

# Electromagnetic Physics I

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slides based on those of Mihaly Novak (CERN)

# Outline

- Electromagnetic (EM) physics overview
  - Introduction, structure of Geant4 EM physics
  - Standard EM physics constructors
  - How to extract EM physics related quantities
- EM processes and stepping
- Multiple scattering and transportation
- Special EM topics:
  - EM models per region
  - Atomic de-excitation
  - Energy loss fluctuation

# Code Location:

## source/processes/electromagnetic/

- /standard
  - $\gamma$ ,  $e^+$  up to 100 TeV
  - hadrons up to 100 TeV
  - ions up to 100 TeV
- /muons
  - up to 1 PeV
  - energy loss propagator
- /xrays
  - Cerenkov, transition, synchrotron radiation
- /highenergy
  - e.g.  $\gamma$  to  $\mu^+ \mu^-$  pairs,  $e^+ e^-$  to  $\pi^+ \pi^-$
- /polarisation
  - models, processes for polarized beams
- /utils
  - model/process interfaces, utilities
- /lowenergy
  - Livermore library:  $\gamma$ ,  $e^-$  [10 eV – 1 GeV]
  - Penelope models (2008):  $\gamma$ ,  $e^+$ ,  $e^-$ , [100 eV – 1 GeV]
  - Livermore polarized processes
  - hadrons and ions up to 1 GeV
  - atomic de-excitation (Auger, fluor.)
- /dna
  - DNA models, processes (0.025 eV to 10 MeV)
  - microdosimetry models for radiology
  - many models are material-specific (water)
- /adjoint
  - reverse Monte Carlo
  - very fast, limited applications

# Standard EM Interactions

- Photon ( $\gamma$ ) :

- conversion to  $e^- e^+$  pairs
- Compton (incoherent) scattering
- photo-electric effect
- Rayleigh (coherent) scattering
- photo-nuclear interaction (see hadronic)

- Electron and positron interactions :

- ionization
- Coulomb (elastic) scattering
- bremsstrahlung photon emission
- positron annihilation
- electron- and positron-nuclear interactions (see hadronic)

# Standard EM Interactions

- Example of photon interactions (from log file)

```
phot:  for gamma  SubType= 12  BuildTable= 0
      LambdaPrime table from 200 keV to 100 TeV in 61 bins
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      LivermorePhElectric : Emin=      0 eV   Emax=    100 TeV  AngularC

compt:  for gamma  SubType= 13  BuildTable= 1
      Lambda table from 100 eV  to 1 MeV, 7 bins per decade, spline: 1
      LambdaPrime table from 1 MeV to 100 TeV in 56 bins
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      Klein-Nishina : Emin=      0 eV   Emax=    100 TeV

conv:  for gamma  SubType= 14  BuildTable= 1
      Lambda table from 1.022 MeV to 100 TeV, 18 bins per decade, spline:
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      BetheHeitler : Emin=      0 eV   Emax=     80 GeV  AngularC
      BetheHeitlerLPM : Emin=    80 GeV  Emax=    100 TeV  AngularC

Rayl:  for gamma  SubType= 11  BuildTable= 1
      Lambda table from 100 eV  to 100 keV, 7 bins per decade, spline: 0
      LambdaPrime table from 100 keV to 100 TeV in 63 bins
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      LivermoreRayleigh : Emin=      0 eV   Emax=    100 TeV  CullenG
```

# Structure of Geant4 EM Physics

- Uniform, coherent design approach covering EM sections
  - standard and low-energy EM models/processes can be combined
- Physical interactions described by processes (e.g. `G4ComptonScattering`)
  - assigned to a particle in the physics list (`G4ComptonScattering` assigned to photon)
- Processes categorized by their interfaces:
  - `G4VEmProcess` for discrete EM processes like Compton
  - `G4VEnergyLossProcess` for continuous-discrete ionization and bremsstrahlung
  - `G4VMultipleScattering` for the condensed history description of multiple Coulomb scattering (along a given step)
- A given EM process can be described by one or more models:
  - an EM model can handle the interaction in a given energy range
  - naming convention: `G4ModelNameProcessNameModel` (e.g. `G4KleinNishinaComptonModel` describes Compton scattering of photons as implemented by the Klein-Nishina differential cross section)
  - each EM model follows the `G4VEmModel` interface:
    - computation of interaction cross section (and stopping power, if any)
    - computation/generation of the interaction final state (kinematics, secondaries, etc.)

