

# **BDSIM User's Manual v0.5**

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# BDSIM v0.5 User's Manual

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## 1 About BDSIM

BDSIM is a Geant4 ([1], page 34) extension toolkit for simulation of particle transport in accelerator beamlines. It provides a collection of classes representing typical accelerator components, a collection of physics processes for fast tracking, procedures of “on the fly” geometry construction and interfacing to ROOT analysis ([2], page 34).

## 2 Obtaining, Installing and Running

BDSIM can be downloaded from <http://ilc.pp.rhul.ac.uk/bdsim.html>. This site also contains some information on planned releases and other issues. Alternatively, a development version is accessible under <http://cvs.pp.rhul.ac.uk>. Download the tarball and extract the source code. Make sure Geant4 is installed and appropriate environment variables defined. Then go through the configuration procedure by running the `./configure` script.

```
./configure
```

It will create a Makefile from template defined in Makefile.in. You may want to edit the Makefile manually to meet your needs (if your CLHEP version is greater than 2.x put `-DCLHEP_VERSION=9`). Then start the compilation by typing

```
make
```

If the compilation is successful the `bdsim` executable should be created in `$(BDSIM)/bin/$(ARCH)` where `$(BDSIM)` is the directory specified during configuration, and `$(ARCH)` is of the form `$(OSTYPE)-$(COMPILER)`, eg `Linux-g++`. Next, set up the `(DY)LD_LIBRARY_PATH` variable to point to the `./parser` directory, and also to the directory where `libbdsim.so` is if building shared libraries.

BDSIM is invoked by the command `bdsim 'options'`

where the options are

```
--file=<filename>      : specify the lattice file
--output=<fmt>          : output format (root|ascii), default ascii
--outfile=<file>        : output file name. Will be appended with _N
                        : where N = 0, 1, 2, 3... etc.
--vis_mac=<file>        : visualization macro script, default vis.mac
--help                 : display this message
--verbose              : display general parameters before run
--verbose_event         : display information for every event
--verbose_step=N        : display tracking information after each step
--verbose_event_num     : display tracking information for event number N
--batch                : batch mode - no graphics
--outline=<file>        : print geometry/optics info to <file>
--outline_type=<fmt>    : type of outline format
```

```

                                where fmt = optics | survey
--materials                    : list materials included in bdsim by default

```

To run `bdsim` one first has to define the beamline geometry in a file which is then passed to `bdsim` via the `--file` command line option, for example

```
bdsim --file=line.gmad --output=root --batch
```

The next section describes how to do it in more detail.

### 3 Lattice description

The beamline, beam properties and physics processes are specified in the input file written in the GMAD language which is a variation of MAD-X language ([3], page 34) extended to handle sophisticated geometry and parameters relevant to radiation transport. GMAD is described in this section. Examples of input files can be found in the `BDSIM` distribution in the `examples` directory. In order to convert a MAD file into a GMAD one, a utility called `mad2gmad.sh` is provided in the `utils` directory.

The following MAD commands are not supported:

- `assign`
- `bmpm`
- `btrns`
- `envelope`
- `optics`<sup>1</sup>
- `title`
- `option`
- `plot`
- `print`
- `return`
- `survey`<sup>2</sup>
- `title`

The following MAD commands:

- `moni`
- `monitor`
- `wire`
- `prof`

are replaced with the `marker` command.

---

<sup>1</sup> To dump the optical properties of the lattice one can invoke `bdsim` with the `--outline=file.txt --outline_type=optics` options.

<sup>2</sup> To compute the coordinates of all machine elements in a global reference system one can invoke `bdsim` with the `--outline=file.txt --outline_type=survey` options

### 3.1 Program structure

A GMAD program consists of a sequence of element definitions and control commands. For example, tracking a 1 GeV electron beam through a FODO cell will require a file like this:

```
mk: marker;
qf: quadrupole, l=0.5*m, k1=0.1*m^-2;
qd: quadrupole, l=0.5*m, k1=-0.1*m^-2;
d: drift, l=0.5*m;
fodo : line=(qf,d,qd,d,mk);
use, period=fodo;
beam, particle="e-",energy=1*GeV;
option, beampipeRadius=5*cm, beampipeThickness=5*mm;
sample, range=mk;
```

Generally, the user has to define a sequence of elements (with **drift**, **quadrupole**, **line** etc.), then select the beamline with the **use** command and specify beam parameters and other options with **beam** and **option** commands. The **sample** and **csample** commands control what sort of information will be recorded during the execution.

The parser is case sensitive. However, for convenience of porting lattice descriptions from MAD the keywords can be both lower and upper case. The GMAD language is discussed in more detail in this section.

### 3.2 Arithmetical expressions

Throughout the program a standard set of arithmetical expressions is available. Every expression is ended with a semicolon, for example:

```
x=1;
y=2.5-x;
z=sin(x) + log(y) - 8e5;
```

Available binary operators are: +, -, \*, /, ^

Available unary operators are: +, -

Available boolean operators are: <, >, <=, >=, <>, ==

Available functions<sup>3</sup> are:

- sqrt
- cos
- sin
- exp
- log
- tan
- asin

---

<sup>3</sup> see `add_func(..)` in `parser/gmad.cc`

- `acos`
- `atan`
- `abs`

### 3.3 Physical elements and Entities

GMAD implements almost all the standard MAD elements, but also allows to define arbitrary geometric entities and magnetic field configurations. The geometry description capabilities are extended by using “drivers” to other geometry description formats, which makes interfacing and standardisation easier. The syntax of a physical element declaration is

```
element_name : element_type, attributes;
```

for example

```
qd : quadrupole, l = 0.1*m, k1 = 0.01;
```

`element_type` can be of basic type or inherited. Allowed basic types are

- `marker`
- `drift`
- `rbend`
- `sbend`
- `quadrupole`
- `sextupole`
- `octupole`
- `multipole`
- `vkick`
- `hkick`
- `rf`
- `rcol`
- `ecol`
- `solenoid`
- `laser`
- `transform3d`
- `element`

All elements except `marker`, `element`, `ecol`, and `rcol` are by default modeled with an inner cylindrical beampipe and an outer cylindrical volume. (FOR MAD COMPATIBILITY `sbend` SHOULD BE A TORUS). The beampipe outer radius and thickness are defined by the global `beampipeRadius` and `beampipeThickness` options; the beampipe outer radius can be redefined for almost every element with the `aper` option. The beampipe material is defined by the global `beampipeMaterial` option (default: “Vacuum”), while the residual gas in the beampipe at the moment cannot be changed by the user and is set to “Vacuum”. The outer volume is represented (with the exception of the `drift` element) by a cylinder with inner radius equal to the beampipe outer radius and with outer radius given by default by the global `boxSize` option, which can usually be overridden with the “outR” option.

An already defined element can be used as a new element type. The child element will have the attributes of the parent.

```
q:quadrupole, l=1*m, k1=0.1;
qq:q,k1=0.2;
```

### 3.3.1 Coordinate system

The usual accelerator coordinate system is assumed (see [3], page 34).

