

# Your First Input and beyond

21st FLUKA Beginner's Course ALBA – Barcelona, Spain 08 – 12 April 2019

# Before starting: FLUKA Manual

in *continuous* development (as the program)! More a User Guide than a Reference Manual (only a short summary about physics)

#### FM.pdf

update of the published CERN yellow report ToC, cross-references, and citations are active links analytical index at the end

#### **ASCII**

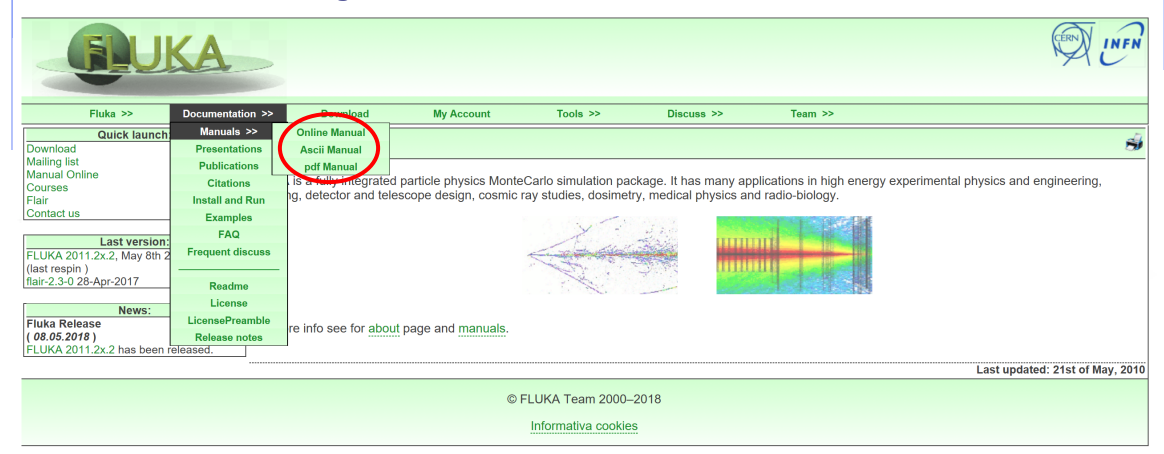
fluka2011.manual (no figures)

Tk interface accessible through FLAIR or from command line /usr/local/bin/fm

(HTML version is available on FLUKA website)

# Before starting: FLUKA Manual

#### **FLUKA** website at fluka.org



### Before starting: FLUKA Manual

#### **Short description of FLUKA**

installation, program and its capabilities, implemented physics models,...

#### **User guide**

- detailed description of input options
- available particles and materials
- combinatorial geometry
- low-energy neutrons library
- how to write/compile/link user routines
- ....and much more!

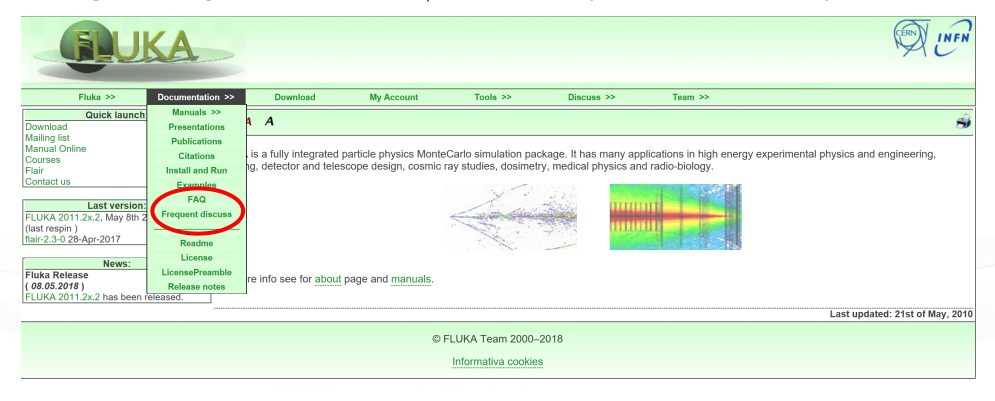
#### **REMEMBER:**

The first place to look at when puzzled

...and the very best friend of a beginner user!

### Before starting: FLUKA FAQ

**FLUKA FAQ and Frequent discuss** The second place to look at when puzzled!



Write at: fluka-discuss@fluka.org

**REMEMBER**: send always your input file together with your question to the fluka-discuss.

### Structure of the input file

### **General definitions**

Beam definition
Materials: definition and assignment
Random number initialization
Start/Stop of simulation

### Physics settings

Defaults
Physical processes
Transport thresholds
Low energy neutrons
Induced radioactivity

