ARIELLA GLADSTEIN, PhD

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https://ariella-gladstein.netlify.app | https://github.com/agladstein | www.linkedin.com/in/ariella-gladstein

Expertise: population genetics, genomics, computational biology, data science, machine learning

EDUCATION

University of Arizona, Tucson, AZ

PhD in Ecology and Evolutionary Biology

Minor in Mathematics

Beloit College, Beloit, WI

May 2011

August 2018

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

Departmental Honors: Mathematical Biology

EXPERIENCE

Research Scientist, Computational Biology

2021 - 2024

Ancestry team

Embark Veterinary, Remote

Developed dog breed ancestry and relative-matching products, enhancing computational efficiency while maintaining rigorous quality standards. Played a key role in improving pipeline automation from 40% to over 95% within one year and scaling processes from hundreds of thousands to millions of samples. Collaborated with a cross-functional team on a science initiative projected to save \$7.5 million annually. Served on the Diversity, Equity, Inclusion, and Belonging (DEIB) committee, co-authoring an Inclusive Norms document, designing and analyzing company-wide DEIB surveys, and contributing to various DEIB initiatives.

Postdoctoral Fellowship

2018 - 2020

Schrider Lab Department of Genetics, University of North Carolina, Chapel Hill Developed deep learning models for population genetics inference, utilizing high-throughput computing for simulations and high-performance GPUs, as well as a Kubernetes cluster, for model training. Contributed to writing documentation and tutorials for the population genomics tskit software ecosystem. Actively participated in the PopSim Consortium, contributing to the development of the stdpopsim library.

Dissertation Research 2011 - 2018

Hammer Lab Ecology and Evolutionary Biology, University of Arizona Developed and applied computational methods for population genetics analysis, addressing ascertainment bias in identifying runs of homozygosity and demographic inference. Created and released software for high-throughput genomic simulations. Applied Approximate Bayesian Computation (ABC) to model the demographic history of Ashkenazi Jews, utilizing over 11 million CPU hours in under six months to run large-scale simulations on the Open Science Grid and multiple HPC systems with a high-throughput workflow.

SKILLS

Software Development HPC/HTC and Cloud Reproducibility Python, R, Bash, unit testing, profiling, multiprocessing, Git Linux/Unix, SLURM, PBS, AWS, JupyterHub, Kubernetes Workflow development (Pegasus, Makeflow, Snakemake), containers

(Singularity, Docker), virtual environments, Jupyter, Knitr

Genomics PLINK, Bedtools, VCFtools, phasing, imputation, haploty

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

structure, genome simulation

Data Science Documentation Numpy, Pandas, Scikit-learn, SciPy, TensorFlow, Keras, Matplotlib, Seaborn, Plotly

IATEX, Markdown, reStructuredText, JupyterBook, Confluence

TEACHING EXPERIENCE

Introduction to using stdpopsim Workshop

2020

Designed and presented a workshop on using the stdpopsim library

Virtual, PopSim Consortium

NSF Cyber Carpentry Workshop

2019

Assisted with 2-week workshop, where participants learned best practices for data-intensive computing, cloud infrastructure, and deep learning through hands-on projects.

University of North Carolina, Chapel
Hill

Research Bazaar Workshop on R

Assisted with hands-on intro R workshop

University of Arizona

CyVerse Container Camp

2018

2018

Assisted with a 3-day workshop teaching participants to use containerized workflows and datasets for more reproducible science.

University of

Arizona

Software Carpentry Workshop

2017

Assisted with Software Carpentry workshop on basic Unix/Bash, Python, and Git University of Arizona

Tucson Womens Hackathon workshop on Git

2017

Assisted with workshop on basic Git

University of Arizona

Fall 2014, 2015, 2016

ECOL 320 Genetics Graduate Teaching Assistant

University of Arizona

ECOL 182L Intro to Ecology and Evolutionary Biology Lab

Spring 2015

 $Graduate\ Teaching\ Assistant$

University of Arizona

BIOL 247 Biometrics

Spring 2011

 $Teaching\ Assistant$

Beloit College

AWARDS

XSEDE Research allocation (250,000 CPU hrs)	2020
XSEDE Supplemental allocation (250,000 CPU hrs)	2018, 2020
XSEDE Startup allocation (100,000 CPU hrs, 6,500 GPU hrs, 5.5 Tb storage)	2019
XSEDE Startup allocation (150,000 CPU hrs)	2017
Open Science Grid User School (travel, lodging, meals, cost of program)	2017
NIH Computational and Mathematical Modeling of Biological Systems Traineeship (\$71,064)	2013-14
NSF Integrative Graduate Education and Research Traineeship in Genomics (\$97,083)	2011-13

WORKSHOPS AND HACKATHONS

SMBE satellite meeting on Speciation Genomics, Tjarno, Sweden (3 days)	06/2019
NCBI RNA-Seq in the Cloud hackathon, Chapel Hill, NC (3 days)	03/2019
Cyber Carpentry, Chapel Hill, NC (2 weeks)	06/2018
XSEDE HPC Workshop: Big Data, Tucson, AZ (2 days)	02/2018
Open Science Grid User School, Madison, WI (1 week)	07/2017

SELECT PUBLICATIONS, POSTERS, & TALKS

Gladstein A.L.*, et al. A Novel Machine Learning Method for Classification of Village Dogs. Probabilistic Modeling in Genomics. 2023. Cold Spring Harbor, NY. (Poster)

Baumdicker, F.*, Bisschop, G.*, Goldstein, D.*, Gower, G.*, Ragsdale, A. P.*, Tsambos, G.*, Zhu, S.*, ..., Gladstein, A. L., et al. 2022. Efficient ancestry and mutation simulation with msprime 1.0. *Genetics*. Volume 220, Issue 3. https://doi.org/10.1093/genetics/iyab229

Adrion, J. R.*, Cole, C. B.*, Dukler, N.*, Galloway, J. G.*, **Gladstein, A. L.**†, Gower, G.*, Kyriazis, C.C.*, Ragsdale, A.P.*, Tsambos, G.*, et al. 2020. A community-maintained standard library of population genetic models. *eLife*. 9:e54967. doi: https://doi.org/10.1101/2019.12.20.885129

Gladstein A.L.[†], Schrider R.D. Demographic model selection with deep learning. Probabilistic Modeling in Genomics. 2019. Aussois, France. (Poster)

Gladstein A.L.* and Hammer M.F. 2019. Substructured population growth in the Ashkenazi Jews inferred with Approximate Bayesian Computation. *Molecular Biology and Evolution*. 36(6): 1162-1171. doi: https://dx.doi.org/10.1093/molbev/msz047

^{*}Presenter

[†]First author

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

Code optimization for research scientists. Research Bazaar. 2018. Tucson, AZ. (Talk)

Gladstein A.L.* et al. 2018. SimPrily: A Python framework to simplify high-throughput genomic simulations. SoftwareX, 7, 335-340. https://doi.org/10.1016/j.softx.2018.09.003

OTHER ACTIVITIES

Circus Arts
Aerial silks, aerial rope, lyra, flying trapeze, tightwire, handbalancing, contortion

Figure Skating 1998 - 2013

 $Singles\ free style$