

## FFR120: Homework 1 by Richard Blücher

## Exercise 1

```
In [11]: import numpy as np
# Parameters for the Lennard-Jones potential.
m = 1 # Mass (units of m0).
m_disc = 10 # Disc mass
r_disc = 10 # Disc radius
sigma = 1 # Size (units of sigma0).
eps = 1 # Energy (unit of epsilon0).
v0 = 10 # Initial speed (units of v0 = sqrt((2 * epsilon0) / m0)).

# Parameters for the simulation.
N_particles = 25**2 # Number of particles.
full_time = 401 # Time for simulation to run

dt = 0.005 # Time step (units of t0 = sigma * sqrt(m0 / (2 * epsilon0))).

L = 260 # Box size (units of sigma0).
x_min, x_max, y_min, y_max = -L/2, L/2, -L/2, L/2

cutoff_radius = 5 * sigma # Cutoff_radius for neighbours list.
```

```
In [12]: # Generate initial positions on a grid and orientations at random.
x0, y0 = np.meshgrid(
    np.linspace(- L / 2, L / 2, int(np.sqrt(N_particles))),
    np.linspace(- L / 2, L / 2, int(np.sqrt(N_particles))),
)
x0 = x0.flatten()[:N_particles]
y0 = y0.flatten()[:N_particles]
phi0 = (2 * np.random.rand(N_particles) - 1) * np.pi

# Initial values of disc
x0_disc = 0
y0_disc = 0
v0_disc = 0

# Initialize the neighbour list.
def list_neighbours(x, y, N_particles, cutoff_radius):
    '''Prepare a neighbours list for each particle.'''
    neighbours = []
    neighbour_number = []
    for j in range(N_particles):
        distances = np.sqrt((x - x[j]) ** 2 + (y - y[j]) ** 2)
        neighbor_indices = np.where(distances <= cutoff_radius)
        neighbours.append(neighbor_indices)
        neighbour_number.append(len(neighbor_indices))
    return neighbours, neighbour_number

neighbours, neighbour_number = list_neighbours(x0, y0, N_particles, cutoff_radius)

# Distance to disc function
def disc_dist(x_disc, y_disc, x, y):

    distances = np.sqrt((x_disc - x) ** 2 + (y_disc - y) ** 2)
    return distances
```

```

# If particles are to close to disc, remove them
disc_distances = disc_dist(x0_disc, y0_disc, x0, y0)
index_to_remove = np.where(disc_distances <= r_disc + 3*sigma)
x0 = np.delete(x0, index_to_remove)
y0 = np.delete(y0, index_to_remove)
phi0 = np.delete(phi0, index_to_remove)
N_particles = len(x0)

# Initialize the variables for the Leapfrog algorithm.
# Current time step.
x = x0
y = y0
x_half = np.zeros(N_particles)
y_half = np.zeros(N_particles)
v = v0
phi = phi0
vx = v0 * np.cos(phi0)
vy = v0 * np.sin(phi0)

x_disc = x0_disc
y_disc = y0_disc
x_disc_half = 0
y_disc_half = 0
v_disc = v0_disc
phi_disc = 0
vx_disc = v0_disc * np.cos(phi_disc)
vy_disc = v0_disc * np.sin(phi_disc)

# Next time step.
nx = np.zeros(N_particles)
ny = np.zeros(N_particles)
nv = np.zeros(N_particles)
nphi = np.zeros(N_particles)
nvx = np.zeros(N_particles)
nvy = np.zeros(N_particles)

nx_disc = 0
ny_disc = 0
nv_disc = 0
nphi_disc = 0
nvx_disc = 0
nvy_disc = 0

```

In [13]:

```

def total_force_cutoff(x, y, N_particles, sigma, epsilon, neighbours):
    ...
        Calculate the total force on each particle due to the interaction with a
        neighbours list with the particles interacting through a Lennard-Jones
        potential.
    ...
    Fx = np.zeros(N_particles)
    Fy = np.zeros(N_particles)
    for i in range(N_particles):
        for j in list(neighbours[i][0]):
            if i != j:
                r2 = (x[i] - x[j]) ** 2 + (y[i] - y[j]) ** 2
                r = np.sqrt(r2)
                Fx[i] += epsilon * sigma * (1 / r2) * (3 * r * r - 1)
                Fy[i] += epsilon * sigma * (1 / r2) * (3 * r * r - 1) * (y[j] - y[i])

```

```

        ka2 = sigma ** 2 / r2

        # Force on i due to j.
        F = 24 * epsilon / r * (2 * ka2 ** 6 - ka2 ** 3) # Modulus.

        Fx[i] += F * (x[i] - x[j]) / r
        Fy[i] += F * (y[i] - y[j]) / r
    return Fx, Fy

```

```

In [14]: def LJ_force(x_disc, y_disc, x, y, epsilon, sigma):
    Fx = np.zeros(N_particles)
    Fy = np.zeros(N_particles)
    r = disc_dist(x_disc, y_disc, x, y) - r_disc
    ka2 = sigma ** 2 / r**2
    F = 24 * epsilon / r * (2 * ka2 ** 6 - ka2 ** 3)

    Fx += F * (x_disc - x) / r
    Fy += F * (y_disc - y) / r

    return -Fx, -Fy

```

```
In [15]: Fx, Fy = LJ_force(x0_disc, y0_disc, x0, y0, eps, sigma)
```

```

In [16]: import time
from scipy.constants import Boltzmann as kB
from tkinter import *
import math

visuals_on = False # To turn on/off visual simulation

if visuals_on:
    window_size = 600

    tk = Tk()
    tk.geometry(f'{window_size + 20}x{window_size + 20}')
    tk.configure(background='#000000')

    canvas = Canvas(tk, background='#ECECEC') # Generate animation window
    tk.attributes('-topmost', 0)
    canvas.place(x=10, y=10, height=window_size, width=window_size)

    disc = canvas.create_oval(
        (x_disc - r_disc / 2) / L * window_size + window_size / 2,
        (y_disc - r_disc / 2) / L * window_size + window_size / 2,
        (x_disc + r_disc / 2) / L * window_size + window_size / 2,
        (y_disc + r_disc / 2) / L * window_size + window_size / 2,
        outline='#000000',
        fill='#000000',
    )

    particles = []
    for j in range(0, N_particles):
        particles.append(
            canvas.create_oval(
                (x[j] - sigma / 2) / L * window_size + window_size / 2,
                (y[j] - sigma / 2) / L * window_size + window_size / 2,
                (x[j] + sigma / 2) / L * window_size + window_size / 2,
                (y[j] + sigma / 2) / L * window_size + window_size / 2,
                outline='#00C0C0',
            )
        )

```

```

        fill='#00C0C0',
    )
)

step = 0

# For plotting disc positions
disc_pos = np.zeros((int(full_time/dt), 2))

def stop_loop(event):
    global running
    running = False
if visuals_on:
    tk.bind("<Escape>", stop_loop) # Bind the Escape key to stop the loop.
running = True # Flag to control the loop.
while running:
    disc_pos[step] = [x_disc, y_disc]

    x_half = x + 0.5 * vx * dt
    y_half = y + 0.5 * vy * dt

    x_disc_half = x_disc + 0.5 * vx_disc * dt
    y_disc_half = y_disc + 0.5 * vy_disc * dt

    # fx, fy = \
#     total_force_cutoff(x_half, y_half, N_particles, sigma, eps, neighbours
fx, fy = LJ_force(x_disc_half, y_disc_half, x_half, y_half, sigma, eps)

nvx = vx + fx / m * dt
nvy = vy + fy / m * dt

nx = x_half + 0.5 * nvx * dt
ny = y_half + 0.5 * nvy * dt

nvx_disc = vx_disc + sum(-fx) / m_disc * dt # Force negative for disc if pos
nvy_disc = vy_disc + sum(-fy) / m_disc * dt

nx_disc = x_disc_half + 0.5 * nvx_disc * dt
ny_disc = y_disc_half + 0.5 * nvy_disc * dt

## Reflecting boundary conditions.
# For disc
if nx_disc < x_min:
    nx_disc = x_min + (x_min - nx_disc)
    nvx_disc = - nvx_disc

if nx_disc > x_max:
    nx_disc = x_max - (nx_disc - x_max)
    nvx_disc = - nvx_disc

if ny_disc < y_min:
    ny_disc = y_min + (y_min - ny_disc)
    nvy_disc = - nvy_disc

if ny_disc > y_max:
    ny_disc = y_max - (ny_disc - y_max)
    nvy_disc = - nvy_disc

```

```

# For particles
for j in range(N_particles):
    if nx[j] < x_min:
        nx[j] = x_min + (x_min - nx[j])
        nvx[j] = - nvx[j]

    if nx[j] > x_max:
        nx[j] = x_max - (nx[j] - x_max)
        nvx[j] = - nvx[j]

    if ny[j] < y_min:
        ny[j] = y_min + (y_min - ny[j])
        nvy[j] = - nvy[j]

    if ny[j] > y_max:
        ny[j] = y_max - (ny[j] - y_max)
        nvy[j] = - nvy[j]

# Update velocities
nv = np.sqrt(nvx ** 2 + nvy ** 2)
for i in range(N_particles):
    nphi[i] = math.atan2(nvy[i], nvx[i])

nv_disc = np.sqrt(nvx_disc ** 2 + nvy_disc ** 2)
nphi_disc = math.atan2(nvy_disc, nvx_disc)

# Update variables for next iteration.
x = nx
y = ny
vx = nvx
vy = nvy
v = nv
phi = nphi

x_disc = nx_disc
y_disc = ny_disc
vx_disc = nvx_disc
vy_disc = nvy_disc
v_disc = nv_disc
phi_disc = nphi_disc

# Update animation frame.
if visuals_on:
    if step % 100 == 0:
        canvas.coords(
            disc,
            (nx_disc - r_disc / 2) / L * window_size + window_size / 2,
            (ny_disc - r_disc / 2) / L * window_size + window_size / 2,
            (nx_disc + r_disc / 2) / L * window_size + window_size / 2,
            (ny_disc + r_disc / 2) / L * window_size + window_size / 2,
        )
        for j, particle in enumerate(particles):
            canvas.coords(
                particle,
                (nx[j] - sigma / 2) / L * window_size + window_size / 2,
                (ny[j] - sigma / 2) / L * window_size + window_size / 2,
                (nx[j] + sigma / 2) / L * window_size + window_size / 2,
                (ny[j] + sigma / 2) / L * window_size + window_size / 2,
            )

```

```

        )

tk.title(f'Time {step * dt:.1f} - Iteration {step}')
tk.update_idletasks()
tk.update()
time.sleep(.001) # Increase to slow down the simulation.

if step == full_time/dt - 1:
    running = False
step += 1

if visuals_on:
    tk.update_idletasks()
    tk.update()
    tk.mainloop() # Release animation handle (close window to finish).