### **Biasing**

Geometry related biasing Interaction/decay biasing

### Geometry

Setup description Voxel phantoms

#### **Output settings**

Estimators / scoring cards

### FLUKA input file commands

Commands aka cards, aka options, aka directives, aka definitions

Card: One keyword (command), 6 floating point numbers (WHATs), one string (SDUM) Example of a FLUKA command (text editor style)

```
*...+...5...+...6...+...7...+...
                       0.0
                               0.0
                                        0.0
                                                0.0
BEAM
            1.E + 04
                                                         0.0PROTON
*keyword
          momentum mom.spread
                            diverg.
                                    X-width
                                           Y-width ignored particle
           WHAT (1)
                   WHAT (2)
                                                      WHAT (6)
                            WHAT (3)
                                    WHAT (4)
                                             WHAT (5)
                                                             SDUM
```

- Command keywords MUST be uppercase, numbers MUST have the decimal point
- Some commands require more than one "card"
- Some special commands (like **TITLE** and **OPEN**) are/may be followed by a text line
- With few exceptions, the order of commands is irrelevant
- Most commands can be repeated several times
- Repeated commands can add themselves or <u>override</u> previous commands
- A line with a \* character in column 1 is a comment
- Text after an exclamation mark (!) is ignored (does not work within the geometry)
- Almost all the WHAT() have a default value
- Commands can be issued in fixed or free format
- Special commands, called #directives, allow input parametrization

### Fixed format

#### Fixed format:

```
*...+...5....+...6....+...7....+...
                   0.0D + 00
                               0.0
                                        0.0
            1.E+04
                                                         0.0PROTON
BEAM
          momentum mom.spread diverg.
*keyword
                                    X-width
                                             Y-width
                                                     ignored particle
           WHAT (1)
                   WHAT (2)
                            WHAT (3)
                                    WHAT (4)
                                             WHAT (5)
                                                      WHAT (6)
                                                             SDUM
```

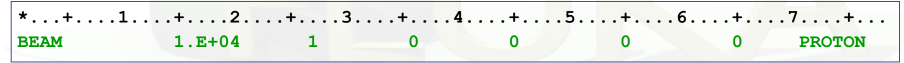
- The "traditional" FLUKA format is (A8, 2X, 6E10.0, A8)
   Numbers: 9 digits at most can be used
- All WHAT fields are in floating point format, <u>even integers</u>
   They <u>must</u> always be written with the decimal point
- Exponential notation numbers (e.g. 1.234E+5), must be right aligned
- Double precision format (e.g. 1.234D+5) is allowed
- Blank numerical fields are read as 0.0
   In most cases (not all!) such values are ignored and the corresponding default values are used
- Blank lines NOT ALLOWED in geometry declaration (tolerated elsewhere)

### Fixed format - Examples

Both lines are correct

```
0.0
                                                0.0
              1.E+04
                            0.0
BEAM
                                                           0.0
                                                                      0.0PROTON
                                   0.0
                                             0.0
              1.E + 04
                         0.0
                                                        0.0
                                                                  0.0
BEAM
                                                                         PROTON
*keyword
            momentum mom.spread
                                  diverg.
                                            X-width
                                                     Y-width ignored particle
*
             WHAT (1)
                                            WHAT (4)
                       WHAT (2)
                                  WHAT (3)
                                                       WHAT (5)
                                                                 WHAT(6)
                                                                          SDUM
```

Incorrect: decimal point is missing



WHAT(2) would be interpreted as 1000!

Incorrect: exponential number not correctly alligned

```
*...+....1....+....2....+....3....+....4....+....5....+....6....+....7....+....

BEAM 1.E+04 0 0 0 0 0 PROTON
```

WHAT(1) might be interpreted as 1.E+4000!

### Fixed vs free format

#### Free format:

- Free format can be made *locally* available using option FREE (without any parameter), until the option FIXED restores the fixed format; the opposite can be done either
- Option GLOBAL provides free format also for the geometry input
- In free format input, the different fields are separated by blanks and/or separators (usually commas). <u>All fields must be present</u> or at least represented by two successive separators
- Character fields (command name, SDUM) must be input without quotes

```
*...+....1....+....2....+....3....+....4....+....5....+....6....+....7....+....

BEAM , 1.234567890E+04 , 0.0 , 0.0 , 0.0 , 0.0 , 0.0 , PROTON

*

*keyword momentum mom.spread diverg. X-width Y-width weight particle

* WHAT(1) WHAT(2) WHAT(3) WHAT(4) WHAT(5) WHAT(6) SDUM
```

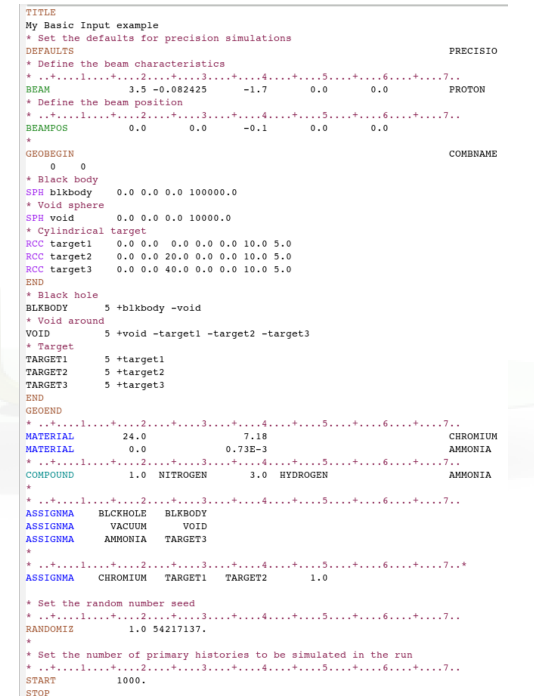
Temporarily switching to FREE format is particularly helpful when more than 10 digits are required for precision reasons !!!

