

# Modern Molecular Simulations using Extended-Lagrangian Approaches

Cameron F. Abrams
Bartlett '81 – Barry '81 Professor and Head cfa22@drexel.edu
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Drexel University, Department of Chemical and Biological Engineering

### Outline

- 1. Introduction and Motivation
- 2. Summary

**Introduction and Motivation** 

## Summary

### Summary

- OTFP combines enhanced sampling and free-energy-profile generation to provide deeper understanding of biomolecular mechanisms
- We recapitulate water-mediated "knock-on" transport of multiple K<sup>+</sup> ions in Kv1.2, but also predict reduced barriers for dry transport
- ullet Demonstrated Markovian milestoning along minimum free-energy pathways to estimate entry and exit rates of  $O_2$  in MSOX
- MSOX: Simulations agree with experiment that modified ping-pong mechanism is more likely than ping-pong because substrate influences O<sub>2</sub> entry and exit kinetics
- MSOX: Substrate-induced gatekeeper closing and active-site desolvation likely underly the substrates effect on O<sub>2</sub>.

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- Climbing String: Dr. Gourav SHRIVASTAV ()
- Other projects:
  - Steven GOSSERT (2021): HIV-1 Entry Inhibitor Design
  - Natasha VERGARA (2022): HIV-1 Entry Inhibitor Design
  - Ming HUANG (2023): Molecular modeling of thermosets
  - Dr. Salman ZARRINI: Molecular modeling of thermoset/glass sizing fluid
  - Dr. Mohammadjavad MOHAMMADI: HIV-1 Entry Inhibitor Design

### Collaborators

- Prof. Eric VANDEN-EIJNDEN (NYU; Mathematics)
- Prof. Irwin CHAIKEN (DUCOM; Biochemistry)
- Prof. Amos SMITH III (UPenn; Organic Chemistry)
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- Resources: www.github.com/cameronabrams/cfacv, www.github.com/cameronabrams/pestifer