# Standard EM Example

- Gamma conversion process described by two EM models

```
phot:  for gamma  SubType= 12  BuildTable= 0
      LambdaPrime table from 200 keV to 100 TeV in 61 bins
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      LivermorePhElectric : Emin=      0 eV   Emax=    100 TeV  AngularGenSauter

compt:  for gamma  SubType= 13  BuildTable= 1
      Lambda table from 100 eV to 1 MeV, 7 bins per decade, spline: 1
      LambdaPrime table from 1 MeV to 100 TeV in 56 bins
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      Klein-Nishina : Emin=      0 eV   Emax=    100 TeV

conv:  for gamma  SubType= 14  BuildTable= 1
      Lambda table from 1.022 MeV to 100 TeV, 18 bins per decade, spline: 1
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      BetheHeitler : Emin=      0 eV   Emax=     80 GeV  AngularGenUrban
      BetheHeitlerLPM : Emin=    80 GeV  Emax=    100 TeV  AngularGenUrban

Rayl:  for gamma  SubType= 11  BuildTable= 1
      Lambda table from 100 eV to 100 keV, 7 bins per decade, spline: 0
      LambdaPrime table from 100 keV to 100 TeV in 63 bins
      ===== EM models for the G4Region  DefaultRegionForTheWorld =====
      LivermoreRayleigh : Emin=      0 eV   Emax=    100 TeV  CullenGenerator
```

# Standard EM Physics Constructors

- Physics processes are assigned to particles in the physics list
- Particles to which EM physics processes can be assigned:
  - $\gamma$ ,  $e^{+/-}$ ,  $\mu^{+/-}$ ,  $\pi^{+/-}$ ,  $p$ ,  $\Sigma^{+/-}$ ,  $\Xi^-$ ,  $\Omega^-$ , anti ( $\Sigma^{+/-}$ ,  $\Xi^-$ ,  $\Omega^-$ )
  - $\tau^{+/-}$ ,  $B^{+/-}$ ,  $D^{+/-}$ ,  $D_S^{+/-}$ ,  $\Lambda_C^+$ ,  $\Sigma_C^+$ ,  $\Sigma_C^{++}$ ,  $\Xi_C^+$ , anti ( $\Lambda_C^+$ ,  $\Sigma_C^+$ ,  $\Sigma_C^{++}$ ,  $\Xi_C^+$ )
  - $d$ ,  $t$ ,  ${}^3\text{He}$ ,  $\alpha$ , generic ion, anti ( $d$ ,  $t$ ,  ${}^3\text{He}$ ,  $\alpha$ )
- Each particle type is a static object which has its own **G4ProcessManager**
  - manager maintains list of assigned processes
- Modular physics lists (**G4VModularPhysicsList**) allow building up a complete physics list from “physics modules”
  - physics module handles well-defined subset of physics (EM physics, decay physics, etc.)
  - **G4VPhysicsConstructor** is the interface class describing such subsets
- Several pre-defined EM physics constructors are available in Geant4

# Standard EM Physics Constructors for HEP

- Description of Coulomb scattering is the same for three of these:
  - $e^{\pm}$ : Urban-MSC model below 100 MeV and the Wentzel-WVI + single scattering model above 100 MeV
  - muons and hadrons: Wentzel-WVI + single scattering model
  - ions: Urban-MSC model
- But different MSC stepping algorithms or parameters are used to optimize either speed or accuracy

Constructor	Components	Comments
<code>G4EmStandardPhysics</code>	Default: nothing or <b>_EM0</b> (QGSP_BERT, FTFP_BERT,...)	for ATLAS and other HEP simulation applications
<code>G4EmStandardPhysics_option1</code>	Fast: due to <b>simpler MSC step limitation</b> , cuts used by photon processes (FTFP_BERT_ <b>EMV</b> )	similar to one used by CMS; good for crystals but not good for sampling calorimeters (i.e. with more detailed geometry)
<code>G4EmStandardPhysics_option2</code>	Experimental: similar to option1 with updated photoelectric model <b>but no-displacement in MSC</b> (FTFP_BERT_ <b>EMX</b> )	similar to one used by LHCb