### Names instead of numbers

• FLUKA also allows the use of keywords (names) - 8 characters maximum length - instead of numbers inside FLUKA commands.

- Examples later (for instance materials, or geometrical region, can be inserted using their name instead of numbers)
- This helps the user, improving the readability of the input FLUKA file
- Internally, FLUKA works always by NUMBERS, and keeps name-to-number bidirectional tables/functions

# A basic input



# A basic input card by card

### Prepare the working space

Remember, don't run inside the \$FLUPRO directory.

Instead, go to your home directory and create a subdirectory named example\_basicinput:

mkdir example\_basicinput cd example\_basicinput

From the USB pen drive copy (cp) the source example file example\_basicinput.inp.

Open the FLUKA input file with your preferred editor program (emacs, vim, kwrite, gedit, ...).

A basic input, step 1: Physics settings

```
My Basic Input example
* Set the defaults for precision simulations
DEFAULTS
                                                                   PRECISIO
* ..+...1....+...2....+...3....+....4....+....5....+....6....+....7..
                3.5 -0.082425
                                   -1.7
* Define the beam position
* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..
                         0.0
                                   -0.1
                                                       0.0
GEOBEGIN
                                                                   COMBNAME
   0 0
* Black body
SPH blkbody
              0.0 0.0 0.0 100000.0
* Void sphere
SPH void
              0.0 0.0 0.0 10000.0
* Cylindrical target
RCC target1
              0.0 0.0 0.0 0.0 0.0 10.0 5.0
              0.0 0.0 20.0 0.0 0.0 10.0 5.0
RCC target3
              0.0 0.0 40.0 0.0 0.0 10.0 5.0
* Black hole
BLKBODY
            5 +blkbody -void
* Void around
VOID
            5 +void -target1 -target2 -target3
* Target
TARGET1
            5 +target1
TARGET2
            5 +target2
TARGET3
            5 +target3
END
GEOEND
* ..+...1....+...2...+...3...+...4....+...5...+...6...+...7..
MATERIAL
               24.0
                                  7.18
                                                                   CHROMIUM
MATERIAL
                0.0
                                0.73E-3
                                                                   AMMONIA
* ..+...1....+....2....+....3....+....4....+....5....+....6....+....7..
COMPOUND
                1.0 NITROGEN
                                   3.0 HYDROGEN
* ..+...1....+...2....+...3....+...4....+...5....+...6...+...7...
           BLCKHOLE
                      BLKBODY
ASSIGNMA
             VACUUM
ASSIGNMA
            AMMONIA TARGET3
* ..+...1....+....2....+....3....+....4....+....5....+....6....+....7...*
ASSIGNMA
           CHROMIUM TARGET1 TARGET2
* ..+...1....+...2....+...3....+....4....+....5....+....6....+....7..
RANDOMIZ
                1.0 54217137.
* Set the number of primary histories to be simulated in the run
* ..+...1....+....2....+....3....+....4....+....5....+....6....+....7...
```

STOP

### Physics settings: **DEFAULTS**

Select predefined physics settings (e.g. transport thresholds) for a specific kind of simulation:

#### SDUM =

**CALORIMEtry**: calorimeter simulations

**EET/TRANsmut**: Energy Transformer or transmutation calculations

**EM-CASCAde**: pure EM cascades

**HADROTHErapy**: hadrotherapy calculations

**ICARUS**: studies related to the ICARUS experiment

**NEUTRONS**: pure low-energy neutron runs

**NEW-DEFAults**: minimal set of generic defaults – set by default

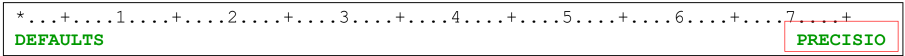
**PRECISIOn**: precision simulations (**recommended**)

**SHIELDINg**: hadron shielding calculations without gammas

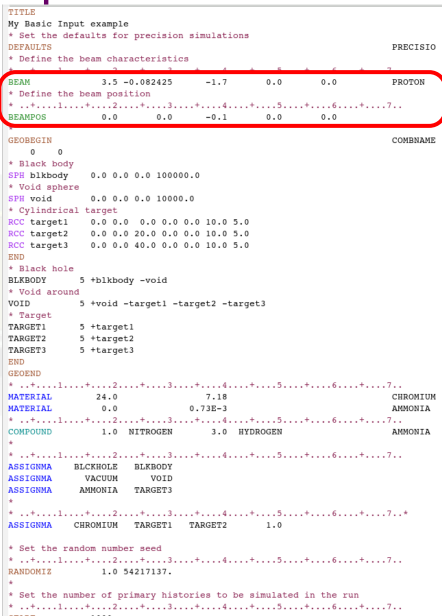
### Physics settings: **DEFAULTS**: **PRECISIO**

