# **Example: Photon Treatment Plan**

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In this example we will show how to load patient data into matRad, how to setup a photon dose calculation based on the VMC++ monte carlo algorithm and how to inversly optimize directly from command window in MatLab.

## **Patient Data Import**

Let's begin with a clear Matlab environment. Next, import the TG119 phantom into your workspace. The phantom is comprised of a 'ct' and 'cst' structure defining the CT images and the structure set. Make sure the matRad root directy with all its subdirectories is added to the Matlab search path.

```
clc,clear,close all
load('BOXPHANTOM.mat');
```

Let's check the two variables, we have just imported. First, the 'ct' variable comprises the ct cube along with some meta information describing properties of the ct cube (cube dimensions, resolution, number of CT scenarios). Please note that mutiple ct cubes (e.g. 4D CT) can be stored in the cell array ct.cube{}

```
ct
ct =
  struct with fields:
       cube: {[160×160×160 double]}
    resolution: [1×1 struct]
```

```
cubeDim: [160 160 160]
numOfCtScen: 1
```

The 'cst' cell array defines volumes of interest along with information required for optimization. Each row belongs to one certain VOI, whereas each column defines different proprties. Specifically, the second and third column show the name and the type of the structure. The tpe can be set to OAR, TARGET or IGNORED. The fourth column depicts a linear index vector depicting voxels in the CT cube that are covered by the VOI. In total, 2 structures are defined in the cst

```
cst
cst =
  2×6 cell array
  Columns 1 through 5
    [0]
            'BODY'
                                            {1×1 cell}
                                                           [1x1 struct]
                               'OAR'
                                            {1×1 cell}
    Γ 1 7
            'OuterTarget'
                               'TARGET'
                                                           [1x1 struct]
  Column 6
    [1x1 struct]
    [1x1 struct]
```

#### **Treatment Plan**

The next step is to define your treatment plan labeld as 'pln'. This structure requires input from the treatment planner and defines the most important cornerstones of your treatment plan.

First of all, we need to define what kind of radiation modality we would like to use. Possible values are photons, protons or carbon. In this case we want to use photons. Then, we need to define a treatment machine to correctly load the corresponding base data. Since we provide generic base data we set the machine to 'Genereric. By this means matRad will look for 'photons\_Generic.mat' in our root directory and will use the data provided in there for dose calculation

Define the flavour of biological optimization for treatment planning along with the quantity that should be used for optimization. Possible values are (none: physical optimization; const\_RBExD: constant RBE of 1.1; LEMIV\_effect: effect-based optimization; LEMIV\_RBExD: optimization of RBE-weighted dose. As we are using photons, simply set the parameter to 'none' thereby indicating the physical dose should be optimized.

```
pln.bioOptimization = 'none';
```

Now we have to set some beam parameters. We can define multiple beam angles for the treatment and pass these to the plan as a vector.matRad will then interpret the vector as multiple beams. In this case, we define one single beam from 0 degree gantry angle and 0 degree couch angle. Moreover, we set the

bixelWidth to 10, which results in a beamlet size of 10 x 10 mm. The number of fractions is set to 30. Be advised that matRad is always optimizing the fraction dose.

```
pln.gantryAngles = [0];
pln.couchAngles = [0];
pln.bixelWidth = 10;
pln.numOfFractions = 30;
```

Obtain the number of beams and voxels from the existing variables and calculate the iso-center which is per default the mass of gravity of all target voxels.

```
pln.numOfBeams = numel(pln.gantryAngles);
pln.numOfVoxels = prod(ct.cubeDim);
pln.voxelDimensions = ct.cubeDim;
pln.isoCenter = ones(pln.numOfBeams,1) *
  matRad_getIsoCenter(cst,ct,0);
```

Enable sequencing and disable direct aperture optimization (DAO) for now. A DAO optimization is shown in a seperate example. The multileaf collimator leaf sequencing algorithm stratifies each beam in N static segments.

```
pln.runSequencing = 0;
pln.runDAO
and et voila our treatment plan is ready. Lets have a look at it:
pln
pln =
  struct with fields:
      radiationMode: 'photons'
             machine: 'Generic'
    bioOptimization: 'none'
       gantryAngles: 0
         couchAngles: 0
          bixelWidth: 10
     numOfFractions: 30
         numOfBeams: 1
         numOfVoxels: 4096000
    voxelDimensions: [160 160 160]
           isoCenter: [240 240 240]
```

## **Generatet Beam Geometry STF**

runDAO: 0

runSequencing: 0

This acronym stands for steering file and comprises the beam geomtry along with the ray and pencil beam positions

```
stf = matRad_generateStf(ct,cst,pln);
```

```
matRad: Generating stf struct... Progress: 100.00 %
```

### **Dose Calculation**

Calculate dose influence matrix for unit pencil beam intensities. using the VMC++ monte carlo algorithm. We define the number of photons simulated per beamlet to be 700. Make sure to download VMC++ files from <a href="http://www.cerr.info/download.php">http://www.cerr.info/download.php</a> into the matRadrootDirectory\vmc++ to run the following function.

```
nCasePerBixel = 700;
dij = matRad_calcPhotonDoseVmc(ct,stf,pln,cst,700,1);
matRad: VMC++ photon dose calculation...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,22.5,22.5)
______
-----
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy :
                             0.447479 MeV
Min. photon transport energy
                          :
                              0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
                              7
CSDA approximation
                              0
                           :
```

```
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step :
                        0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
            : 700
 total histories
 initial rng seeds : 24442 27174
_____
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.317 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
```

```
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.051 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.079 70 70
+++finished batch 10 cpu time: 0.094 70 70
finished simulation, cpu time = 0.101 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze ============
geometry: CT
cpu time: 0.101
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0372868 in region 1803247
   ICRU-2K efficiency: 1066.84 1/s
+++ total
   max dose is 0.0372868 in region 1803247
   ICRU-2K efficiency: 1066.84 1/s
_____
Completed 1 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
               Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,22.5,23)
-----
______
           XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
```

```
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                      0.05 MeV
                   :
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                      1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                      10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                      0.0298764 0.420741
Fano calculation
                    :
Exact Compton
                      Ω
Electron transport mode
                      VMC++
-----
_____
        MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 3810 27402
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
  dump dose: 2
  dose scans:
```

