



# FLUKA Scoring

FLUKA Advanced Course

# Contents

- *fluence*
- the Monte Carlo way
- FLUKA built-in scoring
- user routines [in the dedicated lecture]
- (statistical) errors

# Reaction Rate and Cross Section [1/3]

- We call **mean free path**  $\lambda[cm]$  the average distance travelled by a particle in a material before an interaction. Its inverse,  $\Sigma [cm^{-1}]$  is the probability of interaction per unit distance, and is called **macroscopic cross section**. Both  $\lambda$  and  $\Sigma$  depend on the material and on the particle type and energy.
- For  $N$  identical particles, the number of reactions  $R$  occurring in a given time interval will be equal to the total distance travelled  $Nl$  times the probability per unit distance  $\Sigma$ :  $R = Nl\Sigma$
- The **reaction rate** will be  $\dot{R} = Nd l / dt \Sigma = Nv\Sigma$ , where  $v$  is the average particle velocity.

# Reaction Rate and Cross Section [2/3]

- Assume now that  $n(\mathbf{r}, v) = dN/dV$  [ $cm^{-3}$ ] be the density of particles with velocity  $v = dl/dt$  [ $cm/s$ ], at a spatial position  $\mathbf{r}$ . The reaction rate inside the volume element  $dV$  will be:  $d\dot{R}/dV = n(\mathbf{r}, v)v\Sigma$
- The quantity  $\dot{\Phi}(\mathbf{r}, v) = n(\mathbf{r}, v)v$  is called **fluence rate** or **flux density** and has dimensions [ $cm^{-3} cm s^{-1}$ ] = [ $cm^{-2} s^{-1}$ ].
- The time integral of the flux density  $\Phi(\mathbf{r}, v) = n(\mathbf{r}, v)dl$  is the **fluence** [ $cm^{-2}$ ]
- Fluence is measured in **particles per  $cm^2$**  but in reality it describes the **density of particle tracks**
- The number of reactions inside a volume  $V$  is given by the formula:  $R = \Sigma\Phi V$  (where the product  $\Sigma\Phi$  is integrated over energy or velocity)

# Reaction Rate and Cross Section [3/3]

- Dividing the macroscopic cross section by  $N_0$ , the number of atoms per unit volume, one obtains the **microscopic cross section**  $\sigma$  [*barn* =  $10^{-24} \text{cm}^2$ ].

$$\frac{\text{probability/cm}}{\text{atoms/cm}^3} = \frac{\text{probability} \times \text{cm}^2}{\text{atoms}} = \text{atom effective area}$$

i.e., the **area of an atom weighted with the probability of interaction** (hence the name “cross section”).

- But it can also be understood as the **probability of interaction per unit length, with the length measured in atoms/cm<sup>2</sup>** (the number of atoms contained in a cylinder with a 1 cm<sup>2</sup> base).
- In this way, both microscopic and macroscopic cross section are shown to have a similar physical meaning of “probability of interaction per unit length”, with length measured in different units. Thus, the number of interactions can be obtained from both, by multiplying them by the corresponding particle track-length.

# Fluence estimation [1/2]

- Track length estimation:

USRTRACK

$$\dot{\Phi}(v) dt = n(v) v dt = \frac{dN(v)}{dV} \frac{dl(v)}{dt} dt = \lim_{\Delta V \rightarrow 0} \frac{\sum_i l_i(v)}{\Delta V}$$

- Collision density estimation (NOT IN VACUUM!):

USRCOLL

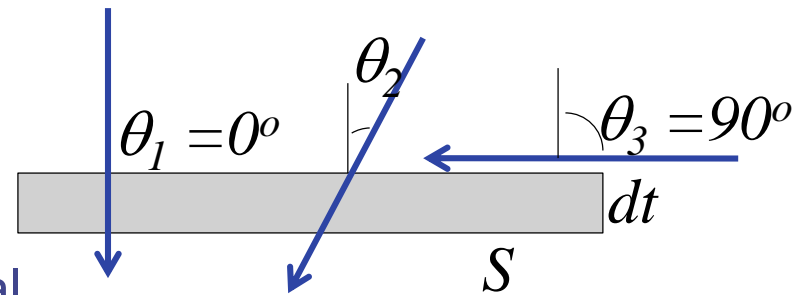
$$\dot{\Phi}(v) = \frac{d\dot{R}(v)}{dV} \lambda(v)$$

