IAN S. ARRIAGA MACKENZIE

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EDUCATION

MS	University of Colorado Denver, Statistics	August 2021
BS	University of Colorado Denver, Mathematics	May 2020

INTERESTS

My recent work has been in statistical genetics and ancestry estimation, including, but not limited to, optimization, cluster computing, algorithm design, and efficient experiment implementation.

I enjoy exploring the intersection between big data and computing, particularly in delivering scalable and adaptable solutions to current research questions.

RESEARCH

University of Colorado Anschutz, Katerina Kechris, PhD 2021 to Present Center for Innovative Design & Analysis Andrew Monte, MD, PhD, Jack Pattee, PhD • Gene and variant association testing for liver damage with acetaminophen use, assessing copy number variants within genome. • Genomic data pre-processing, single and multi-variant models, checking CNVs, analysis of results. University of Colorado Denver, Audrey Hendricks, PhD 2019 to 2021 Hendricks Research • Estimation of ancestry in summary genetic data and adjusting allele frequency data to match ancestry of target population • Genomic data cleaning and merging, sequential quadratic programming, cluster computing and algorithm optimization, Shiny Apps to interactively visualize data, fixation index between populations University of Colorado Denver, Audrey Hendricks, PhD 2020 Women First Project Sarah Borengasser, PhD, Chris Gignoux, PhD, Nancy Krebs, MD, Michael Hambidge, MD • Genome wide association study (GWAS) on birth height in Guatemalan population

• Genomic data pre-processing, correction for kinship (relatedness), linear model with multiple covariates, analysis of results

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University of Colorado Anschutz Medical Campus, Greg Kinney, PhD

MS Consulting Course Project

- Cannabis sleep study to assess differences between intoxicated versus unintoxicated subjects in stage N2 sleep
- Electroencephalogram (EEG) analysis, applying Fast Fourier Transforms, signal processing and cleaning, ANOVA

PUBLICATIONS

Arriaga-MacKenzie IS, Matesi G, Chen S, Ronco A, Marker KM, Hall JR, Scherenberg R, Khajeh-Sharafabadi M, Wu Y, Gignoux CR, Null M, Hendricks AE. Summix: A method for detecting and adjusting for population structure in genetic summary data. *American Journal of Human Genetics* **2021** 108, 1270-1282

Monte AA, **Arriaga-MacKenzie IS**, Pattee J, Kaiser S, Willems E, Rumack B, Reynolds K, Dart RC, Heard KJ. Genetic Variants Associated with ALT Elevation from Therapeutic Acetaminophen. (In Progress, 2022)

Borengasser SJ, Murphy J, Null M, Jambal P, Jones KL, Yang IV, Friedman JE, Waldrop SW, Gilley SP, Saint-Cyr M, **Arriaga-MacKenzie IS**, Kemp J, Westcott J, Garces A, Figueroa L, Greally JM, Hambidge MK, Hendricks AE, Krebs NF. Role of a Preconception Maternal Nutrition Supplement and Pre-pregnancy BMI on Infant DNA Methylation at Birth in Guatemalan Mother-Infant Dyads: The Women First Trial. (In Progress, 2022)

TEACHING

University of Colorado Denver Learning Assistant, Statistical Theory 3382	Fall 2019
• Developed learning activities, assisted students in theory and in	
University of Colorado Denver Grader, Pre-Calculus	Spring/Fall 2019
AWARDS	
CLAS Outstanding MS Graduate Award University of Colorado Denver	2021
C.W. Cotterman Award American Journal of Human Genetics	2021
Undergraduate Research Opportunity Program (UROP) Grant University of Colorado Denver	2019/2020
Education Through Undergraduate Research and Creative Activities	2019

2019

Work Study Award University of Colorado Denver

PRESENTATIONS

Presentations

SUM-MIX: A method for detecting and adjusting for population structure in genetic summary data. Speed Talk Joint Statistical Meetings, Virtual	August 2021	
SUM-MIX: A method for detecting and adjusting for population structure in genetic summary data. The Western North American Region of the International Biometric Soc	June 2021 ciety, Virtual	
Poster Presentations		
SUM-MIX: A Method to Estimate and Adjust for Ancestry in Genetic Summary Data American Society of Human Genetics, Virtual	October 2020	
<i>Estimation of non-Reference Ancestry Proportions in Genotype</i> <i>Frequency Data,</i> Selected for Lighting Presentation International Genetic Epidemiology Society, Virtual	July 2020	
<i>Efficient Estimation of Hidden Ancestry Substructure in Summary</i> <i>Genotype Frequency Data</i> Research and Creative Activities Symposium, Virtual	May 2020	
Estimation of Hidden Ancestry Substructure Using a Least Squares Mixture Model with Summary Genotype Frequency Data	November 2019	
Society for the Advancement of Chicanos/Hispanics and Native Americans in Science, Honolulu HI		
<i>Efficient Estimation of Hidden Ancestry Structure Using Summary</i> <i>Genotype Frequency Data,</i> Selected for Lighting Presentation International Genetic Epidemiology Society, Houston TX	October 2019	
<i>Estimating Hidden Ancestries in Real Data</i> Research and Creative Activities Symposium, Denver CO	April 2019	

PROGRAMMING

Python, R, SQL, Tableau, Bash, git, C++, Matlab, SAS, Julia, Jupyter, Tensorflow, Pytorch, Docker, bedtools, Shiny Apps, Samtools, VCFtools, PLINK, ADMIXTURE, Galaxy, Ingenuity Pathway Analysis, PennCNV

PROFESSIONAL AFFILIATIONS

American Statistical Association (ASA), *Student member* American Society of Human Genetics (ASHG), *Student member* International Genetic and Epidemiology Society, *Student member* The Western North American Region of The International Biometric Society, *Student member* Society for the Advancement of Chicanos/Hispanics and Native Americans in Science, *Student member*

REFERENCES

Audrey E. Hendricks, PhD, Associate Professor Mathematical and Statistical Sciences University of Colorado Denver

P.O. Box 173364, Campus Box 170, Denver, CO 80217-3364 Email: audrey.hendricks@ucdenver.edu

Stephanie A. Santorico, PhD, Professor

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Erin E. Austin, PhD, Assistant Professor

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