CPU time so far: 1.316 seconds

```
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.052 70 70
+++finished batch 7 cpu time: 0.061 70 70
+++finished batch 8 cpu time: 0.071 70 70
+++finished batch 9 cpu time: 0.08 70 70
+++finished batch 10 cpu time: 0.09 70 70
finished simulation, cpu time = 0.096 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.096
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0376496 in region 1880047
   ICRU-2K efficiency: 1086.98 1/s
+++ total
   max dose is 0.0376496 in region 1880047
   ICRU-2K efficiency: 1086.98 1/s
_____
Completed 2 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
                 Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
```

```
Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,22.5,23.5)
______
______
        XYZ Geometry
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
-----
           Monte Carlo Parameter
-----
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                        10%
Max. 1st elastic moment per step :
                       0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                        0.0298764 0.420741
                     :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 18971 2927
_____
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
           = 1
photon split factor = −40
______
______
           Ouasi Random Numbers
______
number of generators: 1
```

```
1st: base = 2 dimensions = 60 warm-up = 1
.-----
______
              Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.323 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.033 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.052 70 70
+++finished batch 7 cpu time: 0.061 70 70
+++finished batch 8 cpu time: 0.071 70 70
+++finished batch 9 cpu time: 0.08 70 70
+++finished batch 10 cpu time: 0.09 70 70
finished simulation, cpu time = 0.096 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.096
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0388407 in region 1956847
   ICRU-2K efficiency: 1081.76 1/s
+++ total
   max dose is 0.0388407 in region 1956847
   ICRU-2K efficiency: 1081.76 1/s
_____
Completed 3 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
```

```
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
             Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,22.5,24)
_____
______
         XYZ Geometry
______
 name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
            Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       : 0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step :
                          0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           0.0298764 0.420741
Fano calculation
Exact Compton
                        : VMC++
Electron transport mode
______
______
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 8355 16407
______
```

```
______
             Variance Reduction
______
               = 0.251
f_repeat
split photons
photon split factor = -40
______
______
              Ouasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.308 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.06
                         70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.079 70 70
+++finished batch 10 cpu time: 0.088 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0374707 in region 2033647
```

```
ICRU-2K efficiency: 1162.9 1/s
+++ total
   max dose is 0.0374707 in region 2033647
   ICRU-2K efficiency: 1162.9 1/s
______
Completed 4 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get region size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,22.5,24.5)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
                              0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
                              7
CSDA approximation
Fractional energy loss/step at Ep :
                              10%
Max. 1st elastic moment per step :
                              0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           :
                              0.0298764 0.420741
Fano calculation
Exact Compton
                              0
```

```
Electron transport mode
                         VMC++
______
_____
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
             : 700
 total histories
 initial rng seeds : 28726 28947
______
.-----
           Variance Reduction
______
f repeat
             = 0.251
split photons = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.33 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70 70
```

```
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0380877 in region 2110447
   ICRU-2K efficiency: 1163.27 1/s
+++ total
   max dose is 0.0380877 in region 2110447
   ICRU-2K efficiency: 1163.27 1/s
_____
Completed 5 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,22.5,25)
_____
______
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
```

```
: 0.447479 MeV
Min. electron transport energy
Min. photon transport energy
                        0.05 MeV
                      :
Local track-end energy deposition :
                        0
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step :
                        0.6 MeV
                        0.0298764 0.420741
alpha and beta
                      :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 4729 29118
______
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 \text{ dimensions} = 60 \text{ warm-up} = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.284 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
```

```
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.051 70 70
+++finished batch 7 cpu time: 0.061 70 70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.08 70 70
+++finished batch 10 cpu time: 0.089 70 70
finished simulation, cpu time = 0.096 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.096
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0386789 in region 2187247
   ICRU-2K efficiency: 1187.56 1/s
   max dose is 0.0386789 in region 2187247
   ICRU-2K efficiency: 1187.56 1/s
______
Completed 6 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
                Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,22.5,25.5)
______
______
           XYZ Geometry
```

```
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
                      0.447479 MeV
Min. electron transport energy :
                    : 0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                       1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step
                      0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                       0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                      VMC++
______
_____
        MC\_Control
______
 will use fixed number of histories
 number of batches : 10
histories per batch : 70
 total histories : 700
 initial rng seeds : 28716 14562
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
```

```
dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.3 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.051 70 70
+++finished batch 7 cpu time: 0.061 70 70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.079 70 70
+++finished batch 10 cpu time: 0.089 70 70
finished simulation, cpu time = 0.095 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ==========
geometry: CT
cpu time: 0.095
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0385553 in region 2264047
   ICRU-2K efficiency: 1109.52 1/s
+++ total
   max dose is 0.0385553 in region 2264047
   ICRU-2K efficiency: 1109.52 1/s
_____
Completed 7 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
```

```
Initializing cross sections ... OK
______
           Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23,22.5)
______
______
        XYZ Geometry
-----
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
           Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                         0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                         1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                         10%
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
-----
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 24009 4257
______
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
            = 1
```

```
photon split factor = -40
-----
______
              Quasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
_____
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.279 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.033 70 70
+++finished batch 5 cpu time: 0.043 70 70
+++finished batch 6 cpu time: 0.053 70 70
+++finished batch 7 cpu time: 0.063 70 70
+++finished batch 8 cpu time: 0.075 70 70
+++finished batch 9 cpu time: 0.088 70 70
+++finished batch 10 cpu time: 0.099 70 70
finished simulation, cpu time = 0.106 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
geometry: CT
cpu time: 0.106
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0391293 in region 1803727
   ICRU-2K efficiency: 1047.97 1/s
+++ total
   max dose is 0.0391293 in region 1803727
   ICRU-2K efficiency: 1047.97 1/s
______
Completed 8 of 49 beamlets...
```

```
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
             Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23,23)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                            0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                            1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                           10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
                            0.0298764 0.420741
alpha and beta
Fano calculation
Exact Compton
                            VMC++
Electron transport mode
_____
           MC Control
______
```

```
will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 12653 27473
______
______
             Variance Reduction
______
f_repeat
               = 0.251
split photons
              = 1
photon split factor = -40
______
______
             Quasi Random Numbers
______
number of generators: 1
            1st: base = 2 dimensions = 60 warm-up = 1
______
______
           Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.352 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.027 70 70
+++finished batch 4 cpu time: 0.039 70 70
+++finished batch 5 cpu time: 0.052 70 70
+++finished batch 6 cpu time: 0.067 70 70
+++finished batch 7 cpu time: 0.078 70 70
+++finished batch 8 cpu time: 0.092
                          70 70
+++finished batch 9 cpu time: 0.108 70 70
+++finished batch 10 cpu time: 0.123 70 70
finished simulation, cpu time = 0.134 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
```

```
geometry: CT
cpu time: 0.134
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0385753 in region 1880527
   ICRU-2K efficiency: 825.064 1/s
+++ total
   max dose is 0.0385753 in region 1880527
   ICRU-2K efficiency: 825.064 1/s
_____
Completed 9 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23,23.5)
______
-----
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with XO = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy :
                             0.447479 MeV
Min. photon transport energy
                           :
                              0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
                           :
```

```
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step :
                        0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
            : 700
 total histories
 initial rng seeds : 23767 28785
_____
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.323 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
```

```
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.08 70 70
+++finished batch 10 cpu time: 0.089 70 70
finished simulation, cpu time = 0.096 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze ============
geometry: CT
cpu time: 0.096
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0378762 in region 1957327
   ICRU-2K efficiency: 1135.9 1/s
+++ total
   max dose is 0.0378762 in region 1957327
   ICRU-2K efficiency: 1135.9 1/s
_____
Completed 10 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
               Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23,24)
-----
______
           XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
```

```
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                      0.05 MeV
                   :
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                      1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                      10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                      0.0298764 0.420741
Fano calculation
                    :
Exact Compton
Electron transport mode
                      VMC++
-----
_____
        MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 19673 1072
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
  dump dose: 2
  dose scans:
```

