Shankara K. Anand



shankara.k.anand@gmail.com • github.com/shankara-a

Updated May 27, 2024

ABOUT

I am a computational biologist and PGY-1 in internal medicine in Stanford's Translational Investigator Pathway for Hematology/Oncology. I am interested in plasma cell dyscrasias, amyloidosis, precision oncology, and early cancer detection. I have a Masters in machine/deep learning for medical applications and 5+ years of genomics expertise from the Broad Institute at MIT & Harvard. My research includes single-cell genomics, multi-omics, structural proteomics, and algorithm development.

CLINICAL

Resident Physician, Internal Medicine, Stanford Health Care, *Stanford*, *CA* 2024 - present

EDUCATION

M.D. (Cum Laude), Boston University, Boston, MA
M.S. Biomedical Informatics, Stanford University, Stanford, CA
B.S. Chemical Engineering, Stanford University, Stanford, CA
2017

LICENSURE

 USMLE Step 1 - Pass
 04/06/2022

 USMLE Step 2CK - 268
 05/26/2023

RESEARCH

Associate Computational Biologist, Broad Institute of MIT & Harvard

• Translational Investigator Pathway (Hematology/Oncology)

2018 - 2023

Supervised by Dr. Gad Getz & the Genotype Tissue Expression Experiment (GTEx)

- Co-led pan-cancer, proteogenomic study, published in *Cell* (CPTAC)
- Mapped single-nucleus, cross-tissue atlas of the human body, published in *Science* (GTEx)
- Risk stratified Smoldering Multiple Myeloma patients with genomics (advised by Dr. Irene Ghobrial)

Researcher, Stanford Dept. of Bioengineering

Jan. - June 2018

Supervised by Dr. Possu Huang

- Designed and tested *de novo*, end-to-end (computational to wet-lab) proteins using Rosetta software
- Predicted ligand binding sites of proteins using deep learning frameworks (GitHub)

Scientific Intern, Novartis

Summer 2017

Computer Aided Drug Discovery Team

- Developed machine learning framework for detecting pKa of in-house compounds
- Published in Journal of Chemical Information & Modeling

Research Intern, Genentech

Summer 2016

Oct. 2018 - Feb. 2020

Early Stage Pharmaceutical Development

• Built molecular dynamics simulation pipeline to investigate antibody degradation

LEADERSHIP

Internal Medicine Student Chief, BUSM Internal Medicine Clerkship	June 2023 - May 2024
Student Leader, BUSM Genetics Interest Group	April 2021 - March 2022
Student Leader, BUSM in. Vision Tech x Medicine Group	April 2021 - March 2022
STEM Tutor, Stanford Academic Resource Center	Summer 2017
Dance Captain, Stanford Basmati Raas (Indian Dance)	Sept. 2013 - June 2018
Resident Assistant, 2x years at Stanford Residential Education	Sept. 2015 - June 2017
Program Coordinator, Stanford LGBT Community Resource Center	Sept. 2015 - June 2017

VOLUNTEER

Fundraising Coordinator & Volunteer, Stanford Camp Kesem (500 hrs)

April 2016 - July 2018

Child Life Volunteer, Boston Children's Hospital Oncology (150 hrs)

2017

2016

SKILLS Advanced: Python, R, PyTorch, TensorFlow, git

Basic: Matlab, LATEX

SOFTWARE

AWARDS

Signature Analyzer: a fast, GPU enabled implementation of bayesian non-negative matrix factorization for extracting mutational and genomic expression signatures (GitHub).

CLUMPS-PTM: an algorithm for identifying 3D clusters of post-translational modifications (GitHub).

CLUMPS-2: an algorithm for identifying 3D clusters of somatic mutations (GitHub).

Genentech Outstanding Student Award, selected Junior in Chemical Engineering

Mudi: python utilities for single-cell RNA-sequencing analyses (GitHub).

Stanford Achievement Award, awarded to 10% of graduating class

SITEFINDER: using deep learning to identify ligand site binding in proteins (Stanford CS231n) (GitHub).

PUBLICATIONS Pre-Prints / Under Review

[1] Rouhana, J. M., Wang, J., Eraslan, G., Anand, S., Hamel, A. R., Cole, B. et al. ECLIPSER: identifying causal cell types and genes for complex traits through single cell enrichment of e/sQTL-mapped genes in GWAS loci. bioRxiv (2021). doi: 10.1101/2021.11.24.469720