```

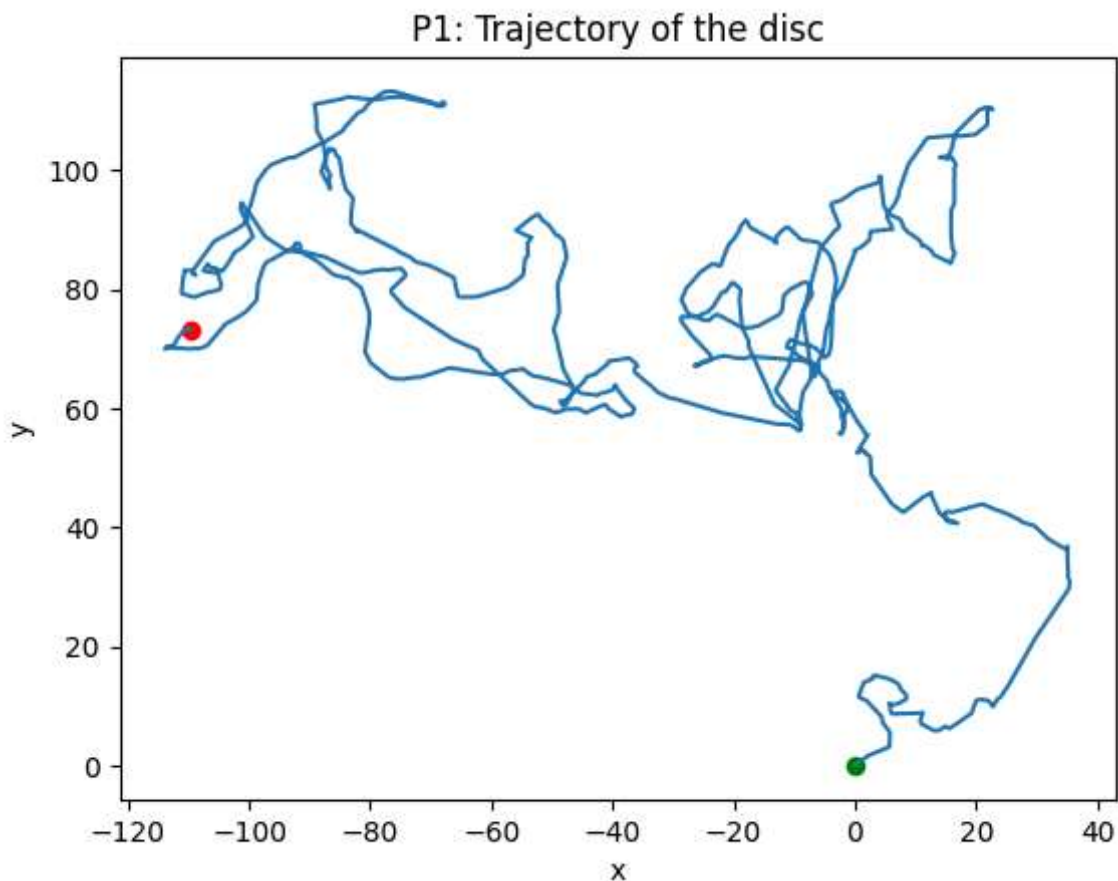
In [17]:

```

from matplotlib import pyplot as plt
plt.plot(disc_pos[:,0], disc_pos[:,1])
plt.scatter(0,0, c='g')
plt.scatter(disc_pos[-1, 0], disc_pos[-1, 1], c='r')
plt.title('P1: Trajectory of the disc')
plt.xlabel('x')
plt.ylabel('y')

```

Out[17]: Text(0, 0.5, 'y')



In [18]:

```

N = step
times = np.linspace(0, full_time, N)
msd2 = np.zeros(N)
disc_pos = disc_pos.T

```

```

for n in range(N):
    msd2[n] = 1/(N-n) * np.sum((disc_pos[0, n+1:N]-disc_pos[0, 1:N-n])**2 + (dis

```

In [19]:

```

msd_fit = np.polyfit(times[:int(150/dt)], msd2[:int(150/dt)], 1)

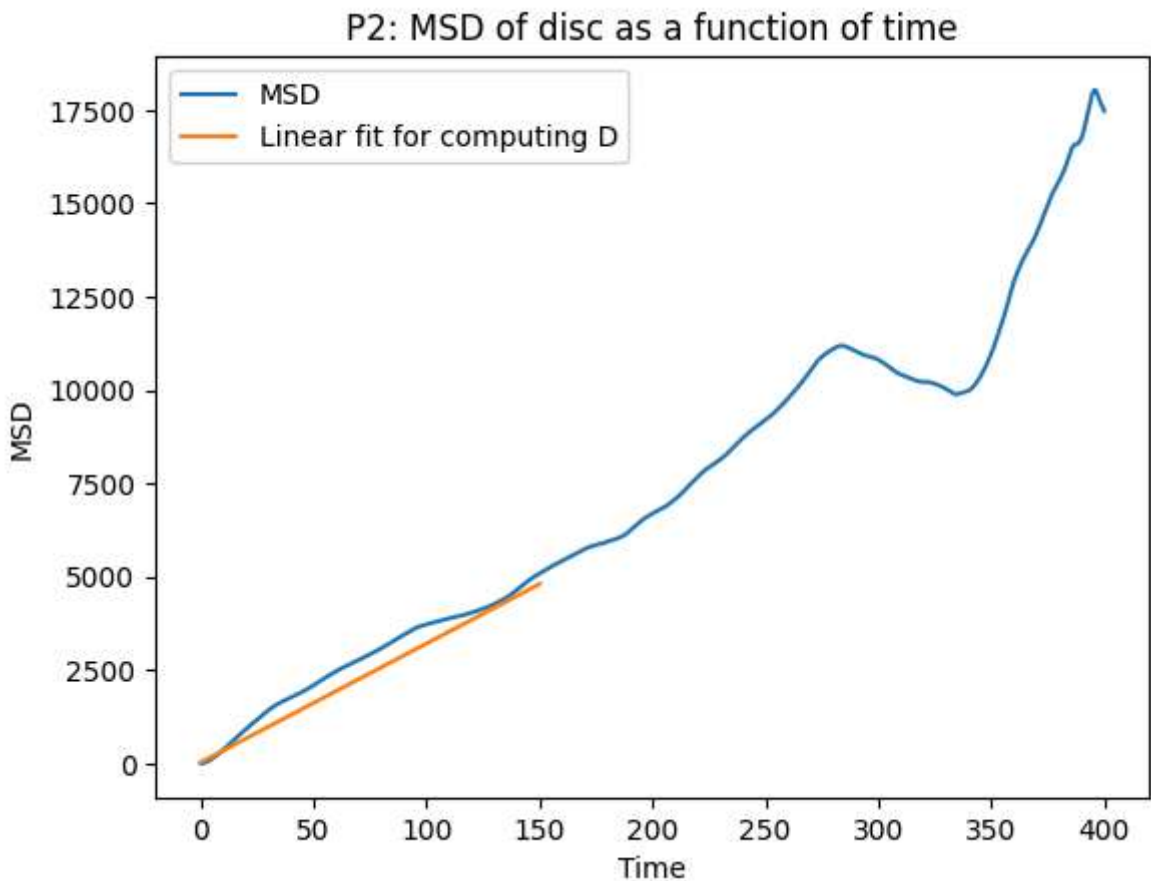
plt.plot(times[:-int(1/dt)], msd2[:-int(1/dt)], label = 'MSD')
plt.plot(times[:int(150/dt)], times[:int(150/dt)]*msd_fit[0] + msd_fit[0], label = 'Linear fit for computing D')

plt.xlabel('Time')
plt.ylabel('MSD')
plt.title('P2: MSD of disc as a function of time')
plt.legend()

print(f'Q1: The diffusion coefficient D = {msd_fit[0]/4:.4f}')

```

Q1: The diffusion coefficient D = 7.953



## Exercise 2

In [ ]:

```

N = 100 # Size of the spin lattice.
H = 0 # External field.
H_list = np.array([-5, -2, -1, -0.5, -0.2, -0.1, 0, 0.1, 0.2, 0.5, 1, 2, 5])
J = 1 # Spin-spin coupling.
T = 5 # Temperature. Temperatura critica ~2.269.
T_list = np.array([0.1, 0.2, 0.5, 1, 2, 2.1, 2.2, 2.3, 2.4, 2.5, 2.6, 2.7, 2.8,
N_iterations = 5000 # Nr of iterations to run simulation for

sl = 2 * np.random.randint(2, size=(N, N)) - 1

N_up = np.sum(sl + 1) / 2
N_down = N * N - N_up

