Philip Fradkin

646-675-5185

EXPERIENCE

Vector Institute

PhD with Brendan Frey & Bo Wang labs

Website: phil-fradkin.github.io

Email: phil.fradkin@gmail.com

MaRS, Toronto CA September 2020 - Present

- To address the challenge of RNA property prediction, we developed IsoCLR, a novel method that extends contrastive learning techniques to genomic data.
- Utilizing biologically inspired RNA augmentations, IsoCLR learns robust RNA isoform representations, particularly effective under low-data conditions.
- Studied properties of deep learning model optima, connecting model loss landscape flatness with the training procedure. This provides insight into a possible contributor to model generalization.

### Memorial Sloan Kettering

New York, USA

Morris Lab visiting student

Februrary 2023 - August 2023

• To overcome challenges in annotation and cell type out-of-distribution detection in single-cell RNA sequencing, developed an energy-based model, that enhanced annotation accuracy and introduced OOD detection, outperforming existing techniques in single-cell data modeling.

### **Deep Genomics**

MaRS, Toronto CA

Machine Learning Graduate Intern

May 2022 - September 2022

- Design self-supervised learning training procedures for effective mRNA representations to improve molecular phenotype prediction.
- Investigated effectiveness of deep learning mRNA half-life models in predicting variant effects. Based on these results demonstrated a richer RNA representation results in learning generalizable signal.

### Deep Genomics

MaRS, Toronto CA

Computational Biologist

May 2016 - August 2020

- Developed deep neural networks to predict cellular biology phenotypes. Implemented a siamese neural network to predict the stability of RNA molecules from sequence surpassing the performance of state of the art.
- Developed software systems that support both wet-lab and computational scientists, such as automation of the design of antisense oligonucleotides short stretches of modified RNA resulting in the automation of a laborious error-prone process.
- Coordinated the experimental validation of computational insights. Based on the results of these investigations, confirmed the regulatory function of polyadenylation on RNA half-life and robustness of in-silico predictions.

# **EDUCATION**

### University of Toronto (Ph.D.)

Computer Engineering, Vector Institute, Brendan Frey and Bo Wang labs

Toronto, ON, CA 2020 - Present

### Teaching Assistant:

- Neural Networks and Deep Learning (CSC2516)
- Introduction to Machine Intelligence (ECE324)

#### **Select Courses:**

- Probabilistic Learning and Reasoning (CSC2506)
- Information Theory (ECE1502)
- Trustworthy Machine Learning (CSC2559)

# University of Toronto (B.Sc.)

Honors Bachelor of Science

Toronto, ON, CA 2012 - 2017

Computer Science Major Computational Biology Major

## Publications & Presentations

• A Graph Neural Network Approach to Molecule Carcinogenicity Prediction:

Publication in Bioinformatics, Oral at ISMB (2022), and LMRL - NeurIPS workshop; *Philip Fradkin*, Adamo Young, Lazar Atanackovic, Brendan Frey, Leo J Lee, Bo Wang Bioinformatics, Volume 38, Issue Supplement\_1, July 2022, Pages i84i91, https://doi.org/10.1093/bioinformatics/btac266 (2022)

• Robustness to Adversarial Gradients: A Glimpse Into the Loss Landscape of Contrastive Pre-training:

Poster at ICML 2022 Pre-Training Workshop; *Philip Fradkin* Lazar Atanackovic, Michael R. Zhang. (2022)

• Energy-based Modelling for Single-cell Data Annotation:

Proceedings of Machine Learning Research:

Tiayni Liu, *Philip Fradkin*, Lazar Atanackovic, Leo J. Lee PMLR 200:94-109 https://proceedings.mlr.press/v200/liu22b.html (2022)

• ATP7B Variant p.Met645Arg Causes Wilson Disease by Promoting Exon 6 Skipping:

Publication in NPJ Genomic Medicine;

Daniele Merico, Carl Spickett, Matthew OHara, Boyko Kakaradov, Amit G. Deshwar, *Philip Fradkin*, Shreshth Gandhi, Jiexin Gao, Solomon Grant, Ken Kron, Frank W. Schmitges, Zvi Shalev, Mark Sun, Marta Verby, Matthew Cahill, James J. Dowling, Johan Fransson, Erno Wienholds, Brendan J. Frey npj Genom. Med. 5, 16. https://doi.org/10.1038/s41525-020-0123-6 (2020)

## Conference Reviews & Frameworks

NeurIPS: LMRL 2022, MetaLearn 2022

ICML: Computational Biology Workshop 2023, Pre-Training Workshop 2022

Python: Pytorch, DGL, Keras, Tensorflow

Julia: Zygote, Distributions.jl

#### Interests

Hackathon Founding member of Toronto bioinformatics hackathon - 2017.

Wilderness orientation Wilderness outdoor orientation program leader.

Outdoor Adventures In my spare time I enjoy rock climbing outdoors, and hiking. I recently completed a 100km backpacking trip in Canadian Rockies.