### Mehdi Foroozandeh Shahraki PhD Student | Graduate Research Assistant

I am a PhD student in the Computing Science department at SFU. My research interests are machine learning, probabilistic modeling, and genomics. I am a graduate research assistant in the Computational Biology lab supervised by Dr. Maxwell Libbrecht.



2022 - 2026 **Ph.D. Computing Science**, Simon Fraser University

MSc. Computing Science, Simon Fraser University, CGPA: 4.0 /4.33,

2020 - 2022 Courses: Machine Learning, Statistical Machine Learning, Design and Analysis of Algorithms, Problem-based Learning in Bioinformatics

BSc. Cell and Molecular Biology- Microbiology, University of Tehran, *GPA (Final Two Years) : 3.68 /4*,

Selected Courses: Biostatistics, Molecular Genetics, Epigenetics, Virology, Evolutionary Biology, Developmental Biology, Animal Cell Differentiation



#### September 2020 Present

#### Research Assistant, SIMON FRASER UNIVERISTY, Department of Computing Sciences

- > At Libbrecht Computational Biology Lab
- > Research on evaluating the reproducibility of predictions of a class of probabilistic graphical models (SAGA methods).
- > Research on building and optimizing deep learning models capable of predicting bacterial antibiotic resistance from genome sequence data.

Machine Learning | Probabilistic Modeling | Genomics | Epigenomics | Deep Learning | Drug Resistance

#### January 2019 September 2020

#### Research Assistant, University of Tehran, IBB- Department of Bioinformatics

- > At Complex Biological Systems and Bioinformatics (CBB) lab, led by Dr. Kavousi.
- > Research on developing machine learning and deep learning models for predicting enzyme properties.
- > Research on the discovery of novel enzymes with particular applications from metagenomic sources.
- > Collaboration in writing and editing academic publications.

Machine Learning Metagenomics Enzymes Bioinformatics

# Publications

Foroozandeh Shahraki, Mehdi, et al. "A computational learning paradigm to targeted discovery of biocatalysts from metagenomic data: a case study of lipase identification" Biotechnology and Bioengineering:21-130.R1 (2022).

Foroozandeh Shahraki, Mehdi, *et al.* "A generalized machine-learning aided method for targeted identification of industrial enzymes from metagenome: A xylanase temperature dependence case study." Biotechnology and Bioengineering (2020).

Foroozandeh Shahraki, Mehdi, *et al.* "MCIC: Automated Identification of Cellulases From Metagenomic Data and Characterization Based on Temperature and pH Dependence." Frontiers in microbiology 11 (2020).

Maleki, Morteza, *et al.* "A novel thermostable cellulase cocktail enhances lignocellulosic bioconversion and biorefining in a broad range of pH." International journal of biological macromolecules 154 (2020): 349-360.

Ariaeenejad, Shohreh, *et al.* "A novel high performance in-silico screened metagenome-derived alkalithermostable endo- $\beta$ -1, 4-glucanase for lignocellulosic biomass hydrolysis in the harsh conditions." BMC biotechnology 20.1 (2020): 1-13.

Foroozandeh Shahraki, Mehdi, et al. "Evaluating the reproducibility of segmentation and genome annotation (SAGA) algorithms" Poster at Machine Learning in Computational Biology 2021 (full submission in progress).

### PROJECTS

#### REPRODUCIBILITY OF SAGA ALGORITHMS

2021 - 2022

Research on evaluating the reproducibility of predictions from a well-known class of probabilistic graphical models known as SAGA algorithms. These models, that are usually based on Hidden Markov Models or Dynamic Bayesian Networks, are essentially clustering algorithms for sequence data (genomic and epigenomic data). Similar to most unsupervised learning algorithms, evaluation of their prediction performance is challenging. In this research project, I use statistical principles of reproducibility analysis to evaluate these probabilistic machine learning algorithms.

probabilistic modeling machine learning reproducibility epigenomics genomics

#### Drug Resistance Prediction Using Machine Learning

2020 - 2021

#### github.com/mehdiforoozandeh/DRML

In a series of related projects, I designed and optimized several machine learning and deep learning frameworks to understand and predict anti-microbial drug resistance from genomic data. In this project, I explored various avenues such as customized feature extraction from high dimensional genomic data, comparing several machine learning algorithms for the same prediction problem, and most importantly, implementing Bayesian hyper-parameter optimization to optimize our models more effectively. Particularly, I applied Bayesian hyper-parameter optimization to optimize a massive deep learning model that could not possibly be optimized using conventional grid-searching method.

Deep Learning Bayesian Optimization Drug Resistance

#### PREIDCTION OF ENZYMATIC PROPERTIES USING MACHINE LEARNING AND ENZYME MINING FROM METAGENOME

2019-2020

github.com/mehdiforoozandeh/MeTarEnz github.com/mehdiforoozandeh/MCIC

Research on developing machine learning and deep learning models for predicting enzyme properties heavily relying on amino acid sequence data for model training and feature extraction. I also designed and developed automated enzyme mining software to facilitate discovery of biocatalysts from metagenome data. Finally, using the developed computational infrastructures, we discovered and mined various novel enzymes with particular applications from metagenomic sources.

Enzyme mining | Metagenomics | Machine Learning | Automation

### **SKILLS**

**Programming** Python, MATLAB

Tools and Software TensorFlow, Keras, Scikit-learn, Scikit-optimize, Numpy, Pandas, Biopython, NCBI Toolkit,

Segway, ChromHMM

Other Unix Shell Scripting, Docker, ŁTFX, Linux, Web Scraping

**Laboratory Skills** Microbial Culture, PCR, DNA extraction, Gene Cloning, Gel Electrophoresis

Languages English, Persian (Native)

# CERTIFICATIONS

2019 Machine Learning by Stanford University on Coursera (Grade: 100/100)

2019 Deep Learning with TensorFlow at Sharif University of Technology

2019 Molecular Cloning Practical Workshop, University of Tehran

2018 International Summer School of Brain and Cognitive Sciences at Royan Institute

2017 PCR, Electrophoresis and Drawing of Phylogenetic Tree Practical Workshop, University of Tehran

## **66** References

#### Maxwell Libbrecht

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#### Kaveh Kavousi

Assistant Professor, UT

#### Shohreh Ariaeenejad

Assistant Professor, ABRII

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