```

```
print(f"Spin lattice created: N_up={N_up} N_down={N_down}")
```

```
Spin lattice created: N_up=5037.0 N_down=4963.0
```

```
In [22]: def neighboring_spins(i_list, j_list, sl):
    """
        Function returning the position of the neighbouring spins of a list of
        spins identified by their positions in the spin lattice.

    Parameters
    =====
    i_list : Spin position first indices.
    j_list : Spin position second indices.
    sl : Spin lattice.
    """

    Ni, Nj = sl.shape # Shape of the spin lattice.

    # Position neighbors right.
    i_r = i_list
    j_r = list(map(lambda x:(x + 1) % Nj, j_list))

    # Position neighbors left.
    i_l = i_list
    j_l = list(map(lambda x:(x - 1) % Nj, j_list))

    # Position neighbors up.
    i_u = list(map(lambda x:(x - 1) % Ni, i_list))
    j_u = j_list

    # Position neighbors down.
    i_d = list(map(lambda x:(x + 1) % Ni, i_list))
    j_d = j_list

    # Spin values.
    sl_u = sl[i_u, j_u]
    sl_d = sl[i_d, j_d]
    sl_l = sl[i_l, j_l]
    sl_r = sl[i_r, j_r]

    return sl_u, sl_d, sl_l, sl_r
```

```
In [23]: def energies_spins(i_list, j_list, sl, H, J):
    """
        Function returning the energies of the states for the spins in given
        positions in the spin lattice.

    Parameters
    =====
    i_list : Spin position first indices.
    j_list : Spin position second indices.
    sl : Spin lattice.
    """

    sl_u, sl_d, sl_l, sl_r = neighboring_spins(i_list, j_list, sl)

    sl_s = sl_u + sl_d + sl_l + sl_r

    E_u = - H - J * sl_s
```

```
E_d = H + J * sl_s

return E_u, E_d
```

```
In [24]: def energies_spins(i_list, j_list, sl, H, J):
    """
        Function returning the energies of the states for the spins in given
        positions in the spin lattice.

    Parameters
    =====
    i_list : Spin position first indices.
    j_list : Spin position second indices.
    sl : Spin lattice.
    """

    sl_u, sl_d, sl_l, sl_r = neighboring_spins(i_list, j_list, sl)

    sl_s = sl_u + sl_d + sl_l + sl_r

    E_u = - H - J * sl_s
    E_d = H + J * sl_s

    return E_u, E_d
```

```
In [25]: def probabilities_spins(i_list, j_list, sl, H, J, T):
    """
        Function returning the energies of the states for the spins in given
        positions in the spin lattice.

    Parameters
    =====
    i_list : Spin position first indices.
    j_list : Spin position second indices.
    sl : Spin lattice.
    """

    E_u, E_d = energies_spins(i_list, j_list, sl, H, J)

    Ei = np.array([E_u, E_d])

    Z = np.sum(np.exp(- Ei / T), axis=0) # Partition function.
    pi = 1 / np.array([Z, Z]) * np.exp(- Ei / T) # Probability.

    return pi, Z
```

```
In [26]: def compute_magnetization(sl):
    m = np.sum(sl)/sl.shape[0]**2
    return m
```

## Task 1

```
In [28]: import random
import time
from tkinter import *
H_iteration = 0
m = np.zeros_like(H_list)
for H in H_list:
```

```

f = 0.05 # Number of randomly selected spins to flip-test.

m_many = 0
n_average = 0

visuals_on = False
if visuals_on:
    N_skip = 10 # Visualize status every N_skip steps.

    window_size = 600

    tk = Tk()
    tk.geometry(f'{window_size + 20}x{window_size + 20}')
    tk.configure(background='#000000')

    canvas = Canvas(tk, background='#ECECEC') # Generate animation window.
    tk.attributes('-topmost', 0)
    canvas.place(x=10, y=10, height=window_size, width=window_size)

Nspins = np.size(sl) # Total number of spins in the spin lattice.
Ni, Nj = sl.shape

S = int(np.ceil(Nspins * f)) # Number of randomly selected spins.

step = 0

def stop_loop(event):
    global running
    running = False
if visuals_on:
    tk.bind("<Escape>", stop_loop) # Bind the Escape key to stop the Loop.
running = True # Flag to control the loop.
while running:
    ns = random.sample(range(Nspins), S)

    i_list = list(map(lambda x: x % Ni, ns))
    j_list = list(map(lambda x: x // Ni, ns))

    pi, Z = probabilities_spins(i_list, j_list, sl, H, J, T)

    rn = np.random.rand(S)
    for i in range(S):
        if rn[i] > pi[0, i]:
            sl[i_list[i], j_list[i]] = -1
        else:
            sl[i_list[i], j_list[i]] = 1

    if visuals_on:
        # Update animation frame.
        if step % N_skip == 0:
            canvas.delete('all')
            spins = []
            for i in range(Ni):
                for j in range(Nj):
                    spin_color = '#FFFFFF' if sl[i,j] == 1 else '#000000'
                    spins.append(
                        canvas.create_rectangle(
                            j / Nj * window_size,
                            i / Ni * window_size,

```

```

        (j + 1) / Nj * window_size,
        (i + 1) / Ni * window_size,
        outline='',
        fill=spin_color,
    )
)

tk.title(f'Iteration {step}')
tk.update_idletasks()
tk.update()
time.sleep(0.1) # Increase to slow down the simulation.

if step >= N_iterations - 200:
    m_many += compute_magnetization(sl)
    n_average += 1

step += 1

if step == N_iterations:
    running = False

m[H_iteration] = m_many/n_average
print(f'For H: {H} -> m = {m[H_iteration]}')
H_iteration += 1

if visuals_on:
    tk.update_idletasks()
    tk.update()
    tk.mainloop() # Release animation handle (close window to finish).

```

For H: -5.0 -> m = -0.9381810000000002  
 For H: -2.0 -> m = -0.7274309999999997  
 For H: -1.0 -> m = -0.49439  
 For H: -0.5 -> m = -0.2746539999999998  
 For H: -0.2 -> m = -0.1159740000000001  
 For H: -0.1 -> m = -0.0595229999999999  
 For H: 0.0 -> m = -0.00816300000000004  
 For H: 0.1 -> m = 0.0509130000000001  
 For H: 0.2 -> m = 0.105925  
 For H: 0.5 -> m = 0.2746990000000001  
 For H: 1.0 -> m = 0.47979700000000014  
 For H: 2.0 -> m = 0.7258960000000001  
 For H: 5.0 -> m = 0.937029

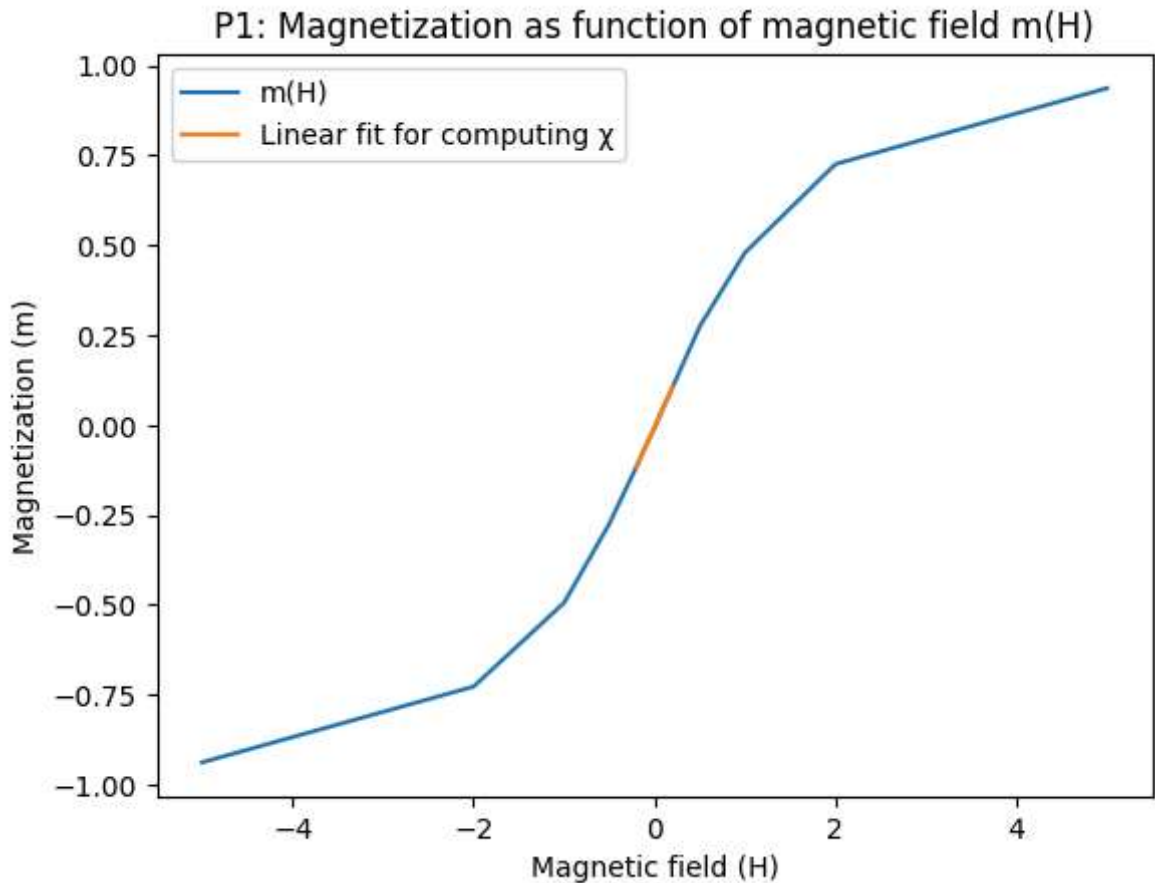
```
In [33]: linfit = np.polyfit(H_list[4:9], m[4:9], 1)
plt.plot(H_list, m, label = 'm(H)')
plt.plot(H_list[4:9], H_list[4:9]*linfit[0]+linfit[1], label = 'Linear fit for c')

print(f'Q1: Magnetic susceptibility \u03c7 = {linfit[0]}')


plt.xlabel('Magnetic field (H)')
plt.ylabel('Magnetization (m)')
plt.title('P1: Magnetization as function of magnetic field m(H)')
plt.legend()
```

