

# MD Simulations with OpenMM



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@giorginolab

*Thesis projects available*

<https://github.com/giorginolab/MD-Tutorial-Data>

University of Padova c/o Prof. Fuxreiter

May 16, 2023

# This class

- Molecular dynamics is a powerful tool for studying molecular systems
- OpenMM is a software library that allows for efficient and customizable MD simulations
- It's exemplary of a modern well-maintained open-source library:
  - CI infrastructure, developed on GitHub
  - C++ w/ Python bindings
- We'll use the latter, testing *live* on Google Colab.

# **Molecular Dynamics**

# What is MD?

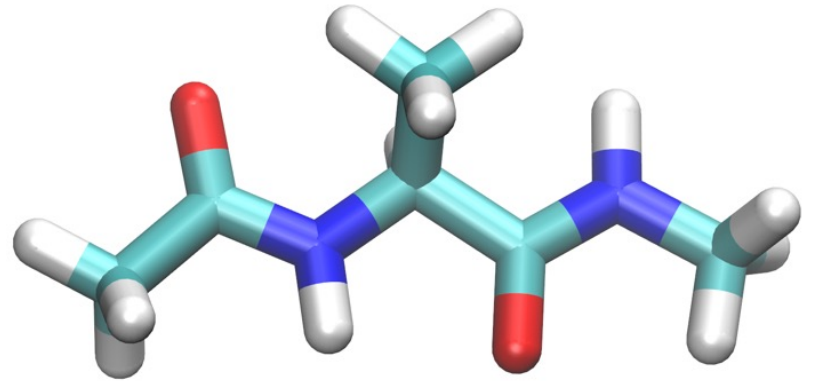
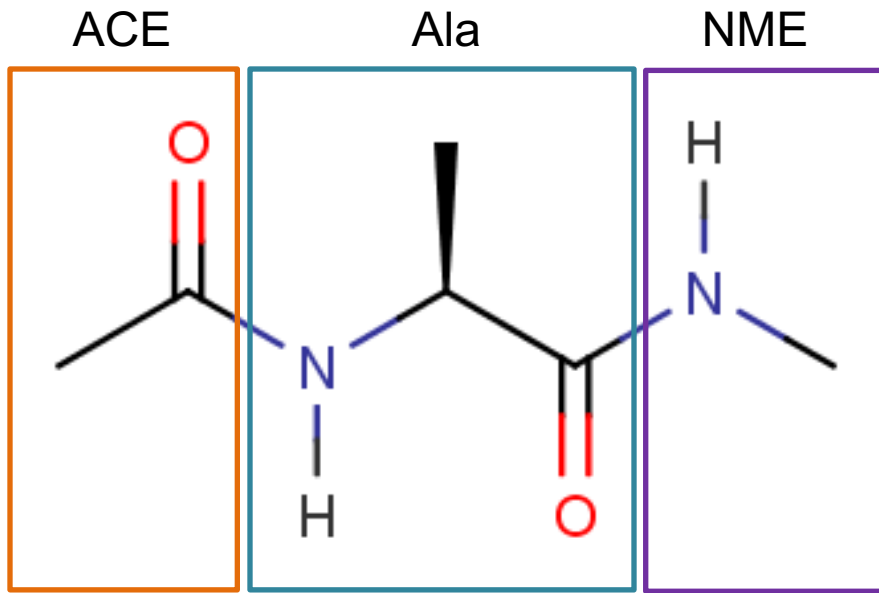
- Attempt the most detailed description of a system which is
  1. atomistic
  2. classical
- Model the internal *forces*...
- ...in order to *integrate* the motion
- Hope in convergent *sampling*

$$\vec{F}_i(\mathbf{x}) = m_i \ddot{\mathbf{x}}_i$$

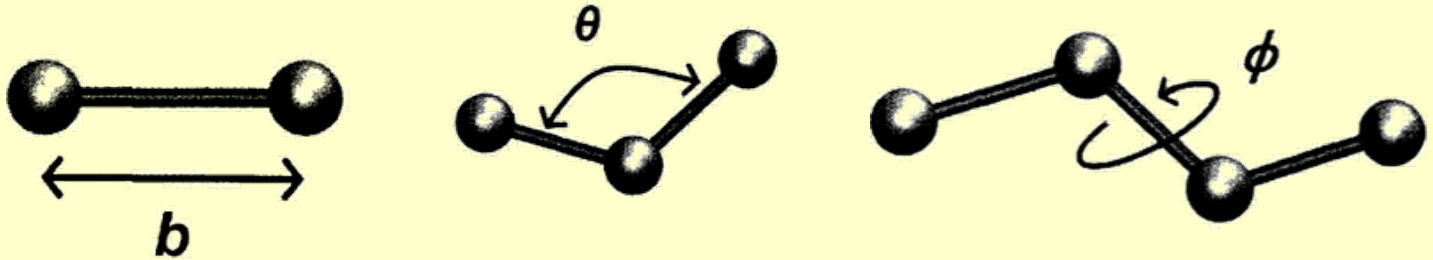
# Assumptions

- In this tutorial we shall deal with **unbiased** sampling approaches with **explicit** solvent, i.e.
  - no added forces except the "physical" ones in your system;
  - all of the system (including water molecules) have atomic resolution.
- Also, current classical MD does not address, by design, the following:
  - Chemical reactions, e.g. catalysis, phosphorylation, ubiquitination etc.
  - Protonation changes
- Finally, small molecules pose distinct challenges and need a separate, expensive **parameterization** step.

