Magnetic Mirror Effect in Magnetron Plasma: Modeling of Plasma Parameters

1 Checks

To verify that our program is working correctly, we perform some checks. We compare results from the program with calculations done by hand; of certain quantities and see if they agree. As of now, we are doing checks on the following aspects of the program:

- 1. Sampling
- 2. Update ✓

1.1 Sampling

As of now we our sampling of particle speeds is based on the Maxwell-Boltzmann distribution, that we have defined previously. We now check if the average speeds of the particles agrees with the calculations done by hand based on the plasma temperature. We recall the Maxwellian density function that we defined earlier.

$$\widehat{f_M} := \widehat{f}(\boldsymbol{x}, \boldsymbol{v}, t) = \left(\frac{m}{2\pi KT}\right)^{\frac{3}{2}} \exp\left(-\frac{\boldsymbol{v}^2}{v_{th}^2}\right)$$
(1)

where

$$v_{th}^2 = \frac{2KT}{m}$$

For our convenience, we use the density function with speed instead of velocity which is defined as:

$$\widehat{f_m} := \widehat{f}(\boldsymbol{x}, v, t) = \left(\frac{m}{2\pi KT}\right)^{\frac{3}{2}} 4\pi v^2 \exp\left(-\frac{v^2}{v_{th}^2}\right)$$
(2)

which is obtained by integrating over the solid angle in the velocity variable.

For this density function we calculate the average speed by with the expression:

$$\langle v \rangle = \int_{v=-\infty}^{v=\infty} dv \, v \, \widehat{f_m} = \int_{v=-\infty}^{v=\infty} dv \, v \, \left(\frac{m}{2\pi KT}\right)^{\frac{3}{2}} 4\pi v^2 \exp\left(-\frac{v^2}{v_{th}^2}\right)$$

$$= 4\pi \left(\frac{m}{2\pi KT}\right)^{\frac{3}{2}} \int_{v=-\infty}^{v=\infty} dv \, v^3 \exp\left(-\frac{v^2}{v_{th}^2}\right)$$

$$= 4\pi \left(\frac{m}{2\pi KT}\right)^{\frac{3}{2}} \frac{v_{th}^4}{2} = 4\pi \left(\frac{1}{\pi v_{th}^2}\right)^{\frac{3}{2}} \frac{v_{th}^4}{2}$$

$$= \frac{2}{\sqrt{\pi}} v_{th} = \frac{2}{\sqrt{\pi}} \sqrt{\frac{2KT}{m}} = \frac{2}{\sqrt{\pi}} \sqrt{\frac{2RT}{M}}$$

For the Hydrogen atom particle distribution, we get

```
\langle v \rangle = \frac{2}{\sqrt{\pi}} \sqrt{\frac{2 \cdot 8.31446261815324 \text{ J K}^{-1} \text{ mol}^{-1} 10000 \text{ K}}{1.008 \times 10^{-3} \text{ kg mol}^{-1}}} = 14492.952993825973 \text{ ms}^{-1}
```

From section 1.1.1 Maxwellian sampling in the **studies.ipynb** notebook, we take the following:

Initialization

```
# c1Maxwell means checking the Maxwellian sampling
c1Maxwell = Run()
# Create 100 particles based on the data available in the files
c1Maxwell.create_batch_with_file_initialization('H+', constants.constants['e'][0],

constants.constants['m_H'][0] * constants.constants['amu'][0], 100, 100, 'H ions',

r_index=0, v_index=1)
```

Inspection

```
# Take the Oth batch of particles
   c1Maxwell_batch = c1Maxwell.batches[0]['H ions']
   # Take the initial positions and velocities of the particles
3
   c1Maxwell_positions = []
4
   c1Maxwell_velocities = []
   for particle in c1Maxwell_batch.particles:
   c1Maxwell_positions.append(particle.r)
   c1Maxwell_velocities.append(particle.v)
   # Let's now look at the velocities
   c1Maxwell velocities
10
   # We need to check if they are really Maxwellian distributed
11
   # Get the speeds
12
   c1Maxwell_speeds = np.sqrt( [ (c1Maxwell_velocities[i][0] ** 2) +
    → range(len(c1Maxwell_velocities)) ] )
   c1Maxwell_meanspeed = np.sum(c1Maxwell_speeds) / c1Maxwell_speeds.size
```

We get an average speed of the distribution to be:

14202.764572898674

We see that the mean average speed of the sampled distribution and that expected from the analytic expression are close; in fact within 2.0431826454479785% error.

