

Stanford



Julia Salzman

Associate Professor of Biomedical Data Science, of Biochemistry and, by courtesy, of Statistics and of Biology

Bio

BIO

The Salzman lab is primarily a statistical and computational biology data science laboratory. We are focusing on a new unifying paradigm, SPLASH (Statistically Primary Alignment Agnostic Sequence Homing), an approach that directly analyzes raw sequencing data to detect a signature of regulation: sample-specific sequence variation. The approach, which includes a new statistical test, is computationally efficient and can be run at scale. SPLASH unifies detection of myriad forms of sequence variation. Our group uses data mining and collaboration in our group and others to experimentally validate predictions on biological systems from single cell analyses of human and non-model organisms to microbes, plants viruses and symbioses. The reference-free nature of SPLASH enable co-detection of genome regulation in and outside of the human genome, including in human variation missing from reference genomes.

ACADEMIC APPOINTMENTS

- Associate Professor, Biomedical Data Science
- Associate Professor, Biochemistry
- Associate Professor (By courtesy), Statistics
- Associate Professor (By courtesy), Biology
- Member, Bio-X
- Member, Institute for Computational and Mathematical Engineering (ICME)
- Faculty Fellow, Sarafan ChEM-H
- Member, Stanford Cancer Institute
- Member, Wu Tsai Neurosciences Institute

HONORS AND AWARDS

- Arc Ignite Investigator, Arc Institute (2023)
- NSF CAREER AWARD, National Science Foundation (2016-2021)
- McCormick-Gabilan Fellowship, Stanford University (2015)
- Alfred P. Sloan Research Fellow, Alfred P. Sloan Foundation (2014)
- Baxter Faculty Scholar Award, Baxter Foundation (2014)
- Pathway to Independence (K99/R00) Award, National Institutes of Health (2012-present)
- Research Grant, National Science Foundation (2009-2012)
- Magna Cum Laude, Princeton University (2002)

PROFESSIONAL EDUCATION

- Ph. D., Stanford University , Statistics (2007)
- A. B., Princeton University , Mathematics (2002)

LINKS

- Lab Website: <http://med.stanford.edu/salzmanlab.html>

Research & Scholarship

CURRENT RESEARCH AND SCHOLARLY INTERESTS

statistical computational biology focusing on splicing, cancer and microbes

Teaching

COURSES

2023-24

- Statistical Genomics for Planetary Health: Oceans, Plants, Microbes and Humans: BIODS 228 (Aut)
- Workshop in Biostatistics: BIODS 260C, STATS 260C (Spr)

2022-23

- Workshop in Biostatistics: BIODS 260A, STATS 260A (Aut)

2021-22

- Introduction to Analysis of RNA Sequence Data: BIOC 239, BIODS 239 (Aut)
- Workshop in Biostatistics: BIODS 260A, STATS 260A (Aut)

2020-21

- Topics in Biomedical Data Science: Large-scale inference: BIODS 215 (Win)

STANFORD ADVISEES

Med Scholar Project Advisor

Ramon Lorenzo Labitigan

Doctoral Dissertation Reader (AC)

Isabel Delwel, Yannick Lee-Yow, Ivan Zheludev

Master's Program Advisor

Katie Ropers, Funmi Solano

GRADUATE AND FELLOWSHIP PROGRAM AFFILIATIONS

- Biochemistry (Phd Program)
- Biomedical Informatics (Phd Program)
- Biophysics (Phd Program)
- Cancer Biology (Phd Program)