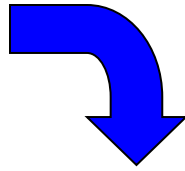
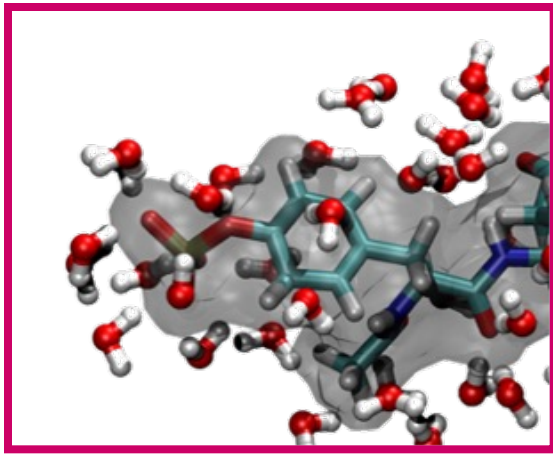
# Alanine “dipeptide”



Bonded  
energy terms  
+ Electrostatics  
+ VdW

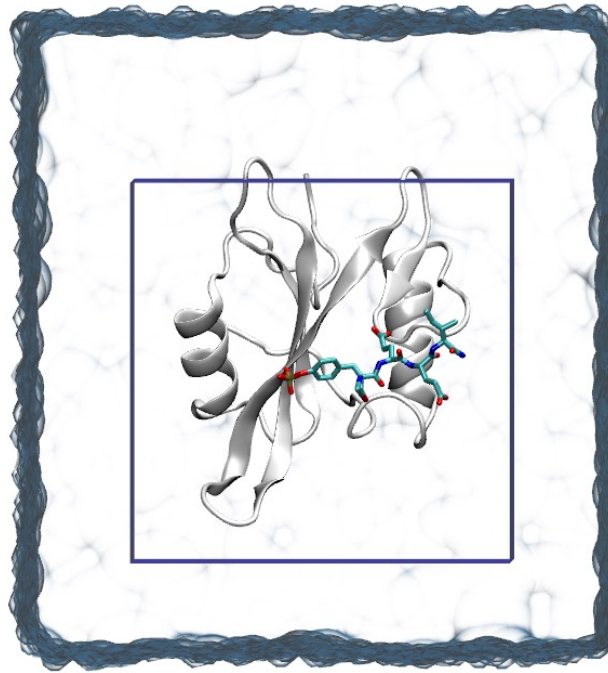


The forcefield is a database of interatomic parameters

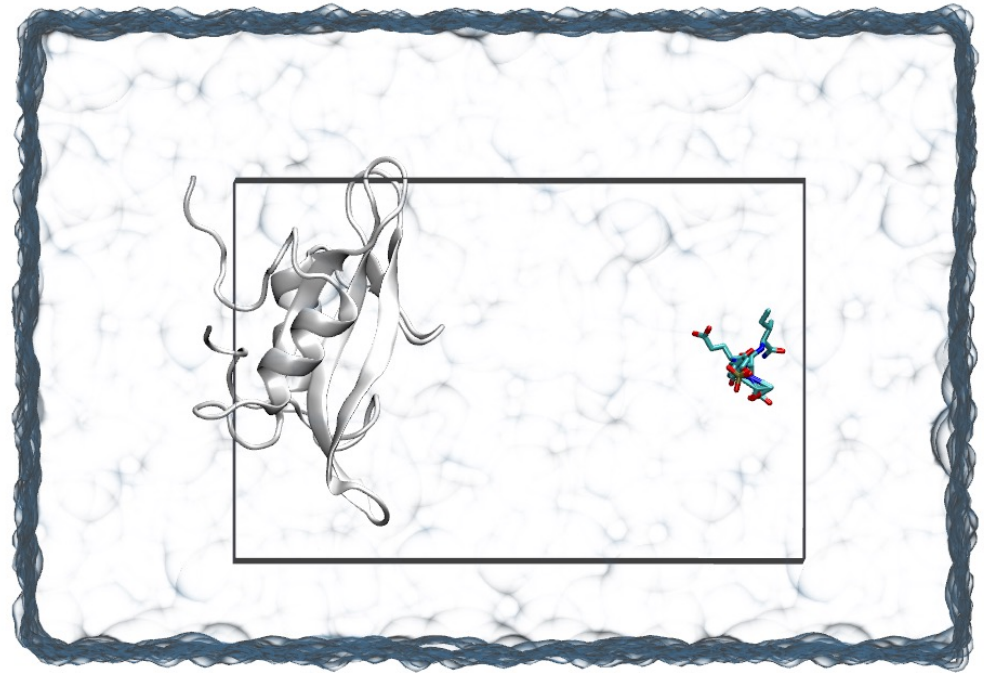


- **Explicit solvation**
- $\rightarrow O(10^5)$  atoms
- **Unbiased dynamics**
- Update every  $10^{-15}$  s (1 fs)

7 nm



7 nm



10 nm

# MD is entirely about timescales

- Your ability to obtain quantitative results is severely limited by the sampling ability you have. You will only be able to reach phenomena occurring on the sampled timescales, or shorter.
  - Sidechain rearrangements, diffusion-limited processes: usually possible \*
  - Local flexibility: usually possible \*
  - Membrane environments: ok-ish
  - Binding: hard but not impossible
  - Folding: very hard but not impossible
    - [\*] Unless there are significant barriers.



# Patience and limits

- The following factors affect the running speed (usually expressed in ns per simulation day, ns/day)
  - System size. Reasonable is 100 AA ~ 30,000 atoms.
  - Computer speed. Forget laptops.
  - Definitely use GPUs.
  - Software.

