Nathan Riley Summers LaPierre

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RECENT POSITIONS

Postdoctoral Scholar

Aug. 2022 - Present

Department of Human Genetics, University of Chicago

Labs of Xin He and Matthew Stephens

Graduate Student Researcher

Sep. 2016 - June 2022

Department of Computer Science, University of California, Los Angeles

Labs of Eleazar Eskin and Wei Wang

EDUCATION

PhD in Computer Science, June 2022 University of California, Los Angeles

M.S. in Computer Science, 3.74 GPA, March 2019

University of California, Los Angeles

B.S. in Applied Computer Science, 3.86 GPA (magna cum laude), December 2015 George Mason University

PUBLICATIONS

A. Khan, J. Bagley, **N. LaPierre**, et al, "Genetic pathways regulating the longitudinal acquisition of cocaine self-administration in inbred and recombinant inbred mice," *Cell Reports*, vol. 42, no. 8, pp. 112856, Aug. 2023.

N. LaPierre*, B. Fu*, S. Turnbull, E. Eskin, and S. Sankararaman, "Leveraging family data to design Mendelian Randomization that is provably robust to population stratification," *Genome Research*, vol. 33, pp. 1032-1041, July 2023.

N. LaPierre and Harold Pimentel, "Accounting for Isoform Expression in eQTL Mapping Substantially Increases Power," *bioRxiv*, June 2023.

B. Fuqua et al (many authors), "The Genetic Architecture of Dietary Iron Overload and Associated Pathology in Mice," bioRxiv, June 2023.

J. Karlin, L. Gai, **N. LaPierre**, et al, "Ensemble neural network model for detecting thyroid eye disease using external photographs," *British Journal of Ophthalmology*, Sep. 2022.

F. Meyer et al (many authors), "Critical Assessment of Metagenome Interpretation-the second round of challenges," *Nature Methods*, vol. 19, pp. 429440, Apr. 2022.

C. Cinelli, **N. LaPierre**, B. L. Hill, S. Sankararaman, and E Eskin, "Robust Mendelian randomization in the presence of residual population stratification, batch effects and horizontal pleiotropy," *Nature Communications*, vol. 13, no 1., pp. 1093, Mar. 2022.

M. Alser et al (many authors), "Packaging, containerization, and virtualization of computational omics methods: Advances, challenges, and opportunities," *arXiv*, Mar. 2022.

- N. LaPierre*, K. Taraszka*, H. Huang, R. He, F. Hormozdiari, and E. Eskin, "Identifying Causal Variants by Fine Mapping Across Multiple Studies," *PLOS Genetics*, vol. 17, no. 9, pp. e1009733, Sept. 2021.
- J. Bloom et al (many authors), "Massively scaled-up testing for SARS-CoV-2 RNA via next-generation sequencing of pooled and barcoded nasal and saliva samples," *Nature Biomedical Engineering*, vol. 5, no. 7, pp. 657-665, July 2021.
- N. LaPierre, M. Alser, E. Eskin, D. Koslicki*, and S. Mangul*, "Metalign: Efficient alignment-based metagenomic profiling via containment min hash," *Genome Biology*, vol. 21, pp. e242, Sep. 2020.
- **N. LaPierre**, R. Egan, W. Wang, and Z. Wang, "De novo Nanopore read quality improvement using deep learning," *BMC Bioinformatics*, vol. 20, no. 1, pp. e552, Dec. 2019.
- N. LaPierre, C. Ju, G. Zhou, and W. Wang, "MetaPheno: A Critical Evaluation of Deep Learning and Machine Learning in Metagenome-Based Disease Prediction," *Methods*, vol. 166, pp. 74-82, Aug. 2019.
- **N. LaPierre***, S. Mangul*, M. Alser, I. Mandric, N.C. Wu, D. Koslicki, and E. Eskin, "MiCoP: Microbial Community Profiling method capable of detecting low abundance viral and fungal organisms in metagenomic samples," *BMC Genomics*, vol. 20, no. 5, pp. e423, June 2019.
- M. A. Rahman, **N. LaPierre**, H. Rangwala, and D. Barbara, "Metagenome sequence clustering with hash-based canopies," *Journal of bioinformatics and computational biology*, vol. 15, no. 6, pp.1740006, Oct. 2017.
- M. A. Rahman, **N. LaPierre**, and H. Rangwala, "Phenotype Prediction from Metagenomic Data Using Clustering and Assembly with Multiple Instance Learning (CAMIL)," *IEEE/ACM transactions on computational biology and bioinformatics*, Oct. 2017.
- N. LaPierre, M. A. Rahman, and H. Rangwala, "CAMIL: Clustering and Assembly with Multiple Instance Learning for Phenotype Prediction," in *IEEE International Conference on Bioinformatics and Biomedicine*, Shenzhen, China, 2016.
- N. LaPierre and H. Rangwala, "Predicting Clinical Phenotype using OTU-based Metagenome Representation," in *IEEE International Conference on Data Mining Workshop on Biological Data Mining and its Applications in Healthcare*, Atlantic City, New Jersey, 2015, pp. 156-163.
- (* Authors contributed equally)

PRESENTATIONS

"Leveraging family data to design Mendelian Randomization that is provably robust to population stratification," in *International Conference on Research in Computational Molecular Biology (RECOMB)*, Apr. 2023.

