## calculation

April 5, 2025

## 1 6 Pair correlation and nearest neighbors

plt.title("Pair correlation functions at triple point")

plt.xlabel('r')
plt.ylabel('g(r)')

#plt.xlim([0,3.5])

plt.legend()

plt.show()

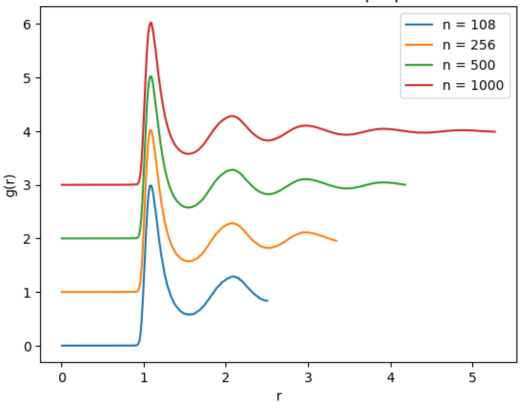
First, let us display the pair correlation function g(r) for each of the states.

```
[193]: import numpy as np
  import matplotlib.pyplot as plt
  import os

[194]: particleNumber = [108,256,500,1000]
  offset = 1 # offset by which each correlation function is offset to see it better

  for i, m in enumerate(particleNumber):
       r,g = np.loadtxt(f'n{m}/amclj.dat', unpack = True)
       plt.plot(r,g+offset*i, label = f"n = {m}")
```

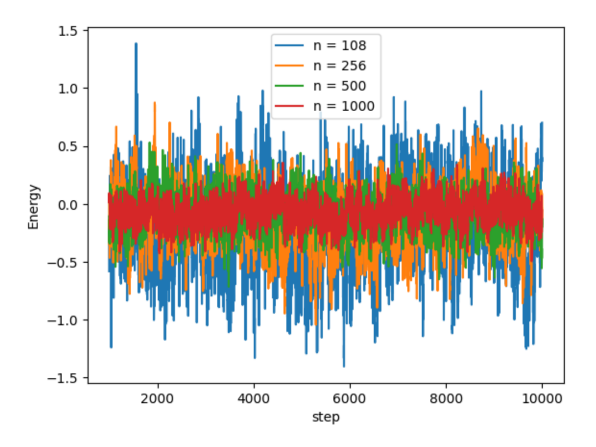




```
[195]: starting_step = 1000

for i, m in enumerate(particleNumber):
    step, acceptance, energy, pressure = np.loadtxt(f"n{m}/data.dat", unpack = True)
    plt.plot(step[starting_step:], pressure[starting_step:], label = f"n = {m}")

plt.legend()
    plt.xlabel("step")
    plt.ylabel("Energy")
    plt.show()
```



```
def calculate_correlation_time(variable):
    n = len(variable)

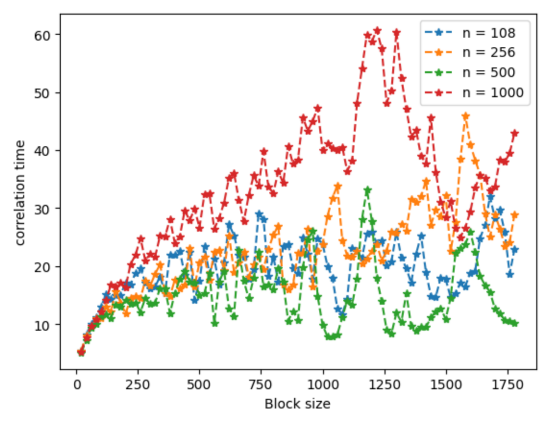
m = np.arange(20,n//5,20)

correlation_time = np.zeros(m.shape)
    var = np.var(variable)

for j, mm in enumerate(m):

# Trim the variable so you can do calculations
    length = n - n % mm
    variable_ = variable.copy()[:length]
    #print(len(variable)-mm)

Am = np.reshape(variable_, (-1,mm))
    Am_average = Am.mean(axis=1)
    correlation_time[j] = np.var(Am_average)*mm/var/2
    return m, correlation_time
```



