0.1 Intermediate Versions

0.1.1 L_{out} Optimization For Single e⁻

Initial step of improving POC towards RhodotronSimulation was to implement L_{out} optimizations to help optimizing magnet designs, as discussed in section ??. First approach was to hang the \mathbf{e}^- outside of the cavity for $t_{out} = L_{out}/v$, then inject it back to the cavity with reversed $\vec{\mathbf{v}}$. Then sweep the t_{out} parameter to find the optimal value. This simple implementation can be found in figure A.1 of $Appendix\ A$.

Although the results from this optimization sweep were promising after they were simulated with CST, simulating one particle would not be sufficiently useful for designing a magnet.

0.1.2 ϕ_{lag} Optimization For Bunches

After successfully accelerating single \mathbf{e}^- , particle bunches were implemented to approximate a real \mathbf{e}^- gun. They were modeled as N electrons fired from an \mathbf{e}^- gun at even time intervals. This approach was taken because the amount of time gun fires, defined as *Gun Active Time*, t_g , is a crucial part of pulsing \mathbf{e}^- gun design.

Addition of bunches would immediately proven useful when finding optimal gun phase lag. ϕ_{lag} for a bunch was defined as the RF phase when the first \mathbf{e}^- of the bunch entered the cavity, it defines the starting time of the current pass. To use the parameter sweep method, as used in L_{out} optimization, relevant bunch characteristics are defined as follows:

- μE : Average energy
- E_{rms} : Root mean square of energy
- R_{rms} : Root mean square of e^- positions

Optimal ϕ_{lag} would produce maximum μE while minimizing E_{rms} & R_{rms} . For the first pass, E_{rms} and R_{rms} would be vaguely dependent of each other; therefore, early implementation of ϕ_{lag} sweep was based on minimizing E_{rms} during simulation (see figure 1). For μE considerations, data from the software would be analyzed either manually or by using external tools such as ROOT.

```
int phase_opt(int phase_sweep_range){
         for(int RFphase = -phase_sweep_range; RFphase <= phase_sweep_range; RFphase++){</pre>
           Bunch bunch1(RFphase);
           double t1 = 0;
           bunch1.bunch_gecis_t(t1);
           bunch1.reset_pos();
10
           if( bunch1.E_rms() < minrms ){</pre>
11
              minrms = bunch1.E_rms();
              opt_phase = RFphase;
12
           }
13
14
         return opt_phase;
```

Figure 1: ϕ_{lag} Optimization For Initial Bunch Design

Since ϕ_{lag} is relatively easy to change after production, another version of figure 1 that was modified for given magnet design parameters was also implemented (see figure A.2 in Appendix A). This version can be useful for optimizing ϕ_{lag} in case of production issues in magnets.

After the bunch and ϕ_{lag} sweep implementations, L_{out} sweep was also updated to minimize E_{rms} . ρ and L calculations, using equations ?? and ??, were also integrated. Two example runs can be found in figures B.1 and B.2 of Appendix B.

0.1.3 Simulation in 3D

After successfully implementing \mathbf{e}^- - $\vec{\mathbf{E}}$ interaction in 1D and confirming the usefulness of this tool, the decision was made to proceed with implementing a 3D version of the *Rhodotron Simulation*. Although complete refactoring of the software was necessary, this upgrade was crucial for implementation of \mathbf{e}^- - $\vec{\mathbf{B}}$ interaction. The refactoring effort included proper implementation of OOP, details of which can be seen in *Appendix A*.

Magnets were modeled as major segments of a circle, defined with $\vec{\mathbf{r}}_{mag}$, \mathbf{R} and $|\mathbf{B}|$. For the initial implementation, $\vec{\mathbf{B}}$ assumed to be uniform and has no leaks outside the magnet boundary (See figure A.4 in Appendix A).

Interaction logic for $e^- - \vec{E}$ and $e^- - \vec{B}$ in 3D can be found in figure 2.

```
vector3d CoaxialRFField::actOn(Electron& e){
        vector3d Efield = getField(e.pos);
2
                                                                      // Calculate E vector
        vector3d F_m = Efield*1E6*eQMratio;
                                                                      // Calculate F/m vector
3
        vector3d acc = (F_m - e.vel*(e.vel*F_m)/(c*c))/e.gamma();
5
6
    }
    vector3d MagneticField::actOn(Electron& e){
        if (isInside(e.pos) == -1)
3
            return vector3d(0,0,0);
        vector3d Bfield = getField(e.pos);
                                                                      // Calculate B vector
        vector3d F_m = (e.vel % Bfield)*eQMratio;
                                                                         Calculate F/m vector
        vector3d acc = (F_m)/e.gamma();
                                                                      // Calculate a vector
6
```

Figure 2: e^- - EM interaction logic from equation ??

