# Haoyun Lei

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# **SUMMARY**

I design algorithm of optimization to study cancer genetics, inferring phylogeny for tumor evolution from multiple types of genomic data. I also work on interdisciplinary projects of machine learning (ML) and deep learning (DL), and their applications to cancer genomics. I am interested in studying cancer or clinical data using bioinformatics, ML and DL.

# **EDUCATION**

**Carnegie Mellon University** 

Aug 2016 - May 2022

Ph.D. in Computational Biology (Mentor: Dr. Russell Schwartz)

(expected)

Computational Biology Department, School of Computer Science

M.Sc in Machine Learning Machine Learning Department, School of Computer Science

Aug 2020 - Dec 2021

**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
- 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
- Recovered CNVs with a weaker signal in a very small panel of targets with over 94% in sensitivity and 90% in specificity

# 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 tumor evolution problem using bulk and single-cell sequencing data
- Developed an efficient coordinate descent algorithm to solve the DNA and RNA deconvolution problem in Python
- Designed a Mixed Integer Linear Programming Model with the popular optimization solvers of Gurobi and SCIP
- Designed a comprehensive simulator for multiple types of mutations in DNA-seq data with phylogenetic progress

Function Specific Representational Similarity Inference in the Brain (PyTorch)

Spring 2021

- Designed probabilistic graphical model for brain fMRI data
- Developed neurons-independent and neurons-dependent network to study function-specific representational similarity
- Recovered similar structure in correlated functional areas in the brain

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

#### TEACHING EXPERIENCE

# **Algorithm and Advanced Data Structure**

Aug 2019 - Dec 2019

Algorithms: Breadth-first Search, Depth-first Search, Binary Search, Quick Sort, Merge Sort etc.

Data Structure: Linked List, Graph, Tree, Stack, Queue, Heap, ArrayList, Hash Table etc.

Concepts: Recursion, Dynamic Programming, Time and Space Complexity, NP-problem etc.

#### **Laboratory Methods for Computational Biologists**

Aug 2018 - Apr 2019

Designed a faster pipeline combining multiple new analysis tools to detect differentially expressed genes in RNA-seq data

## **PUBLICATIONS & TALKS**

## **Articles**

- Lei, H., Guo, A. X., Tao, T., Ding, K., Fu, X., Oesterreich, S., Lee, V. A. and Schwartz, R. (2022) Semi-deconvolution of bulk and single-cell RNA-seq data with application to metastatic progression in breast cancer. (Submitted to ISMB 2022)
- Fu, X., Lei, H., Tao, Y., and Schwartz, R. (2022). Reconstructing clonal lineage trees incorporating single nucleotide variants (SNVs), copy number alterations (CNAs), and structural variations (SVs). (Submitted to ISMB 2022)
- Lei, H., Gertz, E. M., Schäffer, A. A., Fu, X., Tao, Y., Heselmeyer-Haddad, K., ..., and Schwartz, R. (2021). Tumor heterogeneity assessed by sequencing and fluorescence in situ hybridization (FISH) data.

Bioinformatics 37 (24), 4704-4711

- Fu, X., Lei, H., Tao, Y., Heselmeyer-Haddad, K., Li, G., Shi, X., Xu, L., Torres, I., Hou, Y., Wu, K., Dean, M., Ried, T., and Schwartz, R. (2021). Joint clustering of single cell sequencing and fluorescence in situ hybridization data to infer tumor copy number phylogenies.
- Journal of Computational Biology 28 (11), 1035-1051
- Tao, Y., **Lei, H.**, Fu, X., Lee, A. V., Ma, J., and Schwartz, R. (2020). Robust and accurate deconvolution of tumor populations uncovers evolutionary mechanisms of breast cancer metastasis. ISMB2020, *Bioinformatics*, *36*, i407-i416,
- Lei, H., Lyu, B., Gertz, E., Schäffer, A. A., Shi, X., Wu, K., Li, G., Xu, L., Hou, Y., Dean, M., and Schwartz, R. (2020). Tumor Copy Number Deconvolution Integrating Bulk and Single-Cell Sequencing Data.

RECOMB 2019, Journal of Computational Biology, 27(4) 565-598.

- Tao, Y., Lei, H., Lee, A. V., Ma, J., and Schwartz, R. (2020). Neural Network Deconvolution Method for Resolving Pathway-Level Progression of Tumor Clonal Expression Programs with Application to Breast Cancer Brain Metastases. *Frontiers in Physiology*, 11, 1055.
- Tao, Y., Lei, H., Lee, A. V., Ma, J., and Schwartz, R. (2019). Phylogenies derived from matched transcriptome reveal the evolution of cell populations and temporal order of perturbed pathways in breast cancer brain metastases. ISMCO 2019 (pp. 3-28). Springer, Cham.

# **Abstracts & Talks**

- Lei, H., Gertz, E. M., Schäffer, A. A., Fu, X., Tao, Y., Heselmeyer-Haddad, K., ... and Schwartz, R. (2020, July). Tumor heterogeneity assessed by sequencing and fluorescence in situ hybridization (FISH) data.
- Conference on Intelligent System for Molecular Biology (ISMB), virtual
- Fu, X., Lei, H., and Schwartz, R. (2020, July). Joint Clustering of single cell sequencing and fluorescence in situ hybridization data to infer tumor copy number phylogenies. ISMB, virtual.
- Lei, H., Lyu, B., Gertz, E., Schäffer, A. A., Shi, X., Wu, K., Li, G., Xu, L, Hou, Y., Dean, M., and Schwartz, R. (2019,
- May). Tumor Copy Number Deconvolution Integrating Bulk and Single-Cell Sequencing Data. International Conference on Research in Computational Molecular Biology (RECOMB), Washington, DC.
- Lei, H., Lyu, B., Gertz, E. M., Schäffer, A. A., and Schwartz, R. (2018, October). Tumor Copy Number Data Deconvolution Integrating Bulk and Single-cell Sequencing Data. In 2018 IEEE 8<sup>th</sup> International Conference on Computational Advances in Bio and Medical Sciences (ICCABS), Las Vegas, NV.
- Lei, H., Roman, T., Eaton, J., and Schwartz, R. (2018, July). Deconvolution of tumor copy number data using bulk and single-cell sequencing data. ISMB, Chicago, IL.
- Lei, H., Roman, T., Eaton, J., and Schwartz, R. (2018, April). New directions in deconvolving genomics mixtures of copy number variation data. SIAM Conference on Discrete Mathematics, Denver, CO.