Q1: Magnetic susceptibility  $\chi = 0.554234$

Out[33]: <matplotlib.legend.Legend at 0x22e2461f3b0>



## Task 2

```
In [34]: import random
import time
from tkinter import *
T_iteration = 0
H = 0.1
m = np.zeros_like(T_list)
for T in T_list:
    if step >= 300:
        H = 0
    f = 0.05 # Number of randomly selected spins to flip-test.

    m_many = 0
    n_average = 0

    visuals_on = False
    if visuals_on:
        N_skip = 10 # Visualize status every N_skip steps.

    window_size = 600

    tk = Tk()
    tk.geometry(f'{window_size + 20}x{window_size + 20}')
    tk.configure(background="#000000")

    canvas = Canvas(tk, background="#ECECEC") # Generate animation window.
    tk.attributes('-topmost', 0)
    canvas.place(x=10, y=10, height=window_size, width=window_size)

    Nspins = np.size(sl) # Total number of spins in the spin lattice.
    Ni, Nj = sl.shape
```

```

S = int(np.ceil(Nspins * f)) # Number of randomly selected spins.

step = 0

def stop_loop(event):
    global running
    running = False
if visuals_on:
    tk.bind("<Escape>", stop_loop) # Bind the Escape key to stop the Loop.
running = True # Flag to control the Loop.

while running:
    ns = random.sample(range(Nspins), S)

    i_list = list(map(lambda x: x % Ni, ns))
    j_list = list(map(lambda x: x // Ni, ns))

    pi, Z = probabilities_spins(i_list, j_list, sl, H, J, T)

    rn = np.random.rand(S)
    for i in range(S):
        if rn[i] > pi[0, i]:
            sl[i_list[i], j_list[i]] = -1
        else:
            sl[i_list[i], j_list[i]] = 1

    if visuals_on:
        # Update animation frame.
        if step % N_skip == 0:
            canvas.delete('all')
            spins = []
            for i in range(Ni):
                for j in range(Nj):
                    spin_color = '#FFFFFF' if sl[i,j] == 1 else '#000000'
                    spins.append(
                        canvas.create_rectangle(
                            j / Nj * window_size,
                            i / Ni * window_size,
                            (j + 1) / Nj * window_size,
                            (i + 1) / Ni * window_size,
                            outline='',
                            fill=spin_color,
                        )
                    )

    tk.title(f'Iteration {step}')
    tk.update_idletasks()
    tk.update()
    time.sleep(0.1) # Increase to slow down the simulation.

    if step >= N_iterations - 200:
        m_many += compute_magnetization(sl)
        n_average += 1

    step += 1

    if step == N_iterations:
        running = False

m[T_iteration] = m_many/n_average

```

```

print(f'For T: {T} -> m = {m[T_iteration]}')
T_iteration += 1

if visuals_on:
    tk.update_idletasks()
    tk.update()
    tk.mainloop() # Release animation handle (close window to finish).

```

```

For T: 0.1 -> m = 1.0
For T: 0.2 -> m = 1.0
For T: 0.5 -> m = 1.0
For T: 1.0 -> m = 0.9991389999999987
For T: 2.0 -> m = 0.9152119999999997
For T: 2.1 -> m = 0.8425870000000001
For T: 2.2 -> m = 0.7934629999999998
For T: 2.3 -> m = 0.617978
For T: 2.4 -> m = 0.2816760000000001
For T: 2.5 -> m = 0.0433459999999999
For T: 2.6 -> m = -0.000367000000000005
For T: 2.7 -> m = -0.0033520000000000073
For T: 2.8 -> m = 0.017787
For T: 2.9 -> m = 0.01279399999999998
For T: 3.0 -> m = 0.01758299999999999
For T: 5.0 -> m = -0.004646000000000002

```

```

In [49]: plt.plot(T_list, m)
plt.vlines(2.269, 0, 1, 'k', linestyles='dashed', label = 'Theoretical value Tc')

temp_diffs = T_list[1:] - T_list[0:-1]
m_diffs = m[1:] - m[0:-1]
slope = m_diffs/temp_diffs

my_Tc = T_list[np.argmin(slope)]+temp_diffs[np.argmin(slope)]/2
print(f'Q2: My estimate of Tc = {my_Tc:.4f}.')
print('Reason: this is where the slope is steepest --> small changes in T impact')
print('which happens around Tc.')

plt.vlines(my_Tc, 0, 1, 'r', linestyles='dashed', label = f'My estimation of Tc')

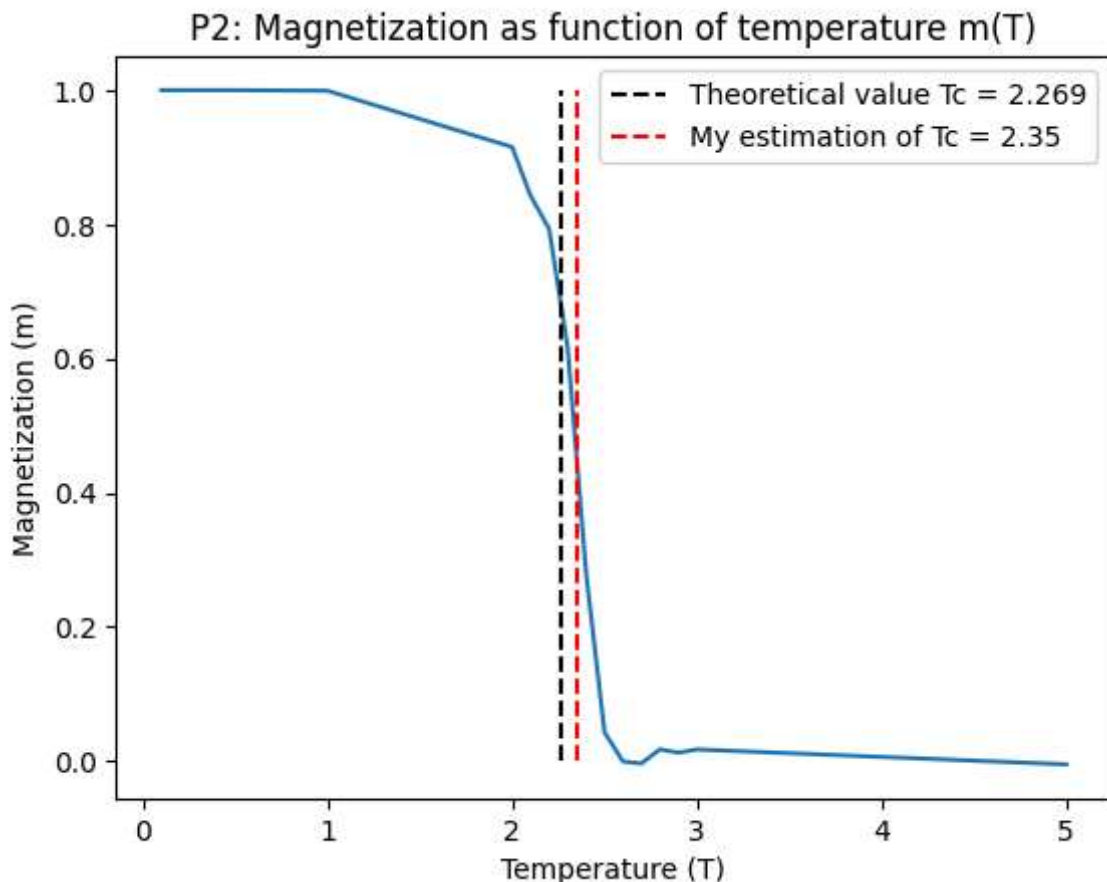
plt.xlabel('Temperature (T)')
plt.ylabel('Magnetization (m)')
plt.title('P2: Magnetization as function of temperature m(T)')
plt.legend()

```

Q2: My estimate of Tc = 2.35.

Reason: this is where the slope is steepest --> small changes in T impact m much, which happens around Tc.