# Hybrid EM Physics Constructors

- Primary goal: best physics accuracy
- Combine standard and low energy EM models to do this
- More accurate  $e^{\pm}$  MSC models (Goudsmit-Saunderson) and/or more accurate stepping algorithms (compared to HEP)
- More stringent continuous step limitations due to ionization
- Recommended for more sensitive applications: medical, space

Constructor	Components	Comments
<code>G4EmStandardPhysics_option3</code>	Urban MSC model for all particles	proton/ion therapy
<code>G4EmStandardPhysics_option4</code>	most accurate combination of models (particle type and energy); GS MSC model with Mott correction and error-free stepping for $e^{\pm}$ )	the ultimate goal is to have the most accurate EM physics description
<code>G4EmLivermorePhysics</code>	Livermore models for $e^-$ , $\gamma$ below 1 GeV and standard above; same GS MSC for $e^{\pm}$ as in <code>option4</code> )	accurate Livermore based low energy $e^-$ and $\gamma$ transport
<code>G4EmPenelopePhysics</code>	PENELOPE models for $e^{\pm}$ , $\gamma$ below 1 GeV and standard above; same GS MSC for $e^{\pm}$ as in <code>option4</code> )	accurate PENELOPE based low energy $e^-$ , $e^+$ and $\gamma$ transport

# Experimental EM Physics Constructors

- Usually used only by developers for validation and model improvement
- Main difference is in description of Coulomb scattering (GS, WVI, SS)

Constructor	Components	Comments
<code>G4EmStandardPhysicsGS</code>	standard EM physics and the GS MSC model for $e^\pm$ with HEP settings	may be considered as an alternative to EM0 i.e. for HEP
<code>G4EmStandardPhysicsWVI</code>	WentzelWVI + Single Scattering mixed simulation model for Coulomb scattering	high and intermediate energy applications
<code>G4EmStandardPhysicsSS</code>	single scattering (SS) model description of the Coulomb scattering	validation and verification of the MSC and mixed simulation models
<code>G4EmLowEPPysics</code>	Monarsh University Compton scattering model, 5D gamma conversion model, WVI-LE model	testing some low energy models
<code>G4EmLivermorePolarized</code>	polarized gamma models	a (polarized) extension of the Livermore physics models

# Extracting EM Physics-related Quantities

- You may want to know cross sections, energy loss, etc.
- Use the G4EmCalculator object
  - include the following lines in your application:

```
#include "G4EmCalculator.hh"
...
G4EmCalculator emCalculator;

G4Material* material = G4NistManager::Instance()->FindOrBuildMaterial(matName);
G4double macXSec = emCalculator.ComputeCrossSectionPerVolume(energy,
                                                             partDefinition,
                                                             procName,
                                                             material);

G4cout << G4BestUnit(macXSec, "1/Length") << G4endl;
```

- make sure physics list is initialized first
- A good example of all the things you can do:
  - /examples/extended/electromagnetic/TestEm0
  - see especially RunAction::BeginOfRun() method

# EM Processes and Stepping

# Continuous-Discrete Processes

- Charged particle processes ionization, bremsstrahlung and multiple scattering are all **continuous-discrete**
  - discrete part has step limit (path length to next interaction) determined by restricted cross section
  - continuous part has a step limit due to maximum allowed energy loss along the step
  - because of along-step energy loss, kinetic energy is different at pre- and post-step points

# Restricted Stopping Power

- Secondary electrons and gammas produced with energies below  $E^{\text{cut}}$  are not simulated explicitly
- They are described as continuous energy loss along the step and are based on a mean value
- This mean value is the restricted stopping power (mean energy loss along a unit step length)

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

# Restricted Cross Section

- Electrons and gammas with energies above  $E^{\text{cut}}$  are simulated explicitly as discrete interactions
- Discrete interaction probability is determined by the restricted cross section

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

- This covers only the interactions in which the secondary has kinetic energy above the production threshold

