# Thomas A. Sasani

## Education

8/15-10/19 **Ph.D, Human Genetics**, *University of Utah*, Salt Lake City, UT.

advisor: Professor Aaron Quinlan

8/11-6/15 BA, Biochemistry, Lawrence University, Appleton, WI, summa cum laude.

advisor: Professor Brian Piasecki

## Experience

5/21-pres. **Senior Data Scientist**, *Recursion Pharmaceuticals*.

- Developing new computational methods to analyze data from massive cellular imaging experiments
- Collaborating with product managers and scientists to disseminate results to wide audiences
- o Leading 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
  - Analyzed single-cell sequencing data to characterize the mutational landscape of spermatogenesis
  - 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 CEPH/Utah families to identify post-zygotic mosaicism and variability in human germline mutation rates
  - Used the Oxford Nanopore Technologies platform to sequence and analyze the genome content of DNA viruses under strong selective pressure during experimental evolution
- 8/12-6/15 Undergraduate Research Assistant, Piasecki Lab, Lawrence University.
  - Cloned and visualized the expression of genes involved in sensory cilia structure and function
  - o Constructed an automated worm tracking instrument to identify behavioral defects in C. elegans

## Preprints

- 2022 Ashbrook DG, **Sasani TA**, Maksimov M, Gunturkun MH, Ma N, Villani F, Ren Y, Rothschild D, Chen H, Lu L, Colonna V, Dumont B, Harris K, Gymrek M, Pritchard JK, Palmer AA, Williams RW. Private and sub-family specific mutations of founder haplotypes in the BXD family reveal phenotypic consequences relevant to health and disease. *bioRxiv*.
- 2020 Fixsen SM, Cone KR, Goldstein SA, **Sasani TA**, Quinlan AR, Rothenburg S, Elde NC. Poxviruses capture host genes by LINE-1 retrotransposition. *bioRxiv*.

## Peer-reviewed manuscripts (also see 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**.
- 2021 Belyeu JR\*, **Sasani TA**\*, Pedersen BS, Quinlan AR. Unfazed: parent-of-origin detection for large and small de novo variants. *Bioinformatics*.
- 2020 Wallace AD, **Sasani TA**, Swanier J, Gates B, Greenland J, Pedersen BS, Varley KT, Quinlan AR. CaBagE: a Cas9-based Background Elimination strategy for targeted, long-read DNA sequencing. *PLoS One*.

- 2020 Cawthon RM, Meeks HD\*, Sasani TA\*, Smith KR, Kerber RA, O'Brien E, Baird L, Dixon MM, Peiffer AP, Leppert MF, Quinlan AR, Jorde LB. Germline mutation rates in young adults predict longevity and reproductive lifespan. Scientific Reports.
- 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.
- 2019 Gao Z, Moorjani P, Sasani TA, Pedersen BS, Quinlan AR, Jorde LB, Amster G, Przeworski M. Overlooked roles of DNA damage and maternal age in generating human germline mutations. *PNAS*.
- 2018 **Sasani TA\***, Cone KR\*, Quinlan AR, Elde NC. Long read sequencing reveals poxvirus evolution through rapid homogenization of gene arrays. *eLife*. **Code**.
- 2018 Belyeu JR, Nicholas TJ, Pedersen BS, **Sasani TA**, Havrilla JM, Kravitz SN, Conway ME, Lohman BK, Quinlan AR, Layer RM. SV-plaudit: A cloud-based framework for manually curating thousands of structural variants. *GigaScience*.
- 2018 Jain M\*, Koren S\*, Miga KM\*, Quick J\*, Rand AC\*, Sasani TA\*, Tyson JR\*, Beggs AD, Dilthey AT, Fiddes IT, Malla S, Marriott H, Nieto T, O'Grady J, Olsen HE, Pedersen BS, Rhie A, Richardson H, Quinlan AR, Snutch TP, Tee L, Paten B, Phillippy AM, Simpson JT, Loman NJ, Loose M. Nanopore sequencing and assembly of a human genome with ultra-long reads. Nature Biotechnology.
- 2017 Feusier J, Witherspoon DJ, Watkins WS, Goubert C, **Sasani TA**, Jorde LB. Discovery of rare, diagnostic Alu Yb8/9 elements in diverse human populations. *Mobile DNA*.
- 2017 Piasecki BP, Sasani TA, Lessenger AT, Huth N, Farrell S. MAPK-15 is a ciliary protein required for PKD-2 localization and male mating behavior in *Caenorhabditis elegans*. *Cytoskeleton*.
  - \* indicates equal contribution

## Awards and Fellowships

- 2020-2021 NIH T32 Postdoctoral Genome Sciences Training Grant, University of Washington.
- 2017-2019 NIH T32 Predoctoral Genetics Training Grant, University of Utah.
- 2017 & 2018 Charles J. Epstein Trainee Award for Excellence in Human Genetics, Semifinalist, American Society for Human Genetics.
  - 2018 Lassonde Institute Student Innovator, University of Utah.
  - 2015 Howard and Helen Russell Award for Excellence in Biological Research, Lawrence University.

## Invited Presentations

- 2021 *Mapping mutator alleles in mice*. Recent Advances in Biology Lecture Series, Lawrence University. Virtual.
- 2021 Mapping mutator alleles in mice. Pritchard Lab Mini-Conference, Stanford University. Virtual.
- 2021 A wild-derived antimutator drives germline mutation spectrum differences in a genetically diverse murine family. Przeworski Lab Meeting, Columbia University. Virtual.
- 2019 A short tale of viral evolution told with long reads. Society for Molecular Biology and Evolution (Annual Meeting). Manchester, UK.

2017 Human immune defense mechanisms drive rapid genome evolution in vaccinia virus. London Calling (Oxford Nanopore Technologies meeting). London, UK.

## Contributed Presentations

- 2018 Directly measuring the dynamics of the human mutation rate by sequencing large, multigenerational pedigrees. American Society of Human Genetics, Annual Meeting. San Diego, CA, USA (Plenary Presentation).
- 2017 Human immune defense mechanisms drive rapid genome evolution in vaccinia virus. American Society of Human Genetics Annual Meeting. Orlando, FL, USA (Platform Presentation).

## Mentorship

#### **Graduate Students**

- 9/20-12/20 **Candice Y.**, *Rotation Project:* Identifying structural variants in a recombinant inbred mouse cross, University of Washington.
- 10/19-12/19 **Bianca A.**, *Rotation Project:* Patterns of recombination in large Utah pedigrees, University of Utah.
  - 1/20-3/20 **Erica H.**, *Rotation Project:* Regional variation in mutation rates and spectra, University of Utah.

## High School Students

7/20-9/20 **Myles F.**, *Summer Project:* Variation in mutation rates and spectra in a recombinant inbred mouse cross, University of Washington.

# Teaching Experience

## Cold Spring Harbor Laboratory

2016 & 2017 **Teaching Assistant**, Advanced Sequencing Technologies and Applications.

#### University of Utah

- 2019-2020 Guest Lecturer, Salt Lake Learners of Biostatistics.
  - 2017 **Teaching Assistant**, Applied Computational Genomics.
  - 2016 **Teaching Assistant**, *Programming for Biomedical Science*.
  - 2016 **Guest Instructor**, Summer Data Science Bootcamp.

## Reviewing

ad hoc Genome Biology, Genome Medicine, Molecular Biology and Evolution, Bioinformatics, eLife

## Courses and Professional Development

2018 Leena Peltonen School of Human Genomics