 2 results in one printing at each beginning of run.

**Example of a PET scanner**

The following example describes how to build a PET scanner based on the `cylindricalPET` system.

```plaintext
# No verbosity
/control/verbose 0
#
# OpenGL online visualization
/vis/open OGLSX
/vis/viewer/reset
/vis/viewer/set/style surface
/vis/drawVolume
/visscene/evAction accumulate
/vis/viewer/viewpointThetaPhi 30 30
/vis/viewer/zoom 2
#
/tracking/storeTrajectory 1
/gate/geometry/enableAutoUpdate
```

The on line visualization is enabled.

```plaintext
# WORLD
/gate/world/geometry/setXLength 40 cm
/gate/world/geometry/setYLength 40. cm
/gate/world/geometry/setZLength 40. cm
```

The *world* is created. Its dimensions should be large enough to contain all the volumes describing the experiment.

```plaintext
# CYLINDRICAL
/gate/world/daughters/name cylindricalPET
/gate/world/daughters/insert cylinder
/gate/cylindricalPET/setMaterial Water
/gate/cylindricalPET/geometry/setRmax 152 mm
/gate/cylindricalPET/geometry/setRmin 130 mm
/gate/cylindricalPET/geometry/setHeight 80 mm
/gate/cylindricalPET/vis/forceWireframe
```
The system is chosen.

```plaintext
# R S E C T O R
/gate/cylindricalPET/daughters/name box1
/gate/cylindricalPET/daughters/insert box
/gate/box1/placement/setTranslation 140 0 0 mm
/gate/box1/geometry/setXLength 20. mm
/gate/box1/geometry/setYLength 19. mm
/gate/box1/geometry/setZLength 76.6 mm
/gate/box1/setMaterial Water
/gate/box1/vis/forceWireframe
# M O D U L E
/gate/box1/daughters/name box2
/gate/box1/daughters/insert box
/gate/box2/geometry/setXLength 20. mm
/gate/box2/geometry/setYLength 19. mm
/gate/box2/geometry/setZLength 19. mm
/gate/box2/setMaterial Water
/gate/box2/vis/forceWireframe
# C R Y S T A L
/gate/box2/daughters/name box3
/gate/box2/daughters/insert box
/gate/box3/geometry/setXLength 20. mm
/gate/box3/geometry/setYLength 2.2 mm
/gate/box3/geometry/setZLength 2.2 mm
/gate/box3/setMaterial Water
/gate/box3/vis/forceWireframe
# L A Y E R L S O
/gate/box3/daughters/name LSO
/gate/box3/daughters/insert box
/gate/LSO/geometry/setXLength 10. mm
/gate/LSO/geometry/setYLength 2.2 mm
/gate/LSO/geometry/setZLength 2.2 mm
/gate/LSO/placement/setTranslation -5 0 0 mm
/gate/LSO/setMaterial LSO
# L A Y E R LuAP
/gate/box3/daughters/name LuAP
/gate/box3/daughters/insert box
/gate/LuAP/geometry/setXLength 10. mm
/gate/LuAP/geometry/setYLength 2. mm
/gate/LuAP/geometry/setZLength 2. mm
/gate/LuAP/placement/setTranslation 5 0 0 mm
/gate/LuAP/setMaterial LuAP
/gate/LuAP/vis/setColor cyan
# R E P E A T C R Y S T A L
/gate/box3/repeaters/insert cubicArray
/gate/box3/cubicArray/setRepeatNumberX 1
/gate/box3/cubicArray/setRepeatNumberY 8
/gate/box3/cubicArray/setRepeatNumberZ 8
/gate/box3/cubicArray/setRepeatVector 10. 2.4 2.4 mm
# R E P E A T M O D U L E
/gate/box2/repeaters/insert cubicArray
/gate/box2/cubicArray/setRepeatNumberX 4
/gate/box2/cubicArray/setRepeatVector 0. 0. 19.2 mm
# R E P E A T R S E C T O R
/gate/box1/repeaters/insert ring
/gate/box1/ring/setRepeatNumber 42
# A T T A C H S Y S T E M
/gate/systems/cylindricalPET/rsector/attach box1
/gate/systems/cylindricalPET/module/attach box2
/gate/systems/cylindricalPET/crystal/attach box3
```

The volumes of the scanner (with names given by the user, e.g. box1) are connected to the cylindricalPET system (to the predefined names of cylindricalPET system, e.g rsector).

The scanner system is completely built. It is composed of 42 \textit{rsectors}, each one made of 8 x 8 LSO and LuAP crystals assembled in phoswich.

The scanner rotates along the Z axis with a rotation speed of 6 deg/s while the phantom has translation movement with a speed of 0.1 cm/s.

The scanner rotates along the Z axis with a rotation speed of 6 deg/s while the phantom has translation movement with a speed of 0.1 cm/s.
The digitizer is set up with a crystal blurring randomly chosen between a minimum and a maximum value. A threshold and an uphold have been respectively set to 250 keV and to 750 keV. The coincidence time window has been set to 10 ns and a coincidence event will be taken into account only if the difference between the two sectors is greater than or equal to 2. A paralyzable dead-time of 250 ns has been introduced at sector level.

At this point, the construction of the detector is over. We can now initialize the simulation.

An important check can be made just after this initialization in order to test that there is no overlap between volumes from the same "family" (the mother volume and her daughters) and that a daughter volume is
inside her *mother* volume. This test is done only once in order to check the geometry.

```plaintext
/geometry/test/recursive_test
#
  V E R B O S I T Y
/control/verbose 0
/grdm/verbose 0
/run/verbose 0
/event/verbose 0
/tracking/verbose 0
/gate/application/verbose 0
/gate/generator/verbose 0
/gate/stacking/verbose 0
/gate/event/verbose 0
/gate/source/verbose 0
#
  S O U R C E S
/gate/source/addSource twogamma
/gate/source/twogamma/setActivity 1000. becquerel
/gate/source/twogamma/setType backtoback
/gate/source/twogamma/gps/particle gamma
/gate/source/twogamma/gps/energytype Mono
/gate/source/twogamma/gps/monoenergy 0.511 MeV
/gate/source/twogamma/gps/centre 0. 2. 0. cm
/gate/source/twogamma/gps/type Volume
/gate/source/twogamma/gps/shape Sphere
/gate/source/twogamma/gps/radius 0.5 cm
/gate/source/twogamma/gps/shape Cylinder
/gate/source/twogamma/gps/radius 10. mm
/gate/source/twogamma/gps/halfz 50. mm
/gate/source/twogamma/gps/confine phantom
/gate/source/twogamma/gps/angtype iso
/gate/source/twogamma/gps/mintheta  0. deg
/gate/source/twogamma/gps/maxtheta 180. deg
/gate/source/twogamma/gps/minphi  0. deg
/gate/source/twogamma/gps/maxphi 360. deg
#
/gate/source/addSource sourceC11
/gate/source/sourceC11/setActivity 10000. becquerel
/gate/source/sourceC11/gps/particle e+
/gate/source/sourceC11/setForcedUnstableFlag true
/gate/source/sourceC11/setForcedHalfLife 1223 s
/gate/source/sourceC11/gps/energytype Carbon11
/gate/source/sourceC11/gps/centre 0. 0. 0. cm
/gate/source/sourceC11/gps/type Volume
/gate/source/sourceC11/gps/shape Cylinder
/gate/source/sourceC11/gps/radius 10. mm
/gate/source/sourceC11/gps/halfz 50. mm
/gate/source/sourceC11/gps/confine phantom
/gate/source/sourceC11/gps/angtype iso
#
/gate/source/list
```

We have defined two sources. The first one called *twogamma* has an activity of 1 kBq and emits two gammas back to back in all directions. The second one, *(sourceC11)*, emits beta+ according to the positron energy spectrum of $^{11}_C$ decays, with an initial activity of 10–kBq and a halflife of 1223 s. The range of the positron is simulated as well as the gamma – gamma acolinearity.

The two sources are confined in the *phantom*. 
The ASCII and LMF outputs are disabled and for the ROOT output, only Gate NTuple, Singles and Coincidences TTrees are stored.

The acquisition duration will last 60 s with slices of 1 s. The geometry will thus be updated every second.

**Important hint**

You can define the complete simulation in one macro. In order to have a more modular simulation you can divide it into several macros (e.g. `vis.mac`, `geometry.mac`, `digi.mac`, `physics.mac`, `sources.mac`, `main.mac`) called from a main macro.
Figure 9.1: GATE simulation architecture

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Users Guide V7.0: Defining a system

From GATE collaborative documentation wiki

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- How to connect the geometry to a system
- Different types of systems
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  - CylindricalPET
  - CPET
  - Ecat
  - ecatAccel
  - OPET
  - SPECTHead
  - OpticalSystem
- How to define a multi-system detector

Definition

A System is a key-concept of GATE. It provides a template of a predefined geometry to simulate a scanner. A system can be used to model several scanners sharing the same general geometrical characteristics. It can be considered as sort of a template described by key components organized in a certain way, what is called a tree level structure, each component having its own specific role or ordering.

For instance, in the cylindricalPET scanner system, the geometrical volumes containing crystals are grouped in matrices, themselves assembled in submodules and modules. At the top level of this structure, the sectors composed of modules are repeated on a cylindrical surface to build up the whole device. Thus, a family of PET scanners obeying this structure can be described using this system, illustrated in figure 4.1, composed of volumes called rsectors, modules, submodules, crystal and finally (crystal) layer.

Different systems are available in GATE: scanner, SPECTHead, cylindricalPET, ecat, CPET, OPET and OpticalSystem, which can be used to simulate most of the existing imaging devices.

Choice of the system

It is possible to use GATE without using a system, but in that case, no information regarding particle interaction in the detector will be available. The reason is that the volumes where the hits (interactions that occur inside the detector parts of the scanner, see Users Guide V6.2: Digitizer and readout parameters) are recorded only for volumes belonging to a defined system (those volumes are declared as crystalSD, SD for sensitive detector, see Users Guide V6.2: Attaching the sensitive detectors). When the user is only testing a scanner geometry, the use of a predefined system is not necessary. But if the user wants to record information related to the particle history inside the detector, the geometry has to be associated with a system. This section explains the elements and rules to associate a geometry with a system.

Geometry constraints

Except for the general system scanner, one should first take into account the geometrical shape of the different components (gantry, sector, bucket, etc.) and also the shape of the crystal or the detector material (e.g. scintillators).

Each level has to be assigned to a physical volume of the geometry. A level volume has to be fully enclosed in the upper level volume.

The number of levels has to be set and must conform to the specifications listed in table 4.1. The numbering of different sensitive volumes is completely set by the choice of the system and conforms to a specific output format.
The maximum number of components in each level depends on the output format since it can be limited by the number of bits used for the numbering of the crystals. See Users Guide V6.2:Data output for further details.

Constraints related to the simulation of the DAQ electronics

Several points have to be considered when designing the simulation of the electronics cards. First, the whole readout electronic components should be analyzed in order to define its main components. This concerns not only the single channel simulation, with effects like thresholder response, but also the crosstalk between different channels including the electronic or the optical crosstalk among components in a same level. For a PET scanner, the coincidence between two channels have to be simulated, based on the single component simulations. In GATE, it is possible to introduce all these signal processing steps through the digitizer modules (see Users Guide V6.2:Digitizer and readout parameters), operating at different levels or depths, as shown in Table 4.2. The depth value is used here to tag a group of similar components operating at a certain level, which could be the scintillator block level (crystal with depth=5, or a group of crystal matrices with depth=1, depth values given in these examples refer to the cylindricalPET system).

To simulate the electronic processing consistently with the system used to model the detector, the following procedure should be used:

- Regroup the detector electronic components in different levels.
- List the signal processing to be used for each of the resulting groups (see adder, readout, dead time in Users Guide V6.2:Digitizer and readout parameters).
- Combine the signals coming from different volumes with, for example, the readout module for the signals summation of a volume, or the crosstalk and/or the coincidence between signals and coincidence.

NOTE : One or several crosstalk processing can be applied to components of different levels, for instance crosstalk between crystals, followed by crosstalk between modules. Such processing involves components at the same level. For PET scanners, coincidences are validated by testing the number difference in the uppermost level (as defined as depth = 1 in table ). This test can reject accidental coincidence between adjacent logic structures. When the user builds a geometry, this logic organisation should correspond to the first level of a system to use this coincidence sorting (see Users Guide V6.2:Digitizer and readout parameters).

How to connect the geometry to a system

The connection between the geometry and a system is performed in several steps:

- The geometrical structure needs first to be defined, keeping in mind that it must fulfill some constraints, as described before.
- The system geometry has then to be introduced, or attached, in the simulation process with the ```attach``` command and a specific keyword argument corresponding to one level of the geometrical structure. The general macro line for this attachment is:

```
 Systems/SystemName/Level/attach UserVolumeName
```

where : SystemName is the specific name of the system (one of the entry in column 1), Level is the specific name of the level (see column 2), UserVolumeName is the name the user gave to a volume, according to the conventions of Users Guide V6.2:Defining a geometry.

- Finally, the specific output corresponding to the system has to be defined for further data analysis (see Users Guide V6.2:Data output).

Table 4.1: Different systems available in GATE and their characteristics. In the second column are listed some of the keyword that are also used at in the macro (see also table 4.2 for a complete list). The shape in the third column describe the mother volume, composed of “daughter” volumes as described in Chap. 3 : a box means a box shaped mother volume containing an array of daughter boxes, a cylinder mother volumes will contains cylinders. Cylinders are understood here as tube sectors defined by an inner and outer radius.

<table>
<thead>
<tr>
<th>System</th>
<th>Components and Shape</th>
<th>Available Outputs</th>
</tr>
</thead>
<tbody>
<tr>
<td>scanner</td>
<td>level1 level2 level3 level4 level5 geometry not fixed</td>
<td>Basic output: ASCII or ROOT, coincidences only for PETscanner</td>
</tr>
<tr>
<td>CTscanner</td>
<td>module cluster pixel box box</td>
<td>Raw Data, ASCII, ROOT</td>
</tr>
</tbody>
</table>

sector | box | cylinder |

cassette | box | cylinder |

## Table 4.2: Keywords corresponding to system components definition to be used with an “attach” command.

At least one level has to be attached to the system. If necessary, these level’s names can be possibly used as input to digitizers modules: for example, different electronic dead times for each level’s electronics can be modelised. The two last lines, listed here for information, are related to “hits” which apply only for “sensitive” volume. Please refer to Chap. 5 for more details on this topic.

<table>
<thead>
<tr>
<th>System</th>
<th>Attach Keyword Argument</th>
<th>Depth for readout segmentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>scanner</td>
<td>&quot;level1&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;level2&quot;</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>&quot;level3&quot;</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>&quot;level4&quot;</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>&quot;level5&quot;</td>
<td>5</td>
</tr>
<tr>
<td>CTscanner</td>
<td>&quot;cluster_0...2&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;pixel_0...2&quot;</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>&quot;pixel_0...2&quot;</td>
<td>3</td>
</tr>
<tr>
<td>cylindricalPET</td>
<td>&quot;module&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;submodule&quot;</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>&quot;crystal&quot;</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>&quot;layer[i], i=0,3&quot;</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>&quot;layer[i], i=0,3&quot;</td>
<td>5</td>
</tr>
<tr>
<td>CPET</td>
<td>&quot;sector&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;cassette&quot;</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>&quot;module&quot;</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>&quot;crystal&quot;</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>&quot;layer[i], i=0,3&quot;</td>
<td>5</td>
</tr>
<tr>
<td>SPECThead</td>
<td>&quot;crystal&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;pixel&quot;</td>
<td>2</td>
</tr>
<tr>
<td>ecat</td>
<td>&quot;block&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;crystal&quot;</td>
<td>2</td>
</tr>
<tr>
<td>ecatAccel</td>
<td>&quot;block&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;crystal&quot;</td>
<td>2</td>
</tr>
<tr>
<td>OPET</td>
<td>&quot;sector&quot;</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>&quot;module&quot;</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>&quot;submodule&quot;</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>&quot;crystal&quot;</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>&quot;layer[i], i=0,7&quot;</td>
<td>5</td>
</tr>
</tbody>
</table>
Different types of systems

![Scanner System Illustration](image)

**Scanner**

**Description**

The *scanner* system is the most generic system in Gate. There is no geometrical constraints on the five different components.

**Use**

Different shapes of the volumes inside the tree level can be chosen, among those listed in table 3.2

Figure 4.4 illustrates the kind of detector that can be simulated with this system without any geometry constraint. On the other hand, there is no specific output format associated with this system and information regarding the hits are only available in ROOT or ASCII format.

**CTscanner**

The CTscanner system allows you to simulate a simple CT scanner. It has three possible levels:

- **module** component, that can be linearly repeated along the Y axis.
- **cluster** component, repeated inside the module, allowing you to simulate many kind of pixels
- **pixel** component, repeated inside the cluster. Raw data are the standard imageCT output to store the simulated CT projections and to produce it at each time slice. The image type is a float matrix (32-bits) of dimension given by the product of the number of pixels in X and Y, the content corresponds to the number of counts per pixel per acquisition (time slice).

Three types of simulations are proposed to the user:
Complete simulation: The modules, the clusters, and the pixels are user defined. All volumes are created by Geant4 and the digitalization can be made at the pixel level (level 3).

Fast simulation: Only the module level is defined. Geant4 creates one volume corresponding to the CT module (only one block possible) and the digitalization is made by the output module. The number of pixels per module are given through the output module messenger. This mode is faster since only one Geant4 volume is simulated, but obviously, only a rather approximated scanner response can be guaranteed.

Complete simulation with a Variance Reduction Technique (VRT): In the same way as the complete simulation, the components (pixels, clusters, and modules) are user defined. Unlike the complete simulation, using Geant4 in the detector, the user handles the particle on the surface of the CT scanner. For more informations see the part below.

This variance reduction technique (VRT) has been developed with the aim to making the simulation time faster. Here are the successive steps of the implementation:

- Generation and Propagation of the particles through the World, then detection of those on the surface of the detector. The propagation of the particles through the detector are 'killed', in order to handle ourself the detection and not by Geant4.
- Computation of the mean free path (MFP) of the particle through the detector with the standard model (the compatibility with the low energy model being not implemented yet)
- Computation of the path of the particle in the detector:

\[ \text{PATH} = \text{MFP} \times - \log(1 - R) \]
R being a distribution uniformly random number between 0 and 1

The user may perform this last step for each particle K times, in order to decrease the simulation time by avoiding a new generation and propagation of the particle. However it has an influence on the variance of the output data. The following scheme shows the differences with a simulation with and without VRT:

Without VRT:

\[ Q_{\text{source}} = N \pm \sqrt{N} \]
\[ p = e^{-px} \]
\[ Np = n \]
\[ Q_{\text{final}} = n \pm \sqrt{n} \]

With VRT:

\[ Q_{\text{source}} = N \pm \sqrt{N} \]
\[ p = e^{-px} \]
\[ NKp = n \]
\[ Q_{\text{final}} = n \pm n \sqrt{1/N \times (K-1/K) + 1/n} \]

K

N: mean of the generated particles, and \( \sqrt{N} \) its standard deviation.
p: binomial probability of detection of the particle.
n = Np: mean of the number of detected particles, and \( \sqrt{n} = \sqrt{Np} \) its standard deviation.

The simulation time decreases linearly with K. But K \( \geq 1/\)1, because of the reduction of the variance should be avoided. For deeper insight, see the following table and graph.
The simulation corresponding to the table is:

- A detector with 10,000 pixels (0.5x0.5x0.5 mm$^3$) in Silicon (Si)
- A monochromatic source (17.6 keV)
- A time of exposition of 1 second
Use

example 1: complete CT simulation

```plaintext
# CT SCANNER #
/gate/world/daughters/name CTscanner
/gate/world/daughters/insert box
/gate/CTscanner/placement/setTranslation 0.00 0.00 100 mm
/gate/CTscanner/geometry/setXLength 100 mm
/gate/CTscanner/geometry/setYLength 100 mm
/gate/CTscanner/geometry/setZLength 0.5 mm
/gate/CTscanner/setMaterial Air
/gate/CTscanner/vis/forceWireframe
/gate/CTscanner/vis/setColor white

# CTSCANNER #   ---->   # MODULE #
/gate/CTscanner/daughters/name module
/gate/CTscanner/daughters/insert box
/gate/module/geometry/setXLength 100 mm
/gate/module/geometry/setYLength 100 mm
/gate/module/geometry/setZLength 0.5 mm
/gate/module/setMaterial Air
/gate/module/vis/forceWireframe
/gate/module/vis/setColor white

# MODULE #   ---->   # CLUSTER_0 #
/gate/module/daughters/name cluster
/gate/module/daughters/insert box
/gate/cluster/geometry/setXLength 100 mm
/gate/cluster/geometry/setYLength 100 mm
/gate/cluster/geometry/setZLength 0.5 mm
/gate/cluster/setMaterial Air
/gate/cluster/vis/forceWireframe
```

```plaintext
# STANDARD DEVIATION vs MODEL #

- Standard Deviation (VRT=10)
- Model (VRT=10)
- Standard Deviation (VRT=5)
- Model (VRT=5)
- Standard Deviation (VRT=2)
- Model (VRT=2)
- Standard Deviation (VRT=0)
- Model (VRT=0)
```
example 2: complete CT simulation with VRT
In the same way as the complete simulation, the difference is the output (K = 5)

Exemple 3 : Fast CT simulation
CylindricalPET

Description

CylindricalPET is a PET system that can describe most of the small animal PET scanners. The main specificity of cylindricalPET is the possibility to record output data in the List Mode Format (LMF) developed by the Crystal Clear Collaboration. A complete description of LMF is can be found in Users Guide V6.2:Data output#LMF output.

A CylindricalPET is based on a cylindrical geometry, and consists of 5 hierarchic levels, from mother to daughter, as defined below:

- **world cylindricalPET** is defined as a cylinder in the world, with a non zero inner radius.
- **rsector** (depth=1) is defined as a box, and repeated with a ring repeater in cylindricalPET.
- **module** (depth=2) is a box inside rsector. It is repeated by a cubicarray repeater with no X repetition (repeatNumberX = 1). This level is optional.
- **submodule** (depth=3) is a box inside module. It is repeated by a cubicarray repeater with no X repetition (repeatNumberX = 1). This level is optional.
- **crystal** (depth=4) is a box inside submodule. It is repeated by a cubicarray repeater with no X repetition (repeatNumberX = 1).
- **layer** (depth=5) is a (or several, in the case of a phoswich system) radially arranged box(es) inside crystal. A repeater should not be used for layers, but the should be build them one by one in the macro. layer must be set as a sensible detector with the macro command:

```
/attachCrystalSD
```

The words in bold characters are dedicated. See also keywords in table 4.2.

Material of layer(s) must be the material of the detector, for instance LSO or BGO + GSO for a double layer phoswich system. Materials of other levels (crystals, submodules, modules, rsectors, cylindricalPET) can be anything else.

**IMPORTANT** : Visualization should help you build this geometry with no overlap. GATE performs a test for detecting volume overlap, but with a limited precision. This test is performed at the end of the initialization of Gate (see Users Guide V6.2:Getting started)

```
/run/initialize
/geometry/test/recursive_test
```

Users should carefully check that volumes are not bigger than the mother volume they are included in.

Use

An example of definition of a PET scanner following the CylindricalPET system structure is given below. The definition of the scanner should be performed at the beginning of the macro, before initializations.

```
# WORLD
/gate/world/geometry/setXLength 40 cm
/gate/world/geometry/setYLength 40. cm
/gate/world/geometry/setZLength 40. cm

# MOUSE
/gate/world/daughters/name mouse
/gate/world/daughters/insert cylinder
/gate/mouse/setMaterial Water
/gate/mouse/vis/setColor red
/gate/mouse/geometry/setRmax 18.5 mm
/gate/mouse/geometry/setRmin 0. mm
/gate/mouse/geometry/setHeight 68. mm

# CYLINDRICAL
/gate/world/daughters/name cylindricalPET
/gate/world/daughters/insert cylinder
/gate/cylindricalPET/setMaterial Water
/gate/cylindricalPET/setRmax 145 mm
/gate/cylindricalPET/geometry/setRmin 130 mm
/gate/cylindricalPET/geometry/setHeight 80 mm
/gate/cylindricalPET/vis/forceWireframe
```

CPET

This system was defined for the simulation of a CPET-like scanner (C-PET Plus, Philips Medical Systems, the Netherlands), with one ring of NaI crystal with a curved shape. For this scanner, a single level in addition to the system level is enough to describe the volume hierarchy.

Description

This system has the particularity to have cylindrical shaped sector components, based on the cylinder shape (see figure 4.3 and Users Guide V6.2: Defining a geometry for definitions), whereas these components are generally boxes in other systems.
Use

Described below is an example of code appropriate for modeling a one ring scanner of NaI crystal with a curved shape.

```
# BASE = CPET SYSTEM
/gate/world/daughters/name CPET
/gate/world/daughters/insert cylinder
/gate/CPET/setMaterial Air
/gate/CPET/geometry/setRmax 60 cm
/gate/CPET/geometry/setHeight 35.0 cm
/gate/CPET/vis/forceWireframe

# FIRST LEVEL = CRYSTAL
/gate/CPET/daughters/name crystal
/gate/CPET/daughters/insert cylinder
/gate/crystal/geometry/setRmax 47.5 cm
/gate/crystal/geometry/setRmin 45.0 cm
/gate/crystal/geometry/setHeight 25.6 cm
/gate/crystal/geometry/setPhiStart 0 deg
/gate/crystal/geometry/setDeltaPhi 60 deg

# REPEAT THE CURVE SECTOR INTO THE WHOLE RING
/gate/crystal/repeaters/insert ring
/gate/crystal/ring/setRepeatNumber 6

# CRYSTAL VOLUME IS MADE OF NAI
/gate/crystal/setMaterial NaI
/gate/crystal/vis/setColor green
```

The object `crystal` is then attached to its corresponding component in the CPET system (level 1: the sector level for CPET system - see previous section for details).

```
/gate/systems/CPET/sector/attach crystal
```

The crystals are set as sensitive detectors (see Users Guide V6.2: Attaching the sensitive detectors#The crystalSD).

```
/gate/crystal/attachCrystalSD
```

The digitizer part (see Users Guide V6.2: Digitizer and readout parameters#Digitizer modules) is made of the `adder` module and some blurring module (see Users Guide V6.2: Digitizer and readout parameters).

Ecat

Description

The ecat system is a simplified version of cylindricalPET and was named ecat because it is appropriate for modelling PET scanners from the
**ECAT** family, from CPS Innovations (Knoxville, TN, U.S.A.). Such scanners are based on the block detector principle, consisting in an array of crystals, typically 8 x 8 read by few photomultipliers, typically 4. The blocks are organized along an annular geometry to yield multi-ring detectors.

An example of macro with an ecat definition is provided in:

```plaintext
$GATEHOME/example_PET_Scanner/PET_Ecat_System.mac
```

The ecat system has only three hierarchical levels: one is for the entire detector (*base*), one for the block (*block*), and one for the crystals within the block (*crystal*).

In addition to the standard output modules (*ASCII* and *root*), two additional output modules are specifically associated to the *ecat* system, and correspond to sinogram formats. These are the *sinogram* and the *ecat7* output modules and are discussed in Users Guide V6.2:Data output#Sinogram output and Users Guide V6.2:Data output#Ecat7 output.

Use

Described below is an example of code for modeling a four block-ring scanner.

