Yuan Xue

BIOINFORMATICS SCIENTIST

▼ xuesoso@gmail.com | In LinkedIn | O GitHub | Xuesoso.github.io | G Google Scholar

EMPLOYMENT

ClearNote Health

San Mateo, California, USA

Bioinformatics Scientist II

Jan 2022 - Present

- contributes to optimization of molecular assays and development of computational methods for cancer detection based on circulating cfDNA in blood
- orchestrates project timelines, designs experimental plans, analyzes data, evaluates results with statistical methods, and presents findings to internal and external stakeholders
- develops methods and bioinformatics pipelines to quantify cfDNA and genomic CNV based on whole-genome sequencing and 5hmC-enriched sequencing data

EDUCATION

Stanford University	Stanford, California, U.S.A.
Ph.D. in Bioengineering	$Sept \ 2015 - Dec \ 2021$
Thesis advisers: Dr. Stephen Quake, Dr. John Boothroyd	
Stanford University	Stanford, California, U.S.A.
M.S. in Bioengineering	Sept 2015 – May 2017
Reed College	Portland, Oregon, U.S.A.
$B.A.\ in\ Biology$	Sept 2010 – May 2014
Thesis adviser: Dr. Jay Mellies	

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Awards & Honors	
Bio-X Travel Award Stanford Bio-X program	2019
Stanford Interdisciplinary Graduate Fellowship Awarded to eleven students to support their doctoral research with an interdisciplinary scope for three years.	2018
Postbaccalaureate Research Fellowship Reed College, Biology Department	2014
Summer Experience Research Award Reed College	2013
Gold Team Medal MIT iGEM Competition	2009

SELECTED PUBLICATIONS

- 1. Yuan Xue*, Yuhong Ning*, Verena Friedl, David Haan, Anna Bergamaschi, Gulfem Guler, Kyle Hazen, Aaron Scott, Tierney Phillips, Erin McCarthy, Christopher K. Ellison, Roger Malta, Albert Nguyen, Vanessa Lopez, William Gibb, Romola Cavet, Shimul Chowdhury, Wayne Volkmuth, Samuel Levy‡. 5-hydroxymethylcytosine analysis reveals stable epigenetic changes in tumor tissue that enable cancer detection in cell-free DNA. under review (2024).
- 2. Dania Nanes Sarfati, Yuan Xue, Eun Sun Song, Ashley Byrne, Daniel Le, Spyros Darmanis, Stephen R. Quake, Adrien Burlacot, James Sikest, Bo Wangt. Coordinated wound responses in a regenerative animal-algal photosymbiotic metaorganism. Nature Communications (2024).
- 3. Trung Pham‡*, Yuan Xue*, Susan Brewer, Kenneth E. Bernstein, Stephen R. Quake‡, Denise Monack‡. Single-cell profiling reveals functional diversity of granuloma macrophages during persistent Salmonella infection. Science Advances (2023). bioRxiv preprint
- 4. Yuan Xue, Ido Braslavsky, Stephen R. Quake. Temperature effect on DNA polymerase fidelity. Journal of Biological Chemistry (2021). bioRxiv preprint

- 5. Pengyang Li, Dania Nanes Sarfati*, **Yuan Xue***, Xi Yu, Alexander Tarashansky, Stephen R. Quake, Bo Wang. Single-cell analysis of Schistosoma mansoni reveals a conserved genetic program controlling germline stem cell fate. Nature Communications (2020). bioRxiv preprint
- 6. Suchita Rastogi, Yuan Xue, Stephen R. Quaket, John Boothroydt. Differential Impacts on Host Transcription by ROP and GRA Effectors from the Intracellular Parasite Toxoplasma gondii. mBio (2020). bioRxiv preprint
- 7. Yuan Xue, Terence Theisen, Suchi Rastogi, Abel Ferrel, Stephen R. Quaket, John Boothroydt. A single-parasite transcriptional atlas of Toxoplasma gondii reveals novel control of antigen expression. eLife (2020). bioRxiv preprint
- 8. Alexander Tarashansky, Yuan Xue, Pengyang Li, Stephen R. Quake, Bo Wang. Self-assembling Manifolds in Single-cell RNA Sequencing Data. Elife (2019). bioRxiv preprint
- 9. The Tabula Muris Consortium, Stephen R. Quake, Tony Wyss-Coray, Spyros Darmanis. Single-cell transcriptomics of 20 mouse organs creates a Tabula Muris. Nature (2018). bioRxiv preprint
- 10. Yuan Xue, Jossef Osborn, Anand Panchal, Jay L. Mellies. The RpoE stress response pathway mediates reduction of enteropathogenic Escherichia coli virulence by zinc. Applied and Environmental Microbiology (2015).
- 11. Jing Zhou, Shi-Hao Tan, Valerie Nicolas, Chantal Bauvy, Nai-Di Yang, Jianbin Zhang, Yuan Xue, Patrice Codogno, Han-Ming Shen. Activation of lysosomal function in the course of autophagy via mTORC1 suppression and autophagosome-lysosome fusion. Cell Research (2013).
 - *equal contributions; ‡corresponding authors

Professional Service

Stanford University
2018
Stanford University
2017
Reed College
2014
Reed College
2011 - 2014

Conferences & Presentations

Oral presentation ESMO Congress 2022

5-Hydroxymethycytosine analysis reveals stable epigenetic changes in tumor tissue that enable cfDNA $cancer\ predictions$

Invited talk National University of Singapore

2019

2019

Building a single-cell atlas of Toxoplasma interactome

Invited talk Cell Symposia Single Cells: From Technology to Biology

Building a single-cell atlas of Toxoplasma interactome

Poster presentation Stanford Bioengineering department retreat

Single-cell co-transcriptomic measurement resolves parasitic life cycle and

2018 $host\ interactions$

Invited talk Stanford Microbiology & Immunology department retreat Building a single-cell atlas of Toxoplasma interactome 2018

Stanford Bioengineering department retreat Poster presentation 2017

Cool biochemistry measured with a hot tool

Poster presentation

Temperature adaptation and polymerase fidelity

Gordon Research Conference: Nucleic Acids

Poster presentation Gordon Research Conference: Microbial Toxins & Pathogenicity

Molecular Mechanism of Zinc Disruption of Enteropathogenic Escherichia coli Pathogenesis

2014

SKILLS

Languages: English, Cantonese, Mandarin, Japanese, Python, R, Bash, LATEX Visualization: matplotlib, plotly, seaborn, bokeh, networkX, graphviz, graph-tool Machine Learning: numpy, pandas, sklearn, scikit-learn, tensorflow, keras, statsmodels

Workflows: Nextflow, Snakemake, cloud computing (e.g. AWS, slurm), Docker

Web development: Flask-REST backend, Apache2

Bioinformatics: STAR, minimap2, htseq-count, bedtools, salmon, velocyto, samtools, scanpy, Kraken

PACKAGE CONTRIBUTIONS

singleCell_snake

A snakemake pipeline for local/Slurm submission of single cell data alignment and transcript counting.

nheatmap

A python package to generate multi-level heatmap with extensive configuration options.

DensityPlot

A python package to generate density scatter plot.

bag_of_velocyto

A bash / python script for parallel submission of RNA velocity alignment on Slurm.