### 3.3.2 Units

In GMAD the SI units are used.

length	[m] (metres)
time	[s] (seconds)
angle	[rad] (radians)
quadrupole coefficient	[m <sup>-2</sup> ]
multipole coefficient 2n poles	[m <sup>-n</sup> ]
electric voltage	[MV] (Megavolts)
electric field strength	[MV/m]
particle energy	[GeV]
particle mass	[GeV/c <sup>2</sup> ]
particle momentum	[GeV/c]
beam current	[A] (Amperes)
particle charge	[e] (elementary charges)
emittances	[pi m mrad]
density	[g/cm <sup>3</sup> ]
temperature	[K] (Kelvin)
pressure	[atm] (atmosphere)
mass number	[g/mol]

There are some predefined numerical values<sup>4</sup> are:

pi	3.14159265358979		
GeV	1	m	1
eV	10 <sup>-9</sup>	cm	10 <sup>-2</sup>
KeV	10 <sup>-6</sup>	mm	10 <sup>-3</sup>
MeV	10 <sup>-3</sup>	um	10 <sup>(-6)</sup>
TeV	10 <sup>3</sup>	nm	10 <sup>(-9)</sup>
MV	1	s	1
Tesla	1	ms	10 <sup>-3</sup>
rad	1	us	10 <sup>-6</sup>
mrاد	10 <sup>-3</sup>	ns	10 <sup>-9</sup>
clight	2.99792458 * 10 <sup>8</sup>		

for example, one can write either 100\*eV or 0.1\*KeV when energy constants are concerned.

---

<sup>4</sup> see add\_var(..) in `parser/gmad.cc`



### 3.3.3 marker

**marker** has no effect (no volume is associated to it) but allows one to identify a position in the beam line (say, where a sampler will be placed). It has no attributes.

Example:

```
m1 : marker;
```

### 3.3.4 drift

**drift** defines a straight drift space. Its volume contains only the vacuum beampipe (no outer iron cylinder).

Attributes:

- **l** - length [m] (default 0)
- **aper** - aperture [m] (default same as **beampipeRadius**)

Example:

```
d13 : drift, l=0.5*m;
```

### 3.3.5 rbend

**rbend** defines a rectangular bending magnet. Attributes:

- **l** - length [m] (default 0)
- **angle** - bending angle [rad] (default 0)
- **B** - magnetic field [T]
- **aper** - aperture [m] (default same as beampipe radius)
- **outR** - external radius [m] of magnet (default set to **aper**+22cm)
- **material** - the magnet material (default set to “Iron”)
- THE CODE ALSO ALLOWS FOR A QUADRUPOLE FIELD GRADIENT K1..

when **B** is set, this defines a magnet with appropriate field strength and **angle** is not taken into account. Otherwise, the value of **B** that corresponds to bending angle **angle** for a particle in use (defined by the **beam** command, with appropriate energy and rest mass) is calculated and used in the simulations.

Example :

```
rb1 : rbend, l=0.5*m, angle = 0.01;
```

### 3.3.6 sbend

**sbend** defines a sector bending magnet. Attributes:

- **l** - length [m] (default 0)
- **angle** - bending angle [rad] (default 0)

- B - magnetic field [T]
- **aper** - aperture [m] (default same as beampipe radius)
- **outR** - external radius [m] of magnet (default set to **aper+22cm**)
- **material** - the magnet material (default set to “Iron”)
- THE CODE ALSO ALLOWS FOR A QUADRUPOLE GRADIENT K1..

The meaning of B and **angle** is the same as for **rbend**.

Example :

```
sb1 : sbend, l=0.5*m, angle = 0.01;
```

### 3.3.7 quadrupole

**quadrupole** defines a quadrupole. Attributes:

- **l** - length [m] (default 0)
- **k1** - normal quadrupole coefficient  $k1 = 1/(B\rho) dB_y/dx$  [ $m^{-2}$ ] Positive **k1** means horizontal focusing of positively charged particles (default 0).  $dB_y/dx$  is the magnetic field gradient, while  $(B\rho)$  is the magnetic “rigidity”:  $B\rho$  (T\*m) =  $p(\text{GeV})/(0.299792458 * |charge(e)|)$
- **ks1** - skew quadrupole coefficient  $ks1 = 1/(B\rho) dB_y/dx$  [ $m^{-2}$ ] where (x,y) is now a co-ordinate system rotated by 45 degrees around s with respect to the normal one.(default 0).
- **tilt** - roll angle [rad] about the longitudinal axis, clockwise.
- **aper** - aperture [m] (default same as beampipe radius)
- **outR** - external radius [m] of magnet (default set to **aper+22cm**)
- **material** - the magnet material (default set to “Iron”)

Example :

```
qf : quadrupole, l=0.5*m , k1 = 0.5 , tilt = 0.01;
```

### 3.3.8 sextupole

**sextupole** defines a sextupole. Attributes:

- **l** - length [m] (default 0)
- **k2** - normal sextupole coefficient  $k2 = 1/(B\rho) d^2B_y/dx^2$  [ $m^{-3}$ ]
- **ks2** - skew sextupole coefficient  $ks2 = 1/(B\rho) d^2B_y/dx^2$  [ $m^{-3}$ ] where (x,y) is now a co-ordinate system rotated by 30 degrees around s with respect to the normal one.(default 0).
- **tilt** - roll angle [rad] about the longitudinal axis, clockwise.
- **aper** - aperture [m] (default same as beampipe radius)
- **outR** - external radius [m] of magnet (default set to **aper+22cm**)
- **material** - the magnet material (default set to “Iron”)

Example :

```
sf : sextupole, l=0.5*m , k2 = 0.5 , tilt = 0.01;
```

### 3.3.9 octupole

**octupole** defines an octupole. Attributes:

- **l** - length [m] (default 0)
- **k3** - normal octupole coefficient  $k3 = 1/(B\rho) d^3 B_y/dx^3$  [m<sup>-4</sup>] Positive **k3** means horizontal focusing of positively charged particles. (default 0)
- **ks3** - skew octupole coefficient  $ks3 = 1/(B\rho) d^3 B_y/dx^3$  [m<sup>-4</sup>] where (x,y) is now a coordinate system rotated by 30 degrees around s with respect to the normal one.(default 0).
- **tilt** - roll angle [rad] about the longitudinal axis, clockwise.
- **outR** - external radius [m] of magnet (default set to **aper+22cm**)
- **material** - the magnet material (default set to “Iron”)

Example :

```
of : octupole, l=0.5*m , k3 = 0.5 , tilt = 0.01;
```

### 3.3.10 multipole

**multipole** defines a multipole. Attributes:

- **l** - length [m] (default 0)
- **kn1** - normal multipole  $kn1[n] = 1/(B\rho) d^n B_y/dx^n$  [m<sup>-(n+1)</sup>]
- **ks1** - skew multipole  $ks1[n] = 1/(B\rho) d^n B_y/dx^n$  [m<sup>-(n+1)</sup>] where (x,y) is now a coordinate system rotated by 30 degrees around s with respect to the normal one.(default 0).
- **tilt** - roll angle [rad] about the longitudinal axis, clockwise.
- **outR** - external radius [m] of magnet (default set to **aper+22cm**)
- **material** - the magnet material (default set to “Iron”)

Example :

```
mul : multipole, l=0.5*m , kn1={ 0,0,1 } , ks1={ 0,0,0 };
```

Note that both **kn1** and **ks1** are required and must contain the same number of parameters.