It has to be named after the selected system (*ecat* here) and is defined as a volume daughter of the *world*. It has a ring shape and should include all detectors (see Figure 4.6).

```plaintext
/gate/world/daughters/name ecat
/gate/world/daughters/insert cylinder
/gate/ecat/setMaterial Air
/gate/ecat/geometry/setRmax 442.0 mm
/gate/ecat/geometry/setRmin 412.0 mm
/gate/ecat/geometry/setHeight 155.2 mm
/gate/ecat/setTranslation 0.0 0.0 0.0 mm
```

The following commands set the size and the position of the first block within the base *ecat*. It is a rectangular parallelepiped and should include all crystals within a block. For a multiple block-ring system centered axially on the base *ecat*, the axial position of this first block should be set to zero (see Figure 4.7).

```plaintext
/gate/ecat/daughters/name block
/gate/ecat/daughters/insert box
/gate/block/placement/setTranslation 427.0 0.0 0.0 mm
/gate/block/geometry/setXLength 30.0 mm
/gate/block/geometry/setYLength 35.8 mm
/gate/block/geometry/setZLength 38.7 mm
/gate/block/setMaterial Air
```

---

The next commands set the size and the position of the first crystal within the block. For a crystal array centered on the block, the position of this first crystal should be at the center of the block (see Figure 4.8).

```
/gate/block/daughters/name crystal
/gate/block/daughters/insert box
/gate/crystal/placement/setTranslation 0.0 0.0 0.0 mm
/gate/crystal/geometry/setXLength 30.0 mm
/gate/crystal/geometry/setYLength 4.4 mm
/gate/crystal/geometry/setZLength 4.75 mm
/gate/crystal/setMaterial BGO
```

Finally, the crystal array is described. The sampling of the crystals within a block is defined, together with the size and the sampling of the crystal array within one block. The crystal array is centered on the position of the original crystal.

```
/gate/crystal/repeaters/insert cubicArray
/gate/crystal/cubicArray/setRepeatNumberX 1
/gate/crystal/cubicArray/setRepeatNumberY 8
/gate/crystal/cubicArray/setRepeatNumberZ 8
/gate/crystal/cubicArray/setRepeatVector 0. 4.45 4.80 mm
```

To create the full scanner, the rings have then to be defined. The following commands set the number of blocks per block-ring and the number of block-rings. Multiple block-ring systems will be centered axially on the axial position of the original block.

```
/gate/block/repeaters/insert linear
/gate/block/linear/setRepeatNumber 4
/gate/block/linear/setRepeatVector 0. 0. 38.8 mm
/gate/block/repeaters/insert ring
/gate/block/ring/setRepeatNumber 72
```

This description results in a 4 block-ring scanner, i.e. a 32 crystal-ring scanner, with 576 crystals per crystal-ring.

Command lines are then used to attach the objects block and crystal to their corresponding components in the ecat system.

```
/systems/ecat/block/attach block
/systems/ecat/crystal/attach crystal
```

To detect events, the crystals are finally set as sensitive detectors (see Users Guide V6.2:Attaching the sensitive detectors#The crystalSD).

```
/gate/crystal/attachCrystalSD
```

The digitizer part (see Users Guide V6.2:Digitizer and readout parameters#Digitizer modules) can be the same as for the cylindricalPET system.

**ecatAccel**

**Description**

The ecatAccel system was introduced to model a new PET scanner family ECAT ACCEL (from CPS Innovations, Knoxville, TN, U.S.A.). The ecatAccel system differs from the ecat system by its geometrical shape: the detection blocks are arranged along a spherical ring whereas they are arranged along annular rings for the ecat system. As data processing and output format are highly dependent on the scanner geometry, it was necessary to introduce a new system even though it has many common features with the ecat system. The same hierarchical levels (base,
block and crystal) as for the ecat system are used to describe the geometry of the ecatAccel system, and the same standard output modules (ASCII and root) and specific outputs (sinogram and ecat7) are also available. Please refer to [Users Guide V6.2:Data output#Sinogram output] and Users Guide V6.2:Data output#Ecat7 output for further information on sinogram and ecat7 outputs for the ecatAccel system.

Use

Described below is an example of code for modeling the ACCEL PET scanner of the BIOGRAPH-LSO (SIEMENS - CTI) PET-CT scanner.

The scanner is named after the selected system (ecatAccel here) and is defined as a volume daughter of the world. As for the ecat system, it has a ring shape and should include all detectors (see Figure 4.6). For the BIOGRAPH, it can be described as follows:

The base is described:

```
/gate/world/daughters/name ecatAccel
/gate/world/daughters/insert cylinder
/gate/ecatAccel/setMaterial Air
/gate/ecatAccel/geometry/setRmax 437.0 mm
/gate/ecatAccel/geometry/setRmin 412.0 mm
/gate/ecatAccel/geometry/setHeight 162. mm
/gate/ecatAccel/setTranslation 0.0 0.0 0.0 mm
```

The block is then described: the size and the position of the first block are set within the base ecatAccel. As for the ecat system, it is a rectangular parallelepiped and should include all crystals within a block. For a multiple block-ring system centered axially on the base ecatAccel, the axial position of this first block should be set to zero.

```
/gate/ecatAccel/daughters/name block
/gate/ecatAccel/daughters/insert box
/gate/block/geometry/setXLength 51.6 mm
/gate/block/geometry/setYLength 25.0 mm
/gate/block/geometry/setZLength 51.6 mm
/gate/block/setMaterial Air
```

The crystal geometry and settings are then specified. The following commands set the size and the position of the first crystal within the block. For a crystal array centered on the block, the position of this first crystal should be at the center of the block.

```
/gate/block/daughters/name crystal
/gate/block/daughters/insert box
/gate/crystal/placement/setTranslation 0.0 0.0 0.0 mm
/gate/crystal/geometry/setXLength 6.45 mm
/gate/crystal/geometry/setYLength 25.0 mm
/gate/crystal/geometry/setZLength 6.45 mm
/gate/crystal/setMaterial LSO
```

Then the crystal array is described within a block. The crystal array will be centered on the position of the original crystal.

```
/gate/crystal/repeaters/insert cubicArray
/gate/crystal/cubicArray/setRepeatNumberX 8
/gate/crystal/cubicArray/setRepeatNumberY 1
/gate/crystal/cubicArray/setRepeatNumberZ 8
/gate/crystal/cubicArray/setRepeatVector 6.45 0.0 6.45 mm
```

Finally, the different rings of the scanner are described. The number of blocks per block-ring (command setRepeatNumberWithTheta) is indicated as well as the number of block-rings (command setRepeatNumberWithPhi). The angle between two adjacent blocks in a block-ring is set with the command setThetaAngle and the angle between two adjacent blocks belonging to two neighbouring rings in the axial direction is set with the command setPhiAngle. Multiple block-ring will be centered axially on the axial position of the original block.

```
/gate/block/repeaters/insert sphere
/gate/block/sphere/setRadius 424.5 mm
/gate/block/sphere/setRepeatNumberWithTheta 3
/gate/block/sphere/setRepeatNumberWithPhi 48
/gate/block/setThetaAngle 7.5 deg
/gate/block/setPhiAngle 7.5 deg
```

This description results in a 3 block-ring scanner, i.e. a 24 crystal-ring scanner, with 384 crystals per crystal-ring.

The objects block and crystal are attached to their corresponding components in the ecatAccel system:

```
/gate/systems/ecatAccel/block/attach block
/gate/systems/ecatAccel/crystal/attach crystal
```
The sensitive detector is set to the crystals as for the ecat system and the digitizer part remains the same as for the cylindricalPET system.

**OPET**

**Description**

The OPET system was introduced to model a new PET prototype.

**Use**

Described below is an example of code for modeling the OPET PET scanner.

```plaintext
/* R E C T O R (Create a box to put the crystals in: one PMT) */
gate/OPET/daughters/name rsector
/gate/OPET/daughters/insert box
/gate/rsector/placement/setTranslation 20.4955 0 0 mm
/gate/rsector/geometry/setXLength 10 mm
/gate/rsector/geometry/setYLength 17.765 mm
/gate/rsector/geometry/setZLength 17.765 mm
/gate/rsector/setMaterial Air
/gate/rsector/vis/setVisible False
/gate/rsector/vis/forceWireframe
/gate/rsector/vis/setColor yellow

/* M O D U L E (Make a box for one row of 8 crystals) */
gate/rsector/daughters/name module
/gate/rsector/daughters/insert box
/gate/module/geometry/setXLength 10 mm
/gate/module/geometry/setYLength 17.765 mm
/gate/module/geometry/setZLength 2.162 mm
/gate/module/setMaterial Air
/gate/module/vis/setVisible False
/gate/module/vis/forceWireframe
/gate/module/vis/setColor black

/* Daughter crystal inside mother crystal */
gate/module/daughters/name crystal0
/gate/module/daughters/insert box
/gate/crystal0/geometry/setXLength 10 mm
/gate/crystal0/geometry/setYLength 2.1620 mm
/gate/crystal0/geometry/setZLength 2.1620 mm
/gate/crystal0/placement/setTranslation 0. -7.8015 0. mm
/gate/crystal0/setMaterial Air
/gate/crystal0/vis/setColor black
/gate/crystal0/vis/setVisible false

/* L A Y E R (Put the LSO in the small box) */
gate/crystal0/daughters/name LSO0
/gate/crystal0/daughters/insert wedge
/gate/LSO0/geometry/setXLength 10 mm
/gate/LSO0/geometry/setYLength 8.921 mm
/gate/LSO0/geometry/setZLength 2.1620 mm
/gate/LSO0/placement/setRotationAxis 0 1 0
/gate/LSO0/placement/setRotationAngle 180 deg
/gate/LSO0/placement/setTranslation 0.2698 0. 0. mm
/gate/LSO0/setMaterial BGO
/gate/LSO0/vis/setColor yellow

/* Daughter crystal inside mom crystal */
gate/module/daughters/name crystal1
/gate/module/daughters/insert box
/gate/crystal1/geometry/setXLength 10 mm
/gate/crystal1/geometry/setYLength 2.1620 mm
/gate/crystal1/geometry/setZLength 2.1620 mm
/gate/crystal1/placement/setTranslation 0. -5.5725 0. mm
/gate/crystal1/setMaterial Air
/gate/crystal1/vis/forceWireframe
/gate/crystal1/vis/setVisible false

/* L A Y E R (Put the LSO in the small box) */
gate/crystal1/daughters/name LSO1
/gate/crystal1/daughters/insert wedge
/gate/LSO1/geometry/setXLength 8.193 mm
/gate/LSO1/geometry/setYLength 8.921 mm
/gate/LSO1/geometry/setZLength 2.1620 mm
/gate/LSO1/placement/setRotationAxis 0 1 0
/gate/LSO1/placement/setRotationAngle 180 deg
/gate/LSO1/placement/setTranslation 0.7215 0. 0. mm
/gate/LSO1/setMaterial BGO
/gate/LSO1/vis/setColor red
```

Daughter crystal inside mom crystal
/gate/module/daughters/name crystal2
/gate/module/daughters/insert box
/gate/crystal2/geometry/setXLength 10 mm
/gate/crystal2/geometry/setYLength 2.1620 mm
/gate/crystal2/geometry/setZLength 2.1620 mm
/gate/crystal2/placement/setTranslation -3.3435 0. 0. mm
/gate/crystal2/setMaterial Air
/gate/crystal2/vis/setColor black
/gate/crystal2/vis/setVisible false

# L A Y E R (Put the LSO in the small box)
/gate/crystal2/daughters/name LSO2
/gate/crystal2/daughters/insert wedge
/gate/LSO2/geometry/setXLength 8.193 mm
/gate/LSO2/geometry/setNarrowerXLength 7.773 mm
/gate/LSO2/geometry/setYLength 2.1620 mm
/gate/LSO2/geometry/setZLength 2.1620 mm
/gate/LSO2/placement/setRotationAngle 180 deg
/gate/LSO2/placement/setTranslation 1.0085 0. 0. mm
/gate/LSO2/setMaterial BGO
/gate/LSO2/vis/setColor green

Daughter crystal inside mom crystal
/gate/module/daughters/name crystal3
/gate/module/daughters/insert box
/gate/crystal3/geometry/setXLength 10 mm
/gate/crystal3/geometry/setYLength 2.1620 mm
/gate/crystal3/geometry/setZLength 2.1620 mm
/gate/crystal3/placement/setTranslation -1.1145 0. 0. mm
/gate/crystal3/setMaterial Air
/gate/crystal3/vis/forceWireframe
/gate/crystal3/vis/setColor black
/gate/crystal3/vis/setVisible false

# L A Y E R (Put the LSO in the small box)
/gate/crystal3/daughters/name LSO3
/gate/crystal3/daughters/insert wedge
/gate/LSO3/geometry/setXLength 7.773 mm
/gate/LSO3/geometry/setNarrowerXLength 7.637 mm
/gate/LSO3/geometry/setYLength 2.1620 mm
/gate/LSO3/geometry/setZLength 2.1620 mm
/gate/LSO3/placement/setRotationAxis 0 0 1
/gate/LSO3/placement/setRotationAngle 180 deg
/gate/LSO3/placement/setTranslation 1.1475 0. 0. mm
/gate/LSO3/setMaterial BGO
/gate/LSO3/vis/setColor blue

Daughter crystal inside mom crystal
/gate/module/daughters/name /gate/crystal4/
/gate/module/daughters/insert box
/gate/crystal4/geometry/setXLength 10 mm
/gate/crystal4/geometry/setYLength 2.1620 mm
/gate/crystal4/geometry/setZLength 2.1620 mm
/gate/crystal4/placement/setTranslation 1.1145 0. 0. mm
/gate/crystal4/setMaterial Air
/gate/crystal4/vis/setColor black
/gate/crystal4/vis/setVisible false

# L A Y E R (Put the LSO in the small box)
/gate/crystal4/daughters/name /gate/LSO4
/gate/crystal4/daughters/insert wedge
/gate/LSO4/geometry/setXLength 7.773 mm
/gate/LSO4/geometry/setNarrowerXLength 7.637 mm
/gate/LSO4/geometry/setYLength 2.1620 mm
/gate/LSO4/geometry/setZLength 2.1620 mm
/gate/LSO4/placement/setRotationAxis 0 0 1
/gate/LSO4/placement/setRotationAngle 180 deg
/gate/LSO4/placement/setTranslation 1.1475 0. 0. mm
/gate/LSO4/setMaterial BGO
/gate/LSO4/vis/setColor blue

Daughter crystal inside mom crystal
/gate/module/daughters/name crystal5
/gate/module/daughters/insert box
/gate/crystal5/geometry/setXLength 10 mm
/gate/crystal5/geometry/setYLength 2.1620 mm
/gate/crystal5/geometry/setZLength 2.1620 mm
/gate/crystal5/placement/setTranslation 3.3435 0. 0. mm
/gate/crystal5/setMaterial Air
/gate/crystal5/vis/setColor black
/gate/crystal5/vis/setVisible false

# L A Y E R (Put the LSO in the small box)
/gate/crystal5/daughters/name LSO5
/gate/crystal5/daughters/insert wedge
/gate/LSO5/geometry/setXLength 8.193 mm
/gate/LSO5/geometry/setNarrowerXLength 7.773 mm
/gate/LSO5/geometry/setYLength 2.1620 mm
/gate/LSO5/geometry/setZLength 2.1620 mm
/gate/LSO5/placement/setRotationAxis 0 0 1
Figure 4.9 shows the final OPET scanner.
Figure 4.9: The OPET scanner

SPECTHead

Description

SPECTHead is a SPECT system appropriate to model SPECT dedicated scanners within GATE. The main reason for specifying SPECTHead is that it can be coupled to the InterFile output which is discussed in [Users Guide V6.2:Data output#Interfile output]. An example macro defining a typical SPECT scanner can be found in:

```
$GATEHOME/exampleSPECT_Scanners/Interfile.mac
```

wherein the specific Interfile output module is called.

A SPECTHead system is a box-shaped geometry element and consists of three hierarchic levels:

- **base** which is always attached to the volume SPECTHead, which is a dedicated word.
- **crystal** which is coupled to the main detector block.
- **pixel** which can be used for modeling a pixelated detector.

If a uniform detector block is being used, then the **crystal** material should be that of the detector. If the detector is pixelated, then the **pixel** material definition should correspond to the detector material, while the crystal material can be anything non-specific.

Use

Below is part of the SPECT benchmark macro, which is distributed with the GATE software, and which involves the SPECTHead system.

```
# World
# Define the world dimensions
/gate/world/ dimensions
/gate/world/geometry/setXLength 100 cm
/gate/world/geometry/setYLength 100 cm
/gate/world/geometry/setZLength 100 cm

# SPECThead is the name of the predefined SPECT system
```

Create the SPECT system, which will yield an Interfile output of the projection data:

```plaintext
/gate/world/daughters/name /gate/SPECThead
```

Define the dimensions:

```plaintext
/gate/SPECThead/geometry/setXLength 7. cm
/gate/SPECThead/geometry/setYLength 21. cm
/gate/SPECThead/geometry/setZLength 30. cm
```

Define the position:

```plaintext
/gate/SPECThead/placement/setTranslation 20.0 0.0 0.0 cm
```

Set the material associated with the main volume:

```plaintext
/gate/SPECThead/setMaterial Air
```

Replicate the head (around the Z axis by default) to get a hypothetical four-headed system:

```plaintext
/gate/SPECThead/repeaters/insert ring
/gate/SPECThead/ring/setRepeatNumber 4
/gate/SPECThead/ring/setAngularPitch 90. deg
```

Define the rotation speed of the head and the orbiting around the Z axis:

```plaintext
/gate/SPECThead/moves/insert orbiting
/gate/SPECThead/orbiting/setSpeed 0.15 deg/s
/gate/SPECThead/orbiting/setPoint1 0 0 0 cm
/gate/SPECThead/orbiting/setPoint2 0 0 1 cm
```

Define visualisation options:

```plaintext
/gate/SPECThead/vis/forceWireframe
```

Collimator:

Create a full volume defining the shape of the collimator (typical for SPECT):

```plaintext
/gate/SPECThead/daughters/name /gate/collimator
/gate/SPECThead/daughters/insert box
```

Define the dimensions of the collimator volume:

```plaintext
/gate/collimator/geometry/setXLength 3. cm
/gate/collimator/geometry/setYLength 19. cm
/gate/collimator/geometry/setZLength 28. cm
```

Define the position of the collimator volume:

```plaintext
/gate/collimator/placement/setTranslation -2.0 0.0 0.0 cm
```

Set the material of the collimator volume:

```plaintext
/gate/collimator/setMaterial Lead
```

Define some visualisation options:

```plaintext
/gate/collimator/vis/setColor red
/gate/collimator/vis/forceWireframe
```

Insert the first hole of air in the collimator:

```plaintext
/gate/collimator/daughters/name /gate/hole
/gate/collimator/daughters/insert hexagon
/gate/hole/geometry/setHeight 3. cm
/gate/hole/geometry/setRadius .15 cm
```

Figure 4.10: Example of a hypothetical four-headed SPECThead system. The detectors are not pixelated in this example.
Modelling the collimator

SPECT systems need collimator. A parameterized collimator setup was developed for both parallel hole collimators and fan beam collimators. It is based on the GEANT4 replica system in which a single volume represents multiple copies of a volume (the air holes) within its mother volume (the collimator itself). SPECT collimator geometries are built using this approach in less than a second.

Example of code for modelling fanbeam collimators:

```/gate/SPECThead/daughters/name fanbeam
/gate/SPECThead/daughters/insert collimator

 set the material for the collimator
/gate/fanbeam/setMaterial Lead

 define the X and Y size of the collimator
/gate/fanbeam/geometry/setDimensionY 53.5 cm
/gate/fanbeam/geometry/setDimensionX 25.0 cm

 specify the focal length
/gate/fanbeam/geometry/setFocalDistanceY 0.0 cm
/gate/fanbeam/geometry/setFocalDistanceX 35.0 cm

 specify the thickness of the collimator
/gate/fanbeam/geometry/setHeight 5.8 cm

 set the septal thickness to the required distance between the holes
/gate/fanbeam/geometry/setSeptalThickness 0.8 cm

 specify the hole radius
/gate/fanbeam/geometry/setInnerRadius 1.70 cm
/gate/fanbeam/placement/setRotationAxis 0.0 1.0
/gate/fanbeam/placement/setRotationAngle -90 deg
/gate/fanbeam/vis/setColor blue
/gate/fanbeam/vis/forceWireframe```

Example for parallel hole collimators:

```/gate/SPECThead/daughters/name colli

 specify that the parallel beam collimator setup must be used
/gate/SPECThead/daughters/insert parallelbeam

 set the collimator material```
Septal Penetration

If one wants to record, for every photon detected, how many times they crossed the collimator septa, the command recordSeptalPenetration must be turned on (default value is false) and the septal volume name must be attached to a PhantomSD:

```
/gate/output/analysis/recordSeptalPenetration true
```

If the septal volume name does not exist, the simulation is aborted.

OpticalSystem

Description

OpticalSystem is appropriate to model Optical Imaging within GATE. An example macro defining a typical Optical Imaging system can be found in:

```
$GATEHOME/example_OPTICAL/Optical_System.mac
```

An OpticalSystem is a box-shaped geometry element and consists of three hierarchic levels:

- **base** which is always attached to the volume OpticalSystem.
- **crystal** which is coupled to the main detector block.
- **pixel** which can be used for modeling a pixelated detector.

Use

Below is part of the Optical Imaging benchmark macro, which is distributed with the GATE software.

```
# World
# Define the world dimensions
/gate/world/geometry/setXLength 100. cm
/gate/world/geometry/setYLength 100. cm
/gate/world/geometry/setZLength 100. cm
/gate/world/setMaterial Air

# Create the Optical Imaging system, which will yield a binary output of the projection data
/gate/world/daughters/name OpticalSystem
/gate/world/daughters/insert box

# Define the dimensions, position and material
/gate/OpticalSystem/geometry/setXLength 10.5 cm
/gate/OpticalSystem/geometry/setYLength 10.5 cm
/gate/OpticalSystem/geometry/setZLength 2.0 cm
/gate/OpticalSystem/placement/setTranslation 0 0 0 cm
/gate/OpticalSystem/setMaterial Air

# Define pixelated detector:
/gate/OpticalSystem/daughters/name crystal
/gate/OpticalSystem/daughters/insert box
/gate/crystal/geometry/setXLength 10.5 cm
/gate/crystal/geometry/setYLength 10.5 cm
/gate/crystal/geometry/setZLength 1.0 cm
/gate/crystal/placement/setTranslation 0 0 -0.5 cm
/gate/crystal/setMaterial Air
/gate/crystal/vis/forceWireframe 1
```

Figure 4.11: Example of a hypothetical OpticalImaging pixelated system. A pixelated camera is simulated in red. The additional volume in yellow could represent some electronic board.
How to define a multi-system detector

To simulate a multi-system device consisting of several detectors (PET, SPECT, CT, ...) you need to add in your macro special commands as explained below. This will allow you to simultaneously register Hits inside every detector.

Defining the systems:

The standard definition of a GATE system is done according to the command:

```
/gate/world/daughters/name  SystemName
```

Where SystemName must be one of available system names in GATE (see table 4.1). Unfortunately, defining a system with this command prevent you to insert more than one system of the same type. Another method has been inserted in GATE to define more than one system at a time. Using this more general method, users can simulate several systems simultaneously. A system is now defined by its own name and its type according to the next two command lines:

```
/gate/world/daughters/name  AnyName
/gate/world/daughters/systemType  SystemType
```

Where AnyName can be any name as for any Geant4 volume name and SystemType must be one the names of GATE systems mentioned in table 4.1.

How to connect the geometry to the systems:

By using more than one system, we have to change the attachment commands to connect the geometrical elements of every system with its defined components, to do that we use the next command:

```
/gate/systems/SystemName/Level/attach  UserVolumeName
```

This command has the same form as for one system, but the essential difference is that SystemName here can be any name gave by the user to the system.

Example of multi-system:

An example for the creation of three systems, one system of type “cylindricalPET” and two systems of type “scanner” is explained below. Note that it is not necessary to use the “ systemType” command for cylindricalPET system because there is only one system of this type.
# MOUSE
/gate/world/geometry/setZLength 60. cm

# SYSTEM 1: cylindricalPET
/gate/world/daughters/name cylindricalPET   # standard definition
/gate/world/daughters/insert cylinder
/gate/cylindricalPET/setMaterial Water
/gate/cylindricalPET/geometry/setRmax 145 mm
/gate/cylindricalPET/geometry/setRmin 130 mm
/gate/cylindricalPET/geometry/setHeight 80 mm
/gate/cylindricalPET/vis/forceWireframe

cylindricalPET => rsector
/gate/cylindricalPET/daughters/name rsector
/gate/rsector/placement/setTranslation 135 0 0 mm
/gate/rsector/geometry/setXLength 10. mm
/gate/rsector/geometry/setYLength 19. mm
/gate/rsector/geometry/setZLength 76.6 mm
/gate/rsector/setMaterial Water
/gate/rsector/vis/forceWireframe

cylindricalPET => module
/gate/rsector/daughters/name module
/gate/rsector/daughters/insert box
/gate/module/geometry/setXLength 10. mm
/gate/module/geometry/setYLength 19. mm
/gate/module/geometry/setZLength 19. mm
/gate/module/setMaterial Water
/gate/module/vis/forceWireframe
/gate/module/vis/setColor gray

cylindricalPET => crystal
/gate/module/daughters/name crystal
/gate/module/daughters/insert box
/gate/crystal/geometry/setXLength 10. mm
/gate/crystal/geometry/setYLength 2.2 mm
/gate/crystal/geometry/setZLength 2.2 mm
/gate/crystal/setMaterial Water
/gate/crystal/vis/forceWireframe
/gate/crystal/vis/setColor magenta

cylindricalPET => LSO
/gate/crystal/daughters/name LSO
/gate/crystal/daughters/insert box
/gate/LSO/geometry/setXLength 10. mm
/gate/LSO/geometry/setYLength 2.2 mm
/gate/LSO/geometry/setZLength 2.2 mm
/gate/LSO/placement/setTranslation 0 0 0 mm
/gate/LSO/setMaterial LSO
/gate/LSO/vis/setColor yellow

# REPEAT CRYSTAL
/gate/crystal/repeaters/insert cubicArray
/gate/crystal/cubicArray/setRepeatNumberX 1
/gate/crystal/cubicArray/setRepeatNumberY 8
/gate/crystal/cubicArray/setRepeatNumberZ 8
/gate/crystal/cubicArray/setRepeatVector 10. 2.4 2.4 mm

# REPEAT MODULE
/gate/module/repeaters/insert cubicArray
/gate/module/cubicArray/setRepeatNumberX 4
/gate/module/cubicArray/setRepeatNumberY 8
/gate/module/cubicArray/setRepeatVector 0. 0. 19.2 mm

# REPEAT RSECTOR
/gate/rsector/repeaters/insert ring
/gate/rsector/ring/setRepeatNumber 42

# SYSTEM 2: Scanner_1
/gate/world/daughters/name Scanner_1       # System name definition for scanner 1
/gate/world/daughters/systemType scanner   # System type definition for scanner 1
/gate/world/daughters/insert box
/gate/Scanner_1/setMaterial Air
/gate/Scanner_1/geometry/setXLength 20 cm
/gate/Scanner_1/geometry/setYLength 20 cm
/gate/Scanner_1/geometry/setZLength 10 cm
/gate/Scanner_1/placement/setTranslation 0 0 -18 cm
/gate/Scanner_1/vis/forceWireframe
/gate/Scanner_1/vis/setVisible 1

Scanner_1 => Cryostat_1
/gate/Scanner_1/daughters/name Cryostat_1
/gate/Scanner_1/daughters/insert box
Notes:

a) The command “systemType” is optional in case of using only one system, but the system name must be one GATE systems (first column in table 4.1) as for standard definition.

b) Same remark, in the case where all systems have different types.

c) In general, one has to use the “systemType” command only for simulating more than one system of the same type.

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Users Guide V7.0: Attaching the sensitive detectors

From GATE collaborative documentation wiki

Table of Contents

- General purpose
- The two types of sensitive detector
  - The crystalSD
  - The phantomSD

General purpose

Once a model has been defined for the scanner through the construction of a system (see Users Guide V6.2: Defining a system), the next step is to attach a sensitive detector (SD) to some volumes of the geometry. As in any Geant4 simulation, these sensitive detectors are used to store information regarding interactions of a particle in the matter (hits) using information from the steps occurring along the particle track. A hit is a snapshot of a physical interaction of a track in a sensitive region of the detector. Figure 6.1 illustrates these notions. Hits contain various pieces of information associated to a step object, such as the energy deposition of a step, geometrical information, position and time of a step, etc.

![Figure 5.1: Particle interactions in a sensitive detector](image)

It is essential to remember that GATE records and stores information related to the hits only for those
volumes that are attached to a sensitive detector. All information regarding the interactions occurring in non-sensitive volumes is lost.

Two sensitive detectors are defined in GATE:

- **The crystalSD** is used to record information regarding interactions inside the volumes belonging to a scanner for instance (crystals or collimators).
- **The phantomSD** is used to record information regarding Compton and Rayleigh interactions taking place in the volumes before the detection in the scanner system (e.g., for a SPECT camera, these volumes can be the table, phantom, and collimator, in which it can be relevant to retrieve information about Compton and Rayleigh interactions).

A complete definition of the simulation context normally involves performing the two sorts of attachments: some volumes are attached to the **phantomSD**, while others are attached to the **crystalSD**.