*0 x ∞*

# Fluence estimation [2/2]

## Surface crossing estimation

- Imagine a surface having an infinitesimal thickness  $dt$ . A particle incident with an angle  $\theta$  with respect to the normal of the surface  $S$  will travel a segment  $dt/\cos\theta$ .



- Therefore, we can calculate an average surface fluence by adding  $dt/\cos\theta$  for each particle crossing the surface, and dividing by the volume  $S dt$

$$\Phi = \lim_{dt \rightarrow 0} \frac{\sum_i \frac{dt}{\cos \theta_i}}{S dt}$$

- While the **current**  $J$  counts the number of particles crossing the surface divided by the surface:

$$J = dN/dS$$

The **fluence** is independent of the orientation of the **surface**  $S$ , while the **current** is NOT!

In an *isotropic field* it can be easily seen that for a flat surface  $J = \Phi/2$

# Complexity

- Simple case: a mono-directional (zero divergence) monochromatic beam attenuated by a uniform shielding layer of thickness  $T$
- The source term: a *fluence*  $\Psi(E, \theta, r) = dN/d\Omega/dE/dS = C \delta(E-E_0) \delta(\theta-\theta_0)$
- Suppose that the particles are absorbed according to a macroscopic absorption cross section  $\Sigma_t =$  interaction probability per cm  $= \sigma_t N_{AP}/A$
- In an infinitesimal thickness  $dt$  the probability to be absorbed is  $P = \Sigma dt$
- $d\Psi(t) = -\Psi(t)P = -\Psi(t) \Sigma_t dt$ , a differential equation that is solved by the exponential function  $\rightarrow$
- $\Psi(T) = \Psi(0) \exp(-\Sigma_t T)$  **EASY !!**
- Now ... suppose that the particles are not absorbed, but scattered according to some energy-dependent distribution  $d\sigma(E', \theta')/d\Omega'$ , exiting with  $E' = g(E, \theta')$  where  $g$  is defined by the kinematics.
- Now ... suppose that the original beam was not monochromatic, or that particles can also be produced or that the geometry is not uniform, and the dimensions of the integral will explode.



# MC mathematical foundation

The Central Limit Theorem is the mathematical foundation of the Monte Carlo method. In words:

Given any observable  $A$ , that can be expressed as the result of a convolution of random processes, the average value of  $A$  can be obtained by sampling many values of  $A$  according to the probability distributions of the random processes.

MC is indeed **an integration method** that allows to solve multi-dimensional integrals by sampling from a suitable stochastic distribution.

The accuracy of MC estimator depends on the number of samples:

$$\sigma \propto \frac{1}{\sqrt{N}}$$

# Integration efficiency

- Traditional numerical integration methods (e.g., Simpson) converge to the true value as  $N^{-1/n}$ , where  $N$  = number of "points" (intervals) and  $n$  = number of dimensions
- Monte Carlo converges as  $N^{-1/2}$ , independent of the number of dimensions
- Therefore:
  - ❑  $n = 1 \rightarrow$  MC is not convenient
  - ❑  $n = 2 \rightarrow$  MC is about equivalent to traditional methods
  - ❑  $n > 2 \rightarrow$  MC converges faster (and the more so the greater the dimensions)
- the dimensions are those of the largest number of "collisions" per history
- Note that the term "collision" comes from low-energy neutron/photon transport theory. Here it should be understood in the extended meaning of "interaction where the particle changes its direction and/or energy, or produces new particles"

