

HEALTH SCIENTIST - INFORMATICS

Respiratory Diseases Laboratory Branch, Centers for Disease Control and Prevention, Atlanta, GA, USA

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I am a scientist with 10+ years of experience in classical genetics, genomics, and evolutionary biology. My training includes using a range of tools - bioinformatics, population genetics, molecular evolution all with experimental design in mind. Over the years as my computational skills increased, the genomes of the organisms I work on have gotten smaller. Now, I work at the interface of public health, application development, and understanding organisms that can cause harm to humans with an empahsis of developing reproduciable research.

Education

University of Georgia Athens, GA

Ph.D. IN GENETICS 2015

University of North Carolina - Asheville

Asheville, NC

B.S. IN BIOLOGY

2010

Appointments

Centers for Disease Control and Prevention

Atlanta, GA

HEALTH SCIENTIST INFORMATICS

June 2022 - Current

November 2020 - June 2022

- * General informatics for Legionella
 - Continued projects from previous position with ASRT, inc.

Centers for Disease Control and Prevention, Contractor with ASRT Inc.

Atlanta, GA

BIOINFORMATICIAN II WITH DIVISION OF BACTERIAL DISEASES

- * Interfacing with state public health laboratories (PHL)
 - Communicated with members of state PHL to provide guidenance regarding bionformatics and genomics for Legionella species
- * Refactored Legionella Species ID tool for broader compatibility and interfacing with Nextflow Tower
 - Removed structural design issues in original code that prevented expansion and adding of new functionalities
- * Development of method validation plan for Legionella Species ID tool
 - Researched and proposed statistically sound validation plan to test tool that uses next generation sequencing data
- * Phylogenetic and phylodynamic analyses for Legionella isolates of public health concern
 - Performing analyses and generating statistical summaries and visualizations
 - Writing and editing manuscripts for publication
 - Identifying scientific questions answerable with existing data
- * Developing or testing scripts and pipelines as necessary
 - Generated script to interact with National Center of Biotechnology Information for downloading all available Legionella genomes
 - Evaluated using Nextflow pipeline for mapping Legionella genomes
- * 60-day detail on Data Analytics and Visualization for Vaccine Task Force (April June 2021)
 - Produced weekly reports and consolidated steps for document creation to be reproducible via one script

Centers for Disease Control and Prevention, Fellow with APHL

Atlanta, GA

ASSOCIATION OF PUBLIC HEALTH LABORATORIES (APHL) BIOINFORMATICS FELLOW

September 2019 - November 2020

- * Developed tinselR an Rshiny application for annotating outbreak trees
 - tinselR is an open-source downloadable package in the R language used for phylogenetic tree analysis and visualization
- * Participated in APHL core competency and leadership training
 - Gained skills and knowledge for a career in public health
- * 30-day detail on Routine Reporting for COVID-19 Response (June July 2020)
 - Produced daily and weekly reports for task force. Consolidated steps for document creation to be reproducible via one script

University of Georgia (UGA)

Athens, GA

October 2017 - August 2019

Bensasson Lab Post-Doctoral Researcher

* Directed collaborative research with UGA graduate and undergraduate students

- Directed Collaborative research with OOA graduate and undergraduate students
- * Collated and streamlined genome mapping for 1000 Saccharomyces cerevisiae isolates as lab resource
- * Used iterative approach to assign 400 genomes to well-defined populations using best practices for population genomics and phylogenetics
- * Resolved haplotypes for Candida albicans genomes with PacBio sequencing

Indiana University (IU) Bloomington, IN

MOYLE LAB POST-DOCTORAL RESEARCHER

June 2015 - September 2017

2017

- * Directed research with faculty, graduate, and undergraduates at IU and with two external universities
- * Genome-wide study of factors that determine the likelihood of introgression for Solanum species
- * Streamlined genotyping to generate biological replicates. Validate data accuracy using technical replicates along with statistical analyses
- * Identified targeting rates and floral trait differences with biological & technical replicates in large-scale crossing project via statistical analyses