- EM transport on (EMF on), production/transport thresholds should always be set by the EMFCUT!
- Inelastic form factor correction to Compton scattering on (EMFRAY on)
- Detailed photoelectric edge treatment and fluorescence photons activated
- Low energy neutron transport on (LOW-NEUT on), threshold 20 MeV, with fully analogue absorption
- All transport threshold = 100keV, but neutrons (10<sup>-5</sup> eV) and neutrinos (0, but they are discarded)
- Multiple Scattering threshold at minimum allowed energy, for both primary and secondary charged particles
- Delta rays production on, threshold 100keV (DELTARAY)
- Restricted ionization energy loss fluctuations for all particles (IONFLUCT)
- Tabulation ratio for hadron/muon dp/dx set at 1.04, fraction of the kinetic energy to be lost in a step set at 0.05, number of dp/dx tabulation points set at 80 (DELTARAY, EMFFIX, FLUKAFIX)
- e+e- pair production and bremsstrahlung by heavy particles on (PAIRBREM)

  Pair threshold =  $2 \text{ m}_e$ , bremsstrahlung threshold = 300 keV
- Muon photonuclear interactions on (MUPHOTON)



# A basic input, step 2: Beam



STOP

# A basic input, step 2: Beam

The card BEAM defines the particle type and energy (or momentum).

The card BEAMPOS controls particle starting position and direction.

For complex particle sources (complex distributions in energy, space and direction) a special user routine, source.f, can be used\*. The card SOURCE has to be added in the input file.

<sup>\*</sup>Some pre-defined cases (volume sources, cosmic ray source, uniform isotropic source etc are already built-in and available via data cards, see the manual)

### Example: **BEAM**

#### Define beam characteristics:

type of particle, energy, divergence, spatial profile....

- [WHAT(1)] = 3.5 GeV/c (average beam momentum)
- [WHAT(2)] = -0.082425 GeV/c FWHM (Gaussian momentum distribution,)
- [WHAT(3)] = -1.7 mrad FWHM (Gaussian angular distribution,)
- [WHAT(4)] = 0.0 (X width)  $\rightarrow$  point-like source
- [WHAT(5)] = 0.0 (X width)  $\rightarrow$  point-like source
- **■** [WHAT(6)] = ignored
- [SDUM] = proton (particle beam)

```
BEAM 3.5 -0.082425 -1.7 0.0 0.0 PROTON
* Define the beam position
* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..
BEAMPOS 0.0 0.0 -0.1 0.0 0.0
```

### Example: **BEAMPOS**

Define beam position and direction:

- [WHAT(1)] = 0.0 cm (x coordinate of the spot center)
- [WHAT(2)] = 0.0 cm (y coordinate of the spot center)
- [WHAT(3)] = -0.1 cm (z coordinate of the spot center)
- [WHAT(4)] = 0.0 (direction cosine of the beam wrt the x-axis)
- [WHAT(5)] = 0.0 (direction cosine of the beam wrt the y-axis)
- [SDUM] = blank (therefore beam towards positive Z)

```
BEAM 3.5 -0.082425 -1.7 0.0 0.0 PROTON
* Define the beam position
* .+...1...+...2...+...3...+...4...+...5...+...6...+...7..
BEAMPOS 0.0 0.0 -0.1 0.0 0.0
```

A basic input, step 3: Geometry

```
My Basic Input example
* Set the defaults for precision simulations
                                                                     PRECISIO
* Define the beam characteristics
* ..+...1....+...2....+...3....+....4....+....5....+....6....+....7...
                3.5 -0.082425
                                   -1.7
* Define the beam position
* ..+...1....+...2....+...3...+...4....+...5...+...6...+....7..
                          0.0
                                   -0.1
GEOBEGIN
                                                                     COMBNAME
    0 0
* Black body
SPH blkbody
               0.0 0.0 0.0 100000.0
* Void sphere
               0.0 0.0 0.0 10000.0
* Cylindrical target
               0.0 0.0 0.0 0.0 0.0 10.0 5.0
              0.0 0.0 20.0 0.0 0.0 10.0 5.0
RCC target3
              0.0 0.0 40.0 0.0 0.0 10.0 5.0
* Black hole
BLKBODY
             5 +blkbody -void
* Void around
VOID
             5 +void -target1 -target2 -target3
* Target
TARGET1
             5 +target1
TARGET2
             5 +target2
TARGET3
             5 +target3
MATERIAL
MATERIAL
                 0.0
                                 0.73E-3
                                                                     AMMONIA
* ..+...1....+....2....+....3....+....4....+....5....+....6....+....7...
```