Where * and % are, dot-product and cross-product respectably. (See figure A.3 of Appendix A)

Simulating in 3D had one other benefit; it was now possible to visualize the results by rendering the interaction data. For this purpose, *gnuplot* was integrated into *Rhodotron Simulation* to produce 2D visualization of acceleration plane. Rendered results could be stored as *gif* animations. Two of such renders can be seen in *figure 3*.

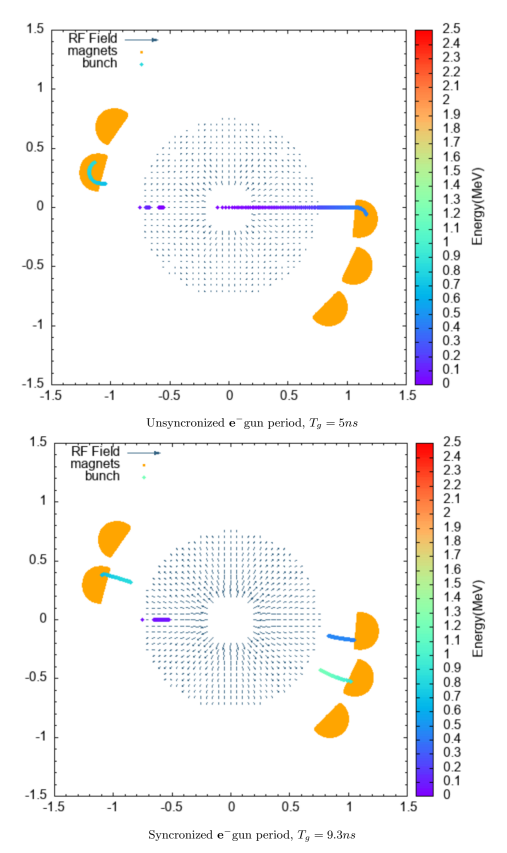


Figure 3: Example gnuplot renders of Rhodotron Simulation $P=12\mathrm{kW},\,f=107.5\mathrm{MHz}$

0.1.4 Acceleration in Magnetic Field

An issue regarding the \mathbf{e}^- - $\vec{\mathbf{B}}$ interaction became apparent when energy gain during these interactions was observed. A setup simulation was implemented in which a bunch of $100\mathbf{e}^-$ at 1MeV was fired to a uniform magnetic field of 0.1T placed in x>0.05m. Initial results at dt=0.01ns proved the suspicion of \mathbf{e}^- - $\vec{\mathbf{B}}$ interaction being broken. However, the energy gain would decrease tremendously as dt decreased.

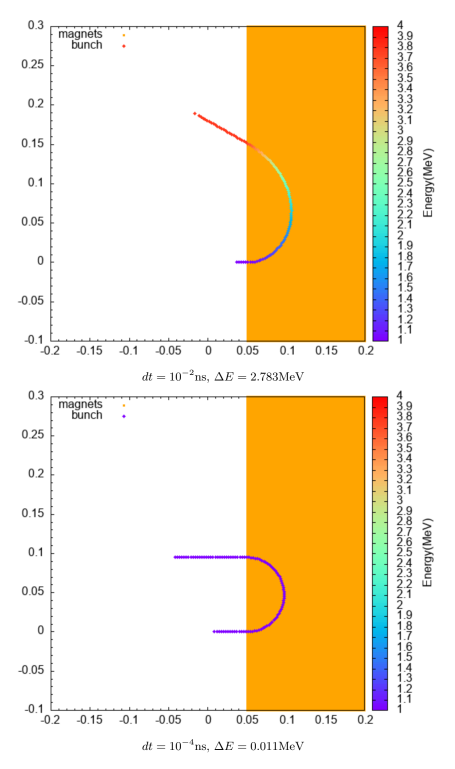


Figure 4: Energy gain of 1MeV bunch in **B**=0.1T

Decreasing dt would be the best way to increase accuracy of the results; however, this is not sustainable because time and computing power. Until this point, *Rhodotron Simulation* have been using *section* ?? for e^- - **EM** interactions. To test newer approaches, two additional version of e^- - **EM** interaction that are using *section* ?? were added.