To be confident that our program indeed samples particles based on Maxwellian distribution as we expect it to, let's check another distribution. For convenience, let's assume Hydrogen molecules to be particles sampled with Maxwellian distribution, and check if the average speed of the sampled distribution agrees with the that expected from analytic calculation. For this we follow a similar procedure.

Initialization

```
# Create 100 particles based on the sampled velocities of H2 gas

# We consider just for this test case that a Hydrogen molecules to be sampled with a

Maxwellian distribution

# We create a new batch on the same Run instance

c1Maxwell.create_batch_with_file_initialization('H2', constants.constants['e'][0], 2 *

constants.constants['m_H'][0] * constants.constants['amu'][0], 100, 100, 'H2 gas',

r_index=0, v_index=2)
```

Inspection

```
# Do the same as before for the second batch
   c1Maxwell_batch2 = c1Maxwell.batches[1]['H2 gas']
2
   c1Maxwell_positions_batch2 = []
3
4
   c1Maxwell_velocities_batch2 = []
   for particle in c1Maxwell_batch2.particles:
5
   c1Maxwell_positions_batch2.append(particle.r)
   c1Maxwell_velocities_batch2.append(particle.v)
   c1Maxwell_speeds_batch2 = np.sqrt( [ (c1Maxwell_velocities_batch2[i][0] ** 2) +
      (c1Maxwell_velocities_batch2[i][1] ** 2) + (c1Maxwell_velocities_batch2[i][2] ** 2)
    → for i in range(len(c1Maxwell_velocities_batch2)) ] )
   c1Maxwell_meanspeed_batch2 = np.sum(c1Maxwell_speeds_batch2) /

→ c1Maxwell_speeds_batch2.size
```

We get an average speed of the distribution to be:

```
10149.754879907316
```

For the Hydrogen molecule particle distribution, we get

$$\langle v \rangle = \frac{2}{\sqrt{\pi}} \sqrt{\frac{2 \cdot 8.31446261815324 \text{ J K}^{-1} \text{ mol}^{-1} 10000 \text{ K}}{2 \times 1.008 \times 10^{-3} \text{ kg mol}^{-1}}} = 10248.06534135222 \text{ ms}^{-1}$$

We see that the mean average speed of the sampled distribution and that expected from the analytic expression are close for this distribution too; in fact within 0.9685993662716086%error. Because the errors are considerably small compared to statistical variation in the sampling, we believe the Maxwellian distribution sampling part of the program to be working as we expect it to.

1.2 Update

We recall the Boris Algorithm, that we use to update the particles in the plasma simulation.

$$\mathbf{v}^{-} = \mathbf{v}_{k} + q' \mathbf{E}_{k}$$

$$\mathbf{v}^{+} = \mathbf{v}^{-} + 2q' \left(\mathbf{v}^{-} \times \mathbf{B}_{k}\right)$$

$$\mathbf{v}_{k+1} = \mathbf{v}^{+} + q' \mathbf{E}_{k}$$

$$\mathbf{x}_{k+1} = \mathbf{x}_{k} + \Delta t \ \mathbf{v}_{k+1}$$
(3)

where $q'=rac{q}{m}rac{\Delta t}{2}.$ To verify that the simulation works as we expect it to, we perform a calculation and compare it to the output of an algorithm. From section 1.2.1 Boris Update in the stud**ies.ipynb** notebook, we take the following:

Initialization

```
# c2Boris means checking the Boris update
c2Boris = Run()
#Create 10 particles
c2Boris.create_batch_with_file_initialization('H+', constants.constants['e'][0],

→ constants.constants['m_H'][0] * constants.constants['amu'][0], 100, 10, 'H ions',

      r_index=0, v_index=1)
```