Three targets: target1 target2 target3

Geometry definition: please, wait for the Geometry lecture on Tuesday morning, but notice the structure

```
* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..*

ASSIGNMA CHROMIUM TARGET1 TARGET2 1.0

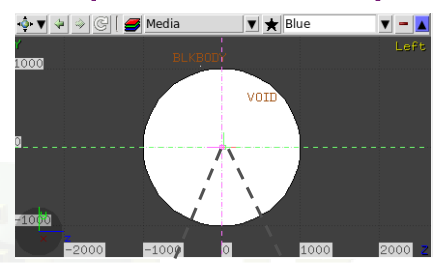
* Set the random number seed
* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..

RANDOMIZ 1.0 54217137.

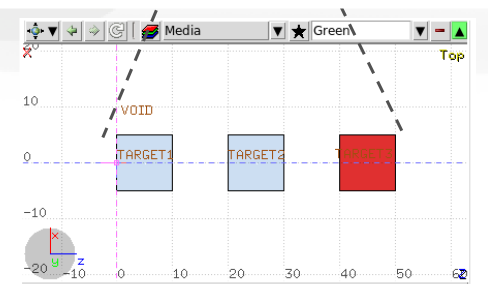
* Set the number of primary histories to be simulated in the run
* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..

START 1000.
```

# A basic input, step 3: Geometry



The whole geometry must be surrounded by a region of "blackhole" limited by a closed body.



# A basic input, step 3: Geometry

The **Combinatorial Geometry** in FLUKA must be preceded by a GEOBEGIN card and followed by a GEOEND card.

For details on the **Combinatorial Geometry** (bodies, regions and optional region volumes) please wait for the Geometry lecture on Tuesday morning.

A basic input, step 4: Materials

STOP

```
My Basic Input example
* Set the defaults for precision simulations
                                                                    PRECISIO
* Define the beam characteristics
* ..+...1....+....2....+....3....+....4....+....5....+....6....+....7...
                                   -1.7
                3.5 -0.082425
* Define the beam position
* ..+...1....+....2....+....3....+....4....+....5....+....6....+....7..
                          0.0
                                   -0.1
                                              0.0
GEOBEGIN
                                                                    COMBNAME
    0 0
* Black body
SPH blkbody
              0.0 0.0 0.0 100000.0
* Void sphere
SPH void
              0.0 0.0 0.0 10000.0
* Cylindrical target
RCC target1
              0.0 0.0 0.0 0.0 0.0 10.0 5.0
RCC target2
              0.0 0.0 20.0 0.0 0.0 10.0 5.0
RCC target3
              0.0 0.0 40.0 0.0 0.0 10.0 5.0
* Black hole
BLKBODY
            5 +blkbody -void
* Void around
VOID
            5 +void -target1 -target2 -target3
* Target
            5 +target1
TARGET1
TARGET2
            5 +target2
TARGET3
            5 +target3
END
                                   7.18
                                                                    CHROMIUM
MATERIAL
                                0.73E-3
                0.0
                                                                    AMMONIA
* ..+...1....+...2....+....3....+....4....+....5....+....6....+....7...
COMPOUND
                1.0 NITROGEN
                                    3.0 HYDROGEN
                                                                    AMMONIA
* ..+...1....+...2....+...3....+...4....+....5....+....6....+....7..
ASSIGNMA
              VACUUM
ASSIGNMA
            AMMONIA
                      TARGET3
* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..*
           CHROMIUM TARGET1 TARGET2
* Set the random number seed
* ..+...1....+...2...+...3...+...4....+...5....+...6...+...7..
RANDOMIZ
                1.0 54217137.
* Set the number of primary histories to be simulated in the run
* ..+...1....+...2....+...3....+....4....+....5....+....6....+....7...
```

### **Materials**

#### FLUKA handles:

- elemental materials (by default natural composition, the user can set a specific isotope, being aware of low energy neutron cross sections availability)
- compounds (chemical molecules, alloys, mixtures...)

Each material is uniquely identified by an index/name. FLUKA has a set of predefined materials (see FLUKA manual). Users can both use/modify these and define their own ones.

#### Basic cards:

**MATERIAL** material declaration

**COMPOUND** compound definition (a **MATERIAL** card is mandatory for a compound

declaration)

**ASSIGNMA** material assignment to regions of geometry

### **Predefined materials**

In FLUKA 2 special materials + 23 natural elements of most common use are predefined, e.g. Carbon, Oxygen, Iron... (check them out in the manual, Chap. 5).

The first two are very important:

- **BLCKHOLE** (mat #1): material with infinite absorbance;
- VACUUM (mat #2)

12 compound materials with the composition suggested by ICRU are predefined as well, e.g. water, PMMA... (see the manual!)

