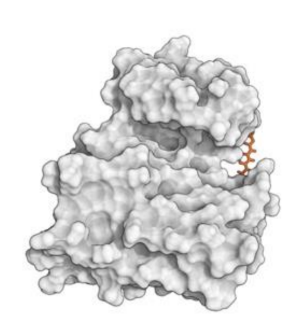
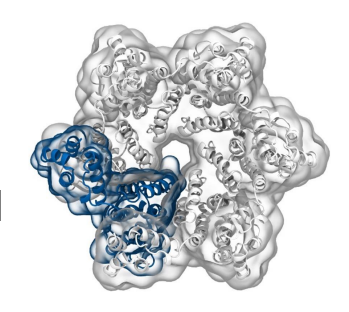
Simulation of Biomolecules



Basic Simulation Analysis

2024 CCP5 Summer School



Dr Matteo Degiacomi

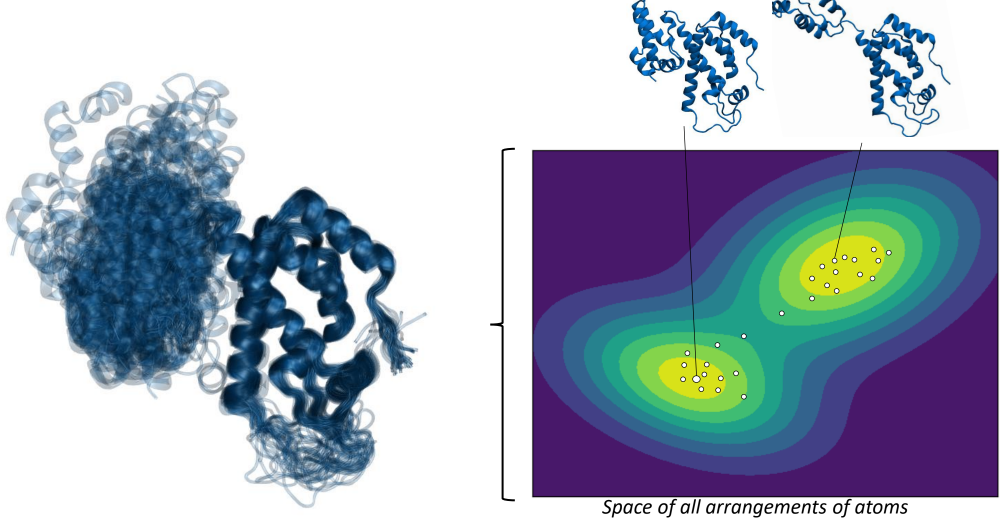
Durham University

matteo.t.degiacomi@durham.ac.uk

Dr Antonia Mey
University of Edinburgh

antonia.mey@ed.ac.uk

Sampling the conformational space



Probability of sampling a conformation is inversely proportional to its energy: $p_i \propto e^{-ct/kT}$

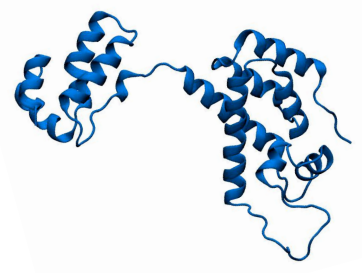
Root Mean Square Deviation (RMSD)

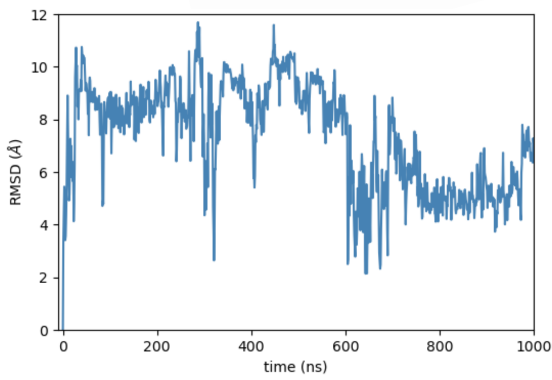
Given a system with N atoms, and a reference arrangement x_0 :

$$RMSD = \sqrt{\frac{1}{N} \sum_{i=0}^{N} (X_i - x_0)^2}$$

In MD, x_0 is often the first conformation in the simulation.