**The two types of sensitive detector**

**The crystalSD**

**Definition and use**

The **crystalSD** may be used to record information regarding interactions taking place inside some volumes of the scanner: energy deposition, positions of interaction, origin of the particle (emission vertex), type of interaction (name of the physical processes involved), etc.

**Attachment of the crystalSD**

A **crystalSD** can be attached only to those volumes that belong to a given system. Once a **crystalSD** has been attached, it is considered as attached to this system. This sensitive detector can be attached using the command **attachCrystalSD**. These volumes are essentially meant to be scintillating elements (crystals) but can also be attached to non-scintillating elements such as collimators, shields or septa.

Below is an example of command lines that should be included in a macro using the **crystalSD**. These command lines must be inserted after the description of the attachment to the system:

The first command is used to attach the scintillation crystal to the detection level **crystal** of the SPECThead system.

```
# ATTACH SYSTEM
/systems/SPECThead/crystal/attach crystal
```

Then, the second command attaches the crystalSD to the volume representing the scintillation crystal in the geometry.

```
# ATTACH SENSITIVE DETECTOR
/gate/crystal/attachCrystalSD
```
The phantomSD

Definition and use

The phantomSD plays a crucial role in GATE simulations, as it is used to detect and tally Compton and Rayleigh interactions taking place in the scanner FOV. These data can then be used to estimate whether a photon reaching a detector is a direct or a Compton-scattered photon. Thus, in PET, the phantomSD is currently the only way to discriminate scattered from true coincidences. To simulate low energy X-ray acquisitions (for example mammography acquisitions from 7 to 28 keV), information concerning Rayleigh interactions is significant.

Using this type of sensitive detector, it is possible to retrieve two pieces of information relating to the hits:

- The number of Compton and Rayleigh interactions occurring in all the volumes attached to the phantomSD: this is stored in the data output variables nPhantomCompton and nPhantomRayleigh. These pieces of information are also available for the crystalSD with the variables nCrystalCompton and nCrystalRayleigh.
- The last volume attached to the phantomSD in which a Compton or a Rayleigh interaction occurred: the data output variables including these volume names are compVolName and RayleighVolName.

Attachment of the phantomSD

- One first needs to define a dummy, air-filled volume covering the whole field-of-view of the scanner.
- Then, all the source volumes should be offsprings (direct or indirect) of this volume.
- Last, all these volumes (FOV and sources) should be attached to the phantomSD using the command attachPhantomSD.

IMPORTANT: To retrieve data output information regarding hits occurring in the phantomSD (nPhantomCompton and compVolName), a crystalSD has to be defined in the simulation. Otherwise, data output variables will be created but will be empty. When all these conditions are satisfied, any interaction taking place within the FOV of the scanner is automatically recorded by the phantomSD, so that the number of Compton interactions for each photon can be accurately computed.

This procedure does not take into account Compton interactions taking place within the detectors, so that inter-crystal cross-talk via Compton interactions is not detected. Here is an example of command-lines that should be included within the macro in order to use the phantomSD. These command lines must be inserted after the description of the attachment to the system. First, commands are used to attach the scattering volumes to the detection level base of the SPECTHead system:

```plaintext
# ATTACH SYSTEM
/systems/SPECTHead/base/attach FOV
/systems/SPECTHead/base/attach head
/systems/SPECTHead/base/attach body
```

Then, commands attach the phantomSD to the volumes representing the scattering volumes in the geometry:

```plaintext
# ATTACH SENSITIVE DETECTOR
/FOV/attachPhantomSD
/head/attachPhantomSD
/body/attachPhantomSD
```
Finally, the last commands are used to attach the scintillation crystal to the detection level crystal of the SPECThead system and to attach the crystalSD to the volume representing the scintillation crystal in the geometry.

```plaintext
# ATTACH SYSTEM AND SENSITIVE DETECTOR CRYSTALSD IN ORDER TO RETRIEVE DATA OUTPUTS ON PHANTOMSD
/systems/SPECThead/crystal/attachCrystalSD
/gate/crystal/attachCrystalSD
```

**IMPORTANT:**

It is impossible to attach two sensitive detectors to the same volume. Thus, to count also the Compton interactions occurring in the scintillating crystal, the variable `nCrystalCompton` has been created: its role is similar to that of the variable `nPhantomCompton`, e.g., it stores the number of Compton interactions in the scintillating crystal.

**In the case of a voxellized matrix:** Previous commands to attach sensitive detectors are used for the volumes created using the geometry commands of GATE (see Users Guide V6.2: Defining a geometry). In order to record the same information concerning the interactions occurring in a voxellized matrix, see Users Guide V6.2: Voxelized Source and Phantom.

Retrieved from

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Users Guide V7.0: Digitizer and readout parameters

From GATE collaborative documentation wiki

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Multi-system approaches: how to use more than one system in one simulation set-up?

General Purpose

The purpose of the digitizer module is to simulate the behavior of the scanner detectors and signal processing chain. In this section, the algorithms used to simulate a scanner electronic readout scheme are described. In the case of PET, an overview of the main steps used to produce coincidences from the simulated particle information is briefly described. A more detailed explanation of how to control the behavior of each of these steps is given. Finally, a complete example of a readout macro file is presented.

From particle detection to coincidences in GATE

GATE uses Geant4 to generate particles and transport them through the different materials. This mimics the physical interactions between particles and matter. The information generated during this process is used by GATE to simulate the detector pulses (digits), which correspond to the observed data. The digitizer represents the series of steps and filters that make up this process.

The typical data-flow for an event is as follows:

- A particle is generated, with its parameters, such as initial type, time, momentum, and energy.
- An elementary trajectory step (referred to in Geant4 simply as a step) is applied. A step corresponds to the trajectory of a particle between discrete interactions (i.e. photoelectric, Compton, pair production, etc). During a step the changes to particle's energy and momentum are calculated. The length of a step depends upon the nature of interaction, the type of particle and material, etc. The calculation of step length is complex and is mentioned here only briefly. For more details, please refer to the Geant4 documentation.
- If a step occurs within a volume corresponding to a sensitive detector, the interaction information between the particle and the material is stored. For example, this information may include the deposited energy, the momentum before and after the interaction, the name of the volume where the interaction occurred, and so on. This set of information is referred to as a Hit.
- Steps 2 and 3 are repeated until the energy of the particle becomes lower than a predefined value, or the particle position goes outside the predefined limits. The entire series of steps form a simulated trajectory of a particle, that is called a Track in Geant4.
- The amount of energy deposited in a crystal is filtered by the digitizer module. The output from the digitizer corresponds to the signal after it has been processed by the Front End Electronics (FEE). Generally, the FEE is made of several processing units, working in a serial and/or in parallel. This process of transforming the energy of a Hit into the final digital value is called Digitization, and is performed by the digitizer portion of the GATE architecture. Each processing unit in the FEE is represented in GATE by a corresponding digitizer module. The final value obtained after filtering by a set of these modules is called a Single. Singles can be saved as output. Each transient value, between two modules, is called a Pulse.

This process is repeated for each event in the simulation in order to produce one or more sets of Singles. These Singles can be stored into an output file (as a ROOT tree, for example).

Once this list is created, a second processing stage can be inserted to sort the Singles list for coincidences in case of PET systems. To do this, the algorithm searches in this list for a set of Singles that are detected within a given time interval (the so called 'coincident events').
Finally, the coincidence data may be filtered-out to mimic any possible data loss which could occur in the coincidence logical circuit or during the data transportation. As for the singles, the processing is performed by specifying a list of generic modules to apply to the coincidence data flow.

**Definition of a hit in Geant4**

A hit is a snapshot of the physical interaction of a track within a sensitive region of a detector. The information given by a hit is:

- Position and time of the step
- Momentum and energy of the track
- Energy deposition of the step
- Interaction type of the hit
- Volume name containing the hit

As a result, the history of a particle is saved as a series of hits generated along the particles trajectory. In addition to the physical hits, Geant4 saves a special hit. This hit takes place when a particle moves from one volume to another (this type of hit deposits zero energy). The hit data represents the basic information that a user has with which to construct the physically observable behavior of a scanner. To see the information stored in a hit, see the file GateCrystalHit.hh.

**Role of the digitizer**

As mentioned above, the information contained in the hit does not correspond to what is provided by a real detector. To simulate the digital values (pulses) that result from the output of the Front End Electronics, the sampling methods of the signal must be specified. To do this, a number of digitizer modules are available and are described below. Moreover, in the case of PET analysis, the trigger logic is based on one or more decisions defined by the user that depend upon physically observable quantities such as energy thresholds and coincidence times.

The role of the digitizer is to build, from the hit information, the physical observables, which include energy, position, and time of detection for each particle. In addition, the digitizer must implement the required logic to simulate coincidences during PET simulations. Typical usage of digitizer module includes the following actions:

- simulate detector response
- simulate readout scheme
- simulate trigger logic

These actions are accomplished by inserting digitizer modules into GATE, as explained in the next sections.

**Disabling the digitizer**

If for any reason you want to disable the digitizer process and all output (that are already disabled by default), you can use the 2 following commands:

```
/gate/output/analysis/disable
/gate/output/digi/disable
```
Digitizer modules

The digitization consists of a series of signal processors. The output at each step along the series is defined as a pulse. At the end of the chain, the output pulses are named singles. These Singles realistically simulate the physical observables of a detector response to a particle interacting with it. An example is shown in figure 8.1.

![Diagram](https://via.placeholder.com/150)

**Figure 8.1:** The digitizer is organized as a chain of modules that begins with the hit and ends with the single which represents the physical observable seen from the detector.

To specify a new signal-processing module i.e. add a new processing unit in the readout scheme, the following command template should be used:

```
/gate/digitizer/insert MODULE
```

where MODULE is the name of the digitizer module. The order of the module declaration should make sense. The data flow follows the same order as the module declaration in the macro. In a typical scanner, the following sequence works well, although it is not mandatory (the module names will be explained in the rest of the section):

- insert adder before readout
- insert readout before thresholder/upholder
- insert blurring before thresholder/upholder

The available modules are explained in the following sections.

Distributions

Definition

Since many of the modules presented below have to deal with functions or probability density, a generic tool is provided to describe such mathematical objects in GATE. Basically, a distribution in GATE is defined by its name, its type (Gaussian, Exponential, etc...) and the parameters specifics to each distribution type (such as the mean and the standard deviation of a Gaussian function). Depending on the context, these objects are used directly as functions, or as probability densities into which a variable is randomly chosen. In the following, the generic term of distribution will be used to describe both of these objects, since their declaration is unified under this term into GATE.
Five types of distribution are available in GATE, namely:

- Flat distributions, defined by the range into which the function is not null, and the value taken within this range.
- Gaussian distributions, defined by a mean value and a standard deviation.
- Exponential distributions, defined by its power.
- Manual distributions, defined by a discrete set of points specified in the GATE macro file. The data are linearly interpolated to define the function in a continuous range.
- File distribution, acting as the manual distribution, but where the points are defined in a separate ASCII file, whose name is given as a parameter. This method is appropriate for large numbers of points and allows to describe any distribution in a totally generic way.

**Usage**

A distribution is declared by specifying its name then by creating a new instance, with its type name:

```
/gate/distributions/name my_distrib
/gate/distributions/insert Gaussian
```

The possible type name available corresponds to the five distributions described above, that is *Flat, Gaussian, Exponential, Manual* or *File*.

Once the distribution is created, the related parameters can be set:

```
/gate/distributions/my_distrib/setMean 350 keV
/gate/distributions/my_distrib/setSigma 30 keV
```

for this example of a Gaussian distribution. All the parameters, for each type of distribution, are summarized in table 8.1

**Table: 8.1: Summary of the parameters for each distribution type**

<table>
<thead>
<tr>
<th>Parameter name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FLAT DISTRIBUTION</strong></td>
<td></td>
</tr>
<tr>
<td>setMin</td>
<td>set the low edge of the range where the function is not null (default is 0)</td>
</tr>
<tr>
<td>SetMax</td>
<td>set the high edge of the range where the function is not null (default is 1)</td>
</tr>
<tr>
<td>setAmplitude</td>
<td>set the value taken by the function within the non null range (default is 1)</td>
</tr>
<tr>
<td><strong>GAUSSIAN DISTRIBUTION</strong></td>
<td></td>
</tr>
<tr>
<td>setMean</td>
<td>set the mean value of the distribution (default is 0)</td>
</tr>
<tr>
<td>setSigma</td>
<td>set the standard deviation of the distribution (default is 1).</td>
</tr>
<tr>
<td>setAmplitude</td>
<td>set the amplitude of the distribution (default is 1).</td>
</tr>
<tr>
<td><strong>EXPONENTIAL DISTRIBUTION</strong></td>
<td></td>
</tr>
<tr>
<td>setLambda</td>
<td>set the power of the distribution (default is 1).</td>
</tr>
<tr>
<td>setAmplitude</td>
<td>set the amplitude of the distribution (default is 1).</td>
</tr>
<tr>
<td><strong>MANUAL DISTRIBUTION</strong></td>
<td></td>
</tr>
<tr>
<td>setUnitX</td>
<td>set the unit for the x axis.</td>
</tr>
<tr>
<td>Command</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>---------------------------------------------------------------------</td>
</tr>
<tr>
<td>setUnitY</td>
<td>set the unit for the y axis.</td>
</tr>
<tr>
<td>insertPoint</td>
<td>insert a new point, giving a pair of (x,y) values.</td>
</tr>
<tr>
<td>addPoint</td>
<td>add a new point, giving its y value, and auto incrementing the x value.</td>
</tr>
<tr>
<td>autoXstart</td>
<td>in case of auto incremental x value, set the first x value to use.</td>
</tr>
</tbody>
</table>

**FILE DISTRIBUTION**

<table>
<thead>
<tr>
<th>Command</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>setUnitX</td>
<td>set the unit for the x axis.</td>
</tr>
<tr>
<td>setUnitY</td>
<td>set the unit for the y axis.</td>
</tr>
<tr>
<td>autoX</td>
<td>specify if the x values are read from file or if they are auto-incremented.</td>
</tr>
<tr>
<td>autoXstart</td>
<td>in case of auto incremental x value, set the first x value to use.</td>
</tr>
<tr>
<td>setFileName</td>
<td>the name of the ASCII file where the data have to be read.</td>
</tr>
<tr>
<td>setColumnX</td>
<td>which column of the ASCII file contains the x axis data.</td>
</tr>
<tr>
<td>setColumnY</td>
<td>which column of the ASCII file contains the y axis data.</td>
</tr>
<tr>
<td>read</td>
<td>do read the file (should be called after specifying all the other parameters).</td>
</tr>
</tbody>
</table>

**Adder**

**Definition**

One particle often creates multiple interactions, and consequently multiple *hits*, within a given crystal. For instance, a photon may interact with a single crystal by two Compton scattering events and a photoelectric absorption. The first step of the digitizer is to sum all the *hits* that occur within the same crystal (i.e. the same volume). This is due to the fact that the electronics always measure an integrated signal, and do not have the time or energy resolution necessary to distinguish between the individual interactions of the particle within a crystal. This digitizer action is completed by a module called the adder. Generally, the adder should be the first module of a digitizer chain.

The adder acts on the lowest level in the system hierarchy, as explained in Users Guide V6.2:Defining a system. As a result:

- A registered system must be used to describe the geometry (also the mother volume name must corresponds to a registered system name),
- The lowest level of this system must be attached to the detector volume and must be declared as a **sensitive detector**.

The adder regroups *hits* per volume into a **pulse**. If one particle that enters a detector makes multiple *hits* within two different crystal volumes before being stopped, the output of the adder module will consists of two **pulses**. Each **pulse** is computed as follows: the energy is taken to be the total of energies in each volume, the position is obtained with an energy-weighted centroid of the different *hit* positions. The time is equal to the time at which the first *hit* occured.

**Command line**

The command to use the adder module is
Adder Compton

Definition

The adderCompton module has a different behavior than the classic adder, which performs an energy-weighted centroid addition of all electronic and photonic hits. Instead, for each electronic energy deposition, the energy is added to the previous photonic hit in the same volume ID (or discarded if none), but the localization remains that of the photonic interaction. That way, the Compton kinematics becomes exact for photonic interactions, enabling further studies. The user must use the classic adder afterwards, to handle multiple photonic interactions in the same crystal.

Command line

The commands to use the adder module are

```
/gate/digitizer/Singles/insert adderCompton
/gate/digitizer/Singles/insert adder
```

Readout

With the exception of a detector system where each crystal is read by an individual photo-detector, the readout segmentation is often different from the basic geometrical structures of the detector. The readout geometry is an artificial geometry that is usually associated with a group of sensitive detectors. This grouping has to be determined by the user through a variable named `depth`. Using this variable, the pulses are summed if their volume ID's are same to this level of depth.

Definition

The readout module regroups pulses per block (group of sensitive detectors). The user needs to specify the block depth to indicate the depth within the volume hierarchy at which pulses are summed together. The results of this module are the total energy in a block, the position of the pulse with the maximum energy (winner-takes-all).

Command line

```
/gate/digitizer/Singles/insert readout
/gate/digitizer/Singles/readout/setDepth 1
```

Figure XXX illustrates the actions of both the `adder` and `readout` modules. The `adder` module transforms the hits into a pulse in each individual volume, and then the `readout` module sums a group of these pulses into a single pulse at the level of depth as defined by the user.
The \textit{setDepth} command line

The importance of this command line is illustrated through the following example from a PET system (see Users Guide V6.2:Defining a system). In a cylindrical PET system, where the first volume level is \textit{rsector}, and the second volume level is \textit{module}, as shown in figure 8.3, the \textit{readout depth} depends upon how the electronic readout functions.

If one PMT reads the four modules in the axial direction, the \textit{depth} should be set with the command:

\begin{verbatim}
/gate/digitizer/Singles/readout/setDepth 1
\end{verbatim}

The energy of this \textit{single} event is the sum of the energy of the pulses inside the white rectangle (\textit{rsector}) of figure 8.3. However, if individual PMTs read each module (group of crystals), the \textit{depth} should be set with
In this case, the energy of the single event is the sum of the energies of the pulses inside the red box (module) of figure 8.3.

The next task is to transform this output pulse from the readout module into a single which is the physical observable of the experiment. This transformation is the result of the detector response and should mimic the behaviors of the photo-detector, electronics, and acquisition system.

**Energy blurring**

**Definition**

The blurring pulse-processor module simulates Gaussian blurring of the energy spectrum of a pulse after the readout module. This is accomplished by introducing a resolution, $R_0$ (FWHM), at a given energy, $E_0$. According to the camera, the energy resolution may follow different laws, such as an inverse square law or a linear law.
Command line

For inverse square law \( R = \frac{R_0 E_0}{\sqrt{E}} \), one must specify the inverse square law and fix the attributes like the energy of reference and the resolution (example of a 15% resolution of @ 511 KeV):

```
/gate/digitizer/Singles/blurring
/gate/digitizer/Singles/blurring/setLaw inverseSquare
/gate/digitizer/Singles/blurring/inverseSquare/setResolution 0.15
/gate/digitizer/Singles/blurring/inverseSquare/setEnergyOfReference 511. keV
```

For linear law, one must specify the linear law and fix the attributes like the energy of reference, the resolution and the slope:

```
/gate/digitizer/Singles/blurring
/gate/digitizer/Singles/blurring/setLaw linear
/gate/digitizer/Singles/blurring/linear/setResolution 0.15
/gate/digitizer/Singles/blurring/linear/setEnergyOfReference 511. keV
/gate/digitizer/Singles/blurring/linear/setSlope -0.055 1/MeV
```

Crystal blurring for a block detector

This type of blurring is used for the scanners where all the detectors are made of the same type of crystal. In this case, it is often useful to assign a different energy resolution for each crystal in the detector block, between a minimum and a maximum value. To model the efficiency of the system, a coefficient (between 0 and 1) can also be set.

Command line

As an example, a random blurring of all the crystals between 15% and 35% at a reference energy of 511 keV, and with a quantum efficiency of 90% can be modelled using the following commands:

```
/gate/digitizer/Singles/insert crystalblurring
/gate/digitizer/Singles/crystalblurring/setCrystalResolutionMin 0.15
/gate/digitizer/Singles/crystalblurring/setCrystalResolutionMax 0.35
/gate/digitizer/Singles/crystalblurring/setCrystalQE 0.9
/gate/digitizer/Singles/crystalblurring/setCrystalEnergyOfReference 511.keV
```

In this example, for each interaction the program randomly chooses a crystal resolution between 0.15 and 0.35. The crystals are not assigned a constant resolution. The crystal quantum efficiency is set using `setCrystalQE` and represents the probability for the event to be detected by the photo-detector. This parameter represents the effect of the transfer efficiency of the crystal and of the quantum efficiency of the photo-detector.

Local energy blurring for a detector module with several types of crystals

The LocalBlurring module is very similar to the energy blurring module, but different energy resolutions are applied to different volumes. This type of blurring is useful for detectors with several layers of different scintillation crystals (e.g. depth of interaction measurement with a phoswich module in a CylindricalPET detector).
system).

**Command line**

To use the *LocalBlurring* module:

- Insert the *LocalBlurring* module,
- choose a valid detector volume name the blurring will be applied to,
- set the resolution for this volume, and set the reference energy for this volume.

For example, if a detector has a resolution of 15.3% @ 511 KeV for a crystal called crystal1 and has a resolution of 24.7% @ 511 KeV for another crystal (crystal2) in a phoswich configuration, the following commands should be used:

```
/gate/digitizer/Singles/insert localBlurring
/gate/digitizer/Singles/localBlurring/chooseNewVolume crystal1
/gate/digitizer/Singles/localBlurring/crystal1/setResolution 0.153
/gate/digitizer/Singles/localBlurring/crystal1/setEnergyOfReference 511 keV
/gate/digitizer/Singles/localBlurring/chooseNewVolume crystal2
/gate/digitizer/Singles/localBlurring/crystal2/setResolution 0.247
/gate/digitizer/Singles/localBlurring/crystal2/setEnergyOfReference 511 keV
```

BEWARE: crystal1 and crystal2 must be valid *Sensitive Detector* volume names !!

**Intrinsic resolution blurring with crystals of different compositions**

**Definition**

This blurring pulse-processor simulates a local Gaussian blurring of the energy spectrum (different for different crystals) based on the following model:

\[
R = \sqrt{2.35^2 \cdot \frac{1 + \bar{\nu}}{N_{ph} \cdot \bar{\epsilon} \cdot \bar{\rho}} + R_i^2}
\]

where \( N_{ph} \) = \( LY \cdot E \) and \( LY, \bar{\rho} \) and \( \bar{\epsilon} \), are the Light Yield, Transfer, and Quantum Efficiency for each crystal.

\( \bar{\nu} \) is the relative variance of the gain of a Photo Multiplier Tube (PMT) or of an Avalanche Photo Diode (APD). It is hard-coded and set to 0.1.

If the intrinsic resolutions, \( R_i \), of the individual crystals are not defined, then they are set to one.

To use this *digitizer* module properly, several modules must be set first. These digitizer modules are *GateLightYield*, *GateTransferEfficiency*, and *GateQuantumEfficiency*.

**Definition of the light yield**

The light yield pulse-processor simulates the crystal light yield. Each crystal must be given the correct light
yield. This module converts the pulse energy into the number of scintillation photons emitted, $N_{ph}$.

**Definition of the transfer efficiency**

The transfer efficiency pulse-processor simulates the transfer efficiencies of the light photons in each crystal. This digitizer reduces the "pulse" energy (by reducing the number of scintillation photons) by a transfer efficiency coefficient which must be a number between 0 and 1.

**Definition of the quantum efficiency**

The quantum efficiency pulse-processor simulates the quantum efficiency for each channel of a photodetector, which can be a Photo Multiplier Tube (PMT) or an Avalanche Photo Diode (APD).

**Command lines for modelling the intrinsic resolution**

The command lines are illustrated using an example of a phoswich module made of two layers of different crystals. One crystal has a light yield of 27000 photons per MeV (LSO crystal), a transfer efficiency of 28%, and an intrinsic resolution of 8.8%. The other crystal has a light yield of 8500 photons per MeV (LuYAP crystal), a transfer efficiency of 24% and an intrinsic resolution of 5.3%

In the case of a cylindricalPET system, the construction of the crystal geometry is truncated for clarity (the truncation is denoted by ...). The digitizer command lines are:

```plaintext
# LSO layer
/gate/crystal/daughters/name LSOlayer ....
# BGO layer
/gate/crystal/daughters/name LuYAPlayer ....
# ATTACH SYSTEM ....
/gate/systems/cylindricalPET/crystal/attach crystal
/gate/systems/cylindricalPET/layer0/attach LSOlayer
/gate/systems/cylindricalPET/layer1/attach LuYAPlayer
# ATTACH CRYSTALS D
/gate/LSOlayer/attachCrystalSD
/gate/LuYAPlayer/attachCrystalSD
# In this example the phoswich module is represented by the crystal volume and is made of two different materials.
# To apply the resolution blurring of equation , the parameters discussed above must be defined for each layer (i.e. Light Yield, Transfer, Intrinsic Resolution, and the Quantum Efficiency).
# DEFINE TRANSFER EFFICIENCY FOR EACH LAYER
/gate/digitizer/Singles/insert transferEfficiency
/gate/digitizer/Singles/transferEfficiency/chooseNewVolume LSOlayer
/gate/digitizer/Singles/transferEfficiency/LSOlayer/setTECoef 0.28
/gate/digitizer/Singles/transferEfficiency/chooseNewVolume LuYAPlayer
/gate/digitizer/Singles/transferEfficiency/LuYAPlayer/setTECoef 0.24
# DEFINE LIGHT YIELD FOR EACH LAYER
/gate/digitizer/Singles/insert lightYield
/gate/digitizer/Singles/lightYield/chooseNewVolume LSOlayer
/gate/digitizer/Singles/lightYield/LSOlayer/setLightOutput 27000
/gate/digitizer/Singles/lightYield/chooseNewVolume LuYAPlayer
/gate/digitizer/Singles/lightYield/LuYAPlayer/setLightOutput 8500
# DEFINE INTRINSIC RESOLUTION FOR EACH LAYER
/gate/digitizer/Singles/insert intrinsicResolutionBlurring
/gate/digitizer/Singles/intrinsicResolutionBlurring/chooseNewVolume LSOlayer
```

Note: A complete example of a phoswich module can be in the PET benchmark.

**Note for Quantum Efficiency**

With the previous commands, the same quantum efficiency will be applied to all the detector channels. The user can also provide lookup tables for each detector module. These lookup tables are built from the user files.

To set multiple quantum efficiencies using files (fileName1, fileName2, ... for each of the different modules), the following commands can be used:

```
/gate/digitizer/Singles/insert quantumEfficiency
/gate/digitizer/Singles/quantumEfficiency/chooseQEVolume crystal
/gate/digitizer/Singles/quantumEfficiency/useFileDataForQE fileName1
/gate/digitizer/Singles/quantumEfficiency/useFileDataForQE fileName2
```

If the crystal volume is a daughter of a module volume which is an array of 8 x 8 crystals, the file fileName1 will contain 64 values of quantum efficiency. If several files are given (in this example two files), the program will choose randomly between these files for each module.

**Important note**

After the introduction of the lightYield (LY), transferEfficiency ($\bar{p}$) and quantumEfficiency ($\bar{\epsilon}$) modules, the energy variable of a pulse is not in energy unit (MeV) but in number of photoelectrons $N_{pe}$.

$$N_{phe} = N_{ph} \cdot \bar{\epsilon} \cdot \bar{p} = LY \cdot E \cdot \bar{\epsilon} \cdot \bar{p}$$

In order to correctly apply a threshold on a phoswich module, the threshold should be based on this number and not on the real energy. In this situation, to apply a threshold at this step of the digitizer chain, the threshold should be applied as explained in #Thresholder & Upholder. In this case, the GATE program knows that these modules have been used, and will apply threshold based upon the number $N_{pe}$ rather than energy. The threshold set with this sigmoidal function in energy unit by the user is translated into number$N_{pe}$ with the lower light yield of the phoswich module. To retrieve the energy it is necessary to apply a calibration module.

**Calibration**

The Calibration module of the pulse-processor models a calibration between $N_{phe}$ and Energy. This is useful when using the class(es) GateLightYield, GateTransferEfficiency, and GateQuantumEfficiency. In addition, a user specified calibration factor can be used.
**Command line**

To set a calibration factor on the energy, use the following commands:

```bash
/gate/digitizer/Singles/insert calibration
/gate/digitizer/Singles/setCalibration VALUE
```

If the calibration digitizer is used without any value, it will correct the energy as a function of values used in GateLightYield, GateTransferEfficiency, and GateQuantumEfficiency.

