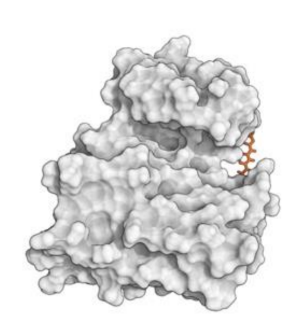
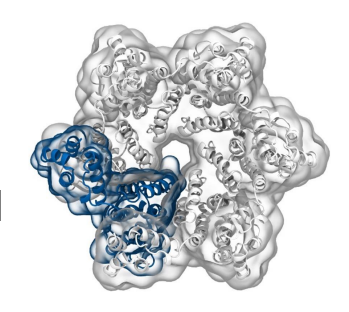
Simulation of Biomolecules



Basic Simulation Analysis

2024 CCP5 Summer School



Dr Matteo Degiacomi

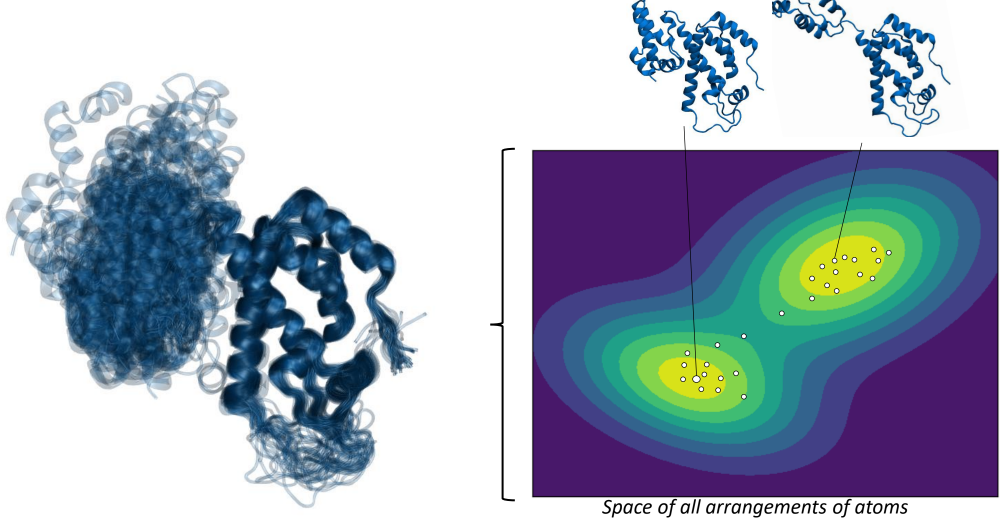
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University of Edinburgh

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Sampling the conformational space



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

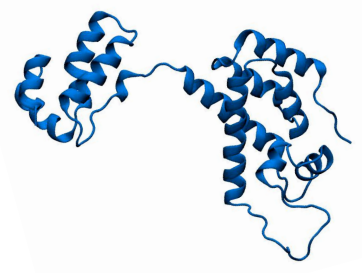
Root Mean Square Deviation (RMSD)

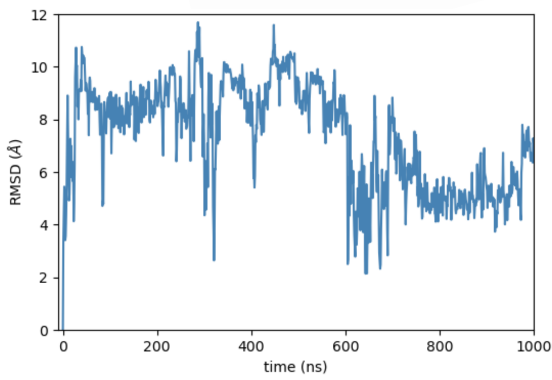
Given a system with N atoms, and a reference arrangement x_0 :

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

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

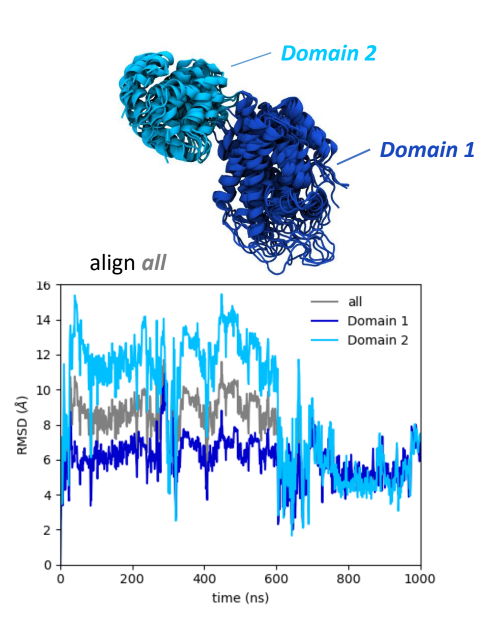
Refrain from using RMSD as a single indicator of simulation convergence.

