Ex8_ThermoACF-Copy1

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1 Exercise 6:

2 Lennard-Jones particles and Velocity Verlet integrator

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```
[1]: %matplotlib inline
import matplotlib.pyplot as plt
from matplotlib import animation, rc
from IPython.display import HTML
import random
import copy
import numpy as np
import scipy as scy
from tqdm.notebook import trange, tqdm
import time
from scipy.stats import maxwell

#used for the video
import subprocess
import glob
import os
```

2.1 Task I: Implementation of Berendsen thermostat

Berendsen thermostat (velocity rescaling factor) Later set to T = 300 K with = 0.2 ps.

$$\lambda = \sqrt{1 + \frac{\Delta t}{\tau} (\frac{T_0}{T} - 1)}$$

```
[2]: def Lambda_fact(Dlt_t, tau, Temp_o, Temp):
    lambda_f = np.sqrt( 1 + (Dlt_t/tau)*(Temp_o/Temp -1))
    return lambda_f
```

Maxwell Boltzmann distribution with a scale parameter $a = \sqrt{\frac{k_b T}{m}}$

```
[3]: def MaxBoltz ():
    a = np.sqrt(k_b * Temp_ini / mass)
    mx = maxwell(scale=a)
    x = np.linspace(mx.ppf(0.01),mx.ppf(0.99), 100)
    velocity_distribution = mx.rvs(size=1000)
    return a, velocity_distribution
```

2.2 Task II: Force autocorrelation and friction coefficient calculation

Force autocorrelation function

With N out total number of steps in the simulation, we need $M + \tau$ to be smaller or equal to N. To avoid having 'overhanging steps' we sum only over $M = N - \tau$ steps.

$$C(\tau) = f(0)f(\tau) \frac{1}{M} \sum_{i=1}^{M} (f(t_i) - \bar{f})(f(t_i + \tau) - \bar{f})$$

```
[65]: def Force_t(t):
    F = 0
    for p_ind in range(Particule_Nbr):
        f = force_BIS(Relative_dist_arrays_X, Relative_dist_arrays_Y, t , p_ind)
        F += np.sqrt( f[0]**2 + f[1]**2)
    return F
```

```
[45]: #calculating f bar
F_bar_list = [0]*steps
for t in trange(steps):
    F_bar_list[t] = Force_t( t)
F_bar = np.mean(F_bar_list)
```

HBox(children=(FloatProgress(value=0.0, max=20000.0), HTML(value='')))

```
[69]: def C_tau(tau, F_bar):
    M = steps - tau -1
    m = np.linspace(0,M, int(M/40))
    m = [int(m[i]) for i in range (len(m))]
    sum_value = 0
    for i in trange(len(m)):
        m_value = m[i]
        sum_value += (Force_t(m_value) - F_bar) * (Force_t(m_value+tau) - F_bar)
    sum_value = sum_value / len(m)
    return sum_value
```

2.3 Task III: Simulation

Simulation of 49 particules in a 5x5nm box (with PBC).

The interparticle interaction is modeled as a Lennard-Jones potential.

The Velocity Verlet integrator is used to calculate the motion of the particles. The following constant are used:

```
[4]: box = (5,5) \#nm^2
     #20000 total time steps in the simulation
     steps = 20000
     #Number of particles in the box
     Particule_Nbr = 49
     mass = 18 \#q/mol
     #Time step (2*e-6 in nm)
     Dlt_t = 2e-6 \# ns = 2fs
     tau = 2e-4 \#ns = 0.2ps
     k b = 8.314462 \#JK^{(-1)}.mol^{(-1)}
     Temp_o = 300 \# K
     #Temperature used for the initial Maxwell-Boltzmann velocity distribution
     Temp_ini = 100 \# K
     #Constant used in the Lennard Jones potential
     C_12 = 9.847044 *10**(-6) #kJ mol^-1 nm^12
     C_6 = 6.2647225 *10**(-3) #kJ mol^-1 nm^6
```

Position

$$x_{k+1} = x_k + v_k \Delta t + \frac{1}{2} a_k \Delta t^2$$

Velocity

$$v_{k+1} = v_k + \frac{1}{2}(a_k + a_{k+1})\Delta t$$

Maybe the units are wrong and my acceleration difference isso smallthat is does not affect my velocity