CPU time so far: 1.411 seconds

```
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.079 70 70
+++finished batch 10 cpu time: 0.088 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0390439 in region 2034127
   ICRU-2K efficiency: 1179.01 1/s
+++ total
   max dose is 0.0390439 in region 2034127
   ICRU-2K efficiency: 1179.01 1/s
_____
Completed 11 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
                 Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
```

```
Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23,24.5)
______
______
        XYZ Geometry
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
-----
           Monte Carlo Parameter
-----
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                       1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                        10%
Max. 1st elastic moment per step :
                       0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                        0.0298764 0.420741
                     :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 25474 28020
_____
______
           Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = −40
______
______
           Ouasi Random Numbers
______
number of generators: 1
```

```
1st: base = 2 dimensions = 60 warm-up = 1
.-----
______
              Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.392 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0378966 in region 2110927
   ICRU-2K efficiency: 1212.35 1/s
+++ total
   max dose is 0.0378966 in region 2110927
   ICRU-2K efficiency: 1212.35 1/s
_____
Completed 12 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
```

```
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
             Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23,25)
_____
______
         XYZ Geometry
______
 name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
            Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       : 0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step :
                          0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           0.0298764 0.420741
Fano calculation
Exact Compton
                        : VMC++
Electron transport mode
______
______
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 20363 22733
______
```

```
______
             Variance Reduction
______
               = 0.251
f_repeat
split photons
photon split factor = -40
______
______
              Ouasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.334 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.052 70 70
+++finished batch 7 cpu time: 0.062
                           70 70
+++finished batch 8 cpu time: 0.072 70 70
+++finished batch 9 cpu time: 0.081 70 70
+++finished batch 10 cpu time: 0.091 70 70
finished simulation, cpu time = 0.1 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.1
number of histories: 700
number of batches: 10
+++ beamlet 0
  max dose is 0.0375397 in region 2187727
```

```
ICRU-2K efficiency: 1195.7 1/s
+++ total
   max dose is 0.0375397 in region 2187727
   ICRU-2K efficiency: 1195.7 1/s
______
Completed 13 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get region size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23,25.5)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
                              0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
                              7
CSDA approximation
Fractional energy loss/step at Ep :
                              10%
Max. 1st elastic moment per step :
                              0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           :
                              0.0298764 0.420741
Fano calculation
Exact Compton
                              0
```

```
Electron transport mode
                         VMC++
______
_____
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
             : 700
 total histories
 initial rng seeds : 22294 11767
______
.-----
           Variance Reduction
______
f repeat
             = 0.251
split photons = 1
photon split factor = -40
______
______
            Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.333 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041
                       70 70
+++finished batch 6 cpu time: 0.052
                       70 70
+++finished batch 7 cpu time: 0.062 70 70
+++finished batch 8 cpu time: 0.072 70 70
+++finished batch 9 cpu time: 0.081 70 70
+++finished batch 10 cpu time: 0.091 70 70
```

```
finished simulation, cpu time = 0.097 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.097
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0376161 in region 2264527
   ICRU-2K efficiency: 1130.64 1/s
+++ total
   max dose is 0.0376161 in region 2264527
   ICRU-2K efficiency: 1130.64 1/s
_____
Completed 14 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23.5,22.5)
_____
______
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
```

```
: 0.447479 MeV
Min. electron transport energy
Min. photon transport energy
                        0.05 MeV
                      :
Local track-end energy deposition :
                        0
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step :
                        0.6 MeV
                        0.0298764 0.420741
alpha and beta
                      :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 19665 5136
______
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 \text{ dimensions} = 60 \text{ warm-up} = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.296 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
```

```
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0382175 in region 1804207
   ICRU-2K efficiency: 1150.93 1/s
   max dose is 0.0382175 in region 1804207
   ICRU-2K efficiency: 1150.93 1/s
______
Completed 15 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
                Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23.5,23)
______
______
           XYZ Geometry
```

```
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
                      0.447479 MeV
Min. electron transport energy :
                    : 0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                       1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step
                      0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                       0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                      VMC++
______
_____
        MC\_Control
______
 will use fixed number of histories
 number of batches : 10
histories per batch : 70
 total histories : 700
 initial rng seeds : 21182 955
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
```

```
dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.317 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022
                                   70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.051 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.069 70 70
+++finished batch 9 cpu time: 0.079 70 70
+++finished batch 10 cpu time: 0.088 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ==========
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0376251 in region 1881007
   ICRU-2K efficiency: 1106.45 1/s
+++ total
   max dose is 0.0376251 in region 1881007
   ICRU-2K efficiency: 1106.45 1/s
_____
Completed 16 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
```

```
Initializing cross sections ... OK
______
           Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23.5,23.5)
______
______
        XYZ Geometry
-----
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 \, Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
           Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                         0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                         1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                         10%
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 8308 1386
______
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
            = 1
```

```
photon split factor = -40
-----
______
              Quasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.283 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.077
                           70 70
+++finished batch 10 cpu time: 0.086 70 70
finished simulation, cpu time = 0.092 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
geometry: CT
cpu time: 0.092
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0386409 in region 1957807
   ICRU-2K efficiency: 1132.18 1/s
+++ total
   max dose is 0.0386409 in region 1957807
   ICRU-2K efficiency: 1132.18 1/s
______
Completed 17 of 49 beamlets...
```

```
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
             Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23.5,24)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 \, Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                            0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                            1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                            10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
                            0.0298764 0.420741
alpha and beta
Fano calculation
Exact Compton
                            VMC++
Electron transport mode
_____
           MC Control
______
```

```
will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 2914 24704
______
______
             Variance Reduction
______
f_repeat
               = 0.251
split photons
              = 1
photon split factor = -40
______
______
             Quasi Random Numbers
______
number of generators: 1
            1st: base = 2 dimensions = 60 warm-up = 1
______
______
           Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.266 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
                         70 70
+++finished batch 8 cpu time: 0.068
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.086 70 70
finished simulation, cpu time = 0.092 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
```

```
geometry: CT
cpu time: 0.092
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0395884 in region 2034607
   ICRU-2K efficiency: 1202.83 1/s
+++ total
   max dose is 0.0395884 in region 2034607
   ICRU-2K efficiency: 1202.83 1/s
_____
Completed 18 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23.5,24.5)
______
-----
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with XO = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy :
                             0.447479 MeV
Min. photon transport energy
                           :
                              0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
                           :
```

```
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step :
                        0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
            : 700
 total histories
 initial rng seeds : 20845 9513
_____
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.292 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
```

```
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.069 70 70
+++finished batch 9 cpu time: 0.079 70 70
+++finished batch 10 cpu time: 0.088 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze ============
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0389809 in region 2111407
   ICRU-2K efficiency: 1147.91 1/s
+++ total
   max dose is 0.0389809 in region 2111407
   ICRU-2K efficiency: 1147.91 1/s
_____
Completed 19 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
               Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23.5,25)
-----
______
           XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
```

```
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                      0.05 MeV
                   :
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                      1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                      10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                      0.0298764 0.420741
Fano calculation
                    :
Exact Compton
Electron transport mode
                      VMC++
-----
_____
        MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 28507 1034
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
  dump dose: 2
  dose scans:
```