# Discrete Part of Step Limit

- Interactions will propose a step length
  - target atom number density is  $N$  and atomic interaction cross section is  $\sigma$  (assumed constant for now)
  - p.d.f. of the interaction length  $x$  is

$$p(x) = N\sigma \exp(-xN\sigma)$$

- the mean (expected value) of the interaction length  $x$  is

$$\mathbb{E}(x) = \frac{1}{N\sigma} \equiv \lambda = \frac{1}{\Sigma}$$

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  $\Sigma_i$  the interaction with the shortest interaction length  $x_i$  is chosen by the simulation

# Discrete Part of Step Limit

- Typically, Monte Carlo simulations will sample the path length to the next discrete interaction point using the distribution  $\exp(-\Sigma_t x)$  where  $\Sigma_t$  is the total macroscopic cross section

$$\Sigma_t = \sum_{i=1}^M \Sigma_i = N \sum_{i=1}^M \sigma_i$$

- Then at the post step point, the type  $i$  of the discrete interaction is sampled according to the discrete probabilities

$$p(\text{proc} = i) = \Sigma_i / \Sigma_t$$

- But in Geant4, each discrete process proposes an interaction length sampled from its own macroscopic cross section:  $\exp(-\Sigma_i x)$ 
  - the process with the shortest interaction length is the one that occurs
- In this way Geant4 already selects at the pre-step point what (if anything) will happen at the post-step point

# Discrete Part of Step Limit

- For particles that have ionization and bremsstrahlung the corresponding restricted macroscopic cross sections  $\Sigma_{\text{rest}}(E, E^{\text{cut}})$  are used to propose the discrete step limit
- Due to the continuous part (along-step energy losses), the energies at the pre-step and post-step points will be different
- The cross section therefore is generally **not constant** along the step. This is accounted for by:

- the addition of a fictitious interaction  $\delta$  with cross section energy dependence such that

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

and  $\Sigma_i^r(E) \leq \Sigma_i^{\text{const}}$  along the step, which implies that  $\Sigma_i^{\text{const}} = \max\{\Sigma_i(E)\}$

- The new constant cross section is used to sample, at the pre-step point, the interaction length to the real or fictitious interaction
- At the post-step point, after energy loss has been accounted for, the probability that the fictitious interaction occurs is

$$p(\delta) = 1 - \Sigma_i^r(E^{\text{post}}) / \Sigma_i^{\text{const}}$$

# Continuous Part of Step Limit

- Up to now, the discrete part has been considered:
  - each process proposed a step length
  - the shortest of these was selected as a **candidate** step length
  - corresponding process was selected as the **candidate** process
  - flag set to indicate that current candidate step length was proposed by discrete part of candidate process
  - a possible energy loss along the step was considered
- Now, G4SteppingManager asks the continuous part of each process to propose its own step limits
  - starting with all the previous (discrete) settings and type flag
  - each proposed continuous limit is compared to the current candidate limit
  - if current continuous step limit is shorter than the current candidate, the candidate step length, process and type flag (continuous) are updated accordingly

# Continuous Part of Step Limit

- For particles that have ionization and bremsstrahlung, the following continuous step limit function is used:

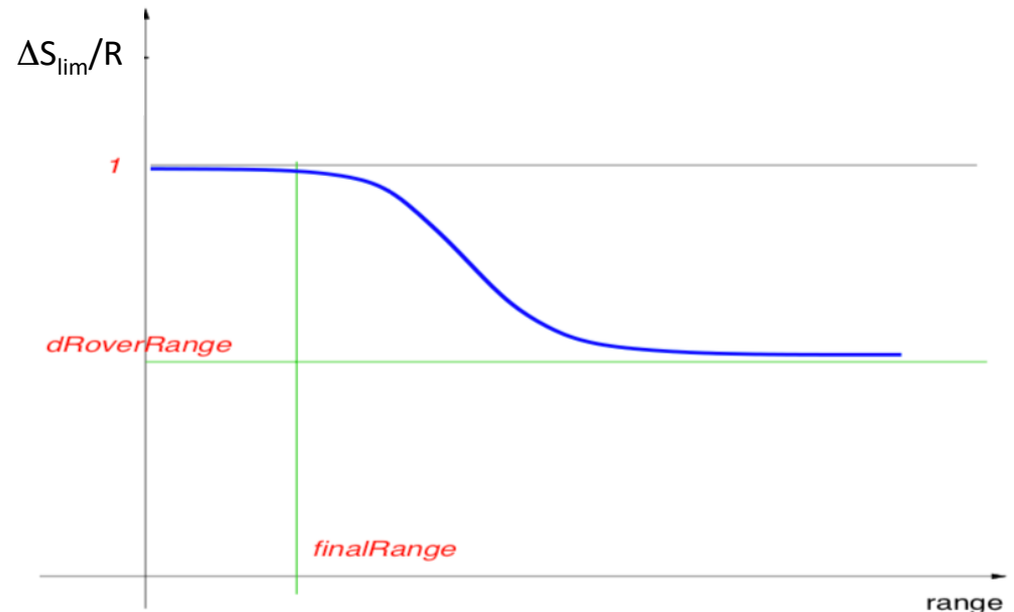
When the particle range  $R > \rho_R \equiv \text{finalRange}$ :