**Crosstalk**

**Definition**

The crosstalk module simulates the optical and/or electronic crosstalk of the scintillation light between neighboring crystals. Thus, if the input pulse arrives in a crystal array, this module creates pulses around it (in the edge and corner neighbor crystals). The percentage of energy that is given to the neighboring crystals is determined by the user.

**BEWARE:** this module works only for a chosen volume that is an array repeater!!

**Command line**

To insert a crosstalk module that distributes 10% of input pulse energy to the adjacent crystals and 5% to the corner crystals, the following commands can be used:

```bash
/gate/digitizer/Singles/insert crosstalk
/gate/digitizer/Singles/crosstalk/chooseCrosstalkVolume crystal
/gate/digitizer/Singles/crosstalk/setEdgesFraction 0.1
/gate/digitizer/Singles/crosstalk/setCornersFraction 0.05
```

In this example, a pulse is created in each neighbor of the crystal that received the initial pulse. These secondary pulses have 10% (5% for each corner crystals) of the initial energy of the pulse.

**Thresholder & Upholder**

**Definition**

The *Thresholder/Upholder* modules allow the user to apply an energy window to discard low and high energy photons. The low energy cut, supplied by the user, represents a threshold response, below which the detector remains inactive. The user-supplied high energy cut is the maximum energy the detector will register. In both PET and SPECT analysis, the proper setting of these windows is crucial to mimic the behavior of real scanners, in terms of scatter fractions and count rate performances for instance.

**Command line**

In a typical PET scanner, the energy selection for the photo-peak is performed using the following
commands. A low threshold of 0 keV allows the user to see all the events, and is often useful for debugging a simulation.

```bash
/gate/digitizer/Singles/insert thresholder
/gate/digitizer/Singles/thresholder/setThreshold 250. keV
/gate/digitizer/Singles/insert upholder
/gate/digitizer/Singles/upholder/setUphold 750. keV
```

## Energy windows

### Definition

In SPECT analysis, subtractive scatter correction methods such as the dual-energy-window or the triple-energy-window method may be performed in post processing on images obtained from several energy windows. If one needs multiple energy windows, several digitizer branches will be created. Furthermore, the projections associated to each energy window can be recorded into one interfile output.

### Command line

In the following example, 3 energy windows are defined separately with their names and thresholds (see #Thresholder & Upholder):

```bash
/gate/digitizer/name Window1
/gate/digitizer/insert singleChain
/gate/digitizer/Window1/setInputName Singles
/gate/digitizer/Window1/insert thresholder
/gate/digitizer/Window1/thresholder/setThreshold 315 keV
/gate/digitizer/Window1/insert upholder
/gate/digitizer/Window1/upholder/setUphold 328 keV

/gate/digitizer/name Window2
/gate/digitizer/insert singleChain
/gate/digitizer/Window2/setInputName Singles
/gate/digitizer/Window2/insert thresholder
/gate/digitizer/Window2/thresholder/setThreshold 328 keV
/gate/digitizer/Window2/insert upholder
/gate/digitizer/Window2/upholder/setUphold 400 keV

/gate/digitizer/name Window3
/gate/digitizer/insert singleChain
/gate/digitizer/Window3/setInputName Singles
/gate/digitizer/Window3/insert thresholder
/gate/digitizer/Window3/thresholder/setThreshold 400 keV
/gate/digitizer/Window3/insert upholder
/gate/digitizer/Window3/upholder/setUphold 416 keV
```

When specifying the interfile output (see Users Guide V6.2:Data output#Interfile output of projection set), the different window names must be added with the following commands:

```bash
/gate/output/projection/setInputDataName Window1
/gate/output/projection/addInputDataName Window2
/gate/output/projection/addInputDataName Window3
```

## Sigmoidal thresholder

**Definition**

The *Sigmoidal thresholder* models a threshold discriminator based on a sigmoidal function. A sigmoidal function is an S-shaped function of the form, \( \sigma(x) = \frac{1}{1 + c \exp(-ax)} \), which acts as an exponential ramp from 0 to 1:

\[
\sigma(E) = \frac{1}{1 + \exp\left(\alpha \frac{E - E_0}{E_0}\right)},
\]

where the parameter \( \alpha \) is proportional to the slope at symmetrical point \( E_0(\sigma(E_0) = 1/2) \).

For this type of threshold discriminator, the user chooses the threshold setThreshold, the percentage of acceptance for this threshold setThresholdPerCent, and the \( \alpha \) parameter setThresholdAlpha. With these parameters and the input pulse energy, the function is calculated. If the result is bigger than a random number generated between 0 and 1, the pulse is accepted and copied into the output pulse-list. On the other hand, if this criteria is not met, the input pulse is discarded.

**Command line**

```
/gate/digitizer/Singles/insert sigmoidalThresholder
/gate/digitizer/Singles/sigmoidalThresholder/setThreshold 250 keV
/gate/digitizer/Singles/sigmoidalThresholder/setThresholdAlpha 60.
/gate/digitizer/Singles/sigmoidalThresholder/setThresholdPerCent 0.95
```

**Time resolution**

**Definition**

The *temporal resolution* module introduces a Gaussian blurring in the time domain. It works in the same manner as the *blurring* module, but with time instead of energy.

**Command line**

To set a Gaussian temporal resolution (FWHM) of 1.4 ns, use the following commands:

```
/gate/digitizer/Singles/insert timeResolution
/gate/digitizer/Singles/timeResolution/setTimeResolution 1.4 ns
```

**Spatial blurring for SPECT**

For SPECT simulations, the spatial resolution is assumed to follow a Gaussian distribution defined by its width \( \sigma \).

**Command line**
Spatial blurring for PET

In PET analysis, coincidence events provide the lines of response (LOR) needed for the image reconstruction. Only the two crystal numbers are transferred by the simulation. The determination of these crystal numbers is based on the crystal in which the highest energy has been deposited. Without additional spatial blurring of the crystal, simulation results will always have a better spatial resolution than experimental measurements. This module is only available for the *ecat* system. The spatial blurring is based on a 2D Gaussian function.

**Command line**

```bash
# E C A T 7
/gate/output/sinogram/enable
/gate/output/sinogram/RadialBins Your_Sinogram_Radial_Bin_Number
/gate/output/sinogram/setTangCrystalBlurring Your_Value_1 mm
/gate/output/sinogram/setAxialCrystalBlurring Your_Value_2 mm
```

**Noise**

**Definition**

Different sources of background noise exist in a PET/SPECT architecture. For example, the electronics can introduce its own noise, or some crystals used for the detection, such as LSO, contains radioactive nucleus, which can contribute to the background detection count rate. Within GATE, the *noise* module adds such background events, in a totally generic way, so that any kind of source of noise can be simulated. To do so, the energy and the inter-event time interval are chosen randomly, for each event, into user defined distributions, by using the mechanism described in #Distributions.

**Command line**

In the following example, a noise source is introduced, whose energy is distributed according to a Gaussian law, and whose time distribution follows a Poisson process. To do this, one first defines the two necessary distributions. Since the noise description uses the distribution of the time interval between consecutive events, one has to define an exponential distribution. Indeed, if the probability of detecting k events in a time interval of t is distributed along a Poisson law

\[ P_1(k, t) = \frac{\lambda^k t^k e^{-\lambda t}}{k!} \]

then the probability density of having a time interval in the range \([t; t + dt]\) between two consecutive events is given by

\[ dP_2(t) = \lambda e^{-\lambda t} dt. \]
The special event ID, \texttt{event\_ID=-2}, is assigned to these noise events.

**Local efficiency**

**Definition**

The different crystals, or groups of crystals, composing a PET/SPECT system can be characterized by their own efficiency. GATE offers a method to describe such efficiency per crystal or volume.

To define the efficiency distribution in the scanner, one can specify which level of the volume hierarchy of the system are differentiated (see the examples in 
Section 5.4). Then the distribution of efficiency, for each differentiated volume, is specified via a generic distribution, as described in \#Distributions.

**Command lines**

In the following examples, one assumes that the system is composed of 8 blocks (level1) of 64 crystals (level2). The first example shows how to specify one efficiency per block, defined in a file named \texttt{eff\_per\_block.dat}, containing 8 values (one per block).

```
/gate/distributions/name block_eff_distrib
/gate/distributions/insert File
/gate/distributions/block_eff_distrib/autoX true
/gate/distributions/block_eff_distrib/setFileName eff_per_block.dat
/gate/distributions/block_eff_distrib/read

/gate/digitizer/Singles/insert localEfficiency
/gate/digitizer/Singles/localEfficiency/enableLevel 1
/gate/digitizer/Singles/localEfficiency/disableLevel 2
/gate/digitizer/Singles/localEfficiency/setEfficiency block_eff_distrib
```

In the second example, one specifies a different efficiency for each crystal inside a block, but the scheme is repeated from one block to another. So a pattern of 64 efficiency values is defined in the file \texttt{eff\_within\_block.dat}.

```
/gate/distributions/name within_block_eff_distrib
/gate/distributions/insert File
/gate/distributions/within_block_eff_distrib/autoX true
/gate/distributions/within_block_eff_distrib/setFileName eff_within_block.dat
/gate/distributions/within_block_eff_distrib/read

/gate/digitizer/Singles/insert localEfficiency
/gate/digitizer/Singles/localEfficiency/enableLevel 1
/gate/digitizer/Singles/localEfficiency/disableLevel 2
/gate/digitizer/Singles/localEfficiency/setEfficiency within_block_eff_distrib
```

Finally, in the next example, each crystal has its own efficiency, described in the file \texttt{eff\_per\_crystal.dat} containing 8 x 64 elements.
Memory buffers and bandwidth

To mimic the effect of limited transfer rate, a module models the data loss due to an overflow of a memory buffer, read periodically, following a given reading frequency. This module uses two parameters, the reading frequency $\nu$ and the memory depth $D$. Moreover, two reading methods can be modelled, that is, in an event per event basis (an event is read at each reading clock tick), or in a full buffer reading basic (at each reading clock tick, the whole buffer is emptied out). In the first reading method, the data rate is then limited to $\nu$, while in the second method, the data rate is limited to $D \cdot \nu$. When the size limit is reached, any new pulse is rejected, until the next reading clock tick arrival which frees a part of the buffer. In such a case, a non null buffer depth allows to manage a local rise of the input data flow.

Command line

To specify a buffer, read at 10 MHz, with a buffer depth of 64 events, in a mode where the whole buffer is read in one clock tick, one can use:

```plaintext
/gate/digitizer/Your_Single_chain/insert buffer
/gate/digitizer/Your_Single_chain/buffer/setBufferSize 64 B
/gate/digitizer/Your_Single_chain/buffer/setReadFrequency 10 MHz
/gate/digitizer/Your_Single_chain/buffer/setMode 1
```

The chain `Your_Single_chain` can be the default chain `Singles` or any of single chain that the user has defined.

The size of the buffer represents the number of elements, 64 Singles in this example, that the user can store in a buffer.

To read the buffer in an event by event basis, one should replace the last line by `setMode = 0`.

Pile-up

Definition

An important characteristic of a detector is its response time, which is the time that the detector takes to form the signal after the arrival of the radiation. The duration of the signal is also important. During this period, if a second event can be accepted, this second signal will pile up on the first. The resulting pulse is a combination in terms of time and energy, of the two signals. If N pulses enter in the time window of the same sensitive volume (set by the depth of the system level), the output pulse of the pile-up module will be a
pulse with an output energy defined by the sum of the energies \( E_{out} = \sum_{i=0}^{N} E_i \) and a time set to the last time of the last pulse participating to the pile-up \( t_{out} = t_N \).

Since multiple events are grouped into a unique event with the pile-up effect, one can consider this as a loss of events occurring during a given time length, which can be seen as a dead time effect. Moreover, since the pile-up end time is always updated with the last single occurring, the effect is more or less represented by a paralysable dead-time.

**Command line**

To insert a pile-up corresponding to a signal formation time of 100 ns in a module corresponding to the crystal group as described by the 4th level of the system, one should use:

```
/gate/digitizer/Singles/insert pileup
/gate/digitizer/Singles/pileup/setDepth 4
/gate/digitizer/Singles/pileup/setPileup 100 ns
```

**Dead time**

Due to the shaping time of signals or for any other reason, each detection of a single event can hide the subsequent single detected on the same electronic module. This loss lasts a certain amount of time, depending on the characteristics of the detectors used as well as of the readout electronics. The dead time can be modelled in GATE as shown below. Two models of the dead-time have been implemented in the digitizer: *paralysable* and *nonparalysable* response. These models can be implemented *event by event* during a simulation. The detailed method underlying these models can be found in Knoll 1979 (Radiation detection and measurement, John Wiley & Sons, New York). The fundamental assumptions made by these two models are illustrated in figure 8.4.
Definition

The dead time module is applied to a specific volume within the Sensitive Detector system hierarchy. All events taking place within this volume level will trigger a dead-time detector response. This action of the digitizer simulates the time during which this detector, busy at processing a particle, will not be able to process the next one. Moreover, one can simulate the case where data are accumulated into a buffer, which is written to a mass storage having a time access, during which no other data can be processed. In such a case, the dead time is not started after the first data, but once the buffer is full. This case can also be simulated in GATE.

Command line

To apply a dead-time to the volume_name (which has to be previously attached to a level of the system), the following commands can be used:

```bash
# ATTACHEMENT TO THE SYSTEM
/gate/systems/system_name/system_level_name/attach volume_name
...
# DEADTIME
/gate/digitizer/Singles/insert deadtime
/gate/digitizer/Singles/deadtime/setDeadTime 100000. ns
/gate/digitizer/Singles/deadtime/setMode paralysable
/gate/digitizer/Singles/deadtime/chooseDTVolume volume_name
```
The name `system_name` and its corresponding `system_level_name` do not exist and have to be chosen in the tables given in Users Guide V6.2:Defining a system.

In the second example, a dead time corresponding to a disk access of 1 µs for a memory buffer of 1 Mbyte is given. The `setMode` command specifies the behavior of the dead time during the disk access. If this mode is set to 0, the memory buffer is assumed to be a shared resource for the computer, and thus is not available during the disk writing. So, no data can fill the buffer during the disk access. On the other hand, in case of model 1, the buffer is immediately freed after being sent to the disk controller. Data are thus not rejected, unless the buffer is filled up again, before the disk access is finished. In such a case, the dead time module will be totally transparent (ie. will not reject any data), unless the counting rate is high enough to fill the buffer in a time lower than the disk access dead time.

```plaintext
# ATTACHEMENT TO THE SYSTEM
/gate/systems/system_name/system_level_name/attach volume_name
...
# DEADTIME
/gate/digitizer/Singles/insert deadtime
/gate/digitizer/Singles/deadtime/setDeadTime 1 mus
/gate/digitizer/Singles/deadtime/setMode nonparalyzable
/gate/digitizer/Singles/deadtime/chooseDTVolume volume_name
/gate/digitizer/Singles/deadtime/setBufferSize 1 MB
/gate/digitizer/Singles/deadtime/setBufferMode 0
```

**Multiple processor chains**

The use of multiple processor chains makes the design of the digitizer and data output system extremely flexible. The manager for the pulse-processors is called the `GatePulseProcessorChain`, and has a messenger called the `GatePulseProcessorChainMessenger`. By default, all the digitizer components are stored in one processor-chain called `digitizer/Singles`. New processor chains can be created that specify the source of their data. For instance, the following sequence of commands will generate three outputs:

- `Singles` with no energy cut
- `LESingles` with a low-energy window
- `HESingles` with a high-energy window

For a standard PET (with BGO crystals), the components of the standard processor chain will consist in the following commands:

```plaintext
/gate/digitizer/Singles/insert adder
/gate/digitizer/Singles/insert readout
/gate/digitizer/Singles/readout/setDepth 1
```

To add the blurring filter to the "Single" branch:

```plaintext
/gate/digitizer/Singles/insert blurring
/gate/digitizer/Singles/blurring/setResolution 0.26
/gate/digitizer/Singles/blurring/setEnergyOfReference 511. keV
```

The following commands create a low-energy chain branching from the output of "Singles" chain:
These next commands create a high-energy chain branching from the output of the "Singles" chain:

```
/gate/digitizer/name HESingles
/gate/digitizer/insert singleChain
/gate/digitizer/HESingles/setInputName Singles
/gate/digitizer/HESingles/insert thresholder
/gate/digitizer/HESingles/thresholder/setThreshold 350. keV
/gate/digitizer/HESingles/insert upholder
/gate/digitizer/HESingles/upholder/setUphold 650. keV
```

### Coincidence sorter

#### Definition

The coincidence sorter searches, into the singles list, for pairs of coincident singles. Whenever two or more singles are found within a coincidence window, these singles are grouped to form a Coincidence event. Two methods are possible to find coincident singles within GATE. In the first method, when a single is detected, it opens a new coincidence window, and search for a second single occurring during the length of the window. In this method, as long as the window opened by the first single is not closed, no other single can open its own coincidence window. In the second method, all singles open their own coincidence window, and a logical OR is made between all the individual signals to find coincidences. The two methods are available in GATE, and can lead to slightly different results, for a given window width. A comparison of the difference of these two behaviors in a real case is sketched in figure 8.5.
Figure 8.5: Comparison between the two methods of coincidence sorting, for a given succession of singles. In the first one (top), the S2 single does not open its own window since its arrival time is within the window opened by S1. With this method, only one coincidence is created, between S1 and S2. With the second method (bottom), where all singles open their own coincidence window, two different coincidences are identified.

To exclude coincidence coming from the same particle that scattered from a block to an adjacent block, the proximity of the two blocks forming the coincidence event is tested. By default, the coincidence is valid.
only if the difference in the block numbers is greater or equal to two, but this value can be changed in GATE if needed.

Delayed coincidences

Each *Single* emitted from a given source particle is stored with an event ID number, which uniquely identifies the decay from which the single is coming from. If two event ID numbers are not identical in a coincidence event, the event is defined as a *Random* coincidence.

An experimental method used to estimate the number of random coincidences consists of using a delayed coincidence window. By default, the coincidence window is opened when a particle is detected (i.e. when a *Single* is created). In this method, a second coincidence window is created in parallel to the normal coincidence window (which in this case is referred to as the prompt window). The second window (usually with the same width) is open, but is shifted in time. This shift should be long enough to ensure that two particles detected within it are coming from different decays. The resulting number of coincidences detected in this delayed window approximates the number of random events counted in the prompt window. GATE offers the possibility to specify an offset value, for the coincidence sorter, so that prompts and/or delayed coincidence lines can be simulated.

Multiple coincidences

When more than two *singles* are found in coincidence, several type of behavior could be implemented. GATE allows to model 9 different rules that can be used in such a case. The list of rules along with their explanation are given in table 8.2, and a comparison of the effects of each processing rule for various cases of multiple coincidences is shown in figure 8.6. If no policy is specified, the default one used is: *keepMultiplesIfAllAreGoods*.

Table 8.2: Available multiple policy and associated meaning. When a multiple coincidence involving n *singles* is processed, it is first decomposed into a list of n*(n−1)* pairs which are analyzed individually. In this table, the term "good" means that a pair of singles are in coincidence and that the 2 singles are separated by a number of blocks greater than or equal to the *minSectorDifference* parameter of the coincidence sorter. The prefix "take" means that 1 or more pairs of coincidences will be stored, while the prefix "keep" means that a unique coincidence, composed of at least three singles will be kept in the data flow and is called "multicoincidence". In the latter case, the multicoincidence will not be written to the disk, but may participate to a possible deadtime or bandwidth occupancy. The user may clear the multicoincidence at any desired step of the acquisition, by using the multipleKiller pulse processor (described in #Multiple coincidence removal). The "kill" prefix means that all events will be discarded and will not produce any coincidence.

<table>
<thead>
<tr>
<th>Policy name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>takeAllGoods</td>
<td>Each good pairs are considered.</td>
</tr>
<tr>
<td>takeWinnerOfGoods</td>
<td>Only the good pair with the highest energy is considered.</td>
</tr>
<tr>
<td>takeWinnerIfIsGood</td>
<td>If the pair with the highest energy is good, take it, otherwise, kill the event.</td>
</tr>
<tr>
<td>takeWinnerIfAllAreGoods</td>
<td>If all pairs are goods, take the one with the highest energy.</td>
</tr>
<tr>
<td>keepIfOnlyOneGood</td>
<td>If exactly one pair is good, keep all the multicoincidence.</td>
</tr>
<tr>
<td>keepIfAnyIsGood</td>
<td>If at least one pair is good, keep all the multicoincidence.</td>
</tr>
</tbody>
</table>
keepIfAllAreGoods | If all pairs are goods, keep all the multicoincidence.
---|---
killAllIfMultipleGoods | If more than one pairs is good, the event is seen as a real "multiple" and thus, all events are killed.
killAll | No multiple coincidences are accepted, no matter how many good pairs are present.

Figure 8.6: Comparison of the behavior of the available multiple processing policies, for various multiple coincidence situtions. The stars represent the detected singles. The size of the star, as well as the number next to it, indicate the energy level of the single no 1 has more energy than single no 2, which has itself more energy than the single no 3. The lines represent the possible good coincidences (ie. with a sector difference higher than or equal to the minSectorDifference of the coincidence sorter). In the minus(-) sign indicates that the event is killed (ie. no coincidence is formed). The ⋆ sign indicates that all the singles are included into a unique multicoincidence, which will not be written to disk, but which might participate to data loss via dead time or bandwidth occupancy. In the other cases, the list of pairs which are written to the disk (unless being removed thereafter by possible filtering) is indicated.

Table associated with Figure 8.6

<table>
<thead>
<tr>
<th>Policy name</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>takeAllGoods</td>
<td>(1,2)</td>
<td>(1,2); (1,3); (2,3)</td>
<td>(1,2); (2,3)</td>
<td>(1,3); (2,3)</td>
</tr>
<tr>
<td>takeWinnerOfGoods</td>
<td>(1,2)</td>
<td>(1,2)</td>
<td>(1,2)</td>
<td>(1,2)</td>
</tr>
<tr>
<td>takeWinnerIfIsGood</td>
<td>(1,2)</td>
<td>(1,2)</td>
<td>(1,2)</td>
<td>(1,2)</td>
</tr>
<tr>
<td>takeWinnerIfAllAreGoods</td>
<td>-</td>
<td>(1,2)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>keepIfOnlyOneGood</td>
<td>*</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>keepIfAnyIsGood</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>keepIfAllAreGoods</td>
<td>-</td>
<td>*</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>killAllIfMultipleGoods</td>
<td>(1,2)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>killAll</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Command line
To set up a coincidence window of 10 ns, the user should specify:

```
/gate/digitizer/Coincidences/setWindow 10. ns
```

To change the default value of the minimum sector difference for valid coincidences (the default value is 2), the command line should be used:

```
/gate/digitizer/Coincidences/minSectorDifference <number>
```

By default, the offset value is equal to 0, which corresponds to a prompt coincidence sorter. If a delayed coincidence sorter is to be simulated, with a 100 ns time shift for instance, the offset value should be set using the command:

```
/gate/digitizer/Coincidences/setOffset 100. ns
```

To specify the depth of the system hierarchy for which the coincidences have to be sorted, the following command should be used:

```
/gate/digitizer/Coincidences/setDepth <system's depth (1 by default)>
```

As explained in figure 8.5, there are two methods for building coincidences. The default one is the method 1. To switch to method 2, one should use:

```
/gate/digitizer/Coincidences/allPulseOpenCoincGate true
```

So when set to false (by default) the method 1 is chosen, and when set to true, this is the method 2. **Be aware that the method 2 is experimental and not validated at all, so potentially containing non-reported bugs.**

Finally, the rule to apply in case of multiple coincidences is specified as follows:

```
/gate/digitizer/Coincidences/setMultiplePolicy <policyName>
```

The default multiple policy is keepIfAllAreGoods.

**Multiple coincidence sorters**

Multiple coincidence sorters can be used in GATE. To create a coincidence sorter, the sorter must be named and a location specified for the input data. In the example below, three new coincidence sorters are created:

- One with a very long coincidence window:

```
/gate/digitizer/name LongCoincidences
/gate/digitizer/insert coincidenceSorter
/gate/digitizer/LongCoincidences/setInputName Singles
```
One for low-energy singles

A schematic view corresponding to this example is shown in figure 8.7.

![Diagram](image)

Figure 8.7: Readout scheme produced by the listing from the sections

**Coincidence processing and filtering**

**Coincidence pulse processors**

Once the coincidences are identified, further processing can be applied to mimic count losses that may occur because of the acquisition limitations, such as dead time. Count loss may also be due to the limited bandwidth of wires or buffer capacities of the I/O interface. The modelling of such effects within GATE is explained below. Moreover, for PET scanners using a delayed coincidence line, data coming from the two types of coincidences (ie. prompts and delayed) can be processed by a unique coincidence sorter. If so, the
rate of a coincidence type can affect the rate of the other. For instance, a prompt coincidence can produce
death time which will hide a delayed coincidence. The prompt coincidence events can also saturate the
bandwidth, so that the random events are partially hidden.

The modelling of such effects consists in grouping the two different coincidence types into a unique one,
which is then processed by a unique filter.

**Definition**

A coincidence pulse processor is a structure that contains the list of coincidence sources onto which a set of
filters will be applied, along with the list of filters themselves. The order of the list of coincidence may
impact the repartition of the data loss between the prompt and the delay lines. For example, if the line of
prompt coincidences has priority over the line of delayed coincidences, then the events of the latter have
more risk to be rejected by a possible buffer overflow than those of the former. This kind of effects can be
suppressed by specifying that, inside an event, all the coincidences arriving with the same time flag are
merged in a random order.

**Command line**

To implement a coincidence pulse processor merging two coincidence lines into one, and apply an XXX
module followed by another YYY module on the total data flow, one should use the following commands,
assuming that the two coincidence lines named *prompts* and *delayed* are already defined:

```
/gate/digitizer/name myCoincChain
/gate/digitizer/insert coincidenceChain
/gate/digitizer/myCoincChain/addSource prompts
/gate/digitizer/myCoincChain/addSource delayed
/gate/digitizer/myCoincChain/insert XXX
  # set parameter of XXX....
/gate/digitizer/myCoincChain/insert YYY
  # set parameter of YYY....
```

To specify that two coincidences arriving with the same time flag have to be processed in random order, one
should use the command:

```
/gate/digitizer/myCoincChain/usePriority false
```

**Coincidence dead time**

The dead time for coincidences works in the same way as that acting on the *singles* data flow. The only
difference is that, for the *single* dead time, one can specify the hierarchical level to which the dead time is
applied on (corresponding to the separation of detectors and electronic modules), while in the coincidence
dead time, the possibility to simulate separate coincidence units (which may exist) is not implemented.
Apart from this limitation, the command lines for coincidence dead time are identical to the ones for *singles*
dead time, as described in #Pile-up. When more than one coincidence can occur for a unique GEANT4 event
(if more than one coincidence line are added to the coincidence pulse processor, or if multiple coincidences
are processed as many coincidences pairs), then the user can specify that the whole event is kept or rejected,
depending on the arrival time of the first coincidence. To do so, one should use the command line:

```
Coincidence buffers

The buffer module for affecting coincidences uses exactly the same command lines and functionalities as the ones used for single pulse lists, and described in section #Local efficiency.

Multiple coincidence removal

Definition

If the multiple coincidences are kept and not splitted into pairs (ie. if any of the keepXXX multiple coincidence policy is used), the multicoincidences could participate to dataflow occupancy, but could not be written to the disk. Unless otherwise specified, any multicoincidence is then cleared from data just before the disk writing. If needed, this clearing could be performed at any former coincidence processing step, by inserting the multipleKiller module at the required level. This module has no parameter and just kill the multicoincidence events. Multiple coincidences split into many pairs are not affected by this module and cannot be distinguished from the normal "simple" coincidences.

Command line

To insert a multipleKiller, one has to use the syntax:

```
/gate/digitizer/myCoincChain/insert multipleKiller
```

Example of a digitizer setting

Here, the digitizer section of a GATE macro file is analyzed line by line. The readout scheme produced by this macro, which is commented on below, is illustrated in Figure 8.8.
Figure 8.8: Readout scheme produced by the listing below. The disk icons represent the data written to the GATE output.