Out[49]: <matplotlib.legend.Legend at 0x22e296c7ec0>



### Exercise 3

```
In [84]: import numpy as np

def grow_trees(forest, p):
    """
    Function to grow new trees in the forest.

    Parameters
    ======
    forest : 2-dimensional array.
    p : Probability for a tree to be generated in an empty cell.
    """

    Ni, Nj = forest.shape # Dimensions of the forest.

    new_trees = np.random.rand(Ni, Nj)

    new_trees_indices = np.where(new_trees <= p)
    forest[new_trees_indices] = 1

    return forest
```

```
In [85]: def propagate_fire(forest, i0, j0):
    """
    Function to propagate the fire on a populated forest.

    Parameters
    ======
    forest : 2-dimensional array.
    i0 : First index of the cell where the fire occurs.
```

```

j0 : Second index of the cell where the fire occurs.
"""

Ni, Nj = forest.shape # Dimensions of the forest.

fs = 0 # Initialize fire size.

if forest[i0, j0] == 1:
    active_i = [i0] # Initialize the list.
    active_j = [j0] # Initialize the list.
    forest[i0, j0] = -1 # Sets the tree on fire.
    fs += 1 # Update fire size.

while len(active_i) > 0:
    next_i = []
    next_j = []
    for n in np.arange(len(active_i)):
        # Coordinates of cell up.
        i = (active_i[n] + 1) % Ni
        j = active_j[n]
        # Check status
        if forest[i, j] == 1:
            next_i.append(i) # Add to list.
            next_j.append(j) # Add to list.
            forest[i, j] = -1 # Sets the current tree on fire.
            fs += 1 # Update fire size.

        # Coordinates of cell down.
        i = (active_i[n] - 1) % Ni
        j = active_j[n]
        # Check status
        if forest[i, j] == 1:
            next_i.append(i) # Add to list.
            next_j.append(j) # Add to list.
            forest[i, j] = -1 # Sets the current tree on fire.
            fs += 1 # Update fire size.

        # Coordinates of cell left.
        i = active_i[n]
        j = (active_j[n] - 1) % Nj
        # Check status
        if forest[i, j] == 1:
            next_i.append(i) # Add to list.
            next_j.append(j) # Add to list.
            forest[i, j] = -1 # Sets the current tree on fire.
            fs += 1 # Update fire size.

        # Coordinates of cell right.
        i = active_i[n]
        j = (active_j[n] + 1) % Nj
        # Check status
        if forest[i, j] == 1:
            next_i.append(i) # Add to list.
            next_j.append(j) # Add to list.
            forest[i, j] = -1 # Sets the current tree on fire.
            fs += 1 # Update fire size.

    active_i = next_i
    active_j = next_j

```

```
    return fs, forest
```

```
In [86]: def complementary_CDF(f, f_max):
    """
        Function to return the complementary cumulative distribution function.

    Parameters
    ======
    f : Sequence of values (as they occur, non necessarily sorted).
    f_max : Integer. Maximum possible value for the values in f.
    """

    num_events = len(f)
    s = np.sort(np.array(f)) / f_max # Sort f in ascending order.
    c = np.array(np.arange(num_events, 0, -1)) / (num_events) # Descending.

    c_CDF = c
    s_rel = s

    return c_CDF, s_rel
```

```
In [87]: N = 100 # Side of the forest.
N_list = np.array([16, 32, 64, 128, 256, 512, 1024])
p = 0.01 # Growth probability.
f = 0.2 # Lightning strike probability.
```

```
alphas = np.zeros(len(N_list))
```

```
target_num_fires = 300
```

```
for a_ind, N in enumerate(N_list):

    alpha_many = 0
    for i in range(10):
        forest = np.zeros([N, N]) # Empty forest.
        Ni, Nj = forest.shape # Sets the variables describing the shape.
        fire_size = [] # Empty list of fire sizes.
        fire_history = [] # Empty list of fire history.

        num_fires = 0
        while num_fires < target_num_fires:

            forest = grow_trees(forest, p) # Grow new trees.

            p_lightning = np.random.rand()
            if p_lightning < f: # Lightning occurs.
                i0 = np.random.randint(Ni)
                j0 = np.random.randint(Nj)
```

```

        fs, forest = propagate_fire(forest, i0, j0)
        if fs > 0:
            fire_size.append(fs)
            num_fires += 1

            fire_history.append(fs)

        else:
            fire_history.append(0)

        forest[np.where(forest == -1)] = 0

#print(f'Target of {target_num_fires} fire events reached')

c_CDF, s_rel = complementary_CDF(fire_size, forest.size)

min_rel_size = 1e-3
max_rel_size = 1e-1

is_min = np.searchsorted(s_rel, min_rel_size)
is_max = np.searchsorted(s_rel, max_rel_size)

# Note!!! The Linear dependence is between the Logarithms
fit = np.polyfit(np.log(s_rel[is_min:is_max]),
                 np.log(c_CDF[is_min:is_max]), 1)

beta = fit[0]
#print(f'The empirical cCDF has an exponent beta = {beta:.4}')

alpha = 1 - beta
alpha_many += alpha

#print(f'The empirical prob. distr. exponent: -alpha')
#print(f'with alpha = {alpha:.4}')
print(f'Size: {N}, iteration: {i+1}/10')
alphas[a_ind] = alpha_many/10.0

```

Size: 16, iteration: 1/10  
Size: 16, iteration: 2/10  
Size: 16, iteration: 3/10  
Size: 16, iteration: 4/10  
Size: 16, iteration: 5/10  
Size: 16, iteration: 6/10  
Size: 16, iteration: 7/10  
Size: 16, iteration: 8/10  
Size: 16, iteration: 9/10  
Size: 16, iteration: 10/10  
Size: 32, iteration: 1/10  
Size: 32, iteration: 2/10  
Size: 32, iteration: 3/10  
Size: 32, iteration: 4/10  
Size: 32, iteration: 5/10  
Size: 32, iteration: 6/10  
Size: 32, iteration: 7/10  
Size: 32, iteration: 8/10  
Size: 32, iteration: 9/10  
Size: 32, iteration: 10/10  
Size: 64, iteration: 1/10  
Size: 64, iteration: 2/10  
Size: 64, iteration: 3/10  
Size: 64, iteration: 4/10  
Size: 64, iteration: 5/10  
Size: 64, iteration: 6/10  
Size: 64, iteration: 7/10  
Size: 64, iteration: 8/10  
Size: 64, iteration: 9/10  
Size: 64, iteration: 10/10  
Size: 128, iteration: 1/10  
Size: 128, iteration: 2/10  
Size: 128, iteration: 3/10  
Size: 128, iteration: 4/10  
Size: 128, iteration: 5/10  
Size: 128, iteration: 6/10  
Size: 128, iteration: 7/10  
Size: 128, iteration: 8/10  
Size: 128, iteration: 9/10  
Size: 128, iteration: 10/10  
Size: 256, iteration: 1/10  
Size: 256, iteration: 2/10  
Size: 256, iteration: 3/10  
Size: 256, iteration: 4/10  
Size: 256, iteration: 5/10  
Size: 256, iteration: 6/10  
Size: 256, iteration: 7/10  
Size: 256, iteration: 8/10  
Size: 256, iteration: 9/10  
Size: 256, iteration: 10/10  
Size: 512, iteration: 1/10  
Size: 512, iteration: 2/10  
Size: 512, iteration: 3/10  
Size: 512, iteration: 4/10  
Size: 512, iteration: 5/10  
Size: 512, iteration: 6/10  
Size: 512, iteration: 7/10  
Size: 512, iteration: 8/10  
Size: 512, iteration: 9/10  
Size: 512, iteration: 10/10

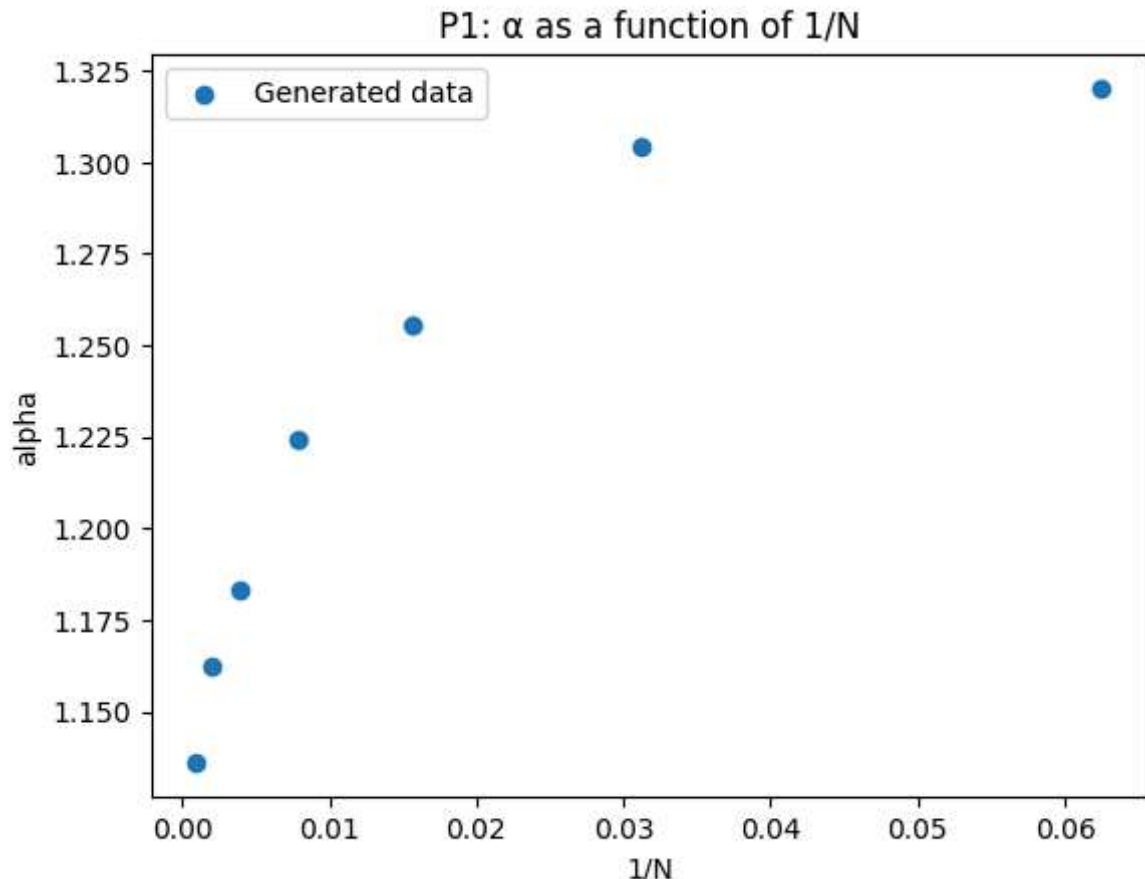
```
Size: 1024, iteration: 1/10
Size: 1024, iteration: 2/10
Size: 1024, iteration: 3/10
Size: 1024, iteration: 4/10
Size: 1024, iteration: 5/10
Size: 1024, iteration: 6/10
Size: 1024, iteration: 7/10
Size: 1024, iteration: 8/10
Size: 1024, iteration: 9/10
Size: 1024, iteration: 10/10
```