"Accounting for Isoform Expression in eQTL Mapping," in CSHL Genome Informatics, virtual, Nov. 2021.

"Metalign: Efficient alignment-based metagenomic profiling via containment min hash," in *Intelligent Systems for Molecular Biology (ISMB) HitSeq*, virtual, July

2020.

"Identifying Causal Variants by Fine Mapping Across Multiple Studies" in *International Conference on Research in Computational Molecular Biology (RECOMB)*, virtual, June 2020.

"Metalign: Efficient alignment-based metagenomic profiling via containment min hash," in *RECOMB-Seq*, virtual, June 2020.

"Identifying Causal Variants by Fine Mapping Across Multiple Studies" in American Society for Human Genetics Annual Meeting, October 2019.

"MiniScrub: de novo long read scrubbing using approximate alignment and deep learning" in Amazon Web Services - UCLA Computational Medicine Symposium, February 2019.

"MiniScrub: de novo long read scrubbing using approximate alignment and deep learning" in American Society for Human Genetics Annual Meeting, October 2018.

"MiCoP: Microbial Community Profiling method capable of detecting low abundance viral and fungal organisms in metagenomic samples" in American Society for Human Genetics Annual Meeting, October 2017.

"CAMIL: Clustering and Assembly with Multiple Instance Learning for Phenotype Prediction" in IEEE International Conference on Bioinformatics and Biomedicine, December 2016.

• Won a conference travel grant sponsored by NSF

"Predicting Clinical Phenotype using OTU-based Metagenome Representation" in IEEE International Conference on Data Mining workshop on Biological Data Mining and its Applications in Healthcare, November 14, 2015

 Won a travel grant from the Undergraduate Student Travel Fund of the Office of Student Scholarship, Creative Activities, and Research at GMU

"Developing a Computational Pipeline for Metagenomic State Classification with Feature Engineering" in Volgenau School of Engineering Undergraduate Research Celebration, April 2015

• Won Outstanding Undergraduate Research Project Award for poster

COMPUTATIONAL Languages / Scripting: Fluent in Python, R, Bash; past experience with C(++) SKILLS and Java

Deep Learning Frameworks: PyTorch, Keras Other Technologies: Git, Docker, LaTeX

Bioinformatics experience: Mendelian randomization; GWAS & statistical fine mapping; alignment; assembly; metagenomics; common software such as Plink, Bedtools, Samtools, Beftools, etc; SGE and Slurm grid/cluster computing systems Machine Learning Algorithms: Deep Learning, Linear & Logistic Regression, SVMs, Random Forests, etc

Operating Systems: Linux, Windows, macOS

RESEARCH EXPERIENCE

Postdoctoral Scholar & Fellow

University of Chicago

Aug. 2022 - Present

• Working with Professors Matthew Stephens and Xin He

• Developing Mendelian Randomization algorithms for robust inference of causal effects of multiple exposures on a single outcome

PhD Student

University of California, Los Angeles

September 2016 - June 2022

- Worked with Professors Wei Wang, Eleazar Eskin, and Harold Pimentel
- Developed multiple Mendelian Randomization methods providing sensitivity analysis and robustness to confounding from population structure
- Developed a method for identifying causal variants (statistical fine mapping) by leveraging information from multiple studies using a Bayesian approach
- Contributed to the effort to develop & deploy a saliva-based COVID-19 test ("Swab-Seq") that is now deployed campus-wide at UCLA
- Developed a method for isoform-aware eQTL mapping
- Developed multiple published methods that perform accurate abundance profiling of microbial organisms based on metagenomic reads
- Developed a method that uses deep learning to improve long sequencing read quality, in collaboration with scientists from the Joint Genome Institute
- Two-time teaching assistant (courses: Computational Genetics and Machine Learning in Genetics), given an average rating of 8 out of 9 by my students
- Mentored four undergraduate summer students, two of whom are now PhD students

Graduate Research Assistant (GRA) / Student Researcher
George Mason University January - August 2016 (June-August as GRA)

- Worked with Professor Huzefa Rangwala
- Developed CAMIL, a pipeline that uses multiple instance learning techniques based on whole metagenome shotgun sequence reads to predict whether or not a patient has a disease.
- CAMIL paper accepted into IEEE BIBM 2016 (19% acceptance rate).
- Second author of paper on using canopy clustering and locality sensitive hashing to reduce clustering time for biological datasets.