RK4-1

First approach for integrating section ?? into \mathbf{e}^- - $\mathbf{E}\mathbf{M}$ interaction was to calculate $\vec{\mathbf{a}}_E$ and $\vec{\mathbf{a}}_B$ from equation ?? using RK4. After $\vec{\mathbf{a}}_{EM} = \vec{\mathbf{a}}_E + \vec{\mathbf{a}}_B$ was calculated, \mathbf{e}^- would move and accelerate using the Leap-frog method. The idea was to produce more refined interaction results, leading to improved accuracy especially in \mathbf{e}^- - $\vec{\mathbf{B}}$. RF field was kept static during the RK4 computation, due to ongoing multithreading implementation efforts. The implementation can be found in figures A.5 and A.6.

RK4-2

Following the implementation of RK4-1, revisions were made to the integration method for RK4 to replace Leap-frog. Instead of calculating $\vec{\mathbf{a}}_{EM}$ using RK4, $\vec{\mathbf{r}}$ and $\vec{\mathbf{v}}$ would be determined directly.

These three methods were then tested in the same setup as figure 4. The results can be found in figure 5.

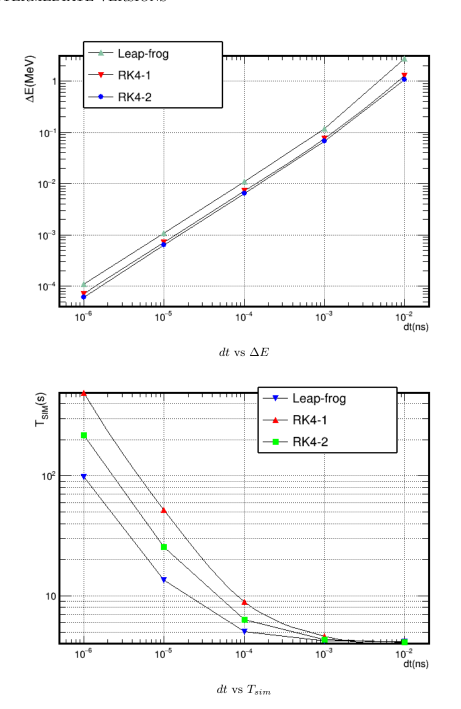


Figure 5: Comparing Leap-frog, RK4-1, RK4-2 performance on ${\bf e}^-$ - $\vec{\bf B}$ interaction $E_{in}=1 {\rm MeV},\, {\bf B}{=}0.1 {\rm T},\, t_{end}=5 {\rm ns}$

The data from these tests can be found in Tables B.1 and B.2 of Appendix B. As anticipated, Leap-frog produced the fastest yet least accurate results, whereas RK4-2 method was proved to be

the best performer for accuracy. When $dt = 10^-5$ ns was taken as reference point due to providing a good balance of accuracy and computational intensity, RK4-2 provides around 41% less energy gain in $\vec{\mathbf{B}}$ with respect to the next performer Leap-frog. The tradeoff is the computational intensity mentioned in section ??; Leap-frog being around 47% faster than RK4-2.

This relation is not linear however; RK4-2 is not as demanding in less computationally intensitive runs and able to keep up with Leap-frog in execution times while increasing the accuracy difference from other methods, whereas Leap-frog excels at simplicity, being able to use higher time resolution for given computational power. Both methods can be used in different situations, therefore both were integrated into $Rhodotron\ Simulation$ for the user to decide. RK4-2 renders from these test can be found in $figure\ B.3$ of $Appendix\ B$.

0.1.5 Acceleration in Electric Field

After the accuracy concerns regarding the \mathbf{e}^- - $\vec{\mathbf{B}}$ interaction were raised, it was decided to test \mathbf{e}^- - $\vec{\mathbf{E}}$ and compare the performance of *Leap-frog* and *RK4*.

As test setups, two simulation configurations were made. They aimed to test the accuracy of acceleration of a beam in parallel and perpendicular static uniform electric fields. Both configurations had an e^- -gunlocated at (-0.753, 0, 0)m, directed at (1, 0, 0) firing electrons with the kinetic energy of 1MeV.

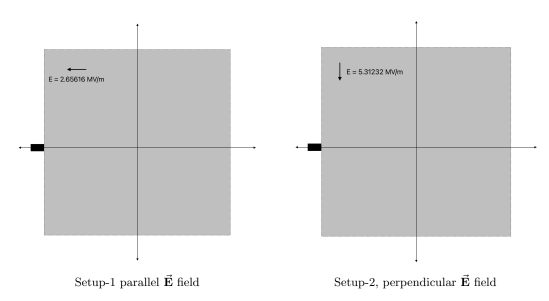


Figure 6: Illustration of test setups.