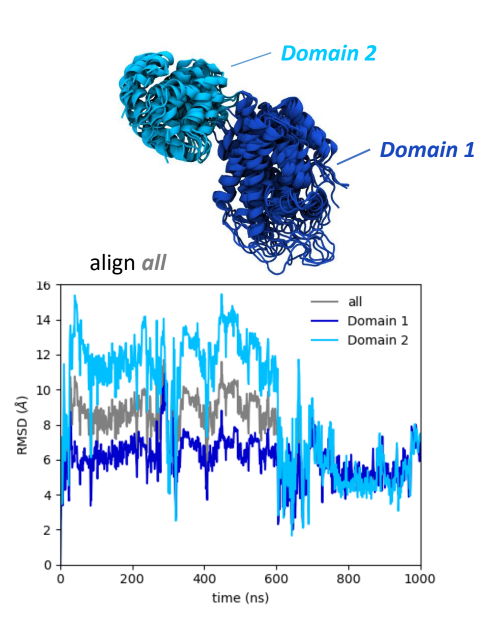
Refrain from using RMSD as a single indicator of simulation convergence.

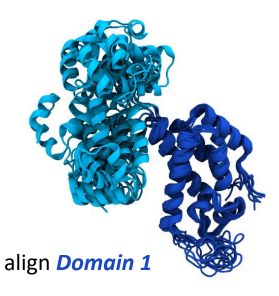


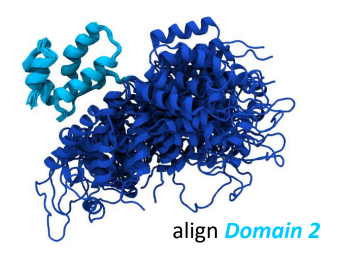


HIV-1 capsomer simulation from: Degiacomi, Structure, 2018

RMSD is alignment- and selection-dependent



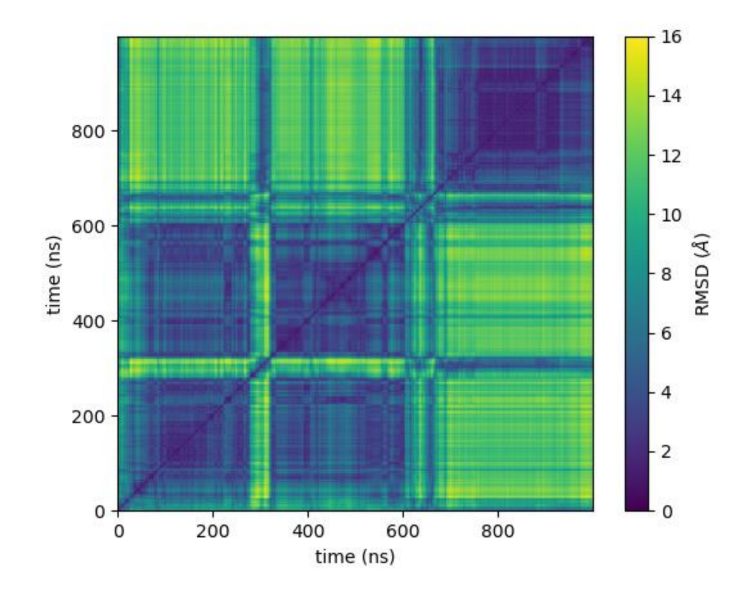




Pairwise RMSD

 Two structures with same RMSD from a reference are not forcefully similar to each other.

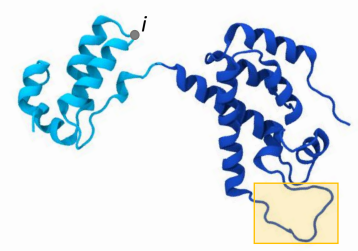
 Pairwise RMSD helps seeing if protein re-visits conformations throughout the simulation.



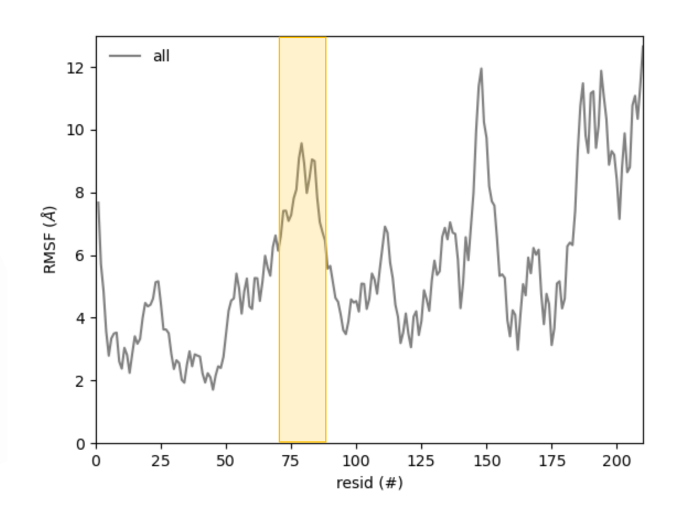
Root Mean Square Fluctuation (RMSF)

