Cole Williams

Curriculum Vitae

Center for Computational Molecular Biology Brown University

✓ cole.w@me.com

Education

Ph.D. in Computational Biology

Expected 2025

Brown University

• Advisor: Dr. Sohini Ramachandran

• Committee: Dr. Emilia Huerta-Sanchez, Dr. Dan Weinreich, Dr. Amy Williams

M.S. in Computational Biology

2024

Brown University

• Earned during Ph.D. studies

• GPA: 4.00/4.00

B.S. in Genetics & Genomics

2019

University of California, Davis

• GPA: 3.99/4.00

• Honors Thesis: "Inferring genetic relatedness in an endogamous African population"

Research Experience

Graduate Research Fellow

2020-Present

Ramachandran Lab, Brown University

- Developed HAPTIC, a novel inter-chromosomal phasing algorithm using spectral clustering of identity-by-descent (IBD) segments, successfully scaling to analyze 500,000+ UK Biobank samples and validated on 23andMe's dataset.
- Engineered and optimized high-performance computing code for large-scale genetic relatedness analysis, enabling efficient processing of genome-wide IBD segments across millions of sample pairs.
- Developed PONDEROSA, a machine learning algorithm for predicting genealogical relationships from IBD segments, with ongoing development of CNN-based approaches for improved accuracy.
- Led research on genetic privacy implications, resulting in forthcoming publication on policy recommendations and best practices.

Professional Research Assistant

2019-2020

Gignoux Lab, University of Colorado, Anschutz Medical Campus

- Developed clustering methods to identify genetic relatives in the San Antonio Mexican American Families Study (SAMAFS).
- Assisted with biobank team to implement relatedness detection algorithms for the Colorado Center for Personalized Medicine biobank.

Undergraduate Researcher

2018-2019

Henn Lab, University of California, Davis

- Investigated genetic relatedness in the Himba population.
- Began development of PONDEROSA, for inferring genealogical relationships using IBD data.

Publications

Journal Articles

- 1. Moody, N.M., **Williams, C.M.**, Ramachandran, S., & Fuxjager, M.J. Social mates dynamically coordinate aggressive behavior to produce flexible territorial strategies. *In revision at PLOS Computational Biology*.
- 2. Williams, C.M., O'Connell, J., Freyman, W.A., 23andMe Research Team, Gignoux, C.R., Ramachandran, S., & Williams, A.L. Phasing millions of samples achieves near perfect accuracy, enabling parent-of-origin classification of variants. *bioRxiv*, May 2024.
- 3. Williams, C.M., Scelza, B.A., Slack, S.D., Mathias, R.A., Watson, H., Barnes, K.C., Lange, E., Johnson, R.K., Gignoux, C.R., Ramachandran, S., & Henn, B.M. A rapid, accurate approach to inferring pedigrees in endogamous populations. *bioRxiv*, July 2024.
- 4. Wiley, L.K., Shortt, J.A., Roberts, E.R., Lowery, J., Kudron, E., Lin, M., Mayer, D., Wilson, M., Brunetti, T.M., Chavan, S., Phang, T.L., Pozdeyev, N., Lesny, J., Wicks, S.J., Moore, E.T., Morgenstern, J., Roth, A.N., Shapowitz, E.L., Stewart, A., Williams, C.M., [...], Barnes, K.C., & Gignoux, C.R. Building a vertically integrated genomic learning health system: The biobank at the Colorado Center for Personalized Medicine. The American Journal of Human Genetics, 111(1):11–23, January 2024.
- Swinford, N.A., Prall, S.P., Gopalan, S., Williams, C.M., Sheehama, J., Scelza, B.A., & Henn, B.M. Increased homozygosity due to endogamy results in fitness consequences in a human population. *Proceedings of the National Academy of Sciences*, 120(43):e2309552120, October 2023.
- 6. Gopalan, S., Berl, R.E.W., Myrick, J.W., Garfield, Z.H., Reynolds, A.W., Bafens, B.K., Belbin, G., Mastoras, M., **Williams, C.**, Daya, M., Negash, A.N., Feldman, M.W., Hewlett, B.S., & Henn, B.M. Hunter-gatherer genomes reveal diverse demographic trajectories during the rise of farming in Eastern Africa. *Current Biology*, 32(8):1852–1860.e5, April 2022.

Conference Abstracts & Presentations

- 1. Williams, C.M., O'Connell, J., 23andMe Research Team, Gignoux, C.R., Ramachandran, S., & Williams, A.L. Phasing of millions of samples achieves near perfect accuracy, enabling parent-of-origin classification of variants. Poster presentation at the American Society for Human Genetics (ASHG) annual meeting. Washington, D.C. November 2023.
- 2. Williams, C.M., 23andMe Research Team, Gignoux, C.R., Ramachandran, S., & Williams, A.L. Parent of origin classification of identity by descent segments. Poster presentation at the Probabilistic Modeling in Genomics meeting. Cold Spring Harbor, NY. March 2023.
- 3. Williams, C.M., Scelza, B.A., Shortt, J.A., Lange, E., Rafaels, N., Crooks, K., Barnes, K.C., Gignoux, C.R., & Henn, B.M. A rapid, accurate approach to inferring pedigrees in endogamous populations. Poster presentation at the American Society for Human Genetics (ASHG) annual meeting. Houston, TX. October 2019.
- 4. Williams, C.M., Scelza, B.A., & Henn, B.M. A rapid, accurate approach to inferring pedigrees in endogamous populations. Talk presented at the Bay Area Population Genomics meeting. Palo Alto, CA. June 2019.

Technical Skills

Programming

• Advanced: Python, Bash

• Intermediate: R

• Basic: C++

Tools & Technologies

• Version Control: Git/GitHub

• Development Tools: Apptainer/Singularity, VS Code, Jupyter Notebooks

• Workflow management: Nextflow, Snakemake

Data Analysis

• Dimensionality reduction: PCA, LDA

• Machine Learning: scikit-learn, TensorFlow

• Visualization: seaborn, matplotlib

Leadership & Service

Academic Service

- Graduate Teaching Assistant (BIOL495: Statistical Analysis of Biological Data)
- Mentorship of Data Science Master's student

Professional Service

• Co-founder: Science on the Hill outreach program