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

- **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 restricted stopping power

# Multiple Scattering and Transportation

# Multiple Scattering and Transportation

- Up to now, a candidate physics step length has been selected which is
  - the current minimum of the step lengths proposed by all discrete and continuous processes
  - and we have assumed that the particle will travel this length in a straight line in its original direction
    - to the post-step point where the selected discrete interaction takes place
    - to the post-step point where no discrete interaction takes place, if a continuous interaction proposed the shortest step length
- However, there are two special continuous processes left: transportation (which always occurs) and multiple scattering (which may occur). The end-of-step limitation depends on one of three conditions:
  - the particle has no Coulomb scattering (A)
  - it has Coulomb scattering and is described by a single-scattering model (discrete) (B)
  - it has Coulomb scattering and is described by a multiple scattering model (continuous process) (C)

# Transportation

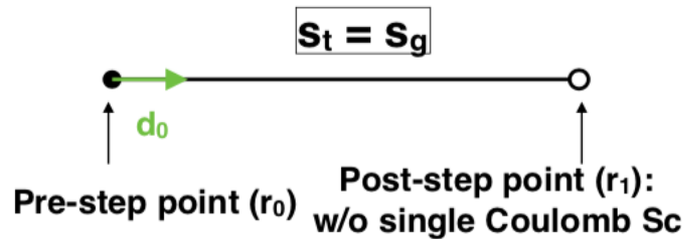
- For particles that do not have Coulomb scattering (A):
  - the only remaining continuous process is transportation
  - it is the last process to propose a step length
  - particle is supposed to be transported the selected distance from the pre-step point along its original direction, according to all the foregoing physics
  - but transportation now gets to propose its step limit:
    - if particle can be propagated to its selected distance without crossing a volume boundary, the transportation process accepts the proposed length
    - otherwise, particle is transported to volume boundary and proposed step length is shortened accordingly

# Single Coulomb Scattering

- For particles that have single Coulomb scattering (B)
  - elastic scattering was already accounted for in the step limit since it is included in the list of discrete processes
  - so everything is the same as in case A, since the only remaining continuous process is transportation

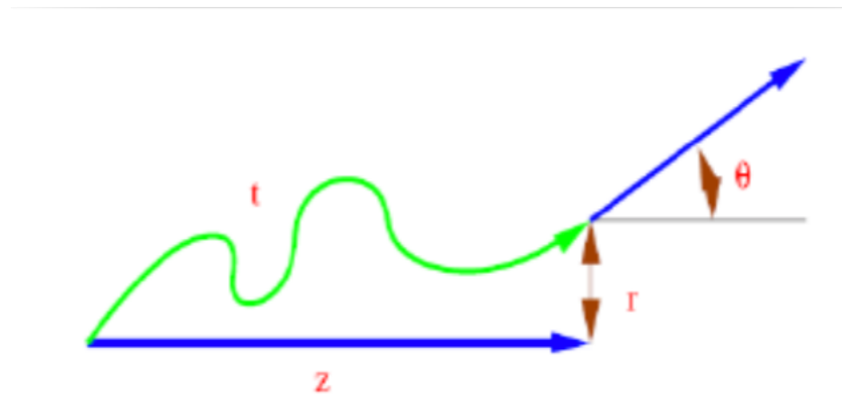
A. and B.