The RMSF σ_i of atom i calculates how much it fluctuates around its mean position $\langle X_i \rangle$.

$$\sigma_i = \sqrt{\langle (X_i - \langle X_i \rangle)^2 \rangle}$$



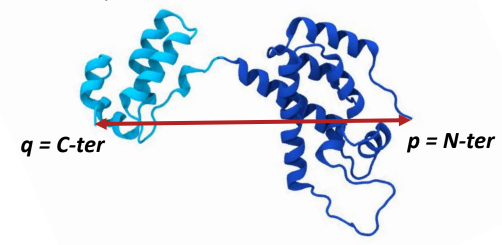
Typically done on C_{α} atoms. Result depends on alignment!



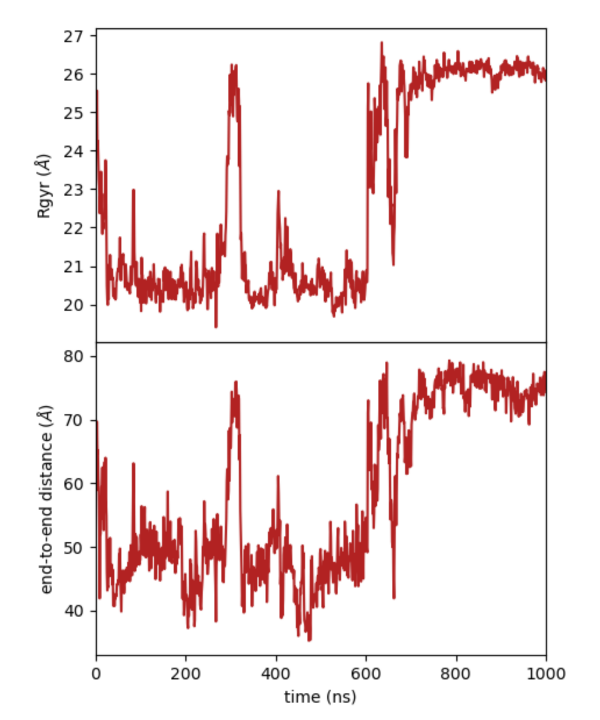
end-to-end distance and Radius of Gyration (Rg)

$$d(p,q) = ||p - q||$$

$$R_g = \sqrt{\frac{1}{N} |r_k - r_{mean}|^2}$$



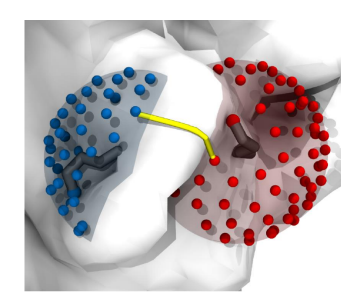
Internal properties do not depend on alignment



Some quantities are not directly measurable

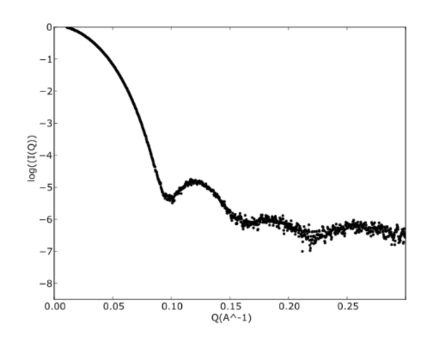
Submit MD conformers to external software simulating experimental data, e.g.:

Chemical cross-linking



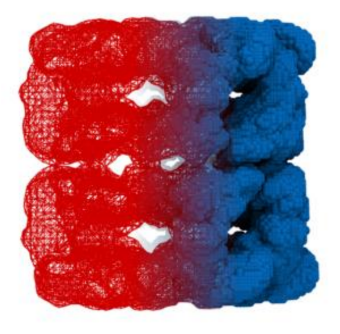
DynamXL, Xwalk, ...

SAXS



CRYSOL, FoXS, ...

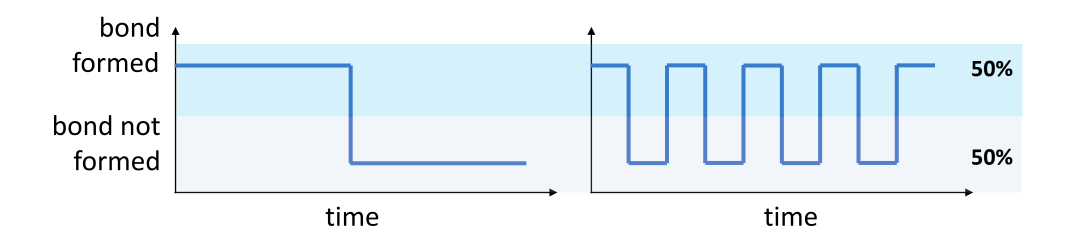
Collision cross-section



IMPACT, MobCal, CollisionCode, ...