CPU time so far: 1.261 seconds

```
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ============
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0374017 in region 2188207
   ICRU-2K efficiency: 1232.57 1/s
+++ total
   max dose is 0.0374017 in region 2188207
   ICRU-2K efficiency: 1232.57 1/s
_____
Completed 20 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
                 Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
```

```
Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,23.5,25.5)
______
______
        XYZ Geometry
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
-----
           Monte Carlo Parameter
-----
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                        10%
Max. 1st elastic moment per step :
                       0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                        0.0298764 0.420741
                     :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 13163 11447
_____
______
           Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = −40
______
______
           Ouasi Random Numbers
______
number of generators: 1
```

```
1st: base = 2 dimensions = 60 warm-up = 1
.-----
______
              Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.29 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0381957 in region 2265007
   ICRU-2K efficiency: 1156.14 1/s
+++ total
   max dose is 0.0381957 in region 2265007
   ICRU-2K efficiency: 1156.14 1/s
_____
Completed 21 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
```

```
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
             Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24,22.5)
_____
______
         XYZ Geometry
______
 name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
            Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       : 0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step :
                          0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           0.0298764 0.420741
Fano calculation
Exact Compton
                        : VMC++
Electron transport mode
______
______
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 22966 23856
______
```

```
______
             Variance Reduction
______
               = 0.251
f_repeat
split photons
photon split factor = -40
______
______
              Ouasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.256 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058
                           70 70
+++finished batch 8 cpu time: 0.068
                          70 70
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.086 70 70
finished simulation, cpu time = 0.092 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ============
geometry: CT
cpu time: 0.092
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0369034 in region 1804687
```

```
ICRU-2K efficiency: 1099.72 1/s
+++ total
   max dose is 0.0369034 in region 1804687
   ICRU-2K efficiency: 1099.72 1/s
______
Completed 22 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get region size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24,23)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
                              0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
                              7
CSDA approximation
Fractional energy loss/step at Ep :
                              10%
Max. 1st elastic moment per step :
                              0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           :
                              0.0298764 0.420741
Fano calculation
Exact Compton
                              0
```

```
Electron transport mode
                         VMC++
______
_____
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories
 initial rng seeds : 5607 14693
______
.-----
           Variance Reduction
______
f repeat
             = 0.251
split photons = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.253 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.03 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.057 70 70
+++finished batch 8 cpu time: 0.067
                      70 70
+++finished batch 9 cpu time: 0.076 70 70
+++finished batch 10 cpu time: 0.085 70 70
```

```
finished simulation, cpu time = 0.091 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.091
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0379846 in region 1881487
   ICRU-2K efficiency: 1158.84 1/s
+++ total
   max dose is 0.0379846 in region 1881487
   ICRU-2K efficiency: 1158.84 1/s
_____
Completed 23 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24,23.5)
______
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
```

```
: 0.447479 MeV
Min. electron transport energy
Min. photon transport energy
                        0.05 MeV
                      :
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step :
                        0.6 MeV
                        0.0298764 0.420741
alpha and beta
                      :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 13368 19390
______
______
           Variance Reduction
______
            = 0.251
f_repeat
split photons
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 \text{ dimensions} = 60 \text{ warm-up} = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.279 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
```

```
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0380079 in region 1958287
   ICRU-2K efficiency: 1124.98 1/s
   max dose is 0.0380079 in region 1958287
   ICRU-2K efficiency: 1124.98 1/s
______
Completed 24 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
                Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24,24)
______
______
           XYZ Geometry
```

```
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
                      0.447479 MeV
Min. electron transport energy :
                    : 0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                       1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step
                      0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                       0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                      VMC++
______
_____
        MC\_Control
______
 will use fixed number of histories
 number of batches : 10
histories per batch : 70
 total histories : 700
 initial rng seeds : 21281 22641
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
```

```
dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.269 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.067 70 70
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.086 70 70
finished simulation, cpu time = 0.092 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ==========
geometry: CT
cpu time: 0.092
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0393287 in region 2035087
   ICRU-2K efficiency: 1185.05 1/s
+++ total
   max dose is 0.0393287 in region 2035087
   ICRU-2K efficiency: 1185.05 1/s
_____
Completed 25 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
```

```
Initializing cross sections ... OK
______
           Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24,24.5)
______
______
        XYZ Geometry
-----
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
           Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                         0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                         1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                         10%
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 8281 20392
______
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
            = 1
```

```
photon split factor = -40
-----
______
              Quasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.279 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0379302 in region 2111887
   ICRU-2K efficiency: 1129.03 1/s
+++ total
   max dose is 0.0379302 in region 2111887
   ICRU-2K efficiency: 1129.03 1/s
______
Completed 26 of 49 beamlets...
```

```
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
             Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24,25)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                            0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                            1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                           10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
                            0.0298764 0.420741
alpha and beta
Fano calculation
Exact Compton
                            VMC++
Electron transport mode
_____
           MC Control
______
```

```
will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 19653 4879
______
______
             Variance Reduction
______
f_repeat
               = 0.251
split photons
              = 1
photon split factor = -40
______
______
             Quasi Random Numbers
______
number of generators: 1
            1st: base = 2 dimensions = 60 warm-up = 1
______
______
           Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.277 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
                         70 70
+++finished batch 8 cpu time: 0.068
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
```

```
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0385447 in region 2188687
   ICRU-2K efficiency: 1150.91 1/s
+++ total
   max dose is 0.0385447 in region 2188687
   ICRU-2K efficiency: 1150.91 1/s
_____
Completed 27 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24,25.5)
______
-----
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with XO = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy :
                             0.447479 MeV
Min. photon transport energy
                           :
                              0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
                           :
```

```
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step :
                        0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
            : 700
 total histories
 initial rng seeds : 3570 14951
_____
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.32 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.033 70 70
```

```
+++finished batch 5 cpu time: 0.043 70 70
+++finished batch 6 cpu time: 0.054 70 70
+++finished batch 7 cpu time: 0.065 70 70
+++finished batch 8 cpu time: 0.076 70 70
+++finished batch 9 cpu time: 0.087 70 70
+++finished batch 10 cpu time: 0.097 70
finished simulation, cpu time = 0.105 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze ============
geometry: CT
cpu time: 0.105
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0382828 in region 2265487
   ICRU-2K efficiency: 975.757 1/s
+++ total
   max dose is 0.0382828 in region 2265487
   ICRU-2K efficiency: 975.757 1/s
_____
Completed 28 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
               Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24.5,22.5)
-----
______
           XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
```

```
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                      0.05 MeV
                   :
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                      1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                      10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                      0.0298764 0.420741
Fano calculation
                    :
Exact Compton
Electron transport mode
                      VMC++
-----
_____
        MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 28793 10212
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
  dump dose: 2
  dose scans:
```