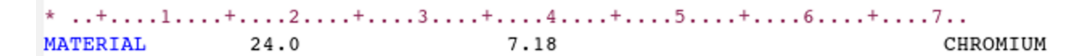
All predefined materials can be used WITHOUT explicit MATERIAL / COMPOUND cards

WARNING: user defined **MATERIAL** cards **OVERRIDE PREDEFINED** materials having the same name

### Example: MATERIAL

Defines a new material or override a previous one

- [SDUM] = CHROMIUM (material name)
- [WHAT(1)] = 24.0 (atomic number Z)
- [WHAT(2)] = leave it empty (atomic weight)
- $[WHAT(3)] = 7.18 \text{ g/cm}^3 \text{ (density)}$
- [WHAT(4)] = leave it empty (material number)
- [WHAT(5)] = normally empty (alternate material to be used for dE/dx)
- [WHAT(6)] = leave it empty unless you want a specific isotope (mass number A)



### Example: **COMPOUND**

Defines a new compound

Each COMPOUND card must be associated to a MATERIAL card More COMPOUND card can be used to define a compound

- [SDUM] = compound name
- [WHAT(1)] = amount of the first component
- [WHAT(2)] = first component material
- [WHAT(3)] = amount of the second component
- [WHAT(4)] = second component material
- [WHAT(5)] = amount of the third component
- [WHAT(6)] = third component material



content > 0	component material > 0
content < 0	component material > 0
content < 0	component material < 0

ATOM content
MASS content
VOLUME content

Names can be preceded by a minus sign!

### Example: compound

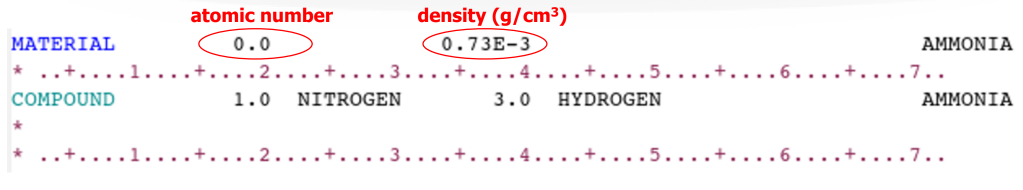
Defines a new compound

Each compound card must be associated to a MATERIAL card.

More COMPOUND card can be used to define a compound.

- [SDUM] compound name
- [WHAT(1)] amount of the first component
- [WHAT(2)] first component material
- [WHAT(3)] amount of the second component
- [WHAT(4)] second component material
- [WHAT(5)] amount of the third component
- [WHAT(6)] third component material

 $AMMONIA = NH_3$ 



### Example: ASSIGNMA

Assign a material to one (or more) region in the geometry

(for the region definition see the geometry lecture or the manual)

A material must be associated to each of the geometry regions, except to those defined as blackhole.

The assigned material could be either a single element material or a compound

- [WHAT(1)] = material index, or material name
- [WHAT(2)] = first region to be "filled" with the material (Default = 2.0)
- [WHAT(3)] = last region to be "filled" with the material (Default = WHAT(2))
- [WHAT(4)] = step length in assigning indices
- [WHAT(5)] = to activate magnetic and electric fields (see manual)
- [WHAT(6)] = assign another material for radioactive decay products transport. As of now, only vacuum and blackhole are allowed.

```
ASSIGNMA
          BLCKHOLE
                    BLKBODY
ASSIGNMA
            VACUUM
                      VOID
ASSIGNMA
           AMMONIA
                    TARGET3
* ..+....5....+....6....+....7...*
ASSIGNMA
                    TARGET1
          CHROMIUM
                             TARGET2
                           FLUKA Beginners' course
                                                             31
```

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### Materials special cards

#### Input card MAT-PROP

Allows to provide extra information about materials

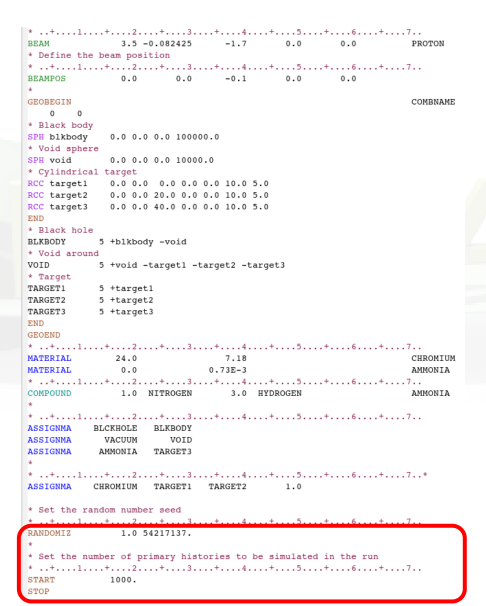
(e.g.: gas pressure, effective density, average ionization potential)

#### Input card CORRFACT

Allows to change material density for dE/dx and nuclear processes on a region-by-region basis

(used in connection with voxel geometries derived from a CT scan) See lecture on FLUKA Medical applications

# A basic input, step 5: Random Seed, START, and STOP



### Example: RANDOMIZ

The random number generator is initialized to read a vector of 97 seeds from an external file. Different numbers input will initialize different and independent random number sequences.

```
[WHAT(1)]: logical file unit from which to read the seeds. Must be 1.0!!

[WHAT(2)]: any number < 9.E8, initialization of the random seed sequences.

Different WHAT(2) lead to different sequences allowing to run parallel jobs.

[WHAT(3-6), SDUM]: not used
```

```
* Set the random number seed

* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..

RANDOMIZ 1.0 54217137.
```

### Example: START AND STOP

A START card at the end of the input file is mandatory. It defines the number of particle histories required.

