TCCM homework 3: A Molecular Dynamics Code

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You can work in groups of up to three people. We expect that you will use Git to work collaboratively on your project. We expect the following files and structure in your submission:

- A LICENSE file specifying the license for your code.
- An AUTHORS file listing the names of all contributors.
- A README.md file providing a brief description of the directory structure and the project.
- An INSTALL.md file with clear instructions on how to compile and run the program.
- A tests directory containing tests to ensure that the program behaves as expected.
- (Optional) A doc directory for additional documentation, if the project requires more detail than the README.md file can provide.
- A src directory containing all the source files of your program.

Molecular dynamics (MD) simulates the movement of atoms based on their initial positions and velocities. In this tutorial, we will develop a molecular dynamics program to illustrate key concepts in MD simulations. Our program will read force field parameters and the initial positions of atoms from an input file. After each small displacement of atoms according to their velocities, the updated coordinates will be saved to an output file, enabling the creation of a video animation with an external tool.

Throughout this tutorial, we will use Argon atoms (mass = 39.948 g/mol) with the following parameters:

• ϵ : 0.0661 j/mol

• $\sigma : 0.3345 \text{ nm}$

Atom coordinates will be provided in nanometers.

You will need to allocate two-dimensional arrays. The following functions should help you allocate and free 2-dimensional arrays:

```
double** malloc_2d(size_t m, size_t n) {
  double** a = malloc(m*sizeof(double*));
  if (a == NULL) {
    return NULL;
  }
  a[0] = malloc(n*m*sizeof(double));
  if (a[0] == NULL) {
    free(a);
    return NULL;
  }
  for (size_t i=1 ; i<m ; i++) {</pre>
    a[i] = a[i-1]+n;
  }
  return a;
}
void free_2d(double** a) {
  free(a[0]);
  a[0] = NULL;
  free(a);
}
```

1 Describing the Atoms

We will create functions to read the atomic data from an input file. The input file follows this format:

- The first line contains the number of atoms (Natoms).
- Each subsequent line contains the x, y, and z coordinates followed by the mass of an atom.

Create a function with the following prototype which reads the number of atoms from an opened input file:

```
size_t read_Natoms(FILE* input_file);
```

Then, write a function with the following prototype:

This function reads the atomic coordinates and masses, and stores the data in the arrays coord and mass provided as parameters. coord is a two dimensional array allocated such that coord[i][2] returns the z coordinate of atom i.

Write another function which takes as input the array of coordinates and returns internuclear distances between each pair in a two-dimensional double array of size (Natoms \times Natoms):

To use the sqrt function, you need to include <math.h>, and you also need to compile using the -lm option to link with the libm.so math library. You might need to put the -lm option at the end of the command line (after your files).

2 The Lennard-Jones potential

Write a function which takes as input ϵ , σ , the number of atoms and the array of distances, and computes the total potential energy

$$V = \sum_{i=1}^{ exttt{Natoms}} \sum_{j>i}^{ exttt{Natoms}} V_{ exttt{LJ}}(r_{ij})$$

where $V_{\rm LJ}(r)$ is the Lennard-Jones potential :

$$V_{\mathrm{LJ}}(r) = 4\epsilon \left[\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^{6} \right].$$

To compute x^n you will need to call the power function located in the libm.so library, defined in math.h.

```
#include <math.h>
x_power_n = pow(x,n);
```

We will now write a function to calculate the total potential energy of the system using the Lennard-Jones potential. This function will take the following inputs: ϵ , σ , the number of atoms (Natoms), and the array containing the distances between each pair of atoms.

The total potential energy V is given by:

$$V = \sum_{i=1}^{\text{Natoms}} \sum_{j>i}^{\text{Natoms}} V_{\text{LJ}}(r_{ij}) \tag{1}$$

where $V_{\rm LJ}(r)$ is the Lennard-Jones potential:

$$V_{\rm LJ}(r) = 4\epsilon \left[\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^{6} \right]. \tag{2}$$

3 Computing the total energy

We will now write functions to compute the total kinetic energy and the total energy of the system.