# Particle transport

- Particle transport is a typical physical process described by probabilities (cross sections = interaction probabilities per unit distance)
- Therefore it lends itself naturally to be simulated by Monte Carlo
- Many applications, especially in **high energy physics and medicine**, are *based on simulations* where the history of each particle (trajectory, interactions) is reproduced in detail
- However in other types of application, typically **shielding design**, the user is interested only in the **expectation values** of some quantities (**fluence** and **dose**) at some space point or region, which are *calculated as solutions of a mathematical equation*
- This equation (the **Boltzmann equation**), describes the **statistical distribution of particles in phase space** and therefore does indeed represent a physical stochastic process
- But in order to estimate the desired expectation values it is not necessary that the Monte Carlo process be identical to it

# Integration without simulation

- In many cases, it is more efficient to replace the actual process by a different one resulting in the same average values but built by sampling from modified distributions
- Such a *biased process*, if based on mathematically correct variance reduction techniques, converges to the same expectation values as the unbiased one
- But it cannot provide information about the higher moments of statistical distributions (fluctuations and correlations)
- In addition, the faster convergence in some user-privileged regions of phase space is compensated by a slower convergence elsewhere

# Analog Monte Carlo

In an **analog** Monte Carlo calculation, not only the mean of the contributions converges to the mean of the actual distribution, but also the variance and **all moments of higher order**:

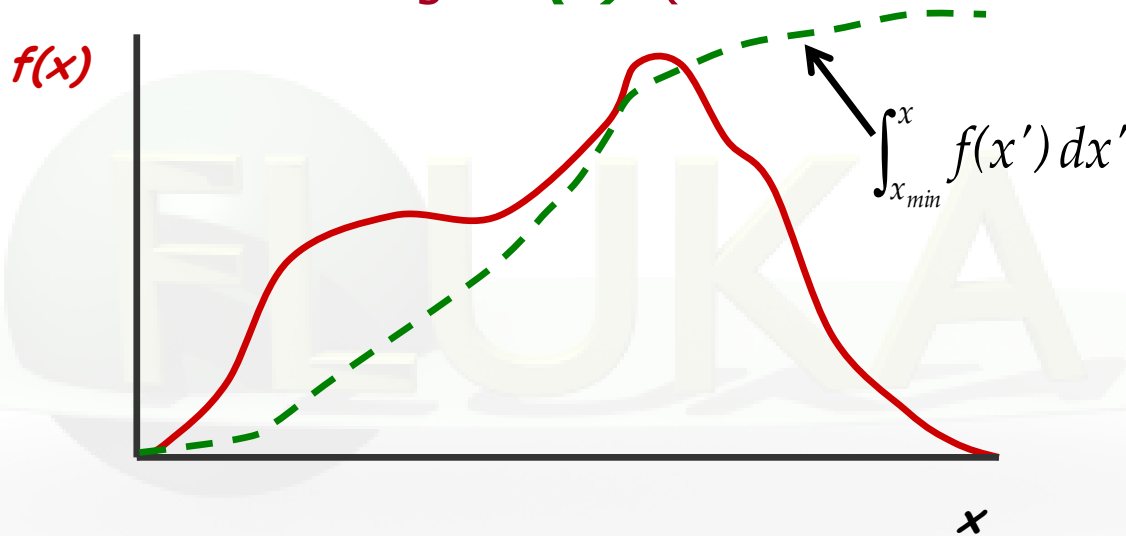
$$\lim_{N \rightarrow \infty} \left[ \frac{\sum_1^N (x - \bar{x})^n}{N} \right]^{\frac{1}{n}} = \sigma_n$$

Then, partial distributions, fluctuations and correlations are all faithfully reproduced: in this case (and in this case only!) we have a real **simulation**

# Random sampling: the key to MC

The central problem of the Monte Carlo method:

Given a Probability Density Function (pdf),  $f(x)$ , generate a sample of  $x$ 's distributed according to  $f(x)$  ( $x$  can be multidimensional)