```
[6]: def velocity (v_k, a_k, a_k1, Dlt_t):
    v_k1 = v_k + (1/2)*(a_k+a_k1)*Dlt_t
    return v_k1
```

Potential (Lennard Jones)

```
V_{IJ}(r_{ij}) = \frac{C_{12}}{r_{ij}^{12}} - \frac{C_6}{r_{ij}^6}
```

```
[7]: def potential(r_ij_vect):
    #distance between the two particules
    r_ij = np.sqrt(r_ij_vect[0]**2 + r_ij_vect[1]**2)

if r_ij == 0:
    return 0

else:
    V_ij = C_12/r_ij**(12) - C_6/r_ij**(6)
    return V_ij
```

```
[8]: def pot_total(time_t):
    V= 0

for our_p_ind in range (Particule_Nbr):
    our_p = Particules_list[our_p_ind]
    for other_p_ind in range(our_p_ind+1 ,Particule_Nbr) :
        other_p = Particules_list[other_p_ind]

        r_ij_vect = get_vect_r(our_p,other_p)
        V += potential(r_ij_vect)

    return V
```

Relative distance between two particles, with periodic boundary conditions

• Using the particle class

```
[10]: def get_vect_r(particul1, particul2):
    x_list =[]
    x_list += [particul2.x - particul1.x]
    x_list += [particul2.x - particul1.x + box[0]]
```

```
x_list += [particul2.x - particul1.x - box[0]]
x_part = min(x_list, key=abs)

y_list =[]
y_list += [particul2.y - particul1.y]
y_list += [particul2.y - particul1.y + box[1]]
y_list += [particul2.y - particul1.y - box[1]]
y_part = min(y_list, key=abs)

r = [x_part, y_part]

return r
```

• Using x and y position of the two particles

```
[11]: def get_vect_r_BIS(x,y, x2, y2):
    x_list =[]
    x_list += [x2 - x]
    x_list += [x2 - x + box[0]]
    x_list += [x2 - x - box[0]]
    x_part = min(x_list, key=abs)

    y_list =[]
    y_list += [y2 - y]
    y_list += [y2 - y + box[1]]
    y_list += [y2 - y - box[1]]
    y_part = min(y_list, key=abs)

    r = [x_part, y_part]
    return r
```

Force / acceleration

```
f(t) = m(t) = mv(t) = ma(t)F_{IJ}(r_{ij}) = \left(12\frac{C_{12}}{r_{ij}^{13}} - 6\frac{C_6}{r_{ij}^7}\right)\frac{\vec{r_{ij}}}{r_{ij}}
```

```
[12]: def force_ij(r_ij_vect):
    #distance between the two particules
    r_ij = np.sqrt(r_ij_vect[0]**2 + r_ij_vect[1]**2)

if r_ij != 0:
    factor = (12*C_12/r_ij**(13) - 6*C_6/r_ij**(7))/r_ij
    Fij_x = factor * r_ij_vect[0]
    Fij_y = factor * r_ij_vect[1]
Fij_vect = [Fij_x, Fij_y]
```

```
return Fij_vect
else :
   return [0,0]
```

• Force on the (p ind)th particle at time t using the Relative dist arrays

```
[13]: def force_BIS(Relative_dist_arrays_X, Relative_dist_arrays_Y, t , p_ind):
    F= [0,0] #total force vector acting on our particle
    if t > (steps-1):
        return F[0,0]

#list of relative distances along x and y with all the other particles
    r_vectx = Relative_dist_arrays_X[t][p_ind,:]
    r_vecty = Relative_dist_arrays_Y[t][p_ind,:]