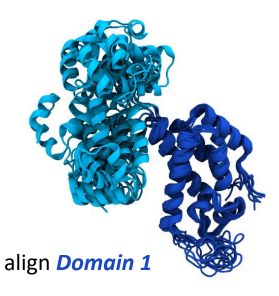


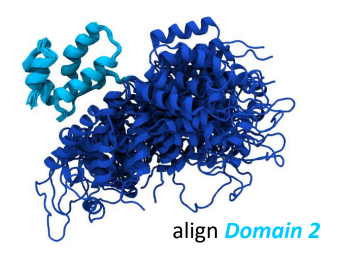


HIV-1 capsomer simulation from: Degiacomi, Structure, 2018

RMSD is alignment- and selection-dependent



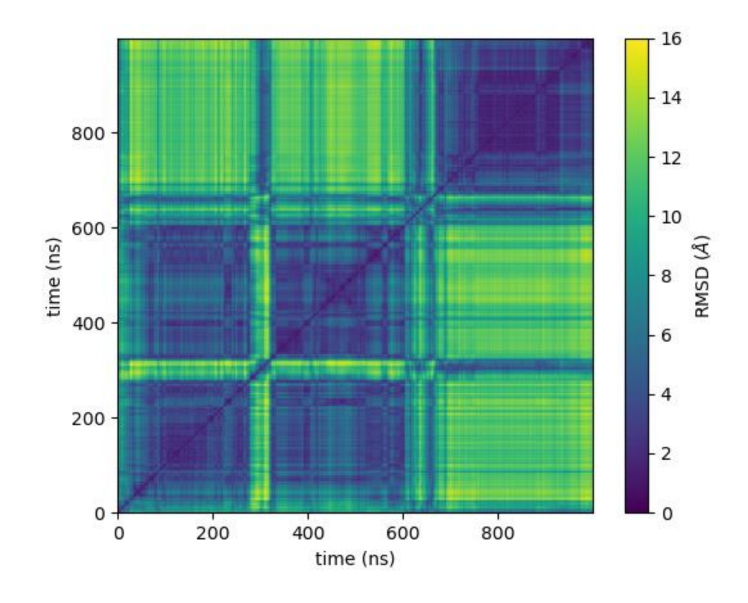




Pairwise RMSD

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

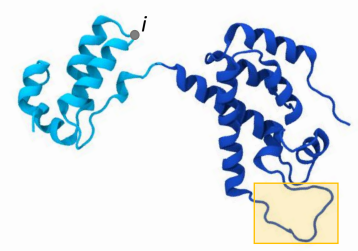
 Pairwise RMSD helps seeing if protein re-visits conformations throughout the simulation.



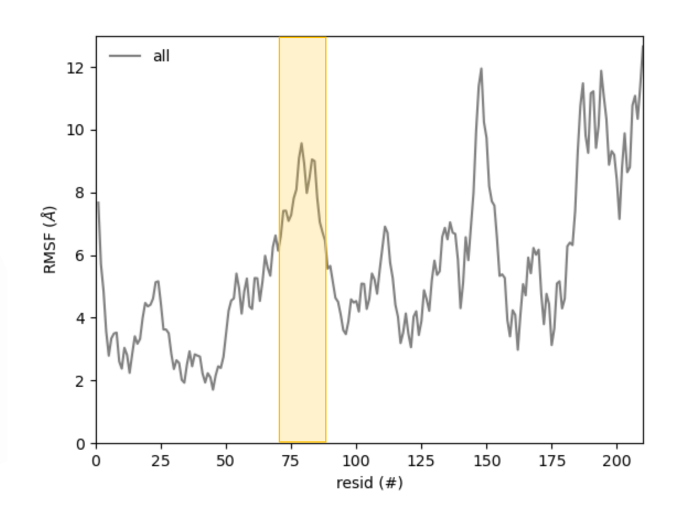
Root Mean Square Fluctuation (RMSF)

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

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



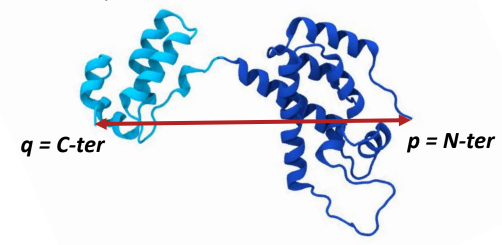
Typically done on C_{α} atoms. Result depends on alignment!



