TENG GAO

tgao@g.harvard.edu | ORCID | GitHub | Google Scholar

EDUCATION

Harvard University (Cambridge, MA, USA)

Sep. 2020 – May 2024 (Expected)

Ph.D. Candidate in Bioinformatics and Integrative Genomics (GPA: 4.00/4.00)

Thesis advisors: Peter V. Kharchenko and Peter J. Park Committee: Nir Hacohen, Vijay Sankaran, Kamila Naxerova

Washington University (St. Louis, MO, USA)

Sep. 2014 – May 2018

B.S. Double Major in Computer Science and Biology (GPA: 3.99/4.00)

Summa Cum Laude

Honor thesis advisor: Christopher Maher

RESEARCH POSITIONS

Harvard Medical School, Department of Biomedical Informatics

Mar. 2021 - Present

Graduate Researcher (Advisors: Peter V. Kharchenko and Peter J. Park)

- Developed Numbat, a novel computational method to infer genome aberrations and single-cell phylogeny from scRNA-seq data
- Applied Numbat to characterize pediatric neuroblastoma and age-related mosaic chromosomal alterations in normal tissues
- Developing new computational methods for the analysis of subcellular RNA organization from spatial transcriptomics data

Memorial Sloan Kettering Cancer Center, Computational Oncology

May 2018 – Aug. 2020

Computational Biologist I-II (Advisors: Elli Papaemmanuil and Kelly Bolton)

• Led large-scale study on clonal hematopoiesis and therapy-related leukemia in 32,000 cancer patients

Washington University in St. Louis, McDonnell Genome Institute

Sep. 2016 - May 2018

Undergraduate researcher (Advisor: Christopher Maher)

- Co-developed INTEGRATE-Vis, a python visualization package for patient-specific gene fusions discovered in RNA-seq data
- Honor thesis: "ceFinder: Machine learning based prediction of novel competing endogenous RNAs."

Roche Molecular Systems

May 2017 – Aug. 2017

Bioinformatics Intern (Manager: Dwight Kuo)

- Developed RNA-Seq analysis pipelines for novel miRNA biomarker discovery and differential expression in prostate cancer
- · Identified a novel antibiotic resistance gene cassette in Methicillin-resistant S. aureus through de novo genome assembly

Wake Forest Institute for Regenerative Medicine

May 2016 – Aug. 2016

Summer Scholar (Advisors: Anthony Atala and Sang Jin Lee)

• Established a mathematical model for biomaterial viscoelasticity in 3D biofabrication

PUBLICATIONS

(*: equal contributions; ‡: co-corresponding authors)

Selected Publications

- Gao T, Eleni MK, Ljungström V, Heinzel A, Tischler AS, Oberbauer R, Loh PR, Adameyko I, Park PJ[‡], Kharchenko PV[‡] (2023). "A pan-tissue survey of mosaic chromosomal alterations in 948 individuals". *Nature Genetics*. In Press. 10.1038/s41588-023-01537-1. [Paper] [Code]
- Gao T, Soldatov R, Sarkar H, Kurkiewicz A, Biederstedt E, Loh PR, and Kharchenko PV (2022). "Haplotype-aware analysis of somatic copy number variations from single-cell transcriptomes". *Nature Biotechnology* 41, 417–426. 10.1038/s41587-022-01468-y. [Paper] [Code]

- 3. <u>Gao T</u>, Ptashkin R, Bolton KL, Sirenko M, Fong C, Spitzer B, Menghrajani K, Ossa JEA, Zhou Y, Bernard E, Levine M, Martinez JSM, Zhang Y, Franch SE, Patel M, Braunstein LZ, Kelly D, Yabe M, Benayed R, Caltabellotta NM, Philip J, Paraiso E, Mantha S, Solit DB, Diaz LA Jr, Berger MF, Klimek V, Levine RL, Zehir A, Devlin SM, Papaemmanuil E (2021). "Interplay between chromosomal alterations and gene mutations shapes the evolutionary trajectory of clonal hematopoiesis". *Nature Communications* 12, 338. 10.1038/s41467-020-20565-7. [Paper] [Code]
- 4. Bolton KL, Ptashkin RN*, <u>Gao T*</u>, Braunstein L, Devlin SM, Kelly D, Patel M, Berthon A, Syed A, Yabe M, Coombs CC, Caltabellotta NM, Walsh M, Offit K, Stadler Z, Mandelker D, Schulman J, Patel A, Philip J, Bernard E, Gundem G, Ossa JEA, Levine M, Martinez JSM, Farnoud N, Glodzik D, Li S, Robson ME, Lee C, Pharoah PDP, Stopsack KH, Spitzer B, Mantha S, Fagin J, Boucai L, Gibson CJ, Ebert BL, Young AL, Druley T, Takahashi K, Gillis N, Ball M, Padron E, Hyman DM, Baselga J, Norton L, Gardos S, Klimek VM, Scher H, Bajorin D, Paraiso E, Benayed R, Arcila ME, Ladanyi M, Solit DB, Berger MF, Tallman M, Garcia-Closas M, Chatterjee N, Diaz LA Jr, Levine RL, Morton LM, Zehir A, Papaemmanuil E (2020). "Cancer therapy shapes the fitness landscape of clonal hematopoiesis". *Nature Genetics* 52, 1219–1226. 10.1038/s41588-020-00710-0. [Paper]
- 5. <u>Gao T</u>*, Gillispie GJ*, Copus JS, Pr AK, Seol YJ, Atala A, Yoo JJ, and Lee SJ (2018). "Optimization of gelatin–alginate composite bioink printability using rheological parameters: a systematic approach". *Biofabrication* 10, 034106. 10.1088/1758-5090/aacdc7. [Paper]

