

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
 - catalyzing reactions with actions
2. Protein-ligand binding
 - neurotransmitters (dopamine), protein is dopamine receptor how does the ligand bind the proper site to open the channel?
 - ligand: how to pronounce?
3. signal transduction
 - bind to other protein to trigger chain of actions
 - release calcium from intercellular stores
4. allosteric regulation
 - a reason why knowing the structure and pockets is important
 - predict allosteric sites
 - similar to non-competitive inhibition?
 - but for dna binding proteins, like the dna transcription inhibitor
 - ligand binds allosteric site and activates the protein

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 millisecond time scales
- i. optimizations
 - A. conformational sampling?
 - retains atomistic representation of the system
 - B. overcome kinetic trapping and thorough 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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1.2.3 | **Voltage Gated Ion Channels**

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 cristologyraphy
 - gold standard
 - now got the structure
 - what does that mean?
 - can we simulate how it interacts?
 - can you then do modeling on that to see if drug molecules work? are useful
- look at some concrete examples?
- tell a biological story alongside with computational relevance piece

1. protien synthase

not as much simulation stuff

2. neurotransmitters

dopamine sodium rushes in, electrochemical and concentration gradient recharge gradient by releasing potassium

(a) nerst equation

electrochemical gradient as battery

(b) goldman-katz equation

- applied to neuro
- takes into account the concentrations of the 4 ions
 - how does the power of the battery work given those components?
 - ligands and pH can change/denature protiens, but there are also voltage gated channels

3. Voltage Driven Things

- Heartbeart
- nervous system

- how do voltage gated ion channels work?

(a) things to know about

- action potential
- voltage gated calcium channels open at depolarization threshold

i. neurotransmitters

- "calcium mediated exocytosis of neurotransmitter vesicles in the synaptic terminal"
- calcium rushes somewhere to allow the neurotransmitters to leave the cell

4. Case study

- why do we care? why is this useful
- knowing the structure can lead to some useful information
- how did it lead to some sort of accelerated understanding?

5. prions

- how to pronounce?

(a) CJD

- is it inheritable?
- one case per million population

i. Cases

- the gene that causes CJD in 5-10% of cases is PRNP
- 87% of cases are sporadic

(b) isoform

- a different set of introns and exons
- spliceosome takes pre-RNA and cuts out introns
 - even if the pre-RNA had 10 exons, the spliceosome might take a subset of those exons and remove the others
- An isoform is a variant of that subset, an abnormal isoform is one that is "bad" and causes problems