CPU time so far: 1.273 seconds

```
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.051 70 70
+++finished batch 7 cpu time: 0.061 70 70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.08 70 70
+++finished batch 10 cpu time: 0.089 70 70
finished simulation, cpu time = 0.095 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ============
geometry: CT
cpu time: 0.095
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.037456 in region 1805167
   ICRU-2K efficiency: 1131.52 1/s
+++ total
   max dose is 0.037456 in region 1805167
   ICRU-2K efficiency: 1131.52 1/s
_____
Completed 29 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
                 Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
```

```
Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24.5,23)
______
______
        XYZ Geometry
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
-----
           Monte Carlo Parameter
-----
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                        10%
Max. 1st elastic moment per step :
                       0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                        0.0298764 0.420741
                     :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 17559 6715
_____
______
           Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = −40
______
______
           Ouasi Random Numbers
______
number of generators: 1
```

```
1st: base = 2 dimensions = 60 warm-up = 1
.-----
______
              Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.277 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.086 70 70
finished simulation, cpu time = 0.092 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.092
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0368158 in region 1881966
   ICRU-2K efficiency: 1158.46 1/s
+++ total
   max dose is 0.0368158 in region 1881966
   ICRU-2K efficiency: 1158.46 1/s
_____
Completed 30 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
```

```
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
             Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24.5,23.5)
_____
______
         XYZ Geometry
______
 name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
            Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       : 0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step :
                          0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           0.0298764 0.420741
Fano calculation
Exact Compton
                        : VMC++
Electron transport mode
______
______
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 22539 7653
______
```

```
______
             Variance Reduction
______
               = 0.251
f_repeat
split photons
photon split factor = -40
______
______
              Ouasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.288 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.069 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ============
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
  max dose is 0.038908 in region 1958767
```

```
ICRU-2K efficiency: 1217.45 1/s
+++ total
   max dose is 0.038908 in region 1958767
   ICRU-2K efficiency: 1217.45 1/s
______
Completed 31 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get region size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24.5,24)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
                              0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
                              7
CSDA approximation
Fractional energy loss/step at Ep :
                              10%
Max. 1st elastic moment per step :
                              0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           :
                              0.0298764 0.420741
Fano calculation
Exact Compton
                              0
```

```
Electron transport mode
                         VMC++
______
_____
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
             : 700
 total histories
 initial rng seeds : 15179 20973
______
.-----
           Variance Reduction
______
f repeat
             = 0.251
split photons = 1
photon split factor = -40
______
_____
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.278 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.067
                       70 70
+++finished batch 9 cpu time: 0.076 70 70
+++finished batch 10 cpu time: 0.086 70 70
```

```
finished simulation, cpu time = 0.092 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.092
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0375922 in region 2035567
   ICRU-2K efficiency: 1239.74 1/s
+++ total
   max dose is 0.0375922 in region 2035567
   ICRU-2K efficiency: 1239.74 1/s
_____
Completed 32 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24.5,24.5)
_____
______
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
```

```
: 0.447479 MeV
Min. electron transport energy
Min. photon transport energy
                        0.05 MeV
                      :
Local track-end energy deposition :
                        0
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step :
                        0.6 MeV
                        0.0298764 0.420741
alpha and beta
                      :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 26728 28779
______
______
           Variance Reduction
______
            = 0.251
f_repeat
split photons
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 \text{ dimensions} = 60 \text{ warm-up} = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.281 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
```

```
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.086 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0389452 in region 2112367
   ICRU-2K efficiency: 1240.86 1/s
   max dose is 0.0389452 in region 2112367
   ICRU-2K efficiency: 1240.86 1/s
______
Completed 33 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
                Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24.5,25)
______
______
           XYZ Geometry
```

```
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
                      0.447479 MeV
Min. electron transport energy :
                    : 0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                       1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step
                      0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                       0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                      VMC++
______
_____
        MC\_Control
______
 will use fixed number of histories
 number of batches : 10
histories per batch : 70
 total histories : 700
 initial rng seeds : 16417 4159
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
```

```
dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.275 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.03 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.069 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.088 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ==========
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0385814 in region 2189167
   ICRU-2K efficiency: 1171.89 1/s
+++ total
   max dose is 0.0385814 in region 2189167
   ICRU-2K efficiency: 1171.89 1/s
_____
Completed 34 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
```

```
Initializing cross sections ... OK
______
           Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,24.5,25.5)
______
______
        XYZ Geometry
-----
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
           Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                         0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                         1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                         10%
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 4479 7726
______
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
            = 1
```

```
photon split factor = -40
-----
______
              Quasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.286 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.033 70 70
+++finished batch 5 cpu time: 0.044 70 70
+++finished batch 6 cpu time: 0.053
                           70 70
+++finished batch 7 cpu time: 0.063 70 70
+++finished batch 8 cpu time: 0.073
                           70 70
+++finished batch 9 cpu time: 0.082 70 70
+++finished batch 10 cpu time: 0.092 70 70
finished simulation, cpu time = 0.098 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
geometry: CT
cpu time: 0.098
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0385168 in region 2265967
   ICRU-2K efficiency: 1123.54 1/s
+++ total
   max dose is 0.0385168 in region 2265967
   ICRU-2K efficiency: 1123.54 1/s
______
Completed 35 of 49 beamlets...
```

```
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
             Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25,22.5)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                            0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                            1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                           10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
                            0.0298764 0.420741
alpha and beta
Fano calculation
Exact Compton
                            VMC++
Electron transport mode
_____
           MC Control
______
```

```
will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 25222 7629
______
______
             Variance Reduction
______
f_repeat
               = 0.251
split photons
              = 1
photon split factor = -40
______
______
             Quasi Random Numbers
______
number of generators: 1
            1st: base = 2 dimensions = 60 warm-up = 1
______
______
           Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.286 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
                         70 70
+++finished batch 8 cpu time: 0.069
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
```

```
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0376269 in region 1805647
   ICRU-2K efficiency: 1145.14 1/s
+++ total
   max dose is 0.0376269 in region 1805647
   ICRU-2K efficiency: 1145.14 1/s
_____
Completed 36 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25,23)
______
-----
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with XO = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy :
                             0.447479 MeV
Min. photon transport energy
                           :
                              0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
                           :
```

```
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step :
                        0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
            : 700
 total histories
 initial rng seeds : 24429 7306
_____
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.275 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
```

```
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.068 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.087 70
finished simulation, cpu time = 0.093 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze ============
geometry: CT
cpu time: 0.093
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0380494 in region 1882447
   ICRU-2K efficiency: 1226.15 1/s
+++ total
   max dose is 0.0380494 in region 1882447
   ICRU-2K efficiency: 1226.15 1/s
_____
Completed 37 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
               Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25,23.5)
-----
______
           XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
```

```
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                      0.05 MeV
                   :
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                      1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                      10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                      0.0298764 0.420741
Fano calculation
                    :
Exact Compton
Electron transport mode
                      VMC++
-----
_____
        MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 27878 10500
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
  dump dose: 2
  dose scans:
```

