## Molecular Dynamics with C++ report

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## 1 Introduction

Molecular dynamics simulations are becoming an essential part of modern technology, because they allow to study the behavior of complex systems in a controlled way. The simulation of molecular systems is a very complex task, because the number of atoms in a system can be very large. Therefore, the simulation of a system requires a lot of computational power. In the last years, the computational power of computers has increased dramatically. This development has led to the fact that molecular dynamics simulations are now possible for systems with millions of atoms. However, the simulation of such systems is still a very demanding task. Therefore, the simulation of such systems is usually done on supercomputers or clusters of computers.

In this report, we will discuss the design and Implementation of a molecular dynamics simulations for atoms and molecules. We will also discuss the different potential forces that are used in the simulations, and how they are implemented, initializing atomic systems in random positions and preserve the atoms from evaporating using Thermostates, how to make the simulation run faster by using only neighbor list search, and how to parallelize the simulation using MPI.

## 2 Methods

Here we will discuss the different methods that we will use in our simulation, then in the next chapter we will discuss the implementation of these methods.

#### 2.1 Velocity-Verlet integrator

The velocity-Verlet algorithm is a numerical method based on the Taylor expansion of the position and velocity of the particles for solving the equations of motion of a system of particles. It is a symplectic integrator, which means that it conserves the total energy of the system that's because the velocity is calculated at the middle of the time step, and the position is calculated at the end of the time step. This means that the velocity is calculated at the same time as the acceleration, and the position is calculated at the same time as the velocity. This means that the total energy of the system is conserved. A disadvantage of the velocity-Verlet algorithm is that it is not very accurate. The error in the position is proportional to  $\Delta t^2$ , and the error in the velocity is proportional to  $\Delta t$ . This means that the velocity-Verlet algorithm is not very accurate for large time steps. The velocity-Verlet algorithm is given by the following equations:

- 1. Calculate the acceleration of the particles at time t.
- 2. Calculate the velocity of the particles at time  $t + \frac{1}{2}\Delta t$ .

- 3. Calculate the position of the particles at time  $t + \Delta t$ .
- 4. Calculate the acceleration of the particles at time  $t + \Delta t$ .
- 5. Calculate the velocity of the particles at time  $t + \Delta t$ .

where  $\Delta t$  is the time step.

#### 2.2 Lennard-Jones potential force

The Lennard-Jones potential is a potential that is used in molecular dynamics simulations to model the interatomic interactions between atoms and molecules. It models both the attractive and repulsive forces between atoms and molecules in a system in a simple way. It is a simple model because it only depends on the distance between the atoms and molecules. It is not a very accurate model though because it does not take into account the charge of the particles, and the different types of interactions between the particles. This potential is very popular to use in molecular dynamics simulations because it is simple to use, and it is computationally cheap. It can also be used to model different types of interactions between atoms and molecules, like the interaction between the electrons of an atom with the electrons of another atom, the interaction between the electrons of an atom with the nuclei of another atom, and the interaction between the nuclei of an atom with the nuclei of another atom.

The Lennard-Jones potential is given by:

$$U(r) = 4\epsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{6} \right] \tag{1}$$

where  $\epsilon$  is the depth of the potential well, and  $\sigma$  is the distance at which the potential is at a minimum. The force is given by the following equation:

$$F(r) = -\frac{dV(r)}{dr} = 24\epsilon \left[ \frac{2}{r} \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{6} \right]$$
 (2)

The force is calculated by using the following steps:

- 1. Calculate the distance between all pairs of atoms.
- 2. Calculate the potential between all pairs of atoms.
- 3. Calculate the force between all pairs of atoms.

#### 2.3 Thermostat

The thermostat is an algorithm that is used to preserve the atoms from evaporating. The idea is when we initialize the atoms in random positions, the atoms will try to get to the equilibrium position. This means that the atoms will move very fast and will collide with each other and the temperature of the system will increase very fast. This will lead to the atoms evaporation. The thermostat is used to prevent this from happening. The thermostat is used to slow down the atoms and make them move slower until they reach the equilibrium position and the temperature of the system is stable.

