

Dr. Alex Albaugh

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Education

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|------------------------------------|--------|----------------------|---------------|-----------|
| University of California, Berkeley | Ph.D. | Chemical Engineering | Berkeley, CA | 2012-2018 |
| University of Michigan | B.S.E. | Chemical Engineering | Ann Arbor, MI | 2008-2012 |

Professional Experience

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|----------------------------------|----------------------------|-------------------------|--------------|
| Postdoctoral Fellow | Todd Gingrich Group | Northwestern University | 2018-Present |
| Quantitative Researcher | Property Options Group | Freelance | 2019-Present |
| Quantitative Researcher | Partial <i>f</i> | Freelance | 2018-2019 |
| Graduate Student Researcher | Teresa Head-Gordon Group | UC Berkeley | 2012-2018 |
| Graduate Student Instructor | Chemical Engineering Dept. | UC Berkeley | 2012-2017 |
| Undergraduate Research Assistant | Ronald Larson Group | University of Michigan | 2011-2012 |

Teaching Experience

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|--------------------------------------|-------------|------------|
| Transport Processes | UC Berkeley | 2017 (GSI) |
| Process Dynamics and Control | UC Berkeley | 2015 (GSI) |
| Chemical Kinetics and Reactor Design | UC Berkeley | 2013 (GSI) |
| Introduction to Chemical Engineering | UC Berkeley | 2012 (GSI) |

Selected Coursework

Computer Science: Machine learning, parallel computing

Mathematics: Numerical differential equations

Physical science: Statistical mechanics, quantum mechanics, fluid mechanics, heat transfer, petroleum geology

Engineering: Process design, process control, reactor engineering, separation processes, biomolecular engineering

Computer Proficiencies

Environments: Mac, Linux, Windows

Applications: MATLAB, Aspen, Mathematica, COMSOL, Microsoft Office, Git, LaTeX, TensorFlow

Languages: Fortran, Python, C++, C, Bash, OpenMP/MPI

Honors and Awards

Outstanding Graduate Student Instructor, UC Berkeley, 2017

Summa cum laude, University of Michigan, 2012

Julius F. Bartus Memorial Scholarship, University of Michigan, 2011-2012

James B. Angell Scholar, University of Michigan, 2011-2012

Award of Excellence Scholarship, University of Michigan, 2008-2012

Dean's List and University Honors, University of Michigan, 2008-2012

Publications

1. **A. Albaugh**, T. Gingrich (2020). Simulating a Chemically-Fueled Molecular Motor with Nonequilibrium Molecular Dynamics. (*in review*)
2. **A. Albaugh**, T. Gingrich (2020). Estimating Reciprocal Partition Functions to Enable Design Space Sampling. *Journal of Chemical Physics*, 153, 204102.
3. **A. Albaugh**, M. Tuckerman, T. Head-Gordon (2019). Combining Iteration-Free Polarization with Large Time Step Stochastic-Isokinetic Integration. *Journal of Chemical Theory and Computation*, 15, 2195.
4. **A. Albaugh** (2018). Improved Methods for Polarizable Classical Molecular Dynamics Simulations. *UC Berkeley Dissertation published by ProQuest.*