**OpenMM**



# OpenMM

High performance, customizable molecular simulation.

.org

- OpenMM is a molecular dynamics simulation toolkit that allows for high-performance simulations of biomolecules.
- Allows for simulation of a variety of molecular systems, including proteins, nucleic acids, and small molecules
- OpenMM supports a wide range of force fields and integrators and can run on CPUs and GPUs.
- Open source, written in C++ with Python and other language bindings available

# Basic Workflow (object-oriented)

1. Download, complete and edit the structure:
  - **Topology** (i.e. the identity of atoms, bonds, etc)
  - **Positions** (i.e. the starting coordinates)
2. Create the **system** object.
3. Create the **integrator** object.
4. Create and add custom **forces** to system if needed.
5. Define the **simulation** object.
6. Set the initial positions and velocities.
7. Minimize.
8. Run the simulation.
9. (Analyze the results.)

# Integrators

- ...are algorithms that solve the equations of motion for a system
- OpenMM includes several integrators, e.g. Langevin dynamics, Verlet integrator, and Monte Carlo barostat
- Different integrators are appropriate for different types of simulations and conditions (e.g.: NPT vs NVT)

# Simulating a system

- Once a system has been defined and the force field and integrator selected, it can be simulated
- The simulation (run) involves running a series of steps, where each step involves calculating the forces on each atom, integrating the equations of motion, and updating the system's coordinates
- After the simulation, data analysis can be performed to obtain information about the system's behavior and properties

**Let's pick a test system**

# 6H1F: Gelsolin G2+nanobody

Structure Summary

3D View

Annotations

Experiment

Sequence

Genome

Versions

Biological Assembly 1 ?



**3D View:** [Structure](#) | [1D-3D View](#) | [Electron Density](#) | [Validation Report](#) | [Ligand Interaction](#)

**Global Symmetry:** Asymmetric - C1 ⓘ

**Global Stoichiometry:** Hetero 2-mer - A1B1 ⓘ

[Find Similar Assemblies](#)

Biological assembly 1 assigned by authors and generated by PISA (software)

**Biological Assembly Evidence:** gel filtration

## Macromolecule Content

- Total Structure Weight: 28.49 kDa ⓘ
- Atom Count: 1,896 ⓘ
- Modelled Residue Count: 229 ⓘ
- Deposited Residue Count: 259 ⓘ
- Unique protein chains: 2

## 6H1F

Structure of the nanobody-stabilized gelsolin D187N variant (second domain)

**PDB DOI:** [10.2210/pdb6H1F/pdb](#)

**Classification:** [STRUCTURAL PROTEIN](#)

**Organism(s):** [Lama glama](#), [Homo sapiens](#)

**Expression System:** [Escherichia coli](#)

**Mutation(s):** Yes ⓘ

**Deposited:** 2018-07-11 **Released:** 2019-01-23

**Deposition Author(s):** [Hassan, A.](#), [Milani, M.](#), [Mastrangelo, E.](#), [de Rosa, M.](#)

**Funding Organization(s):** [Amyloidosis Foundation](#)

## Experimental Data Snapshot

**Method:** X-RAY DIFFRACTION

**Resolution:** 1.90 Å

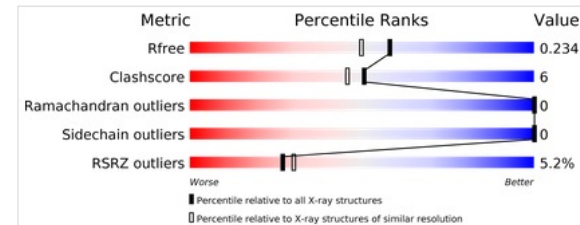
**R-Value Free:** 0.233

**R-Value Work:** 0.199

**R-Value Observed:** 0.202

## wwPDB Validation ⓘ

[3D Report](#) [Full Report](#)



This is version 1.0 of the entry. See complete [history](#).

## Literature

[Download Primary Citation](#)

**Nanobody interaction unveils structure, dynamics and proteotoxicity of the Finnish-type amyloidogenic gelsolin variant.**

[Giorgino, T.](#), [Mattioni, D.](#), [Hassan, A.](#), [Milani, M.](#), [Mastrangelo, E.](#), [Barbiroli, A.](#), [Verhelle, A.](#), [Gettemans, J.](#), [Barzago, M.M.](#), [Diomede, L.](#), [de Rosa, M.](#)

(2019) *Biochim Biophys Acta Mol Basis Dis* **1865**: 648-660

**PubMed:** [30625383](#) [Search on PubMed](#)

**DOI:** [10.1016/j.bbdis.2019.01.010](#)

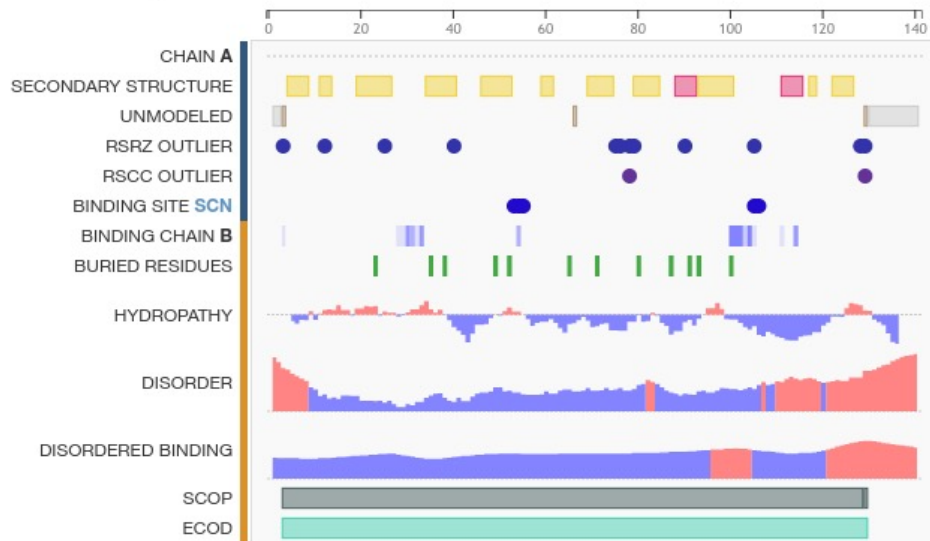
Primary Citation of Related Structures:



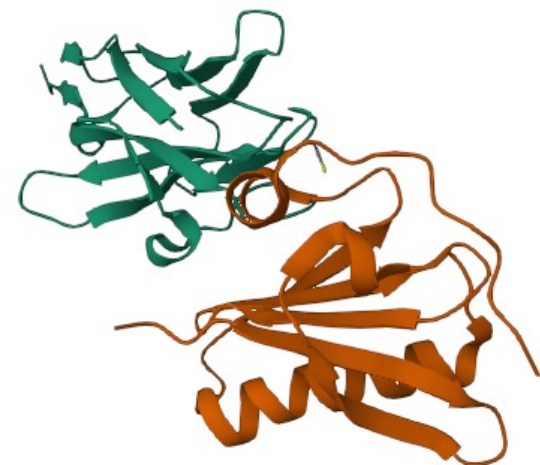
Structure of the nanobody-stabilized gelsolin D187N variant (second domain)

Chain

A
 ▼
 Gelsolin nanobody - Lama glama



Residue



Giorgino T, Mattioni D, Hassan A, Milani M, Mastrangelo E, Barbiroli A, et al.  
Nanobody interaction unveils structure, dynamics and proteotoxicity of the Finnish-type amyloidogenic gelsolin variant. *Biochimica et Biophysica Acta (BBA) - Molecular Basis of Disease*. 2019 Mar 1;1865(3):648–60.

[Journal link.](#)

[Preprint.](#)

BBA - Molecular Basis of Disease 1865 (2019) 648–660



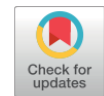
Contents lists available at [ScienceDirect](#)

**BBA - Molecular Basis of Disease**

journal homepage: [www.elsevier.com/locate/bbadis](http://www.elsevier.com/locate/bbadis)



## Nanobody interaction unveils structure, dynamics and proteotoxicity of the Finnish-type amyloidogenic gelsolin variant



Toni Giorgino<sup>a,b</sup>, Davide Mattioni<sup>a,c,1</sup>, Amal Hassan<sup>b,1</sup>, Mario Milani<sup>a,b</sup>, Eloise Mastrangelo<sup>a,b</sup>, Alberto Barbiroli<sup>d</sup>, Adriaan Verhelle<sup>e</sup>, Jan Gettemans<sup>f</sup>, Maria Monica Barzago<sup>c</sup>, Luisa Diomede<sup>c</sup>, Matteo de Rosa<sup>a,b,\*</sup>

<sup>a</sup> Istituto di Biofisica, Consiglio Nazionale delle Ricerche, Milano, Italy

<sup>b</sup> Dipartimento di Bioscienze, Università degli Studi di Milano, Milano, Italy

<sup>c</sup> Department of Molecular Biochemistry and Pharmacology, Istituto di Ricerche Farmacologiche Mario Negri IRCCS, 20156 Milan, Italy

<sup>d</sup> Dipartimento di Scienze per gli Alimenti, la Nutrizione e l'Ambiente, Università degli Studi di Milano, Milano, Italy

<sup>e</sup> Department of Molecular Medicine, Department of Molecular and Cellular Neuroscience, Dorris Neuroscience Center, The Scripps Research Institute, La Jolla, CA 92037, USA

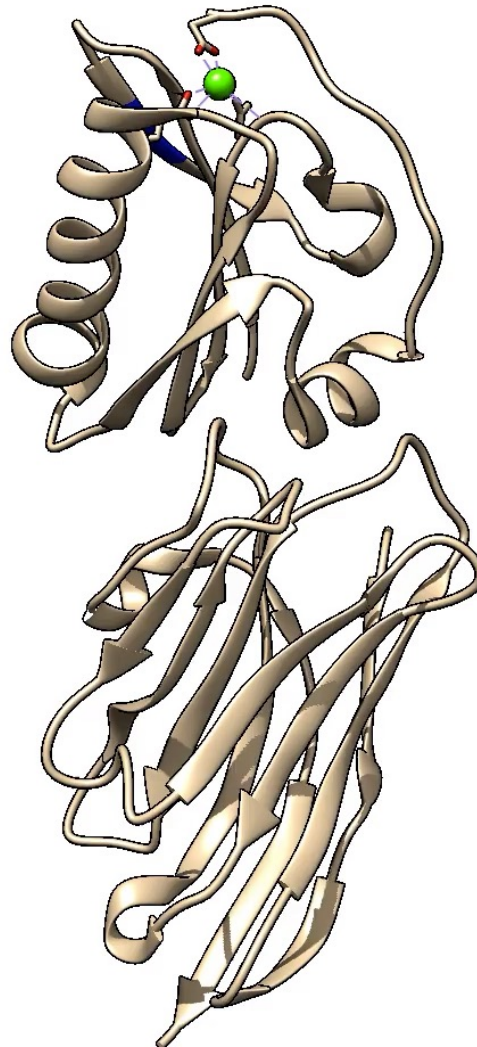
<sup>f</sup> Nanobody Lab, Department of Biochemistry, Faculty of Medicine and Health Sciences, Ghent University, Ghent, Belgium