#Calculating the total force on (p_ind)th particle
for i in range (len(r_vectx)):
    Fij_vect = force_ij([r_vectx[i],r_vecty[i]])

#sum
    F[0] += Fij_vect[0]
    F[1] += Fij_vect[1]

return F
```

Kinetic energy

```
E_{kin} = \frac{1}{2}m < v^2 >
```

```
def Kinetic(Data_traj_list, time_t):
    K_list = []
    for p_ind in range (Particule_Nbr):
        p_vx = Data_traj_list[p_ind][2,time_t]
        p_vy = Data_traj_list[p_ind][3,time_t]
        v = np.sqrt(p_vx**2 + p_vy**2)
        K_list += [v**2]
    K = (1/2)*mass*np.mean(K_list)
    #print(time_t, v)
    return K
```

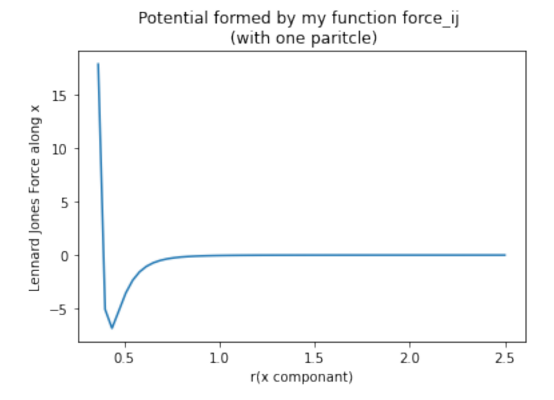
Initialisation

• Check that the force function looks correct:

```
[15]: a = np.linspace(0,2.5, 70)
f =[]
for i in range (70):
    f += [force_ij([a[i],0])[0]]
```

```
plt.plot(a[10:],f[10:])
plt.title('Potential formed by my function force_ij \n (with one paritcle)')
plt.ylabel('Lennard Jones Force along x')
u = plt.xlabel('r(x componant)')
mi = f.index(min(f)) #index of the minimum
print('There is a strong repulsive force if a particle gets closer then ', \_
\to a[mi], 'nm')
```

There is a strong repulsive force if a particle gets closer then 0.4347826086956522 nm



2.3.1 Relative distance between the particles

Creating a gereral list of arrays, with one array per time steps storing the relative distance of the particles. With r_{ij} being the vector r from particle i to particle j:

$$\begin{pmatrix} r_{11} = 0 & r_{12} & \dots & r_{1N} \\ r_{21} & r_{22} = 0 & \dots & r_{2N} \\ \dots & \dots & \ddots & \dots \\ r_{N1} & r_{N2} & \dots & r_{NN} = 0 \end{pmatrix}_{t}$$

From the data traj list information it calculates the relative distance between all the particles of

the system at the time t given in argument. And updates the gereral Relative_distance_arrays, adding the new arrays for time t

-Takes a bit of time to run-

```
[16]: def Relative dist(Relative_dist_arrays X, Relative_dist_arrays Y, t):
          Relative_dist_arrays_X += [np.zeros((Particule_Nbr,Particule_Nbr))]
          Relative_dist_arrays_Y += [np.zeros((Particule_Nbr,Particule_Nbr))]
          for p in range (Particule_Nbr):
              xp= Data_traj_list[p][0,:]
              yp= Data_traj_list[p][1,:]
              for p2 in range (p+1,Particule_Nbr):
                  xp2 = Data_traj_list[p2][0,:]
                  yp2 = Data_traj_list[p2][1,:]
                  #vector r from particle p to particle p2
                  r_vector = get_vect_r_BIS(xp[t],yp[t], xp2[t], yp2[t])
                  Relative_dist_arrays_X[t][p,p2] = r_vector[0]
                  Relative_dist_arrays_Y[t][p,p2] = r_vector[1]
                  #opposite vector r from p2 to p
                  Relative_dist_arrays_X[t][p2,p] = -r_vector[0]
                  Relative_dist_arrays_Y[t][p2,p] = -r_vector[1]
          return (Relative_dist_arrays_X, Relative_dist_arrays_Y)
          \#i\_lower = np.tril\_indices(n, -1)
          #matrix[i_lower] = matrix.T[i_lower]
```

Initialise simulation

The particles are placed on a regular grid of 7x7 particles. They are assigned their initial velocity |v(x,y)| following the Maxwell-Boltzmann distribution at T = 100K, with random directions.