The Berendsen thremostate is the thermostat that we will use in our simulation. It's one of the most popular thermostats. It is a simple thermostat that is easy to implement. To maintain the temperature, the system is coupled to an external heat bath with fixed Temperature  $T_0[1]$ . The velocities of the particles are rescaled with a

factor  $\lambda$  at each time step:

$$\lambda = \sqrt{1 + \frac{\Delta t}{\tau} \left(\frac{T_0}{T} - 1\right)} \tag{3}$$

where  $\tau$  is the relaxation time of the thermostat, and T is the temperature of the system. The relaxation time is the time it takes for the system to reach the equilibrium temperature. The relaxation time is an important parameter in the thermostat. If the relaxation time is too small, there will be a strong coupling between the system and the heat bath, this means that the system will reach the equilibrium temperature very fast. And if the relaxation time is too big, there will be a weak coupling between the system and the heat bath, this means that the system will take a long time to reach the equilibrium temperature. And in any case it needs to be bigger than the time step, otherwise the system will not reach the equilibrium temperature.

The algorithm is as follows:

- 1. Calculate the temperature of the system.
- 2. Calculate the temperature ratio  $\frac{T_0}{T}$ .
- 3. Calculate the scaling factor  $\sqrt{\frac{T_0}{T}}$ .
- 4. Rescale the velocity of the particles in the system with the scaling factor.

where T is the temperature of the system,  $T_0$  is the target temperature, and  $\frac{T}{T_0}$  is the temperature ratio.

### 2.4 Neighbor list search

The problem of using potential forces in molecular dynamics simulations is that the force is calculated between all pairs of atoms. This means that the computational

cost of the simulation is proportional to  $N^2$ , where N is the number of atoms in the system. This is a very big computational cost, and it is not very efficient. To solve this problem, we can use a neighbor list search. It's an algorithm that is used to speed up the calculation of the forces that act on the particles. It is based on the idea of only calculating the force between atoms that are close to each other. This means that we will only calculate the force between atoms that are in the same cell, and the neighboring cells. Therefor the computational cost of the simulation becomes proportional to N instead of  $N^2$ . And this is a very big improvement in the computational cost of the simulation.

The neighbor list search is calculated by using the following steps:

- 1. Calculate the distance between all particles.
- 2. If the distance between two particles is smaller than the threshold value, then calculate the force between them.
- 3. If the distance between two particles is larger than the threshold value, then do not calculate the force between them.

#### 2.5 Embedded-atom method potential force

The embedded-atom method (EAM) is a method for calculating the forces between atoms. It is based on the idea that the forces between atoms are dependent on the density of the atoms. EAM is a potential force that is used in the simulation of metals, alloys, and intermetallic compounds.

The EAM potential force is as follows:

## 2.6 Parallelization

1. Initialize the MPI environment.

The parallelization of the simulation is done using MPI. The algorithm is as follows:

2.	Get the number of processes.
3.	Get the rank of the process.
4.	Initialize the atoms.
5.	Calculate the forces.
6.	Calculate the total energy.
7.	Calculate the total momentum.
8.	Calculate the temperature.
9.	Calculate the pressure.
10.	Print the results.
11	Finaliza the MPI environment

## 3 Implementation

### 3.1 Velocity-Verlet integrator

#### 3.1.1 Test strategy for Verlet integrator

We test the Verlet integrator by comparing the results of the Verlet integrator with the results of the analytical solution of the equations of motion of a particle. the analytical solution of the equations of motion of a particle is given by the following equations:

$$x_{i}(t+dt) = x_{i}(t) + v_{i}(t)dt + \frac{1}{2m}f_{i}(t)dt^{2}$$

$$v_{i}(t+dt) = v_{i}(t) + \frac{1}{2m_{i}}(f_{i}(t) + f_{i}(t+dt))dt$$
(4)

If we assume that the there is no acting forces (constant and equal to zero) on the particles, then the analytical solution of the equations of motion of a particle after N time steps is given by the following equations:

$$x_{i}(t + N * dt) = x_{i}(t) + \sum_{i=0}^{N} v_{i}(t)dt$$

$$v_{i}(t + N * dt) = v_{i}(t)$$
(5)

where dt is the time step, N is the number of time steps,  $x_i(t)$  is the position of the particle i at time t,  $v_i(t)$  is the velocity of the particle i at time t,  $f_i(t)$  is the force acting on the particle i at time t, and  $m_i$  is the mass of the particle i.

