

# Special EM processes and their role in Geant4-stepping (energy-loss processes and multiple scattering)

Mihály Novák  
CERN EP-SFT  
(simulation group)

# 1 The **G4SteppingManager** and its responsibilities

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- 3 Pre-step point: step limitation
  - discrete part of the step limit
  - continuous part of the step limitation
  - special continuous processes: transportation, multiple scattering

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## General stepping: G4SteppingManager

- A G4Track encapsulates both static (G4ParticleDefinition) and dynamic properties (energy, momentum, direction, etc.) of a particle
- Geant4 propagates such tracks: one track at a time, step-by-step:
  - from the beginning: first step with a primary or secondary track
  - till the end: particle left the world, particle kinetic energy became zero, dropped below the (user defined) tracking limit or the particle had a destructive interaction (, the user requested to kill the particle)
  - properties of the track (currently under tracking) are updated after each step
  - secondary tracks are pushed into a track stack at each step
- The G4SteppingManager is responsible to coordinate one step
  - implements a **general stepping** algorithm: independent from the type of the particle and processes associated to the particle
- We will have a closer look at the step limitation part of this stepping by focusing on charged particles with EM interactions: *Ionization, Bremsstrahlung, Multiple Scattering* (Why?)

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*Ionization* and *Bremsstrahlung* are described as *continuous-discrete* interactions:

- secondary electron and gamma production with energy **below** the given production energy threshold  $E^{\text{cut}}$  are **not simulated explicitly**
- instead, these low energy losses described as **continuous** energy losses along the step (i.e. between the pre- and post-step points) based on a mean value
- this mean value is the *restricted stopping power* i.e. mean energy loss in unit step length

$$-\frac{dE}{dX_{\text{rest}}}(E; E^{\text{cut}}, \dots) = \int_0^{E^{\text{cut}}} k \frac{d\sigma}{dk}(E, \dots) dk \quad (1)$$

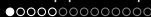
- secondary electron and gamma production with energy **above** the given production energy threshold  $E^{\text{cut}}$  are **simulated explicitly** as **discrete** interactions
- this discrete interaction probability is determined by the *restricted cross section*

$$\sigma_{\text{rest}}(E; E^{\text{cut}}, \dots) = \int_{E^{\text{cut}}}^E \frac{d\sigma}{dk}(E, \dots) dk \quad (2)$$

that covers interactions only in which the secondary particle will have an energy above the production energy threshold

- the discrete part of the interaction will imply a **discrete step limit** i.e. path length till the next discrete interaction, determined by the **restricted cross section**  $\sigma_{\text{rest}}(E; E^{\text{cut}}, \dots)$
- the continuous part of the interaction will also imply a **continuous step limit** due to a maximum allowed "**along step energy loss**" value
- due to this "along step energy loss", the particle **kinetic energy** will be **different at the pre- and post-step points**

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First the interactions, that have discrete part, will propose a step length:

- if the target atom density is  $N$  and the **atomic interaction cross section** of a process is  $\sigma$  (**assumed to be constant along a step at the moment!**)
- according to the Beer-Lambert law, the p.d.f. of the corresponding **interaction length**  $x$  (i.e. the probability that the interaction will happen at  $x, x + dx$  is  $p(x)dx$ )

$$p(x) = N\sigma \exp(-xN\sigma) \implies x \in \mathbb{Exp}(N\sigma) \quad (3)$$

- the mean or expected value of the interaction length  $x$

$$\mathbb{E}(x) = \frac{1}{N\sigma} \equiv \lambda = \frac{1}{\Sigma} \leftarrow x \in \mathbb{Exp}(N\sigma) \quad (4)$$

where  $\lambda$  is the **mean free path** and  $\Sigma = N\sigma = 1/\lambda$  is the **macroscopic cross section**

