Haoyun Lei

E-mail: <u>haoyunl@andrew.cmu.edu</u> | Phone: +1(412)969-3798

LinkedIn: <u>linkedin.com/in/haoyunlei/</u> | Website: <u>https://leovam.github.io/</u>

EDUCATION

Carnegie Mellon University

Aug 2016 – May 2021

Ph.D. in Computational Biology

(expected)

Joint Carnegie Mellon-University of Pittsburgh Ph.D. Program in Computational Biology Computational Biology Department, School of Computer Science

Huazhong University of Science and Technology

Sep 2008 – Jun 2012

B.S. in Biological Science

College of Life Science and Technology

SKILLS

Programming Languages: Python (proficient), R (fluent), MATLAB (fluent), Shell (fluent), Java (familiar)

Technical Skills and Tools: Machine Learning (scikit-learn), Deep Learning (PyTorch, TensorFlow), Bioinformatics (GATK, SAMtools, bedtools CNVkit etc.), Data Analysis (Numpy, Scipy, Pandas), Data Visualization (Matplotlib, Seaborn), Combinatorial Optimization (Gurobi, SCIP), Cloud Computing (AWS), Web Development (HTML/CSS/JS)

WORK EXPERIENCE

Laboratory Corporation of America Holdings (LabCorp)

May 2020 - Jul 2020

Bioinformatics Summer Intern

Westborough, MA

Converting Free-text Patient Data to ICD Codes using Natural Language Processing (PyTorch, TensorFlow)

- Explored language tools (BioBERT, medaCy) to annotate and chunk the important information in medical text
- Fine-tuned **BERT** model on ICD-10 code classification at chapter and block (first three characters) level
- Designed a two-step BERT model to predict multiple ICD-10 codes in LabCorp's patient medical text
- Managed to work on a small dataset and reached 84% on multi-label clarification at chapter level

Benchmarking CNV Detection Tools (Python, R, Perl)

- Tested and compared public CNV detection tools for calling CNVs in targeted NGS data with a very small panel
- Explored combinations of parameters of tools to increase true positive detection in CNVkit, DECoN & CoNVaDING
- Designed algorithms to rescue and recover CNVs with a weaker signal in a very small panel of targets
- Reached over 94% in sensitivity while kept specificity around 90%

RESEARCH EXPERIENCE

Ph.D. Thesis: Integrating Multiple Data Types to Infer Tumor Evolution (Python, R, MATLAB)

May 2017 - Present

- Created a mixed membership model for the Non-negative Matrix Factorization (NMF) problem
- Developed an efficient coordinate descent algorithm to solve the NMF problem in Python
- Designed a Mixed Integer Linear Programming Model with the popular optimization solvers of Gurobi and SCIP
- Reached ~95% accuracy, surpassing exisiting methods

Detection of Cancer Types and Relevant Features using Deep Learning with RNA-seq Data (PyTorch)

Spring 2020

- Designed and fine-tuned 1D CNN, 2D CNN and a hybrid CNN models to detect cancer types
- Designed a Stacked Denoising Autoencoder Classifier to improve the detections (~96% accuracy)
- Applied embedding method to find implicit relationships between cancer samples and genes

Footprint Match and Pattern Detection using Machine Learning (scikit-learn)

Spring 2017

- Classified ~ 10,000 footprint images with Neural Network and SVM using scikit-learn (~95% accuracy)
- Applied the Scale-invariant feature transform (SIFT) algorithm to the match of saved and new images
- Extracted the image patterns with K-Means and Gaussian Mixture Model

Predict Proto Genes using **Logistic Regression**, **Naïve Bayes Classifier** and **Decision Tree**Spring 2017

Copy Number Extraction from DNA Sequencing Data with **Numpy**, **Scipy** and **Regular Expression**Fall 2016

Model Gene Regulatory Network by combining **Boolean network** and **Ordinary Differential Equation** models

Fall 2016