```plaintext
# ADDER
/gate/digitizer/Singles/insert adder

# READOUT
/gate/digitizer/Singles/insert readout
/gate/digitizer/Singles/readout/setDepth

# ENERGY BLURRING
/gate/digitizer/Singles/insert blurring
/gate/digitizer/Singles/blurring/setResolution 0.26
/gate/digitizer/Singles/blurring/setEnergyOfReference 511. keV

# LOW ENERGY CUT
/gate/digitizer/Singles/insert thresholder
/gate/digitizer/Singles/thresholder/setThreshold 50. keV

/gate/digitizer/name cutLowSingles
/gate/digitizer/insert singleChain
/gate/digitizer/cutLowSingles/setInputName Singles

# NOISE
/gate/distributions/name energy_distrib
/gate/distributions/insert Gaussian
```
Lines 1 to 15: The branch named "Singles" contains the result of applying the adder, readout, blurring, and threshold (50 keV) modules.
Lines 17 to 20: A new branch (line 18) is defined, named "cutLowSingles" (line 17), which follows the "Singles" branch in terms of data flow (line 19).

Lines 21 to 35: Two distributions are created, which will be used for defining a background noise. The first distributions, named energy_distribution (line 23) is a Gaussian centered on 450 keV and of 30 keV standard deviation, while the second one is an exponential distribution with a power of 7.57 \( \mu s \). These two distributions are used to add noise. The energy distribution of this source of noise is Gaussian, while the exponential distribution represents the distribution of time interval between two consecutive noise events (lines 32-34).

Lines 37 to 40: A paralysable (line 39) dead time of 2.2 \( \mu s \) is applied on the resulting signal+noise events.

Lines 43 to 62: Another branch (line 44) named "cutSingles" (line 43) is defined. This branch contains a subset of the "cutLowSingles" branch (line 45) (after dead-time has been applied), composed of those events which pass through the 350 keV/700 keV threshold/uhold window (lines 46-50). In addition, the events tallied in this branch must pass the two dead-time cuts (lines 52 to 61) after the energy window cut.

Lines 65 to 68: The "default" coincidence branch consists of data taken from the output of the high threshold and two dead-time cuts ("cutSingles") (line 65). At this point, a 24 ns window with no delay is defined for this coincidence sorter.

Lines 70 to 75: A second coincidence branch is defined (line 71), which is named "delayedCoincidences". This branch takes its data from the same output ("cutSingles"), but is defined by a delayed coincidence window of 24 ns, and a 100 ns delay (line 73).

Lines 77 to 89: The delayed and the prompts coincidence lines are grouped (lines 79-80). Between two coincidences coming from these two lines and occurring within a given event, the priority is set to the delayed line, since it is inserted before the prompt line, and the priority is used (line 81). A non-paralysable dead time of 60 ns is applied on the delayed+prompt coincidences (lines 82-85). If more than one coincidence occur inside a given event, the dead time can kill all of them or none of them, depending on the arrival time of the first one. As a consequence, if a delay coincidence is immediately followed by a prompt coincidence due to the same photon, then, the former will not hide the latter (line 85). Finally, a memory buffer of 32 coincidences, read at a frequency of 14.45 MHz, in an event-by-event basis (line 89) is applied to the delayed+prompt sum (lines 86-89).

**Digitizer optimization**

In GATE standard operation mode, primary particles are generated by the source manager, and then propagated through the attenuating geometry before generating *hits* in the detectors, which feed into the digitizer chain. While this operation mode is suited for conventional simulations, it is inefficient when trying to optimize the parameters of the digitizer chain. In this case, the user needs to compare the results obtained for different sets of digitizer parameters that are based upon the same series of hits. Thus, repeating the particle generation and propagation stages of a simulation is unnecessary for tuning the digitizer setting.

For this specific situation, GATE offers an operation mode dedicated to digitizer optimization, known as *DigiGATE*. In this mode, *hits* are no longer generated: instead, they are read from a hit data-file (obtained from an initial GATE run) and are fed directly into the digitizer chain. By bypassing the generation and propagation stages, the computation speed is significantly reduced, thus allowing the user to compare
various sets of digitizer parameters quickly, and optimize the model of the detection electronics. *DigiGATE* is further explained in chapter 13.

**Angular Response Functions (ARFs) to speed-up single photon (planar and SPECT) simulations**

The ARF is a function of the incident photon direction and energy and represents the probability that a photon will either interact with or pass through the collimator, and be detected at the intersection of the photon direction vector and the detection plane in an energy window of interest.

The use of ARF can significantly speed up planar or SPECT simulations. The use of this functionality involves 3 steps.

**Calculation of the data needed to derive the ARF tables**

In this step, the data needed to generate the ARF tables are computed from a rectangular source located at the center of FOV. The SPECT head is duplicated twice and located at roughly 30 cm from the axial axis.

The command needed to compute the ARF data is:

```
/gate/systems/SPECThead/arf/setARFStage generateData
```

The ARF data are stored in ROOT format as specified by the GATE command output:

```
/gate/output/arf/setFileName testARFdata
```

By default the maximum size of a ROOT file is 1.8 Gbytes. Once the file has reached this size, ROOT automatically closes it and opens a new file name testARFdata_1.root. When this file reaches 1.8 Gb, it is closed and a new file testARFdata_2.root is created etc.

A template macro file is provided in the folder /examples/example_ARF/ generateARFdata.mac which illustrates the use of the commands listed before.

**Computation of the ARF tables from the simulated data**

Once the required data are stored in ROOT files, the ARF tables can be calculated and stored in a binary file:

```
# COMPUTE THE ARF TABLES FROM ARF DATA
# /gate/systems/SPECThead/arf/setARFStage computeTables
```

The digitizer parameters needed for the computation of teh ARF table are defined by:

```
# DIGITIZER PART OF THE ARF SENSITIVE DETECTOR
#/gate/systems/SPECThead/ARFTables/setEnergyDepositionThreshHold 328. keV
#/gate/systems/SPECThead/ARFTables/setEnergyDepositionUpHold 400. keV
```
In this example, we shot photons with 364.5 keV as kinetic energy. We chose an energy resolution of 10% @ 140 keV and the energy window was set to [328-400] keV. The simulated energy resolution at an energy Edep will be calculated by:

\[
\text{fwhm} = 0.10 \times \sqrt{140 \times E_{\text{dep}}}, \quad \text{where } E_{\text{dep}} \text{ is the photon deposited energy.}
\]

If we want to discard photons which deposit less than 130 keV, we may use:

```bash
/gate/systems/SPECThead/ARFTables/setEnergyDepositionThreshold 130. keV
```

The ARF tables depend strongly on the distance from the detector to the source used in the previous step. The detector plane is set to be the half-middle plan of the detector part of the SPECT head. In our example, the translation of the SPECT head was 34.5 cm along the X direction (radial axis), the detector was 2 cm wide along X and its center was located at x = 1.5 cm with respect to the SPECThead frame. This is what we call the detector plane (x = 1.5 cm) so the distance from the source to the detector plane is \(34.5 + 1.5 = 36\) cm.

```bash
# DISTANCE FROM SOURCE TO DETECTOR PLANE
# TAKEN TO BE THE PLANE HALF-WAY THROUGH THE CRYSTAL RESPECTIVELY
# TO THE SPECHEAD FRAME
# here it is 34.5 cm + 1.5 cm
#
/gate/systems/SPECThead/ARFTables/setDistanceFromSourceToDetector 36 cm
```

The tables are then computed from a text file which contains information regarding the incident photons called ARFData.txt which is provided in the release folder /examples/example_ARF:

```bash
# NOW WE ARE READY TO COMPUTE THE ARF TABLES
#
/gate/systems/SPECThead/ARFTables/ComputeTablesFromEnergyWindows ARFData.txt
```

The text file reads like this:

```
# this file contains all the energy windows computed during first step
# with their associated root files
# it has to be formatted the following way
# [Emin,Emax] is the energy window of the incident photons
# the Base FileName is the the name of the root files name.root, name_1.root name_2.root ...
# the number of these files has to be given as the last parameter
#
# enter the data for each energy window
# Incident Energy Window: Emin - Emax (keV) | Root FileName | total file number
0.     365.         test1head            20
```

Here we have only one incident energy window for which we want to compute the corresponding ARF tables. The data for this window are stored inside 20 files whose base file name is test1head. These ARF data files were generated from the first step and were stored under the names of test1head.root, test1head_1.root ... test1head_19.root.
Finally the computed tables are stored to a binary file:

```
/gate/systems/SPECThead/ARFTables/list
# SAVE ARF TABLES TO A BINARY FILE FOR PRODUCTION USE
/gate/systems/SPECThead/ARFTables/saveARFTablesToBinaryFile ARFSPECTBench.bin
```

### Use of the ARF tables

The command to tell GATE to use ARF tables is:

```
/gate/systems/SPECThead/arf/setARFStage useTables
```

The ARF sensitive detector is attached to the SPECThead:

```
/gate/SPECThead/attachARFSD
```

These tables are loaded from binary files with:

```
/gate/systems/SPECThead/ARFTables/loadARFTablesFromBinaryFile ARFTables.bin
```

### Multi-system approaches: how to use more than one system in one simulation set-up?

Singles arriving from different systems request different treatment in the digitizer. So we have to use multiple digitizer chains and to separate between these singles according to their systems.

#### SystemFilter: definition

The systemFilter module separates between the singles coming from systems. This module have one parameter which is the name of the system.

**Command line**

```
/gate/digitizer/SingleChain/insert systemFilter
/gate/digitizer/SingleChain/systemFilter/selectSystem SystemName
```

SingleChain is the singles chain, Singles by default, and SystemName is the name of the selected system. Note that the natural order of this module is the first.

**Example: modification in your Digitizer macro file**

Suppose that we have two systems with the names “cylindricalPET” and “Scanner_1”, so to select singles in cylindricalPET system we use:
Note we didn't insert a singles chain because we have the default chain “Singles”, on the other side for “Scanner_1” we to insert a new singles chain “Singles_S1”:

```
gate/digitizer/name Singles_S1
/gate/digitizer/insert singleChain
```

Then we insert the system filter.

```
gate/digitizer/Singles_S1/insert systemFilter
/gate/digitizer/Singles_S1/systemFilter/selectSystem Scanner_1
```

How to manage the coincidence sorter with more than one system: the Tri Coincidence Sorter approach

Description:

The aim of this module is to obtain the coincidence between coincidence pulses and singles of another singles chain (between TEP camera and scanner for example). In this module we search the singles, of the concerned singles chain, which are in coincidence with the coincidence pulses according to a time window. In fact, this module, from the point of view of the code, is coincidence pulses processor. So we have a coincidence chain as input with a singles chain. We have also to define a time window to search the coincident coincidence-singles within this window. We test the time difference between the average time of the tow singles of the coincidence pulse and the time of every single of the singles chain in question. We have as output of this module two trees of ROOT: one tree contain the coincidences which have at first one coincident single with every one of them. The second tree contain the coincident singles and it is generated automatically with name of coincidence tree+“CSingles”. For example if we call the coincidence “triCoinc”, so the name of the singles tree will be “triCoincCSingles”. This singles tree contain the same branches as any singles tree with an additional branch named “triCID” from (tri coincidence ID) and it has the same value for all singles which are in coincidence with one coincidence pulse.

Strategy:

In this module we store the singles event by event in a singles buffer that have a semi-constant size that the user can adjust. When we have a coincidence pulse we compare between the average time of this coincidence pulse and the time of each single pulse in the buffer. When we have coincident singles with the coincidence pulse, we store them in two trees as mentioned above.

Commands and example:

Define the classical coincidence sorter applied on the cylindrical PET system:
Define now a "coincidenceChain" where you will plug the first coincidence sorter (named 'Coincidences' in that case):

```
/gate/digitizer/Coincidences/setWindow 30. ns
```

Finally, we call the 'triCoincProcessor' module and we plug on it the second system which defines in that case by the 'Singles_S1' chain:

```
/gate/digitizer/TriCoinc/name TriCoinc
/gate/digitizer/TriCoinc/insert coincidenceChain
/gate/digitizer/TriCoinc/addInputName Coincidences

/gate/digitizer/TriCoinc/insert triCoincProcessor
/gate/digitizer/TriCoinc/triCoincProcessor/setSinglesPulseListName Singles_S1
/gate/digitizer/TriCoinc/triCoincProcessor/setWindow 15 ns
/gate/digitizer/TriCoinc/triCoincProcessor/setSinglesBufferSize 40
```

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Users Guide V7.0 : Data output

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Data output is a key point for a software intended to be used for various applications, in various scientific communities. In GATE, there are several types of output format, which can be enabled. By default all outputs are disabled. And moreover there is no default output file name. You _must_ give a file name, otherwise the output module will be automatically disabled. The following chapter describes the various output formats, such as ASCII, Root, Interfile, LMF, and ECAT.

The ASCII output

Introduction

The GateToASCII class of GATE enables the ASCII file output, which is the easiest possible output. It allows you to process your raw data with your own tools. On the other hand, this output is not compressed and the output files are very large.

If the ASCII files are not needed for analysis, it is strongly recommended not using this output to speed up the simulation and save space.

How to enable this output in your macro?

All the output commands

```
/gate/output/...
```

must always be written after the initialization line.

As in most of the output modules of GATE, you can enable ASCII output files for Hits, Singles (at the end of the digitizer chain), Coincidences, but also for Singles after the different steps of the digitizer chain.

As a first step, enable the ASCII output and give an output file name:
In your macro, the different flags should be set to 1:

```gate/output/ascii/enable
/gate/output/ascii/setFileName test
```

The names of the digitizer module are: *Adder, Readout, Spblurring, Blurring, Thresholder, Upholder*. Their actions are explained in Users Guide V6.2: Digitizer and readout parameters.

How to disable this output in your macro?

This output, as for all output is disabled by default, but if it is enabled, some ASCII files will be created, namely: *gateHits.dat, gateSingles.dat, gateRun.dat*. In addition, if coincidences are processed in the simulation, the file *gateCoincidences.dat* will be generated. To disable these ASCII files which can be large, the macro should contain the following lines:

```gate/output/ascii/setOutFileHitsFlag 0
/gate/output/ascii/setOutFileSinglesFlag 0
/gate/output/ascii/setOutFileCoincidencesFlag 0
```

Only the file *gateRun.dat* which contain the number of decay per run will then be created.

Description of the ASCII file content

In all files, the units are:

- MeV (energy)
- mm (position)
- s (time)
- deg (angle)

**Hits file (gateHits.dat)**

Each line is a hit and the columns represent:

- Column 1: ID of the run (i.e. time-slice)
- Column 2: ID of the event
- Column 3: ID of the primary particle whose descendant generated this hit
- Column 4: ID of the source which emitted the primary particle
- Columns 5 to N+4: Volume IDs at each level of the hierarchy of a system, so the number of columns
depends on the system used.

For cylindricalPET system N=6 :

- Column 5 : ID of volume attached to the "base" level of the system
- Column 6 : ID of volume attached to the "rsector" level of the system
- Column 7 : ID of volume attached to the "module" level of the system
- Column 8 : ID of volume attached to the "submodule" level of the system
- Column 9 : ID of volume attached to the "crystal" level of the system
- Column 10 : ID of volume attached to the "layer" level of the system

For SPECTHead system N=3 :

- Column 5 : ID of volume attached to the "base" level of the system
- Column 6 : ID of volume attached to the "crystal" level of the system
- Column 7 : ID of volume attached to the "pixel" level of the system
- Column N+5 : Time stamp of the hit
- Column N+6 : Energy deposited by the hit (which may be given as a percentage of the initial particle)
- Column N+7 : Range of particle which has generated the hit
- Column N+8, N+9 ,N+10 : XYZ position of the hit in the world referential
- Column N+11 : Geant4 code of the particle which has generated the hit (11 for Electrons & 22 for Photons)
- Column N+12 : ID of the particle which has generated the hit
- Column N+13 : ID of the mother of the particle which has generated the hit
- Column N+14 : ID of the photon giving the particle which has generated the hit
- Column N+15 : Number of Compton interactions in phantoms before reaching the detector
- Column N+16 : Number of Rayleigh interactions in phantoms before reaching the detector
- Column N+17 : Name of the process which has generated the hit
- Column N+18 : Name of the last volume where a Compton effect occurred
- Column N+19 : Name of the last volume where a Rayleigh effect occurred

For the next sections, the system will be set as a cylindricalPET system, so that the number of lines concerning the Volume ID of each level will always be 5.

Singles files (gateSingles.dat)

Each line is a single and the columns are :

- Column 1 : ID of the run (i.e. time-slice)
- Column 2 : ID of the event
- Column 3 : ID of the source
- Column 4, 5, 6 : XYZ position of the source in world referential
- Column 7 to 12 : Volume IDs*(cf. columns 5-10 of sec 11.)
- Column 13 : Time stamp of the single
- Column 14 : Energy deposited by the single
- Column 15 to 17 : XYZ position of the single in the world referential
- Column 18 : Number of Compton interactions in phantoms before reaching the detector
- Column 19 : Number of Compton interactions in detectors before reaching the detector
- Column 20 : Number of Rayleigh interactions in phantoms before reaching the detector
- Column 21 : Number of Rayleigh interactions in detectors before reaching the detector
- Column 22 : Name of the phantom where a Compton effect occurred
- Column 23 : Name of the phantom where a Rayleigh effect occured

## Coincidences files (gateCoincidences.dat)

Each line is a coincidence created with two singles and the columns are:

- Column 1 : ID of the run (i.e. time-slice) (first single)
- Column 2 : ID of the event (first single)
- Column 3 : ID of the source (first single)
- Column 4 to 6 : XYZ position of the source in world referential (first single)
- Column 7 : Time stamp (first single)
- Column 8 : Deposited energy (first single)
- Column 9 to 11 : XYZ position in the world referential (first single)
- Column 12 to 17 : volume IDs* (cf. Users Guide V6.2:Data output#Description of the ASCII file content) (first single)
- Column 18 : Number of Compton interactions in phantoms before reaching the detector (first single)
- Column 19 : Number of Compton interactions in detectors before reaching the detector (first single)
- Column 20 : Number of Rayleigh interactions in phantoms before reaching the detector (first single)
- Column 21 : Number of Rayleigh interactions in detectors before reaching the detector (first single)
- Column 22 : Scanner axial position (first single)
- Column 23 : Scanner angular position (first single)
- Column 24 : ID of the run (i.e. time-slice) (second single)
- Column 25 : ID of the event (second single)
- Column 26 : ID of the source (second single)
- Column 27 to 29 : XYZ position of the source in world referential (second single)
- Column 30 : Time stamp (second single)
- Column 31 : Energy deposited (second single)
- Column 32 to 34 : XYZ position in the world referential (second single)
- Column 35 to 40 : volume IDs.

The number of different volumeIDs depends on the complexity of the system geometry (6 IDs for cylindricalPET system, 3 for ecat system, ...). Then, the number of column of your ASCII file is not constant, but system-dependent.

- Column 41 : Number of Compton interactions in phantoms before reaching the detector (second single)
- Column 42 : Number of Compton interactions in detectors before reaching the detector (second single)
- Column 41 : Number of Rayleigh interactions in phantoms before reaching the detector (second single)
- Column 42 : Number of Rayleigh interactions in detectors before reaching the detector (second single)
- Column 45 : Scanner axial position (second single)
- Column 46 : Scanner angular position (second single)

## Selection of the variables in Singles/Coincidences ASCII output

The user can select with a macro command which variables he/she wants in the ASCII file. The mechanism is based on a mask with a mask, i.e. a series of 0/1, one for each variable. By default all variables are enabled, but one can choose to enable only some of the variables listed in 10.4.1. One example is:
Note: the VolumeID variables are enabled/disabled together, as a group. The component of the 3D vectors, instead, like the positions (components x,y,z), are enabled/disabled one by one.

Large files: automatic file swap for the ASCII output

When a user defined limit is reached by the Coincidence or Single ASCII output file, by default Gate closes the file and opens another one with the same name but a suffix _1 (and then _2, and so on). By default the file limit is set to 200000000 bytes. One can change the number of bytes with a command like

```
/gate/output/ascii/setOutFileSizeLimit 30000
```

If the value is < 10000, no file swapping is made (to avoid creating thousands of files by mistake).

For example, if one does not have any limit in the Operating System, one can put the number to 0, and there will be only one large (large) file at the end.

In case of high statistics applications, one might consider enabling only the ROOT output (see #The Root output), which contains the same information as the ASCII one, but automatically compressed and ready for analysis.

What is the file gateRun.dat?

This file is the list of the number of decays generated by the source for each run (one by line). The Output manager is called for each event, even if the particle(s) of the decay do not reach the detector. Note that the number of processed decays can be slightly different from the expected number \( N = A \times \Delta t \) where \( A \) is the activity and \( \Delta t \) is the time of the acquisition, due to the random character of the decay which governs the event generation (Poisson law). Gate generates the time delay from the previous event, if it is out of the time slice it stops the event processing for the current time slice and if needed it starts a new time slice.

The Binary output

Introduction

The GateToBinary class of GATE enables the Binary file output, which is the easiest possible output. It allows you to process your raw data with your own tools. On the other hand, this output is not compressed and the output files are very large.

If the binary files are not needed for analysis, it is strongly recommended not using this output to speed up the simulation.

How to enable this output in your macro?

All the output commands
must always be written after the initialization line.

As in most of the output modules of GATE, you can enable binary output files for Hits, Singles (at the end of the digitizer chain), Coincidences, but also for Singles after the different steps of the digitizer chain.

As a first step, enable the binary output and give an output file name:

```plaintext
/gate/output/binary/enable /gate/output/binary/setFileName test
```

In your macro, the different flags should be set to 1:

```plaintext
# enable binary output for hits
/gate/output/binary/setOutFileHitsFlag 1
#
# enable binary output for Singles (end of digitizer chain) 
/gate/output/binary/setOutFileSinglesFlag 1
#
# enable binary output for coincidences
/gate/output/binary/setOutFileCoincidencesFlag 1
#
# enable binary output for singles (after a digitizer module)
/gate/output/binary/setOutFileSingles< name of the digitizer module >Flag 1
```

The names of the digitizer module are: *Adder, Readout, Spblurring, Blurring, Thresholder, Upholder*. Their actions are explained in Users Guide V6.2: Digitizer and readout parameters.

**How to disable this output in your macro?**

This output, as for all output is disabled by default, but if it is enabled, some binary files will be created, namely: *gateHits.bin, gateSingles.bin, gateRun.bin*. In addition, if coincidences are processed in the simulation, the file *gateCoincidences.bin* will be generated. To disable these binary files which can be large, the macro should contain the following lines:

```plaintext
/gate/output/binary/setOutFileHitsFlag 0
/gate/output/binary/setOutFileSinglesFlag 0
/gate/output/binary/setOutFileCoincidencesFlag 0
```

Only the file *gateRun.bin* which contain the number of decay per run will then be created.

**Description of the bin file content**

In all files, the units are:

- MeV (energy)
- mm (position)
- s (time)
- deg (angle)

**Hits file (gateHits.bin)**
Each line is a hit and the columns represent:

- **Column 1**: ID of the run (i.e. time-slice) (4-bytes, G4int)
- **Column 2**: ID of the event (4-bytes, G4int)
- **Column 3**: ID of the primary particle whose descendant generated this hit (4-bytes, G4int)
- **Column 4**: ID of the source which emitted the primary particle (4-bytes, G4int)
- **Columns 5 to N+4**: Volume IDs at each level of the hierarchy of a system, so the number of columns depends on the system used.

For cylindricalPET system N=6:

- **Column 5**: ID of volume attached to the "base" level of the system (4-bytes, G4int)
- **Column 6**: ID of volume attached to the "rsector" level of the system (4-bytes, G4int)
- **Column 7**: ID of volume attached to the "module" level of the system (4-bytes, G4int)
- **Column 8**: ID of volume attached to the "submodule" level of the system (4-bytes, G4int)
- **Column 9**: ID of volume attached to the "crystal" level of the system (4-bytes, G4int)
- **Column 10**: ID of volume attached to the "layer" level of the system (4-bytes, G4int)

For SPECTHead system N=3:

- **Column 5**: ID of volume attached to the "base" level of the system (4-bytes, G4int)
- **Column 6**: ID of volume attached to the "crystal" level of the system (4-bytes, G4int)
- **Column 7**: ID of volume attached to the "pixel" level of the system (4-bytes, G4int)
- **Column N+5**: Time stamp of the hit (8-bytes, G4double)
- **Column N+6**: Energy deposited by the hit (8-bytes, G4double)
- **Column N+7**: Range of particle which has generated the hit (8-bytes, G4double)
- **Column N+8, N+9 ,N+10**: XYZ position of the hit in the world referential (8-bytes, G4double)
- **Column N+11**: Geant4 code of the particle which has generated the hit (4-bytes, G4int)
- **Column N+12**: ID of the particle which has generated the hit (4-bytes, G4int)
- **Column N+13**: ID of the mother of the particle which has generated the hit (4-bytes, G4int)
- **Column N+14**: ID of the photon giving the particle which has generated the hit (4-bytes, G4int)
- **Column N+15**: Number of Compton interactions in phantoms before reaching the detector (4-bytes, G4int)
- **Column N+16**: Number of Rayleigh interactions in phantoms before reaching the detector (4-bytes, G4int)
- **Column N+17**: Name of the process which has generated the hit (8-bytes, G4string)
- **Column N+18**: Name of the last volume where a Compton effect occured (8-bytes, G4string)
- **Column N+19**: Name of the last volume where a Rayleigh effect occured (8-bytes, G4string)

For the next sections, the system will be set as a cylindricalPET system, so that the number of lines concerning the Volume ID of each level will always be 5.

**Singles files (gateSingles.bin)**

Each line is a single and the columns are:

- **Column 1**: ID of the run (i.e. time-slice) (4-bytes, G4int)
- **Column 2**: ID of the event (4-bytes, G4int)
- **Column 3**: ID of the source (4-bytes, G4int)
- **Columns 4, 5, 6**: XYZ position of the source in world referential (8-bytes, G4double)
- **Column 7 to 12**: Volume IDs*(cf. columns 5-10 of sec 11.)* (4-bytes, G4int)
| Column 13 | Time stamp of the single (8-bytes, G4double) |
| Column 14 | Energy deposited by the single (8-bytes, G4double) |
| Column 15 to 17 | XYZ position of the single in the world referential (8-bytes, G4double) |
| Column 18 | Number of Compton interactions in phantoms before reaching the detector (4-bytes, G4int) |
| Column 19 | Number of Compton interactions in detectors before reaching the detector (4-bytes, G4int) |
| Column 20 | Number of Rayleigh interactions in phantoms before reaching the detector (4-bytes, G4int) |
| Column 21 | Number of Rayleigh interactions in detectors before reaching the detector (4-bytes, G4int) |
| Column 22 | Name of the phantom where a Compton effect occurred (8-bytes, G4string) |
| Column 23 | Name of the phantom where a Rayleigh effect occurred (8-bytes, G4string) |

**Coincidences files (gateCoincidences.bin)**

Each line is a coincidence created with two singles and the columns are:

| Column 1 | ID of the run (i.e. time-slice) (first single) (4-bytes, G4int) |
| Column 2 | ID of the event (first single) (4-bytes, G4int) |
| Column 3 | ID of the source (first single) (4-bytes, G4int) |
| Column 4 to 6 | XYZ position of the source in world referential (first single) (8-bytes, G4double) |
| Column 7 | Time stamp (first single) (8-bytes, G4double) |
| Column 8 | Deposited energy (first single) (8-bytes, G4double) |
| Column 9 to 11 | XYZ position in the world referential (first single) (8-bytes, G4double) |
| Column 12 to 17 | volume IDs* (cf. Users Guide V6.2:Data output#Description of the ASCII file content) (first single) (8-bytes, G4double) |
| Column 14 | (8-bytes, G4double) |
| Column 15 to 17 | (4-bytes, G4int) |
| Column 18 | Number of Compton interactions in phantoms before reaching the detector (first single) (4-bytes, G4int) |
| Column 19 | Number of Compton interactions in detectors before reaching the detector (first single) (4-bytes, G4int) |
| Column 20 | Number of Rayleigh interactions in phantoms before reaching the detector (first single) (4-bytes, G4int) |
| Column 21 | Number of Rayleigh interactions in detectors before reaching the detector (first single) (4-bytes, G4int) |
| Column 22 | Scanner axial position (first single) (8-bytes, G4double) |
| Column 23 | Scanner angular position (first single) (8-bytes, G4double) |
| Column 24 | ID of the run (i.e. time-slice) (second single) (4-bytes, G4int) |
| Column 25 | ID of the event (second single) (4-bytes, G4int) |
| Column 26 | ID of the source (second single) (4-bytes, G4int) |
| Column 27 to 29 | XYZ position of the source in world referential (second single) (8-bytes, G4double) |
| Column 30 | Time stamp (second single) (8-bytes, G4double) |
| Column 31 | Energy deposited (second single) (8-bytes, G4double) |
| Column 32 to 34 | XYZ position in the world referential (second single) (8-bytes, G4double) |
| Column 35 to 40 | volume IDs. (8-bytes, G4double) |
| Column 37 | (8-bytes, G4double) |
| Column 38 to 40 | (4-bytes, G4int) |
The number of different volumeIDs depends on the complexity of the system geometry (6 IDs for cylindricalPET system, 3 for ecat system, ...). Then, the number of column of your ASCII file is not constant, but system-dependent.

- Column 41: Number of Compton interactions in phantoms before reaching the detector (second single) (4-bytes, G4int)
- Column 42: Number of Compton interactions in detectors before reaching the detector (second single) (4-bytes, G4int)
- Column 41: Number of Rayleigh interactions in phantoms before reaching the detector (second single) (4-bytes, G4int)
- Column 42: Number of Rayleigh interactions in detectors before reaching the detector (second single) (4-bytes, G4int)
- Column 45: Scanner axial position (second single) (8-bytes, G4double)
- Column 46: Scanner angular position (second single) (8-bytes, G4double)

**Selection of the variables in Singles/Coincidences binary output**

The user can select with a macro command which variables he/she wants in the binary file. The mechanism is based on a mask with a mask, i.e. a series of 0/1, one for each variable. By default all variables are enabled, but one can choose to enable only some of the variables listed in 10.4.1. One example is:

```
/gate/output/binary/setCoincidenceMask 1 0 1 0 1 1
/gate/output/binary/setSingleMask 0 0 1 1
```

**Note:** the VolumeID variables are enabled/disabled together, as a group. The component of the 3D vectors, instead, like the positions (components x,y,z), are enabled/disabled one by one.

**Large files: automatic file swap for the binary output**

When a user defined limit is reached by the Coincidence or Single binary output file, by default Gate closes the file and opens another one with the same name but a suffix _1 (and then _2, and so on). By default the file limit is set to 2000000000 bytes. One can change the number of bytes with a command like

```
/gate/output/binary/setOutFileSizeLimit 30000
```

If the value is < 10000, no file swapping is made (to avoid creating thousands of files by mistake).

For example, if one does not have any limit in the Operating System, one can put the number to 0, and there will be only one large (large) file at the end.

In case of high statistics applications, one might consider enabling only the ROOT output (see #The Root output), which contains the same information as the binary one, but automatically compressed and ready for analysis.

**What is the file gateRun.bin ?**

This file is the list of the number of decays generated by the source for each run (one by line). The Output manager is called for each event, even if the particle(s) of the decay do not reach the detector. Note that the
number of processed decays can be slightly different from the expected number \( N = A \times \Delta t \) where \( A \) is the activity and \( \Delta t \) is the time of the acquisition, due to the random character of the decay which governs the event generation (Poisson law). Gate generates the time delay from the previous event, if it is out of the time slice it stops the event processing for the current time slice and if needed it starts a new time slice.

**The Root output**

**How to enable this output in your macro ?**

If you need to generate the root output file, this can be done by adding the following lines in the macro:

```
/gate/output/root/enable
/gate/output/root/setFileName FILE_NAME
```

which will provide you with a FILE_NAME.root file.

By default, this root file will contain : 2 Trees for SPECT systems (Hits and Singles) or 3 Trees for PET systems (Coincidences, Hits and Singles) in which several variables are stored.

When launching ROOT with the command:

```
root file.root
root [1] TBrowser t
```

you can easily see the content of your ROOT data file.

**How to disable this output in your macro ?**

If needed, and for a matter of file size, you could choose not to generate all trees. In this case, just add the following lines in your macro:

```
/gate/output/root/setRootHitFlag 0
/gate/output/root/setRootSinglesFlag 0
/gate/output/root/setRootCoincidencesFlag 0
/gate/output/root/setRootNtupleFlag 0
```

By turning to 1 (or 0) one of this tree flag, you will fill (or not) the given tree.

In a debug mode, it can be useful to store in a Tree the informations after the action of one particular module of the digitizer chain. The following flags exist to turn on or off these intermediate Trees.