$$\mathbf{r}_1 = \mathbf{r}_0 + \mathbf{d}_0 S_t$$



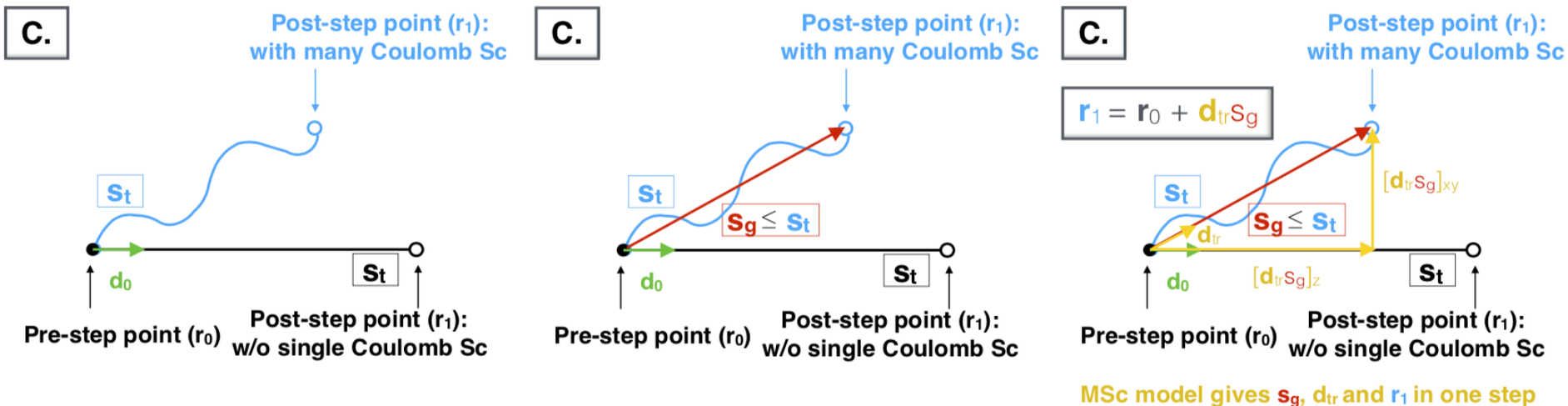
# Multiple Coulomb Scattering

- Elastic scattering of charged particles by the atomic potential
- Event-by-event modeling of elastic scattering is feasible only if mean number of interactions per track is less than a few hundred
- Detailed simulation therefore limited to electrons with low kinetic energies ( < 100 keV) or thin targets
- Electrons with  $E_{\text{kin}} > 100$  keV undergo a large number of elastic while slowing down in thick targets → **condensed history approach**
- MSC models simulate each particle by allowing individual steps which are much larger than average step length between two successive elastic scatterings → **only summed effects are modeled**



# Multiple Coulomb Scattering

- For particles that have the continuous process multiple Coulomb scattering (C)
  - elastic scattering is not included in the list of discrete interactions  $\rightarrow$  cannot propose an elastic step size
  - with elastic scattering there would be many scatterings and changes of direction along ( $s_t$ ) the proposed step length  $\rightarrow$  zig-zag trajectory instead of straight line
  - the multiple scattering model provides the real step length ( $s_g$ ) and final direction of travel  $d_{tr}$



# Multiple Coulomb Scattering

- So, with transportation the last, multiple scattering is next to last 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 multiple scattering, transportation invokes its step limitation by providing the transport distance  $s_g$  instead of the true step length  $s_t$
- From this point on, everything is identical to cases A and B