The START card is optionally followed by a STOP card, which stops the execution of the program.

```
[WHAT(1)] = maximum number of primary histories simulated in the run [WHAT(2)] = not used [WHAT(3-6), SDUM] = see manual
```

```
* Set the number of primary histories to be simulated in the run

* ..+...1...+...2...+...3...+...4...+...5...+...6...+...7..

START 1000.

STOP
```

### Exercise: Basic Input

Run example\_basicinput.inp. In the terminal type: \$FLUPRO/flutil/rfluka -N0 -M4 example\_basicinput

Look at the .out file with less ex1001.out or any text editor e.g. emacs, vi

(FLUKA mode available for emacs and vi on the web page http://www.fluka.org/fluka.php?id=tools&mm2=5)

# The FLUKA Output Files

### The FLUKA output consists of:

- A main (standard) output, written on logical output unit LUNOUT (predefined as 11 by default) [\*.out]
- A file with the last random number seeds, unit LUNRAN (2 by default) [ran\*]
- A file of error messages, unit LUNERR (15 by default) [.err]
- Any number (including zero) of estimator output files. Their logical unit number is defined by the user [\*\_fort.xx] (see scoring lecture)
- The available range of logical output numbers is: 21-99
- Generally, the user can choose between formatted (ASCII) and unformatted (binary) scoring (negative or positive sign in the logical unit number). Unformatted scoring is mandatory for the use of provided post-processing utilities.
- Several estimators can be output on the same file (same logical unit) provided they are of the same type
- Possible additional output generated by the user in any user routine

### **FLUKA Preprocessor**

- FLUKA supports preprocessing instructions like those used in C or C++
- This useful feature allows to keep different setups and configurations in a single input file, selecting the desired one when starting a run

#### Definition of constants:

```
#define VARIABLE1
or #define VARIABLE1 Value
#undef VARIABLE1
```

One can refer to VARIABLE1 inside the input file (geometry included) using \$VARIABLE1

### Example:

```
#define Ekbeam 100.0
#define Beampart PROTON
...
*...+....1....+...2....+...3...+...4...+...5...+...6...+...7....+...8
BEAM -$Ekbeam $BeamPart
```

### **FLUKA Preprocessor**

- FLUKA supports preprocessing instructions like those used in C or C++
- This useful feature allows to keep different setups and configurations in a single input file, selecting the desired one when starting a run
- The #include directive can ease the handling of large input files

#### Definition of constants:

#define VARIABLE1
or #define VARIABLE1 Value
#undef VARIABLE1

One can refer to VARIABLE1 inside the input file (geometry included) using \$[VARIABLE1]

#### Conditional directives

#if VARIABLE1
#elif VARIABLE2
#else

#endif

Up to 10 nested levels of conditional statements (#if/#else/#endif) are supported

#### Include directive:

# FLUKA Preprocessor example

```
#define DUMP COPPER
*#define DUMP CARBON
#if DUMP COPPER
* Select copper as material for the dump
*...+....4....+....5....+....6
ASSIGNMA
             COPPER BEAMDUMP
#elif DUMP CARBON
* Select carbon as material for the dump
ASSTGNMA
            CARBON BEAMDUMP
#else
* Use default material for the dump
ASSIGNMA
              IRON BEAMDUMP
#endif
```

# Depending on the active define (DUMP\_COPPER or DUMP\_CARBON) different ASSIGNMA card are used

```
#define can be either commented out or undefined
e.g.: #undef DUMP COPPER
```

### Extra Exercise: Basic Input

### Extra:

- Change the type of primary particles from proton to neutrons
- $\square$  Move the beam position in (0.0, 0.0, -10.0) cm
- □ Change the material CHROMIUM in WATER with density equal to 0.99669 g/cm<sup>3</sup>
- Use the card MAT-PROP to set the ionization potential for WATER equal to 78.0 eV
- □ Change the number of primary particles from 1000. to 10000.

Run your input file and see how the .out file has changed.