In [88]: `print(alphas)`

```
[1.3202274 1.30413252 1.25533152 1.22404548 1.18335414 1.16246048
1.13614403]
```

In [ ]: `from matplotlib import pyplot as plt
plt.scatter(1/N_list, alphas, label = 'Generated data')
#plt.xscale('log')
#plt.xticks(1/N_list, ['1/16', '1/32', '1/64', '1/128', '1/256', '1/512', '1/1024'])
plt.xlabel('1/N')
plt.ylabel('alpha')
plt.title('P1: alpha as a function of 1/N')
plt.legend()`

Out[ ]: <matplotlib.legend.Legend at 0x22e2ce26a80>



In [ ]: `a_fit = np.polyfit(np.log(1/N_list), alphas, 1)`

```
plt.plot(np.log(1/N_list), np.log(1/N_list)*a_fit[0] + a_fit[1], 'k--', label =
```

```

plt.scatter(np.log(1/N_list), alphas, label = 'Generated data')

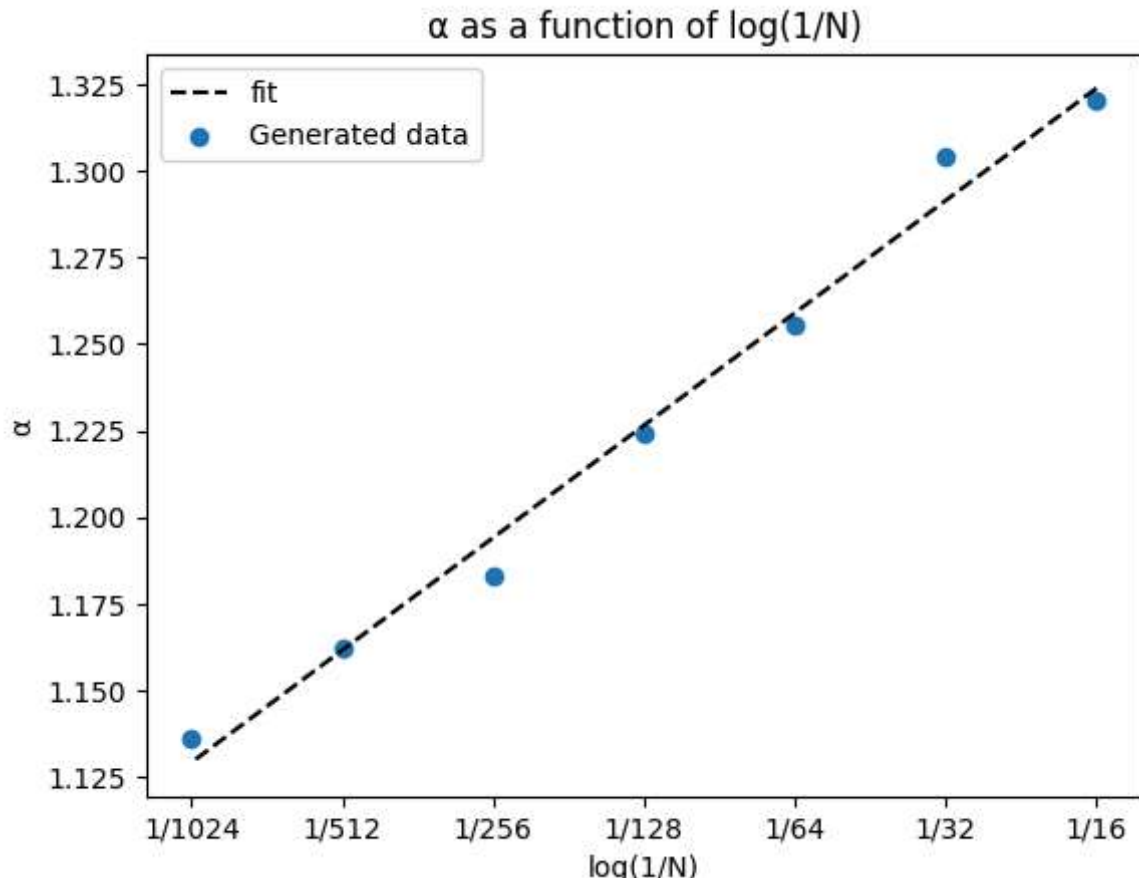
plt.xticks(np.log(1/N_list), ['1/16', '1/32', '1/64', '1/128', '1/256', '1/512'],

plt.xlabel('log(1/N)')
plt.ylabel('α')
plt.title('α as a function of log(1/N)')
plt.legend()

print('Using linear fit of log(1/N):')
print(f'Q1: α when N^(-1) -> 0: -inf')

```

Using linear fit of log(1/N):  
Q1:  $\alpha$  when  $N^{-1} \rightarrow 0$ : -inf



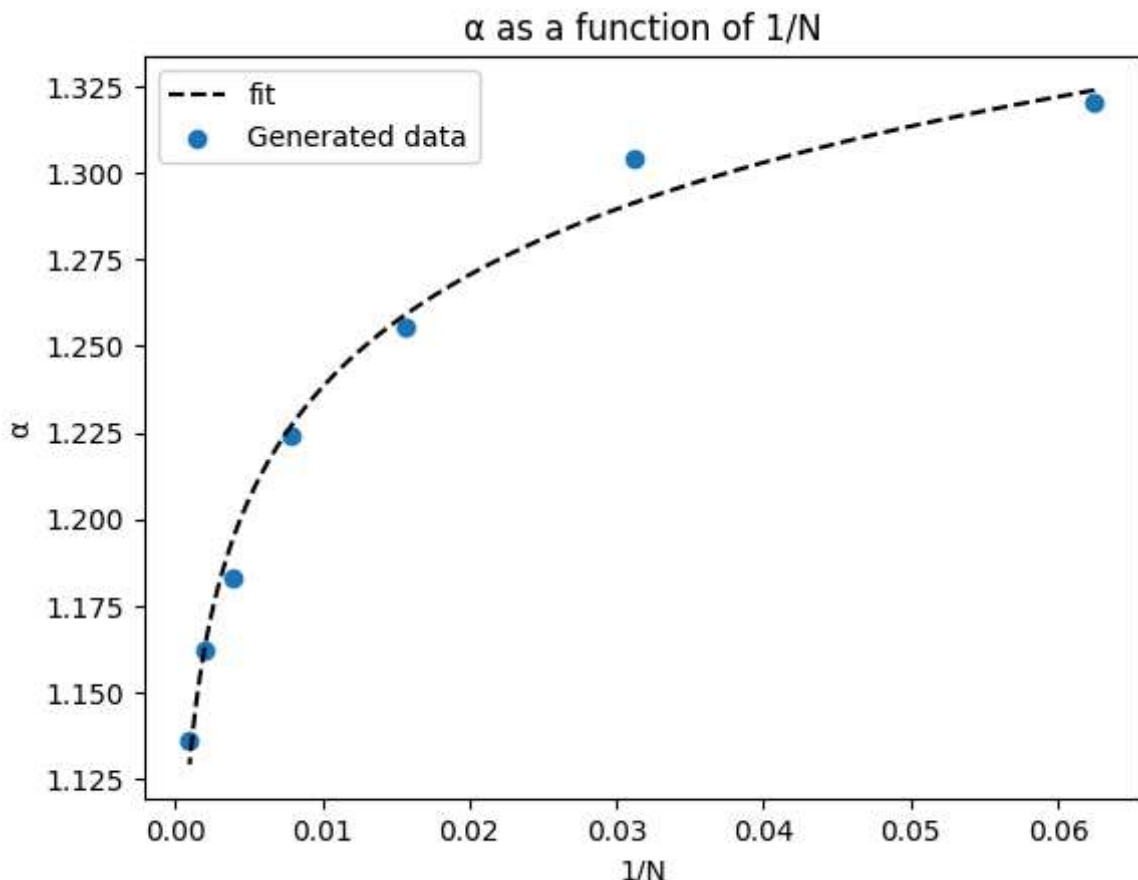
In [107]:

```

N_space = np.linspace(N_list[0], N_list[-1], 1000)
plt.plot(np.exp(np.log(1/N_space)), np.log(1/N_space)*a_fit[0] + a_fit[1], 'k--'
plt.scatter(np.exp(np.log(1/N_list)), alphas, label = 'Generated data')
plt.xlabel('1/N')
plt.ylabel('α')
plt.title('α as a function of 1/N')
plt.legend()

