



# Mehrad Ansari





Personal website: https://mehradans92.github.io

**Summary:** 

A detail-oriented chemical engineer with a strong background in computational modeling, data science and scientific software development, supported by research and professional work experience in process design and optimization.



#### **Education**



### Doctor of Philosophy in Chemical Engineering Master of Science in Chemical Engineering

(May 2023) (Oct 2021)

University of Rochester, Rochester, NY

Thesis: "Physics-informed Machine Learning in Chemical Engineering"

### **Master of Science in Environmental Engineering**

(May 2018)

Missouri University of Science and Technology (UMR), Rolla, MO

Thesis: "Numerical Modeling of Capillary-driven Flow in Open Microchannels: An Implication of Optimized Wicking Fabric Design"

### **Bachelor of Science in Chemical Engineering**

(July 2015)

University of Tehran, Iran

Thesis: "Experimental Setup and Optimization for Electro-catalytical Generation of Hydroxyl Radicals in Wastewater Treatment"



### Work Experience and Practical Training



Research Assistant

(2019 - present)

University of Rochester, Rochester, NY

- Developing <u>edge-computing deep learning frameworks</u> for semi-supervised classification of the activity of antimicrobial peptides via positive-unlabeled learning
- Developed a disease modeling framework to predict future disease spreads and infer location of patient-zero
- Developed an automated <u>framework</u> in CFD modeling that reduces the number of simulations using active learning and generates a symbolic equation for the system of interest via symbolic regression
- Contributed to development of a <u>simulation-based inference framework</u> via maximum entropy re-weighting
- Contributed to development of a plugin with TensorFlow GPU-accelerated operations combined with HOOMD-Blue molecular dynamics simulation engine (HOOMD-TF)
- Developed a web-app for peptide-based gelator transparency classification using Kernel ridge regression
- Developed an <u>automated module</u> on a Raspberry-Pi for real-time monitoring of HPC using Python, JS and HTML
- Implemented finite difference analysis in Python to study 2D shallow water dynamics
- Implemented Monte Carlo simulations in MATLAB to study evolution of spin configurations of a ferromagnet using the Ising model

### **Energy & Materials Research Intern**

(May 2022 - present)

Toyota Research Institute, Los Altos, CA

> Developing deep learning frameworks to infer cell degradation in battery informatics

# Teaching Assistant of "Advanced Transport Phenomena"

(Aug - Dec 2020)

University of Rochester, Rochester, NY

> Tutored students on homework related problems

## Teaching Assistant of "Fundamentals of Fluid Mechanics"

(Jan - May 2020)

University of Rochester, Rochester, NY

> Tutored students on homework related problems and organized laboratory experiments

## Lead CFD Analyst at Missouri S&T Solar Car Design Team

(2016 - 2018)

 ${\it Missouri\ University\ of\ Science\ and\ Technology,\ Rolla,\ MO}$ 

- > Developed validated wind tunnel simulations in STAR-CCM+ for aerodynamic optimization of the solar car
- Improved aerodynamic design efficiency prior to manufacturing

# **Manufacturing Process Modeling Intern**

(May-Dec 2017)

The Goodyear Tire & Rubber Company, Akron, OH

- Phase-change heat transfer modeling and optimization of tire vulcanization process in ANSYS
- Model verification based on plant data and analytical solution
- > Utilized assets more efficiently through MATLAB post processing and automating the simulation process using OPTIMUS
- Provided faster simulation results using Adaptive Mesh Refinement and High-Performance Computing
- GUI development and coupling ANSYS with MATLAB for time-effective post processing

Research Assistant (2016 - 2018)

Missouri University of Science and Technology, Rolla, MO

- > Numerical modeling of multiphase flow in open microfluidics using ANSYS and STAR-CCM +
- Reduced simulation run-time by developing an algorithm for Adaptive Mesh Refinement (AMR)
- Increased solver's stability by developing an algorithm for Adaptive Time Step





# Mehrad Ansari





(Jan-May 2017)

(2014-2015)

(July-Sept 2014)



