

Monte Carlo Simulation of X-ray Transport

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Abstract. An article usually includes an abstract, a concise summary of the work covered at length in the main body of the article. It is used for secondary publications and for information retrieval purposes.

INTRODUCTION

TRANSPORT THEORY

In order to represent the computational domain discretely, space is broken up into a grid of voxels (a pixel with volume), with each voxel being assigned a particular material ID depending on the geometries and compounds/elements in the domain. With our discrete space, given a photon position \vec{r} , the corresponding voxel in which the photon resides can be calculated. Therefore, a particular material can be associated with all possible \vec{r} 's in the domain.

A photon's position in space after taking the n -th step in the domain \vec{r}_n is represented by the following parametric ray equation:

$$\vec{r}_n = \vec{r}_{n-1} + \hat{d}t_n, \quad (1)$$

where \vec{r}_{n-1} is the initial position before the n -th step, \hat{d} is a unit vector in the direction of the step, and t_n is the distance of the n -th step.

In order to sample t_n , we utilize the following probability distribution function (PDF) $p(t)$ of the distance traveled t by a photon of energy E through material M before interacting:

$$p(t) = n\sigma \exp[-t(n\sigma)], \quad (2)$$

where n is the number density of M and $\sigma = \sigma(E, M)$ is the microscopic cross-section of M at E .

Using the inversion method for sampling a PDF, it follows that random values of the free path t can be generated with the following equation:

$$t = -\frac{1}{n\sigma} \ln \gamma, \quad (3)$$

where γ is a uniformly distributed random number in the interval $(0, 1)$.

SURFACE AND DELTA-TRACKING

If, after taking a step, the photon lands in a voxel with a different material, then the corresponding free path for the new material to be accounted for. This method, called surface-tracking, requires photons to be stopped at voxel boundaries and intersections with surrounding voxels to be calculated, which can be computationally intensive for materials that have a large average free path.

Alternatively, the delta-tracking algorithm offers a solution by sampling the maximum cross-section σ_{\max} in the computational domain. This, in turn, brings down the average free path to the minimum in the domain. To account for this decrease in free path, the algorithm introduces delta interactions as an alternative to real interactions, resulting in no change to the energy or direction. The probability of delta interaction P_δ is given by the following equation:

$$P_\delta = \frac{\sigma_{\max}(E) - \sigma(E, M)}{\sigma_{\max}(E)}, \quad (4)$$

where E is the energy of the photon undergoing the step, and M is the material corresponding to the position of the photon at the end of the step. Note that when the photon lands in the material corresponding to the maximum cross-section, $\sigma(E, M) = \sigma_{\max}$ and $P_\delta = 0$. On the contrary, if the photon landed in air and the domain's maximum cross-section corresponded to lead, then $\sigma(E, M) \ll \sigma_{\max}$, making $P_\sigma \approx 1$.

Overall, delta-tracking is significantly more computationally efficient for domains with similar cross-sections and can be shown to yield equivalent results to surface-tracking.

PHOTON INTERACTIONS

If a delta interaction does not occur, then a real interaction is sampled. Therefore, the probability of a real interaction P_r is directly related to P_δ by

$$P_r = 1 - P_\delta, \quad (5)$$

where P_δ is given by Eq. 4.

If a real interaction occurs in material M , then probability of interaction i occurring is

$$P_i = \frac{\sigma_i(E, M)}{\sigma(E, M)}, \quad (6)$$

where σ_i is the cross-section of interaction i .

If there are N possible interactions for a particular E and M , then $\sigma(E, M)$ is calculated as so

$$\sigma(E, M) = \sum_{i=1}^N \sigma_i(E, M). \quad (7)$$

For x-rays, there are three possible photon interactions:

1. Photoelectric Effect

In the photoelectric effect model used in MIDSX, a rather simple approach is taken. When a photon interacts with an atom's electron, the photon is terminated and all energy is deposited at the location of interaction. In general purpose particle transport code systems, when a photoelectric interaction occurs, a photon of energy E is absorbed by an electron in subshell i , causing the electron to leave the atom with energy $E_e = E - U_i$, where U_i is the binding energy of the i th subshell. In addition, photons are emitted due to atomic relaxations. For photon energies in the medical imaging range (30 - 120 keV), the energy of the released electrons does not allow for significant traversal through typically used materials, such as tissue, bone, and fat. This limited traversal results in a localized dose distribution, in turn, validating the model used by MIDSX.

