Seth Temple, Ph.D. in Statistics in 🖓 😵

Modeling in evolution and epidemiology using rigorous statistics

About Me: I have developed several interpretable, computationally efficient, and robust statistical methods for challenging problems in biology. Much of my work involves <u>modeling spatiotemporal processes in limited data</u> scenarios or in the face of complex correlation structures. I am interested in extending my expertise to problems at the <u>intersection of deep learning and health</u>. In internships, collaborations, and consultations, I have quickly learned new application areas to offer valuable statistical insights to research teams. I have been recognized for my ability to <u>communicate complex scientific topics clearly</u> and effectively.

EXPERIENCE

Post-doctoral Fellow, Schmidt AI in Science, University of Michigan	2024–2025
 Developing probabilistic deep learning models to simulate publicly available get that is representative of a population sample but protects individual privacy Working group on physics-informed neural networks in basic science 	genetic data
Researcher, University of Washington	2019–2024
 Proposed <u>multiple-testing</u> corrections in detecting evidence of natural selection and phenotypic case-control associations Proved <u>central limit theorems</u> in light of correlations from unobserved genealed Developed a suite of methods to model selection with uncertainty and infer ca Developed a linear runtime algorithm versus worse-case quadratic runtime Assistantship on Gaussian mixture models for genetic ancestry Researcher, Fred Hutchinson Cancer Research Center 	ogies
 Developed a novel method to detect and <u>cluster</u> adaptive SARS-CoV-2 varian 	
 Developed a novel method to detect and <u>cluster</u> adaptive SARS-Cov-2 variant based on amino acid <u>mutation counts over time</u> Provided feedback on machine learning models for non-neutral synonymous r 	
Researcher, Los Alamos National Laboratory	06/30-10/20
 Extended Gibbs sampler for <u>Bayesian spatiotemporal regression</u> in specific <u>missing data</u> mechanism in mosquito-borne epidemiology Developed and visualized <u>predictive maps</u> for mosquito prevalence 	
Actuary Trainee, Liberty Mutual Insurance	06/17-09/17, 07/18-08/19
EDUCATION	
 Ph.D., M.S., Statistics, University of Washington <u>NDSEG Fellow</u>, NIH Trainee in Statistical Genetics Advisors: Sharon Browning and Elizabeth Thompson 	2019–2024
B.S., Mathematics, Summa cum laude, University of Oregon	2014-2018
SKILLS [*] python [*]	

Excellent presentations: Z.W. Birnbaum Award, WNAR conference award, Seattle Science Slam

6+ years teaching/tutoring, including as the lead instructor of a graduate course

Coding: Python + Snakemake and R (expert), C++, SQL, and bash (intermediate)

Computing: slurm and qsub clusters, terabytes of genetic sequence data

SOFTWARE

<u>https://github.com/sdtemple/isweep</u>: extensively documented Python package <u>supporting 5 research articles</u> and containing <u>4 automated pipelines</u> for large-scale genomic analyses of natural selection.

<u>https://github.com/sdtemple/flare-pipeline</u>: automatic pipeline using Python and R for preliminary analyses in genomic analyses that require <u>Tb's disk memory</u>, <u>100's Gb RAM</u>, and <u>10's CPUs</u> on slurm clusters.

https://github.com/sdtemple/btvoccu: R for Bayesian logistic regression in epidemiology with missing data.

https://github.com/sdtemple/pblas: R for efficient simulation and estimation in SIR epidemiology model.

PUBLICATIONS

- **Temple, S.D.,** Waples, R.K., & Browning, S.R. Modeling recent positive selection using identity-by-descent segments. *The American Journal of Human Genetics* (2024). <u>https://doi.org/10.1016/j.ajhg.2024.08.023</u>.
- Temple, S.D., Manore, C.A. & Kaufeld, K.A. Bayesian time-varying occupancy model for West Nile virus in Ontario, Canada. *Stoch Environ Res Risk Assess* (2022). <u>https://doi.org/10.1007/s00477-022-02257-4</u>.
- **Temple, S.D.,** & Thompson, E.A. Identity-by-descent in large samples. *bioRxiv* (2024). <u>https://www.biorxiv.org/content/10.1101/2024.06.05.597656v1</u>.
- **Temple, S.D.,** Browning, S.R., & Thompson, E.A. Fast simulation of identity-by-descent segments. *bioRxiv* (2024). <u>https://www.biorxiv.org/content/10.1101/2024.12.13.628449v1</u>
- **Temple, S.D.** Statistical Inference using Identity-by-Descent Segments: Perspectives on Recent Positive Selection. PhD thesis. University of Washington (2024). <u>https://www.proquest.com/docview/3105584569</u>.
- **Temple, S.D.** The Tweedie Index Parameter and Its Estimator. Bachelor's thesis. University of Oregon (2018). <u>https://scholarsbank.uoregon.edu/xmlui/handle/1794/29040.</u>

Writing in progress:

- **Temple, S.D.**, & Browning, S.R. Multiple-testing corrections in selection studies using identity-by-descent segments. *bioRxiv* (December 2024). Coauthor reviewing.
- **Temple, S.D.**, Thornton, T.A., Wijsman, E.M., & Blue, E.E. Multiple-testing corrections in case-control studies using identity-by-descent segments. *bioRxiv* (estimated Winter 2025).

Collaborations:

- Gorris, M.E., Bartlow, A.W., **Temple, S.D.**, et al. Updated distribution maps of predominant Culex mosquitoes across the Americas. *Parasites & Vectors* 14, 547 (2021). <u>https://doi.org/10.1186/s13071-021-05051-3</u>.
- Haddox, H.K., Angehrn, G., Sesta, L., Jennings-Shaffer, C., Temple, S.D., Galloway, J.G., DeWitt, W.S., Matsen IV, F.A., & Neher, R.A. SARS-CoV-2's mutation rate is highly variable between sites and is influenced by sequence context, genomic region, and RNA structure. *bioRxiv*. To be submitted in late December 2024.
- Horimoto, A.R.V.R., Boyken, L.A., Blue, E.E., et al. Admixture mapping implicates 13q33.3 as ancestry-of-origin locus for Alzheimer disease in Hispanic and Latino populations. *HGG Advances* 4 (3) (2023): 100207.