```
class Particle:
    def __init__(self, x, y, vx, vy, ax, ay):
        self.x = x
        self.y = y
        self.vx = vx
        self.vy = vy
        self.ax = ax
        self.ay = ay

def __repr__(self):
```

```
return str("This is a particle at %0.2f, %0.2f with v=%0.2f,%0.2f" %⊔

⇔(self.x,self.y,self.vx,self.vy))
```

```
[18]: def Particules initialise (Particule Nbr, steps):
          Particules_list = []
          Data_traj_list = []
          a, velocity_distribution = MaxBoltz()
          #regurlar grid 7*7
          nx, ny = (7, 7)
          x_grid = np.linspace(0, 5, nx+1)
          y_grid = np.linspace(0, 5, ny+1)
          d = x_grid[2]-x_grid[1]
          x grid = x grid[0:len(x grid)-1] +d/2
          y_grid = y_grid[0:len(y_grid)-1] +d/2
          print('The space between two particle is : ', d, 'nm')
          p_count = 0
          for i in range (nx):
              for j in range (ny):
                  #Initial position on a uniform grid
                  angle = np.random.uniform(0,2*np.pi)
                  x_p = x_{grid}[i]
                  y_p = y_{grid}[j]
                  #Initial velocities : Maxwell-Boltzmann distribution with random
       \rightarrow direction
                  vel = velocity_distribution[np.random.randint(0,__
       →len(velocity_distribution))]
                  vx_p = np.sin(angle)*vel
                  vy_p = np.cos(angle)*vel
                  Particules_list += [Particle(x_p, y_p, vx_p, vy_p, 0, 0)]
                  Data_traj_list += [np.zeros((4,steps))]
                  Data_traj_list[p_count][:,0] = [x_p, y_p, vx_p, vy_p] #[particle_
       → indice] [data type, time step]
                  p_count += 1
          print('Number of particles = ', p_count)
          return Particules_list, Data_traj_list
```

[19]: Particules_list, Data_traj_list = Particules_initialise (Particule_Nbr, steps)

```
The space between two particle is: 0.7142857142857143 nm Number of particles = 49
```

Run Simulation

At the start of the simulation we already have the initial positions, velocities and acceleration/force calculated. At each step we calculate:

FIRST LOOP: * Position at time t+1

Calculate all the relative distance at time t+1

SECOND LOOP: * Force at time t+1 * Velocities at time t+1

END OF TIME STEP * Rescaling the velocities with Berendsen thermostat

```
[20]: #arrays of relative distance between the particles
      Relative_dist_arrays_X = []
      Relative dist arrays Y = []
      Potential_list = [0]*steps
      # At time = 0
      Relative_dist_arrays_X, Relative_dist_arrays_Y = __
      →Relative_dist(Relative_dist_arrays_X, Relative_dist_arrays_Y, 0)
      Potential list[0] = pot total BIS(Relative dist arrays X,___
      →Relative_dist_arrays_Y, 0)
      for i in trange (steps-1):
          for p_ind in range (Particule_Nbr):
              our_P = Particules_list[p_ind]
              # calculating the next position
              x_1 = position (our_P.x, our_P.vx , our_P.ax, Dlt_t)%box[0]
              y_1 = position (our_P.y, our_P.vy, our_P.ay, Dlt_t)%box[1]
              # updating the particule position
              our_P.x = x_1
              our_P.y = y_1
              Data_traj_list[p_ind][0,i+1] = x_1
              Data_traj_list[p_ind][1,i+1] = y_1
          # Calculating the relative distance between all the particles at time i+111
       \rightarrow---> the one that takes the most time !
          Relative_dist_arrays_X, Relative_dist_arrays_Y =
       →Relative_dist(Relative_dist_arrays_X, Relative_dist_arrays_Y, i+1)
          Potential_list[i+1] = pot_total_BIS(Relative_dist_arrays_X,__
       →Relative_dist_arrays_Y, i+1)
          for ind in range (Particule_Nbr):
              our_P = Particules_list[ind]
              # calculating the force/acceleration at the next step
```