```
/gate/output/root/setOutFileSinglesAdderFlag 0
/gate/output/root/setOutFileSinglesReadoutFlag 0
/gate/output/root/setOutFileSinglesSpblurringFlag 0
/gate/output/root/setOutFileSinglesBlurringFlag 0
/gate/output/root/setOutFileSinglesThresholderFlag 0
/gate/output/root/setOutFileSinglesUpholderFlag 0
```

And if you want to disable the whole ROOT output, just do not call it, or use the following command:
How to merge Root files?

Two or more Root files can be merged into one single file by using the `hadd` utility on the command line:

```
hadd result.root file1.root file2.root ... filen.root
```

How to analyze the Root output

You can either plot the variables directly from the browser, or through a macro file (e.g. called `analysis.C`). In this case, type:

```
root [0] .x analysis.C
```

You may also use the root class called MakeClass:

```
root [0] Coincidences->MakeClass("test");
```

Please consult the ROOT Homepage: http://root.cern.ch/ for more details.

The ROOT Online plotter

Along with standard output for post-processing (such as root, LMF, ecat), GATE provides a very convenient tool called the online plotter, which enables online display of several variables. This online analysis is available even if the root output is disabled in your macro, for instance because the user does not want to save a large root file. But Gate have to be compiled with certain options to have this output available.

The online plotter can be easily used with the following macro:

```
/gate/output/plotter/enable
/gate/output/plotter/showPlotter
/gate/output/plotter/setNColumns 2
/gate/output/plotter/setPlotHeight 250
/gate/output/plotter/setPlotWidth 300
/gate/output/plotter/addPlot hist Ion_decay_time_s
/gate/output/plotter/addPlot hist Positron_Kinetic_Energy_MeV
/gate/output/plotter/addPlot tree Singles comptonPhantom
/gate/output/plotter/addPlot tree Coincidences energy1
/gate/output/plotter/listPlots
```

The command:

```
addPlot hist NAME_of_the_histo
```

plots an histogram previously defined in GATE, and:
addPlot tree NAME_of_the_tree NAME_of_the_variable

plots a variable from one of the GATE trees.

The command \texttt{setNColumns} sets the number of display windows to be used.

Figure 10.2 presents an example of online plotter, obtained with the above macro.

![Figure 10.2: The Online Plotter](http://wiki.opengatecollaboration.org/index.php/Users_Guide_V7.0:Data_output)

Interfile output of projection set

Description

The Interfile format is especially suited for acquisition protocol using a multiple headed rotating gamma camera. The total description of the Interfilev3.3 format can be found on the Interfile website: http://www.medphys.ucl.ac.uk/interfile/index.htm.

When images are acquired in multiple windows (e.g. energy windows, time windows, multiple heads), the images are recorded according to the order in which the corresponding keys are defined. Thus if multiple energy windows are used, all image data for the first window must be given first, followed by the image data for the second window, etc. This loop structure is defined in the Interfile syntax by the use of the 'for' statement. Two files are created when using the Interfile/Projection output: \texttt{your_file.hdr} and \texttt{your_file.sin}.

The header file contains all information about the acquisition while the \texttt{your_file.sin} file contains the binary
information. An example of such a header is:

```plaintext
!INTERFILE :=
  imaging modality := nucmed
  version of keys := 3.3
  date of keys := 1992:01:01
;
!GENERAL DATA :=
  data description := GATE simulation
  data starting block := 0
  name of data file := your_file.sin
;
!GENERAL IMAGE DATA :=
  type of data := TOMOGRAPHIC
  total number of images := 64
  study date := 2003:09:15
  study time := 11:42:34
  imagedata byte order := LITTLEENDIAN
  number of energy windows := 1
;
!SPECT STUDY (general) :=
  number of detector heads := 2
;
!number of images/energy window := 64
!process status := ACQUIRED
!number of projections := 32
!matrix size [1] := 16
!number format := UNSIGNED INTEGER
!number of bytes per pixel := 2
!scaling factor (mm/pixel) [1] := 1
!scaling factor (mm/pixel) [2] := 1
!extent of rotation := 180
!time per projection (sec) := 10
!study duration (elapsed) sec := 320
!maximum pixel count := 33
;
!SPECT STUDY (acquired data) :=
  direction of rotation := CW
  start angle := 0
  first projection angle in data set := 0
  acquisition mode := stepped
  orbit := circular
  camera zoom factor := 1
;
!number of images/energy window := 64
!process status := ACQUIRED
!number of projections := 32
!matrix size [1] := 16
!number format := UNSIGNED INTEGER
!number of bytes per pixel := 2
!scaling factor (mm/pixel) [1] := 1
!scaling factor (mm/pixel) [2] := 1
!extent of rotation := 180
!time per projection (sec) := 10
!study duration (elapsed) sec := 320
!maximum pixel count := 36
;
!SPECT STUDY (acquired data) :=
  direction of rotation := CW
  start angle := 180
  first projection angle in data set := 180
  acquisition mode := stepped
  orbit := circular
  camera zoom factor := 1
;
!SPECT STUDY (acquired data) :=

DGEGEOMETRY :=
  head x dimension (cm) := 30
```
head y dimension (cm) := 80
head z dimension (cm) := 70
head material := Air
head x translation (cm) := -25
head y translation (cm) := 0
head z translation (cm) := 0
crystal x dimension (cm) := 1.5
crystal y dimension (cm) := 60
crystal z dimension (cm) := 50
crystal material := NaI

GATE SIMULATION :=
number of runs := 32

END OF INTERFILE :=

Use

To use the Interfile output, the following lines have to be added to the macro:

# PROJECTION
/gate/output/projection/enable
/gate/output/projection/setFileName your_file
/gate/output/projection/projectionPlane YZ
/gate/output/projection/pixelSizeY 1. mm
/gate/output/projection/pixelSizeX 1. mm
/gate/output/projection/pixelNumberY 16
/gate/output/projection/pixelNumberX 16

The projectionPlane should be chosen correctly, according to the simulated experiment. The pixelSize and the pixelNumber are always described in a fixed XY-axes system.

Read .sin with ImageJ

There are several way of editing an interfile image with ImageJ.

The Interfile Output is available as a "sin" and "hdr" files directly into the folder of concern. Several software may be used to read the data, among them the software ImageJ is quite often used. The procedure to use is the following:

Once ImageJ is opened, click on the thumb File and select Import -> Raw. A window appears into which the name.sin can be selected.

Once the image is selected, select the following information:

- Image Type: 16-bit Unsigned
- Width & Height & Number of Images can be read into the .hdr files if unknown.
- Tick the case: Little Endian byte Order
- Tick the case: Use Virtual Stack if the data had multiple projection windows.
However one must be careful with this editing. Some users complained that the image in tomographic views provided image in stack in a strange fashion.

A second way to read Interfile images is to use this plugin with ImageJ Interfile Plugin Decoder (http://www.med.harvard.edu/jpnm/ij/plugins/Interfile.html). The advantage is that the plugin seeks all the information in the .hdr files by itself.

**Read .sin with IDL**

For a planar projection, the image projections created with GATE may also be read with IDL with the function `Read_Binary``". In the example below, the projection `name.sin` has to be inserted into the IDL main folder. The image size must be detailed into the `READ_BINARY` function which might lead to a false image if not specified properly. If in doubt, the image size information is to be obtained in the .hdr files.

- **IDL>** file = 'name.sin'
- **IDL>** SizeImageX = 128
- **IDL>** SizeImageZ = 128
- **IDL>** data=READ_BINARY(file,DATA_DIMS=[SizeImageX,SizeImageY],DATA_TYPE=12,ENDIAN='Little')

**Sinogram output**
If the ecat system or the ecatAccel system have been selected (see Users Guide V6.2: Defining a system#Ecat), the sinogram output module can be enable with the following commands:

For the ecat system:

```
/gate/output/sinogram/enable
/gate/output/sinogram/setFileName MySinogramFileName
```

For the ecatAccel system:

```
/gate/output/sinoAccel/enable
/gate/output/sinoAccel/setFileName MySinogramFileName
```

Using this format, the coincident events are stored in an array of 2D sinograms. There is one 2D sinogram per pair of crystal-rings. For example, for the ECAT EXACT HR+ scanner (32 crystal-rings) from CPS Innovations (Knoxville, TN, U.S.A.), there are 1024 2D sinograms. The number of radial bins is specified using the command:

For the ecat system:

```
/gate/output/sinogram/RadialBins 256
```

For the ecatAccel system:

```
/gate/output/sinoAccel/RadialBins 256
```

There is a one-to-one correspondence between the sinogram bins and the lines-of-response (LOR) joining two crystals in coincidence. The sinogram bin assignment is not based on the true radial and azimuthal position of the LOR, but on the indexing of the crystals. This means that the sinograms are subject to curvature effects. By default, all coincident events are recorded, regardless of their origin (random, true unscattered, or true scattered coincidence). It is possible to discard random events:

For the ecat system:

```
/gate/output/sinogram/TruesOnly true
```

For the ecatAccel system:

```
/gate/output/sinoAccel/TruesOnly true
```

In the trues, both scattered and unscattered coincidences are included. There is no simulation of a delayed coincidence window. At the beginning of each run, the content of the 2D sinograms is reset to zero. At the end of each run, the contents of the 2D sinograms can be optionally written to a raw file (one per run). This feature has to be enabled:

For the ecat system:
For the ecatAccel system:

```
/gate/output/sinoAccel/RawOutputEnable
```

Three files are written per run:

- the raw data (unsigned short integer) in MySinogramFileName.ima;
- a mini ASCII header in MySinogramFileName.dim;
- an information file in MySinogramFileName.info.

MySinogramFileName.dim contains the minimal information required to read the flat file *MySinogramFileName.ima*.

Here is an example with the default setting for the ECAT EXACT HR+ scanner:

```
288 288 1024
-type U16
dx 1.0
dy 1.0
dz 1.0
```

The first line specifies the size of the matrix: 1024 2D sinograms (third coordinate) with 288 radial bins (first coordinate) and 288 azimuthal bins (second coordinate). The second line specifies the format: unsigned short integer. The next three lines specify the size of each bin; there are set arbitrarily to 1.

'MySinogramFileName.info' describes the ordering of the 2D sinograms in the flat file MySinogramFileName.ima.

Here is an example with the default setting for the ECAT EXACT HR+ scanner:

```
1024 2D sinograms
[RadialPosition;AzimuthalAngle;AxialPosition;RingDifference]
RingDifference varies as 0,+1,-1,+2,-2, ...,+31,-31
AxialPosition varies as |RingDifference|,...,62-|RingDifference| per increment of 2
AzimuthalAngle varies as 0,...,287 per increment of 1
RadialPosition varies as 0,...,287 per increment of 1
Date type : unsigned short integer (U16)
```

Each 2D sinogram is characterized by the two crystal-rings in coincidence ring1 and ring2. Instead of indexing the 2D sinograms by ring1 and ring2, they are indexed by the ring difference ring2 – ring1 and the axial position ring2 + ring1:

```
ring_1 = (AxialPosition - RingDifference)/2
ring_2 = RingDifference + (AxialPosition - RingDifference)/2
Write Sinogram(ring_1;ring_2)
```
In addition to the sinogram output module, there is a conversion of the 2D sinograms to an ecat7 formatted 3D sinogram in the ecat7 output module. This 3D sinogram is then written to an ecat7 matrix file.

**Ecat7 output**

If and only if both the ecat system and the sinogram output module have been selected, the ecat7 output module can be enable using the following commands:

```
/gate/output/ecat7/enable
/gate/output/ecat7/setFileName MySinogramFile
```

This module writes the content of the 2D sinograms defined in the sinogram output module to an ecat7 formatted matrix scan file, the native file format from CPS Innovations (Knoxville (TN), U.S.A.) for their ECAT scanner family. Due to the large size of a full 3D PET data set, the data set size is reduced before writing it to disk. Therefore it is not possible to go back from an ecat7 formatted 3D sinogram to the original 2D sinograms set.

**Installation**

In order to compile the ecat7 output module of Gate, the ecat library written at the PET Unit of the Catholic University of Louvain-la-Neuve (UCL, Belgium) is required. It can be downloaded from their web site: http://www.topo.ucl.ac.be/ecat_Clib.html

Three files are required: the library file libecat.a and the two header files matrix.h and machine_indep.h.

To compile Gate with the ecat7 library without changing the env_gate.csh and GNUmakefile files, the environment variable ECAT7_HOME has to be defined and set to the name of the home directory where the ecat7 library is installed (for example, /usr/local/ecat7). In this ecat7 home directory, two subdirectories should be created: lib and include. The header files are put in the ${ECAT7_HOME}/include directory. For each system, a specific subdirectory named after the G4SYSTEM environment variable value should be created in the ${ECAT7_HOME}/lib directory. The corresponding library file libecat.a has to be located in this ${ECAT7_HOME}/lib/${G4SYSTEM} directory. The `matrix.h` file has to be modified to add the declaration of the `mh_update()` function. The following line can be added in the "high level user functions" part of matrix.h:

```c
int mh_update(MatrixFile*);
```

**Data reduction**

The polar coordinate of a LOR is approximately defined by the crystal-ring index difference between the 2 rings in coincidence. For a scanner with N crystal rings, the total number of polar samples is given by $2 \times N - 1$. Usually, on ecat systems, not all crystal-ring differences are recorded. Only absolute crystal-ring differences up to a given value, referred to as the maximum ring difference, are recorded. In Gate, this maximum ring difference is defined using:
The value of the maximum ring difference should be smaller than \( N \).

A polar mashing is applied to group 2D sinograms with adjacent polar coordinates. The size of this grouping is called the span [reference]. Its minimum value is 3 and it should be an odd integer. The span value can be set using:

\[
\text{/gate/output/ecat7/span 9}
\]

The Michelogram represented in Figure 10.3 graphically illustrates mashing in the polar coordinate for a 16 crystal-ring scanner with a maximum ring difference set to 12 and a span factor of 5, resulting to 5 polar samples instead of 31. Each dot represents a 2D sinogram for a given pair of crystal-rings. The grouped 2D sinograms are connected by diagonal lines.

By default, the maximum ring difference is set to \( N - 1 \) and the span factor to 3. After choosing a maximum ring difference value \( \text{MaxRingDiff} \), only certain \textit{span} factors are possible as the resulting number of polar samples must be an integer:

\[
\frac{2 \times \text{MaxRingDiff}}{\text{span}} + 1
\]

In addition to the polar mashing, the number of azimuthal samples can also be reduced from \( N_{\text{azi}} = N_{\text{cryst}} / 2 \) to \( N_{\text{azi}} / m \) where \( m \) is the mashing factor. The mashing factor can be set using:

\[
\text{/gate/output/ecat7/mashing 2}
\]
The default mashing value is 1.

**Sinogram file**

At the end of each run, a new 3D sinogram is written with an incremental frame indexing. For example, with the following configuration:

```plaintext
/gate/application/setTimeSlice 60 s
/gate/application/setTimeStart 0 s
/gate/application/setTimeStop 300 s
```

5 frames of 60 seconds each will be generated.

The *ECAT* code of the scanner model is specified by

```plaintext
/gate/output/ecat7/system 962
```

This information can be needed by some *ecat7* based reconstruction routines.

It should be noted that not all fields of the main- or sub-header are filled. In particular, the `coincidence_sampling_mode` field of the main-header is always set to *Prompts and Delayed* (1), regardless of the value of the `/gate/output/sinogram/TruesOnly` tag.

For the scan sub-header, the value of the *prompts* field is correctly filled and the value of the *delayed* field is set to the actual number of random coincidences, and not to the number of delayed coincidences (not simulated).

The radial bin size in the scan sub-header is set to half the value of the crystal transverse sampling and does not take into account the arc and depth-of-interaction (DOI) effects. After arc correction, the radial bin size should be slightly increased to account for the DOI effect. Note that this correction is included in the reconstruction software provided with the *ECAT* scanners.

![Figure 10.4: Increase of the radial bin size due to the DOI effect.](image)

**LMF output**

**Introduction**
The Crystal Clear Collaboration has developed a List Mode Format (LMF) to store the data of ClearPET prototypes. Monte Carlo data generated by GATE can also be stored under the same format using the class GateToLMF. This format is only available for the cylindricalPET system (see Users Guide V6.2:Defining a system) and GATE can only store single events.

Several tools enabling the reading of this format and the processing of events are implemented in the LMF library. As an example, coincidences can be created from GATE single events. It is also possible to apply different deadtimes, and even to generate sinograms in the Interfile format as used by the STIR library, which implements several image reconstruction algorithms.

The LMF library and its documentation are available on the OpenGate web site.

### Size of information to be stored in LMF.

<table>
<thead>
<tr>
<th>Information</th>
<th>Size (bytes/single)</th>
<th>Real machines</th>
<th>GATE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>8</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>Energy</td>
<td>1</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>detector ID</td>
<td>2</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>PET's axial position</td>
<td>2</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>PET's angular position</td>
<td>2</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>run ID</td>
<td>4</td>
<td>NO</td>
<td>YES</td>
</tr>
<tr>
<td>event ID</td>
<td>4</td>
<td>NO</td>
<td>YES</td>
</tr>
<tr>
<td>source ID</td>
<td>2</td>
<td>NO</td>
<td>YES</td>
</tr>
<tr>
<td>source XYZ Position</td>
<td>6</td>
<td>NO</td>
<td>YES</td>
</tr>
<tr>
<td>global XYZ Position</td>
<td>6</td>
<td>NO</td>
<td>YES</td>
</tr>
<tr>
<td>number of Compton in phantomSD</td>
<td>1</td>
<td>NO</td>
<td>YES</td>
</tr>
<tr>
<td>number of Compton in crystalSD</td>
<td>1</td>
<td>NO</td>
<td>YES</td>
</tr>
</tbody>
</table>

### Usage

LMF data are composed of two files with the same base-name, but different extensions:

- An ASCII file with a .cch extension contains general information about the scan and about the scanner, like the scan duration, the sizes of the detectors, or the angular rotation speed.
- A binary file with a .ccs extension contains headers, which set the topology of the scanner, followed by fixed size records.

The user can generate these two output files automatically by using the macro scripting. Scripting also allows to select the kind of information to be stored. All pieces of information are optional, except time, which makes the ClearPET LMF quite versatile. Table 10.7.2 lists all options and memory requirements that can be stored in the LMF event record when using the cylindricalPET system.

The binary output file size depends on its content. It amounts to 11 MB for 1 million single events stored with their time, energy and detector ID for a small animal PET scanner including about 1500 crystals.

Macros commands (available only once initialisation has been performed) used to configure the LMF output.
are:

```
/gate/output/lmf/enable
```

to enable LMF output.

```
/gate/output/lmf/disable
```

to disable LMF output (but it is disable by default).

```
/gate/output/lmf/setFileName myFirst
```

to set the LMF files name. Here the output files will be myFirst.ccs and myFirst.cch.

```
/gate/output/lmf/setDetectorIDBool 1
```

to store (1) or to not store (0) the detector ID.

```
/gate/output/lmf/setEnergyBool 1
```

to store (1) or to not store (0) the energy.

```
/gate/output/lmf/setGantryAxialPosBool 0
```

to store (1) or to not store (0) the axial position.

```
/gate/output/lmf/setGantryAngularPosBool 0
```

to store (1) or to not store (0) the angular position.

The following lines must always be included as they appear below with option set to 0:

```
/gate/output/lmf/setSourcePosBool 0
/gate/output/lmf/setNeighbourBool 0
/gate/output/lmf/setNeighbourhoodOrder 0
/gate/output/lmf/setCoincidenceBool 0
```

All information that is not available in real acquisitions is stored in a `GateDigi record` using the command:

```
/gate/output/lmf/setGateDigiBool 1
```

If the option 0 is used, the following commands are ignored

```
/gate/output/lmf/setComptonBool 1
```
to store (1) or to not store (0) the number of Compton scattering that occurred in a *phantomSD* and a *crystalSD*.

```xml
/gate/output/lfm/setSourceIDBool 0
```

to store (1) or to not store (0) the source ID.

```xml
/gate/output/lfm/setSourceXYZPosBool 0
```

to store (1) or to not store (0) the source XYZ position.

```xml
/gate/output/lfm/setGlobalXYZPosBool 0
```

to store (1) or to not store (0) the real XYZ position.

The information regarding the gantry position, translation or rotation speed(s), or the position of the eccentric rotation axis are automatically passed from the macro scripting to the LMF output.

```xml
/gate/output/lfm/setEventIDBool 1
```

to store (1) or to not store (0) the event ID.

```xml
/gate/output/lfm/setRunIDBool 1
```

to store (1) or to not store (0) the run ID.

**Limitations**

The LMF format was originally designed for the development of small animal PET scanners for which the number of crystals is smaller than for clinical PET scanners. Consequently, the user should carefully read the LMF specifications and make sure that this format allows him to model his scanner design. In particular, the maximum number of sub-volumes in a volume (e.g. the maximum number of submodules in a module) is set by the number of bits used to encode the sub-volume ID. The final ID encoding the position of an event has to be stored on 16, 32, or 64 bits only.

**Image CT output**

The *imageCT* output is a binary matrix of float numbers that stores the simulated CT image and is produced for each time slice.

The output is enabled by:

```xml
/gate/output/imageCT/enable
```
The output file name is set by:

```
/gate/output/imageCT/setFileName test
```

The output file name is "test_xxx.dat", where xxx is the corresponding time slice number.

In the case of the fast simulation mode, the number of pixels is set by:

```
/gate/output/imageCT/numPixelX 80
/gate/output/imageCT/numPixelY 80
```

In the case of VRT simulation mode (see Users Guide V6.2:Defining a system#CTscanner), the VRT K factor is set by:

```
/gate/output/imageCT/vrtFactor 10
```

Finally the random seed can be defined using:

```
/gate/output/imageCT/setStartSeed 676567
```


- This page was last modified on 9 January 2014, at 08:50.
Users Guide V7.0:Generating and tracking optical photons

From GATE collaborative documentation wiki

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- Bibliography

!!! Important if using Geant4.9.5 and Geant4.9.5.p01 !!!

All propertyvector in both Surface.xml and Materials.xml entries have to be given in increasing order of energy.

(see http://hypernews.slac.stanford.edu/HyperNews/geant4/get/opticalphotons/458/1/1.html)

Introduction

This chapter explains how Gate can be used to generate and track optical photons in order to investigate, for example, the energy resolution or spatial resolution of detectors.

To use the optical photon capabilities of GATE, the GATE_USE_OPTICAL variable has to be set to ON in the configuration process using ccmake.

Before discussing how to use the optical photon tracking, it has to be mentioned that there are a few disadvantages in using optical transport. First, the simulation time will increase dramatically. For example, most scintillators used in PET generate in the order of 10,000 optical photons at 511 keV, which means that approximately 10,000 more particles have to be tracked for each annihilation photon that is detected. Although the tracking of optical photons is relatively fast, a simulation with optical photon tracking can easily be a factor thousand slower than one without. Finally, in order to perform optical simulations, many parameters are needed for the materials and surfaces, some of which may be difficult to determine.

Optical Photon Generation

```
/gate/source/addSource                           Mysource gps
/gate/source/Mysource/gps/particle               opticalphoton
/gate/source/Mysource/gps/energytype             Mono
/gate/source/Mysource/gps/angtype                iso
```

An optical photon with a wave length of 530nm corresponds to an optical photon of energy=2.34eV [approximation: 1240/E(eV) = wavelength (nm)]:

```
/gate/source/Mysource/gps/monoenergy             2.34 eV
```
An optical photon which is not assigned a polarization at production may not be Rayleigh scattered:

```
gate/source/Mysource/gps/polarization 1 0 0
```

**Optical System**

The GATE OpticalSystem is appropriate to model Optical Imaging systems. This system is defined in the following section of the Users Guide: "OpticalSystem" ([http://wiki.opengatecollaboration.org/index.php/Users_Guide_V6.2:Defining_a_system#OpticalSystem](http://wiki.opengatecollaboration.org/index.php/Users_Guide_V6.2:Defining_a_system#OpticalSystem)).

**Defining material properties**

The optical properties of materials are stored in a material property table. In this table each of the properties of a material is identified by a name. There are two kinds of properties. The first are constant properties, these contain only one value. The second are property vectors, these contain properties that depend on the energy of the optical photon. Such a vector is a list of energy-value pairs.

The property tables for the materials used in a simulation are to be stored in a file separate from the material database. This makes it easier to change the properties without having to change the material database. This file should be named Materials.xml. When Gate reads in a material from the materials database, it also checks if the Materials.xml file contains a property table for this material. If so, this table is read in and coupled to the material.

**Scintillation**

A scintillator is characterized by its photon emission spectrum. The scintillation follows an exponential decay with two time constants, a fast and a slow one. The relative strength of the fast component FASTCOMPONENT as a fraction of total scintillation yield is given by the YIELDRATIO. The emission spectra of both decays are given by the property vectors FASTCOMPONENT and SLOWCOMPONENT and the time constants FASTTIMECONSTANT and SLOWTIMECONSTANT. These vectors specify the probability that a photon with the given energy is emitted. The sum of each of the vectors should therefore be one.

In order to have scintillation in a material, the first parameter that has to be specified is the SCINTILLATIONYIELD (1/Mev, 1/keV), which gives the number of photons that is emitted per amount of energy absorbed, or, more precisely, it gives the expectation value of this number, since the real number of emitted photons follows a normal distribution. The variance of this normal distribution is RESOLUTION-SCALE times this expectation value. Thus, for example, when a gamma photon deposits $E$ amount of energy in the scintillator, $N$ optical photons are emitted with an expectation value of $\bar{N} = E \cdot \text{SCINTILLATIONYIELD}$ and a standard deviation of $\sigma_N = \text{RESOLUTIONSCALE} \cdot \sqrt{E \cdot \text{SCINTILLATIONYIELD}}$.

The parameters RESOLUTIONSCALE can be calculated from the energy resolution of the scintillator. The energy resolutions specified in the literature may contain contributions of electronic noise. The energy resolution needed to calculate the RESOLUTIONSCALE should be the intrinsic energy resolution of the scintillator.

$$\text{RESOLUTIONSCALE} = \frac{R}{2.35} \cdot \sqrt{E \cdot \text{SCINTILLATIONYIELD}},$$

where $R$ is the energy resolution (FWHM - Full width at half maximum) at energy $E$.