That means if we compare the result after all the integration steps of the two Verlet steps with the expected output of the analytical solution, then we can be sure that the Verlet integrator is working correctly.

#### 3.2 Lennard-Jones potential force

## 3.2.1 Derivation of the analytical expression for the forces of the Lennard-Jones potential

#### 3.3 Berendsen thermostat

talk a little bit about the Berendsen thermostat implementation

#### 3.3.1 Test strategy for Berendsen thermostat

We have implemented two test cases for the Berendsen thermostat. The first test case is a test case where we test the Berendsen thermostat on a system with a single particle. The second test case is a test case where we test the Berendsen thermostat on a cubic lattice of size 5x5x5 with lattice constant=1.12. The test cases are implemented in the file.

#### Test case 1: Berendsen thermostat on a system with a single particle

Here, we make use of the derived equation in the lecture notes:

$$T(t) = T_0 + (T_1 - T_0) * exp(-\frac{t}{\tau})$$
(6)

where T(t) is the temperature of the system at time t,  $T_1$  is the initial temperature of the system,  $T_0$  is the target temperature of the system, and  $\tau$  is relaxation time constant. and using this equation we can calculate the temperature of the system at every iteration t before and after the thermostat, and then make sure that the temperature relaxes exponentially to the target temperature.

# Test case 2: Berendsen thermostat on a cubic lattice of size 5x5x5 with lattice constant=1.12

This test is confirming if the temperature really converges to the target temperature after running the simulation for some time. So, the test is as follows:

- 1. Create a cubic lattice of size 5x5x5 with lattice constant=1.12.
- 2. using time\_step=0.01, target temperature=1.0, and relaxation time constant= 10\* time\_step.
- 3. Run the simulation for 10000 time steps with the Berendsen thermostat.
- 4. In the last half of the simulation (5000 time steps), calculate the system temperature in every time step and make sure that the temperature converges to the target temperature.

#### 3.4 Embedded atom method

## 3.5 Units and specification of the time unit

To work with EAM potential, we need to convert the units of the system to the units of the EAM potential. So, typically in Molecular dynamics simulations, the energy and the length are constants. and here we use E = eV and L = Angstrom. Where

$$1eV = 1.602176634 \times 10^{-19} J$$

$$1Angstrom = 10^{-10} m$$
(7)

and forces are typically calculated in units of eV/Angstrom, which is the natural force unit for this system of units. To conclude, the units of the EAM potential used are:

• Energy: eV

• Distance: Angstrom

• Mass: amu

• Temperature: K

• Time: fs

#### 3.5.1 time step for the gold potential

We have to fix either the mass unit or the time unit. And in this implementation we fixed the time unit of 1 fs for the gold potential. because using this time step is small enough to keep the energy conservation and also large enough to get to the melting point of gold in a reasonable time for clusters of layers from 3:12. To convert to a 1 fs time step we use the equation below:

$$m = \frac{[E][t^2]}{[L^2]} = 1.66 * 10^{-29} kg$$
 (8)

where m is the mass of the particle, E is the energy unit, t is the time unit, and L is the length unit. and the commonly used mass unit is amu, so we can convert the

mass unit to amu using the following equation:

$$m = 1.66 * 10^{-29} kg * 6.022 * 10^{23} kg/mol * 1000 = 0.009648 amu$$
 (9)

And accordingly, we multiply all atoms masses by 103.642 to get a time step of 1 fs.