### 3.3.11 rf

**rf** defines an rf cavity. Attributes:

- **l** - length [m] (default 0)
- **gradient** - field gradient [MV / m]

- **material** - the cavity material (default set to “Iron”)

Example :

```
rf1 : rf,l=5*m, gradient = 10 * MV / m;
```

### 3.3.12 rcol

**rcol** defines a rectangular collimator (the aperture is a rectangle, the external profile in the transverse plane is a square). The longitudinal collimator structure is not taken into account. To do this the user has to describe the collimator with the generic type **element**. Attributes:

- **l** - length [m] (default 0)
- **xsize** - horisontal aperture [m] (default set to **boxSize**)
- **ysize** - vertical aperture [m] (default set to **boxSize**)
- **outR** - external extent [m] in x and y of the collimator (default set to **boxSize**)
- **material** - collimator material (default set to “Graphite”)

Example :

```
col1 : rcol,l=0.4*m, xsize=2*mm, ysize=1*mm, material="W"
```

### 3.3.13 ecol

**ecol** defines an elliptical collimator (the aperture is an ellipse, the external profile in the transverse plane is a square). Here, again, the longitudinal collimator structure is not taken into account. Attributes:

- **l** - length [m] (default 0)
- **xsize** - horisontal aperture [m] (default set to **boxSize**)
- **ysize** - vertical aperture [m] (default set to **boxSize**)
- **outR** - limits external extent [m] in x and y of the collimator (default set to **boxSize**)
- **material** - collimator material (default set to “Graphite”)

Example :

```
col2 : ecol,l=0.4*m, xsize=2*mm, ysize=1*mm, material="W"
```

### 3.3.14 muspoiler

**muspoiler** defines a muon spoiler, which is a rotationally megnetised iron cylinder with an inner radius, outer radius, magnetic field strength and length. Attributes:

- **l** - length [m] (default 0)
- **inR** - inner radius [m] (default is outer beam pipe radius)
- **outR** - outer radius [m] (default set to **boxSize**)

- B - magnetic field [T] (default set to 1)

Example :

```
musp1 : muspoiler,l=5*m, inR=1*cm, outR=60*cm, B=1.5
```

### 3.3.15 solenoid

Not yet implemented

### 3.3.16 hkick and vkick

**hkick** and **vkick** are equivalent to a **rbend** and an **rbend** rotated by 90 degrees respectively. However, **hkick** and **vkick** do not rotate the frame of reference.

### 3.3.17 transform3d

An arbitrary 3-dimensional transformation of the coordinate system is done by placing a **transform3d** element in the beamline. Attributes:

- x = <x offset>
- y = <y offset>
- z = <z offset>
- phi = <phi Euler angle>
- theta = <theta Euler angle>
- psi = <psi Euler angle>

Example:

```
rot : transform3d, psi=pi/2
```

### 3.3.18 element

All the elements are in principle examples of a general type **element** which can represent an arbitrary geometric entity with arbitrary B field maps. Attributes:

- geometry = <geometry\_description>
- bmap = <bmap\_description>
- outR - limits external extent component box size (default set to **tunnelRadius/2**)

Descriptions are of the form

**format:filename**

where **filename** is the path to the file with the geometry description and **format** defines the geometry description format. The possible formats are given in Appendix A [Geometry], page 23.

Example :

```
qq : element, geometry ="mokka:qq.sql", bmap ="mokka:qq.bmap";
```

### 3.3.19 line

Elements are grouped into sequences by the `line` command.

```
line_name : line=(element_1,element_2,...);
```

where *element\_n* can be any element or another line. Lines can also be reversed using `line_name : line=-(line_2)`, or within another line by `line=(line_1,-line_2)`. Reversing a line also reverses all nested lines within.

Example :

A sequence of FODO cells can be defines as

```
qf: quadrupole, l=0.5, k1=0.1;
qd: quadrupole, l=0.5, k1=-0.1;
d: drift, l=0.5;
fodo : line=(qf,d,qd,d);
section : line=(fodo,fodo,fodo);
beamline : line=(section,section,section);
```

### 3.3.20 matdef

To define a material the `matdef` keyword must be used.

If the material is composed by a single element, it can be defined using the following syntax:<sup>5</sup>

```
<material> : matdef, Z=<int>, A=<double>, density=<double>, T=<double>,
P=<double>, state=<char*>;
```

Attributes

- Z - atomic number
- A - mass number [g/mol]
- density - density in [g/cm<sup>3</sup>]
- T - temperature in [K] (default set to 300)
- P - pressure [atm] (default set to 1)
- state - “solid”, “liquid” or “gas” (default set to “solid”)

Example:

```
iron : matdef, Z=26, A=55.845, density=7.87
```

---

<sup>5</sup> In this case, in `src/BDSDetectorConstruction.cc` the `BDSMaterials::AddMaterial(name, Z, A, density)` method is called, which in turns (`src/BDSMaterials.cc`) invokes the Geant4 `G4Material` constructor: `G4Material(name, Z, A, density);`

If the material is made up by several components, first of all each of them must be specified with the `atom` keyword:<sup>6</sup>

```
<element> : atom, Z=<int>, A=<double>, symbol=<char*>;
```

Attributes:

- `Z` - atomic number
- `A` - mass number [g/mol]
- `symbol` - atom symbol

Then the compound material can be specified in two manners:

1) If the number of atoms of each component in material unit is known, the following syntax can be used:<sup>7</sup>

```
<material> : matdef, density=<double>, T=<double>, P=<double>,
             state=<char*>, components=<[list<char*>]>,
             componentsWeights=<{list<int>}>;
```

Attributes

- `density` - density in [g/cm<sup>3</sup>]
- `T` - temperature in [K] (default set to 300)
- `P` - pressure in [atm] (default set to 1)
- `state` - “solid”, “liquid” or “gas” (default set to “solid”)
- `components` - list of symbols for material components
- `componentsWeights` - number of atoms of each component in material unit, in order

Example:

```
niobium : atom, symbol="Nb", z=41, a=92.906;
titanium : atom, symbol="Ti", z=22, a=47.867;
NbTi : matdef, density=5.6, temperature=4.0, ["Nb","Ti"], {1,1}
```

2) On the other hand, if the mass fraction of each component is known, the following syntax can be used:<sup>8</sup>

<sup>6</sup> In this case, in `src/BSDDetectorConstruction.cc` the `BDSMaterials::AddElement(name, symbol, Z, A)` method is called, which in turns (`src/BDSMaterials.cc`) invokes the Geant4 `G4Element` constructor: `G4Element(name, symbol, Z, A)`;

<sup>7</sup> In this case, in `src/BSDDetectorConstruction.cc` the `BDSMaterials::AddMaterial(name, density, state, temp, pressure, list<char*> itsComponents, list<G4int> itsComponentsWeights)` method is called, which in turns (`src/BDSMaterials.cc`) invokes the Geant4 `G4Material` constructor: `G4Material(name, density, (G4int)itsComponents.size(), state, temp, pressure)`. Then each component is added with a call to the `G4Material::AddElement(G4string, G4int)` method.

<sup>8</sup> In this case, in `src/BSDDetectorConstruction.cc` the `BDSMaterials::AddMaterial(name, density, state, temp, pressure, list<char*> itsComponents, list<G4double> itsComponentsFractions)` method is called, which in turns (`src/BDSMaterials.cc`) invokes the Geant4 `G4Material` constructor: `G4Material(name, density, (G4int)itsComponents.size(), state, temp, pressure)`. Then each component is added with a call to the `G4Material::AddElement(G4string, G4double)` method.

```
<material> : matdef, density=<double>, T=<double>, P=<double>,
            state=<char*>, components=<[list<char*>]>,>,
            componentsFractions=<{list<double>}>;
```

#### Attributes

- **density** - density in [g/cm<sup>3</sup>]
- **T** - temperature in [K] (default set to 300)
- **P** - pressure in [atm] (default set to 1)
- **state** - “solid”, “liquid” or “gas” (default set to “solid”)
- **components** - list of symbols for material components
- **componentsFractions** - mass fraction of each component in material unit, in order

Example:

```
samarium : atom, symbol="Sm", z= 62, a=150.4;
cobalt : atom, symbol="Co", z= 27, a=58.93;
SmCo : matdef, density=8.4, temperature=300.0, ["Sm","Co"],
      {0.338,0.662}
```

The second syntax can be used also to define materials which are composed by other materials (and not by atoms).

Nb: Square brackets are required for the list of element symbols, curly brackets for the list of weights or fractions.