In the first test, the beam would be injected into a static uniform electric field $\vec{\mathbf{E}} = (-2.65616, 0, 0) \text{ MV/m}$ where -0.753 < x < 0.753 and -0.753 < y < 0.753, $\vec{\mathbf{E}} = 0$ elsewhere. Considering the $\vec{\mathbf{E}}$ is parallel to the beam path, potential difference V in the trejectory until

(-0.753, 0, 0, i, s)

$$\Delta V^1 = -\int \vec{\mathbf{E}} \cdot \vec{\mathbf{ds}}$$
 (1)

$$= -\int_{-0.753}^{0.753} -2.65616 \times dx \tag{2}$$

$$= 2.65616 \times 1.506 \tag{3}$$

$$\Delta V^1 = 4MV \tag{4}$$

$$\Delta E^1 = 4MeV \tag{5}$$

$$E_{exitTH}^{1} = 5MeV ag{6}$$

In the second test on the other hand, the beam would be injected into a different static uniform electric field,

 $\vec{\mathbf{E}} = (0, -5.31232, 0) \text{ MV/m where } -0.753 < x < 0.753 \text{ and } -0.753 < y < 0.753, \vec{\mathbf{E}} = 0 \text{ elsewhere.}$

$$\Delta V^2 = -\int \vec{\mathbf{E}} \cdot \vec{\mathbf{ds}} \tag{7}$$

$$= -\int_0^{0.753} -5.31232 \times dy \tag{8}$$

$$= 5.31232 \times 0.753 \tag{9}$$

$$\Delta V^2 = 4MV \tag{10}$$

$$\Delta E^2 = 4MeV \tag{11}$$

$$E_{exitTH}^2 = 5MeV (12)$$

Therefore, in the both tests, the beam was expected to exit $\vec{\mathbf{E}}$ with $E_{exitTH}=5$ MeV. To also measure the variance in simulation completion times, T_{sim} , set of 10 runs were completed at the configuration for each dt value.

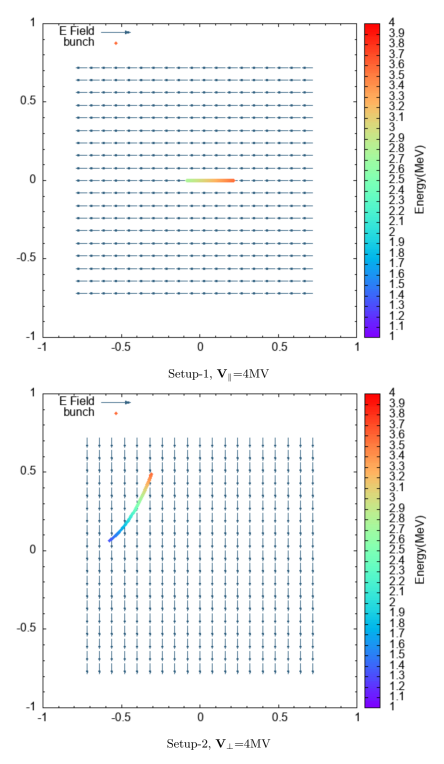
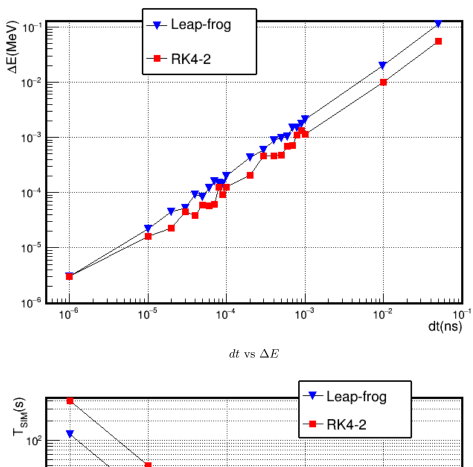


Figure 7: Render of the test setups. $E_{in} = 1 \ {\rm MeV}, \, t_{end} = 6 {\rm ns}$



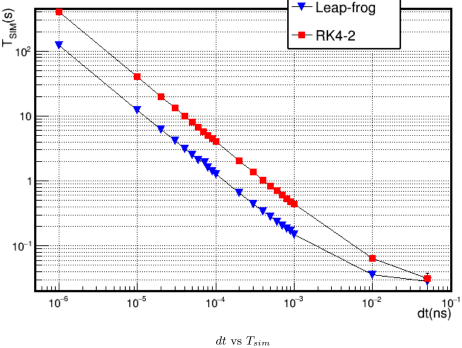


Figure 8: Comparing Leap-frog, RK4-2 performance on ${\bf e}^-$ - $\vec{\bf E}$ interaction $E_{in}=1{\rm MeV},~{\bf V}_\parallel{=}4{\rm MV},~t_{end}=6{\rm ns}$