- if there are  $M$  **independent interactions** with the corresponding atomic cross sections  $\{\sigma_i\}_{i=1}^M$ , mean free paths  $\{\lambda_i = 1/N\sigma_i\}_{i=1}^M$ , macroscopic cross sections  $\{\Sigma_i = N\sigma_i\}_{i=1}^M$  and the  $M$  interaction lengths as stochastic variables  $\{s_i\}_{i=1}^M$  such that  $s_i \in \mathbb{Exp}(\Sigma_i)$ , then the **shortest interaction length** i.e.  $\eta = \min\{s_1, \dots, s_M\}$ ,  $\eta \in \mathbb{Exp}(\Sigma_t)$ , where  $\Sigma_t = \sum_{i=1}^M \Sigma_i = \sum_{i=1}^M 1/\lambda_i$  is the **total macroscopic cross section**

## Theorem

If  $\{s_1, \dots, s_M\}$  are independent random variables and  $\forall i \in 1, \dots, M$   $s_i \in \text{Exp}(1/\lambda_i)$ , then  $\eta = \min\{s_1, \dots, s_M\} \in \text{Exp}(\sum_{i=1}^M 1/\lambda_i)$


## Proof.

By definition, the cumulative distribution function of  $\eta$  as a random variable is equal to the probability that

$$\begin{aligned} P(\eta < x) &= 1 - P(\eta \geq x) = 1 - P(s_1 \geq x, \dots, s_M \geq x) = 1 - \prod_{i=1}^M P(s_i \geq x) \\ &= 1 - \prod_{i=1}^M (1 - P(s_i < x)) = 1 - \prod_{i=1}^M (1 - (1 - e^{-x/\lambda_i})) = \\ &1 - e^{-x \sum_{i=1}^M 1/\lambda_i} \implies \eta = \min\{s_1, \dots, s_M\} \in \text{Exp}(\sum_{i=1}^M 1/\lambda_i) \end{aligned} \quad (5)$$



- usually, the **total macroscopic cross section**  $\Sigma_t = \sum_{i=1}^M \Sigma_i = N \sum_{i=1}^M \sigma_i$  is computed and used to sample the path length  $\eta$  to the next discrete interaction point from the exponential distribution,  $\eta \in \mathbb{Exp}(\Sigma_t)$
- then at the post-step point, the type of the discrete interaction is sampled based on the discrete probabilities  $p(\text{proc} = i) = \Sigma_i / \Sigma_t$
- in Geant4, **each discrete process will propose an interaction length**  $s_i \in \mathbb{Exp}(\Sigma_i \equiv 1/\lambda_i)$  based on their own **macroscopic cross section**  $\Sigma_i$
- then **the shortest**  $\eta = \min\{s_1, \dots, s_M\}$  **will be selected as discrete step limit**, which is (based on the proof given in the previous slide) equivalent to the previous one
- moreover, with this Geant4 already **selects the discrete interaction at the pre-step point** that will(if any<sup>1</sup>) happen at the post-step point i.e. after travelling the path  $\eta$

<sup>1</sup> there are several cases when the discrete interaction does not happen at all, we will discuss them soon one by one 

In case of particles, that have *ionization* and *Bremsstrahlung*:

- due to the secondary production threshold, the corresponding **restricted macroscopic cross sections**  $\Sigma_{\text{rest}}(E; E^{\text{cut}}, \dots)$  are used to propose the discrete step limit
- due to the continuous part, i.e. the "along step energy losses", the pre-step point  $E^{\text{pre}}$  and post-step point  $E^{\text{post}}$  kinetic energies will be different
- the **cross section is not constant along the step**, that must be accounted:
  - a fictive, delta interaction is introduced for each interaction such that, the sum of the real  $\Sigma_i^r(E)$  and this delta interaction  $\Sigma_i^\delta(E)$  cross section is constant along the step i.e. does not depend on the particle energy

$$\Sigma_i^r(E) + \Sigma_i^\delta(E) = \Sigma_i(E) \equiv \Sigma_i^{\text{const}} \implies \text{constant along the step} \quad (6)$$