# Three puzzles!

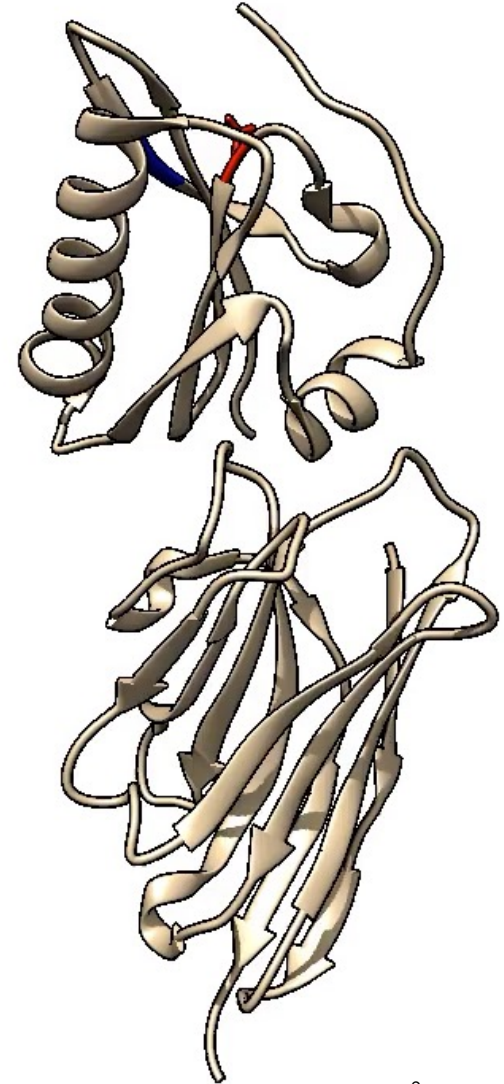
*WT:NbII complex compared to D187N:NbII.*

1. WT and **D187N** are **virtually identical\***: same structure, different function
2. NbII binds far from the furin cleavage site...
3. ...and far from the **Ca<sup>2+</sup>** ion

\* Except Ca<sup>2+</sup> binding



WT: 4S10, 2.6 Å



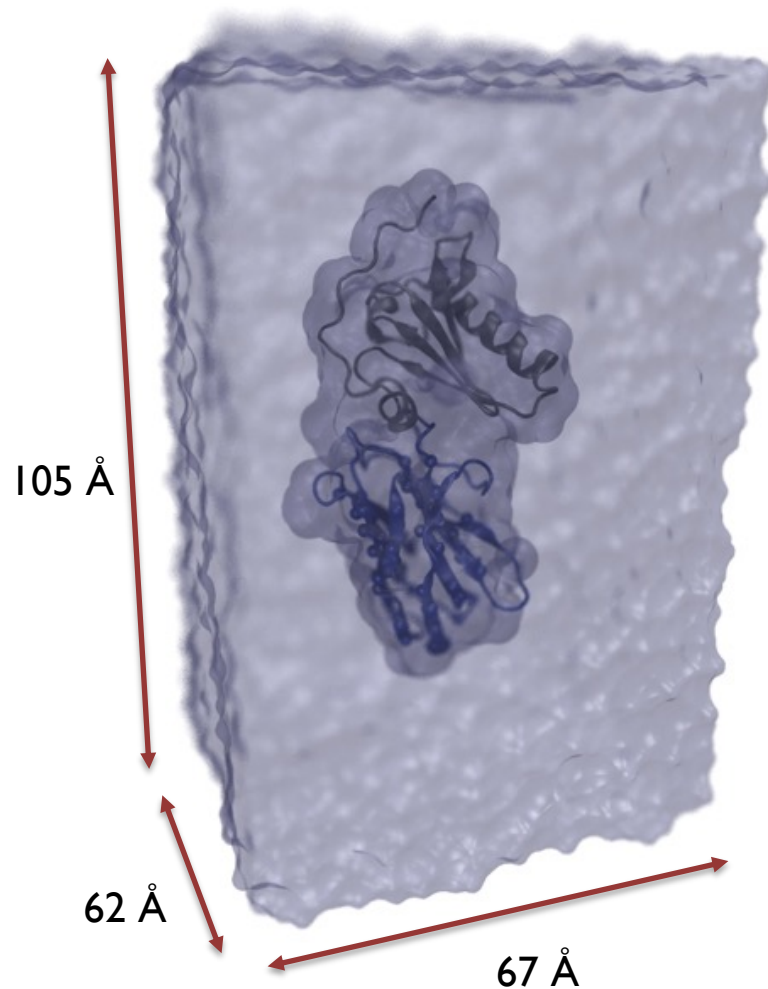
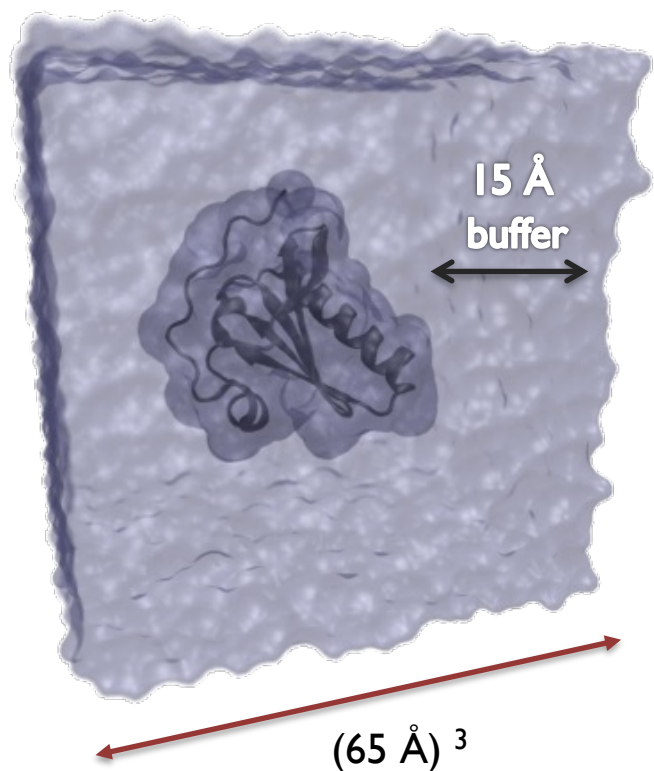
D187N: **6HIF**, 1.9 Å