CPU time so far: 1.3 seconds

```
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.059 70 70
+++finished batch 8 cpu time: 0.069 70 70
+++finished batch 9 cpu time: 0.079 70 70
+++finished batch 10 cpu time: 0.088 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0373512 in region 1959247
   ICRU-2K efficiency: 1128.34 1/s
+++ total
   max dose is 0.0373512 in region 1959247
   ICRU-2K efficiency: 1128.34 1/s
_____
Completed 38 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
                 Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
```

```
Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25,24)
______
______
        XYZ Geometry
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
-----
           Monte Carlo Parameter
-----
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                        10%
Max. 1st elastic moment per step :
                       0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                        0.0298764 0.420741
                     :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 5898 7533
_____
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
           = 1
photon split factor = −40
______
______
           Ouasi Random Numbers
______
number of generators: 1
```

```
1st: base = 2 dimensions = 60 warm-up = 1
.-----
______
              Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.29 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.067
+++finished batch 9 cpu time: 0.077 70 70
+++finished batch 10 cpu time: 0.086 70 70
finished simulation, cpu time = 0.092 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.092
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0394999 in region 2036047
   ICRU-2K efficiency: 1247.74 1/s
+++ total
   max dose is 0.0394999 in region 2036047
   ICRU-2K efficiency: 1247.74 1/s
_____
Completed 39 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
```

```
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
             Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25,24.5)
_____
______
         XYZ Geometry
______
 name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 \, Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
            Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       : 0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step :
                          0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           0.0298764 0.420741
Fano calculation
Exact Compton
                        : VMC++
Electron transport mode
______
______
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 18482 14199
______
```

```
______
             Variance Reduction
______
               = 0.251
f_repeat
split photons
photon split factor = -40
______
______
              Ouasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.277 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.021 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058
                           70 70
+++finished batch 8 cpu time: 0.068
                          70 70
+++finished batch 9 cpu time: 0.079
                          70 70
+++finished batch 10 cpu time: 0.09 70 70
finished simulation, cpu time = 0.096 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.096
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0376824 in region 2112847
```

```
ICRU-2K efficiency: 1101.39 1/s
+++ total
   max dose is 0.0376824 in region 2112847
   ICRU-2K efficiency: 1101.39 1/s
______
Completed 40 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get region size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25,25)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
                              0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
                              7
CSDA approximation
Fractional energy loss/step at Ep :
                              10%
Max. 1st elastic moment per step :
                              0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           :
                              0.0298764 0.420741
Fano calculation
Exact Compton
                              0
```

```
Electron transport mode
                         VMC++
______
_____
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
             : 700
 total histories
 initial rng seeds : 10550 24925
______
.-----
           Variance Reduction
______
f repeat
             = 0.251
split photons = 1
photon split factor = -40
______
_____
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.303 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.031 70 70
+++finished batch 5 cpu time: 0.04 70 70
+++finished batch 6 cpu time: 0.049 70 70
+++finished batch 7 cpu time: 0.058 70 70
+++finished batch 8 cpu time: 0.069 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.088 70 70
```

```
finished simulation, cpu time = 0.095 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.095
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0372181 in region 2189647
   ICRU-2K efficiency: 1160.01 1/s
+++ total
   max dose is 0.0372181 in region 2189647
   ICRU-2K efficiency: 1160.01 1/s
_____
Completed 41 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25,25.5)
_____
______
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
```

```
: 0.447479 MeV
Min. electron transport energy
Min. photon transport energy
                        0.05 MeV
                      :
Local track-end energy deposition :
                        0
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step :
                        0.6 MeV
                        0.0298764 0.420741
alpha and beta
                      :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 17558 16492
______
______
           Variance Reduction
______
            = 0.251
f_repeat
split photons
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 \text{ dimensions} = 60 \text{ warm-up} = 1
______
______
          Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.278 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
```

```
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.014 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.033 70 70
+++finished batch 5 cpu time: 0.043 70 70
+++finished batch 6 cpu time: 0.052 70 70
+++finished batch 7 cpu time: 0.062 70 70
+++finished batch 8 cpu time: 0.072 70 70
+++finished batch 9 cpu time: 0.081 70 70
+++finished batch 10 cpu time: 0.09 70 70
finished simulation, cpu time = 0.097 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.097
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0382575 in region 2266447
   ICRU-2K efficiency: 1060.1 1/s
   max dose is 0.0382575 in region 2266447
   ICRU-2K efficiency: 1060.1 1/s
______
Completed 42 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
                Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25.5,22.5)
______
______
           XYZ Geometry
```

```
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
                      0.447479 MeV
Min. electron transport energy :
                    : 0.05 MeV
Min. photon transport energy
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                       1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step
                      0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                       0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                      VMC++
______
_____
        MC\_Control
______
 will use fixed number of histories
 number of batches : 10
histories per batch : 70
 total histories : 700
 initial rng seeds : 27516 8576
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
```

```
dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.286 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.014 70 70
+++finished batch 3 cpu time: 0.024
                                   70 70
+++finished batch 4 cpu time: 0.034 70 70
+++finished batch 5 cpu time: 0.044 70 70
+++finished batch 6 cpu time: 0.056 70 70
+++finished batch 7 cpu time: 0.066 70 70
+++finished batch 8 cpu time: 0.076 70 70
+++finished batch 9 cpu time: 0.085 70 70
+++finished batch 10 cpu time: 0.095 70 70
finished simulation, cpu time = 0.101 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ==========
geometry: CT
cpu time: 0.101
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0365287 in region 1806127
   ICRU-2K efficiency: 1074.27 1/s
+++ total
   max dose is 0.0365287 in region 1806127
   ICRU-2K efficiency: 1074.27 1/s
_____
Completed 43 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
```

```
Initializing cross sections ... OK
______
           Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25.5,23)
______
______
        XYZ Geometry
-----
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 \, Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
           Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                         0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                         1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                         10%
Max. 1st elastic moment per step :
                         0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
-----
______
         MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 22717 22612
______
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
            = 1
```

```
photon split factor = -40
-----
______
              Quasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.286 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.052 70 70
+++finished batch 7 cpu time: 0.063 70 70
+++finished batch 8 cpu time: 0.072 70 70
+++finished batch 9 cpu time: 0.082 70 70
+++finished batch 10 cpu time: 0.091 70 70
finished simulation, cpu time = 0.097 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
geometry: CT
cpu time: 0.097
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0383494 in region 1882927
   ICRU-2K efficiency: 1082.79 1/s
+++ total
   max dose is 0.0383494 in region 1882927
   ICRU-2K efficiency: 1082.79 1/s
______
Completed 44 of 49 beamlets...
```

```
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
OK
Initializing cross sections ... OK
______
             Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25.5,23.5)
______
______
         XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 \, Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
             Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                            0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                            1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                            10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
                            0.0298764 0.420741
alpha and beta
Fano calculation
Exact Compton
                            VMC++
Electron transport mode
_____
           MC Control
______
```

```
will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 11414 17035
______
______
             Variance Reduction
______
f_repeat
               = 0.251
split photons
              = 1
photon split factor = -40
______
______
             Quasi Random Numbers
______
number of generators: 1
            1st: base = 2 dimensions = 60 warm-up = 1
______
______
           Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.275 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.012 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.033 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.053 70 70
+++finished batch 7 cpu time: 0.062 70 70
+++finished batch 8 cpu time: 0.072 70 70
+++finished batch 9 cpu time: 0.082 70 70
+++finished batch 10 cpu time: 0.091 70 70
finished simulation, cpu time = 0.098 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
```

```
geometry: CT
cpu time: 0.098
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0368155 in region 1959727
   ICRU-2K efficiency: 1038.24 1/s
+++ total
   max dose is 0.0368155 in region 1959727
   ICRU-2K efficiency: 1038.24 1/s
_____
Completed 45 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
              Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25.5,24)
______
-----
          XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with XO = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
              Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy :
                              0.447479 MeV
Min. photon transport energy
                           :
                              0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                              1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
                           :
```

```
Fractional energy loss/step at Ep :
Max. 1st elastic moment per step :
                        0.5
Max. acceptable energy loss/step :
                        0.6 MeV
alpha and beta
                        0.0298764 0.420741
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
______
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
            : 700
 total histories
 initial rng seeds : 2276 1619
_____
______
           Variance Reduction
______
f_repeat
            = 0.251
split photons
           = 1
photon split factor = -40
______
______
           Quasi Random Numbers
______
number of generators: 1
           1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.274 seconds
Will run approximately 700 particle sets
 with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.023 70 70
+++finished batch 4 cpu time: 0.032 70 70
```

```
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.051 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.07 70 70
+++finished batch 9 cpu time: 0.08 70 70
+++finished batch 10 cpu time: 0.09 70 70
finished simulation, cpu time = 0.097 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze ============
geometry: CT
cpu time: 0.097
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0382325 in region 2036527
   ICRU-2K efficiency: 1135.07 1/s
+++ total
   max dose is 0.0382325 in region 2036527
   ICRU-2K efficiency: 1135.07 1/s
_____
Completed 46 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
Parsing input file ... construct mmc geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
               Beamlet Source
______
     charge = 0
virtual source position = (-76,24,24)
     Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25.5,24.5)
-----
______
           XYZ Geometry
______
  name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 \, Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
```

```
______
          Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                      0.05 MeV
                   :
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                      1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                      10%
Max. 1st elastic moment per step : 0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                      0.0298764 0.420741
Fano calculation
                    :
Exact Compton
Electron transport mode
                      VMC++
-----
_____
        MC Control
_____
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 15924 23376
______
______
          Variance Reduction
______
f repeat
           = 0.251
split photons
           = 1
photon split factor = -40
______
______
          Quasi Random Numbers
______
number of generators: 1
          1st: base = 2 dimensions = 60 warm-up = 1
______
______
         Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
  dump dose: 2
  dose scans:
```