```
F_1 = force_BIS (Relative_dist_arrays_X, Relative_dist_arrays_Y,i+1_
\hookrightarrow, ind)
       ax_1, ay_1 = -F_1[0]*1/(mass*1), -F_1[1]*1/(mass*1)
       #velocity
       vx 1 = velocity (our P.vx, our P.ax, ax 1, Dlt t)
       vy_1 = velocity (our_P.vy, our_P.ay, ay_1, Dlt_t)
       #updating velocity and acceleration
       our_P.ax = ax_1
       our_P.ay = ay_1
       our_P.vx = vx_1
       our_P.vy = vy_1
       Data_traj_list[ind][2,i+1] = vx_1
       Data_traj_list[ind][3,i+1] = vy_1
  #rescaling the velocities
  K = Kinetic(Data_traj_list, i+1)
  Temp = K / k b
  lbda = Lambda_fact(Dlt_t, tau, Temp_o, Temp)
  for ind in range(Particule Nbr):
       Rs_vx = (Data_traj_list[ind][2,i+1]) * lbda
       Rs_vy = (Data_traj_list[ind][3,i+1]) * lbda
       P = Particules_list[ind]
       P.vx, Data_traj_list[ind][2,i+1] = Rs_vx, Rs_vx
       P.vy, Data_traj_list[ind][3,i+1] = Rs_vy, Rs_vy
```

HBox(children=(FloatProgress(value=0.0, max=19999.0), HTML(value='')))

```
[21]: for i in trange(1,desc= 'Plot the graph of the last simulation'):

#def plot_simulation (Particule_Nbr, Data_traj_list, Particules_list):
for parti in range (Particule_Nbr):
    data_traj = Data_traj_list[parti]
    #Particule = Particules_list[parti]
    plt.plot(data_traj[0,:],data_traj[1,:], marker='.', markersize='10', use of the plot.ylabel('position x')
    plt.xlabel('position y')
    plt.xlim(0,5)
```

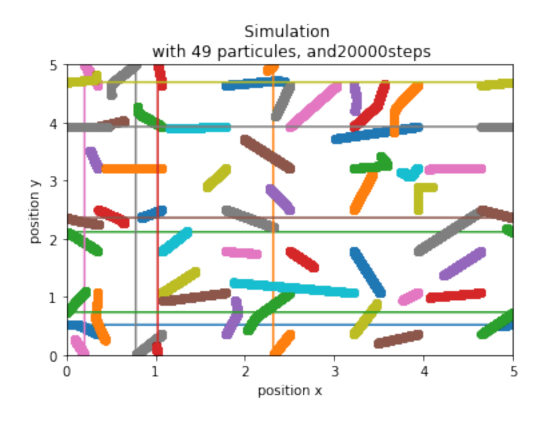
```
plt.ylim(0,5)

Titles_graph1 = 'Simulation \n with ' + str(Particule_Nbr)+ ' particules,⊔

→and' + str(steps) + 'steps'

plt.title(Titles_graph1)
```

HBox(children=(FloatProgress(value=0.0, description='Plot the graph of the last simulation', make the state of the last simulation', make the state of the last simulation of the last



• Analyse the force of the first particles coliding