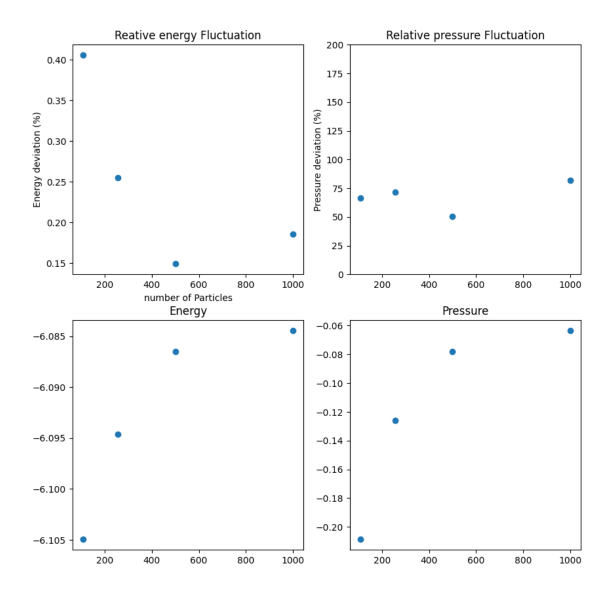
```
[197]: # Calculate the stderrors:
starting_step = 2000
```

```
minimalAcceptedBlockSize = 400
energyDeviations = []
pressureDeviations = []
energies = []
pressures = []
print(f"{'_'*80}")
print(f"{'Energy':^80}")
print(f"{'_'*80}")
print(f"{'n':<10}{f'Mean':<30}{f'correlated stdev':<20}{f'relative Error':}\n")
for i, m in enumerate(particleNumber):
    step, acceptance, energy, pressure = np.loadtxt(f"n{m}/data.dat", unpack = u
→True)
    stepTruncated = step[starting_step:]
    energyTruncated = energy[starting_step:]
    pressureTruncated = pressure[starting_step:]
    ### Energy
    binSize, correlationTime = calculate_correlation_time(energyTruncated)
    averageEnergyCorrelationTime = np.mean(correlationTime[binSize >=_
→minimalAcceptedBlockSize])
    energyMean = np.mean(energyTruncated)
    energyStdev = np.std(energyTruncated, ddof = 1)
    energyCorrStdev = energyStdev * np.sqrt(2 * averageEnergyCorrelationTime / ___
 →minimalAcceptedBlockSize)
    energies.append(energyMean)
    energyDeviations.append(energyCorrStdev/abs(energyMean)*100)
```

```
### pressure
           binSize, correlationTime = calculate_correlation_time(pressureTruncated)
           averagePressureCorrelationTime = np.mean(correlationTime[binSize >=_
   →minimalAcceptedBlockSize])
           pressureMean = np.mean(pressureTruncated)
           pressureStdev = np.std(pressureTruncated, ddof= 1)
           pressureCorrStdev = pressureStdev * np.sqrt(2 *__
   →averagePressureCorrelationTime / minimalAcceptedBlockSize)
           pressures.append(pressureMean)
           pressureDeviations.append(pressureCorrStdev/abs(pressureMean)* 100)
           ###
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  \rightarrowabs(energyMean) * 100 :>0.2f} \frac{\ln n}{\ln n}
print(f"{'_'*80}")
#### Plotting it:
fig, axes = plt.subplots(2, 2, figsize=(10,10))
[ax1, ax2], [ax3, ax4] = axes[:,:]
ax1.set_xlabel('number of Particles')
ax1.set_ylabel('Energy deviation (%)')
ax1.set_title("Reative energy Fluctuation")
ax1.scatter(particleNumber, energyDeviations)
ax2.scatter(particleNumber, pressureDeviations)
ax2.set_title("Relative pressure Fluctuation")
ax2.set_ylabel("Pressure deviation (%)")
ax2.set_ylim((0,200))
ax3.scatter(particleNumber, energies)
ax3.set_title("Energy")
```

```
ax4.scatter(particleNumber, pressures)
ax4.set_title("Pressure")
plt.show()
```

Energy			
n	Mean	correlated stdev	relative Error
108	-6.10e+00	2.48e-02	0.41 %
256	-6.09e+00	1.56e-02	0.26 %
500	-6.09e+00	9.08e-03	0.15 %
1000	-6.08e+00	1.13e-02	0.19 %
1000	-6.08e+00	1.13e-02	0.19 %



```
[198]: from scipy.signal import argrelmin

nOffset = 5 # offset for nice displaying

rho = 0.84

for i, m in enumerate(particleNumber):
    r,g = np.loadtxt(f'n{m}/amclj.dat', unpack = True)

    integrand = r**2 * g

    n = []
    coordination_number = []
```

Coordination number for n = 108: 12.5975

R\_min: 1.5450

Coordination number for n = 256: 12.5413

R\_min: 1.5450

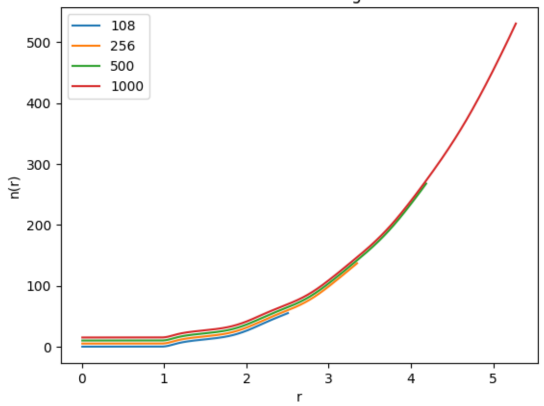
Coordination number for n = 500: 12.6760