# **Work Experience and Practical Training**



Teaching Assistant of "Applied Numerical Methods in CFD" Missouri University of Science and Technology, Rolla, MO

- Lectured on Finite Difference Analysis in fluid dynamics, heat and mass transfer using MATLAB
- Organized CFD and programming workshops for ANSYS and Star-CCM +

### Teaching Assistant of "Process Control"

University of Tehran, Iran

Tutored undergraduate students in process control using MATLAB and VisSim

### **Engineering Intern**

Emden-Leer University of Applied Sciences, Emden, Germany

Design of experiments in advanced oxidation process (AOP) for wastewater treatment

**President of IAESTE Iran** (Apr 2013-July 2014)

International Association for the Exchange of Students for Technical Experience

Led a team of college students that organized technical internships internationally



#### **Computer Skills**



**Scientific Softwares Developed:** 

- MaxEnt AL-CFD
- Py0 • Peptide.bio
- HOOMD-TF • <u>GTP</u>

Other tools: MATLAB, STAR-CCM+, ANSYS, OPTIMUS, CATIA

Languages: Python, JavaScript, HTML, CSS



## **Honors and Awards**



1st place winner at Battery Informatics & ML Competition (Dec 2022) Materials Research Society, Boston, MA Kwang-Yu and Lee-Chien Wang Fellowship (Nov 2021) Department of Chemical Engineering, University of Rochester Earl W. Costich Graduate Fellowship (May 2020) Department of Chemical Engineering, University of Rochester 1st place winner: 2017 Mike Alizadeh Scholarship (Aug 2017) American Society of Civil Engineers (ASCE) Recognized reviewer: Journal of Environmental Chemical Engineering (May 2016) (Mar 2014) **MATLAB Programming Contest** Sharif Computer-Aided Chemical Engineering Contest (SC<sub>3</sub>), Sharif University of Technology, Iran

#### **Publications and Patents**



History-agnostic Battery Life-cycle Inference 1.

M Ansari, S Sun, S Torrisi, A Trewartha

(Patent in progress)

Do Large Language Models Know Chemistry?

AD White, GM Hocky, HA Gandhi, M Ansari, S Cox, GP Wellawatte, S Sasmal, Z Yang, K Liu, Y Singh, WJ Peña Ccoa

(Submitted Dec 2022)

(Submitted Dec 2022)

Serverless Prediction of Peptide Properties with Recurrent Neural Networks

Journal of Chemical Information and Modeling

**M Ansari**, AD White

Book chapter: Hyper-parameter Optimization in Deep Learning (July 2022)

Deep Learning for Molecules and Materials, Living Journal of Computational Molecular Science

M Ansari, AD White.

Inferring Spatial Source of Disease Outbreaks using Maximum Entropy (July 2022)

American Physical Society, Physical Review E M Ansari, D Soriano-Paños, G Ghoshal, AD White.

**Iterative Symbolic Regression for Learning Transport Equations** 

(March 2022)

AIChE Journal, Special Edition for AI M Ansari, HA Gandhi, DG Foster, AD White

Simulation-based Inference with Approximately Correct Parameters via Maximum Entropy

(Apr 2022)

Machine Learning in Science and Technology R Barrett, M Ansari, G Ghoshal, AD White



### **Presentations and Talks**



1.	Maximum Entropy Inference in Chemical Reaction Networks with Unknown Kinetic Parameters	(Nov 2022)
	AIChE, Phoenix, AZ	

- Wang Lecture: Rescuing Physics-based Model with Maximum Entropy Reweighting (Sept 2022) Department of Chemical Engineering, Rochester, NY
- Positive Unlabeled Learning of Peptide Properties (July 2022) Fundamentals of Molecular Modeling and Simulation, Delavan, WI
- Serverless Prediction of Peptide Properties with Recurrent Neural Networks (June 2022) Middle Atlantic Regional Meeting of the American Chemical Society, Ewing Township, NJ
- Simulation-based Inference with Approximately Correct Parameters via Maximum Entropy (Dec 2020) Advances in Neural Information Processing Systems 33: Workshop on Machine Learning for Structural Biology