```
<material name="LSO">
  <propertiestable>
    <property name="SCINTILLATIONYIELD" value="26000" unit="1/MeV"/>
    <property name="RESOLUTIONSCALE" value="4.41"/>
    <property name="FASTTIMECONSTANT" value="40" unit="ns"/>
    <property name="YIELDRATIO" value="1"/>
    <propertyvector name="FASTCOMPONENT" energyunit="eV">
      <ve energy="2.95167" value="1"/>
    </propertyvector>
    <propertyvector name="ABSLENGTH" unit="m" energyunit="eV">
      <ve energy="1.84" value="50"/>
      <ve energy="4.08" value="50"/>
    </propertyvector>
    <propertyvector name="RINDEX" energyunit="eV">
      ...
    </propertyvector>
  </propertiestable>
</material>
```
Absorption

This process kills the particle. It requires the Material.xml properties filled by the user with the Absorption length $\text{ABSLENGTH}$ (average distance traveled by a photon before being absorbed by the medium).

```
<gate/physics/addProcess OpticalAbsorption
```

Mie/Rayleigh Scattering

Mie Scattering is an analytical solution of Maxwell’s equations for scattering of optical photons by spherical particles. It is significant only when the radius of the scattering object is of order of the wave length. The analytical expressions for Mie Scattering are very complicated. One common approximation (followed by Geant4) made is called Heney-Greenstein (HG). For small size parameter (scattering particle diameter) regime the Mie theory reduces to the Rayleigh approximation.

```
<gate/physics/addProcess OpticalRayleigh
<gate/physics/addProcess OpticalMie
```

For Rayleigh or Mie scattering, we require the final momentum, initial polarization and final polarization to be in the same plane. Mie/Rayleigh processes require material properties to be filled by the user with Mie/Rayleigh scattering length data: $\text{MIEHG/RAYLEIGH}$, which is the average distance traveled by a photon before it is Mie/Rayleigh scattered in the medium. In the case of the Mie scattering, the user also needs to provide parameters of the HG approximation: $\text{MIEHG_FORWARD}$ (forward anisotropy), $\text{MIEHG_BACKWARD}$ (backward anisotropy), and $\text{MIEHG_FORWARD_RATIO}$ (ratio between forward and backward angles). Geant4 code allows the forward and backward angles to be treated separately. If your material characteristics provides only one number for the anisotropy (= average cosine of the scattering angle), below is an example of how (part of) the Materials.xml file could look like:

```
<material name="Biomimic">
  <propertiestable>
    <propertyvector name="ABSLENGTH" unit="cm" energyunit="eV">
      <ve energy="1.97" value="0.926"/>
      <ve energy="2.34" value="0.847"/>
    </propertyvector>
    <propertyvector name="RINDEX" energyunit="eV">
      <ve energy="1.97" value="1.521"/>
      <ve energy="2.34" value="1.521"/>
    </propertyvector>
    <property name="MIEHG_FORWARD" value="0.62" />
    <property name="MIEHG_BACKWARD" value="0.62" />
    <property name="MIEHG_FORWARD_RATIO" value="1.0" />
    <propertyvector name="MIEHG" unit="cm" energyunit="eV">
      <ve energy="1.97" value="0.043"/>
      <ve energy="2.34" value="0.043"/>
    </propertyvector>
  </propertiestable>
</material>
```

Fluorescence

Fluorescence is a 3 step process: The fluorophore is in an excited state after the absorption of an optical photon provided by an external source (laser, lamp). The life time of the excited state is of order of 1-10ns during which the fluorophore interacts with its environment and ends-up in a relaxed-excited state. The last step is the emission of a fluorescent photon which energy/wave length...
is smaller(larger) than the one of the excitation optical photon.

Geant4 simulates the WaveLength Shifting (WLS) fibers that are used in High Energy Physics experiments. As an example, the CMS hadronic EndCap calorimeter is made of scintillator tiles with WLS fibers embedded. These fibers collect/absorb blue light produced in tiles and re-emit green light so that as much light reaches the PMTs. A new class in Gate has been implemented as a physics builder class that inherits from the G4OpWLS class. The following command line enables the optical photon fluorescence:

```
/gate/physics/addProcess OpticalWLS
```

Gate user needs to provide four parameters/properties to define the fluorescent material: RINDEX, WLSABSLENGTH, WLSCOMPONENT and WLSTIMECONSTANT. The WLSABSLENGTH defines the fluorescence absorption length which is the average distance travelled by a photon before it is absorbed by the fluorophore. This distance could be very small but probably not set to 0 otherwise the photon will be absorbed immediately upon entering the fluorescent volume and fluorescent photon will appear only from the surface. The WLSCOMPONENT describes the emission spectrum of the fluorescent volume by giving the relative strength between different photon energies. Usually these numbers are taken from measurements (i.e. emission spectrum). The WLSTIMECONSTANT defines the time delay between the absorption and re-emission.

**Simulation of the Fluorescein** [see http://en.wikipedia.org/wiki/Fluorescein]

We define the refractive index of the fluorophore's environment (water or alcohol):

```
<material name="Fluorescein">
<propertytable>
<vector name="RINDEX" energyunit="eV">
<ve energy="1.0" value="1.4"/>
<ve energy="4.13" value="1.4"/>
</vector>

We describe the fluorescein absorption length taken from measurements or literature as function of the photon energy:

```
<vector name="WLSABSLENGTH" unit="cm" energyunit="eV">
<ve energy="3.19" value="2.81"/>
<ve energy="3.20" value="2.82"/>
<ve energy="3.21" value="2.81"/>
</vector>

We describe the fluorescein Emission spectrum taken from measurements or literature as function of the photon energy:

```
<vector name="WLSCOMPONENT" energyunit="eV">
<ve energy="1.771" value="0.016"/>
<ve energy="1.850" value="0.024"/>
<ve energy="1.901" value="0.040"/>
<ve energy="2.003" value="0.111"/>
<ve energy="2.141" value="0.325"/>
<ve energy="2.210" value="0.540"/>
<ve energy="2.250" value="0.683"/>
<ve energy="2.343" value="0.873"/>
<ve energy="2.384" value="0.968"/>
<ve energy="2.484" value="0.817"/>
<ve energy="2.749" value="0.008"/>
<ve energy="3.099" value="0.008"/>
</vector>
```
Boundary Processes

When a photon arrives at a medium boundary its behavior depends on the nature of the two materials that join at that boundary.

```
/gate/physics/addProcess OpticalBoundary
```

In the case of two dielectric materials, the photon can undergo total internal reflection, refraction or reflection, depending on the photon’s wavelength, angle of incidence, and the refractive indices on both sides of the boundary. In the case of an interface between a dielectric and a metal, the photon can be absorbed by the metal or reflected back into the dielectric.

Defining (or Not) the Surface

When simulating a perfectly smooth surface, the user doesn't have to provide a G4Surface. The only relevant property is the refractive index (RINDEX) of the two materials on either side of the interface. Geant4 will calculate from Snell's Law the probabilities of refraction and reflections. However, should the user want to include possible irregularities of the surface, e.g. surface roughness that isn't specifically defined in the geometrical definition, then the user may want to provide probability options for various reflection types. The UNIFIED model allows the user to control the radiant intensity of the surface: Specular lobe, Specular spike, Backscatter spike (enhanced on very rough surfaces) and Reflectivity (Lambertian or diffuse distribution). The sum of the four constants is constrained to unity. In Gate, the simulation model that is used by the boundary process is ALWAYS the Geant4 UNIFIED model (see: source/geometry/src/GateSurface.cc). In that model, the probability of micro-facet normals that populates the annulus of solid angle \( \sin(\alpha) \, d\alpha \) is proportional to a Gaussian of \( \sigma_{\alpha} \) given in degrees. This parameter defines the standard deviation of the Gaussian distribution of micro-facets around the average surface normal. In the case of a perfectly polished surface, the normal used by the G4BoundaryProcess is the normal to the surface.

Defining surfaces

On this image, the photon travels through the surface between Volume1 and Volume2. To create a surface with the name Surface-From-Volume1-To-Volume2 between the two volumes with the names Volume1 and Volume2, the following commands should be used:

```
/gate/Volume2/surfaces/name Surface-From-Volume1-To-Volume2
```

Specular lobe, Specular spike, Backscatter spike (enhanced on very rough surfaces), and Reflectivity (Lambertian or diffuse distribution).
The surface between Volume1 and Volume2 is NOT the same surface as that between Volume2 and Volume1. When there is optical transport in both directions, both surfaces should be created.

To load the surface properties stored under rough_teflon_wrapped in the Surface.xml file:

```
/gate/Volume2/surfaces/Surface-From-Volume1-To-Volume2/SetSurface rough_teflon_wrapped
```

An example of a surface definition looks like:

```xml
<surface name="rough_teflon_wrapped" type="dielectric_dielectric" sigmaalpha="0.1" finish="groundbackpainted">
    <propertytable>
        <propertyvector name="SPECULARLOBECONSTANT" energyunit="eV">
            <ve energy="4.08" value="1"/>
            <ve energy="1.84" value="1"/>
        </propertyvector>
        <propertyvector name="RINDEX" energyunit="eV">
            <ve energy="4.08" value="1"/>
            <ve energy="1.84" value="1"/>
        </propertyvector>
        <propertyvector name="REFLECTIVITY" energyunit="eV">
            <ve energy="1.84" value="0.95"/>
            <ve energy="4.08" value="0.95"/>
        </propertyvector>
        <propertyvector name="EFFICIENCY" energyunit="eV">
            <ve energy="1.84" value="0"/>
            <ve energy="4.08" value="0"/>
        </propertyvector>
    </propertytable>
</surface>
```

The attribute type can be either dielectric_dielectric or dielectric_metal, to model either a surface between two dielectrics or between a dielectricum and a metal. The attribute sigma-alpha models the surface roughness and is discussed in the next section. Finally, the attribute finish can have one of the following values: ground, polished, ground-back-painted, polished-back-painted, ground-front-painted and polished-front-painted. It is therefore possible to cover the surface on the inside or outside with a coating that reflects optical photons using Lambertian reflection. In case the finish of the surface is polished, the surface normal is used to calculate the probability of reflection. In case the finish of the surface is ground, the surface is modeled as consisting of small micro-facets. The surface normals of the facets are distributed around the average surface normal, following a normal distribution with standard deviation \( \sigma_\alpha \), which can be specified in degrees in the attribute sigmaalpha. When an optical photon reaches a surface, a random angle \( \alpha \) is drawn for the micro facet that is hit by the optical photon. Using the angle of incidence of the optical photon with respect to this micro facet and the refractive indices of the two media, the probability of reflection is calculated.

In case the optical photon is reflected, four kinds of reflection are possible. The probabilities of the first three are given by the following three property vectors:

- **SPECULARSPIKECONSTANT** gives the probability of specular reflection about the average surface normal
- **SPECULARLOBECONSTANT** gives the probability of specular reflection about the surface normal of the micro facet
- **BACKSCATTERCONSTANT** gives the probability of reflection in the direction the optical photon came from

LAMBERTIAN (diffuse) reflection occurs when none of the other three types of reflection happens. The probability of Lambertian reflection is thus given by one minus the sum of the other three constants.

![Specular, diffuse, and spread reflection from a surface.](image_url)
When an optical photon reaches a painted layer, the probability of reflection is given by the property vector \textbf{REFLECTIVITY}. In case the paint is on the inside of the surface, the refractive indices of the media are ignored, and when the photon is reflected, it undergoes Lambertian reflection.

When the paint is on the outside of the surface, whether the photon is reflected on the interface between the two media is calculated first, using the method described in the previous section. However, in this case the refractive index given by the property vector \textit{RINDEX} of the surface is used. When the photon is refracted, it is reflected using Lambertian reflection with a probability \textbf{REFLECTIVITY}. It then again has to pass the boundary between the two media. For this, the method described in the previous section is used again and again, until the photon is eventually reflected back into the first medium or is absorbed by the paint.

A dielectric-dielectric surface may have a wavelength dependent property \textbf{TRANSMITTANCE}. If this is specified for a surface it overwrites the Snell's law's probability. This allows the simulation of anti-reflective coatings.

**Detection of optical photons**

Optical photons can be detected by using a dielectric-metal boundary. In that case, the probability of reflection should be given by the \textit{REFLECTIVITY} property vector. When the optical photon is reflected, the UNIFIED model is used to determine the reflection angle. When it is absorbed, it is possible to detect it. The property vector \textit{EFFICIENCY} gives the probability of detecting a photon given its energy and can therefore be considered to give the internal quantum efficiency. Note that many measurements of the quantum efficiency give the external quantum efficiency, which includes the reflection: external quantum efficiency = efficiency*(1-reflectivity).

The hits generated by the detection of the optical photons are generated in the volume from which the optical photons reached the surface. This volume should therefore be a sensitive detector.

**Digitizer**

The hits generated in the sensitive detector are first processed by \textit{analysis}. Unfortunately \textit{analysis} is quite slow when there are a large number of hits, as is the case when there is optical transport. Therefore, an alternative has been created that is faster and is therefore called fast\textit{analysis}.

```
gate/output/analysis/disable
/gate/output/fastanalysis/enable
```

Switching both on has no effect on the results, but only affects the speed of the simulation. After processing the hits with one of the analysis routines, the singles should be created from the hits. This is usually done using the \textit{opticaladder} which adds all hits generated by optical photons. In this way, it is possible to create a digitizer chain containing the singles generated by optical photons.

```
gate/digitizer/Singles/insert opticaladder
/gate/digitizer/Singles/insert readout
/gate/digitizer/Singles/readout/setDepth your_detector_readout_level
```

Digitizer modules like \textit{threshold} or \textit{uphold} can be used (see "Thresholder & Upholder" (http://wiki.opengatecollaboration.org/index.php/Users_Guide_V6.2:Digitizer_and_readout_parameters#Thresholder_26_Upholder)). This is crucial when you do a fluorescence experience for example. If you want to detect only fluorescent photons you need to apply an energy cut (upholder) in order to discard high energy photons (non-fluorescent photons have higher energy than fluorescent photons).

```
gate/digitizer/Singles/insert upholder
/gate/digitizer/Singles/upholder/setUphold 2.0 eV
/gate/digitizer/Singles/insert thresholder
/gate/digitizer/Singles/thresholder/setThreshold 1.0 eV
```

The \textit{projection} (see Projection set) associated to this digitizer records only photons corresponding to the defined energy window. The projection image is therefore the fluorescence image.
Optical Imaging Simulation Outputs

Root output

When working with optical photons, an additional ROOT tree is created: OpticalData. You can decide to fill this tree or not by using the following command:

```
/gate/output/root/setRootOpticalFlag 0 or 1
```

OpticalData tree is generated with the following information:

- `CrystalLastHitEnergy`  `CrystalLastHitPos_X`  `CrystalLastHitPos_Y`  `CrystalLastHitPos_Z`  
  Energy and Positions of the photon last hit in the Crystal (Detected photon position)

- `PhantomLastHitEnergy`  `PhantomLastHitPos_X`  `PhantomLastHitPos_Y`  `PhantomLastHitPos_Z`  
  Energy and Positions of the photon last hit in the Phantom

- `NumCrystalWLS`  
  Number of Fluorescence processes per event(photon) in the Crystal

- `NumPhantomWLS`  
  Number of Fluorescence processes per event(photon) in the Phantom

- `NumScintillation`  
  Number of Scintillation processes per event(photon) in the Crystal

- `CrystalProcessName`  `PhantomProcessName`  
  List of process names that occurred in the Crystal or in the Phantom

- `MomentumDirectionx`  `MomentumDirectiony`  `MomentumDirectionz`  
  Optical photon momentum direction

Binary output of projection set

In order to create a projection set (see Interfile output of projection set) using the Optical System in GATE, the following lines have to be added to the macro:

```
/gate/output/projection/enable
/gate/output/projection/setFileName        your_name
/gate/output/projection/projectionPlane    XY
/gate/output/projection/pixelSizeX        0.105 cm
/gate/output/projection/pixelSizeY        0.105 cm
/gate/output/projection/pixelNumberX      100
/gate/output/projection/pixelNumberY      100
```

The result of projection set is saved in a binary file (.bin). A header file (.hdr) is also provided with the following information:

```
!INTERFILE :=
!imaging modality := optical imaging

!GENERAL DATA :=
data description := GATE simulation
!name of data file := ./OpticalSimulationProjection.bin

!GENERAL IMAGE DATA :=
type of data := OPTICAL
!total number of images := 1

!OPTICAL STUDY (general) :=
number of detector heads := 1
```

EXAMPLE: Optical Imaging Simulation

In the GATE software you will find simple examples of a bioluminescence/fluorescence experiment. All macros are located in `example/example_OPTICAL`. In addition, a ROOT macro `[DrawBranches.C]` is available and draws all branches of the OpticalData tree into a postscript file.

The optical imaging system is composed of an array of pixels, an electronic board and an angular aperture that limits the range of angles over which the optical system can accept light. The phantom is composed of a box of water and two layers made of either water, hypodermis or epidermis. In case of a bioluminescence experiment, the tumor is described as a voxelized source of optical photons and is positioned under the inner layer of the phantom. In case of a fluorescence experiment, we assigned the Rhodamine B fluorophore to each voxel of a voxelized tumor and positioned it under the inner layer of the phantom. The fluorophore is excited by two external beam light sources emitting optical photons towards the tumor.

These two experiments are available in `example/example_OPTICAL` through the following macros: `bioluminescence.mac` and `fluorescence.mac`. The voxelized source or phantom is available in `example/example_OPTICAL/voxelized-source-phantom` with an attenuation file and an optical-flux file. These macros will generate a root output file with the OpticalData tree enabled and a binary file which corresponds to the GATE ProjectionSet on the XY plane (i.e. detection plane). Using the root macros `MakeBioluminescencePlots.C` and `MakeFluorescencePlots.C`, you can read the root output file and draw the bioluminescent/fluorescent light that is detected by the optical system. In case of the fluorescence experiment, two plots are drawn: all detected light (any wavelength) and the fluorescent light (wavelength cut). The projection binary file (.bin and .hdr) can be
viewed directly using Anatomist or ImageJ. In case of the fluorescence experiment, an Upholder (uphold cut) was applied through the digitizer so the binary image illustrates the fluorescent light.

The Materials.xml file is updated with several tissues properties at specific wavelengths (from literature): brain, kidney, epidermis and hypodermis but also with the emission spectra of the Fluorescein and Rhodamine B.

Bibliography


- Optical Absorption of Water (http://omlc.ogi.edu/spectra/water/index.html) with all currently available data (presented in terms of wavelength and absorption coefficient) - Scott Prahl, Oregon Medical Laser Center.


- This page was last modified on 13 May 2014, at 08:17.
Users Guide V7.0: Architecture of the simulation for radiotherapy

From GATE collaborative documentation wiki

Main rules

GATE simulations are based on the execution of scripted commands (Users Guide V6.2: Getting started) gathered in macros. We describe here the most important points which are specific to define simulation for radiotherapy applications. The general set-up should be generally divided into 6 steps as follows:

1. Verbosity and Visualization (Users Guide V6.2: Visualization)
3. Actor and output management (Users Guide V6.2: Readout parameters for Radiotherapy applications: Actors)
4. Physics (Users Guide V6.2: Setting up the physics)
5. Sources (Users Guide V6.2: Activity, source, voxelized phantoms)
6. Experiment - How to run the simulation? (Users Guide V6.2: Getting started)

from gate/application/setTotalNumberOfPrimaries [Nparticle]
gate/application/start

The first 4 steps correspond to the PreInit mode of GEANT4 whereas the last 3 occur after the initialization of the simulation. The first 4 steps are validated by the GEANT4 command:

gate/run/initialize

Once this phase is completed, the sources can be inserted in the setup and the simulation can be run.

Retrieved from

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Users Guide V7.0: Readout parameters for Radiotherapy applications: Actors

From GATE collaborative documentation wiki

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- Filters
  - Filter on particle type
  - Filter on particle ID
  - Filter on volume
  - Filter on energy
  - Filter on direction

Actors are tools which allow to interact with the simulation. They can collect information during the simulation, such as energy deposit, number of particles created in a given volume, etc. They can also modify the behavior of the simulation. Actors use hooks in the simulation: run (begin/end), event (begin/end), track (begin/end), step. There are different types of actors which collect different types of information, however some commands and behavior are common to all actors. To use selection criteria, it is possible to add filters.

General Purpose

Add an actor

To add an actor, the command is:

```
//gate/actor/addActor [Actor Type] [Actor Name]
```

Attach to a volume

Tells that the actor is attached to the volume [Volume Name]. For track and step levels, the actor is activated for step inside the volume and for tracks created in the volume. If no attach command is provided then the actor is activated in any volume. The
children of the volume inherit the actor.

\[
\text{/gate/actor/[Actor Name]/attachTo [Volume Name]}
\]

### Save output

This command allow to save the data of the actor to the file [File Name]. The particular behaviour (format, etc.) depends on the type of the actor.

\[
\text{/gate/actor/[Actor Name]/save [File Name]}
\]

It is possible to save the output every N events with the command:

\[
\text{/gate/actor/[Actor Name]/saveEveryNEvents [N]}
\]

It is possible to save the output every N seconds with the command:

\[
\text{/gate/actor/[Actor Name]/saveEveryNSeconds [N]}
\]

### 3D matrix actor (Image actor)

Some actors, such as the dose actor, can store some information into a 3D rectangular image (or matrix) according to the spatial position of the hit. User can specify the resolution of the 3D matrix (in this case, the size is equal to the size of the bounding box of the attached volume). Alternatively, user can specify the size to allow larger of smaller matrices.

- "attachTo". the main principle is that scoring value is stored in the 3D matrix only when a hit occur in the attached volume. If the size of the volume is greater than the 3D matrix, hit occurring out of the matrix are not recorded. Conversely, if the 3D matrix is larger than the attached volume, part which are outside the volume will never record hit (even if it occurs) because hit is performed out of the volume.
- "type". In Geant4, when a hit occur, energy is deposited along a step line. A step is defined by two positions the PreStep and the PostStep. The user can choose at which position the actor have to stored the information (edep, dose ...) : it can be at PreStep (pre), at PostStep (post), at the middle between PreStep and PostStep (middle) or distributed from PreStep to PostStep (random). According to the matrix size, such line can be located inside a single dosel or cross several dosels. Preferred type of hit is "random", meaning that a random position is computed along this step line and all the energy is deposited inside the dosel that contains this point.
- the attached volume can be a voxelized image the scoring matrix volume (dosels) are thus different from the geometric voxels describing the image.

\[
\text{/gate/actor/[Actor Name]/attachTo waterbox}
\]
\[
\text{/gate/actor/[Actor Name]/setSize 5 5 5 cm}
\]
\[
\text{/gate/actor/[Actor Name]/voxelsize 10 20 5 mm}
\]
\[
\text{/gate/actor/[Actor Name]/setPosition 1 0 0 mm}
\]
\[
\text{/gate/actor/[Actor Name]/stepHitType random}
\]

### Actor list

#### Simulation statistic

This actor counts the number of steps, tracks, events, runs in the simulation. If the actor is attached to a volume, the actor counts the number of steps and tracks in the volume. The output is an ASCII file.

\[
\text{/gate/actor/addActor SimulationStatisticActor MyActor}
\]
\[
\text{/gate/actor/MyActor/save MyOutput.txt}
\]

### Electromagnetic (EM) properties
This actor allows extracting EM properties for all materials defined in a simulation, as listed below:

- **Density** (mass density in g/cm$^3$)
- **$e$-density** (electronic density in e-/mm$^3$)
- **RadLength** (radiation length in mm)
- **$I$** (ionization potential in eV)
- **EM-DEDX** (EM mass stopping power in MeV.cm$^2$/g)
- **Nucl-DEDX** (nuclear mass stopping power in MeV.cm$^2$/g)
- **Tot-DEDX** (total mass stopping power in MeV.cm$^2$/g)

EM properties are calculated relative to a specific particle type and energy, as defined by the user. For instance, EM properties corresponding to a 30 MeV neutron can be calculated using the following command lines:

```
/gate/actor/addActor EmCalculatorActor MyActor
/gate/actor/MyActor/setParticleName proton
/gate/actor/MyActor/setEnergy 150 MeV
/gate/actor/MyActor/save output/MyOutput.txt
```

**Dose measurement**

This actor builds 3D images of the energy deposited (edep), dose deposited and the number of hits in a given volume. It takes into account the weight of particles. It can store multiple information into a 3D grid, each information can be enable by using:

```
/gate/actor/[Actor Name]/enableEdep             true
/gate/actor/[Actor Name]/enableUncertaintyEdep  true
/gate/actor/[Actor Name]/enableSquaredEdep      true
/gate/actor/[Actor Name]/enableDose             true
/gate/actor/[Actor Name]/enableUncertaintyDose  true
/gate/actor/[Actor Name]/enableSquaredDose      true
/gate/actor/[Actor Name]/enableNumberOfHits     true
```

Informations can be disable by using "false" instead of "true" (by default all states are false):

```
/gate/actor/[Actor Name]/enableEdep             false
```

The unit of edep is MeV and the unit of dose is Gy. The dose/edep squared is used to calculate the uncertainty when the output from several files are added. The uncertainty is the relative statistical uncertainty. "SquaredDose" flag allows to store the sum of squared dose (or energy). It is very useful when using GATE on several workstations with numerous jobs. To compute the final uncertainty, you only have to sum the dose map and the squared dose map to estimate the final uncertainty according to the uncertainty equations.

It is possible to normalize the maximum dose value to 1:

```
/gate/actor/[Actor Name]/normaliseDose   true
```

For the output, the suffixes Edep, Dose, NbOfHits, Edep-Uncertainty, Dose-Uncertainty, Edep-Squared or Dose-Squared are added to the output file name given by the user. You can use several files types: ASCII file (.txt), root file (.root), Analyze (.hdr/.img) and MetaImage (.mhd/.raw) (mhd is recommended!). The root file works only for 1D and 2D distributions.

