







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


Education/Research

- 2020 –  **Postdoctoral Fellow, Emory University**
Investigating the behavior of the molecule-solvent interface in Fang Liu's group.
- 2014 – 2020  **Ph.D., Physics, Rice University**
Thesis title: *Adaptive sampling of Conformational Dynamics* Advisor: *Cecilia Clementi*

Publications selected

- 1 **Hruska, Eugen**, Ariel Gale, and Fang Liu. "Bridging the Experiment-Calculation Divide: Machine Learning Corrections to Redox Potential Calculations in Implicit and Explicit Solvent Models". In: *J. Chem. Theory Comput.* (2022).
- 2 **Hruska, Eugen** et al. "AutoSolvate: A Toolkit for Automating Quantum Chemistry Design and Discovery of Solvated Molecules". In: *submitted* (2022).
- 3 **Hruska, Eugen** et al. "Extensible and Scalable Adaptive Sampling on Supercomputers". In: *J. Chem. Theory Comput.* (2020).
- 4 **Hruska, Eugen** et al. "Quantitative comparison of adaptive sampling methods for protein dynamics". In: *J. Chem. Phys.* 149.24 (2018), p. 244119.

Talks

- 2021  **Benchmarking the accuracy of free energy landscapes generated by adaptive sampling strategies**, CECAM, Mixed-gen Session 6: Activated Events
-  **Reducing the error of redox potential calculations in implicit and explicit solvents with machine learning**, ACS Fall
- 2020  **Deep learning of molecular dynamics representations**, Emory Machine Learning in Chemistry Journal Club

Awards

2009: **Gold medal, International Physics Olympiad, top 50 in world**, 2011: Gold medal and Best Experiment, World Physics Olympiad, 2007-2008: **Gold medal**, International Junior Science Olympiad, 2012: Student award, German Physical Society, 2009: **Scholarship**, German Academic Scholarship Foundation