```

Out[107]: <matplotlib.legend.Legend at 0x22e2d1645f0>



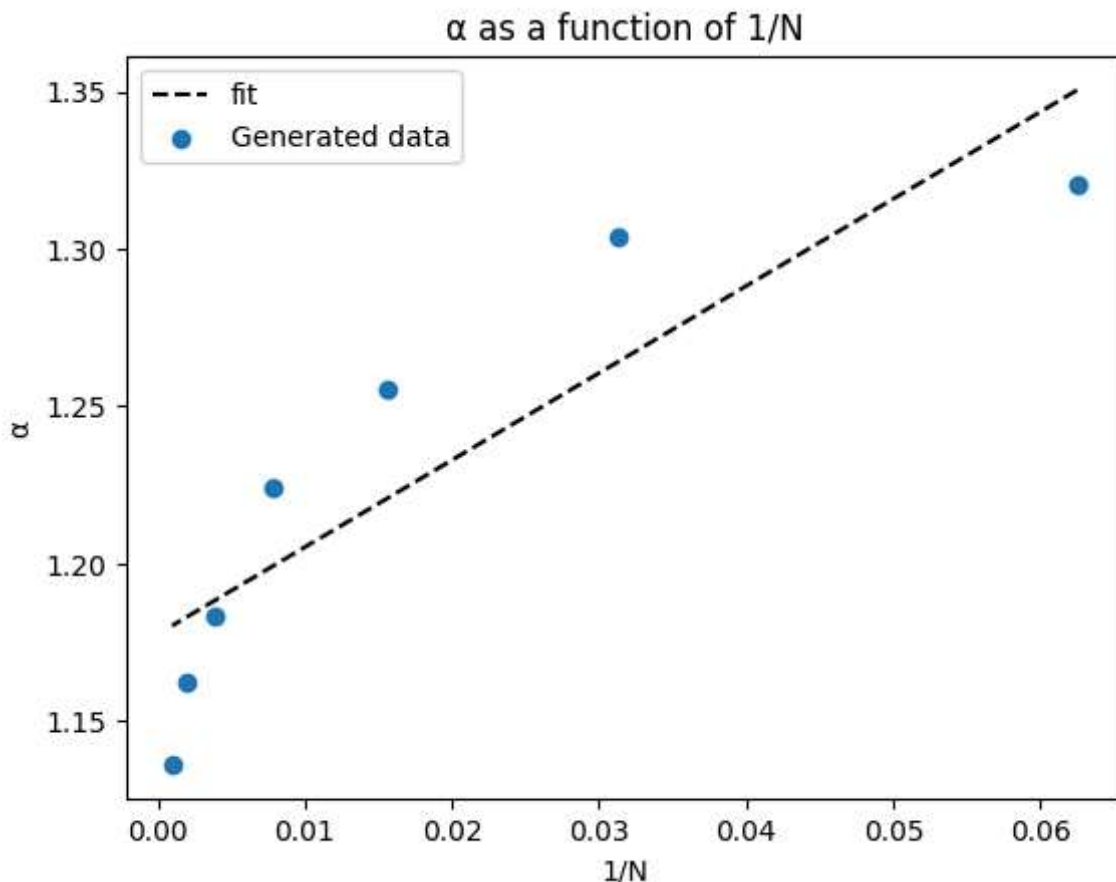
```
In [113]: a1_fit = np.polyfit(1/N_list, alphas, 1)

plt.plot(1/N_list, 1/N_list*a1_fit[0] + a1_fit[1], 'k--', label = 'fit')
plt.scatter(1/N_list, alphas, label = 'Generated data')

plt.xlabel('1/N')
plt.ylabel('alpha')
plt.title('alpha as a function of 1/N')
plt.legend()

print('When using a linear fit to unmodified data:')
print(f'Q1: alpha when N^(-1) -> 0: {a1_fit[1]:.4f}')
```

When using a linear fit to unmodified data:  
Q1:  $\alpha$  when  $N^{-1} \rightarrow 0$ : 1.178



Q2: My first thought when seeing the data was that it looked logarithmic so after plotting  $\alpha$  as a function of  $\log(1/N)$  it seemed obvious that a linear fit to  $\alpha(\log(1/N))$  would work well. As seen in the plots this fit is good but the result when extrapolating to  $1/N \rightarrow 0$  gives  $\alpha = -\infty$ . This is far from the estimate of  $\alpha = 1.15$  given in Chapter 3 of the book. When realizing this I tried a linear fit to  $\alpha(1/N)$  as seen in Fig 3.6 in the book. This fit looks much worse than both my previous one and the one in the book. However, extrapolating to  $1/N \rightarrow 0$  gives  $\alpha = 1.178$  which is quite close to the estimated  $\alpha = 1.15$ .

#### Exercise 4

```
In [50]: def neighbors_Moore(status):
    """
    Function to return the number of neighbors for each cell in status.

    Parameters
    ======
    status : Current status.
    """

    # Initialize the neighbor count array
    n_nn = (
        np.roll(status, 1, axis=0) +  # Up.
        np.roll(status, -1, axis=0) +  # Down.
        np.roll(status, 1, axis=1) +  # Left.
        np.roll(status, -1, axis=1) +  # Right.
        np.roll(np.roll(status, 1, axis=0), 1, axis=1) +  # Up-Left.
        np.roll(np.roll(status, 1, axis=0), -1, axis=1) +  # Up-Right
        np.roll(np.roll(status, -1, axis=0), 1, axis=1) +  # Down-Left
        np.roll(np.roll(status, -1, axis=0), -1, axis=1)  # Down-Right
    )
```

```
)  
return n_nn
```

```
In [51]: def apply_rule_2d(rule_2d, status):  
    """  
        Function to apply a 2-d rule on a status. Return the next status.  
  
    Parameters  
    ======  
    rule_2d : Array with size [2, 9]. Describe the CA rule.  
    status : Current status.  
    """  
  
    Ni, Nj = status.shape # Dimensions of 2-D lattice of the CA.  
    next_status = np.zeros([Ni, Nj])  
  
    # Find the number of neighbors.  
    n_nn = neighbors_Moore(status)  
    for i in range(Ni):  
        for j in range(Nj):  
            next_status[i, j] = rule_2d[int(status[i, j]), int(n_nn[i, j])]  
  
    return next_status
```

```
In [72]: import time  
from tkinter import *  
  
visuals_on = False  
N = 100  
full_time = 1000  
alive = np.zeros([5, full_time])  
c = np.zeros([5, full_time])  
  
for alive_ind in range(5):  
    # Random initial state.  
    gol = np.random.randint(2, size=[N, N])  
  
    rule_2d = np.zeros([2, 9])  
  
    # Game of Life's rules.  
    rule_2d[0, :] = [0, 0, 0, 1, 0, 0, 0, 0, 0] # New born from empty cell.  
    rule_2d[1, :] = [0, 0, 1, 1, 0, 0, 0, 0, 0] # Survival from living cell.  
  
    Ni, Nj = gol.shape # Sets the variables describing the shape.  
    N_skip = 1 # Visualize status every N_skip steps.  
    window_size = 600  
  
    if visuals_on:  
        tk = Tk()  
        tk.geometry(f'{window_size + 20}x{window_size + 20}')  
        tk.configure(background='#000000')  
  
        canvas = Canvas(tk, background='#ECECEC') # Generate animation window.  
        tk.attributes('-topmost', 0)  
        canvas.place(x=10, y=10, height=window_size, width=window_size)  
  
        step = 0
```

```

def stop_loop(event):
    global running
    running = False
if visuals_on:
    tk.bind("<Escape>", stop_loop) # Bind the Escape key to stop the Loop.
running = True # Flag to control the loop.
while running:
    alive[alive_ind, step] = np.sum(gol) # Count how many calls are alive
    prev_gol = gol # Save previous gol to compute C(t)

    gol = apply_rule_2d(rule_2d, gol)

    gol_switch = abs(prev_gol - gol) # Compare to know if cells switch
    nr_of_switches = np.sum(gol_switch)
    c[alive_ind, step] = nr_of_switches
    if visuals_on:
        # Update animation frame.
        if step % N_skip == 0:
            canvas.delete('all')
            gol_cells = []
            for i in range(Ni):
                for j in range(Nj):
                    gol_cell_color = '#FFFFFF' if gol[i, j] == 1 \
                        else '#000000'
                    gol_cells.append(
                        canvas.create_rectangle(
                            j / Nj * window_size,
                            i / Ni * window_size,
                            (j + 1) / Nj * window_size,
                            (i + 1) / Ni * window_size,
                            outline='',
                            fill=gol_cell_color,
                        )
                    )

    tk.title(f'Iteration {step}')
    tk.update_idletasks()
    tk.update()
    time.sleep(0.1) # Increase to slow down the simulation.
    if step == full_time-1:
        running = False

    step += 1

if visuals_on:
    tk.update_idletasks()
    tk.update()
    tk.mainloop() # Release animation handle (close window to finish).

```

## Task 1

In [115...]

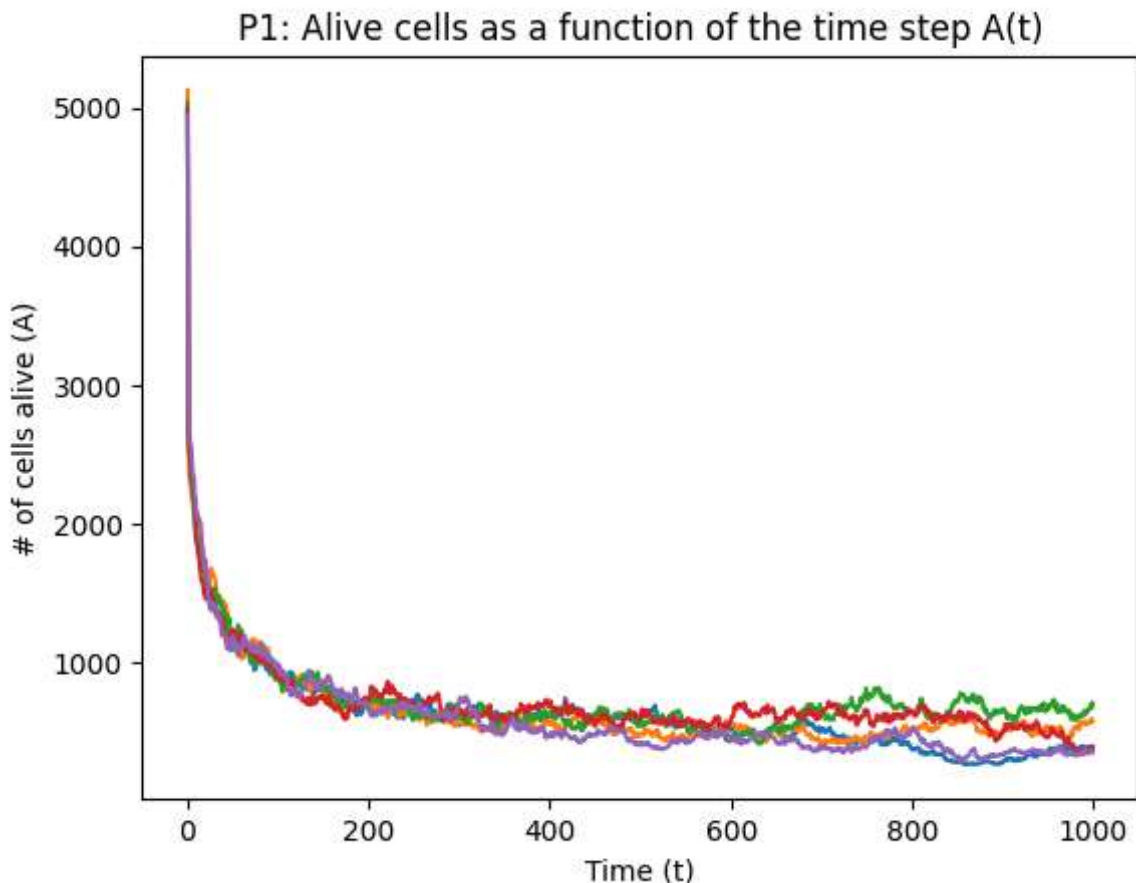
```

for i in range(alive.shape[0]):
    plt.plot(alive[i])

plt.xlabel('Time (t)')
plt.ylabel('# of cells alive (A)')
plt.title('P1: Alive cells as a function of the time step A(t)')