Predicting Clinical Phenotype using OTU-based Metagenome Representation George Mason University January 2015 - November 2015

- Worked with Professor Huzefa Rangwala
- Developed a computational pipeline that uses clustering and classification methods to quickly and accurately predict whether a patient has a disease based on a case/control metagenomic dataset
- Paper accepted for publication, and poster presentation won Outstanding Undergraduate Project Award (see publications and presentations sections)

TEACHING EXPERIENCE

Graduate Teaching Assistant

Spring 2020

 $\operatorname{CS/BIOINFO}$ 122/222 - Algorithms in Bioinformatics, Computer Science Department, UCLA

Graduate Teaching Assistant

Winter 2020

CS/BIOINFO 124/224 - Machine Learning in Genomics, Computer Science Department, UCLA

Undergraduate Teaching Assistant

Fall 2014 - Spring 2015

CS 306 - Computer Law and Ethics, Computer Science Department, George Mason University

- Received Outstanding Undergraduate Teaching Assistant Award
- Assisted students with legal research, writing, and oral communication

• Responsible for grading student assignments worth 25% of their grade; one of the only Undergraduate Teaching Assistants entrusted with this responsibility

INTERNSHIPS

Graduate Student Researcher

Summer 2017

DOE Joint Genome Institute / Lawrence Berkelely National Lab

- Developed a method that uses a Convolutional Neural Network (Deep learning method) to improve Oxford Nanopore long read quality de novo
- Method improves both read accuracy and downstream de novo assembly
- Helped pioneer use of deep learning within the lab

Security Engineering Intern

Summers 2013-2015

Sony Corporation of America

- Wrote secure and scalable software and worked with Big Data in order to help analyze, detect, and prevent attacks on Sony's networks
- Used Python, Javascript (Node, Express, Meteor), MongoDB, X/HTML, CSS
- Developed a workplace communications system using Meteor.js; solo project
- Assisted in the development of a web application based on a searchable database system using Node.js, Express.js, and MongoDB
- Assisted in the development of a network forensics system; wrote backend python scripts and XML web layouts

AWARDS, FELLOWSHIPS, SCHOLARSHIPS

Selected Honors

- Honorable Mention in NSF Graduate Research Fellowship, 2015-16
- Outstanding Undergraduate Student Award, given to overall best undergraduate student in Computer Science at George Mason University, May 2016
- Outstanding Academic Achievement Award, given for outstanding performance in Computer Science, May 2016
- Outstanding Undergraduate Research Project for presentation of my research project at Volgenau School of Engineering Undergraduate Research Symposium, April 2015
- Outstanding Undergraduate Teaching Assistant for two semesters of excellence as a teaching assistant, April 2015
- Dean's List every semester

Fellowships and Merit-Based Scholarships

- ModEling and uNdersTanding human behaviOR (MENTOR) NSF Training Grant, 2018-19 academic year at UCLA
- NIH T32 Doctoral Training Fellowship, 2016-18 academic years at UCLA
- Bersoff Endowed Scholarship, Outstanding Academic Achievement, Awarded twice in 2015 and 2014
- SWIFT Scholarship, Outstanding Academic Achievement, Awarded in 2015

Honors Societies and Organizations

- Honors College at George Mason University
- Alpha Lambda Delta Honor Society
- Golden Key International Honour Society

Technical Competitions

- Top 5 at VTHacks, a software development competition at Virginia Tech with over 45 teams, April 2014
- 2nd Place in the Technical Innovation Challenge, a week-long competition at George Mason University to design a viable software product, jointly refereed by Computer Science and Business Departments, October 2014

VOLUNTEER EXPERIENCE

Executive Curriculum Planner and Mentor, Community Programming Initiative George Mason University Honors College and SRCT 2014 - 2015

- Volunteer effort to teach basic programming to local 5th-8th grade students
- Planned and developed parts of the curriculum for the sessions, such as designing games and hints to help the students create them
- Mentored the elementary and middle school students during the sessions

MEMBERSHIPS AND LEADERSHIP

Student-Run Computing and Technology (SRCT)

2013 - 2016

- Student organization at George Mason University that works on software projects and competitions to benefit the university and broader local community
- Secretary and Member of Executive Board, Fall 2015 Semester