Skills

Analytical

DATA SCIENCE, EXPERIMENTAL DESIGN, NEXT GENERATION SEQUENCING, GENOMICS, HIGH PERFORMANCE COMPUTING, PROJECT MANAGEMENT, QUALITY CONTROL, REPRODUCIBLE RESEARCH, STATISTICS, TECHNICAL WRITING AND

DOCUMENTATION

Programming

AWK - PROFICIENT, BASH - EXPERT, PYTHON - EXPERT, R - EXPERT, RMARKDOWN - EXPERT, NEXTFLOW - PROFICIENT, GITHUB -

Publications

JAP Hamlin, T Nakov, A Williams-Newkirk	Microbiology Resource
	Announcements
TINSELR—AN R SHINY APPLICATION FOR ANNOTATING PHYLOGENETIC TREES	2021

JAP Hamlin, MS Hibbins, LC Moyle **Evolution Letters** Assessing biological factors affecting postspeciation introgression

JAP Hamlin, GB Dias, CM Bergman, D Bensasson G3PHASED DIPLOID GENOME ASSEMBLIES FOR THREE STRAINS OF CANDIDA ALBICANS FROM OAK TREES 2019

JAP Hamlin, NA Sherman, LC Moyle G3

TWO LOCI CONTRIBUTE EPISTASTICALLY TO HETEROSPECIFIC POLLEN REJECTION, A POSTMATING ISOLATING BARRIER BETWEEN

SPECIES

Biological Journal of the Linnean JAP Hamlin, TJ Simmonds, ML Arnold Society

NICHE CONSERVATISM FOR ECOLOGICAL PREFERENCE IN THE LOUISIANA IRIS SPECIES COMPLEX

JAP Hamlin, ML Arnold Journal of Heredity NEUTRAL AND SELECTIVE PROCESSES DRIVE POPULATION DIFFERENTIATION FOR IRIS HEXAGONA

Reticulate Evolution: ML Arnold, AN Brothers, JAP Hamlin, SJ Taylor, NH Martin Symbiogenesis, Lateral Gene

Transfer, Hybridization ... DIVERGENCE-WITH-GENE-FLOW—WHAT HUMANS AND OTHER MAMMALS GOT UP TO 2015

JAP Hamlin, ML Arnold Ecology and Evolution

DETERMINING POPULATION STRUCTURE AND HYBRIDIZATION FOR TWO IRIS SPECIES.

Rapidly evolving genes and genetic ML Arnold, JAP Hamlin, AN Brothers, ES Ballerini, RS Singh, J Xu, ...

systems

ML Arnold, ES Ballerini, AN Brothers, JAP Hamlin, CDA Ishibashi, ...

Plant Ecology & Diversity 2012

THE GENOMICS OF NATURAL SELECTION AND ADAPTATION: CHRISTMAS PAST, PRESENT AND FUTURE (?).

