

# Exercise Session 3 MDMC Spring 2024

Salomé Guilbert, Qihao Zhang, Thibault Kläy, Evan Vasey, Sophia Johnson, Andrea Levy

March 25, 2025



### Exercise Check-In

How did you feel during the process of completing, turning in, interviewing, and receiving the comments for Ex 1?

Course feedback: Theoretical background assumed; Theoretical concepts missing?

Remember: Exercises contribute to 1/2 of final grade! We count the best 5 out of the 6 reports for your exercise grade.



### Notebooks Reminder

- Always access the notebooks via the rocket button on the top right of the code files and choose JupyterHub to launch noto.epfl.ch
- Make sure to access noto this way each time you begin the exercise to ensure you have the latest version!



Launch on JupyterHub JupyterHub



### Exercise Structure



Understand importance sampling

Learn importance of detailed balance

Apply the Metropolis Monte Carlo algorithm to calculate properties of a model gas

#### Chapter in script

Chapter 3 - Monte Carlo 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 3 - Intro & Tips

Today we'll be writing and executing Monte Carlo code.

## Tips!

- The theoretical part is about basic Monte Carlo simulations.
   Be sure to know what we mean by:
  - random sampling vs importance sampling
  - configurational space
  - transition or Markov matrix
  - detailed balance
  - Metropolis algorithm
- In the practical part we will run MC code for two systems:

Exercise Session 3

- A photon gas in which the energy states are quantized meaning we can calculate the ensemble average of state occupancy analytically
- A gas in which we test different ensembles (NVT vs NPT) and use the Lennard-Jones potential to describe pairwise interactions



## Exercise 3 - Intro & Tips

- Photon Gas
  - You'll need to write a loop of code to define the estimatedOccupancy function (read the hints and ask questions)
  - NB: randint(0,1) function will generate either 0 or 1. However, in our loop we need either 1 or -1
  - Recall that beta is the inverse of the product of the Boltzmann constant and simulation temperature. Varying beta is a way of varying simulation T



## Exercise 3 - Intro & Tips

#### LJ Potential

- Lots of helper functions to import and functions to define. You
  don't need to modify anything there but it is good to have a
  look to understand what those functions do!
- While there is quite a bit of code to execute, the goal is to see how we incorporate the system ensemble and its interactions when generating sample configurations for MC moves