

Exercise Session 4 MDMC Spring 2024

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Reports Reminder

- We encourage you to work together, but the content in your report should be original
- If you use an image or wording from an external source please cite it correctly



Exercise 4 Structure

- Learning goals
 - Derive a time evolution integrator (e.g. Verlet)
 - Understand importance of periodic boundary conditions
 - Run a molecular dynamics simulation for a small molecule (CO₂)
- Chapter in script
 - Chapter 4 Molecular Dynamics Simulations
- Resources
 - Understanding Molecular Simulation, Frenkel & Smit, 2nd Edition - Chapter 3 & Chapter 5 (extra)
 - Computer Simulation of Liquids, Allen & Tildesley, 2nd Edition -Chapter 4



Exercise 4 - Intro & Tips

Today we will provide you a simple Molecular Dynamics (**Toy MD**) code in python and you will extend it to run a MD simulation.

- The theoretical part introduces you to MD Be sure to understand what we mean by:
 - ergodicity
 - phase space sampling
 - MD propagation algorithm
 - periodic boundary conditions
- In the practical part you will implement:
 - Velocity verlet algorithm
 - Periodic boundary conditions (PBC)

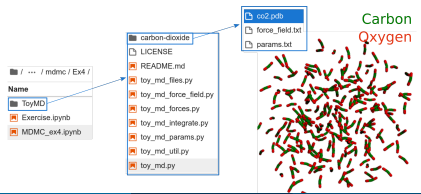
N.B.: Question 8 is no longer a bonus question in case you worked on Ex4 earlier this semester



Exercise 4 - ToyMD structure

ToyMD code structure:

- main code (propagation of MD steps) in toy_md.py script
- additional code for specific tasks, i.e. toy_forces.py
- parameters for the system in separate files (carbon-dioxide folder)



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Exercise 4 - Run ToyMD

ToyMD is a python script, which can be run

- 1. via terminal
- 2. via jupyter notebook

(see instructions in the exercise). In both cases, you will execute a bash command, passing files as arguments to the toy_md.py script with the following structure (paths may change):

```
python3 toy_md.py -c co2.pdb -p params.txt -ff
force_field.txt -o traj.pdb -w co2-output.pdb
```

If you'd like to re-run the code with different parameters, please delete the previous traj.pdb file or rename it