### 3.3.21 laser

**laser** defines a drift section with a laser beam inside. The laser is considered to be the intersection of the laser beam with the volume of the drift section. Attributes:

- **l** - length of the drift section [m]
- **x,y,z** - components of the laser direction vector
- **waveLength** - laser wave length [m]

```
laserWire : laser, l=1*um,x=1,y=0,z=0,waveLength=532*nm
```

### 3.3.22 gas

To be implemented in v0.5

### 3.3.23 spec keyword

This was removed in v0.4 and no longer has an effect. For setting the outer radius of a quadrupole, use the **outR** parameter in the same way as for other elements.



### 3.3.24 Element number

When several elements with the same name are present in the beamline they can be accessed by their number in the sequence. In the next example the sampler is put before the second drift<sup>9</sup>

```
bl:line=(d,d,d);
sample,range=d[2];
```

### 3.3.25 Element attributes

Element attributes such as length, multipole coefficients etc, can be accessed by putting square brackets after the element name, e.g.

```
x=d[1];
```

### 3.3.26 Editing apertures

Apertures can be set after an element has already been defined by writing the element name followed by a semicolon followed by the attributes. For example, if quadrupole **qf** has already been defined then its aperture can be set to 4 mm using:

```
qf: aper=4*mm;
```

### 3.3.27 Material table

There is a set of predefined materials for use in elements such as collimators, e.g.

"Air"	"LiquidHelium"
"Aluminium"	"NbTi"
"BeamGasPlugMat"	"Niobium"
"Beryllium"	"Silicon"
"CarbonMonoxide"	"SmCo"
"CarbonSteel"	"Soil"
"Concrete"	"Titanium"
"Copper"	"TitaniumAlloy"
"Graphite"	"Tungsten"
"Invar"	"Vacuum"
"Iron"	"Vanadium"
"LaserVac"	"Water"
"Lead"	"WeightIron"
"LeadTungstate"	

Currently "Air", "CarbonMonoxide" and "Vacuum" are gas at T=300K, p=10<sup>-12</sup> bar: both "Air" and "Vacuum" are a N(80):O(20) mixture, "CarbonMonoxide" is composed of CO molecules.

There are also predefined elements (i.e. atoms) that can be used for building composite materials: "H", "He", "Be", "C", "N", "O", "Al", "Si", "P", "S", "Ca", "Ti", "V", "Mn", "Fe", "Co", "Ni", "Cu", "Nb", "Sm", "W", "Pb".

---

<sup>9</sup> See Appendix D [Known Issues], page 33

For more details see the file `src/BDSMaterials.cc` or run the command `bdsim --materials` from the command line.

## 3.4 Run control and output

The execution control is performed in the GMAD input file through `option` and `sample` commands. How the results are recorded is controlled by the `sample` command. When the visualization is turned on, it is also controlled through Geant4 command prompt

### 3.4.1 option

Most of the options in `bdsim` are set up by the command

```
option, <name>=value, ...;
```

The following options influence the geometry:

<code>beampipeRadius</code>	default beampipe outer radius [m]
<code>beampipeThickness</code>	default beampipe thickness [m]
<code>pipeMaterial</code>	default beampipe material
<code>boxSize</code>	default accelerator component size [m]
<code>vacMaterial</code>	the beam pipe gas material (default air)
<code>vacuumPressure</code>	the pressure of the beam pipe gas in bar (default 1e-12)
<code>buildTunnel</code>	whether to build a tunnel (default=0)
<code>buildTunnelFloor</code>	whether to add a floor to the tunnel (default=0)
<code>tunnelRadius</code>	tunnel radius [m]
<code>tunnelThickness</code>	the thickness of the tunnel wall [m]
<code>tunnelSoilThickness</code>	the thickness of the soil surrounding the tunnel [m]
<code>tunnelMaterial</code>	the material of the tunnel (default concrete)
<code>soilMaterial</code>	the material of the soil surrounding the tunnel (default soil)
<code>tunnelOffsetX</code>	the horizontal offset of the tunnel with respect to the beam line
<code>tunnelOffsetY</code>	the vertical offset of the tunnel with respect to the beam line
<code>tunnelFloorOffset</code>	the offset of the tunnel floor from the centre of the tunnel
<code>samplerDiameter</code>	the diameter of the sampler planes (default is 2 times <code>tunnelRadius</code> )
<code>blmRad</code>	the radius of the beam loss monitor cylinders
<code>blmLength</code>	the lengths of the beam loss monitor cylinders
<code>includeIronMagFields</code>	whether to include the magnetic fields in the magnet iron (default=1)

The following options influence the tracking:

<code>deltaChord</code>	chord finder precision
<code>deltaIntersection</code>	boundary intersection precision
<code>chordStepMinimum</code>	minimum step size
<code>lengthSafety</code>	element overlap safety

<code>minimumEpsilonStep</code>	minimum relative error acceptable in stepping
<code>maximumEpsilonStep</code>	maximum relative error acceptable in stepping
<code>deltaOneStep</code>	set position error acceptable in an integration steps
<code>synchPhotonMultiplicity</code>	multiplies the number of synchrotron radiation photons

The following options influence the physics:

<code>physicsList</code>	determines the set of physics processes used
<code>thresholdCutCharged</code>	charged particle cutoff energy
<code>thresholdCutPhotons</code>	photon cutoff energy
<code>stopTracks</code>	if set, tracks are terminated after interaction with material and energy deposit recorded
<code>synchRadOn</code>	turn on Synchrotron Radiation process
<code>srTrackPhotons</code>	whether to track the SR photons
<code>srLowX</code>	sets lowest energy of SR to $X \cdot E_{\text{critical}}$
<code>srLowGamE</code>	lowest energy of propagating SR photons
<code>synchPhotonMultiplicity</code>	a factor multiplying the number of synchrotron radiation photons
<code>prodCutPhotons</code>	standard overall production cuts for photons
<code>prodCutPhotonsP</code>	precision production cuts for photons in <b>element</b>
<code>prodCutElectrons</code>	standard overall production cuts for electrons
<code>prodCutElectronsP</code>	precision production cuts for electrons in <b>element</b>
<code>prodCutPositrons</code>	standard overall production cuts for positrons
<code>prodCutPositronsP</code>	precision production cuts for positrons in <b>element</b>
<code>turnOnCerenkov</code>	if set, Cerenkov radiation is turned on
<code>defaultRangeCut</code>	the default predicted range at which a particle is cut. Default is 0.7mm
<code>gammaToMuFe</code>	the cross section enhancement factor for the gamma to muon process
<code>annihiToMuFe</code>	the cross section enhancement factor for the electron-positron annihilation to muon process
<code>eetoHadronsFe</code>	the cross section enhancement factor for the electron-positron annihilation to hadrons process
<code>useEMLPB</code>	if set, electromagnetic lead particle biasing is used. Default is 0
<code>LPBFraction</code>	the fraction of EM processes in which electromagnetic lead particle biasing is used, from 0.0=never to 1.0=always

The following options influence the generation:

<code>randomSeed</code>	seed for the random number generator; setting to -1 uses the system clock to generate the seed
<code>ngenerate</code>	number of primary particles fired when in batch mode

The following options influence the output

<code>eLossHistoBinWidth</code>	bin width in metres for the energy loss histogram
<code>sensitiveBeamlineComponents</code>	if set, energy losses in beamline components are recorded in the energy loss histogram. Set by default
<code>sensitiveBeamPipe</code>	if set, energy losses in the beam pipe are recorded in the energy loss histogram. Set by default
<code>sensitiveBLMs</code>	if set, energy losses in the beam loss monitors are recorded in the energy loss histogram. Set by default
<code>storeTrajectory</code>	if set, the trajectories are stored in the root file
<code>storeMuonTrajectories</code>	if set, the muon trajectories are stored in the root file
<code>storeNeutronTrajectories</code>	if set, the neutron trajectories are stored in the root file
<code>trajCutGTZ</code>	do not store any trajectories who end less than this z distance
<code>trajCutLTR</code>	do not store any trajectories who end outside of this radius