```
[22]: np.savez('save02.npz', Data_traj_list, Potential_list, Relative_dist_arrays_X, ⊔ →Relative_dist_arrays_Y)
```

```
[25]: #npzfile = np.load('save_array5.npz')

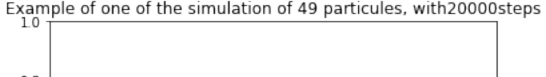
#Data_traj_list = npzfile['arr_0']

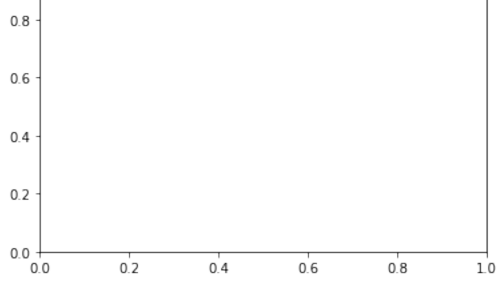
#Potential_list = npzfile['arr_1']
```

2.3.2 Video

```
[37]: def creat files():
         os.chdir("Image_storing_video")
         #def plot simulation (Particule Nbr, Data traj list, Particules list):
         for t in trange (int(18000), steps) :
             for parti in range (Particule_Nbr):
                data_traj = Data_traj_list[parti]
                plt.plot(data_traj[0,t],data_traj[1,t], marker='.',__
      →markersize='10', linestyle = '-', color = 'black')
                plt.xlabel('position x')
                plt.ylabel('position y')
                plt.xlim(0,box[0])
                plt.ylim(0,box[1])
            plt.savefig("File%02d.png" % t)
             #plt.show()
             #plt.savefig("file.png")
            plt.close()
         Titles graph1 = 'Example of one of the simulation of ' +11
      plt.title(Titles_graph1)
         os.chdir("../")
[38]: def creat_video():
         os.chdir("Image storing video")
         subprocess.call(['ffmpeg', '-framerate', '8', '-i', 'File%02d.png', '-r', _
      #subprocess.call(['ffmpeq', '-framerate', '8', '-i', Titles_files, '-r', __
      \rightarrow '30', '-pix_fmt', 'yuv420p', Filename])
         for file_name in glob.glob("*.png"):
             os.remove(file_name)
         os.chdir("../")
[39]: #os.chdir("ThermoACF")
     retval = os.getcwd()
     print ("Current working directory %s" % retval)
     creat_files()
     creat_video()
```

Current working directory
/home/lea/Bureau/Fac/Master/Simulating_the_physical_world/Simulations/ThermoACF
HBox(children=(FloatProgress(value=0.0, max=2000.0), HTML(value='')))





2.4 Task III: Potential and Kinetic energy

```
[101]: time_list = np.linspace(0, steps, steps)
```

Total potential energy (per step)

Calculating the potential using the relative distance arrays

```
from scipy.linalg import sqrtm
def pot (t):
    # All the other particles relative distances with the p_ind th particle at_
    itime t
        r_vector_x = Relative_dist_arrays_X[t]
        r_vector_y = Relative_dist_arrays_Y[t]

#Calculating the distance with all the other particuls (including over the_
    independence)
    rx_sqrd = np.linalg.matrix_power(r_vector_x , 2)
```

```
ry_sqrd = np.linalg.matrix_power(r_vector_y , 2)
norme_sqrd = np.add(rx_sqrd, ry_sqrd)
radial_distance_list = sqrtm(norme_sqrd) # SQUARE ROOT
#radial_distance_list = [np.sqrt(r_vector_x[i]**2 + r_vector_y[i]**2) for iu
in range (len(r_vector_x))]

