# Saurabh Sivakumar

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Interests: Multi-scale molecular simulations, Computational chemistry and materials modelling, Geometric deep learning, Fine-tuning ML models, High-performance computing and Automated workflow development

#### Education

University of California	Davis, CA
Ph.D. in Chemical Engineering	Sep 2021 – Present
Carnegie Mellon University	Pittsburgh, PA
M.S in Chemical Engineering	<b>Aug 2019 – Dec 2020</b>
Thesis: Applications of machine learning interatomic potentials in catalysis	Advisor: Zachary Ulissi
National Institute of Technology	Tiruchirappalli, India
B.Tech in Chemical Engineering, Minor in Economics	Jul 2015 – May 2019
Thesis: Plant design for Trichloroethylene manufacturing	Advisor: S. Saravanan

#### SKILLS

- Languages: Python, Bash, R, C++, LATEX
- Python Packages: PyTorch, ASE, RDKit, scikit-learn, scipy, Pandas, Seaborn, Pymatgen, VASP-kit, MDanalysis, OpenMM, CuPy, Numba, PyG, WandB, Parsl, Deap
- Technologies/Frameworks: Git, Plotly
- Simulation Tools: VASP, ORCA, QuantumEspresso, LAMMPS, PLUMED, NWChem, Psi4, jDFTx, GROMACS, Gaussian, Zacros, PySCF, COMSOL, Cantera, OVITO, VMD

#### PUBLICATIONS

An Automated Pynta-based Curriculum for ML-Accelerated Calculation of Transition States Trevor Price, Saurabh Sivakumar, Matthew S. Johnson, Judit Zador and Ambarish Kulkarni Journal of Physical Chemistry C 2025 [Paper Link] Toward an ab Initio Description of Adsorbate Surface Dynamics Saurabh Sivakumar and Ambarish Kulkarni Journal of Physical Chemistry C 2024 [Paper Link] Cluster-MLP: An Active Learning Genetic Algorithm Framework for Accelerated Discovery of Global Minimum Configurations of Pure and Alloyed Nanoclusters Rajesh K. Raju, Saurabh Sivakumar, Xiaoxiao Wang and Zachary W Ulissi Journal of Chemical Information and Modeling 2023 [Paper Link] Enabling robust offline active learning for machine learning potentials using simple physics-based priors Muhammed Shuaibi, Saurabh Sivakumar, Rui Qi Chen and Zachary W Ulissi Machine Learning: Science and Technology 2020 [Paper Link]

#### EXPERIENCE

#### Graduate Researcher

University of California

Sep 2021 - Present

Davis, CA

• Working under the guidance of *Ambarish Kulkarni* on studying novel thermal and electro-catalysts for energy and chemical transformations.

#### Machine Learning Accelerated Catalysis and Materials Design

• Led ML-enabled simulation efforts for DOE and industry-funded projects on catalyst deactivation, electrochemical separations, and near surface gas phase reactions.

- Engineered machine learned interatomic potentials(MLIPs) to model adsorption, diffusion, and reactions on catalytic surfaces—achieving up to 1000x speedups over density functional theory (DFT) and enabling mechanistic insights into reactions like methanol partial oxidation on silver and the reverse water-gas shift.
- Developed automated frameworks for transition state discovery and kinetic modeling by integrating graph and descriptor-based models with open-source Python libraries.
- Leveraged these MLIPs to simulate diverse catalytic and materials systems and support theory-experiment collaboration through advanced techniques including Microkinetic Modeling, Kinetic Monte Carlo, Metadynamics, and On-the-fly Probability Enhanced Sampling.
- Co-developed and deployed active learning pipelines for model training and data generation Code

## Data-Driven Screening of Transition Metal Complexes for Oxygen Separation

- Designed computational workflows and curated datasets for high-throughput DFT screening of transition metal complexes and their  $O_2$  binding properties for electrochemical oxygen separation yielding over 300 new candidate materials.
- Benchmarked DFT functionals against coupled cluster methods to evaluate accuracy and performance in property prediction.

## **Research Assistant**

## Carnegie Mellon University

## Development of ML-Enhanced Frameworks for Atomistic Simulations

- Worked on an ARPA-E-funded project on surface segregation using deep reinforcement learning, focusing on dataset preparation and Bayesian hyperparameter optimization.
- Contributed to the development of open-source tools and sampling strategies to build an active learning framework leveraging physics-informed Behler-Parrinello Neural Networks to accelerate Nudged elastic band calculations and molecular dynamics. Code
- Built and validated software for global optimization of nanoclusters using a modified genetic algorithm enhanced with neural networks and Gaussian processes Code

## **Research Intern**

Singapore University of Technology and Design

- Worked on building and studying lightweight solar PV modules with a poly-carbonate substitute under supervision of Arief S Budiman

## Awards

NREL Kestrel compute grant 50,000+ node hrs	Oct 2024 - Present
NSF ACCESS Discover and Maximize compute grants 8,000,000 Credits	Oct 2021 - Jan 2025
Multiple NERSC compute grants 50,000+ CPU and GPU node hrs	Nov 2021 - Present

## Projects

- Enhanced multi-modal transformers for predicting density of states in crystalline materials, improving training efficiency and model performance. Code
- Accelerated the Chebyshev Interaction Model by porting from OpenMP to GPU with Nsight Compute profiling, boosting performance for large-scale simulations. Code
- Applied ML techniques (NNs, k-NN, decision trees, logistic regression, PCA, t-SNE) to classification and time-series analysis, achieving  $\geq 96\%$  accuracy on the UC Irvine Musk dataset and uncovering trends in COVID-19 data from the New York Times.

Jan 2020 – Jul 2021 Pittsburgh, PA

May 2018 – Aug 2018 Singapore

## Conferences & Posters/Talks

An Active Learning Framework for Accelerating Saddle Point Searches	Nov 2020
AIChE Annual Meeting 2020 [Poster Abstract Link]	(Virtual)
Developing integrated experiment-theory approaches for heterogeneous catalysis	April 2021
UC Davis - Sandia Symposium [Poster Link ]	Davis, CA
Microkinetic modeling for integrated near surface gas phase kinetics	May 2022
CeRCAS Meeting 2022 San	Francisco, CA
<b>Optimizing Chebyshev Interaction Model with Parallel Programming</b>	Oct 2024
NERSC Annual User Group Meeting	Berkeley, CA
Other professional activities	

Judge, UC Davis Chemical Engineering symposium	$Sep \ 2023$
Teaching Assistant, Chemical Reaction Engineering	Fall 2023
Head, Media Relations – Pragyan (NITT's Technical Organization)	Jan 2017 - Jul 2019