CPU time so far: 1.325 seconds

```
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.032 70 70
+++finished batch 5 cpu time: 0.041 70 70
+++finished batch 6 cpu time: 0.05 70 70
+++finished batch 7 cpu time: 0.06 70 70
+++finished batch 8 cpu time: 0.069 70 70
+++finished batch 9 cpu time: 0.078 70 70
+++finished batch 10 cpu time: 0.088 70 70
finished simulation, cpu time = 0.094 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.094
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.036899 in region 2113327
   ICRU-2K efficiency: 1140.63 1/s
+++ total
   max dose is 0.036899 in region 2113327
   ICRU-2K efficiency: 1140.63 1/s
_____
Completed 47 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam temp 1.vmc
Reading MS data ... OK
Parsing input file ... construct_mmc_geometry: urs = 1
DE_GeometryFactory::get_region_size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
                 Beamlet Source
______
      charge = 0
virtual source position = (-76,24,24)
```

```
Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25.5,25)
______
______
        XYZ Geometry
______
 name: CT id: 0
 global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
-----
           Monte Carlo Parameter
-----
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       0.05 MeV
Local track-end energy deposition :
Cut-off energy for KERMA approx. :
                        1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep :
                        10%
Max. 1st elastic moment per step :
                       0.5
Max. acceptable energy loss/step :
                       0.6 MeV
alpha and beta
                        0.0298764 0.420741
                     :
Fano calculation
Exact Compton
Electron transport mode
                        VMC++
______
_____
         MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 28021 3898
_____
______
           Variance Reduction
______
f repeat
            = 0.251
split photons
           = 1
photon split factor = −40
______
______
           Ouasi Random Numbers
______
number of generators: 1
```

```
1st: base = 2 dimensions = 60 warm-up = 1
.-----
______
              Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
    dump dose: 2
    dose scans:
CPU time so far: 1.263 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.003 70 70
+++finished batch 2 cpu time: 0.013 70 70
+++finished batch 3 cpu time: 0.022 70 70
+++finished batch 4 cpu time: 0.033 70 70
+++finished batch 5 cpu time: 0.042 70 70
+++finished batch 6 cpu time: 0.052 70 70
+++finished batch 7 cpu time: 0.062 70 70
+++finished batch 8 cpu time: 0.071 70 70
+++finished batch 9 cpu time: 0.08 70 70
+++finished batch 10 cpu time: 0.089 70 70
finished simulation, cpu time = 0.096 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE_ScoreDose::analyze =============
geometry: CT
cpu time: 0.096
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0377187 in region 2190127
   ICRU-2K efficiency: 1146.38 1/s
+++ total
   max dose is 0.0377187 in region 2190127
   ICRU-2K efficiency: 1146.38 1/s
_____
Completed 48 of 49 beamlets...
Input file is: C:\Users\wieserh\Documents\matlab\matRad\vmc++\runs/
MCpencilbeam_temp_1.vmc
Reading MS data ... OK
```

```
Parsing input file ... construct_mmc_geometry: urs = 1
DE GeometryFactory::get region size():
get_region_size: urs = 1 ignore = 1
region size: 0.3
geometry 1 region size: 0.3
Initializing cross sections ... OK
______
             Beamlet Source
______
    charge = 0
virtual source position = (-76,24,24)
    Energy = 6 MeV
number of beamlets = 1
beamlet 1: size = 0.5x0.5 cm**2 midpoint = (-26,25.5,25.5)
_____
______
         XYZ Geometry
______
 name: CT id: 0
  global smax: 1e+030
 Number of x-planes: 160 uniform with Xo = 0.15 \, Dx = 0.3
 Number of y-planes: 160 uniform with Yo = 0.15 Dx = 0.3
 Number of z-planes: 160 uniform with Zo = 0.15 Dx = 0.3
______
            Monte Carlo Parameter
______
Delta particle production threshold: 0.447479 MeV
Bremsstrahlung production threshold: 0.05 MeV
Min. electron transport energy : 0.447479 MeV
Min. photon transport energy
                       : 0.05 MeV
Local track-end energy deposition : 0
Cut-off energy for KERMA approx. : 1.10239 MeV
Bremsstrahlung transport mode
CSDA approximation
Fractional energy loss/step at Ep : 10%
Max. 1st elastic moment per step :
                          0.5
Max. acceptable energy loss/step : 0.6 MeV
alpha and beta
                           0.0298764 0.420741
Fano calculation
Exact Compton
                        : VMC++
Electron transport mode
______
______
          MC Control
______
 will use fixed number of histories
 number of batches : 10
 histories per batch : 70
 total histories : 700
 initial rng seeds : 17065 14082
______
```

```
______
             Variance Reduction
______
               = 0.251
f_repeat
split photons
photon split factor = -40
______
______
              Ouasi Random Numbers
______
number of generators: 1
             1st: base = 2 dimensions = 60 warm-up = 1
______
______
            Scoring and output options
______
 number of dose scoring objects: 1
 dose output options for geometry CT
   dump dose: 2
   dose scans:
CPU time so far: 1.286 seconds
Will run approximately 700 particle sets
  with 1 particles per set on average
will use 2 quasi numbers to sample the source
Starting MC simulation
Running part 1 of 1 ...
+++finished batch 1 cpu time: 0.004 70 70
+++finished batch 2 cpu time: 0.014 70 70
+++finished batch 3 cpu time: 0.024 70 70
+++finished batch 4 cpu time: 0.034 70 70
+++finished batch 5 cpu time: 0.043 70 70
+++finished batch 6 cpu time: 0.053 70 70
+++finished batch 7 cpu time: 0.063
                           70 70
+++finished batch 8 cpu time: 0.072 70 70
+++finished batch 9 cpu time: 0.082 70 70
+++finished batch 10 cpu time: 0.093 70 70
finished simulation, cpu time = 0.099 seconds
total particle fluence from the source: 2800
total number of particle sets: 700
total number of particles: 700
average sampled energy: 6 +/- 0
======= DE ScoreDose::analyze ===========
geometry: CT
cpu time: 0.099
number of histories: 700
number of batches: 10
+++ beamlet 0
   max dose is 0.0367436 in region 2266927
```