Miscellaneous options:

<code>nperfile</code>	number of events recorded per file in ROOT output
<code>nlinesIgnore</code>	number of lines to skip when reading bunch files

For a more detailed description of how the option influence the tracking see Chapter 5 [Physics], page 21

### 3.4.2 beam

The parameters related to the beam are set with the `beam` command

```
beam, <name>=value, ...;
```

There is a set of predefined distribution types that can be generated<sup>10</sup>. In this case one needs to specify the following parameters:

- `particle` - particle name, "e-", "e+", "gamma", "proton", etc
- `energy` - particle energy
- `distrType` - type of distribution

and, in addition, other parameters that depend on the distribution type that has been chosen:

1. Global options:

- `X0` - Offset of distribution centre in x[m]
- `Y0` - Offset of distribution centre in y[m]
- `Z0` - Offset of distribution centre in z[m]
- `Xp0` - Angular offset from nominal axis in x-z plane
- `Yp0` - Angular offset from nominal z axis in y-z plane
- `Zp0` - Directional flag: `Zp0 < 0` points the particle back up the beamline
- `T0` - Global time offset [s]

---

<sup>10</sup> see `src/BDSBunch.cc` for more details

2. `distrType="gauss"`: a gaussian in  $x, x', y, y'$ , energy and time, with given widths:
  - `sigmaX` - RMS of  $x$  distribution in [m]
  - `sigmaXp` - RMS of  $x'$  distribution in [rad]
  - `sigmaY` - RMS of  $y$  distribution in [m]
  - `sigmaYp` - RMS of  $y'$  distribution in [rad]
  - `sigmaE` - RMS of energy distribution divided by nominal beam kinetic energy
  - `sigmaT` - RMS of time distribution in [s]
3. `distrType="gausstwiss"`: a gaussian bunch defined by twiss parameters [4], page 34, emittance, energy and time:
  - `betx` -  $\beta_x$  in [m]
  - `bety` -  $\beta_y$  in [m]
  - `alfx` -  $\alpha_x$
  - `alfy` -  $\alpha_y$
  - `emitx` -  $\epsilon_x$  in [m]
  - `emity` -  $\epsilon_y$  in [m]
  - `sigmaE` - RMS of energy distribution divided by nominal beam kinetic energy
  - `sigmaT` - RMS of time distribution in [s]
4. `distrType="eshell"`: a thin elliptic shell in  $x, x'$  and  $y, y'$  with given semiaxes:
  - `shellX`
  - `shellXp`
  - `shellY`
  - `shellYp`
  - `sigmaE`
5. `distrType="ring"`: in the  $x, y$  plane the particles are uniformly distributed in  $r$  and in  $\phi$  inside a ring with inner radius `Rmin` and outer radius `Rmax`.  $x', y'$  and time are exactly `Xp0, Yp0` and `T0` respectively for each generated particle. The kinetic energy distribution is a gaussian of width `sigmaE` centered about the nominal beam kinetic energy.
  - `Rmin, Rmax` - inner and outer radius in [m]
  - `sigmaE` - RMS energy spread [GeV]

Example:

```
beam, particle="e+", energy=100*MeV, distrType="gauss", sigmaX=0.01,
      sigmaXp=0.1, sigmaY=0.01, sigmaYp=0.1;
```

In alternative, one can pass to the simulation a file containing a list of particles to be generated. For more details see Appendix C [Bunch description formats], page 33.

### 3.4.3 sample and csample

To record the tracking results one uses the `sample` and `csample` commands. To insert a sampling plane before `<element>` the following command should be used:

```
sample, range=<element>;
```

Example:

```
sample, range=d;
```

To put a cylindrical sampler of length 10 (in [m]) around element `<element>` at distance `r0` (in [m]) the following command should be used:

```
csample, range=<element>, r=r0, l=10;
```

Samplers output the following parameters at the specified location:

E	Energy[GeV]	E0	Energy at last scatter[GeV]
X	Global X position	s	path length
Y	Global Y position	t	time of flight
Z	Global Z position	t0	time of flight at last scatter
Xp	Global angle in x-z	trackID	trackID of particle
Yp	Global angle in y-z	weight	weight of track
Zp	$1-\sqrt{Xp^2+Yp^2}$	parentID	trackID of parent particle
x	Relative x position	x0	x at last scatter
y	Relative y position	y0	y at last scatter
z	Relative z position <sup>11</sup>	z0	z at last scatter
xp	Relative angle in x-z	xp0	xp at last scatter
yp	Relative angle in y-z	yp0	yp at last scatter
zp	$1-\sqrt{xp^2+yp^2}$	zp0	xp at last scatter
nEvent	Event number	partID	PDG particle identifier

### 3.4.4 dump

Used in conjunction with `option,fifo=<filename>` to output the bunch distribution at a given point. If the specified output file is a fifo, the distribution can be modified by an external program before being piped back in to continue tracking. This is useful for including multi-particle effects such as wakefields at given points in the lattice.

```
dump, range=dumpMarker1
option, fifo="/tmp/temp.dat"
```

Output is in the standard Guineapig format, with a header line stating the number of particles to be output. The file to be read back should be in the same format as this.

### 3.4.5 use

`use` command selects the beam line for study

```
use, period=l1, range=q1/q2
```

---

<sup>11</sup> See Appendix D [Known Issues], page 33

## 4 Visualization

When BDSIM is invoked in interactive mode, the run is controlled by the Geant4 shell. A visualization macro should be then provided. A simple visualization macro is include with the distribution, and is outlined below.

```
# Invoke the OGLSX driver
# Create a scene handler and a viewer for the OGLSX driver
/vis/open OGLIX

# Create an empty scene
/vis/scene/create

# Add detector geometry to the current scene
/vis/scene/add/volume

# Attach the current scene handler
# to the current scene (omittable)
/vis/sceneHandler/attach

# Add trajectories to the current scene
# Note: This command is not necessary in exampleN03,
#       since the C++ method DrawTrajectory() is
#       described in the event action.

/vis/viewer/set/viewpointThetaPhi 90 90
# /vis/drawVolume
#/vis/scene/add/trajectories
# /tracking/storeTrajectory 0
#/vis/viewer/zoom
/tracking/storeTrajectory 1
#
# for BDS:
#/vis/viewer/zoom 300
#/vis/viewer/set/viewpointThetaPhi 3 45
```

By default the macro is read from the file named `vis.mac` located in the current directory. The name of the file with the macro can also be passed via the `vis_mac` switch.

```
bdsim --file=line.gmad --vis_mac=my_macro.mac
```

In interactive mode all the Geant4 interactive comamnds are available. For instance, to fire 100 particles type

```
/run/beamOn 100
```

and to end the session type

```
exit
```

To display help menu

```
/help;
```

For more details see [1], page 34.

## 5 Physics

BDSIM can exploit all physics processes that come with Geant4. In addition fast tracking inside multipole magnets is provided. More detailed description of the physics is given below.

### 5.1 physicsList option

Depending on for what sort of problem BDSIM is used, different sorts of physics processes should be turned on. These processes are grouped into so called “physics lists”. The physics list is specified by the `physicsList` option in the input file, e.g.

```
option, physicsList="em_standard";
```

Several predefined physics lists are available. Some physics lists allow biasing and re-weighting for some processes e.g. muon production. To set the amount of biasing see Section 3.4.1 [option], page 15. Further details of the QGSP, FTFP and BERT hadronic physics lists can be found in [5], page 34.