R\_min: 1.5550

Coordination number for n = 1000: 12.3817

R\_min: 1.5350

# Mean number of neighbors

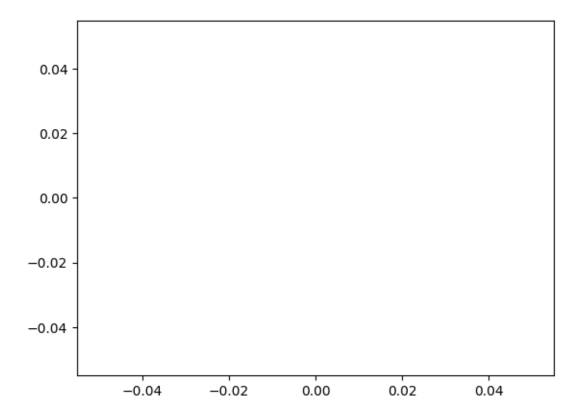


```
[199]: def extractPositionFromTraj(fileName, particleNumber, steps, saveEvery):
           data = np.loadtxt(fileName, delimiter="0", dtype=str)
           # Total number of rows
           numberOfRows = data.shape[0]
           skipRows = 2
           # Select every (particleNumber + skipRows) block, keeping only the first
        →particleNumber rows of each
           cleanStringList = []
           i = skipRows
           while i < numberOfRows:</pre>
               cleanStringList.append(data[i:i+particleNumber]) # take particleNumber_
        →rows
               i += particleNumber + skipRows
                                                         # skip next 'skipRews' rows
           del data
           # Stack the result into one array
```

[]:

```
[200]: showEvery = 100
       particleNumber = 108
       density = 0.84
       volume = density * particleNumber
       boxLength = volume**(1/3)
       print(boxLength)
       #boxLength = 3.904483
       boxLength = 5.047173
       positions = extractPositionFromTraj(f"n{particleNumber}/traj.xyz",__
       →particleNumber, 10000, 1)
       steps = 10000
       starting_step = 1000
       for step in range(starting_step,steps):
           centralParticlePosition = positions[step, 0, :]
           positions[step,:,:] -= centralParticlePosition
           positions[step,:,:] -= np.round(positions[step,:,:] / boxLength) * boxLength
       zThickness = 1.6
       zMask = np.all(positions[:, :, -1] < zThickness, axis=1)</pre>
       filteredPositions = positions[zMask]
       plt.plot(filteredPositions[starting_step:,:,0], filteredPositions[starting_step:
       \rightarrow,:,1], 'o', markersize = 0.01, color = 'blue')
       plt.show()
```

#### 4.4933234323311035



### 2 7 The Structure Factor

We will calculate the structure factor S(k)

$$S(k) = 1 + \frac{4\pi\rho}{k} \times \int_0^\infty dr \ r \sin(kr)[g(r) - 1]$$

for our case with particleNumber = 1000. Technically, the integral goes to  $\infty$ , but the maximum r is at max(r)

[202]: max(r)

[202]: 5.275

We will just integrate using the trapezoidal rule and we define S(k) as a function of k.

```
integrand = r * np.sin(kr) * (g - 1)  # broadcasts over r
integrals = np.trapz(integrand, x=r, axis=1) # integrate over r for each k
return 1 + 4 * np.pi * rho * integrals / (k + 1e-12)

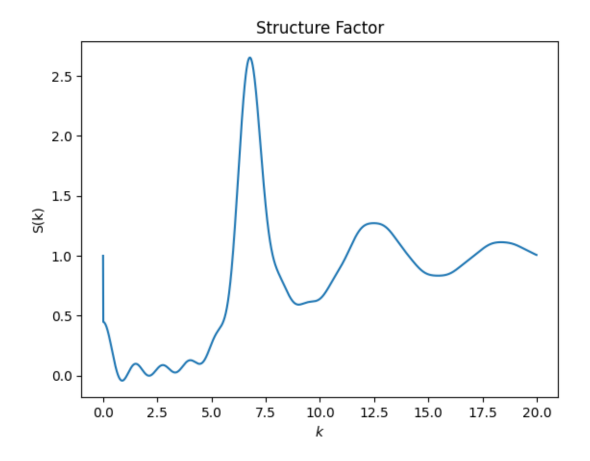
structureFactor(1)
```

[203]: array([-0.02799143])

```
[204]: k = np.arange(0,20,.01)

plt.plot(k, structureFactor(k))
plt.xlabel("$k$")
plt.ylabel("S(k)")
plt.title("Structure Factor")
```

[204]: Text(0.5, 1.0, 'Structure Factor')



Now we will try to recreate the Pair correlation function using the integral

$$g(r) = 1 + \frac{1}{2\pi^2 \rho r} \times \int_0^\infty dk \ k \sin(kr) [S(k) - 1]$$

Because we cannot calculate the integral to infinity, we will use a cutoff (The standard cutoff is  $20/\sigma$ ): cutoff

To discretize k, we will use a kStep:

np.arange(0, cutoff, kStep)

```
[205]: cutoff = 50 # sigma = 1
kStep = 0.01
k = np.linspace(0, cutoff, 528)

def gReconstructed(r):
    # kr: shape (len(r), len(k)) = outer product of r and k
    kr = np.outer(r, k)

    # integrand: shape (len(r), len(k))
    integrand = k * np.sin(kr) * (structureFactor(k) - 1) # S_k is shape
    integrate along k-axis (axis=1) for each r
    integrals = np.trapz(integrand, x=k, axis=1)

# final g(r)
    denominator = 2 * np.pi**2 * rho * r + 1e-12
    result = 1 + integrals / denominator
    return result
```