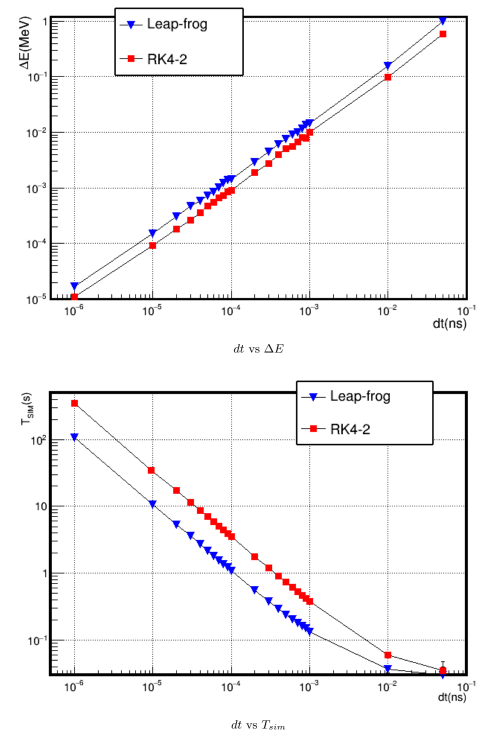


Figure 9: Comparing Leap-frog, RK4-2 performance on ${\bf e}^-$ - $\vec{\bf E}$ interaction $E_{in}=1{\rm MeV},\,{\bf V}_\perp{=}4{\rm MV},\,t_{end}=6{\rm ns}$

0.1.6 Multithreading

As already mentioned before, multithreading implementation efforts were ongoing since right after the start of this project. There have been a number of different approaches for implementation. First implementation was done when the *Rhodotron Simulation* was only capable of 1D simulations. After the sizable refactoring done for 3D capabilities, this implementation was obselete.

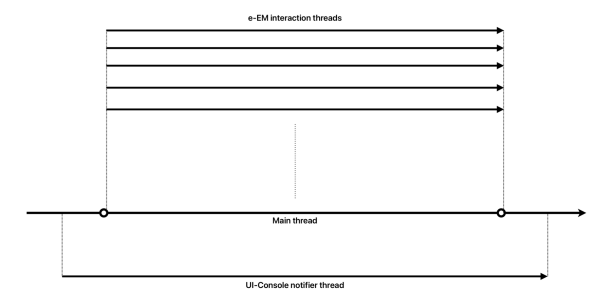


Figure 10: An Illustration of the multithreading architecture.

For the current version, a main thread that spawns and manages several other worker threads would be used as can be seen in *figure 10*. The UI-Console thread would handle incoming and outgoing communication, notifying the user about the status of simulation (see *figure B.5* in *Appendix*??) or communicating with the *GUI* that will be disscussed in the next section.

Focus of this section is the worker threads, also known as e^- - **EM** interaction threads. There were four competing architecture for these worker threads,

- 1. Have a thread pool that calculates e^- EM in a queue
- 2. Assign a thread to each bunch
- 3. Assign random electrons to each thread and calculate e^- EM with global time
- 4. Assign random electrons to each thread and calculate e^- EM with local thread time

Architecture 1 would require constant waiting in worker threads to get mutexes of $\vec{\mathbf{E}}$ field, especially in RK4.

Architecture 2 was performing well in configurations with a large number of bunches, but was not increasing performance in lower bunch count configurations as expected.

Architecture 3 was inefficient and wasteful since all the worker threads would wait the main thread to get the next time after they finish calculation, while the main thread would be waiting for the slowest worker thread.

Architecture 4 was thought to be the best performer. It would give freedom to calculate the whole simulation to each thread while giving up the global time. This would also mean thread-safety is ensured since there is no shared data between the worker threads. However, this architecture can cause issues if e^- - e^- interactions were decided to be implemented in the future.

Another caveat is that a copy operation of $\vec{\mathbf{E}}$ and $\vec{\mathbf{B}}$ objects for each worker thread would be needed. This would lead to larger memory allocations and more time spent setting up simulations. Therefore, *Architecture* 4 is not ideal for fast calculations and when the memory is an important constraint. An implementation that can use both **Architecture** 2 & **Architecture** 4 when necessary would be a better approach considering these methods; nevertheless, **Architecture** 4 was chosed to be implemented.