Publications

PUBLICATIONS

- **Cell types of origin of the cell-free transcriptome.** *Nature biotechnology*
Vorperian, S. K., Moufarrej, M. N., Tabula Sapiens Consortium, Quake, S. R., Jones, R. C., Karkanias, J., Krasnow, M., Pisco, A. O., Quake, S. R., Salzman, J., Yosef, N., Bulthaup, B., Brown, P., et al
2022
- **RNA splicing programs define tissue compartments and cell types at single cell resolution.** *eLife*
Olivieri, J. E., Dehghannasiri, R., Wang, P. L., Jang, S., de Morree, A., Tan, S. Y., Ming, J., Ruohao Wu, A., Tabula Sapiens Consortium, Quake, S. R., Krasnow, M. A., Salzman, J.
2021; 10
- **Specific splice junction detection in single cells with SICILIAN.** *Genome biology*
Dehghannasiri, R., Olivieri, J. E., Damljanovic, A., Salzman, J.
2021; 22 (1): 219
- **Circular RNA Expression: Its Potential Regulation and Function in Abdominal Aortic Aneurysms** *OXIDATIVE MEDICINE AND CELLULAR LONGEVITY*
Han, Y., Zhang, H., Bian, C., Chen, C., Tu, S., Guo, J., Wu, Y., Boeckler, D., Zhang, J.
2021; 2021: 9934951
- **High-throughput SARS-CoV-2 and host genome sequencing from single nasopharyngeal swabs.** *medRxiv : the preprint server for health sciences*
Gorzynski, J. E., De Jong, H. N., Amar, D., Hughes, C. R., Ioannidis, A., Bierman, R., Liu, D., Tanigawa, Y., Kistler, A., Kamm, J., Kim, J., Cappello, L., Neff, et al
2020
- **Molecular sampling at logarithmic rates for next-generation sequencing.** *PLoS computational biology*
Horn, C., Salzman, J.
2019; 15 (12): e1007537
- **Improved detection of gene fusions by applying statistical methods reveals oncogenic RNA cancer drivers.** *Proceedings of the National Academy of Sciences of the United States of America*
Dehghannasiri, R., Freeman, D. E., Jordanski, M., Hsieh, G. L., Damljanovic, A., Lehnert, E., Salzman, J.
2019
- **Hyperammonemia after capecitabine associated with occult impairment of the urea cycle.** *Cancer medicine*
Chu, G., Salzman, J.
2019
- **Discovery of gene regulatory elements through a new bioinformatics analysis of haploid genetic screens.** *PloS one*
Patel, B. B., Lebensohn, A. M., Pusapati, G. V., Carette, J. E., Salzman, J. n., Rohatgi, R. n.
2019; 14 (1): e0198463
- **Ambiguous splice sites distinguish circRNA and linear splicing in the human genome.** *Bioinformatics (Oxford, England)*
Dehghannasiri, R., Szabo, L., Salzman, J.
2018
- **cIRS-7 exonic sequence is embedded in a long non-coding RNA locus** *PLOS GENETICS*
Barrett, S. P., Parker, K. R., Horn, C., Mata, M., Salzman, J.
2017; 13 (12): e1007114
- **Statistical algorithms improve accuracy of gene fusion detection.** *Nucleic acids research*
Hsieh, G., Bierman, R., Szabo, L., Lee, A. G., Freeman, D. E., Watson, N., Sweet-Cordero, E. A., Salzman, J.
2017
- **Extensive site-directed mutagenesis reveals interconnected functional units in the alkaline phosphatase active site.** *eLife*
Sunden, F., Peck, A., Salzman, J., Ressl, S., Herschlag, D.
2015; 4