<code>standard</code>	transportation of primary particles only
<code>em_standard</code>	transportation of primary particles, ionization, bremsstrahlung, Cerenkov, multiple scattering
<code>em_low</code>	the same but using low energy electromagnetic models
<code>em_muon</code>	<code>em_standard</code> plus muon production processes with biased muon cross-sections
<code>lw</code>	list for laser wire simulation - standard electromagnetic physics and "laser wire" physics which is Compton Scattering with total cross-section renormalized to 1.
<code>merlin</code>	transportation of primary particles, and the following processes for electrons: multiple scattering, ionisation, and bremsstrahlung
<code>hadronic_standard</code>	<code>em_standard</code> plus fission, neutron capture, neutron and proton elastic and inelastic scattering
<code>hadronic_muon</code>	<code>hadronic_standard</code> plus muon production processes with biased muon cross-sections
<code>hadronic_QGSP_BERT</code>	<code>em_standard</code> plus hadron physics using the quark gluon string plasma (QGSP) model and the Bertini cascade model (BERT)
<code>hadronic_QGSP_BERT_muon</code>	<code>hadron_QGSP_BERT</code> plus muon production processes with biased muon cross-sections
<code>hadronic_QGSP_BERT_HP_muon</code>	<code>hadron_QGSP_BERT_muon</code> with high precision neutron tracking



<code>hadronic_FTFP_BERT</code>	<code>em_standard</code> plus hadron physics using the Fritiof model followed by Reggion cascade and Precompound and evaporation models for the nucleus de-excitation (FTFP) model and the Bertini cascade model (BERT)
<code>hadronic_FTFP_BERT_muon</code>	<code>hadronic_FTFP_BERT</code> plus muon production processes with biased muon cross-sections

By default the `standard` physics List is used

## 5.2 Transportation

The transportation follows the scheme: the step length is selected which is defined either by the distance of the particle to the boundary of the “logical volume” it is currently in (which could be, e.g. field boundary, material boundary or boundary between two adjacent elements) or by the mean free path of the activated processes. Then the particle is pushed to the new position and secondaries are generated if necessary. Each volume has an associated transportation algorithm. For an on-energy particle travelling close to the optical axis of a quadrupole, dipole or a drift, standard matrix transportation algorithms are used [4], page 34. For multipoles of higher orders and for off-axis/energy particles Runge-Kutta methods are used.

## 5.3 Tracking accuracy

The following options influence the tracking accuracy

<code>chordStepMinimum</code>	minimum chord length for the step
<code>deltaIntersection</code>	determines the precision of locating the point of intersection of the particle trajectory with the boundary and hence the error in the path length in each volume. This may influence the results especially in the case when EM fields are present.
<code>deltaChord</code>	
<code>lengthSafety</code>	all volumes will have an additional overlap of this length
<code>thresholdCutCharged</code>	energy below which charged particles are not tracked
<code>thresholdCutPhotons</code>	energy below which photons are not tracked

# 6 Output Analysis

During the execution the following things are recorded:

- energy deposition along the beamline
- sampler hits

If the output format is ASCII i.e. if BDSIM was invoked with the `--output=ascii` option, then the output file “output.txt” containing the hits will be written which has rows like

```
#hits PDGtype p[GeV/c] x[micron] y[micron] z[m] x'[microrad] y'[microrad]
11 250 -4.72907 -5.86656 5.00001e-06 0 0
11 250 -8.17576 -4.99729 796.001 0.320334 -0.126792
```

If ROOT output is used then the root files `output_0.root`, `output_1.root` etc. will be created with each file containing the number of events given by `nperfile` option. The file contains the energy loss histogram and a tree for every sampler in the line with self-explanatory branch names.

## Appendix A Geometry description formats

The element with user-defined physical geometry is defined by

```
<element_name> : element, geometry=format:filename, attributes
```

for example,

```
colli : element, geometry="gmad:colli.geo"
```

### A.1 gmad format

`gmad` is a simple format used as G4geometry wrapper. It can be used for specifying more or less simple geometries like collimators. Available shapes are:

```
Box {
  x0=x_origin,
  y0=y_origin,
  z0=z_origin,
  x=xsize,
  y=ysize,
  z=zsize,
  phi=Euler angle for rotation,
  theta=Euler angle for rotation,
  psi=Euler angle for rotation,
  material=MaterialName
}
```

```
Tubs {
  x0=x_origin,
  y0=y_origin,
  z0=z_origin,
  rmin=inner radius,
  rmax=outer radius,
  z=zsize,
  phi=Euler angle for rotation,
  theta=Euler angle for rotation,
  psi=Euler angle for rotation,
  material=MaterialName
}
```

```

Cons {
  x0=x_origin,
  y0=y_origin,
  z0=z_origin,
  rmin1=inner radius at start,
  rmax1=outer radius at start,
  rmin2=inner radius at end,
  rmax2=outer radius at end,
  z=zsize,
  material=MaterialName,
  phi=Euler angle for rotation,
  theta=Euler angle for rotation,
  psi=Euler angle for rotation,
  phi0=angle for start of sector,
  dphi=angle swept by sector
}

Trd {
  x0=x_origin,
  y0=y_origin,
  z0=z_origin,
  x1=half length at wider side,
  x2=half length at narrower side,
  y1=half length at wider side,
  y2=half length at narrower side,
  z=zsize,
  phi=Euler angle for rotation,
  theta=Euler angle for rotation,
  psi=Euler angle for rotation,
  material=MaterialName
}

```

A file can contain several objects which will be placed sequentially into the volume, A user has to make sure that there is no overlap between them.

## A.2 mokka

As well as using the gmad format to describe user-defined physical geometry it is also possible to use a Mokka style format. This format is currently in the form of a dumped MySQL database format - although future versions of BDSIM will also support online querying of MySQL databases. Note that throughout any of the Mokka files, a # may be used to represent a commented line. There are three key stages, which are detailed in the following sections, that are required to setting up the Mokka geometry:

- Describing the geometry
- Creating a geometry list
- Defining a Mokka Element to load geometry descriptions from a list

### A.2.1 Describing the geometry

An object must be described by creating a MySQL file containing commands that would typically be used for uploading/creating a database and a corresponding new table into a MySQL database. BDSIM supports only a few such commands - specifically the **CREATE TABLE** and **INSERT INTO** commands. When writing a table to describe a solid there are some parameters that are common to all solid types (such as **NAME** and **MATERIAL**) and some that are more specific (such as those relating to radii for cone objects). A full list of the standard and specific table parameters, as well as some basic examples, are given below with each solid type. All files containing geometry descriptions must have the following database creation commands at the top of the file:

```

DROP DATABASE IF EXISTS DATABASE_NAME;
CREATE DATABASE DATABASE_NAME;
USE DATABASE_NAME;

```

A table must be created to allow for the insertion of the geometry descriptions. A table is created using the following, MySQL compliant, commands:

```
CREATE TABLE TABLE-NAME_GEOMETRY-TYPE (

TABLE-PARAMETER                                VARIABLE-TYPE,

TABLE-PARAMETER                                VARIABLE-TYPE,

TABLE-PARAMETER                                VARIABLE-TYPE

);
```

Once a table has been created values must be entered into it in order to define the solids and position them. The insertion command must appear after the table creation and must be the MySQL compliant table insertion command:

```
INSERT INTO TABLE-NAME_GEOMETRY-TYPE VALUES(value1, value2, "char-value",
...);
```

The values must be inserted in the same order as their corresponding parameter types are described in the table creation. Note that ALL length types must be specified in mm and that ALL angles must be in radians.