# GSN ± Nb I MD simulations

- Unbiased sampling @300 °K
- 100 mM NaCl
- Harmonic restraints:  
SS Nb I @ 0.03 kcal/mol/Å<sup>2</sup>

CHARMM36

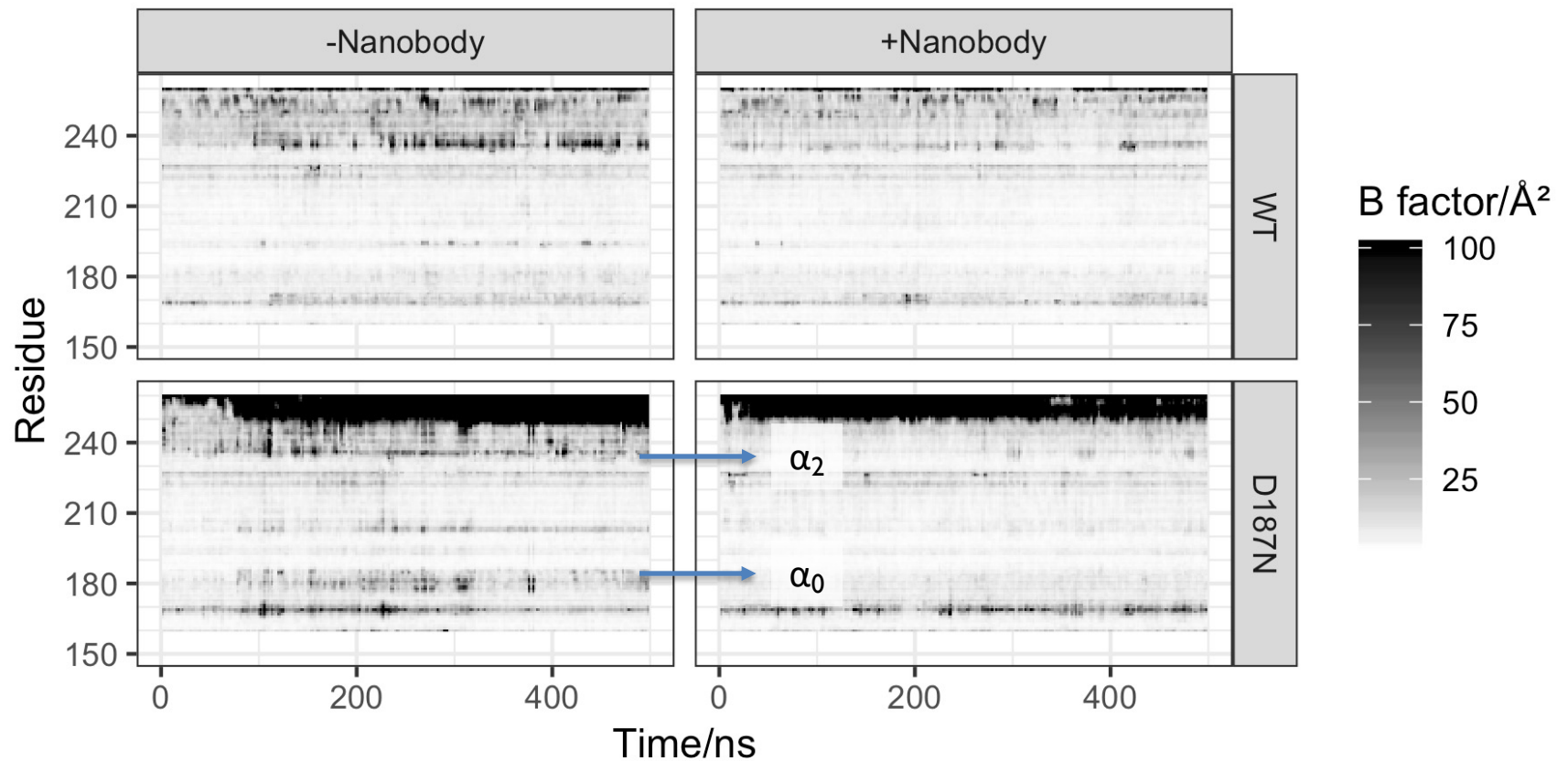
~3 μs tot. ~25k/43k atoms



# MD results



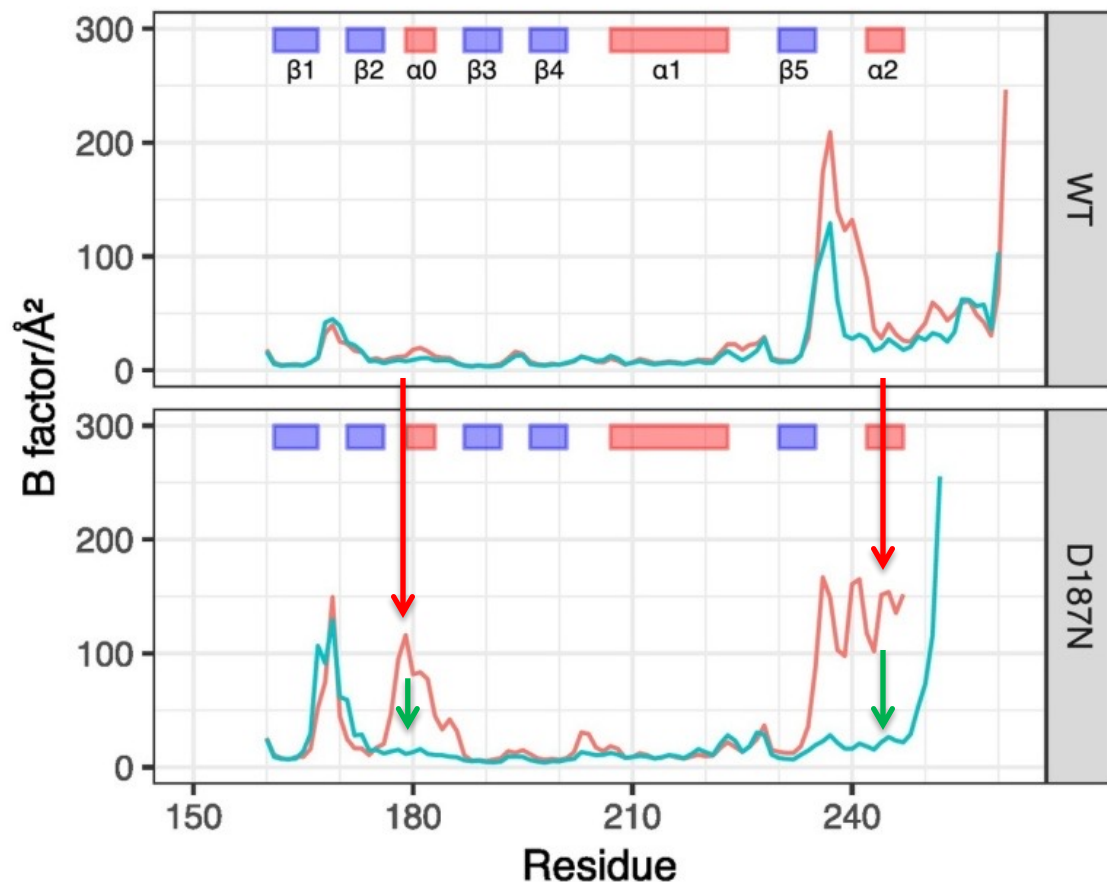
Sample	Nb11	Ca <sup>2+</sup>	Simulated time (ns)	C-terminal disorder onset
WT <sub>G2</sub>	—	+	800	Not observed
WT <sub>G2</sub>	+	+	750	Not observed
D187N <sub>G2</sub>	—	—	748	After 83 ns
D187N <sub>G2</sub>	+	—	512	After 40 ns





# A matter of dynamics?

$$B = (8\pi^2/3) \text{RMSF}^2$$

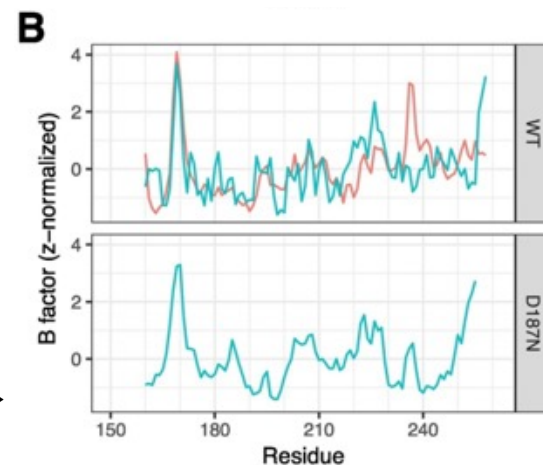


← WT: Nb I I has ~ no effect on fluctuation profile (control)

← Mutant: destabilized  $\alpha_0, \alpha_2$

← Nb I I: complete stability recovery

MD vs exp. B-factors →



**In practice**


# Using OpenMM on Google Colab

- We'll test OpenMM on Google Colab to run molecular dynamics simulations without the need for installing any software on your local machine.
- **Google Colab** is a free Jupyter environment that allows you to run Python code in the cloud. GPUs runtimes are available.
- To use OpenMM on Google Colab or locally, open the provided notebook (read the comments)






<https://github.com/giorginolab/MD-Tutorial-Data>





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
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 **tonigi** Created using Colaboratory

6a4dcdf 7 hours ago 5 commits

 GSN	import	3 weeks ago
 HIVPR	import	3 weeks ago
 notebooks	Created using Colaboratory	7 hours ago
 README.md	Initial commit	3 weeks ago

README.md



# MD-Tutorial-Data

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Data for various MD analysis tutorials



giorginolab / MD-Tutorial-Data Public Edit Pins UI

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**Code**

main + 🔍


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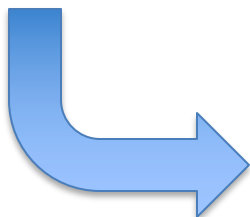
- > GSN
- > HIVPR
- ▼ notebooks