NATURAL HYBRIDIZATION AS A CATALYST OF RAPID EVOLUTIONARY CHANGE

Presentations

Talks

mashWrapper Atlanta, GA

GENOMICS INTEREST GROUP - TECHINCAL 2022

tinselR - An RShiny App for Phylogenetic Tree Annotation	Atlanta, GA
CDC DATA VIZ DAY	2020
Start Writing Manuscripts in Rmarkdown	Atlanta, GA
GENOMICS INTEREST GROUP - TECHINCAL	2020
Build Shiny Apps Warnell School of Forestry & Natural Resources Data Manipulation and Management course	Athens, GA 2018
Population Genomics of wild and domesticated yeast University of Georgia Enthusiasts of Diversity Genetics and Evolution	Athens, GA 2018
No Easy D: Little Evidence for Introgression from Whole-Genome Data in Wild Solanum	Portland, OR
Society for the Study of Evolution	2017
Epistasis for Postmating Prezygotic Isolation	Austin, TX
SOCIETY FOR THE STUDY OF EVOLUTION	2016
Spatial, temporal, and genome-wide patterns of differentiation for the Louisiana iris species complex	Bloomington, IN
Indiana Unviversity Brown Bag	2015
What Drives Phenotypic and Genetic Divergence for Iris hexagona	Raleigh, NC
SOCIETY FOR THE STUDY OF EVOLUTION	2014
Comparative Phylogeography of Two Louisiana Irises	Snowbird, UT
SOCIETY FOR THE STUDY OF EVOLUTION	2013
Posters	
A Community Outbreak of Legionnaires' Disease Possibly Associated with an Outdoor Hotel Spa – California, 2021	Salt Lake City, Utah
Council of State and Territorial Epidemiologists	2023
 Burmester J, Evans S, Ferguson D, Lunsforward R, Rutschmann S, Kimura A, Flores E, Mukhopadhyay R, Hanna Miller, G 	
Combining Mash and NCBI Datasets via Nextflow for a Legionella Species Identification	Atlanta, GA
Tool CDC Laboratory Science Symposium	2023
Hamlin JAP, Willby MJ, and Winchell JM	2025
Spread of Sequence Type ST222 from the Great Lakes Area to the US West Coast and	
Europe	Atlanta, GA
CDC Laboratory Science Symposium	2023
Kozak-Muiznieks NA, Hamlin JAP, Lucas C, and Willby MJ	
Combining Mash and NCBI Datasets via Nextflow for a Legionella Species Identification	Baltimore, MD
Tool American Society for Microbiology	2022
Hamlin JAP, Willby MJ, and Winchell JM	2022
Spread of Sequence Type ST222 from the Great Lakes Area to the US West Coast and	
Europe	Yokohama, Japan
International Conference on Legionella	2022
Kozak-Muiznieks NA, Hamlin JAP, Lucas C, and Willby MJ	
Tinsel - an Rshiny app for phylogenetic tree annotation	Virtual
ASSOCIATION OF PUBLIC HEALTH LABORATORIES ANNUAL MEETING	2020
Hamlin JAP and Williams-Newkirk A	
Tinsel - an Rshiny application for annotating outbreak trees Integrated Foodborne Outbreak Response and Management	Canceled - Atlanta, GA
INTEGRATED FOODBORNE OUTBREAK RESPONSE AND MANAGEMENT	2020
Hamlin JAP and Williams-Newkirk A	
	Pacific Grove CA
 Hamlin JAP and Williams-Newkirk A Genetic admixture and the origins of clinical Saccharomyces cerevisiae yeast Fungal Genetics Conference 	Pacific Grove, CA 2019

Clinical Saccharomyces cerevisiae are admixed domesticated yeast Holderness, NH CELLULAR AND MOLECULAR FUNGAL BIOLOGY: GORDON RESEARCH CONFERENCE 2018 • Hamlin JAP and Bensasson D Higher Prevalence of Genetic Exchange Given Geographic Proximity Laural Hill, NC SOUTHEASTERN POPULATION ECOLOGY & EVOLUTIONARY GENETICS. 2017 Hamlin JAP and Movle I C. Two Loci Contribute Epistastically to Heterospecific Pollen Tube Rejection, a Postmating Davis, CA **Isolating Barrier Between Species** SOLANACEAE CONFERENCE 2016 · Hamlin JAP and Moyle LC What Drives Phenotypic and Genetic Divergence for Iris hexagona Boise, ID **BOTANICAL SOCIETY OF AMERICA** 2014 · Hamlin JAP and Arnold ML Comparative Phylogeography of Two Louisiana Iris Species: Iris fulva and Iris brevicaulis Guanajuato, Mexico AMERICAN GENETIC ASSOCIATION • Hamlin JAP, Brothers AN, Ballerini ES, and Arnold ML Packages ____ tinselR - An R package for visulizing and annotating phylogenetic trees https://github.com/jennahamlin/tinselR AUTHOR AND MAINTAINER 2019 mashwrapper - a Python tool for parsing mash results https://github.com/jennahamlin/mashwrapper AUTHOR AND MAINTAINER **Mentoring University of Georgia graduate students** E. Scopel Ferreira Da Costa J. Pena **University of Georgia undergraduate students** T. Simmonds K. Miller J. Foley S. Duque B. Webb **Indiana University graduate students** C. Jewell M. Gibson **Indiana University undergraduate students** A. Huh J. Breisch

S. Henderson L. Howser C. Plasterer D. Shukla S. Seo