An example of two simple boxes with no visual attributes set is shown below. The first box is a simple vacuum cube whilst the second is an iron box with length\_x = 10mm, length\_y = 150mm, length\_z = 50mm, positioned at x=1m, y=0, z=0.5m and with zero rotation.

```
CREATE TABLE mytable_BOX (

NAME                                VARCHAR(32),

MATERIAL                            VARCHAR(32),

LENGTHX                             DOUBLE(10,3),

LENGTHY                             DOUBLE(10,3),

LENGTHZ                             DOUBLE(10,3),

POSX                                DOUBLE(10,3),

POSY                                DOUBLE(10,3),

POSZ                                DOUBLE(10,3),

ROTPSI                              DOUBLE(10,3),
```

```

ROTTHETA                                DOUBLE(10,3),

ROTPHI                                  DOUBLE(10,3)

);

INSERT INTO mytable_BOX VALUES("a_box","vacuum", 50.0, 50.0, 50.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0);
INSERT INTO mytable_BOX VALUES("another_box","iron", 10.0, 150.0, 50.0,
1000.0, 0.0, 500.0, 0.0, 0.0, 0.0);

```

Further examples of the Mokka geometry implementation can be found in the examples/Mokka/General directory. See the common table parameters and solid type sections below for more information on the table parameters available for use.

### A.2.1.1 Common Table Parameters

The following is a list of table parameters that are common to all solid types either as an optional or mandatory parameter:

- **NAME**

Variable type: `VARCHAR(32)`

This is an optional parameter. If supplied, then the Geant4 LogicalVolume associated with the solid will be labelled with this name. The default is set to be the table's name plus an automatically assigned volume number.

- **MATERIAL**

Variable type: `VARCHAR(32)`

This is an optional parameter. If supplied, then the volume will be created with this material type - note that the material must be given as a character string inside double quotation marks(""). The default material is set as Vacuum.

- **PARENTNAME**

Variable type: `VARCHAR(32)`

This is an optional parameter. If supplied, then the volume will be placed as a daughter volume to the object with ID equal to **PARENTNAME**. The default parent is set to be the Component Volume. Note that if **PARENTID** is set to the Component Volume then **POSZ** will be defined with respect to the start of the object. Else **POSZ** will be defined with respect to the center of the parent object.

- **INHERITSTYLE**

Variable type: `VARCHAR(32)`

This is an optional parameter to be used with **PARENTNAME**. If set to "SUBTRACT" then the instead of placing the volume within the parent volume as an inherited object, it will be subtracted from the parent volume in a boolean solid operation. The default for this value is set to "" - which sets to the usual mother/daughter volume inheritance.

- **ALIGNIN**

Variable type: `INTEGER(11)`

This is an optional parameter. If set to 1 then the placement of components will be rotated and translated such that the incoming beamline will pass through the z-axis of this object. The default is set to 0.

- **ALIGNOUT**

Variable type: `INTEGER(11)`

This is an optional parameter. If set to 1 then the placement of the next beamline component will be rotated and translated such that the outgoing beamline will pass through the z-axis of this object. The default is set to 0.

- **SETSENSITIVE**

Variable type: `INTEGER(11)`

This is an optional parameter. If set to 1 then the object will be set up to register energy depositions made within it and to also record the z-position at which this deposition occurs. This information will be saved in the ELoss Histogram if using ROOT output. The default is set to 0.

- **MAGTYPE**

Variable type: `VARCHAR(32)`

This is an optional parameter. If supplied, then the object will be set up to produce the appropriate magnetic field using the supplied K1 or K2 table parameter values . Three magnet types are available - “QUAD”, “SEXT” and “OCT”. The default is set to no magnet type. Note that if **MAGTYPE** is set to a value whilst K1/K2/K3 are not set, then no magnetic field will be implemented.

- **K1**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set to a value other than zero, in conjunction with **MAGTYPE** set to “QUAD” then a quadrupole field with this K1 value will be set up within the object. Default is set to zero.

- **K2**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set to a value other than zero, in conjunction with **MAGTYPE** set to “SEXT” then a sextupole field with this K2 value will be set up within the object. Default is set to zero.

- **K3**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set to a value other than zero, in conjunction with **MAGTYPE** set to “OCT” then a sextupole field with this K3 value will be set up within the object. Default is set to zero.

- **POSX, POSY, POSZ**

Variable type: `DOUBLE(10,3)`

These are required parameters. They are form the position in mm used to place the object in the component volume. **POSX** and **POSY** are defined with respect to the center of the component volume and with respect to the component volume’s rotation. **POSZ** is defined with respect to the start of the component volume. Note that if the object

is being placed inside another volume using PARENTNAME then the position will refer to the center of the parent object.

- ROTPSI, ROTTHETA, ROTPHI

Variable type: DOUBLE(10,3)

These are optional parameters. They are the Euler angles in radians used to rotate the object before it is placed. The default is set to zero for each angle.

- RED, BLUE, GREEN

Variable type: DOUBLE(10,3)

These are optional parameters. They are the RGB colour components assigned to the object and should be a value between 0 and 1. The default is set to zero for each colour.

- VISATT

Variable type: VARCHAR(32)

This is an optional parameter. This is the visual state setting for the object. Setting this to “W” results in a wireframe displayment of the object. “S” produces a shaded solid and “I” leaves the object invisible. The default is set to be solid.

- FIELDX, FIELDY, FIELDZ

Variable type: DOUBLE(10,3)

These are optional parameters. They can be used to apply a uniform field to any volume, with default units of Tesla. Note that if there is a solenoid field present throughout the entire element then this uniform field will act in addition to the solenoid field.

### A.2.1.2 'Box' Solid Types

Append \_BOX to the table name in order to make use of the G4Box solid type. The following table parameters are specific to the box solid:

- LENGTHX, LENGTHY, LENGTHZ

Variable type: DOUBLE(10,3)

These are required parameters. Their values will be used to specify the box's dimensions.

### A.2.1.3 'Trapezoid' Solid Types

Append \_TRAP to the table name in order to make use of the G4Trd solid type - which is defined as a trapezoid with the X and Y dimensions varying along z functions. The following table parameters are specific to the trapezoid solid:

- LENGTHXPLUS

Variable type: DOUBLE(10,3)

This is a required parameter. This value will be used to specify the x-extent of the box's dimensions at the surface positioned at +dz.

- LENGTHXPMINUS

Variable type: DOUBLE(10,3)

This is a required parameter. This value will be used to specify the x-extent of the box's dimensions at the surface positioned at -dz.

- **LENGTHYPLUS**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the y-extent of the box's dimensions at the surface positioned at +dz.

- **LENGTHYMINUS**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the y-extent of the box's dimensions at the surface positioned at -dz.

- **LENGTHZ**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the z-extent of the box's dimensions.

### A.2.1.4 'Cone' Solid Types

Append `_CONE` to the table name in order to make use of the `G4Cons` solid type. The following table parameters are specific to the cone solid:

- **LENGTH**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the z-extent of the cone's dimensions.

- **RINNERSTART**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set then this value will be used to specify the inner radius of the start of the cone. The default value is zero.

- **RINNEREND**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set then this value will be used to specify the inner radius of the end of the cone. The default value is zero.

- **ROUTERSTART**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the outer radius of the start of the cone.

- **ROUTEREND**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the outer radius of the end of the cone.

- **STARTPHI**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set then this value will be used to specify the starting angle of the cone. The default value is zero.



- DELTAPHI

Variable type: DOUBLE(10,3)

This is an optional parameter. If set then this value will be used to specify the delta angle of the cone. The default value is  $2\pi$ .

### A.2.1.5 'Torus' Solid Types

Append `_TORUS` to the table name in order to make use of the G4Torus solid type. The following table parameters are specific to the torus solid:

- RINNER

Variable type: DOUBLE(10,3)

This is an optional parameter. If set then this value will be used to specify the inner radius of the torus tube. The default value is zero.

- ROUTER

Variable type: DOUBLE(10,3)

This is a required parameter. This value will be used to specify the outer radius of the torus tube.