First, we will write a function to calculate the total kinetic energy of the system T. This function will take as input the number of atoms, the array of velocities $(3 \times \texttt{Natoms})$ and masses (Natoms). The velocities are initialized to zero. The total kinetic energy T is given by:

$$T = \frac{1}{2} \sum_{i=1}^{\text{Natoms}} m_i \mathbf{v}_i^2 \tag{3}$$

Next, we will write a function to compute the total energy of the system, which is the sum of the total kinetic energy T and the total potential energy V. The total energy E is given by:

$$E = T + V. (4)$$

4 Computing the acceleration

The acceleration vector for each atom is given by:

$$\mathbf{a}_i = -\frac{1}{m_i} \nabla_i V$$

Using a finite-difference approximation, the components of the acceleration vector can be written as:

$$\begin{cases} a_{xi} = -\frac{1}{m_i} \frac{V(x_i + \delta) - V(x_i - \delta)}{2\delta} \\ a_{yi} = -\frac{1}{m_i} \frac{V(y_i + \delta) - V(y_i - \delta)}{2\delta} \\ a_{zi} = -\frac{1}{m_i} \frac{V(z_i + \delta) - V(z_i - \delta)}{2\delta} \end{cases}$$

where δ is a small step size.

We need to write a function that computes the acceleration vector for each atom and stores it in a double precision array. This function will call the Lennard-Jones potential function V to compute the necessary potential energy values for the finite-difference approximation.

5 Implementing the molecular dynamics

The molecular dynamics simulation is based on the Verlet algorithm, which updates the position and velocity of each atom at each time step. The algorithm consists of two main equations:

1. Position update:

$$\mathbf{r}^{(n+1)} = \mathbf{r}^{(n)} + \mathbf{v}^{(n)} \Delta t + \mathbf{a}^{(n)} \frac{(\Delta t)^2}{2}$$
 (5)

2. Velocity update:

$$\mathbf{v}^{(n+1)} = \mathbf{v}^{(n)} + \frac{1}{2} \left(\mathbf{a}^{(n)} + \mathbf{a}^{(n+1)} \right) \Delta t \tag{6}$$

where

- (n) denotes the index of the current step.
- **r** is the position vector.
- **v** is the velocity vector.
- a is the acceleration vector.
- Δt is the time step.

We will write a subroutine that implements the Verlet algorithm. The process involves several steps for each iteration, starting with initial velocities set to zero and initial coordinates read from a file.

5.1 Steps for Each Iteration:

Initialize the calculation with initial coordinates and compute the acceleration vector. At each step (n):

- 1. Compute the coordinates $\mathbf{r}^{(n+1)}$ using equation (5)
- 2. Update the part of $\mathbf{v}^{(n+1)}$ which depends on the current acceleration vector $\mathbf{a}^{(n)}$ (in equation (6))
- 3. Update the acceleration $\mathbf{a}^{(n+1)}$ using the finite-difference approximation for the acceleration (equation (4))
- 4. Finalize the update of the $\mathbf{v}^{(n+1)}$ using the $\mathbf{a}^{(n+1)}$ (in equation (6))

We will consider the following parameters for this simulation:

- $\Delta t = 0.2$
- $\delta = 10^{-4}$
- Total number of steps is 1000.

Write into a file the coordinates of the trajectory in xyz format. This corresponds to writing every M steps:

- A line containing the number of atoms.
- A line containing a comment. This line can contain the kinetic, potential and total energy for plotting, and checking that the total energy is conserved.

ullet For each atom, a line containing the atomic Symbol (Ar for Argon atoms), followed by its $x,\,y$ and z coordinates

The xyz can be opened with software like Molden or Jmol to play the animation of the dynamics.