2. Coherent Scattering

Thomson scattering is defined as an incoming photon of energy E elastically scattering with a free electron at rest, resulting in a scattered photon of same energy E . The atomic DCS per unit solid angle Ω for the interaction can be derived with classical electrodynamics, and is given by

$$\frac{d\sigma_T}{d\Omega} = r_e^2 \frac{1 + \cos^2(\theta)}{2}, \quad (8)$$

where r_e^2 is the classical electron radius.

In an atom, photons scatter off bound electrons rather than the free electrons described by Thomson scattering, resulting in what is known as coherent (Rayleigh) scattering at . The DCS per unit solid angle Ω of the interaction, ignoring absorption edge effects, is given by

$$\frac{d\sigma_{Co}}{d\Omega} = \frac{d\sigma_T}{d\Omega} F(x, Z), \quad (9)$$

where x is the momentum transfer between the photon and atom, Z is the atomic number of the atom, and $F(x, Z)$ is the atomic form factor. x is related to the scattering angle θ by

$$x = ak\sqrt{1 - \cos\theta}, \quad (10)$$

where

$$a = \frac{m_e c^2}{\sqrt{2}hc}, \quad (11)$$

and

$$k = \frac{E}{m_e c^2}, \quad (12)$$

where m_e is the mass of an electron, c is the speed of light, and h is Planck's constant.

The DCS per unit solid angle Ω can be integrated over ϕ to obtain the DCS per unit polar angle θ :

$$\frac{d\sigma_{Co}}{d\theta} = \pi r_e^2 \sin\theta (1 + \cos^2\theta) F(x, Z)^2. \quad (13)$$

The PDF of the polar angle θ is then given by

$$p(\theta)d\theta = \frac{d\sigma_{Co}}{d\theta} \frac{1}{\sigma_{Co}} d\theta = \frac{\pi r_e^2}{\sigma_{Co}} \sin\theta (1 + \cos^2\theta) F(x, Z)^2 d\theta. \quad (14)$$

The PDF of θ can be transformed into a PDF of $\mu = \cos\theta$, resulting in

$$p(\mu) = \frac{\pi r_e^2}{\sigma_{Co}} (1 + \mu^2) F(x, Z)^2. \quad (15)$$

To then sample μ for a particular scattering event, the inversion method is used. In particular, a look up table for the CDF of $P(\mu)$ is generated for each material in the domain for a grid of μ values. The details of this algorithm are discussed in Appendix ...

3. Incoherent Scattering

Incoherent (Compton) scattering is defined as an incoming photon of energy E interacting with an atom's electron, resulting in a scattered photon of energy E' and an released electron with energy $E_e = E - E' - U_i$, where U_i is the binding energy of the interacting subshell. While coherent scattering effectively interacts with atom itself, incoherent scattering interacts with the electron. The DCS per unit solid angle Ω of the interaction was derived by Klein and Nishina in 1929, making it one of the first findings of quantum electrodynamics. The Klein-Nishina formula is given by

$$\frac{d\sigma_{KN}}{d\Omega} = \frac{r_e^2}{2} \left(\frac{E'}{E} \right)^2 \left(\frac{E'}{E} + \frac{E}{E'} - \sin^2\theta \right), \quad (16)$$

Note that when $E' = E$, the KN DCS is equal to the Thomson DCS, showing that incoherent scattering is a generalization of coherent scattering for inelastic interactions.