- and  $\Sigma_i^r(E) \leq \Sigma_i^{\text{const}}$  along the step, which implies that  $\Sigma_i^{\text{const}} = \max\{\Sigma_i(E)\}$  i.e. maximum along the step
- this constant macroscopic cross section  $\Sigma_i^{\text{const}}$  is used to sample the discrete interaction length(to a real or delta interaction) at the pre-step point
- at the post-step point, after the along step energy loss is already accounted, the probability that the delta interaction happen is  $p(\delta) = 1 - \Sigma_i^r(E^{\text{post}})/\Sigma_i^{\text{const}}$



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So far, the discrete interaction step limits have been considered:

- each discrete process (or discrete part of the process) proposed a step length
- the shortest among these was selected as **the current candidate step length**
- the corresponding process was selected as **the current candidate process**
- **a flag to indicate** that the current candidate step length was proposed by the **discrete part** of the current candidate process
- possible energy loss along the step was considered

At this point, `G4SteppingManager` will ask each continuous part of the processes to propose their own step limits one by one:

- starts with the previous settings regarding the candidate step length, process, and type flag (discrete)
- **each proposed continuous step limit is compared to the current candidate one**
- if the current continuous step limit is shorter than the current candidate, **the candidate step length, process and type flag (continuous)** are updated

In case of particles, that have *Ionization* and *Bremsstrahlung* the following continuous step limit function is used:

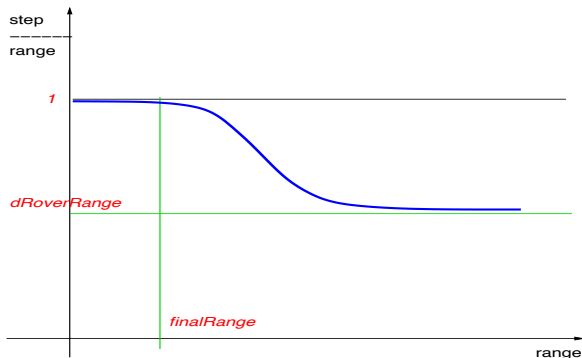
When the particle range  $R > \rho_R \equiv \text{finalRange}$ :

$$\Delta S_{lim} = \alpha_R R + \rho_R (1 - \alpha_R) \left( 2 - \frac{\rho_R}{R} \right)$$

- **default value:**  $\rho_R = 1.0[\text{mm}]$
- $\alpha_R \equiv dRoverRange$
- **default value:**  $\alpha_R = 0.2$
- **at high energies:**  $\Delta S_{lim} \approx \alpha_R R$

When the particle range  $R < \rho_R$ :

- **low energies:**  $\Delta S_{lim} = R$



Based on **restricted range**, computed from the **restricted stepping power**!

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So far a candidate physics step length has been selected that is:

- **the current minimum of the step lengths, proposed by all discrete and continuous processes, was selected as candidate step length**
- **it is assumed at this point, that the particle will travel this path length as a straight line along it's original direction:**
  - to the post-step point where the selected discrete interaction takes place ← that discrete interaction proposed the shortest step length
  - to the post-step point where no discrete interaction takes place ← if a continuous interaction proposed the shortest step length

However, there are one or two **special continuous processes left: transportation (always), multiple scattering (possible)**. The end of the step limitation depends:

- if the particle **does not have Coulomb scattering (A.)**
- if the particle has **Coulomb scattering** and its **described by a single scattering model** i.e. as a **discrete process (B.)**
- if the particle has **Coulomb scattering** and its **described by a multiple scattering model** i.e. as a **continuous process (C.)**

