Source:

1 | Preliminary Research

1.1 | Sources

https://www.frontiersin.org/articles/10.3389/fchem.2019.00540/full

1.2 | **Notes**

1.2.1 | Target Processes

- 1. Enzyme catalysis
- 2. Protein-ligand binding
- 3. signal transduction
- 4. allosteric regulation

1.2.2 | Folding Simulation Methods

- 1. all-atom molecular dynamics (MD)
 - · Obtains all desired information regarding the kinetics and thermodynamics
 - (a) Time scale bottleneck
 - very slow (supercomputers -> microseconds of simulation)
 - · require microsecond to milisecond time scales
 - i. optimizations
 - A. conformational sampling?
 - · retains atomistic representation of the system
 - B. overcome kinetic trapping and thourough sampling of conformational space techniques
 - umbrella sampling
 - · multicanonical algorithms
 - · simulated tempering
 - · transition path sampling
 - · targeted molecular dynamics
 - replica exchange method molecular dynamics (REMD)
 - · accelerated molecular dynamics (AMD)
 - · see below
- 2. Accelerated molecular dynamics (AMD)

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1.3 | **Meetings**

1.3.1 **| 12 oct 2020**

- · computational prediction modeling
 - · trying to predict the crystal structure
 - why?
 - · to analyze would this fit?
 - · does it work with this target
- · solving the structure
 - · xray cristolography
 - gold standard
- · look at some concrete examples?

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