0.1.7 Graphical User Interface

Until this point, Rhodotron Simulation could be used with a configration file, defining the problem that would be simulated. An example of this configuration file can be found in figure B.4 of Appendix B. This approach was simple and fast; however, it was not suitable for the average target user since required basic knowledge of command line interface and was not up to modern standards. For this reason, a GUI was decided to be built, using ROOT framework. This would also enable Rhodotron simulation to make use of analysis tools offered by ROOT.

The initial design of the Rhodotron Simulation GUI can be observed in figure 11.

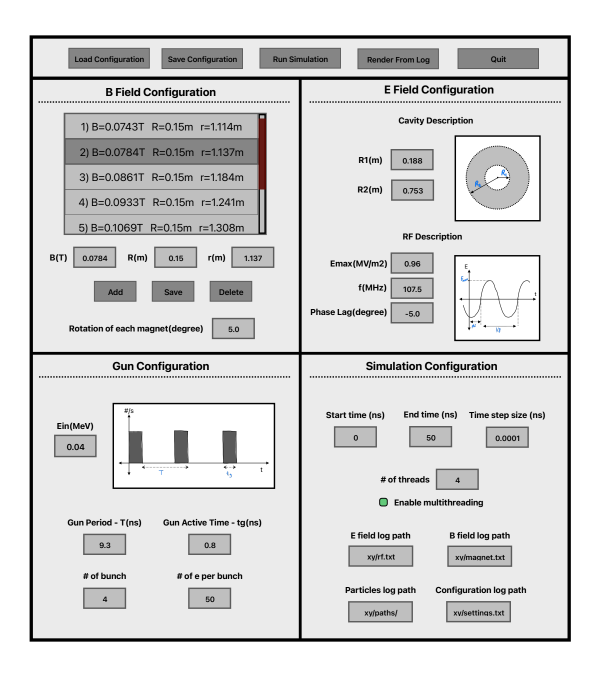


Figure 11: An Illustration of first GUI design.

The GUI would be a standalone application, running the $Rhodotron\ Simulation$ as a service when needed. For this reason, the now called $simulation\ engine$ was updated to be able to run as a background service of GUI. By this approach, one could also ignore the GUI altogether and use the $simulation\ engine$ as before, as these are two separate products.

Appendix A

Intermediate Codes

```
for(double i = 2; i < 9; i += dT_out){</pre>
           double enow = gecis(r_pos, vel, Et, t_dum);
if( enow > maxE ){
  maxE = enow;
              t_opt = i;
           t_dum = t;
 9
      double gecis(double r_pos, double vel, double Et, double &t){
           for(; r_pos >= -R2 && r_pos <= R2 ; t+=dT){
    vel = c*sqrt(Et*Et-E0*E0)/Et;</pre>
                double RelBeta = vel/c;
double RelGamma = 1.0 / sqrt(1.0-RelBeta*RelBeta);
                double ef=Eradial(r_pos*1000,t,RFphase*deg_to_rad);
                double acc=ef*1E6*eQMratio/(RelGamma*RelGamma*RelGamma);
10
                r_{pos} = r_{pos} + vel * dT*ns + 1/2*acc*(dT*ns)*(dT*ns);
11
                relevel+acc*dT*ns;
RelBeta = vel/c;
RelGamma = 1.0 / sqrt(1.0-RelBeta*RelBeta);
12
13
15
                Et=RelGamma*E0;
           return Et;
17
```

Figure A.1: L_{out} Optimization For Single e^-

```
int phase_opt(const vector<double>& Louts, int phase_sweep_range){
2
          double minrms = 1;
         int opt_phase;
for(int RFphase = -phase_sweep_range; RFphase <= phase_sweep_range; RFphase++){</pre>
3
              Bunch bunch1(RFphase);
              bunch1.bunch_gecis_t(t1);
              bunch1.reset_pos();
9
              for(int i = 0; i < Louts.size(); i++){</pre>
10
                  bunch1.bunch_gecis_d(Louts[i]);
11
                  bunch1.reset_pos();
              if( bunch1.E_rms() < minrms ){</pre>
15
                  minrms = bunch1.E_rms();
16
                  opt_phase = RFphase;
17
         }
20
         return opt_phase;
     }
21
```

