ARIELLA GLADSTEIN

 $(202) \cdot 258 \cdot 4408 \diamond algladstein@gmail.com \diamond github.com/agladstein$

Expertise: Human population genetics, genomics, computational biology, bioinformatics, statistics

EDUCATION

University of Arizona, Tucson, AZ

August 2018

PhD in Ecology and Evolutionary Biology

Minor in Mathematics

Beloit College, Beloit, WI

May 2011

B.S. in Mathematical Biology & Russian, Cum Laude

Departmental Honors: Mathematical Biology

RESEARCH EXPERIENCE

Postdoctoral Fellowship

2018 - present

Schrider Lab Department of Genetics, University of North Carolina, Chapel Hill

Project: Inference of demographic history with deep learning

Dissertation Research

2011 - 2018

Hammer Lab Ecology and Evolutionary Biology, University of Arizona Dissertation: Inference of recent demographic history of population isolates using genome-wide high density SNP arrays and whole genome sequences

Functional Genomics Research Rotation

Spring 2012

Restifo Lab Neuroscience, University of Arizona

Computational Genomics Research Rotation

Fall 2011

Kececioglu Lab Computer Science, University of Arizona

Lab technician intern

Sp, Fall 2009

Laboratory of Population Genetics Russian Academy of Medical Sciences, Moscow, Russia

SKILLS

Programming	Python Roch R Por	l (including unit testing and	profiling)
i rogramming	I y thon, Dasn, It, I cr.	i (including and testing and	proming

Other Computer Linux command line, Git, HPC/HTC (with HTCondor, PBS, Slurm),

cloud computing (OpenStack, AWS)

Genomics PLINK, Bedtools, VCFtools, phasing, imputation, haplotype detection,

population structure, genome simulation

Data Science Numpy, Scikit-learn, Keras, Pandas, Spark, Tidyverse, Matplotlib, ggplot

Reproducibility scripting, workflow development (Pegasus, Makeflow), containers

(Singularity, Docker), Jupyter, Knitr

Documentation LATEX, Markdown, reStructuredText

Language Skills English (native speaker), Russian (fluent), Spanish (basic)

AWARDS

XSEDE Startup allocation (100,000 CPU hrs, 6,500 GPU hrs, 5.5 Tb storage)

2019

XSEDE Supplemental allocation (200,000 CPU hrs)

2018

XSEDE Startup allocation (150,000 CPU hrs)

2017

Open Science Grid User School (travel, lodging, meals, cost of program)

2017

GPSC Travel Grant (\$761)

2015, 2016, 2017

University of Arizona Galileo Circle Scholarship (\$1,000)

2015

NIH Computational and Mathematical Modeling of Biological Systems Traineeship (\$71,064) 2013-2014

NSF Integrative Graduate Education and Research Traineeship in Genomics (\$97,083) 2011-2013

Society for Learning Unlimited Grant (\$2,000)

Fall 2009

Study Abroad Enhancement Grant, Beloit College (\$250)

Spring 2009

Beloit College Presidential Scholar (\$60,000)

2007-2011

PUBLICATIONS

Gladstein A.L. and Hammer M.F. 2019. Substructured population growth in the Ashkenazi Jews inferred with Approximate Bayesian Computation. *Molecular Biology and Evolution*, msz047; doi: https://dx.doi.org/10.1093/molbev/msz047

Gladstein A.L. et al. 2018. SimPrily: A Python framework to simplify high-throughput genomic simulations. SoftwareX, 7, 335340. https://doi.org/10.1016/j.softx.2018.09.003

Gladstein A. and Hammer M.F. 2016. Population Genetics of the Ashkenazim. In: eLS. John Wiley & Sons, Ltd: Chichester. pp. 1-8. https://doi.org/10.1002/9780470015902.a0020818.pub2

Behar et al. 2013. No evidence from genome-wide data of a Khazar origin for the Ashkenazi Jews. *Human Biology.* 85.6:859-900. https://doi.org/10.3378/027.085.0604

Gladstein A.L. 2011. Split decomposition analysis groups Jewish populations together between European and Middle Eastern populations. *The Beloit Biologist*. 30:29-36.

POSTERS AND PRESENTATIONS

Invited Talks

Inference of evolutionary history with Approximate Bayesian Computation. Open Science Grid All-Hands Meeting. 2018. Salt Lake City, Utah.

Posters

Gladstein A.L., Hammer M.F. Substructured population growth in the Ashkenazi Jews inferred with Approximate Bayesian Computation. Probabilistic Modeling in Genomics. 2018. Cold Spring Harbor, NY.

Gladstein A.L., et al. Efficient pipeline for whole genome simulation and summary statistic calculation with flexible demographic models. Meeting of the American Society for Human Genetics. 2017. Orlando, FL.

Gladstein A.L., et al. The effect of SNP array ascertainment bias on the distribution of runs of homozygosity lengths. Annual Meeting of the American Society for Human Genetics. 2015. Baltimore, MD.

PROJECT MANAGEMENT

Managed team of 5, including computer scientists, software engineer, and mathematician, working on code development and high throughput computing for bioinformatics

WORKSHOPS AND HACKATHONS

NCBI RNA-Seq in the Cloud hackathon (3 days)	03/2019	
Cyber Carpentry (2 weeks)	06/2018	
XSEDE HPC Workshop: Big Data (2 days)	02/2018	
Open Science Grid User School (1 week)	07/2017	
COMMUNITY SERVICE AND OUTREACH		
Poster presentation at UNC Women in Computing Research Symposium	2019	
Helped with Research Bazaar workshop on R	2018	
Teaching assistant for CyVerse Container Camp	2018	
Active participant in the Research Bazaar	2017, 2018	
Helped with Software Carpentry on Unix/Bash, Python, and Git	2017	
Helped with the Tucson Womens Hackathon workshop on Git	2017	
Judge for Graduate & Professional Student Council Travel Grants	2012, 2016, 2017, 2018	
Judge for Tucson Magnet High School Science Fair	2012, 2015	
Taught Population Genetics Module at the Kino School	2013	
Arizona Assurance Mentor	2012	
Judge for EEB Undergraduate Research Poster Session	2012	
TEACHING EXPERIENCE		
ECOL 320 Genetics Graduate Teaching Assistant	Fall 2014, 2015, 2016 University of Arizona	
ECOL 182L Intro to Ecology and Evolutionary Biology Lab Graduate Teaching Assistant	Spring 2015 University of Arizona	
BIOL 247 Biometrics Teaching Assistant	Spring 2011 Beloit College	
OTHER ACTIVITIES		
Circus Arts Aerial silks, aerial rope, lyra, flying trapeze, tightwire, handbalancing, conto	2012 - present	
Figure Skating Singles freestyle	1998 - 2013	