The use of random sampling techniques is the distinctive feature of Monte Carlo

Solving the integral Boltzmann transport equation by Monte Carlo consists of:

- Geometry and material description of the problem
- Random sampling from probability distributions of the outcome of physical events

# Particle transport Monte Carlo

Application of Monte Carlo to particle transport and interaction:

- Each particle is followed on its path through matter
- At each step the occurrence and outcome of interactions are decided by random selection from the appropriate probability distributions
- All the secondaries issued from the same primary are stored in a "stack" or "bank" and are transported before a new history is started
- The accuracy and reliability of a Monte Carlo depend on the models or data on which the probability distribution functions are based
- Statistical precision of results depends on the number of "histories"
- Statistical convergence can be accelerated by "biasing" techniques.

# Assumptions

- **Static, homogeneous, isotropic, amorphous** media and geometry  
Problems: e.g. moving targets\*, atmosphere [must be represented by discrete layers of uniform density], radioactive decay taking place in a geometry different from that in which the radionuclides were produced\*, crystal channeling\*.  
\* These restrictions have been (\* are being) overcome in FLUKA
- **Markovian process**: the fate of a particle depends **only on its actual present properties**, not on previous events or histories
- **Particles do not interact with each other**  
Problems: e.g. the Chudakov effect (charges cancelling in  $e^+e^-$  pairs)
- **Particles interact with individual electrons / atoms / nuclei / molecules**  
Problems: invalid at low energies (X-ray mirrors)
- **Material properties are not affected by particle reactions**  
Problems: e.g. burnup



# Built-in scoring

- Several **pre-defined estimators** can be activated in FLUKA.
- One usually refers to these estimators as **“scoring”** capabilities
- Users have also the possibility to build their own scoring through user routines (some of which mentioned afterwards), HOWEVER:
  - **Built-in scoring** covers most of the **common needs**
  - **Built-in scoring** has been **extensively tested**
  - **Built-in scoring** takes BIASING **weights automatically into account**
  - **Built-in scoring** has **refined algorithms** for track subdivision
  - **Built-in scoring** comes with **utility programs** that allow to evaluate statistical errors
- Geometry dependent and geometry independent scoring both available
- FLUKA can score **particle fluence (tracklength), current, energy spectra, angular distributions, energy deposition, activity ...**
- Either integrated over the **“run”**, with proper normalization, OR **event-by event**
- Standard scoring can be weighted by means of **simple user routines**

# Scoring cards [1]

see the Beginners' Course

- **SCORE** scores energy deposited (or star density) in each region *[table in the .out file, not automatically merged over cycles]*
- **RESNUCLEi** scores residual nuclei (or their activity) in a given region
- **USRTRACK (USRCOLL)** scores average differential fluence  $d\Phi/dE$  of a given type or family of particles over a given region
- **USRBDX** scores average double differential fluence (or current)  $d^2\Phi/dEd\Omega$  of a given type or family of particles over a given surface
- **USRBIN** scores the spatial distribution of deposited energy density, dose, integrated fluence, star density, *dose equivalent*, net charge, specific activity, ... in a regular mesh (cylindrical or Cartesian) described by the user
- **USRYIELD** scores a double differential yield *[do not ask for cross section, like by default]* of particles escaping from a surface. The distribution can be with respect to energy and angle (wrt the beam direction), but also many other more "exotic" quantities

Remember that *low energy (<20MeV) neutrons* have a pre-defined energy binning

# Warnings [I]

- USRBIN scoring algorithm:

By selecting **WHAT(1)≥10**, *energy deposition, dose, ...* are distributed along the particle track (recommended!)

- \*\*\* Activity/fission/neutron balance binnings cannot be track-length!!!