Figure A.2: ϕ_{lag} Optimization For Initial Bunch Design

```
double vector3d::operator* (const vector3d& other){
    double dot = 0;
    dot += this->x * other.x;
    dot += this->y * other.y;
    dot += this->z * other.z;
    return dot;
}

vector3d vector3d::operator% (const vector3d& other){
    double x_ = (this->y * other.z) - (this->z * other.y);
    double y_ = (this->z * other.x) - (this->x * other.z);
    double z_ = (this->x * other.y) - (this->y * other.x);
    vector3d crossed(x_, y_, z_);
    return crossed;
}
```

Figure A.3: * and % operators of vector3d class

```
bool isInsideHalfSphere(vector3d e_position, double r, vector3d hs_position){

vector3d relative = e_position - hs_position;

// r/5 can be changed, use this for now

if ( relative.magnitude() <= r && relative * hs_position.direction() >= -r/5){

return true;

}

return false;

}
```

Figure A.4: Logic of is e⁻inside the shape of magnet

```
vector3d Electron2D::interactB_RK(const MagneticField& B, double time_interval){
          if (B.isInside(pos) == -1){
    return vector3d(0,0,0);
 2
3
           Electron2D e_dummy;
 5
           e_dummy.Et = Et;
           e_dummy.pos = pos;
e_dummy.vel = vel;
 9
           double time_halved = time_interval*0.5;
           // get k1
10
           vector3d F_m = (e_dummy.vel % B.getField(pos))*eQMratio;
11
           vector3d k1 = F_m * e_dummy.gamma_inv();
           // get k2
14
           e_dummy.move(time_halved);
          e_dummy.accelerate(k1, time_halved);
F_m = (e_dummy.vel % B.getField(pos))*eQMratio;
vector3d k2 = F_m * e_dummy.gamma_inv();
15
16
17
           // get k3
18
19
           e_dummy.vel = vel;
           e_dummy.accelerate(k2, time_halved);
           vector3d k3 = F_m * e_dummy.gamma_inv();
21
           // get k4
22
           e_dummy.vel = vel;
23
           e_dummy.move(time_halved);
24
           e_dummy.accelerate(k3, time_interval);
          F_m = (e_dummy.vel % B.getField(pos))*eQMratio;
vector3d k4 = F_m * e_dummy.gamma_inv();
26
27
28
           return (k1 + k2*2 + k3*2 + k4)/6;
29
30
```

Figure A.5: RK4-1 implementaion of e^- - \vec{B}

```
void Electron2D::interactRK_ActorE(const RFField& E, const MagneticField& B, double time_interval){
    vector3d run_kut_E = interactE_RK(E, time_interval);
    vector3d run_kut_B = interactB_RK(B, time_interval);

    vector3d acc = run_kut_E + run_kut_B;

    move(acc, time_interval/2);
    accelerate(acc, time_interval);
    move(acc, time_interval/2);
}
```

Figure A.6: RK4-1 implementaion of e^- - EM

Figure A.7: RK4-2 implemenation of \mathbf{e}^- - $\mathbf{E}\mathbf{M}$

Appendix B

Example Simulation Runs

```
Optimal phase with the least RMS : -5
     Simulation settings :
ph = -5 deg, gt = 1 ns, enum = 1000
      dT = 0.001 \text{ ns}, dT_{out} = 0.01 \text{ ns}
     For the 1th magnet:
Optimum out path = 0.81044 m
     Magnet guide = 0.25852 m
Rho = 0.088477 m
      Drift time of the first electron in the bunch : 7.688 ns
     Drift time of the last electron in the bunch : 7.487 ns
     Max energy = 0.47581 MeV
RMS = 0.0058165 MeV
15
     For the 2th magnet:
16
      Optimum out path = 1.0833 m
17
      Magnet guide = 0.37766 m
      Rho = 0.098898 m
     Drift time of the first electron in the bunch : 5.597 ns Drift time of the last electron in the bunch : 5.617 ns Max energy = 0.89172 MeV
20
      RMS = 0.0099018 MeV
      For the 3th magnet:
     Optimum out path = 1.1705 m
Magnet guide = 0.41573 m
      Rho = 0.10223 m
      Drift time of the first electron in the bunch : 5.314 ns
      Drift time of the last electron in the bunch : 5.325 \text{ ns}
      Max energy = 1.298 MeV
32
      RMS = 0.013879 MeV
      Electron with the most energy : 623) 1.6999 MeV,
                                                                           RMS of bunch: 0.017981 MeV
      Total steps calculated: 12468052652
      Simulation finished in : 632050015 us
                                                          (632.1 s)
```

