# **Donald Loveland**

Email: dlovelan@umich.edu | Website: donaldloveland.com | LinkedIn: donald-loveland

## RESEARCH INTERESTS

I am a second year PhD student at the University of Michigan, Ann Arbor. My research focuses on fundamental questions in **machine learning**, **graph representation learning** and **graph neural networks (GNNs)**. Broadly, I study how to improve the interpretability, robustness, and fairness of GNNs in real-world settings such as social networks.

## **EDUCATION**

## University of Michigan, Ann Arbor

Ph.D. in Computer Science (3.93/4.00 GPA)

• Advisor: Danai Koutra - Graph Exploration and Mining at Scale (GEMS) Lab

• Fellowships: Rackham Merit Fellow

## California Polytechnic State University, San Luis Obispo

B.S. in Physics (3.61/4.00 GPA)

• Minors: Computer Science, Mathematics, Astronomy, Ethnic Studies

Aug. 2014 - June 2018 San Luis Obispo, CA

Aug. 2021 - May 2026

Ann Arbor, MI

# RECENT RESEARCH EXPERIENCE

## **MIT Lincoln Laboratory**

Research Intern - Advised by Dr. Rajmonda Caceres

May 2022 - Aug. 2022

Developed a gradient-based optimization method to efficiently edit graph structures and improve task performance.
Demonstrated the capabilities of this method through shortest path and substructure manipulation. Proposed method extended to Hyperbolic GNNs, and tested on a variety of other GNN architectures, demonstrating improvements of up to 60% over baseline heuristics. Paper in Submission

## **Lawrence Livermore National Lab**

Feb. 2019 - Sept. 2021

Applied Machine Learning Researcher

- Developed an XAI method for counterfactual introspection of DL models in PyTorch. Method was applied to materials science problems to extract domain-specific concepts learned by convolution neural networks. US patent recieved.
- Explored the relationship between adversarial training and post-hoc XAI methods for GNNs in PyTorch Geometric. Findings were presented at ICML XAI Workshop and used to elucidate important molecular substructures.
- Created an inverse design method for molecular graphs through a GNN-based RL model. New molecules were designed through an interpretable graph editing policy. Work was used to aid scientists in molecule discovery tasks.

#### PREPRINTS AND IN PROGRESS

- D. Loveland, R. Caceres, "EdgeEdit: Graph Neural Network-guided Edge Editing for Task Optimization", To submit to ECML-PKDD, 2023
- **D. Loveland**, J. Zhu, M. Heimann, B. Fish, M. Schaub, D. Koutra, "Characterizing Performance Discrepancies Across Local Homophily Ratios in Graph Neural Networks", *Submitted to KDD 2023*

## **PUBLICATIONS**

- **D. Loveland**, J. Zhu, M. Heimann, B. Fish, M. Schaub, D. Koutra, "On Graph Neural Network Fairness in the Presence of Heterophilous Neighborhoods", *KDD Workshop on Deep Learning on Graphs*, 2022
- J. Zhu, J. Jin, D. Loveland, M. Schaub, D. Koutra, "On the Relationship Between Heterophily and Robustness of Graph Neural Network", KDD, 2022

- **D. Loveland**, S. Liu, B. Kailkhura, A. Hiszpanski, Y. Han, "Reliable Graph Neural Network Explanations Through Adversarial Training", *ICML Workshop on Theoretic Foundation, Criticism, and Application Trend of Explainable AI*, 2021
- P. Nguyen\*, D. Loveland\*, J. Kim, P. Karande, A. M. Hiszpanski, Y. Han, "Predicting Energetics Materials' Crystalline Density from Chemical Structure by Machine Learning", *Journal of Chemical Information and Modeling*, 2021
- **D. Loveland**, B. Kailkhura, P. Karande, A. M. Hiszpanski, Y. Han, "Automated Identification of Molecular Crystals' Packing Motifs", *Journal of Chemical Information and Modeling*, 2020
- B. Gallagher, M. Rever, D. Loveland, T. N. Mundhenk, B. Beauchamp, et al, "Predicting Compressive Strength of Consolidated Molecular Solids using Computer Vision and Deep Learning", *Materials and Design*, 2020
- S. Liu, B. Kailkhura, J. Zhang, A. M. Hiszpanski, E. Robertson, **D. Loveland**, Y. Han. "Actionable attribution maps for scientific machine learning", *ICML Workshop on ML Interpretability for Scientific Discovery*, 2020
- S. Liu, B. Kailkhura, **D. Loveland**, Y. Han, "Generative Counterfactual Introspection for Explainable Deep Learning", *IEEE Global Conference on Signal and Information Processing (GlobalSIP*), 2019
- V. N. Bennert, **D. Loveland**, E. Donohue, M. Cosens, S. Lewis, et al, "Studying the [OIII] $\lambda$  5007Å emission-line width in a sample of 80 local active galaxies: a surrogate for  $\sigma^*$ ?", *Monthly Notices of the Royal Astronomical Society*, 2018

## **TECHNICAL SKILLS**

- Programming Languages: Python, Java (Basic)
- Packages: Pytorch, Pytorch-Geometric, Scikit-Learn, NetworkX

#### ADDITIONAL EXPERIENCE

## **Cambridge Coaching**

Dec. 2021 - Present

Computer Science and Math Tutor

• Tutoring students at various academic stages in courses such as high school AP computer science, undergraduate calculus, and graduate machine learning.

## Cal Poly College of Science and Math

August. 2016 - Dec. 2017

Peer Mentor/Advisor

• Mentored first-year students on academic probation with the goals of improving study skills, promoting time management, and helping with general adjustment to college.

<sup>\*</sup>Denotes equal contribution