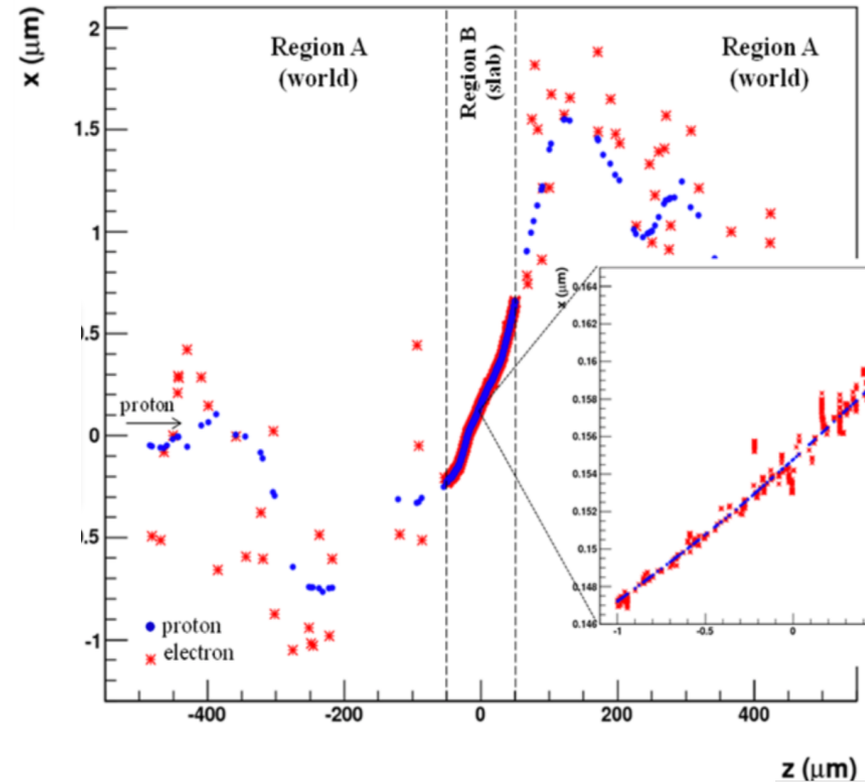
# Geant4 Multiple Scattering Models

Model	Particle type	Energy limit	Specifics and applicability
Urban (L. Urban 2006)	Any	-	Default model (i.e. in EM-opt0) for electrons and positrons below 100 MeV, (Lewis 1950) approach, tuned to data, <u>used for LHC production</u> .
Screened Nuclear Recoil (Mendenhall and Weller 2005) TestEm5	p, ions	< 100 MeV/A	Theory based, providing simulation of nuclear recoil for sampling of radiation damage, focused on precise simulation of in case of <b>space applications</b>
Goudsmit-Saunderson	e <sup>+</sup> , e <sup>-</sup>	-	Theory based angular distributions (Goudsmit and Saunderson 1950). Mott correction and several stepping option including error-free (Kawrakov et al. 1998), <b>precise electron transport</b>
Coulomb scattering (2008)	Any	-	Theory based (Wentzel 1927) single scattering model, uses nuclear form-factors (Butkevich et al. 2002), focused on <b>muons and hadrons</b>
WentzelVI (2009) LowEnergyWentzelVI (2014)	Any	-	Mixed simulation model: MSC for small angles, Coulomb Scattering (Wentzel 1927) for large angles. Focused on simulation for <b>muons and hadrons</b> ; low-energy model is applicable for low-energy e <sup>-</sup>
Ion Coulomb scattering (2010) Electron Coulomb scattering (2012)	Ions e <sup>+</sup> , e <sup>-</sup>	-	Model based on Wentzel DCS + relativistic effects + screening effects for projectile & target.

# EM Special Topics

# EM Models per Region

- Special models may be used in a particular G4Region, while all other parts of detector use general models
- Example: Geant4-DNA physics in a slice of volume (Region B) nested in World (Region A) which uses standard EM
- Use G4EmConfigurator to select special model, set energy limits within region
- UI commands allow optimization and easy configuration of models
- Can be used on top of any EM constructor:
  - `/process/em/AddPAIRegion proton MYREGION pai`
  - `/process/em/AddMicroElecRegion MYREGION`
  - `/process/em/AddDNARegion MYREGION opt0`

