# Thomas A. Sasani

#### Education

- 2019 Ph.D, Human Genetics, University of Utah, Salt Lake City, UT
- 2015 B.A, Biochemistry, Lawrence University, Appleton, WI, summa cum laude

### Experience

- 10/22-pres. Staff Research Scientist, Quinlan Lab, Univ. of Utah, Dept. of Human Genetics
  - O Developing new methods to analyze germline mutation and genome evolution
- 5/21-10/22 **Senior Data Scientist**, Recursion Pharmaceuticals
  - O Developed new computational methods to analyze data from massive cellular imaging experiments
  - Collaborated with product managers and scientists to disseminate results to wide audiences
  - Led regular sprint planning sessions and delegating project goals to teams of ICs
- 3/20-5/21 **Postdoctoral Fellow**, *Harris Lab*, Univ. of Washington, Dept. of Genome Sciences
  - Discovered alleles that influence the germline mutation rate using whole-genome sequencing data
  - O Analyzed single-cell sequencing data to characterize the mutational landscape of spermatogenesis
  - O Developed reproducible workflows for processing large DNA sequencing datasets
- 4/16-3/20 **Graduate Research Assistant**, *Quinlan Lab*, Univ. of Utah, Dept. of Human Genetics
  - Analyzed whole-genome sequencing data from large multi-generational families to identify post-zygotic mosaicism and variability in human germline mutation rates
  - Used the Oxford Nanopore Technologies platform to sequence DNA virus genomes under strong selective pressure during experimental evolution

## Skills

Programming Python (proficient), R (familiar), SQL (familiar)

Computing Unix, git, Sun Grid Engine, Amazon EC2

Data analysis numpy, scipy, sklearn, pandas, jupyter, unit testing frameworks (pytest)

Visualization matplotlib, ggplot2, plotly + Dash

## Selected Academic Publications (full list at Google Scholar)

- 2022 **Sasani TA**, Ashbrook DG, Beichman AC, Lu L, Palmer AA, Williams RW, Pritchard JK, Harris K. A natural mutator allele shapes mutation spectrum variation in mice. *Nature*. **Code**.
- 2019 **Sasani TA**, Pedersen BS, Gao Z, Baird L, Przeworski M, Quinlan AR, Jorde LB. Large, three-generation human families reveal post-zygotic mosaicism and variability in germline mutation accumulation. *eLife*. **Code**. Interview on the Naked Scientists podcast.

## Funding and Awards

- 2020 2021 NIH T32 Postdoctoral Genome Sciences Training Grant
- 2017 2019 NIH T32 Predoctoral Genetics Training Grant
  - 2017 Lassonde Student Innovator, University of Utah