Applying conservation of energy and momentum to free electron at rest, the following equation can be derived relating the scattered photon energy E' to the scattering angle θ and the incident photon energy E :

$$E' = \frac{E}{1 + k(1 - \cos \theta)}. \quad (17)$$

Similar to the Thomson DCS, the KN DCS assumes a free electron at rest. In an atom, the electron is bound, resulting in a modified DCS. In the case of incoherent scattering, the KN DCS is modified by the Coherent Scattering Function, making the DCS per unit solid angle Ω of the interaction

$$\frac{d\sigma_{In}}{d\Omega} = \frac{d\sigma_{KN}}{d\Omega} S(x, Z). \quad (18)$$

Instead of directly sampling the PDF of the DCS, μ is first sampled using the acceptance-rejection method developed by Ozmutlu, then $S(x, Z)$ is sampled once again with the acceptance-rejection method. The details of this algorithm are discussed in Appendix ...

METHODS

In order to simulate the transport of x-rays through a domain, the following is required:

1. The geometry of the domain
2. The materials in the domain
3. The associated cross-sections, form factors, and scattering functions of the materials
4. The source of the x-rays (energy spectrum, position, and direction)
5. The number of photons to simulate

and to retrieve information about the performed simulation, the following is required:

1. Geometries to check for intersection
2. Quantities to tally
3. Derived quantities to calculate from tallied quantities

GEOMETRY

In MIDSX, the geometry of the domain is represented by a 3D array of voxels, with each voxel being assigned a material ID. The domain is assigned a particular size, indicated by its spatial extent along the x, y, and z dimensions. In addition, the domain is assigned a background material ID. Geometries inside the domain are specified by NIFTI files. These NIFTI files are assigned material IDs, spatial size, voxel size, and an origin, which is the location of the voxel in the domain that corresponds to the origin of the NIFTI file. After constructing the NIFTI files, the domain is defined by supplying the background material ID, domain size, and a list of NIFTI files into a custom .domain file, which is then read by the MIDSX executable.

In the code, both a VoxelGrid and ComputationalDomain object are created, with the ComputationalDomain consisting of the specified dimensions, background material ID, and a vector of VoxelGrid objects which are created via the provided NIFTI files. To determine the current material of a photon, it is first determined if the photon is inside the ComputationalDomain. If so, it then checks if the photon is inside any of the VoxelGrid objects. If so, it is then determined which Voxel the photon is located inside the Voxel, then the corresponding material ID is returned. If the photon is not inside the ComputationalDomain, then the photon is terminated.

MATERIALS & DATA

In order to simulate the transport of x-rays through a domain, the materials in the domain must be specified. In MIDSX, materials are all defined in an SQLite database, which is read by the MIDSX executable. The database contains the following information for each element:

1. Symbol
2. Atomic Number
3. Mass
4. Mass Density
5. Number Density
6. Mass Number

All of the above data was obtained from the periodictable and mendelev python packages. In addition, the database contains the following data from the EPDL database for each element:

1. Total Microscopic Cross-Section
2. Photoelectric Microscopic Cross-Section
3. Coherent Scattering Microscopic Cross-Section
4. Incoherent Scattering Microscopic Cross-Section
5. Atomic Form Factor
6. Scattering Function

In MIDSX, all the above data is initialized upon creation of the `InteractionData` object with a vector of strings of materials names. These material names correspond to entries in the SQLite database, which contains a table of material compositions and mass densities which were obtained from NIST's XCOM database. The `InteractionData` object contains a map of `Material` objects with their names, along with additional computed data, such as the maximum cross-section which is used for delta-tracking. The `Material` objects construct the above data for the specified material by performing an additivity approximation of the data for each element in the material (ref). The data is separated into two further objects: `MaterialData` and `MaterialProperties`. `MaterialData` contains the microscopic cross-sections, form factors, and scattering functions, while `MaterialProperties` contains the mass density, number density, and mass.

In addition to retrieving and storing the above data, the `MaterialData` object constructs interpolators for all its data. The interpolators for each type of data vary depending on its shape. For example, the photoelectric and total cross-sections are interpolated with a log-log linear interpolator, while the incoherent and coherent cross-sections are interpolated with a log-log cubic spline interpolator. Below is a figure of .

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REFERENCES