- RSWEPT

Variable type: DOUBLE(10,3)

This is a required parameter. This value will be used to specify the swept radius of the torus. It is defined as being the distance from the center of the torus ring to the center of the torus tube. For this reason this value should not be set to less than `ROUTER`.

- STARTPHI

Variable type: DOUBLE(10,3)

This is an optional parameter. If set then this value will be used to specify the starting angle of the torus. The default value is zero.

- DELTAPHI

Variable type: DOUBLE(10,3)

This is an optional parameter. If set then this value will be used to specify the delta swept angle of the torus. The default value is  $2\pi$ .

### A.2.1.6 'Polycone' Solid Types

Append `_POLYCONE` to the table name in order to make use of the G4Polycone solid type. The following table parameters are specific to the polycone solid:

- NZPLANES

Variable type: INTEGER(11)

This is a required parameter. This value will be used to specify the number of z-planes to be used in the polycone. This value must be set to greater than 1.

- PLANEPOS1, PLANEPOS2, ..., PLANEPOSN

Variable type: DOUBLE(10,3)

These are required parameters. These values will be used to specify the z-position of the corresponding z-plane of the polycone. There should be as many `PLANEPOS` parameters set as the number of z-planes. For example, 3 z-planes will require that `PLANEPOS1`, `PLANEPOS2`, and `PLANEPOS3` are all set up.

- **RINNER1, RINNER2, ..., RINNERN**

Variable type: `DOUBLE(10,3)`

These are required parameters. These values will be used to specify the inner radius of the corresponding z-plane of the polycone. There should be as many **RINNER** parameters set as the number of z-planes. For example, 3 z-planes will require that **RINNER1**, **RINNER2**, and **RINNER3** are all set up.

- **ROUTER1, ROUTER2, ..., ROUTERN**

Variable type: `DOUBLE(10,3)`

These are required parameters. These values will be used to specify the outer radius of the corresponding z-plane of the polycone. There should be as many **ROUTER** parameters set as the number of z-planes. For example, 3 z-planes will require that **ROUTER1**, **ROUTER2**, and **ROUTER3** are all set up.

- **STARTPHI**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set then this value will be used to specify the starting angle of the polycone. The default value is zero.

- **DELTAPHI**

Variable type: `DOUBLE(10,3)`

This is an optional parameter. If set then this value will be used to specify the delta angle of the polycone. The default value is  $2\pi$ .

### A.2.1.7 'Elliptical Cone' Solid Types

Append `_ELLIPTICALCONE` to the table name in order to make use of the `G4Ellipticalcone` solid type. The following table parameters are specific to the elliptical cone solid:

- **XSEMIAXIS**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the Semiaxis in X.

- **YSEMIAXIS**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the Semiaxis in Y.

- **LENGTHZ**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the height of the elliptical cone.

- **ZCUT**

Variable type: `DOUBLE(10,3)`

This is a required parameter. This value will be used to specify the upper cut plane level.

Note that the above parameters are used to define an elliptical cone with the following parametric equations (in the usual Geant4 way):

$$x = XSEMIAXIS * (LENGTHZ - u) / u * \cos(v)$$

$$Y = YSEMIAXIS * (LENGTHZ - u) / u * \sin(v)$$

$$z = u$$

where  $v$  is between 0 and  $2\pi$  and  $u$  between 0 and  $h$  respectively.

### A.2.2 Creating a geometry list

A geometry list is a simple file consisting of a list of filenames that contain geometry descriptions. This is the file that should be passed to the GMAD file when defining the mokka element. An example of a geometry list containing 'boxes.sql' and 'cones.sql' would be:

```
# '#' symbols can be used for commenting out an entire line

/directory/boxes.sql

/directory/cones.sql
```

### A.2.3 Defining a Mokka element in the gmad file

The Mokka element can be defined by the following command:

```
<element_name> : element, geometry=format:filename, attributes
```

where `format` must be set to `mokka` and `filename` must point to a file that contains a list of files that have the geometry descriptions.

for example,

```
collimator : element, geometry=mokka:coll_geomlist.sql
```

## A.3 gdml

GDML is a XML schema for detector description. GDML will be supported as an external format starting from next release.

## Appendix B Field description formats

The element with user-defined magnetic field map is defined by the command

```
<element_name> : element, bmap=format:filename, attributes
```

for example,

```
colli : element, bmap=XY:colli.bmap
```

Supported formats are “mokka” and “XY”. In the latter case a text files must be specified, where each rows must have the following format:  $x$   $y$   $B_x$   $B_y$   $B_z$

## Appendix C Bunch description formats

For compatibility with other simulation codes following bunch formats can be read. For example, to use the file `distr.dat` as input the beam definition should look like

```
beam, particle="e-",distrType="guineapig_bunch",distrFile="distr.dat"
```

The formats currently supported are listed below:<sup>12</sup>

- `guineapig_bunch` : E[GeV] x[mum] y[mum] z[mum] x'[murad] y'[murad]
- `guineapig_slac` : E[GeV] x'[rad] y'[rad] z[mum] x[nm] y[nm]
- `guineapig_pairs` : E[GeV] x'[rad] y'[rad] z'[rad] x[nm] y[nm] z[nm] (here a particle with E>0 is assumed to be an electron and with E<0 a positron.)
- `cain` : ....

A custom distribution file format can be specified in the form

```
distrType="field1[unit1]:field1[unit1]:...
```

The allowed values for fields/units are: .....

For instance:

```
beam, particle="e-",
      energy=ener * GeV,
      distrType="pt[1]:E[GeV]:xp[rad]:yp[rad]:z[mum]:x[nm]:y[nm]",
      distrFile="bunches/beam.dat";
```

## Appendix D Known Issues

A bug is present where a sampler attached to a bending magnet (RBend/SBend) will cause the magnetic field to fail to be set. The reference frame rotates correctly but the particle trajectory does not follow. To work around this issue, samplers should be attached to a marker rather than directly to the magnet. For example:

```
dip: sbend,l=1*m, angle=0.1;
temp: line=(dip);
use, period=temp;
sample, range=dip;
```

should be replaced by:

```
dip: sbend,l=1*m, angle=0.1;
dipMark: marker;
temp: line=(dipMark,dip);
use, period=temp;
sample, range=dipMark;
```

Samplers attached to multiple instances of the same element incorrectly register hits only from the first instance in all such samplers. For example:

---

<sup>12</sup> see `src/BDSBunch.cc` for more details

```

drift1: drift, l=1*m;
mark1: marker;
line1: line=(mark1,drift1,mark1,drift1);
sample, range=mark1[1];
sample, range=mark1[2];

```

will incorrectly record hits at `mark1[1]` in the sampler attached to `mark1[2]`. To avoid this, samplers should be attached to uniquely named elements.

There is a known issue with ROOT output using ROOT versions higher than 5.10. To use ROOT output, BDSIM should be compiled with ROOT version 5.10 or lower.

There is a known issue with the `z` parameter output to samplers. As particle data is output at the `z` location of the sampler, when the global position is transformed from global to relative coordinates `z` is identically zero. For a description of a particle's longitudinal position in the bunch, please use the parameter `s` instead.

## 7 References

- [1] Geant4 User's Guide, <http://geant4.cern.ch/support/userdocuments.shtml>
- [2] Root User's Guide, <http://root.cern.ch/root/doc/RootDoc.html>
- [3] MAD-X User's Guide, <http://mad.home.cern.ch/mad/uguide.html>
- [4] for example 'Basic course on Accelerator optics' by Schmuesser, Rossbach, CERN Accelerator school
- [5] A. Ribon et. al., Status of GEANT4 hadronic physics for the simulation of LHC experiments at the start of LHC physics program, CERN-LCGAPP-2010-02, July 20 2010