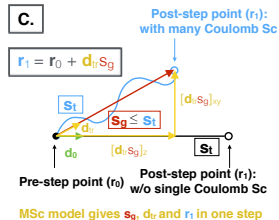
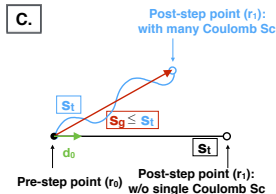
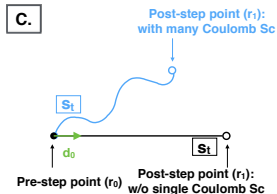
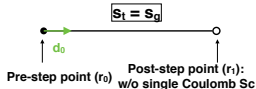
Particles, that do not have Coulomb scattering (**A.**):

- the only remaining (continuous) process is the **transportation** (the last in the list!)
- the above **candidate physical step** length will be **selected** as proposed by the physics since all discrete and all but one (transportation) continuous processes have already proposed their step lengths
- regarding the physics interactions (transportation is still to be called), the particle is supposed to be transported the **selected distance** from the pre-step point **along its original direction**
- **the last continuous process, transportation** will be asked to propose its step limit:
  - if the **particle can be propagated** along its original direction to the selected distance **without hitting volume boundary**, then the **transportation process will accept the proposed step length** (and it will propagate the particle to its proposed post-step point)
  - **otherwise**, the **particle will be transported to the volume boundary** and the **proposed step length** will be **shortened** by transportation process accordingly

Particles, that have Coulomb scattering and its described by a single scattering model i.e. as a discrete process (**B.**):

- elastic scattering was already accounted in the step limit since it is included in the list of discrete processes
- therefore, everything is the same as in the previous case (**A.**) since the only remaining (continuous) process is the **transportation** (the last in the list!)

**A. and B.**  
 $\mathbf{r}_1 = \mathbf{r}_0 + \mathbf{d}_0 S_t$



Particles, that have Coulomb scattering and its described by a multiple scattering model i.e. as a continuous process (**C.**):

- elastic scattering was removed from the list of discrete interactions  $\implies$  **elastic scattering is not included in the current step limit  $s_t$**
- therefore, the current candidate step length  $s_t$  does not contains the effects of (possible many) elastic scattering
- with elastic scattering, the particle would stop (probably many times) along the currently selected path length  $s_t$  and the direction of propagation would change each time
- therefore, the particle would travel the currently selected path length to the next interaction along a curved, zig-zagged trajectory instead of a straight line
- the real post-step point location, the distance (straight line) between the pre- and post step points (**geometrical step length  $s_g$** ) and final direction of propagation is provided by the Multiple Scattering model



- therefore, before the **transportation (the last), the last but one continuous process, multiple scattering** is asked to provide its step limitation:
  - multiple scattering can further limit the current candidate path length  $s_t$
  - after its own step limitation, multiple scattering will change the current **true step length  $s_t$**  to the **geometrical step length  $s_g$**  by computing the corresponding transport distance and **transport direction  $d_{tr}$**
- after the **multiple scattering (last but one)**, the **last continuous process, transportation**'s continuous step limitation is invoked, by providing **the transport distance  $s_g$  instead of the true step length  $s_t$**
- from this point everything is the same as in case of **A.** and **B.**

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- using production cuts is **NOT mandatory** in Genat4: interactions (e.g. *ionization*) can be described as pure discrete interaction (the appropriate model, process need to be selected)
- using multiple scattering model is **NOT mandatory** neither: single scattering models, processes for elastic scattering are available in Geant4 that will be pure discrete
- using pure discrete processes in the simulation gives exact solution of the transport problem: but **feasible only at low energies!** (below few 100 keV)
- in general, at high energies we need to use condensed history approximation and production cuts to be able to solve the transport problem within acceptable computational time
- these approximations make the transport simulation more complex and naturally introduce some user defined parameters, settings
- understanding more details of these approximations helps the user to provide settings that are the most suitable for their applications
- Geant4 provides several predefined physics lists prepared and maintained by expert physics developers for special application areas

# Questions...