

# VAIDISH SUMARIA

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## EDUCATION

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**University of California Los Angeles**

Ph.D. Chemical Engineering

*Aug. 2018-Aug. 2022*  
(Dissertation Year Fellowship Awardee) GPA: 4.0

**Carnegie Mellon University, Pittsburgh**

MS Chemical Engineering

*Jun. 2016-Dec. 2017*

GPA: 4.0

**University of Petroleum and Energy Studies, Dehradun, India**

B.Tech. Chemical Engineering with Specialization in Refining and Petrochemicals

*Jun. 2012-May 2016*

(Silver Medalist) GPA: 3.57

**Relevant Graduate Courses:** AI & Machine Learning, Neural Networks and Deep Learning, Data Management and Statistical Computing, Statistical Thermodynamics, Mathematical Modeling, Statistical Thermodynamics, Electrochemical Energy Systems, Industrial Chemical Technology, Interface Engineering.

## TECHNICAL SKILLS

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**Programming:** Python, Matlab, C++, SAS, R, Scikit-Learn, PyTorch

**Modeling:** VASP, GPAW, Atomic Simulation Environment, LAMMPS, Comsol, Aspen Plus

**Experimental:** Gas Chromatography, Temperature Programmed Desorption, BET Surface Area Analyzer

## RESEARCH EXPERIENCE

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**Research Assistant, Sautet Lab, UCLA, USA**

*Aug. 2018- Present*

- Developed a C-O bond distance based energy correction for the DFT energies to solve "Pt/CO Puzzle".
- Using thermodynamic analysis, generated stability diagram to understand Pt-CO system in real reaction conditions.
- Developed High Dimensional Neural Network potential for the system to investigate surface reconstruction.
- Designed modified Grand Canonical Basin Hopping to efficiently explore configurations.
- Model sophisticated Pt reconstruction induced by CO using nudged elastic band theory and meta-dynamic calculations. **(Publication in preparation)**

**Research Assistant, Viswanathan Lab, CMU, Pittsburgh**

*Sep. 2017-Jul. 2018*

- Developed a rigorous framework to propagate uncertainty within thermodynamic catalytic activity-prediction models.
- Defined a computational measures: prediction efficiency and confidence values to quantitatively the correct energy descriptors for a given reaction scheme and degree of confidence in the predicted surface phase diagrams respectively.
- Studied electrochemical oxygen, hydrogen and chlorine evolution reactions: activity and selectivity predictions

**Mitacs Globalinks Intern, University of Saskatchewan, Canada**

*Jun. 2017-Aug. 2017*

- Optimized fixed bed adsorption by functional polymers for the removal of refractory nitrogen species from bitumen derived gas oil. Effects of temperature and flow rates were also studied and used to develop a mathematical model.

**Graduate Researcher, Gopalakrishnan Lab, UPES, India**

*Aug. 2013-Apr. 2014*

- Optimized the synthesis parameters of ZSM-12 catalyst for the direct conversion of n-Butane to iso-octane.
- Devised a seed assisted synthesis technique to avoid the use of expensive and toxic organic structure directing agents.

## INDUSTRIAL EXPERIENCE

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**Computational Materials Science Intern, Robert Bosch LLC**

*Jun. - Sep. 2022*

- Utilized computational methods to understand electrode degradation under oxidizing conditions.

- Developed a method to computationally generate Cyclic Voltammetry results to understand the the surface reconstructions.

#### Computational Chemistry Intern, Phaseshift Technologies Inc.

Jan. - Sep. 2021

- Created a database for Bulk Metallic glass (BMG) alloy properties to train machine learning models for mechanical property prediction
- Developed framework for creating interatomic potentials for BMG with different compositions for rapid exploration.
- The potentials were used to estimate the liquid and glass transition temperatures.

#### Intern, Reliance Industries Limited, Patalganga, India

Jun.-Jul. 2014

- Designed Prefractionation column for Para-xylene unit to combine the operations of two columns.
- Developed Aspen model had the Naphtha feed separated into the lighters (C1-C6), the heaviers (C10 onwards) and the heart-cut (C7-C9) was obtained from a side-stream with the required concentrations.

#### PUBLICATIONS

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- Uncovering the mechanism of Pt catalyst restructuring under a pressure of gas. **ChemRxiv**, 2022.
- CO organization at ambient pressure on stepped Pt surfaces: First principle modeling accelerated by neural networks. **Chem. Sci.**, 2021.
- Optimal packing of CO at high coverage on Pt(100) and Pt(111) surfaces, **ACS Catal.**, 2020.
- Maximal predictability approach for identifying the right descriptors for electrocatalytic reactions, **J. Phys. Chem. Lett.**, 2018, 9 (3), pp 588595.
- Quantifying Confidence in DFT Predicted Surface Pourbaix Diagrams and Associated Reaction Pathways for Chlorine Evolution, **ACS Catal.**, 2018, 8 (10), pp 90349042.
- Quantifying robustness of DFT predicted pathways and activity determining elementary steps for electrochemical reactions, **J. Chem. Phys.** 150, 041717 (2019).

#### CONFERENCE

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- Sumaria, V., Sautet, P. and Tao, F.F., *Modeling CO-induced Pt reconstruction using DFT-trained Neural Network Potential*, **International Conference on Theoretical Aspects of Catalysis 2022, Lyon, France**
- Sumaria, V. and Sautet, P., *CO Organization at Ambient Pressure on Stepped Pt Surfaces: First Principle Modeling Accelerated By Neural Networks*, **The 27th North American Catalysis Society Meeting, 2022, NYC** (Poster)
- Sumaria, V., Nguyen, L., Sun Geng, Tao, F.F. and Sautet, P., *Optimal Packing of CO at High Coverage on Pt(100) and Pt(111) Surfaces*, **AIChE Annual Meeting 2020** (Virtual Talk)
- Sumaria, V., Krishnamurthy, D., and Viswanathan V., *Self-Formed Catalysts Using Electrochemical (de)-Lithiation for Oxygen Evolution Reaction* **232<sup>nd</sup> Electrochemical Society Meeting, 2017**
- Sumaria, V., Krishnamurthy, D., and Viswanathan V., *How to choose the right descriptor for electrochemical reactions?* **Suncat Summer Institute, 2017** (Poster Presentation)
- Sumaria, V., Krishnamurthy, D., and Viswanathan V., *Quantification of Uncertainty in Activity Relationships for Oxygen Evolution Reaction*. **Quantum Revolutions, Pittsburgh Quantum Institute, 2017** (Best Poster Award)
- Deshpande, S., Krishnamurthy, D., Sumaria, V., Viswanathan, V. *A framework for Identifying Optimal Descriptors and Quantifying Uncertainty DFT-predicted Electrocatalytic Activity*. **25<sup>th</sup> North American Catalysis Society Meeting, 2017**