*Point-wise quantities* have to be scored at a point (select **WHAT(1)<10**)

- Badly defined USRBIN limits

```
***** Fluka stopped in Usrbin: "usr/eventbin" n. 1 *****  
***** with zero width 0.000 for axis R *****
```

- Never use unit numbers smaller than 20 or higher than 99  
<20 reserved by FLUKA >99 FORTRAN limitation

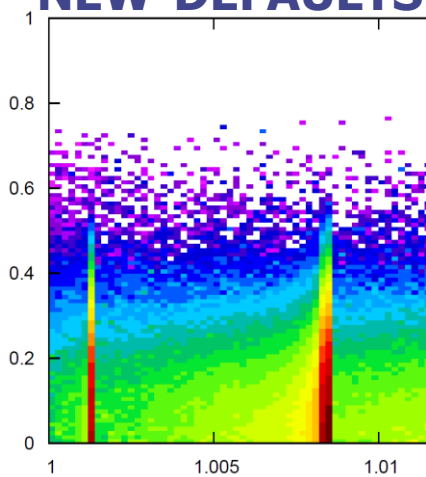
- Never mix the output of different scoring cards in the same unit

- Verify that you didn't merge cycles referring to different input versions  
(change the name of the input file for every new problem!)

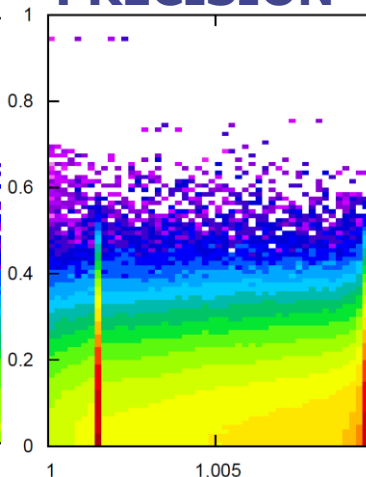
# Example

- Thin window with low-E (5MeV) electron beam
- Energy deposition profile in the window (for radiation damage studies)
- Observation of 'strange peaks'
- **Trying to understand:** lower e<sup>-</sup>-thresholds help
- **Real-Problem:** point-wise scoring requested