RDist_12 = np.linalg.matrix_power(radial_distance_list, -12)*C_12
RDist_6 = np.linalg.matrix_power(radial_distance_list, -6)*C_6

pot = np.sum(np.subtract(RDist_12,RDist_6))
return pot
```

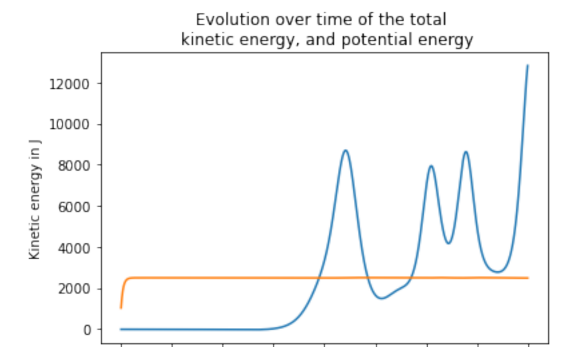
```
[106]: plt.plot(time_list[1:-1], Potential_list[1:-1])
    #plt.title('Evolution over time of the total potential energy')
    #plt.ylabel('Potential energy in J')

Kin_list = []
for i in trange (steps):
    Kin_list += [Kinetic(Data_traj_list,i)]

plt.plot(time_list, Kin_list)
    plt.title('Evolution over time of the total \n kinetic energy, and potential_\sum_\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\t
```

HBox(children=(FloatProgress(value=0.0, max=20000.0), HTML(value='')))

```
[106]: Text(0, 0.5, 'Kinetic energy in J')
```



Time in fs

7500 10000 12500 15000 17500 20000

Total kinetic energy (per step)

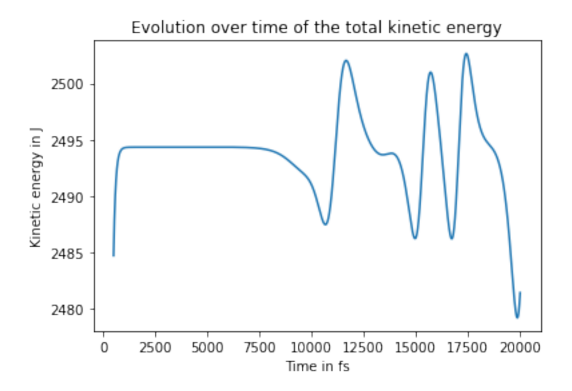
2500

0

5000

```
[105]: plt.plot(time_list[500:], Kin_list[500:])
   plt.title('Evolution over time of the total kinetic energy')
   plt.xlabel('Time in fs')
   plt.ylabel('Kinetic energy in J')
```

[105]: Text(0, 0.5, 'Kinetic energy in J')



Interpretation of the results

There seems to be some problems in the calculation, as the Kinetic and potential energy do have similar and opposite shapes but they are in different order of magnitude.

2.5 Task V: Calculation and interpretation of friction coefficient

```
[]: time_list = list(np.linspace(0,200,200))
C_list = []

count = 0
for tau in trange(0, 15000):
    #I had running time issues so I collected 1 out of 10 values for tau < 200
    # and 1 out of 20 values other tau
    if tau%20==0:
        count += 1
        C_list2 += [C_tau(tau, F_bar)]
        time_list += [tau]</pre>
```

```
[]: np.savez('save_Ctau02.npz', C_list2, time_list)
np.savez('save_Fbar.npz', F_bar)
```

I had issues with running the code on my computer so I ran it using the code in the above cells and saved the results in arrays.

```
[96]: C_tau5000_File = np.load('save_Ctau03_5000.npz')
    C_tau5000to15000_File = np.load('save_Ctau04_5000to15000.npz')
    C_tau_list = C_tau5000_File['arr_0']+C_tau5000to15000_File['arr_0']
    time_list_C_tau = C_tau5000_File['arr_1']+C_tau5000to15000_File['arr_1']

[100]: plt.plot(time_list_C_tau, C_tau_list)
    plt.title('Evolution of the force autocorrelation function')
    plt.xlabel('Lag time = Tau')
    a = plt.ylabel('C(tau)')
```