- **Circular RNA biogenesis can proceed through an exon-containing lariat precursor.** *eLife*
Barrett, S. P., Wang, P. L., Salzman, J.
2015; 4
- **Statistically based splicing detection reveals neural enrichment and tissue-specific induction of circular RNA during human fetal development** *Genome biology*
Szabo1, L.
2015
- **Circular RNA Is Expressed across the Eukaryotic Tree of Life.** *PloS one*
Wang, P. L., Bao, Y., Yee, M., Barrett, S. P., Hogan, G. J., Olsen, M. N., Dinneny, J. R., Brown, P. O., Salzman, J.
2014; 9 (3)
- **Best permutation analysis** *JOURNAL OF MULTIVARIATE ANALYSIS*
Rajaratnam, B., Salzman, J.
2013; 121: 193-223
- **Cell-type specific features of circular RNA expression.** *PLoS genetics*
Salzman, J., Chen, R. E., Olsen, M. N., Wang, P. L., Brown, P. O.
2013; 9 (9)
- **Association between living environment and human oral viral ecology** *ISME JOURNAL*
Robles-Sikisaka, R., Ly, M., Boehm, T., Naidu, M., Salzman, J., Pride, D. T.
2013; 7 (9): 1710-1724
- **Improved discovery of molecular interactions in genome-scale data with adaptive model-based normalization.** *PloS one*
Salzman, J., Klass, D. M., Brown, P. O.
2013; 8 (1)
- **A penalized likelihood approach for robust estimation of isoform expression** *eprint arXiv:1310.0379*
Jiang, H., Salzman, .
2013
- **Statistical properties of an early stopping rule for resampling-based multiple testing** *BIOMETRIKA*
Jiang, H., Salzman, J.
2012; 99 (4): 973-980
- **Extensive Gene-Specific Translational Reprogramming in a Model of B Cell Differentiation and Abl-Dependent Transformation** *PLOS ONE*
Bates, J. G., Salzman, J., May, D., Garcia, P. B., Hogan, G. J., McIntosh, M., Schlissel, M. S., Brown, P. O.
2012; 7 (5)
- **Evidence of a robust resident bacteriophage population revealed through analysis of the human salivary virome** *ISME JOURNAL*
Pride, D. T., Salzman, J., Haynes, M., Rohwer, F., Davis-Long, C., White, R. A., Loomer, P., Armitage, G. C., Relman, D. A.
2012; 6 (5): 915-926
- **Widespread mRNA Association with Cytoskeletal Motor Proteins and Identification and Dynamics of Myosin-Associated mRNAs in *S. cerevisiae*** *PLOS ONE*
Casolari, J. M., Thompson, M. A., Salzman, J., Champion, L. M., Moerner, W. E., Brown, P. O.
2012; 7 (2)
- **Circular RNAs Are the Predominant Transcript Isoform from Hundreds of Human Genes in Diverse Cell Types** *PLOS ONE*
Salzman, J., Gawad, C., Wang, P. L., Lacayo, N., Brown, P. O.
2012; 7 (2)
- **Comparisons of CRISPRs and viromes in human saliva reveal bacterial 3 adaptations to salivary viruses** *Environmental Microbiology*
Pride, D.
2012
- **ESRRA-C11orf20 Is a Recurrent Gene Fusion in Serous Ovarian Carcinoma** *PLOS BIOLOGY*
Salzman, J., Marinelli, R. J., Wang, P. L., Green, A. E., Nielsen, J. S., Nelson, B. H., Drescher, C. W., Brown, P. O.

2011; 9 (9)

● **Statistical Modeling of RNA-Seq Data** *STATISTICAL SCIENCE*

Salzman, J., Jiang, H., Wong, W. H.

2011; 26 (1): 62-83

● **Analysis of streptococcal CRISPRs from human saliva reveals substantial sequence diversity within and between subjects over time** *GENOME RESEARCH*

Pride, D. T., Sun, C. L., Salzman, J., Rao, N., Loomer, P., Armitage, G. C., Banfield, J. F., Relman, D. A.

2011; 21 (1): 126-136

● **Proteome-Wide Search Reveals Unexpected RNA-Binding Proteins in *Saccharomyces cerevisiae*** *PLOS ONE*

Tsvetanova, N. G., Klass, D. M., Salzman, J., Brown, P. O.

2010; 5 (9)

● **Reliable concurrent calling of multiple genetic alleles and 24-chromosome ploidy without embryo freezing using parental support™ technology (PS)** *Fertility and Sterility*

Rabinowitz, M.

2008

● **Limits on the ability of quantum states to convey classical messages** *JOURNAL OF THE ACM*

Nayak, A., Salzman, J.

2006; 53 (1): 184-206

● **An improved upper bound for the pebbling threshold of the n-path** *Discrete Mathematics*

Wierman, A.

2004