Other Publications

- 6. Franch SE, Mehine M, Ptashkin RN, Bolton KL, Bandlamudi C, Srinivasan P, Zhang L, Goodell MA, Gedvilaite E, Menghrajani K, Sánchez-Vela P, Mandelker D, Comen E, Norton L, Benayed R, <u>Gao T</u>, Papaemmanuil E, Taylor B, Levine R, Offit K, Stadler Z, Berger MF, Zehir A (2023). "Associations between cancer predisposition mutations and clonal hematopoiesis in patients with solid tumors". *JCO Precision Oncology* 2023:7. 10.1200/PO.23.00070. [Paper]
- 7. Spitzer B, Rutherford KD, Gundem G, McGovern EM, Millard NE, Arango Ossa JE, Cheung IY, <u>Gao T</u>, Levine MF, Zhang Y, Medina JSM, Feng Y, Ptashkin RN, Bolton KL, Farnoud N, Zhou Y, Patel MA, Asimomitis G, Cobbs CC, Mohibullah N, Huberman KH, Arcilla ME, Kushner BH, Modak S, Kung AL, Zehir A, Levine RL, Armstrong SA, Cheung NKV, Papaemmanuil E (2022). "Bone marrow surveillance of pediatric cancer survivors identifies clones that predict therapy-related leukemia". *Clinical Cancer Research* 28, 1614–1627. 10.1158/1078-0432.CCR-21-2451. [Paper]
- 8. Bolton KL, Koh Y, Foote MB, Im H, Jee J, Sun CH, Safonov A, Ptashkin R, Moon JH, Lee JY, Jung J, Kang CK, Song K-H, Choe PG, Park WB, Kim HB, Oh M-D, Song H, Kim S, Patel M, Derkach A, Gedvilaite E, Tkachuk KA, Wiley BJ, Chan IC, Braunstein LZ, <u>Gao T</u>, Papaemmanuil E, Esther NB, Pessin MS, Kamboj M, Diaz LA Jr, Ladanyi M, Rauh MJ, Natarajan P, Machiela MJ, Awadalla P, Joseph V, Offit K, Norton L, Berger MF, Levine RL, Kim ES, Kim NJ, Zehir A (2021). "Clonal hematopoiesis is associated with risk of severe Covid-19". *Nature Communications* 12, 5975. 10.1038/s41467-021-26138-6. [Paper]
- 9. Zhang J, <u>Gao T</u>, Maher CA (2017). "INTEGRATE-Vis: a tool for comprehensive gene fusion visualization". *Scientific Reports* 7. 10.1038/s41598-017-18257-2. [Paper] [Code]

PRESENTATIONS & INVITED TALKS

New York University Applied Bioinformatics Laboratory Single-Cell Club - New York, NY	Sep. 2022
Single-Cell Cancer Biology Gordon Research Conference – Easton, MA	Jun. 2022
Broad Institute Single-Cell Seminar Series – Cambridge, MA	Feb. 2022
American Society of Hematology (ASH) 62nd Annual Meeting and Exhibition – San Diego, CA	Dec. 2020

HONORS & AWARDS

TEACHING EXPERIENCES

Harvard Medical School (Cambridge, MA, USA)

Sep. 2022 – Dec. 2022

Teaching Fellow, Concepts in Genome Analysis (BMIF 201)

Washington University (St. Louis, MO, USA)

Teaching Assistant, Elementary to Intermediate Statistics and Data Analysis (Math 3200)