## 3.6 Neighbor List

## 3.7 Parallelization using MPI

## 4 Results

# 4.1 Total energy as a function of time for different time steps

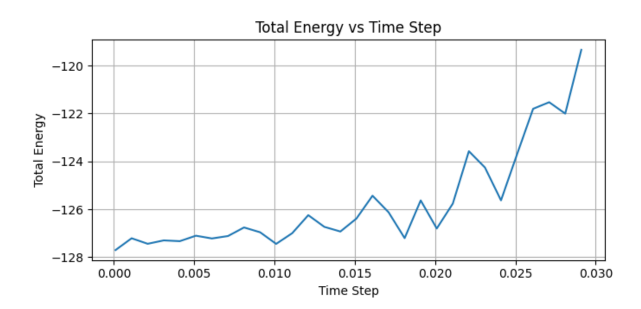


Figure 1: Total energy vs Time steps using parameters:  $\begin{array}{c} start\_time\_step=1e\text{-}4,\ end\_time\_step=30e\text{-}3,step=1e\text{-}3,\\ sigma=1,\ mass=1,\ epsilon=1,\ total\_time=5000 \end{array}$ 

## 4.2 Snapshots sequence of LJ simulation

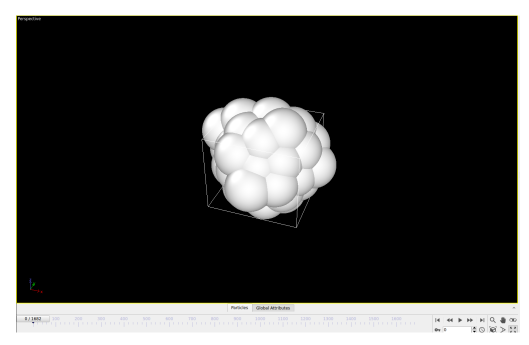


Figure 2: LJ simulation snapshot1

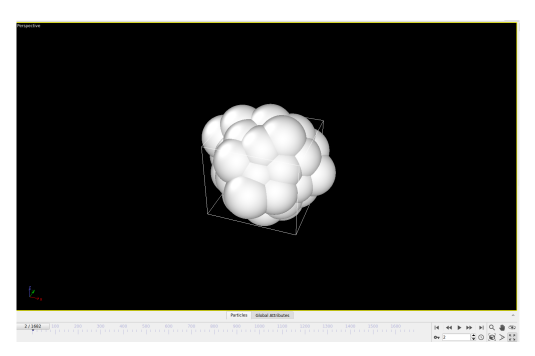


Figure 3: LJ simulation snapshot2

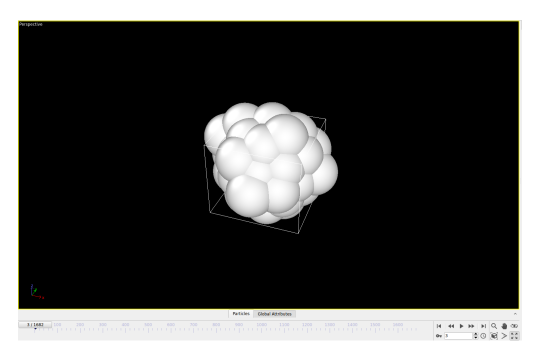
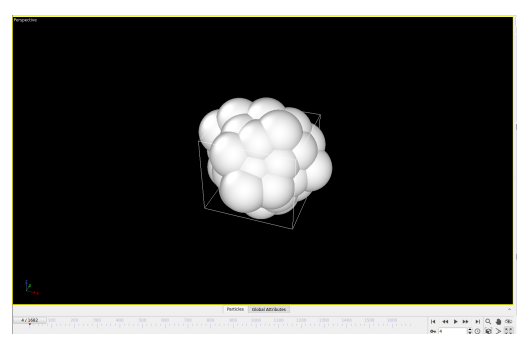