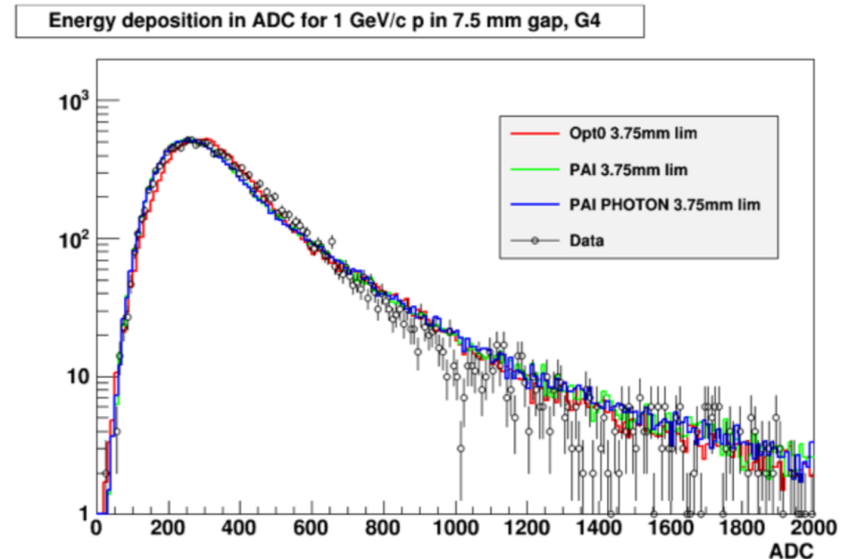
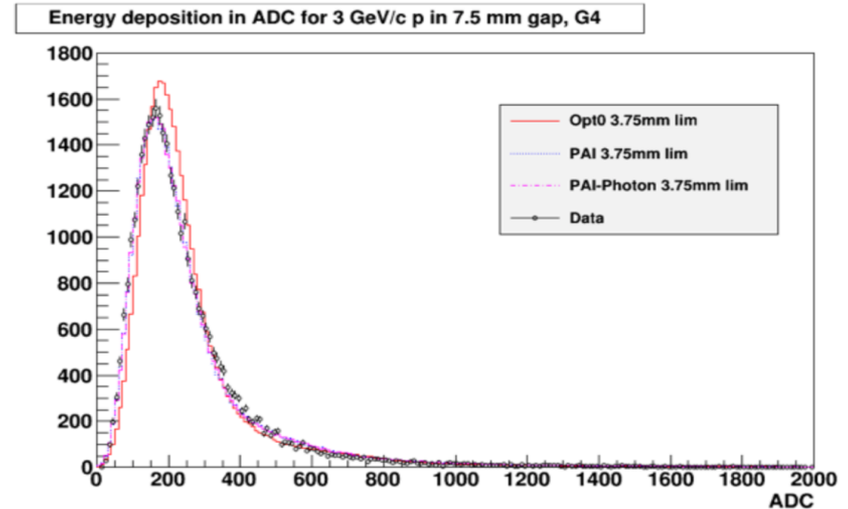


# Atomic De-excitation

- Initiated by other physics interactions
  - e.g. photoelectric effect, ionization, radioactive decay
  - interactions leave target atom in excited state
- Evaluated Atomic Data Library (EADL) contains transition probabilities for:
  - radiative transitions (fluorescence photon emission)
  - Auger e- emission (initial and final vacancies are in different shells)
  - Coster-Kronig e- emission (initial and final vacancies are in same shell)
- Due to a common interface, Geant4 atomic de-excitation is compatible with both standard and low energy EM categories
  - can be enabled and controlled by UI commands (before initialization):
    - `/process/em/fluor true`
    - `/process/em/auger true`
    - `/process/em/augerCascade true`
    - `/process/em/pixar true`
    - `/run/initialize`
  - fluorescence transition is active by default in some EM constructors

# Energy Loss Fluctuation

- For condensed history models:
  - secondary photons (e-) with initial energy below the photon (e-) production threshold, are not generated in bremsstrahlung (ionization)
- Corresponding energy loss (that would have been taken away by these secondaries) is counted as continuous energy loss of primary particle along its step
- Mean value of energy loss along step (due to these sub-threshold secondaries) can be calculated using the restricted stopping power
- So we have the mean – what about the distribution?
  - energy loss fluctuation models will tell us
- Urban and PAI models available in Geant4



# Summary

- EM processes and models are available to cover all “long-lived” charged particles and photons
- Energy range covered: few eV up to ~PeV
  - often more than one model required for this coverage
- EM physics constructors build the models, cross sections and processes
  - many pre-packaged constructors have been prepared and tested by Geant4 developers – pick one
  - but you can still build your own!
- Possible to use different EM physics constructors in different regions of your geometry
- Ionization, bremsstrahlung, multiple scattering and transportation couple in a complicated way to limit the final step length and interaction at the post-step point, with process sampling and step limit proposals in the following order:
  - bremsstrahlung and ionization
  - multiple scattering
  - transportation