## Cole Williams

### Curriculum Vitae

Center for Computational Molecular Biology Brown University

✓ cole.w@me.com

## Education

#### Ph.D. in Computational Biology

Expected August 2025

Brown University

- Thesis: Population-Genetic Methods and Insights Across the Spectrum of Relatedness
  - Defended: April 2025
- Advisor: Dr. Sohini Ramachandran
- Committee: Dr. Emilia Huerta-Sanchez, Dr. Daniel Weinreich, Dr. Amy Williams
- National Science Foundation Graduate Research Fellow

## 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

- Thesis: Inferring Genetic Relatedness in an Endogamous African Population
- GPA: 3.99/4.00

# 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. 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. Preprint: 10.1101/2020.02. 25.965376. Accepted at *GENETICS*.
- 2. Williams, C.M., O'Connell, J., Freyman, W.A., 23andMe Research Team, Gignoux, C.R., Ramachandran, S.<sup>†</sup>, & Williams, A.L.<sup>†</sup> Phasing millions of samples achieves near perfect accuracy, enabling parent-of-origin classification of variants. Preprint: 10.1101/2024.05.06. 592816. In revision at *HGG Advances*.
- 3. Moody, N.M., Williams, C.M., Ramachandran, S., & Fuxjager, M.J. Social mates dynamically coordinate aggressive behavior to produce strategic territorial defense. *PLOS Computational Biology*, 21(1):e1012740, January 2025. doi: 10.1371/journal.pcbi.1012740.
- 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.L., 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. doi: 10.1016/j.ajhg.2023.12.001.
- 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. doi: 10.1073/pnas.2309552120.

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

<sup>&</sup>lt;sup>†</sup> Denotes equal senior author contribution

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. doi: 10.1016/j.cub.2022.020.050.

## Manuscript Preprints

1. Ramanan, V.\*, Vinod, R.\*, **Williams, C.\***, Ramachandran, S.<sup>†</sup>, Venkatasubramanian, S.<sup>†</sup> Genetic Data Governance in Crisis: Policy Recommendations for Safeguarding Privacy and Preventing Discrimination. *arXiv* 10.48550/arXiv.2502.09716. February 2025.

#### Conferences

- 1. Ramanan V.\*, Vinod R.\*, **Williams C.\***, Ramachandran S.<sup>†</sup>, Venkatasubramanian S.<sup>†</sup> Genetic Data Governance in Crisis: Policy Recommendations for AI Applications. Poster presentation at Sociotechnical AI Governance workshop. Yokohama, Japan. April 2025. [Paper]
- 2. 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.
- 3. 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.
- 4. 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.
- 5. 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

## Awards & Fellowships

• UC Davis College of Biological Sciences Undergraduate of the Year (2019)

- NIH T32 Predoctoral Training Grant (2021-2022)
- NSF Graduate Research Fellowship (2022-2025)

# Leadership & Service

#### Academic Service

- Graduate Teaching Assistant (BIOL495: Statistical Analysis of Biological Data)
- Guest Lecturer in BIOL495 (Statistical Analysis of Biological Data): Spring 2023 & Spring 2025

#### **Professional Service**

• Co-founder: Science on the Hill outreach program