Update input data

```
c2Boris_index_update = 0 # Update the first batch in this Run instance run_Boris_check
2
    c2Boris_particle_track_indices = [i for i in range(10)] # Track all 10 particles
3
    c2Boris dT = 10**(-6) # 1 microseconds
    c2Boris_stepT = 10**(-7) # 0.1 microseconds time step
5
    c2Boris_E0 = 1000 # say 1000 Volts (voltage) per meter (size of chamber)
6
    c2Boris_Edirn = [1,0,0] #in the x-direction [1,0,0]
    c2Boris_B0 = 10 * (10**(-3)) # Meant to say 10 mT
    c2Boris_Bdirn = [0,1,0] #in the y-direction [0,1,0]
9
    c2Boris_argsE = [element * c2Boris_E0 for element in c2Boris_Edirn] # currently the
10
    → uniform E field configuration is used
    c2Boris_argsB = [element * c2Boris_B0 for element in c2Boris_Bdirn] # currently the
11
    → uniform_B_field configuration is used
```

Update

```
c2Boris_positions_and_velocities =

→ c2Boris_update_batch_with_unchanging_fields(c2Boris_index_update, c2Boris_dT,
       c2Boris_stepT, c2Boris_argsE, c2Boris_argsB, c2Boris_particle_track_indices)
2
    #Let's inspect the positions and velocities of the particles at index 1 and 7
3
    c2Boris_p1 = c2Boris_positions_and_velocities[1]
4
    c2Boris_p7 = c2Boris_positions_and_velocities[7]
5
    #Let's take the positions and velocities of the particle 1 after the 3rd and the 4th
7
    \hookrightarrow time steps.
    c2Boris_p134_p = [c2Boris_p1[3][1], c2Boris_p1[4][1]]
8
    c2Boris_p134_v = [c2Boris_p1[3][2], c2Boris_p1[4][2]]
9
10
    #Similary for particle 7
11
    c2Boris_p734_p = [c2Boris_p7[3][1], c2Boris_p7[4][1]]
12
   c2Boris_p734_v = [c2Boris_p7[3][2], c2Boris_p7[4][2]]
13
```

The program gives the following output. For example, we can check the particles 1 and 7 in the update. After the 3^{rd} step of the update, the velocity of particle 1 was:

```
[39771.83801956555, -5029.491038, 4533.343932509505]
```

and after the 4^{th} update, it changed to:

```
[48909.86581773698, -5029.491038, 8798.399243869582]
```

Its position after the 3^{rd} step of the update was:

[-0.48985112694809785, -0.0020117964152, 0.00017005183797547688]

which was updated after the 4^{th} step of the update to:

[-0.4849601403663242, -0.0025147455189999997, 0.001049891762362435]

We now check if the same if true, with a calculation of the update done by hand.

$$q' = \frac{q}{m} \frac{\Delta t}{2} = \frac{1.602176634 \times 10^{-19} \,\mathrm{C}}{1.008 \,\mathrm{a.m.u.} \times 1.6605390666 \times 10^{-27} \,\mathrm{kg \,a.m.u.}^{-1}} \frac{10^{-7} \,\mathrm{s}}{2} = 4.78597877761177 \,\mathrm{Cskg}^{-1}$$

$$\boldsymbol{v}^- = \boldsymbol{v}_3 + q'\mathbf{E} = \begin{bmatrix} 39771.83801956555 \\ -5029.491038 \\ 4533.343932509505 \end{bmatrix} + 4.78597877761177 \begin{bmatrix} 1000 \\ 0 \\ 0 \end{bmatrix} = \begin{bmatrix} 44557.81679718 \\ -5029.491038 \\ 4533.34393251 \end{bmatrix}$$

$$\boldsymbol{v}^{+} = \boldsymbol{v}^{-} + 2q' \left(\boldsymbol{v}^{-} \times \mathbf{B} \right) = \begin{bmatrix} 44557.81679718 \\ -5029.491038 \\ 4533.34393251 \end{bmatrix} + 2 \cdot 4.78597877761177 \left(\begin{bmatrix} 44557.81679718 \\ -5029.491038 \\ 4533.34393251 \end{bmatrix} \times \begin{bmatrix} 0.0 \\ 0.01 \\ 0.0 \end{bmatrix} \right)$$

$$= \begin{bmatrix} 44123.88704013 \\ -5029.491038 \\ 8798.39924387 \end{bmatrix}$$

$$\mathbf{v}_4 = \mathbf{v}^+ + q'\mathbf{E} = \begin{bmatrix} 44123.88704013 \\ -5029.491038 \\ 8798.39924387 \end{bmatrix} + 4.78597877761177 \begin{bmatrix} 1000 \\ 0 \\ 0 \end{bmatrix} = \begin{bmatrix} 48909.86581774 \\ -5029.491038 \\ 8798.39924387 \end{bmatrix}$$