# Beyond a basic input

# Special sources: 3D distributions

**BEAMPOS** card allows the user to define some 3D *spatial* distributions of source particles

SDUM = SPHE-VOL:

defines a spatially extended source in a spherical shell

SDUM = CART-VOL:

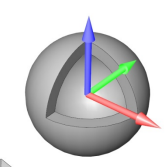
defines a spatially extended source in a Cartesian shell with the sides parallel to the beam frame axes

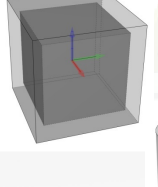
#### SDUM = CYLI-VOL:

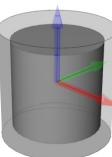
defines a spatially extended source in a cylindrical shell with the height parallel to the z-axis of the beam frame

#### SDUM = FLOOD:

defines a source distribution on a spherical surface, such as to produce a uniform and isotropic fluence within the sphere FLUKA Beginners' course







## Special sources: SPECSOUR

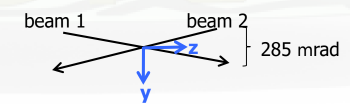
FLUKA allows the definitions of special sources for:

- two colliding beams
- galactic cosmic rays
- solar particles events

Various SDUM allow plenty of opportunities:

```
PPSOURCE, CROSSASY, CROSSSYM; GCR-IONF, GCR-SPEC, GCR-ALLF; SPE-SPEC, SPE-2003, SPE-2005
```

*Example:* LHC proton-proton collision 7 TeV/c, full crossing angle of 285 mrad in yz-plane



For all these special cases of source distributions, please, refer to the FLUKA manual!

# More Physics settings

### Input card PHYSICS

Allows to override standard FLUKA defaults for some processes:

- activates coalescence (critical for calculation of residual nuclei)
- activates the new fragmentation model ("evaporation" of fragments up to A=24, critical for calculation of residual nuclei)
- activates PEANUT above 5 GeV
- activates electromagnetic dissociation of heavy ions
- activates charmed particle transport

#### Input card PHOTONUC

- activates photo-nuclear interactions
- activates muon pair production by photons

The use of the LAM-BIAS card (see Biasing lecture) coupled to the **PHOTONUC** card is recommended, in order to artificially *increase* the probability to have photonuclear reactions

### Transport thresholds

#### Input card PART-THR

- Defines transport cut-offs for hadrons, muons, and neutrinos
- Setting done by particle type, overriding the selected **DEFAULTS**
- For **neutrons**, a <20.0 MeV cut-off is internally translated into the corresponding group energy; On a region basis, the neutron cut-off can be *increased* by the **LOW-BIAS** card (see Neutrons lecture)
- Charged particles (but electrons) are not stopped, but ranged out to rest in an approximate way (if the threshold is < 100 MeV)</li>

#### Input card **EMFCUT**

For electron, positron, and photon, sets:

- Energy thresholds for production in the selected materials
- Transport cut-offs in the selected regions.
- Use STRONGLY recommended

#### Input card DELTARAY

- Activates delta ray production by muons and charged hadrons
- Sets energy threshold for their production

# Low energy neutrons (E < 20.0 MeV)

FLUKA transports neutrons with energies lower than 20 MeV by means of a multi-group algorithm, based on 260 groups (See Neutrons lecture)

#### Input card LOW-NEUT

- Activates low-energy neutron transport (by default off only in EM-CASCA)
- Requests point-wise cross sections

   (only available for a few elements, see manual)

### Input card LOW-MAT

- Sets the correspondence between FLUKA materials and low-energy neutron cross-sections
- By default, the link is between the FLUKA material and the first material of the same name present in the library. Therefore, the option is not needed in many cases

## Heavy ions: beams and transport

#### Input card HI-PROPE

• When **BEAM**'s SDUM is **ISOTOPE**...

...specifies the isotope of a radioactive source

...requires a RADDECAY card

When BEAM'S SDUM is HEAVYION...

... specifies the properties of an ion beam:

in the **BEAM** card, the beam energy is given in GeV/nmu

(**n**uclear **m**ass **u**nit, i.e. 1/12 of the <sup>12</sup>C nucleus mass)

<sup>2</sup>H, <sup>3</sup>He, and <sup>4</sup>He beams have dedicated SDUM in the **BEAM** card

## Heavy ions: beams and transport

#### Input card IONTRANS

- Is not required when using an heavy ion beam HEAVYION
- Activates the ions transport
- Allows to limit it to a subset of light ions (A < 5)</li>
- Switches between approximate and full transport
   ...(including nuclear interactions)
- Nucleus-nucleus interactions above 125 MeV/n
  - ...can be performed only if the event generators
    - DPMJET and RQMD are linked to the FLUKA executable
- Below 125 MeV/n...
  - ...the BME event generator is already linked in the standard executable

## Induced radioactivity

#### Input card RADDECAY

- Activates the simulation of the decay of generated radioactive nuclides
- Allows to set biasing for radioactive decay products

### Input card IRRPROFI

Defines an irradiation profile (i.e. irradiation time and intensity)

### Input card DCYTIMES

Defines the decay (cooling) time

#### Input card DCYSCORE

 Associates scoring detectors (radio-nuclides, fluence, dose) with different cooling times

