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MC-GPU [1-4] is a Monte Carlo simulation code that can generate synthetic radiographic images and computed tomography (CT) scans of realistic models of the human anatomy using the computational power of commodity Graphics Processing Unit (GPU) cards. The code implements a massively multi-threaded Monte Carlo simulation algorithm for the transport of x rays in a voxelized geometry. The x ray interaction models and material properties have been adapted from **PENELOPE 2006** [5].

MC-GPU was developed using the CUDA programming model from NVIDIA [6] to achieve maximum performance on NVIDIA GPUs. The code can also be compiled with a standard C compiler to be executed in a regular CPU. In a typical medical imaging simulation, the use of GPU computing with MC-GPU has been shown to provide a speed up of between 20 and 40 times, compared to the execution on a single CPU core.

The MC-GPU code has been described in different scientific publications [1-4]. The main reference of this work, which the users should cite, is the following [1]:

Andreu Badal and Aldo Badano, "Accelerating Monte Carlo simulations of photon transport in a voxelized geometry using a massively parallel Graphics Processing Unit", Medical Physics 36, pp. 4878-4880 (2009)

The main developer of MC-GPU is **Andreu Badal**, PhD, at the U.S. **Food and Drug Administration**. The source code is free and open software in the public domain, as explained in the Disclaimer section below. The software distribution website is: http://code.google.com/.

This documentation has been automatically generated by **Doxygen** parsing the comments in the MC-GPU source code. This code is still in development, please report to

the author any issue/bug that you may encounter. Feel free to suggest improvements to the code too!

#### 1.1 Disclaimer

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### 1.2 Code features

In this section we provide a brief description of the features of the MC-GPU code. For more details, consult our published articles and the full Doxygen documentation of the source code and the input file of the sample simulation provided with the MC-GPU package.

The basic operation of the code consists in adapting the simulation input file to describe the location and characteristics of the x ray source, define the CT trajectory (if any), list the materials to be used in the simulation, define the geometry of the x ray detector and, finally, specify the voxelized object file to be used as the simulation material universe. In the first line of the input file, the user can fix the total number of x rays that have to be simulated (> 1e5 histories) or the total simulation time (maximum 1e5 seconds).

The coordinate system of the simulated world is determined by the input voxelized geometry. The origin of coordinates is assumed to be located at the lower-back corner of the voxelized volume, and the axis are located on the vertices of the voxelized volume. This means that the lower-back corner of the first voxel is on the origin and the following voxels are located along the positive X, Y and Z axis (first quadrant).

The code includes two tally options: an **image tally** that creates projection x-ray images, and a radiation **dose tally** that estimates the dose deposited inside the patient model. MC-GPU does not currently simulate the transport of electrons and therefore

1.2 Code features 3

the dose deposition tally (KERMA tally rigorously) will not be accurate for high energies or near material interfaces and small voxels. In the image tally the images are formed by counting the energy that enters a user-defined 2D grid of pixels, which is a simple approximation to a noise-free flat-panel detector with 100% detection efficiency. The pixel values have units of eV/cm<sup>2</sup>. Four different images are reported at the end of the simulation, corresponding to the signal produced by x rays that did not interact between the source and the detector (non-scattered), x rays that suffered a single Compton (inelastic) interaction, a single Rayleigh (elastic) interaction, and multi-scattered x rays. The dose tally counts the energy deposited by each x ray track inside each voxel of the geometry, within a user-defined volumetric region-of-interest (ROI). The average dose deposited inside each voxel and in each material (and the associated statistical uncertainties) are reported at the end of the simulation.

MC-GPU can simulate a single projection image or a full CT scan. The CT is simulated generating many projection images around the static voxelized geometry. Currently, the code is limited to perform a simple CT trajectory rotating around the Z axis. The user can specify the angular shift and longitudinal translation (pitch) of the source between each projection and also the distance between the source and the axis of rotation (the axis is assumed to be parallel to the Z axis). By now, the code does not simulate some relevant components of a CT scanner such as the anti-scatter grid, a bow-tie filter or a curved detector (flat-panel detector only).

The x ray source is defined as a point source emitting x rays with an energy randomly sampled from the user-provided energy spectrum. The polyenergetic spectrum is efficiently sampled using the Walker aliasing algorithm. The emitted cone beam is computationally collimated to produce a rectangular field on the detector plane, within the azimuthal and polar angles specified by the user. The detector plane is automatically located at the specified distance right in front of the source focal spot, with the collimated cone beam pointing towards the geometric center of the detector.

In order to optimize the particle tracking algorithm (ray-tracing) and minimize the accesses to the slow GPU main memory, the photon trajectories across the voxels are computed using the Woodcock tracking algorithm. With this technique the photons perceive the geometry as a uniform medium composed of the material of the most attenuating voxel. In this way, the voxel boundaries do not have to be explicitly calculated and multiple voxels can be crossed in a single step. To keep the simulation unbiased, some of the interactions are considered "virtual" (i.e., do not change the photon energy or direction of movement), depending on the x ray energy and the actual material at the interaction site. In typical medical imaging simulations where the most attenuating material is cortical bone, the Woodcock tracking algorithm gives an speed up of almost one order of magnitude compared to computing voxel boundaries all the time. However, if the geometry includes a high density voxel, such as a metallic implant, the performance of the code can be severely reduced because a large fraction of the sampled interactions will be virtual.