Figure B.1: $\phi_{lag},~\rho~\&~L$ optimization at $P=12{\rm KW},~R_1=0.188{\rm m},~R_2=0.753{\rm m},~t_g=1{\rm ns},~E_{in}=40{\rm KeV}$

```
Optimal phase with the least {\tt RMS} : {\tt O}
     Simulation settings :  ph = 0 \ deg, \ gt = 0.8 \ ns, \ enum = 1000 \\ dT = 0.001 \ ns, \ dT\_out = 0.01 \ ns 
      For the 1th magnet:
      Optimum out path = 0.80787 m
      Magnet guide = 0.2574 \text{ m}
      Rho = 0.088379 \text{ m}
10
      Drift time of the first electron in the bunch : 7.629 ns
11
      Drift time of the last electron in the bunch : 7.48 \text{ ns}
      Max energy = 0.47579 MeV
      RMS = 0.0038689 MeV
15
      For the 2th magnet:
16
     Optimum out path = 1.0833 m
Magnet guide = 0.37765 m
17
      Rho = 0.098898 m
      Drift time of the first electron in the bunch : 5.589 \ \mathrm{ns}
21
      Drift time of the last electron in the bunch : 5.605 \ \text{ns}
     Max energy = 0.89169 MeV
RMS = 0.0068848 MeV
22
23
      For the 3th magnet:
      Optimum out path = 1.1705 m
      Magnet guide = 0.41573 m
      Rho = 0.10223 m
28
      Drift time of the first electron in the bunch : 5.311 ns
29
      Drift time of the last electron in the bunch : 5.318 ns
      Max energy = 1.298 MeV
      RMS = 0.0096887 MeV
      Electron with the most energy : 629) 1.6999 MeV,
                                                                        RMS of bunch: 0.012318 MeV
35
      Total steps calculated : 12455378454
36
      Simulation finished in: 631136046 us
                                                        (631.1 s)
```

Figure B.2: ϕ_{lag} & ρ & L optimization at $P=12{\rm KW},~R_1=0.18{\rm 8m},~R_2=0.753{\rm m},~t_g=0.8{\rm ns},~E_{in}=40{\rm KeV}$

dt(ns)	$T_{simLF}(s)$	$T_{simRK1}(s)$	$T_{simRK2}(s)$	ΔT_{LF-RK2}
10^{-2}	4.14092	4.146	4.10945	0.77%
10^{-3}	4.23817	4.61703	4.30225	-1.51%
10^{-4}	5.04679	8.93529	6.3065	-24.96%
10^{-5}	13.5135	51.9024	25.4279	-88.17%
10^{-6}	97.6229	486.378	217.704	-123.01%

Table B.1: dt vs T_{sim} of Leap-frog, RK4-1, RK4-2 $E_{in}=1 \text{MeV}, \ \mathbf{B}{=}0.1 \text{T}, \ t_{end}=5 \text{ns}$

dt(ns)	$\Delta E_{LF}({ m MeV})$	$\Delta E_{RK1}({ m MeV})$	$\Delta E_{RK2}({ m MeV})$	ΔE_{LF-RK2}
10^{-2}	2.78323	1.26503	1.04713	165.8%
10^{-3}	0.11712	0.07469	0.06691	42.87%
10^{-4}	0.01101	0.00716	0.00644	41.5%
10^{-5}	0.00109	0.00071	0.00064	41%
10^{-6}	0.00011	0.00007	0.00006	45%

Table B.2: dt vs ΔE of Leap-frog, RK4-1, RK4-2 $E_{in} = 1 \mathrm{MeV}, \, \mathbf{B}{=}0.1 \mathrm{T}, \, t_{end} = 5 \mathrm{ns}$

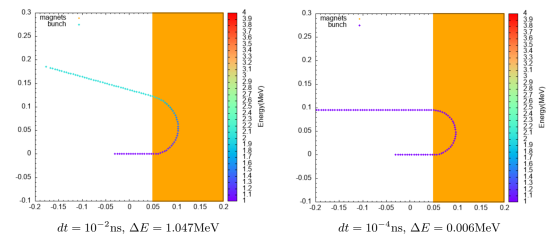


Figure B.3: Energy gain of 1MeV bunch in $\mathbf{B}{=}0.1\mathrm{T}$ using RK4-2

Figure B.4: ϕ_{lag} & ρ & L optimization at $P=12{\rm KW},\,R_1=0.188{\rm m},\,R_2=0.753{\rm m},\,t_g=0.8{\rm ns},\,E_{in}=40{\rm KeV}$

Figure B.5: Example of console output while simulation is running.