```
/gate/actor/addActor DoseActor MyActor
/gate/actor/MyActor/save output/MyOutputFile.mhd
/gate/actor/MyActor/attachTo MyVolume
/gate/actor/MyActor/setHitType random
/gate/actor/MyActor/setSize 5 5 5 m
/gate/actor/MyActor/setResolution 1 1 3000
/gate/actor/MyActor/enableEdep true
/gate/actor/MyActor/enableUncertaintyEdep true
/gate/actor/MyActor/enableSquaredEdep false
/gate/actor/MyActor/enableDose false
/gate/actor/MyActor/normaliseDose false
```
Water equivalent doses (or dose to water) can be also calculated, in order to estimate doses calculated using water equivalent path length approximations, such as in Treatment Planning Systems (TPS). Command previously presented for the "dose" also work for the "dose to water" as shown below:

```
/gate/actor/[Actor Name]/enableDoseToWater      true
/gate/actor/[Actor Name]/enableUncertaintyDoseToWater   true
/gate/actor/[Actor Name]/normalizeDoseToWater   true
```

**New image format : MHD**

Gate now can read and write mhd/raw image file format. This format is similar to the previous hdr/img one but should solve a number of issues. To use it, just specify .mhd as extension instead of .hdr. The principal difference is that mhd store the 'origin' of the image, which is the coordinate of the 0.0,0 pixel expressed in the physical world coordinate system (in general in millimetres). Typically, if you get a dicom image and convert it into mhd (vv (http://vv.creatis.insa-lyon.fr) can conveniently do this), the mhd will keep the same pixels coordinate system than the dicom.

In Gate, if you specify the macro "TranslateTheImageAtThisIsoCenter" with the coordinate of the isocenter that is, for example read, in a dicom-rt-plan file, the image will be placed such that this isocenter is at position 0,0,0 of the mother volume (often the world). This is very useful to precisely position the image as indicated in a RT plan. Also, when using a DoseActor attached to a mhd file, the output dose distribution can be stored also in mhd format. In this case, the origin of the dose distribution will be such that it corresponds the the attached image, thus make it possible easy superimposition display in vv.

Note however, that the mhd module is still experimental and not complete. It is thus possible that some mhd images cannot be read. Use and enjoy at your own risk, please contact us if you find bugs and be warmly acknowledged if you correct bugs.

**Kill track**

This actor kills tracks entering the volume. The output is the number of tracks killed. It is stored an ASCII file.

```
/gate/actor/addActor KillActor MyActor
/gate/actor/MyActor/save  MyOutputFile.txt
/gate/actor/MyActor/attachTo  MyVolume
```

**Stop on script**

This actor gets the output of a script and stop the simulation if this output is true. The script is called with the command:

```
/gate/actor/[Actor Name]/save  MyScript
```

It is possible to save all the others actor before stopping the simulation with the command:

```
/gate/actor/[Actor Name]/saveAllActors  true
```

Example:

```
/gate/actor/addActor StopOnScriptActor MyActor
/gate/actor/MyActor/save  MyScript
/gate/actor/MyActor/saveAllActors  true
```

**Track length**

This actor stores the length of each tracks in a root file. It takes into account the weight of particles. They are three commands to define the boundaries and the binning of the histogram.
Energy spectrum

This actor builds four histograms: the initial kinetic energy of each track (energySpectrum), the energy deposition per event (edepHisto), the energy deposition per track (edepTrackHisto), the energy loss per track (eLossHisto). These histograms are stored in a root file. They take into account the weight of particles. They are three commands to define the boundaries and the binning of the energy spectrum and three commands to define the boundaries and the binning of the energy loss histograms (edepHisto, edepTrackHisto, eLossHisto).

Production and stopping particle position

This actor stores in a 3D image the position where particles are produced and where particles are stopped. For the output, the suffixes Prod and Stop are added to the output file name given by the user. You can use several file types: ASCII file (.txt), root file (.root), (.hdr/.img). The root file works only for 1D and 2D distribution.

Secondary production

This actor creates a root file and stores the number of secondaries in function of the particle type. Ionisation electrons are dissociated from electrons produced by other processes. Decay positrons are dissociated from positrons produced by other processes. Gammas are classified in four categories: gammas produced by EM processes, gammas produced by hadronic processes, gammas produced by decay processes and other gammas.

Delta kinetic energy

This actor sums the relative and absolute $\Delta$ (kinetic energy) and stores the results in two files (with suffixes "-RelStopPower" and "-StopPower"). It also stores the map of the hits to allow users to calculate the mean values.

Number of particles entering volume
This actor builds a map of the number of particles produced outside of the actor volume and interacting in the volume. The particle is recorded once in each voxel where it interacting.

```yaml
/gate/actor/addActor ParticleInVolumeActor MyActor
/gate/actor/MyActor/save MyOutputFile.hdr
/gate/actor/MyActor/attachTo MyVolume
/gate/actor/MyActor/setResolution 10 10 100
/gate/actor/MyActor/setHitType post
```

**Q-value**

This actor calculate the Q-values of interactions.

```yaml
/gate/actor/addActor QvalueActor MyActor
/gate/actor/MyActor/save MyOutputFile.hdr
/gate/actor/MyActor/attachTo MyVolume
/gate/actor/MyActor/setResolution 10 10 100
/gate/actor/MyActor/setHitType random
```

**CrossSectionProductionActor**

The CrossSectionProductionActor derives the production of C-11, or O-15 from the equation proposed by (Parodi et al, 2007). The cross section data are provided by directly in the class code. By default, only the production of the C-11 is activated.

**WARNING**: The size of the image has to be given in mm

The current limit in cross section data is 199 MeV, other data can be added in the class.

```yaml
/gate/actor/addActor CrossSectionProductionActor beta
/gate/actor/beta/attachTo volume
/gate/actor/beta/save output_dump/test_small.hdr
/gate/actor/beta/addO15 true
/gate/actor/beta/addC11 true
/gate/actor/beta/setVoxelSize 1 1 1 mm
/gate/actor/beta/saveEveryNEvents 100000
```

**Biological Washout: WashOutActor**

The biological washout processes were incorporated into the GATE code. The Mizuno model (H. Mizuno et al. Phys. Med. Biol. 48, 2003), is implemented. For this purpose, a new actor, which allows the management of particles during the tracking, was developed. This actor is named WashoutActor and it is attached to a certain volume, in which the washout processes are applied. In particular, the activity distributions of the associated volume are continuously modified as a function of the acquisition time in terms of the following equation:

\[
C_{washout}(t) = M_f \exp\left(-\frac{t}{T_f \ln 2}\right) + M_m \exp\left(-\frac{t}{T_m \ln 2}\right) + M_s \exp\left(-\frac{t}{T_s \ln 2}\right)
\]

Where 3 components are defined (fast, medium and slow) with two parameters for each: the half life \(T\) and the fraction \(M\) (\(M_f + M_m + M_s = 1\)).

Users should provide a table as an ASCII file with the washout parameters values for any radioactive source in the associated volume. In order to take into account the physiological properties of each tissue, it is important to highlight that one independent radioactive source should be defined per each material involved in the simulation.

Example: How to define a washout actor module

```yaml
/gate/actor/addActor WashOutActor [ACTOR NAME]
/gate/actor/[ACTOR NAME]/attachTo [VOLUME NAME]
/gate/actor/[ACTOR NAME]/readTable [TABLE FILE NAME]
```
Example of [TABLE FILE NAME]: How to specify different parameters which are associated to the washout model - This ASCII file will be used by the washout Actor.

<table>
<thead>
<tr>
<th>SOURCE 1 NAME</th>
<th>MATERIAL 1 NAME</th>
<th>Mf VALUE</th>
<th>Tf VALUE IN SEC</th>
<th>Mm VALUE</th>
<th>Tm VALUE IN SEC</th>
<th>Ms VALUE</th>
<th>Ts VALUE IN SEC</th>
</tr>
</thead>
<tbody>
<tr>
<td>SOURCE 2 NAME</td>
<td>MATERIAL 2 NAME</td>
<td>Mf VALUE</td>
<td>Tf VALUE IN SEC</td>
<td>Mm VALUE</td>
<td>Tm VALUE IN SEC</td>
<td>Ms VALUE</td>
<td>Ts VALUE IN SEC</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Filters

Filters are used to add selection criteria on actors. They are also used with reduction variance techniques. They are filters on particle type, particle ID, energy, direction....

Filter on particle type

With this filter it is possible to select particle with the name [Particle Name]:

```
/gate/actor/[Actor Name]/addFilter particleFilter
/gate/actor/[Actor Name]/particleFilter/addParticle [Particle Name]
```

User can select various particles. It is also possible to select particles which has a parent with the name [Particle Name]:

```
/gate/actor/[Actor Name]/addFilter particleFilter
/gate/actor/[Actor Name]/particleFilter/addParentParticle [Particle Name]
```

For ions, user should use the Geant4 nomenclature (C12[0.0], c11[0.0]...). These names are different from those used for physics.

To select all ions except alpha, deuton and triton, there is the key word 'GenericIon'.

Example: To kill electrons and positrons in the volume MyVolume:

```
/gate/actor/addActor KillActor MyActor
/gate/actor/MyActor/save MyOutputFile.txt
/gate/actor/MyActor/attachTo MyVolume
/gate/actor/MyActor/addFilter particleFilter
/gate/actor/MyActor/particleFilter/addParticle e-
/gate/actor/MyActor/particleFilter/addParticle e+
```

Filter on particle ID

In an event, each track has an unique ID. IDs are reset at each event. The incident particle has an ID equal to 1. This filter select particles with the ID [Particle ID] or particles which has a parent with the ID [Particle ID]. As for particle filter, user can select many IDs.

```
/gate/actor/[Actor Name]/addFilter IDFilter
/gate/actor/[Actor Name]/IDFilter/selectID [Particle ID]
```

```
/gate/actor/[Actor Name]/addFilter IDFilter
/gate/actor/[Actor Name]/IDFilter/selectParentID [Particle ID]
```

Example: To kill all particle except the incident particle in the volume MyVolume (all particles are the children of the incident particle except the incident particle itself):

```
/gate/actor/addActor KillActor MyActor
/gate/actor/MyActor/save MyOutputFile.txt
/gate/actor/MyActor/attachTo MyVolume
/gate/actor/MyActor/addFilter IDFilter
/gate/actor/MyActor/IDFilter/selectParentID 1
```
Filter on volume

This actor is especially useful for reduction variance techniques or for selections on daughter volumes.

Example: To kill particles in volume A and B, children of the volume MyVolume:

```plaintext
/gate/actor/addActor KillActor MyActor
/gate/actor/MyActor/save MyOutputFile.txt
/gate/actor/MyActor/attachTo MyVolume
/gate/actor/MyActor/addFilter volumeFilter
/gate/actor/MyActor/volumeFilter/addVolume A
/gate/actor/MyActor/volumeFilter/addVolume B
```

Filter on energy

This filter allows to select particles with a kinetic energy above a threshold Emin and/or below a threshold Emax.

Example: To kill particles with an energy above 5 MeV:

```plaintext
/gate/actor/[Actor Name]/addFilter energyFilter
/gate/actor/[Actor Name]/energyFilter/setEmin [Value] [Unit]
/gate/actor/[Actor Name]/energyFilter/setEmax [Value] [Unit]
```

Filter on direction

This filter is used to select particle with direction inside a cone centered on the reference axis. The angle between the axis and the edge of the cone is in degree. The axis is defined with the (x,y,z) directions.

Example: To kill particles in a cone of 20 degrees around x axis:

```plaintext
/gate/actor/[Actor Name]/addFilter angleFilter
/gate/actor/[Actor Name]/angleFilter/setAngle [Value]
/gate/actor/[Actor Name]/angleFilter/setDirection [x] [y] [z]
```

Retrieved from


- This page was last modified on 14 April 2014, at 14:49.
Users Guide V7.0: Phase space concept

From GATE collaborative documentation wiki

Table of Contents

- How to create a phase space?
- How to use a phase space?

How to create a phase space?

This actor records information about particles entering the volume which the actor is attached to. They are two file types for the output: root file (.root) and IAEA file (.IAEAphsp and .IAEAnheader). The name of the particle, the kinetic energy, the position along the three axes, the direction along the three axes, the weight are recorded. In a IAEA file, each particle is designated by an integer while the full name of the particle is recorded in the root file. Particles in IAEA files are limited to photons, electrons, positrons, neutrons and protons. The root file has two additional pieces of information: the name of the volume where the particle was produced and the name of the process which produced the particle. It is possible to disable some information in the phase space file:

```
/gate/actor/source/enableEkine              false
/gate/actor/source/enableXPosition          false
/gate/actor/source/enableYPosition          false
/gate/actor/source/enableZPosition          false
/gate/actor/source/enableXDirection         false
/gate/actor/source/enableYDirection         false
/gate/actor/source/enableZDirection         false
/gate/actor/source/enableProductionVolume   false
/gate/actor/source/enableProductionProcess  false
/gate/actor/source/enableParticleName       false
/gate/actor/source/enableWeight             false
```

By default the frame used for the position and the direction of the particle is the frame of the world. To use the frame of the volume which the actor is attached to, the following command should be used:

```
/gate/actor/source/useVolumeFrame
```

By default, the phase space stores particles entering the volume. To store particles exiting the volume, the following command should be used:

```
/gate/actor/MyActor/storeOutgoingParticles
```
To store all secondary particles created in the volume, use the command:

```
/gate/actor/MyActor/storeSecondaries true
```

Phase spaces built with all secondaries should not be used as source because some particles could be generated several times.

**With ROOT files**, to avoid very big files, it is possible to restrict the maximum size of the phase space. If a phase space reaches the maximum size, the files is closed and a new file is created. The new file has the same name and a suffix is added. The suffix is the number of the file. For instance, instead of one file of 10 GB, user may prefer 10 files of 1 GB. The value of the maximum size is not exactly the size of the file (value is the size of the TTree).

```
/gate/actor/MyActor/setMaxFileSize [Value] [Unit (B, kB, MB, GB)]
```

### How to use a phase space?

The source of the simulation could be a phase space. Gate read two types of phase space: root files and IAEA phase spaces. Both can be created with Gate. However, Gate could read IAEA phase spaces created with others simulations.

```
/gate/source/addSource [Source name] phaseSpace
```

User can add several phase space files. All files should have the same informations about particles. The files are chained:

```
/gate/source/[Source name]/addPhaseSpaceFile [File name 1]
/gate/source/[Source name]/addPhaseSpaceFile [File name 2]
```

If particles in the phase space are defined in the world frame, user has to used the command:

```
/gate/source/[Source name]/setPhaseSpaceInWorldFrame
```

If the particle type is not defined in the phase space file, user have to give the particle name. It is supposed that all particles have the same name:

```
/gate/source/[Source name]/setParticleType [Particle name]
```

If user have several phase space sources, each source have the same intensity. User can also choose to give at each source an intensity proportionnal to the number of particles in the files attach to the source:

```
/gate/source/[Source name]/useNbOfParticleAsIntensity true
```
For each run, if the number of events is higher than the number of particles in file, each particle is used several times with the same initial conditions. However, it is possible to rotate the particle position and direction around the z axis of the volume (make sure your phase space files have a rotational symmetry). The regular rotation is a rotation with a fixed angle:

\[ \alpha = \frac{2\pi}{N_{used}} \]

where \( N_{used} \) is the number of time the particle is used.

The random rotation is a rotation with a random angle.

By default, all particles in a phase space are used. The particles in the the phase space can be preselected in function of their position in the \((x,y)\) plan. For instance, a particle with a origin far from the collimator aperture is not useful and should be ignored. Particles in a \( r \) cm-radius circle are selected. The particles outside the circle are ignored.

Examples are available here (http://wiki.opengatecollaboration.org/index.php/GateRT)


- This page was last modified on 4 September 2013, at 18:23.
GateRT

From GATE collaborative documentation wiki

GATE for Radiation Therapy applications

GATE was initially developed for PET and SPECT applications. From version 6.0, some specific tools have been added for radiation therapy (RT) applications. A set of examples illustrates the use of Gate for various photon, proton and carbon ion applications.

Disclaimer: the tools dedicated to radiation therapy simulations provided in this GATE release are provided "as is" and on an "as available" basis without any representation or endorsement made and without warranty of any kind. In clear, it is a first attempt in this field and the authors are pretty sure that there are bugs. We will be happy if you can report bugs and hence improve this collaborative software.

The examples are located in the source code into to the folder: examples/example_Radiotherapy/. There are 10 examples:

- Example 1: Example of Carbon beam in water tank or in patient CT image. Output is a 3D dose distribution map (with associated statistical uncertainty) and map of produced C11. (This example is similar to Novice_1 in Gate V6.1)

- Example 2: Example of photon beam in patient CT image. Output is a 3D dose distribution map (with associated uncertainty). Two different navigators are tested NestedParameterized and Regionalized, with two number of materials. (This example is similar to the example1 presented on the wiki web site of Gate V6.1)

- Example 3: Example of photon beam in patient CT image with IMRT irradiation. 100 slices with different MLC positions. (This example is similar to the example2 presented on the wiki web site of Gate V6.1)

- Example 4: Example of proton beam in water box with Pencil Beam Scanning source and Phase Space output. You can use root -x PhS-Analysis.C to analyse the produced phase spaces. (This example is similar to Novice_4 in Gate V6.1)

- Example 5: Example of proton beam in water box with Pencil Beam Scanning source with a treatment plan obtained from a TPS. (This example is similar to Novice_5 in Gate V6.1)

- Example 6: Example of proton beam in heterogeneous phantom (water, bones, Lung) with Pencil Beam Scanning source: comparison between dose to water and dose to medium (conventional one).

- Example 7: Example to use repeater/mover and both at the same time. (This example is similar to the example5 presented on the wiki web site of Gate V6.1)
Example 8: Photon beam from a Linac into a box with water/alu/lung. See Figure 4 from [Jan et al PMB 2011].

Example 9: Electron beam from a Linac into a box with water/alu/lung. See Figure 5 from [Jan et al PMB 2011].

Example 10: Radiography of a thorax phantom. Outputs are 3D dose distribution maps computed with the classical method and the accelerated (TLE) method.

Older examples are still available here: [1](http://www.creatis.insa-lyon.fr/gate) but are no more maintained.

Retrieved from "http://wiki.opengatecollaboration.org/index.php/GateRT"

- This page was last modified on 13 July 2012, at 08:05.
- [2 watching users]
Users Guide V7.0: How to use Gate on a Cluster

From GATE collaborative documentation wiki

Table of Contents

- Installation of the job splitter (gjs)
- Installation of the file merger (gjm)
- Preparing your macro
- Using the job splitter
- Using the file merger
- Alternative to the file merger
- What about errors?

To reduce the overall computing time of GATE experiments, a parallel computing platform for running simulations in a cluster of computers was developed which significantly shortens the setup time and provides fast data output handling. To use Gate in cluster mode you need 3 components:

- The job splitter (gjs)
- The file merger (gjm)
- A cluster aware version of Gate

Installation of the job splitter (gjs)

The job splitter can be installed in the same directory as Gate. Two environment variables are already added to the environment file used to compile Gate (but you can customize them):

```bash
export GC_DOT_GATE_DIR=/somedir/
export GC_GATE_EXE_DIR=/somedir/bin/Linux-g++/
```

The first variable indicates the location of a hidden directory called .Gate. The directory will contain the split macros for each part of the simulation. Even when splitting the same macro several times, a new directory will be created for each instance (with an incremental number). In normal circumstances, one does not need to look into it. In case of an error, it can be used to run only a specific part of a simulation again (See #What about errors?). The second environment variable indicates the location of the job splitter executable. As the Gate environment file will be used to compile the job splitter source code, the executable will likely be located in the same directory as the Gate executable.

To install, load the Gate/Geant4 environment variables, go to the job splitter directory (bash example):

```bash
source env_gate.sh
cd jobsplitter
make
```
By default, the executable will be created in the jobsplitter directory. If the GATEHOME variable is correctly defined, the executable will also be copied in the same directory as the Gate executable (same for the dynamic library).

**Installation of the file merger (gjm)**

To install, it is the same way, go to the file merger directory and compile (bash example):

```bash
cd filemerger
make
```

The file merger executable is located in the current directory (and will also be copied in the Gate bin directory as for the gjs program).

**Preparing your macro**

The cluster software should be able to handle all GATE macros. However, only ROOT is currently supported as an output format for the gjm program. So be aware that other output formats cannot yet be merged with the gjm program and you will have to do this on your own (but it is usually quite simple ~ addition or mean most of the time).

If an isotope with a shorter half life than the acquisition time is simulated, then it may be useful to specify the half life in your macro as follows:

```
/gate/cluster/setTimeSplitHalflife 6600. s
```

This way, the CPU time will be approximately equal for each job.

In planning simulation time, it is important to be aware that Gate simulations currently seem to benefit only from the addition of physical CPUs. A computer with 8 hyper-threaded physical CPU cores (16 logical CPUs) will have the same computational efficiency if 8 processes are run simultaneously as it would with 16 simultaneous processes.

**Using the job splitter**

To view information regarding general usage, you can run the job splitter executable without any options:

```
+-------------------------------------------+
| gjs -- The GATE cluster job macro spliter |
+-------------------------------------------+
Usage: gjs [-options] your_file.mac

Options (in any order):
-a value alias : use any alias
-numberofsplits, -n n : the number of job splits; default=1
-clusterplatform, -c name : the cluster platform, name is one of the following:
  openmosix - condor - openPBS - xgrid
This executable is compiled with condor as default
```

-openPBSscript, os : template for an openPBS script  

see the example that comes with the source code (script/openPBS.script)  
overrules the environment variable below

-condorscript, cs : template for a condor submit file  

see the example that comes with the source code (script/condor.script)

-v : verbosity 0 1 2 3 - 1 default

Environment variables:

GC_DOT_GATE_DIR : indicates the .Gate directory for splitted mac files

GC_GATE_EXE_DIR : indicates the directory with the Gate executable

GC_PBS_SCRIPT : the openPBS template script (!optionnal variable!)

Usage (bash):

export GC_DOT_GATE_DIR=/home/user/gatedir/
export GC_GATE_EXE_DIR=/home/user/gatedir/bin/Linux-g++/

Examples:

gjs -numberofsplits 10 -clusterplatform openmosix macro.mac

gjs -numberofsplits 10 -clusterplatform openmosix -a /somedir/rootfilename ROOT_FILE macro.mac

gjs -numberofsplits 10 -clusterplatform openPBS -openPBSscript /somedir/script macro.mac

gjs -numberofsplits 10 -clusterplatform xgrid macro.mac

gjs -numberofsplits 10 /somedir/script macro.mac

The supported platforms are currently: openMosix, openPBS, Condor and Xgrid.

Let's take openMosix as an example:

```
gjs -numberofsplits 5 -clusterplatform openmosix macro.mac
```

The job splitter will subdivide the simulation macro into fully resolved, non-parameterized macros. In this case there are 5 such macros. They are located in the .Gate directory, as specified by the GC_DOT_GATE_DIR environment variable.

A list of all the data output options is given after successful completion, as well as a list of all activated actors. The user is asked to clearly enable each needed output module and to give them an output file name. It is the same for actors. Remember that by default, no output module nor actor is enabled.

If an alias was expected for output files and it was not supplied, then this will be mentioned in the output options list. A standard name will be supplied automatically, as well as appropriate numbering.

The time of each sub-macro is manage using a virtual timeStart and a virtual timeStop calculated by the gjs and used by the command /gate/application/startDAQCluster. All defined runs and geometry updates will be totally respected. The only inconsistency in the use of gjs is when using the projection output: the virtualStop minus virtualStart time have to be a multiple of timeSlice, otherwise the GateToProjectionSet output will lead to an error.

The .Gate directory will have a subdirectory called as the macro name, that contains the following files:

```
macro1.mac
macro2.mac
macro3.mac
macro4.mac
macro5.mac
macro.split
```
The 5 macros are listed as well as well as the .split file that contains information about the splitted simulation and that will be used to merge the data after the simulation (using the gjm program). The current directory, from which the jobsplitter was called, now contains the cluster submit file. In order to run the split simulation on the cluster, one only needs to execute or call this file with a certain program (depending on the cluster platform used).

The .Gate directory supports automatic numbering. If the same macro is used repeatedly, then the subsequent directories will be numbered using an incremental number.

### Using the file merger

The file merger have to be run giving the split file as input. To view information on general usage, just run the file merger executable without any options:

```bash
+-------------------------------------------+
| gjm -- The GATE cluster job output merger |
+-------------------------------------------+

Usage: gjm [-options] your_file.split

You may give the name of the split file created by gjs (see inside the .Gate directory). !! This merger is only designed to ROOT output. !!

Options:
-outdir path : where to save the output files default is PWD
-v : verbosity 0 1 2 3 - 1 default
-f : forced output - an existing output file will be overwritten
-cleanonly : do only a the cleanup step i.e. no merging
-cleanonlyTest : just tells you what will be erased by the -cleanonly
-clean : merge and then do the cleanup automatically
-fastMerge : correct the output in each file, to be used with a TChain (only for Root output)

Environment variable:
GC_DOT_GATE_DIR : points to the .Gate directory

To merge the output files into a single file, just supply the split file to the file merger. The output file could be used as a usual single CPU output file.

```bash
gjm macro.split

Combining: ./rootf1.root ./rootf2.root ./rootf3.root ./rootf4.root ./rootf5.root $->$ ./rootf.root
```

In case a single output file is not required, it is possible to use the option fastMerge. This way, the eventIDs in the output files are corrected locally. Figure 13.1 shows the newly created tree in each ROOT file.
Figure 13.1: Example of ROOT file with added cluster eventIDs

A ROOT chain, which is a list of files containing the same tree, is then required to link the output files together for analysis. A chain for the Singles could be made as follows (in a file called chain.c):

```c
{
  gROOT->Reset();
  TChain chain("Singles");
  chain.Add("rootf1.root");
  chain.Add("rootf2.root");
  chain.Add("rootf3.root");
  chain.Add("rootf4.root");
  chain.Add("rootf5.root");
}
```

Once all files are added to the chain, one can use the chain as a regular Ttree, and the normal ROOT prompt is returned:

```bash
$root chain.c
```
FreeType Engine v2.1.3 used to render TrueType fonts.
Compiled for linux with thread support.
CINT/ROOT C/C++ Interpreter version 5.15.94, June 30 2003
Type ? for help. Commands must be C++ statements.
Enclose multiple statements between \{ \}.
root [0]
> Processing chain.c...
root [1]
root [1] Singles->Draw("energy")

Alternative to the file merger

Root files can also be merged by using the hadd utility on the command line:

hadd result.root file1.root file2.root ... filen.root

What about errors?

If something went wrong during a simulation and a ROOT file is corrupted or incomplete, then this will be detected by the file merger. There are two options. First, one can restart only the specific part of the simulation that went wrong. This can be easily done, as the ROOT files are numbered and one can edit the submit file so it only launches that specific part. Alternatively, one can find the macro file that was used to start that part of the simulation in the .Gate directory and start the simulation directly with the macro file and its corresponding seed file. The second option is to edit the split file, located in the .Gate directory. Once the reference to the corrupted root file is removed from it, it is possible to merge the files again. At this point, the eventIDs will not be valid anymore.

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Users Guide V7.0: How to use Gate on a GPU

From GATE collaborative documentation wiki

Table of Contents

- Installation of CUDA tools
- PET applications
- CT applications

Installation of CUDA tools

IMPORTANT NOTE:

!! GPU modules can be used ONLY with NVIDIA hardware and CUDA tools !!

For details, see the link on the wiki installation guide page:
http://wiki.opengatecollaboration.org/index.php/NewCompilationProcedureV7.0#GPU_.26_CUDA_tools

PET applications

For PET applications, the GPU manages the particle tracking within the voxelized phantom and this is why the source type is defined as a [GPUEmisTomo]. Examples are provided within the GATE source package and users can find a complete example about How To define and run a complete PET simulation set-up by using a GPU architecture.

IMPORTANT NOTES:

1/ By using the GPU, phantom scatter informations are NOT available in the output files

2/ The sources radioactive decay time is NOT available with the GPU

```
#==================================================
# VOXELIZED SOURCES
#==================================================
/gate/source/addSource srcvoxel GPUEmisTomo
/gate/source/srcvoxel/attachPhantomTo ncat
/gate/source/srcvoxel/setGPUBufferSize 1000000
/gate/source/srcvoxel/setGPUDeviceID 1
# Read the phantom as usual
/gate/source/srcvoxel/reader/insert interfile
/gate/source/srcvoxel/reader/translator/insert range
/gate/source/srcvoxel/reader/translator/readTable data/activities.dat
/gate/source/srcvoxel/reader/translator/describe 1
/gate/source/srcvoxel/reader/translator/verbose 0
/gate/source/srcvoxel/reader/translator/readFile data/thorax_phantom.hdr
/gate/source/srcvoxel/setType backtoback
/gate/source/srcvoxel/gps/particle gamma
/gate/source/srcvoxel/gps/energytype Mono
/gate/source/srcvoxel/gps/monoenergy 0.511 MeV
/gate/source/srcvoxel/setPosition 0 0 0 cm
```

CT applications

For CT applications, the GPU tracking is defined as an Actor.

Examples are provided within the GATE source package and users can find a complete example about How To define and run a complete CT simulation set-up by using a GPU architecture.

IMPORTANT NOTE:

The scannerCT system, the digitizer and the CT dedicated output are NOT available with the GPU CT module. The CT detector must be define by using an actor as it is shown in the complete example provided with the source.