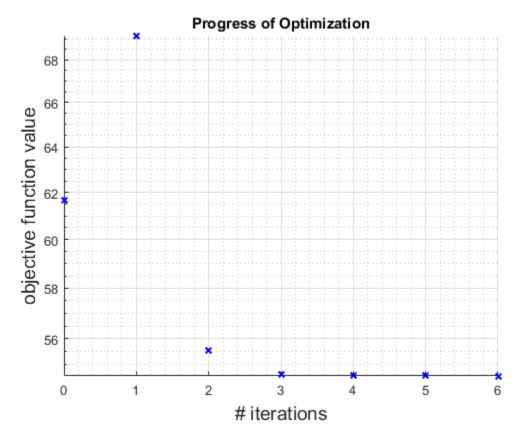
## **Inverse Optimization for IMRT**

The goal of the fluence optimization is to find a set of bixel/spot weights which yield the best possible dose distribution according to the clinical objectives and constraints underlying the radiation treatment.

```
resultGUI = matRad_fluenceOptimization(dij,cst,pln);
This is Ipopt version 3.11.8, running with linear solver ma57.
Number of nonzeros in equality constraint Jacobian...:
Number of nonzeros in inequality constraint Jacobian .:
                                                           0
Number of nonzeros in Lagrangian Hessian....:
                                                           0
Total number of variables....:
                                                          49
                    variables with only lower bounds:
                                                          49
               variables with lower and upper bounds:
                                                           0
                    variables with only upper bounds:
                                                           0
Total number of equality constraints....:
                                                           0
Total number of inequality constraints....:
       inequality constraints with only lower bounds:
                                                           0
   inequality constraints with lower and upper bounds:
       inequality constraints with only upper bounds:
iter
       objective
                    inf pr
                            inf_du lg(mu) ||d|| lg(rg) alpha_du
 alpha pr ls
  0 6.1682705e+001 0.00e+000 1.67e+001
                                        0.0 0.00e+000
                                                        - 0.00e
+000 0.00e+000
   1 6.9151634e+001 0.00e+000 2.06e+001 0.8 7.05e+000
 5.32e-001 1.51e-002f 5
  2 5.5544138e+001 0.00e+000 1.21e+001 0.3 5.67e-002
                                                           1.00e
+000 1.00e+000f 1
   3 5.4632309e+001 0.00e+000 1.53e+000 -1.0 2.01e-002
 9.98e-001 1.00e+000f 1
   4 5.4600721e+001 0.00e+000 1.07e+000 -2.7 4.81e-003
 9.98e-001 1.00e+000f 1
   5 5.4570186e+001 0.00e+000 3.25e-001 -4.3 1.28e-002
                                                        - 1.00e
+000 1.00e+000f 1
  6 5.4569142e+001 0.00e+000 4.20e-001 -5.9 4.60e-003
                                                        - 1.00e
+000 1.00e+000f 1
Number of Iterations....: 6
                                 (scaled)
                                                         (unscaled)
                         5.4569141774764034e+001
Objective....:
 5.4569141774764034e+001
Dual infeasibility....: 4.1999906083566546e-001
 4.1999906083566546e-001
```

```
Number of objective function evaluations = 15
Number of objective gradient evaluations = 7
Number of equality constraint evaluations = 0
Number of inequality constraint evaluations = 0
Number of equality constraint Jacobian evaluations = 0
Number of inequality constraint Jacobian evaluations = 0
Number of Lagrangian Hessian evaluations = 0
Total CPU secs in IPOPT (w/o function evaluations) = 0.313
Total CPU secs in NLP function evaluations = 0.559
```

EXIT: Solved To Acceptable Level. Calculating final cubes...

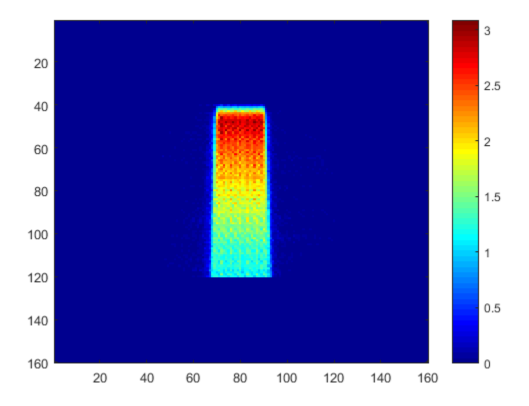


## **Plot the Resulting Dose Slice**

Just let's plot the transversal iso-center dose slice

```
slice = round(pln.isoCenter(1,3)./ct.resolution.z);
```

figure,
imagesc(resultGUI.physicalDose(:,:,slice)),colorbar, colormap(jet)



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