



# Mehrad Ansari





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 web-based deep learning frameworks 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 framework 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 simulation-based inference framework 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 automated module 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)

Tovota Research Institute, Los Altos, CA

Developing deep learning frameworks to infer degradation in battery informatics

#### Teaching Assistant of "Advanced Transport Phenomena"

University of Rochester, Rochester, NY

Tutored students on homework related problems

#### Teaching Assistant of "Fundamentals of Fluid Mechanics"

(Jan - May 2020)

(Aug - Dec 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)

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





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



#### **Work Experience and Practical Training**



(Jan-May 2017)

(2014-2015)

(July-Sept 2014)

(Apr 2013-July 2014)

(July-Sept 2011)

(May 2020)

(Submitted June 2022)

(July 2022)

(July 2022)

(March 2022)

(Apr 2022)

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** 

University of Tehran, Iran

Emden-Leer University of Applied Sciences, Emden, Germany

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

**President of IAESTE Iran** 

International Association for the Exchange of Students for Technical Experience

Led a team of college students that organized technical internships internationally

**MATLAB Programming Tutor** 

Organized advanced programming workshops for engineering students

Scientific Softwares Developed:

### **Computer Skills**

 MaxEnt Py0 • AL-CFD • Peptide.bio

• HOOMD-TF GTP

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

Languages: Python, JavaScript, HTML, CSS

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#### **Honors and Awards**



Kwang-Yu and Lee-Chien Wang Fellowship (Nov 2021) Department of Chemical Engineering, University of Rochester

Earl W. Costich Graduate Fellowship

Department of Chemical Engineering, University of Rochester

First place winner: 2017 Mike Alizadeh Scholarship (Aug 2017)

American Society of Civil Engineers (ASCE)

Recognized reviewer: Journal of Environmental Chemical Engineering (May 2016) MATLAB programming contest (Mar 2014)

University of Sharif Computer-Aided Chemical Engineering Contest (SC<sub>3</sub>)

Sharif University of Technology, Iran

# **Publications**



(Submitted July 2022) Do Large Language Models Know Chemistry? 1. AD White, GM Hocky, HA Gandhi, M Ansari, S Cox

Serverless Prediction of Peptide Properties with Recurrent Neural Networks

**Bioinformatics** M Ansari, AD White

Book chapter: Hyper-parameter Optimization in Deep Learning

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

Inferring Spatial Source of Disease Outbreaks using Maximum Entropy

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

<u>Iterative Symbolic Regression for Learning Transport Equations</u>

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

Simulation-based Inference with Approximately Correct Parameters via Maximum Entropy

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

## **Presentations and Talks**



Positive Unlabeled Learning of Peptide Properties 1. (July 2022) Fundamentals of Molecular Modeling and Simulation, Delavan, WI

Serverless Prediction of Peptide Properties with Recurrent Neural Networks Middle Atlantic Regional Meeting of the American Chemical Society, Ewing Township, NJ

3.

Simulation-based Inference with Approximately Correct Parameters via Maximum Entropy Advances in Neural Information Processing Systems 33: Workshop on Machine Learning for Structural Biology

(Dec 2020)

(June 2022)