MD-Tutorial-Data / notebooks / 1\_OpenMM\_build.ipynb

tonigi Created using Colaboratory

Preview Code Blame 573 lines (573 loc) · 15.2 KB

 Open in Colab



OpenMM\_2023.ipynb Convidi Settings

File Modifica Visualizza Inserisci Runtime Strumenti Guida

+ Codice + Testo Copia su Drive Connetti

Colab-specific instructions start here

```
[ ] # Here we use a Conda environment inside Google Colab. Blocks specific for Colab
# (like this one) mention "condacolab". On "normal" platforms the procedure
# for installation may be different - you need to check the system's documentation.

# Colab notebooks are "brittle": in the course of time Colab is updated
# and dependencies no longer work properly. Proper HPC platforms are more
# stable (and supported)

# After executing this cell, Colab restarts.

!pip install -q condacolab
import condacolab
condacolab.install_miniforge()

[ ] # Verify Python version
import sys
sys.version

[ ] import condacolab
condacolab.check()

[ ] # Colab-specific workaround for a weird error upon shell escape:
# NotImplementedError: A UTF-8 locale is required. Got ANSI_X3.4-1968
import locale
def getpreferredencoding(do_setlocale = True):
    return "UTF-8"
locale.getpreferredencoding = getpreferredencoding
```

**...when done...**

# Visualize

- After you have done the simulation, load the minimized PDB and output.dcd in PyMOL
- What about PBCs? Fix with: `pbs_unwrap` ...



# Questions

- How many atoms?
- How many residues?
- Disulfide bridges?
- How many trajectory frames?
- Simulation length in *actual* time?

# More questions

- Does density change? Should it?
- What is the box size? Is it appropriate?
- Relaxation time?
- Plot the log file

# Conclusion

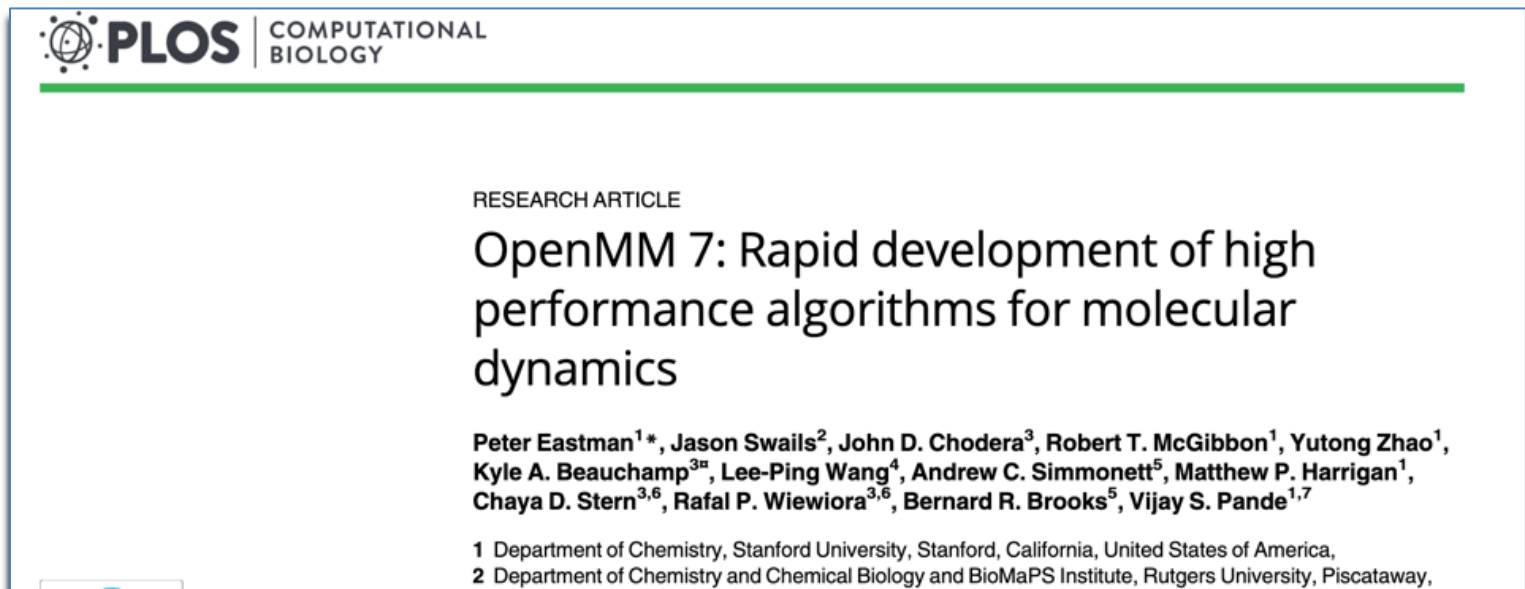
# Conclusion

- OpenMM is a powerful tool for molecular dynamics simulations
- Good, if fragmented, documentation
- With its customizable force fields and integrators, it can be used to study a wide range of atomistic systems, e.g.
  - “toy” polymers
  - all-atom MD with major FFs
  - ANN potentials



# Resources for learning OpenMM

- OpenMM.org website and documentation
- GitHub repository with examples and tutorials
- Community forums and mailing lists for support and discussion
- See also
  - OpenMMtools
  - <https://openforcefield.org/>
  - HTMD, ACEMD
  - <https://github.com/openmm/pdbfixer>
  - Charmm-GUI



**End**