5. **A. Albaugh**, T. Head-Gordon, A. Niklasson (2018). Higher-order Extended Lagrangian Born-Oppenheimer Molecular Dynamics for Classical Polarizable Models. *Journal of Chemical Theory and Computation*, 14, 499.
6. **A. Albaugh**, T. Head-Gordon (2017). A New Method for Treating Drude Polarization in Classical Molecular Simulation. *Journal of Chemical Theory and Computation*, 13, 5207.
7. **A. Albaugh**, A. Niklasson, T. Head-Gordon (2017). Accurate Classical Polarization Solution with No Self-Consistent Field Iterations. *Journal of Physical Chemistry Letters*, 8, 1714.
8. V. Vitale, J. Dziedzic, **A. Albaugh**, A. Niklasson, T. Head-Gordon, C.-K. Skylaris (2017). Performance of Extended Lagrangian Schemes for Molecular Dynamics Simulations with Classical Polarizable Force Fields and Density Functional Theory. *Journal of Chemical Physics*, 146, 124115.
9. **A. Albaugh**, H. Boateng, R. Bradshaw, O. Demerdash, J. Dziedzic, Y. Mao, D. Margul, J. Swails, Q. Zeng, D. Case, P. Eastman, L.-P. Wang, J. Essex, M. Head-Gordon, V. Pande, J. Ponder, Y. Shao, C.-K. Skylaris, I. Todorov, M. Tuckerman, T. Head-Gordon (2016). Advanced Potential Energy Surfaces for Molecular Simulation. *Journal of Physical Chemistry B*, 120, 9811.
10. **A. Albaugh**, O. Demerdash, T. Head-Gordon (2015). An Efficient and Stable Hybrid Extended Lagrangian/Self-Consistent Field Scheme for Solving Classical Mutual Induction. *Journal of Chemical Physics*, 143, 174104.
11. I.S. Dalal, C.-C. Hsieh, **A. Albaugh**, R.G. Larson (2014). Effects of Volume and Hydrodynamic Interactions on the Behavior of Isolated Bead-Rod Polymer Chains in Shearing Flow. *AIChE Journal*, 60, 1400.
12. I.S. Dalal, **A. Albaugh**, N. Hoda, R.G. Larson (2012). Tumbling and Deformation of Isolated Polymer Chains in Shearing Flow. *Macromolecules*, 45, 9493.

Presentations

1. *Estimating Reciprocal Partition Functions to Enable Design Space Sampling*. Berkeley Statistical Mechanics Meeting in Berkeley, CA January 10, 2020.
2. *Sampling Designs by Estimating Reciprocal Partition Functions*. Northwestern University Department of Chemistry Theory Seminar in Evanston, IL November 25, 2019.
3. *Improved Methods for Polarizable Classical Molecular Dynamics Simulations*. Northwestern University Department of Chemistry Theory Seminar in Evanston, IL December 3, 2018.
4. *Improved Methods for Polarizable Classical Molecular Dynamics Simulations*. Argonne National Laboratory Institute for Molecular Engineering Seminar in Lemont, IL May 8, 2018.
5. *Novel Methods for Classical Polarizable Molecular Dynamics Simulations*. UC Berkeley Chemical Engineering Department Colloquium in Berkeley, CA March 1, 2017.
6. *Novel Methods for Polarizable Classical Molecular Dynamics Simulations* (poster). Berkeley Statistical Mechanics Meeting in Berkeley, CA January 13, 2017.
7. *Efficient Solutions of Classical Polarization Using Hybrid Extended Lagrangian/Self-Consistent Field Methods*. American Chemical Society National Meeting in Philadelphia, PA August 21, 2016.
8. *New Methods for Polarizable Molecular Dynamics Simulations*. Professor Ali Mesbah Research Group Meeting in Berkeley, CA April 15, 2016.
9. *A Hybrid Extended Lagrangian/Self-Consistent Field Scheme for Solving Classical Polarization*. National Institutes of Health Laboratory of Computational Biology Weekly Seminar in Rockville, MD January 7, 2016.
10. *Novel Methods and Models for AMOEBA Polarization*. Telluride Science Research Center Advanced Potential Energy Surfaces Workshop in Telluride, CO June 16, 2015.
11. *Increasing the Efficiency of the AMOEBA Force Field Using Drude Oscillators* (poster). American Chemical Society Annual Meeting in San Francisco, CA August 12, 2014.
12. *A Drude Polarization Model for the AMOEBA Force Field*. Telluride Science Research Center Many-Body Interaction Workshop in Telluride, CO June 18, 2014.
13. *Non-monotonic Stretch of Isolated Semi-Flexible Polymer Chains* (poster). Macromolecular Science and Engineering Symposium in Ann Arbor, MI October 27, 2011.
14. *Non-monotonic Stretch of Isolated Semi-Flexible Polymer Chains* (poster). Society of Rheology Annual Meeting in Cleveland, OH October 12, 2011.