Figure 4: LJ simulation snapshot3



 $\textbf{Figure 5:} \ \, \text{LJ simulation snapshot4}$ 

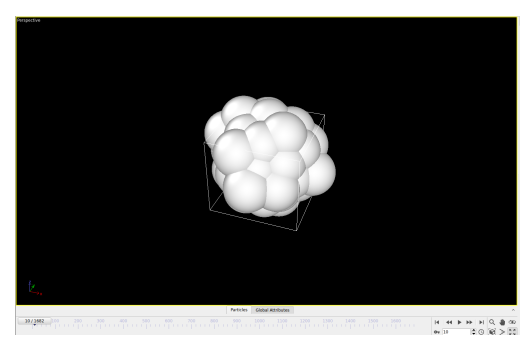
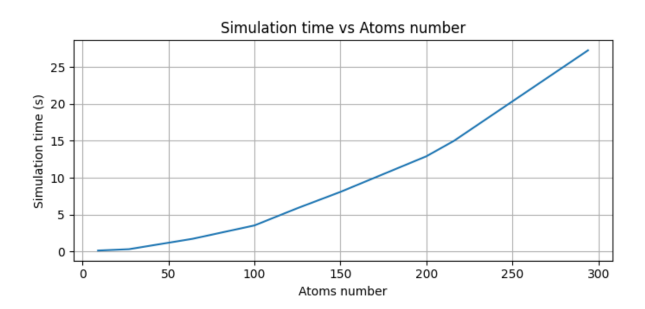


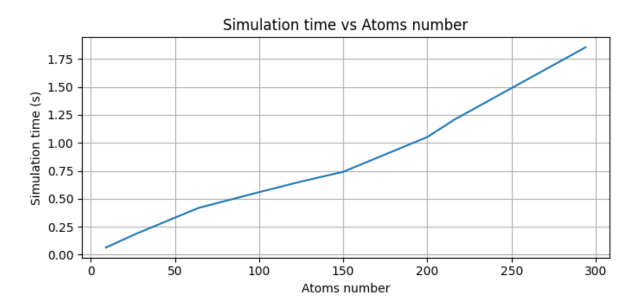
Figure 6: LJ simulation snapshot5

# 4.3 simulation time as a function of the size (number of atoms) without neighbor list



Here we notice that the simulation time increases quadratically with the number of atoms in the system. This is because in the energy update step in the LJ simulation we have to calculate the energy between all the pairs of atoms in the system. So the time complexity of the energy update step is  $O(N^2)$  where N is the number of atoms in the system. For example, In the figure above, The simulation time for 100 atoms is 3.5s and for 200 atoms is 12.9s which is almost 4 times more than the simulation time for 100 atoms.

# 4.4 simulation time as a function of the size (number of atoms) with neighbor list



Here, it's very clear the linearity of the simulation time with the number of atoms in the system. By introducing the concept of just calculating the energy between the atoms that are close to each other, we have reduced the time complexity of the energy update step from  $O(N^2)$  to O(N) where N is the number of atoms in the system. For example, In the figure above, The simulation time for 100 atoms is 0.56s and for 200 atoms is 1.05s which is almost 2 times more than the simulation time for 100 atoms.

### 4.5 total energy vs temperature

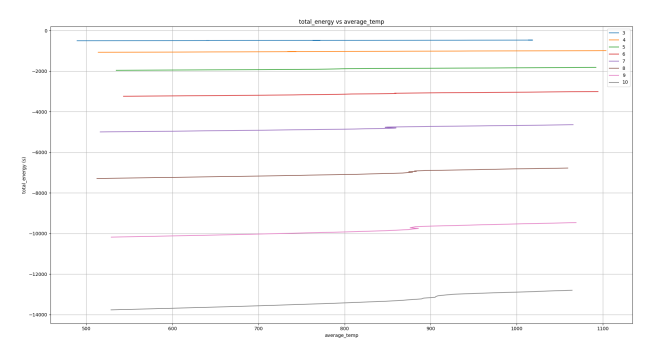


Figure 9: Total energy vs Temp using parameters: time\_step=10fs,  $mass=20405.7294, initial\_simulation\_time=10000, \\ relaxation\_time\_start = 10* time\_step, then after 500 step, it \\ increases to 1e10*time\_step, and cutoff\_radius = \\ 10, target\_temp = 500, number\_experiments=26, \\ experiment\_simulation\_time=500, \\ added\_energy\_in\_each\_experiment=0.01*no\_atoms, \\ cluster\_sizes=3:10, and atomic\_distance = 2.885$ 