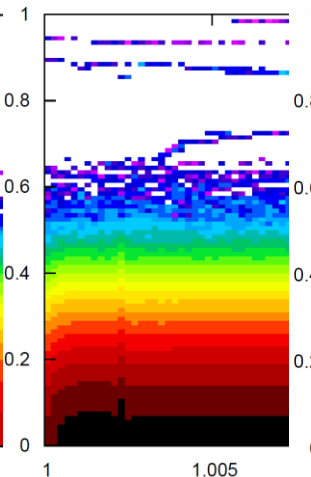
**NEW-DEFAULTS**



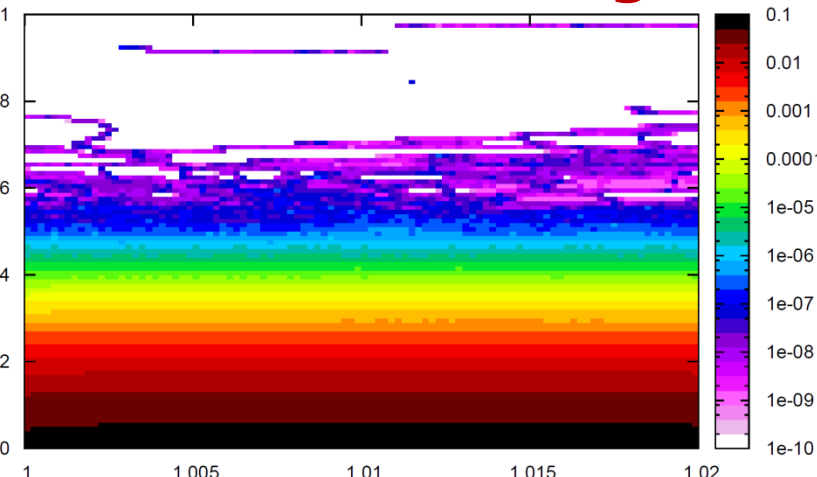
**PRECISION**



**Low e- Thr.**



**CORRECT Scoring**



# Scoring cards [2]

- **EVENTBIN** is like **USRBIN**, but prints the binning output **after each event** instead of an average over histories
- **ROTPRBIN** sets the **storage precision** (single or double) and assigns **rotations/translations** for a given user-defined binning (**USRBIN** or **EVENTBIN**). Quite useful in case of **LATTICE**
- **USERDUMP** defines the events to be written onto a **"collision tape"** file  
Coupled to the **mgdraw** user routine
- **AUXSCORE** defines **filters** and **conversion coefficients**
- **TCQUENCH** sets scoring **time cut-offs** and/or **Birks quenching** parameters for binnings (**USRBIN** or **EVENTBIN**) indicated by the user
- **DETECT** scores **energy deposition in coincidence** or anti-coincidence with a trigger, separately for each "event" (primary history). Dedicated post-processing routine available



# "FILTER" : AUXSCORE

There is the possibility to **filter** the estimators, restricting the scoring to a selected subset of particles.

For instance: USRBIN energy deposition by muons only

USRBIN	11.0	ENERGY	-40.0	10.0	15.0	TargEne
USRBIN	0.0		-5.0	100.0	200.0	&
AUXSCORE	USRBIN	MUONS		TargEne		

ionization (+NIEL) by the selected particle, critically depending on the delta rays threshold!  
[doubtful physical meaning]

Another example: score the yield [vs polar angle and kinetic energy] of 56-Iron ions (there is no separate name for each ion specie, except light ones. HEAVYION scores all isotopes heavier than alpha together!)

USRYIELD	124.0	HEAVYION	-87.	TARGS3	INAIR	1.0Fe56
USRYIELD	180.0	0.0	18.	10.0	0.0	3.0&
AUXSCORE	USRYIELD	-5602600.		Fe56		

The requested ion is coded in  $WHAT(2) = - (100 * \mathbf{Z} + 100000 * \mathbf{A} + \mathbf{m} * 100000000)$  according to its **A**, **Z** and (optionally) isomeric state **m** with 0==all, i.e. -2600 == all Iron isotopes

ACTIVITY!

# Routines associated to FLUKA scoring

- comscw.f      weighting energy deposition and star production
- fluscw.f      weighting fluence, current and yield
- mgdraw.f      *general event interface*
- usrrnc.f      intercepting produced residual nuclei (at the end of their path)
- endscp.f      shifting energy deposition
- fldscp.f      shifting fluence
- musrbr.f      special USRBIN binning (lattice): returns region #
- lusrbl.f      special USRBIN binning (lattice): returns lattice #
- fusrbv.f      special USRBIN binning (lattice): returns zero
  
- mdstck.f      }  
● stuprf.f      } intercepting particle stack  
● stupre.f      }



# Statistical Errors [1]

- Can be calculated for **single histories** (not in FLUKA), or for **batches** of several histories
- Distribution of scoring contributions **by single histories** can be very asymmetric (many histories contribute little or zero)
- Scoring distribution **from batches** tends to Gaussian for  $N \rightarrow \infty$ , **provided  $\sigma^2 \neq \infty$**  (thanks to Central Limit Theorem)
- The standard deviation of an estimator calculated from batches or from single histories is **an estimate of the standard deviation of the actual distribution** ("error of the mean")
- How good is such an estimate depends on the type of estimator and on the particular problem (but it converges to the true value for  $N \rightarrow \infty$ )

# Statistical Errors [2]

- The **variance of the mean** of an estimated quantity  $x$  (e.g., fluence), calculated in  $N$  batches, is:

$$\sigma_{\langle x \rangle}^2 = \frac{1}{N - 1} \left[ \frac{\sum_1^N n_i x_i^2}{n} - \left( \frac{\sum_1^N n_i x_i}{n} \right)^2 \right]$$

mean of squares - square of means  
N - 1

where:

$n_i$  = number of histories in the  $i^{\text{th}}$  batch

$n = \sum n_i$  = total number of histories in the  $N$  batches

$x_i$  = average of  $x$  in the  $i^{\text{th}}$  batch:  $x_i = \sum_{j=1}^{n_i} \frac{x_{ij}}{n_i}$

$x_{ij}$  is the contribution to  $x$  of the  $j^{\text{th}}$  history in the  $i^{\text{th}}$  batch

In the limit  $N = n$ ,  $n_i = 1$ , the formula applies to single history statistics

# Statistical Errors [3]

Practical tips:

- Use always at least 5-10 **batches** of comparable size (it is not at all mandatory that they be of equal size)
- Never forget that **the variance itself is a stochastic variable** subject to fluctuations
- Be careful about the way convergence is achieved: often (particularly with biasing) **apparent good statistics** with few isolated spikes could point to a lack of sampling of the most relevant phase-space part
- Plot **2D and 3D distributions!** Looking at them the eye is the best tool in judging the quality of the result

# Statistical Errors [4]

*from an old version of the MCNP Manual:*

<u>Relative error</u>	<u>Quality of Tally</u>
50 to 100%	Garbage
20 to 50%	Factor of a few
10 to 20	Questionable
< 10%	Generally reliable

- Why does a 30%  $\sigma$  mean an uncertainty of a “factor of a few”?  
Because  $\sigma$  in fact corresponds to the sum (in quadrature) of two uncertainties: one associated to the **fraction of histories which don't give a zero contribution** and the other reflecting the **spread of the non-zero contributions**
- The MCNP guideline is empirically based on experience, not on a mathematical proof. But it has been generally confirmed as working also with other codes
- **Small penetrations and cracks** are very difficult to handle by MC, because the “detector” is too small and too few non-zero contributions can be sampled, even by biasing

# Systematic Errors

- **physics**: different codes are based on different physics models. Some models are better than others. Some models are better in a certain energy range. Model quality is best shown by benchmarks at the **microscopic** level (e.g. thin targets)
- **artifacts**: due to imperfect algorithms, e.g., energy deposited in the middle of a step\*, inaccurate path length correction for multiple scattering\*, missing correction for cross section and  $dE/dx$  change over a step\*, etc. Algorithm quality is best shown by benchmarks at the **macroscopic** level (thick targets, complex geometries)
- **data uncertainty**: results can never be better than allowed by available experimental data!
- **material composition**: not always well known. In particular *concrete/soil* composition (how much water content? Can be critical). *Air* contains humidity and pollutants, has a density variable with pressure
- **beam losses**: most of the time these can only be guessed
- presence of **additional material**, not well defined (cables, supports...)
- **geometries** cannot be reproduced exactly (or would require too much effort)  
*Is it worth doing a very detailed simulation when some parameters are unknown or badly known?*

# Mistakes

- **mis-typing the input:** Flair is good at checking, but the final responsibility is the user's
- **error in user code:** use the built-in features as much as possible!
- **wrong units**
- **wrong normalization:** quite common
- **unfair biasing:** energy/space cuts cannot be avoided, but must be done with much care
- forgetting to check that **gamma production** is available in the low energy neutron library (e.g., Ba cross sections)
- ...

# Biasing Mean Free Paths

## Multiplicity Tuning

## BIASING

- Multiplicity tuning is meant to be to **hadrons** what **LPB** is for electrons and photons.
- A hadronic nuclear interaction at LHC energies can end in **hundreds of secondaries**. Except for the leading particle, many secondaries are of the same type and have **similar energies** and other characteristics
- The user **can tune the average multiplicity** in different regions

## Interaction Length

## LAM-BIAS

- **Mean life / average decay length** of unstable particles can be artificially shortened
- Can **increase generation rate** of decay products without discarding the parent
- For **hadrons** the **mean free path for nuclear inelastic interactions** can be artificially decreased. Useful for very thin targets, and also for photonuclear reactions where the cross section is relatively small