Warning: time averaging may hide processes

Thought experiment: typically hydrogen bond is considered established if donor-acceptor distance <2.5 Å, and donor-acceptor-hydrogen angle <20°.



Reporting % time a bond is established in simulation can be misleading!

Large ecosystem of software and packages for data analysis



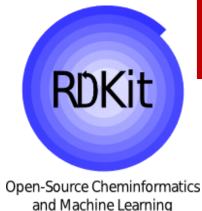


















And many more...

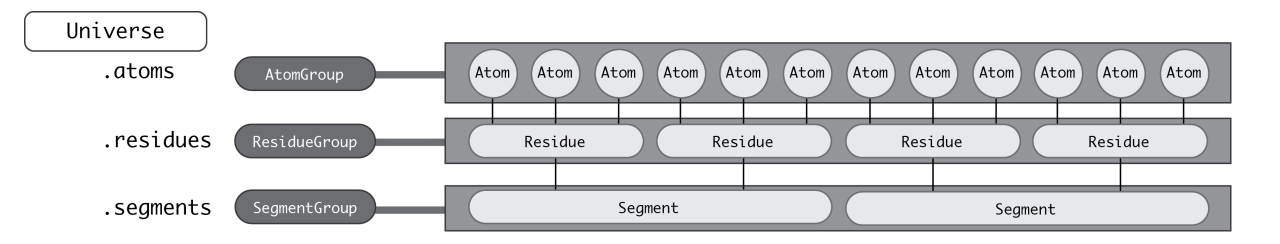


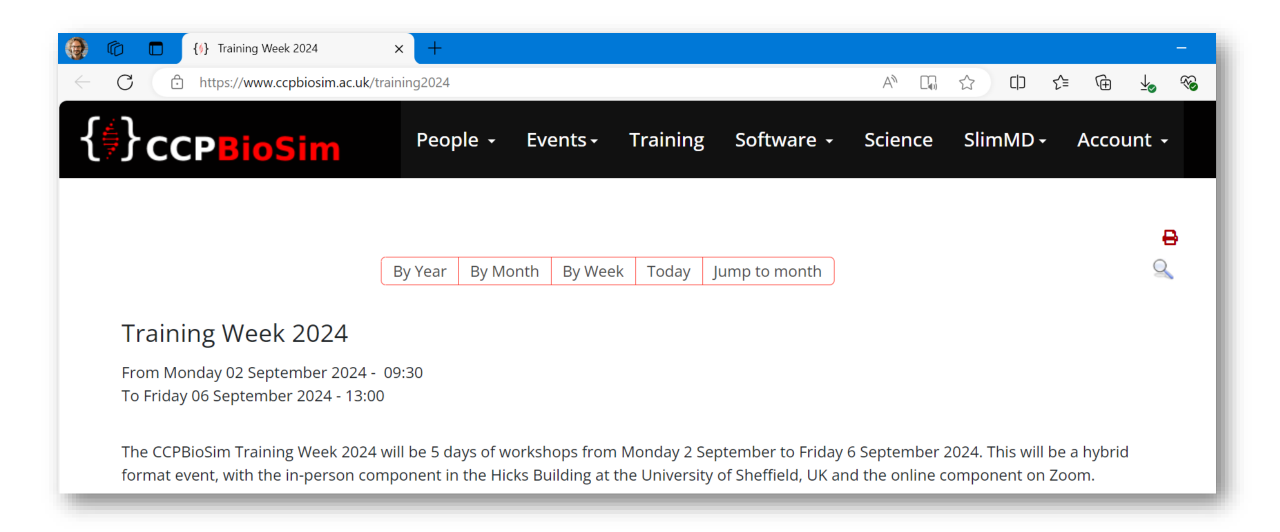


The Universe contains everything about a MD system

- Static information: atoms and their connectivities
- Dynamic information: The trajectory

Data accessible via a hierarchy of containers





Thursday 03 September 2024, 9:30am – 5pm: **Intermediate and Advanced MDAnalysis**This workshop will build on basic MDAnalysis skills to learn about building more complex analysis scripts. Please feel free to bring your own systems and problems to get help from the experts.