## 4.6 melting point versus cluster size

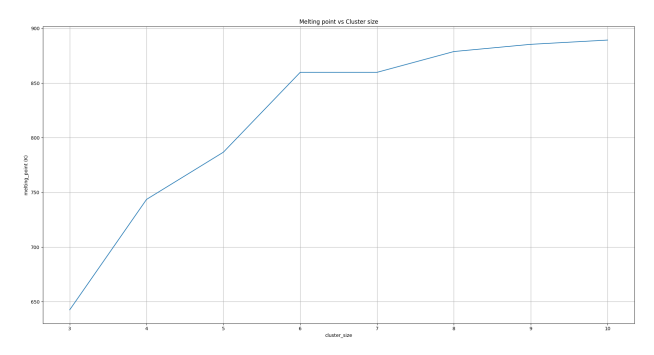


Figure 10: Melting point vs cluster size using parameters:  $time\_step=10fs, mass=20405.7294,$   $initial\_simulation\_time=10000, relaxation\_time\_start=10* time\_step, then after 500 step, it increases to <math display="block">1e10*time\_step, and cutoff\_radius=10, target\_temp=500,$   $number\_experiments=26,$   $experiment\_simulation\_time=500,$   $added\_energy\_in\_each\_experiment=0.01*no\_atoms,$   $cluster\_sizes=3:10, and atomic\_distance=2.885$ 

## 4.7 heat capacity versus cluster size

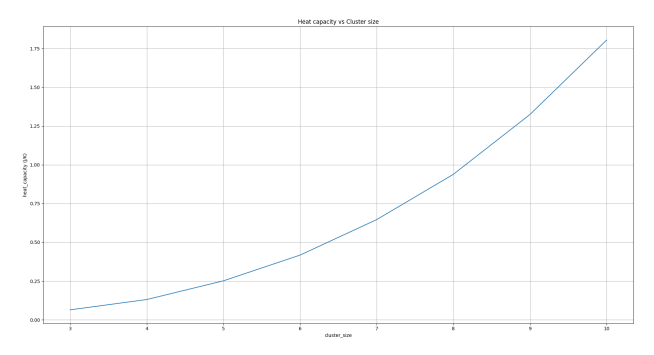
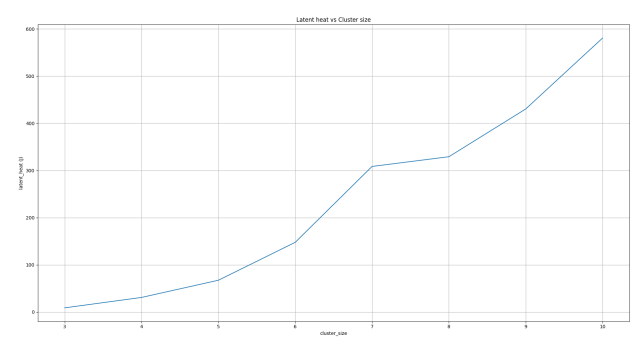


Figure 11: Heat capacity vs cluster size using parameters:  $time\_step=10fs, mass=20405.7294,$   $initial\_simulation\_time=10000, relaxation\_time\_start=10^* time\_step, then after 500 step, it increases to <math display="block">1e10^*time\_step, and cutoff\_radius=10, target\_temp=500,$   $number\_experiments=26,$   $experiment\_simulation\_time=500,$   $added\_energy\_in\_each\_experiment=0.01^*no\_atoms,$   $cluster\_sizes=3:10, and atomic\_distance=2.885$ 

#### 4.8 latent heat versus cluster size



## 4.9 Energy conservation with MPI parallelization

#### 4.10 Nanowire defects

## 5 Conclusion

## Bibliography

[1] H. J. Berendsen, J. v. Postma, W. F. Van Gunsteren, A. DiNola, and J. R. Haak, "Molecular dynamics with coupling to an external bath," *The Journal of chemical physics*, vol. 81, no. 8, pp. 3684–3690, 1984.