```

Out[115...]: Text(0.5, 1.0, 'P1: Alive cells as a function of the time step A(t)')



In [74]: `alive_density = alive/(N**2)`

In [75]: `ensemble_mean_alive_density = np.mean(alive_density, axis=0)`

I think Q1 here is ambiguous so forgive me for giving several answers.

```
In [114...]: plt.plot(ensemble_mean_alive_density)

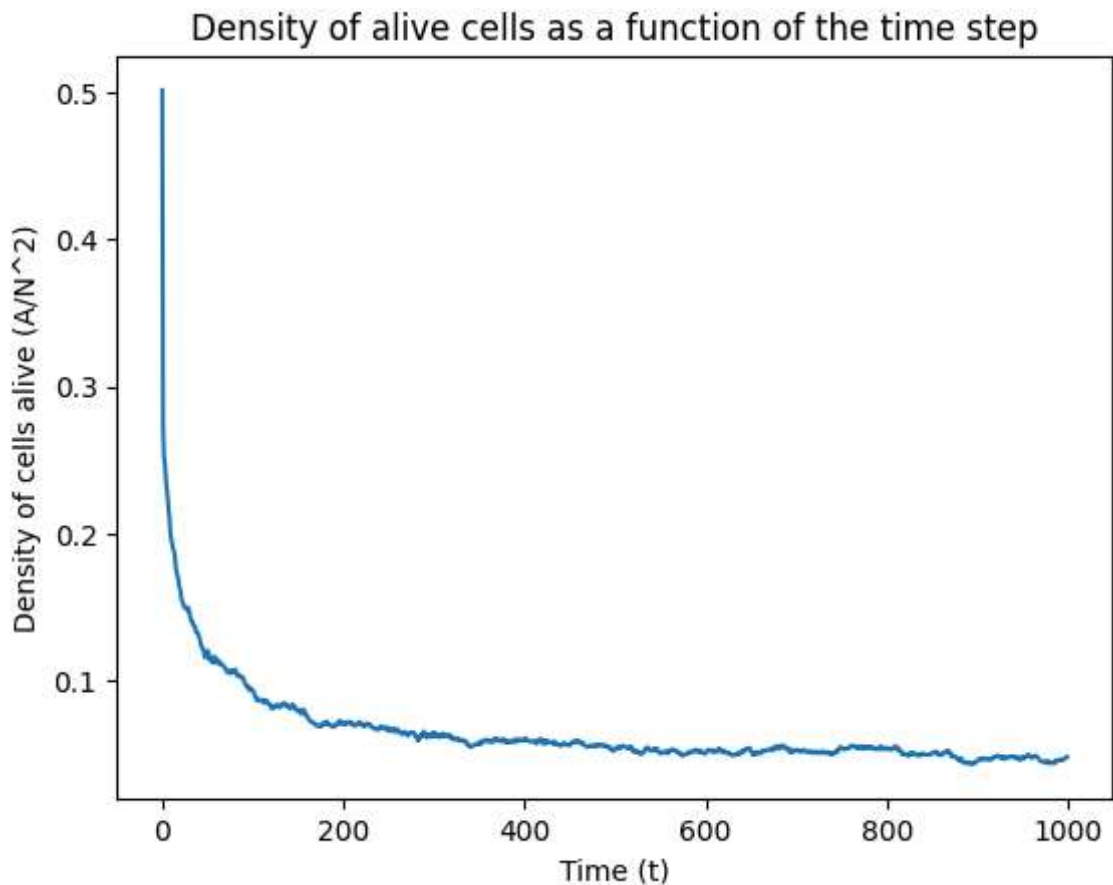
plt.xlabel('Time (t)')
plt.ylabel('Density of cells alive (A/N^2)')
plt.title('Density of alive cells as a function of the time step')

time_average_density = np.mean(ensemble_mean_alive_density)
time_average_density_stable = np.mean(ensemble_mean_alive_density[400:])

print(f'Q1: The average density of alive cell per unit area is {time_average_den')
print(f'Q1: The average density of alive cell per unit area after stabilizing is
```

Q1: The average density of alive cell per unit area is 0.06553.

Q1: The average density of alive cell per unit area after stabilizing is 0.05196.

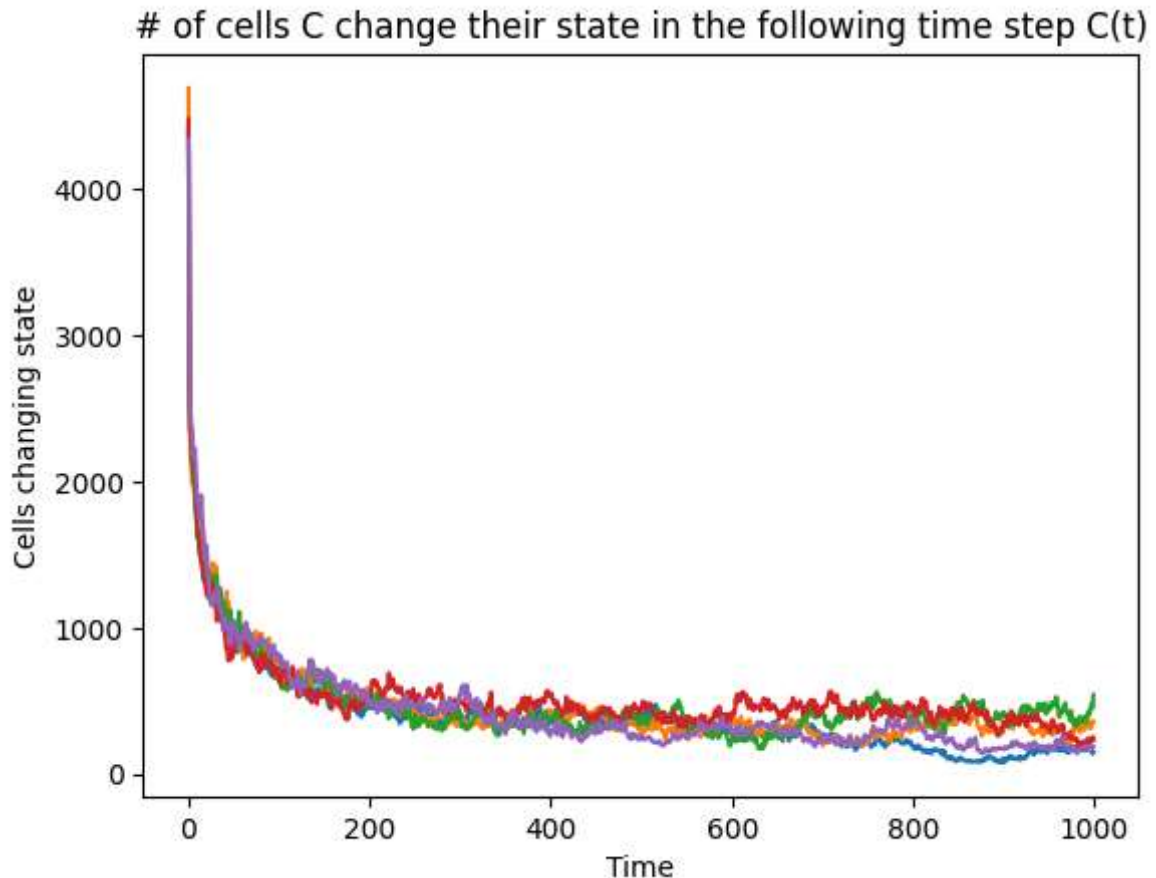


Q2: After approximately 400 iterations a steady state is reached with a density around 0.05.

#### Task 2

```
In [81]: for i in range(c.shape[0]):  
    plt.plot(c[i])  
  
plt.xlabel('Time')  
plt.ylabel('Cells changing state')  
plt.title('# of cells C change their state in the following time step C(t)')
```

```
Out[81]: Text(0.5, 1.0, '# of cells C change their state in the following time step C(t)')
```



```
In [82]: for i in range(c.shape[0]):  
    plt.plot(c[i, 400:])  
  
plt.xlabel('Time after stabilizing')  
plt.ylabel('Cells changing state')  
plt.title('P2: # of cells C change their state in the following time step C(t)')
```

```
Out[82]: Text(0.5, 1.0, 'P2: # of cells C change their state in the following time step  
C(t)')
```

P2: # of cells C change their state in the following time step C(t)