The random number generator used in PENELOPE [5], RANECU, is also used in the GPU program. To ensure that the simulated tracks are not correlated, each thread initializes the generator to a unique position in the random sequence, far enough from

the other threads, using the algorithm implemented in the seedsMLCG code [7].

In a typical simulation, several thousand threads are launched simultaneously in the GPU, each one of them simulating a batch of several x ray tracks. If the code is compiled with MPI support (see below), multiple GPUs can be used in parallel. The code will perform a short speed test to estimate the relative speed of each GPU used in the simulation and then distribute the number of particles among the available GPUs correspondingly. If the user specified a time limit in the simulation, all the GPUs will simulate in parallel for the allowed time. Since the code is already optimized to scale well in thousands of GPU threads, it scales almost linearly with the number of GPUs in most situations, with only a few seconds of overhead in the initialization of the multiple GPUs and in the reduction of the final results.

## 1.3 Code output

At the end of the simulation the code reports the tallied 3D dose distribution and the final simulated images in RAW binary form, as 32-bits float values. The image data is provided as a collection of five consecutive images corresponding to: total image (scatter+primaries), primary particles, Compton, Rayleigh and multi-scatter. The dose data is reported as two RAW files with the mean dose and twice the standard deviation of the dose in each voxel of the geometry respectively, within the input ROI. The average dose deposited in each material of the geometry is also reported to the standard output. Organ doses can be obtained by post-processing the output dose file, knowing which voxel corresponds to each organ. The pixel and voxel dose data values are stored with the X coordinate incrementing first, the Y coordinate incrementing second, and the Z coordinate incrementing last.

The program also reports the simulated images and the dose at the Z plane at the level of the x ray source as ASCII text files. The ASCII output can be readily visualized with the GNUPLOT scripts distributed with MC-GPU. The header section at the beginning of these text files provides the information required to easily read the RAW binary files with IMAGEJ, OCTAVE or other programs.

## 1.4 Code compilation and execution

MC-GPU has been developed and tested only in the Linux operating system. A Makefile script is provided to compile the MC-GPU code in Linux. The CUDA libraries and the GNU GCC compiler must be previously installed. The Makefile may have to be edited to modify the library path.

MC-GPU uses CUDA to access NVIDIA GPUs but all the actual computations are coded in standard C and the CUDA-specific commands are enclosed within preprocessor "if" statements. Defining the pre-processor variable "USING\_CUDA" (i.e., compiling with "-DUSING\_CUDA") the particle transport routines are compiled to simulate

many x ray histories in parallel in an NVIDIA GPU using CUDA. Otherwise, the code is sequentially executed in the CPU. The same coding approach has been used to allow the use of multiple GPUs. Defining the pre-processor variable "USING\_MPI" (i.e., compiling with "-DUSING\_MPI"), Message Passing Interface (MPI) library calls are used to share information between multiple CPU threads in different computers. Each MPI thread gets a unique id in the CPU and addresses a unique GPU. At the end of the simulation the images and doses tallied by the different GPUs are reduced to form single output file equivalent to a sequential simulation of the same number of particles.

This is an example shell command that would compile the code using CUDA and MPI (openMPI library in this case):

```
nvcc -DUSING_CUDA -DUSING_MPI -O3 MC-GPU_v1.2.cu -o MC-GPU_v1.2.x
-use_fast_math -I./ -I/usr/include/openmpi -I/usr/local/cuda/include
-I/opt/cuda_SDK_4.0/C/common/inc/ -L/opt/cuda_SDK_4.0/C/lib/
-L/usr/local/cuda/lib64/ -lmpi -lcutil_x86_64 -lcudart -lm
```

The same source code could also be compiled for a regular CPU using:

```
gcc -x c -03 MC-GPU_v1.2.cu -o MC-GPU_v1.2_CPU.x -I./ -lm
```

To run a simulation (and keep the information reported to the standard output in an external file) the compiled code and its input file can be executed as:

```
./MC-GPU_v1.2.x MC-GPU_v1.2.in | tee MC-GPU_v1.2.out
```

To run the simulation in parallel in multiple GPUs with MPI (5 GPUs in this case) the user can execute:

```
mpirun -n 5 /shared_drive/MC-GPU_v1.2.x /shared_drive/MC-GPU_v1.2.in -v -x LD_LIBRARY_PATH -hostfile hostfile
```

The text file 'hostfile' lists the IP addresses and number of computing slots (GPUs) of the computers collaborating in the simulation. This file is not necessary when using multiple GPUs in a single workstation. When using multiple computers, the simulation files should be located in a shared drive to make sure every node can access the input data. The different workstations must have different host names in order to be differentiated by the MPI threads. The multiple threads communicate to each other to make sure they don't use the same GPU in the same workstation.

# 1.5 List of changes in code version 1.2

- Implemented the dose tally.
- Polyenergetic source model.
- MPI support for simulating individual projections.
- Simulation by time limit.
- Improved flexibility of the CT trajectories.

### 1.6 Known issues

We have observed a small (around 2%) difference between the voxel doses estimated with MC-GPU and PENELOPE (with photons only). This discrepancy is currently under investigation.

## 1.7 References

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