$$\boldsymbol{x}_4 = \boldsymbol{x}_3 + \Delta t \ \boldsymbol{v}_4 = \begin{bmatrix} -4.89851127 \times 10^{-01} \\ -2.01179642 \times 10^{-03} \\ 1.70051838 \times 10^{-04} \end{bmatrix} + 10^{-7} \ \begin{bmatrix} 48909.86581774 \\ -5029.491038 \\ 8798.39924387 \end{bmatrix} = \begin{bmatrix} -0.48496014 \\ -0.00251475 \\ 0.00104989 \end{bmatrix}$$

We see that the values for x_3 , x_4 , v_3 and v_4 for particle 1 are close (the latter digits differ due to the differing precision of calculations); i.e. the same when done by hand and in the program.

Similarly, for particle 7. After the 3^{rd} step of the update, the velocity of the particle was:

[38895.85871094631, -8670.854777, 13914.969423620274]

and after the 4^{th} update, it changed to:

[47135.881299118584, -8670.854777, 18096.176367366777]

Its position after the 3^{rd} step of the update was:

[-0.4896669752432719, -0.0034683419108, 0.0038875496903932644]

which was updated after the 4^{th} step of the update to:

[-0.48495338711336006, -0.0043354273885, 0.0056971673271299424]

$$\begin{aligned} \boldsymbol{v}^- &= \boldsymbol{v}_3 + q'\mathbf{E} = \begin{bmatrix} 38895.85871095 \\ -8670.854777 \\ 13914.96942362 \end{bmatrix} + 4.78597877761177 \begin{bmatrix} 1000 \\ 0 \\ 0 \end{bmatrix} = \begin{bmatrix} 43681.83748856 \\ -8670.854777 \\ 13914.96942362 \end{bmatrix} \\ \boldsymbol{v}^+ &= \boldsymbol{v}^- + 2q' \left(\boldsymbol{v}^- \times \mathbf{B} \right) = \begin{bmatrix} 43681.83748856 \\ -8670.854777 \\ 13914.96942362 \end{bmatrix} + 2 \cdot 4.78597877761177 \left(\begin{bmatrix} 43681.83748856 \\ -8670.854777 \\ 13914.96942362 \end{bmatrix} \times \begin{bmatrix} 0.0 \\ 0.01 \\ 0.0 \end{bmatrix} \right) \\ &= \begin{bmatrix} 42349.90252151 \\ -8670.854777 \\ 18096.17636737 \end{bmatrix} \end{aligned}$$

$$\boldsymbol{v}_4 = \boldsymbol{v}^+ + q'\mathbf{E} = \begin{bmatrix} 42349.90252151 \\ -8670.854777 \\ 18096.17636737 \end{bmatrix} + 4.78597877761177 \begin{bmatrix} 1000 \\ 0 \\ 0 \end{bmatrix} = \begin{bmatrix} 47135.88129912 \\ -8670.854777 \\ 18096.17636737 \end{bmatrix}$$

$$\boldsymbol{x}_4 = \boldsymbol{x}_3 + \Delta t \ \boldsymbol{v}_4 = \begin{bmatrix} -0.48966698 \\ -0.00346834 \\ 0.00388755 \end{bmatrix} + 10^{-7} \ \begin{bmatrix} 47135.88129912 \\ -8670.854777 \\ 18096.17636737 \end{bmatrix} = \begin{bmatrix} -0.48495339 \\ -0.00433543 \\ 0.00569717 \end{bmatrix}$$

We see that the values for x_3 , x_4 , v_3 and v_4 for particle 7 are close (the latter digits differ due to the differing precision of calculations); i.e. the same when done by hand and in the program.

Based on these calculations, we believe that our particle update works as expected; according to the Boris Algorithm, and this part of the program works correctly.

2 Plasma Stream

- Maxwellian distribution
- maybe Parabolic distribution
- Change Voltage of electrode to change electric field
- Change current in Helmholtz coil to change the magnetic field

while the updates are running. on different batches.

We now create our first plasma system, where we can change certain parameters of the fields; so as to simulate controlling plasma in a chamber by changing the electric and magnetic fields as required.

3 The Magnetic Mirror configuration

need parabolic B along z-axis configuration for $\nabla \mathbf{B} \parallel \mathbf{B}$