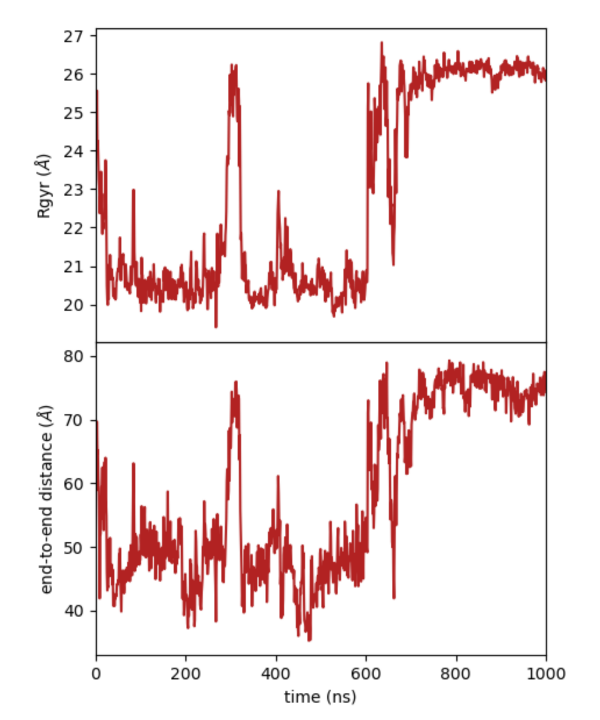
end-to-end distance and Radius of Gyration (Rg)

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

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



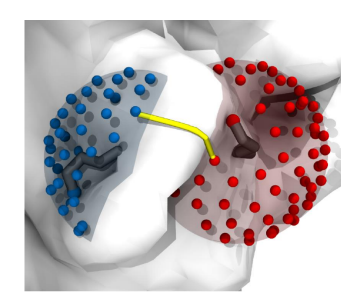
Internal properties do not depend on alignment



Some quantities are not directly measurable

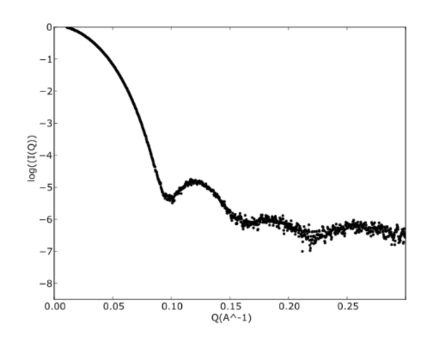
Submit MD conformers to external software simulating experimental data, e.g.:

Chemical cross-linking



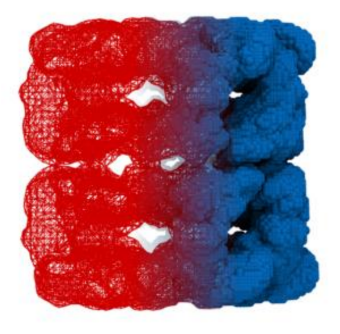
DynamXL, Xwalk, ...

SAXS



CRYSOL, FoXS, ...

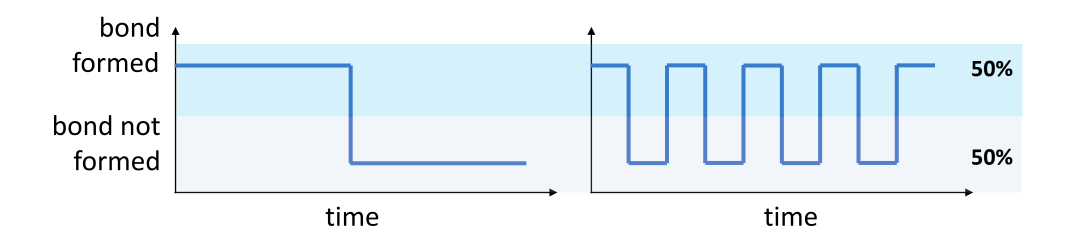
Collision cross-section



IMPACT, MobCal, CollisionCode, ...

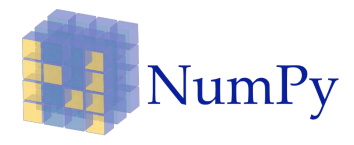
Warning: time averaging may hide processes

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



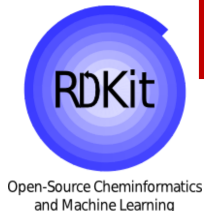


















And many more...





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