Peer-Reviewed Publications

- [1] **Anand, S.**[†], Verma, A., Sanchorawala, V. Systemic Amyloidosis and Kidney Transplantation: An Update. **Seminars in Nephrology**, (2024). doi: 10.1016/j.semnephrol.2024.151496
- [2] Geffen, Y.[†], **Anand**, S.[†], Akiyama, Y.[†], Yaron, T.[†], Song, Y.[†], et al. Pan-Cancer Analysis of Post-Translational Modifications Reveals Shared Patterns of Protein Regulation. **Cell**, 186, 1-24 (2023). doi: 10.1016/j.cell.2023.07.013
- [3] Li, Y., Porta-Pardo, E., Tokheim, C., Bailey, M.H., Yaron, T.M., Stathias, V., Geffen, Y., Imbach, K.J., Cao, S., Anand, S., Akiyama, Y., et al. Pan-cancer proteogenomics connects oncogenic drivers to functional states. Cell, 186, 1–24 (2023). doi: 10.1016/j.cell.2023.07.014
- [4] Li, Y., Dou, Y., Leprevost, F. D. V., Geffen, Y., Calinawan, A. P., Aguet, F., Akiyama, Y., **Anand, S.**, et al. Proteogenomic data and resources for pan-cancer analysis. **Cancer Cell**, 41(8), 1397-1406 (2023). doi: 10.1016/j.ccell.2023.06.009
- [5] Boiarsky, R., Haradhvala, N. J., Alberge, J. B., Sklavenitis-Pistofidis, R., Mouhieddine, T. H., Zavidij, O. et al. Single Cell Characterization of Myeloma and its Precursor Conditions Reveals Transcriptional Signatures of Early Tumorigenesis. Nature Communications, 13, 7040 (2022). doi: 10.1038/s41467-022-33944-z
- [6] Sklavenitis-Pistofidis, R., Aranha, M. P., Redd, R. A., Baginska, J., Haradhvala, N. J., Hallisey, M., et al. Immune biomarkers of response to immunotherapy in patients with high-risk smoldering myeloma. Cancer Cell, 40(11), 1358-1373 (2022). doi: 10.1016/j.ccell.2022.10.017
- [7] Yaron, T. M., Heaton, B. E., Levy, T. M., Johnson, J. L., Jordan, T. X., Cohen, B. M. et al. Host protein kinases required for SARS-CoV-2 nucleocapsid phosphorylation and viral replication. **Science**

[†] indicates co-first authorship

[‡] indicates co-second authorship

- **Signaling**, 15(757) (2022). doi: 10.1126/scisignal.abm0808
- [8] Roh, W., Geffen, Y., Cha, H., Miller, M., Anand, S., Kim, J. et al. High-resolution profiling of lung adenocarcinoma identifies expression subtypes with specific biomarkers and clinically relevant vulnerabilities. Cancer Research (2022). doi: 10.1158/0008-5472.CAN-22-0432
- [9] Knisbacher, B. A., Lin, Z., Hahn, C. K., Nadeu, F., Duran-Ferrer, M., Stevenson, K. E. et al. The CLL-1100 project: towards complete genomic characterization and improved prognostics for CLL. Nature Genetics, 1-11 (2022). doi: 10.1038/s41588-022-01140-w
- [10] Bustoros, M.[†], **Anand, S.[†]**, Sklavenitis-Pistofidis, R. et al. Genetic subtypes of smoldering multiple myeloma are associated with distinct pathogenic phenotypes and clinical outcomes. **Nature Communications**, 13, 3449 (2022). doi: 10.1038/s41467-022-30694-w (GitHub)
- [11] Eraslan, G.[†], Drokhlyansky, E.[†], **Anand, S.[‡]**, Fiskin, E.[‡], Subramanian, A.[‡], Sylper, M.[‡], Wang, J.[‡] et al. Single-nucleus cross-tissue molecular reference maps toward understanding disease gene function. **Science**, 376(6594) (2022). doi: 10.1126/science.abl4290
- [12] Beauchamp, E. M., Leventhal, M., Bernard, E., Hoppe, E. R., Todisco, G., Creignou, M. et al. ZBTB33 is mutated in clonal hematopoiesis and myelodysplastic syndromes and impacts RNA splicing. **Blood Cancer Discovery**, 2(5), 500-517 (2021). doi: 10.1158/2643-3230.BCD-20-0224
- [13] Satpathy, S., Krug, K., Beltran, P. M. J., Savage, S. R., Petralia, F., Kumar-Sinha, C. et al. A proteogenomic portrait of lung squamous cell carcinoma. **Cell**, 184(16), 4348-4371 (2021). doi: 10.1016/j.cell.2021.07.016
- [14] de Goede, O. M., Nachun, D. C., Ferraro, N. M., Gloudemans, M. J., Rao, A. S., Smail, C. et al. Population-scale tissue transcriptomics maps long non-coding RNAs to complex disease. **Cell**, 184(10), 2633-2648 (2021). doi: 10.1016/j.cell.2021.03.050
- [15] Jiang, L., Wang, M., Lin, S., Jian, R., Li, X., Chan, J. et al. A quantitative proteome map of the human body. **Cell**, 183(1), 269-283 (2020). doi: 10.1016/j.cell.2020.08.036
- [16] Ferraro, N. M., Strober, B. J., Einson, J., Abell, N. S., Aguet, F., Barbeira, A. N. et al. Transcriptomic signatures across human tissues identify functional rare genetic variation. **Science**, 369(6509) (2020). doi: 10.1126/science.aaz5900
- [17] GTEx Consortium. The GTEx Consortium atlas of genetic regulatory effects across human tissues. **Science**, 369(6509), 1318-1330 (2020). doi: 10.1126/science.aaz1776
- [18] Bustoros, M., Sklavenitis-Pistofidis, R., Park, J., Redd, R., Zhitomirsky, B., Dunford, A. J. et al. Genomic profiling of smoldering multiple myeloma identifies patients at a high risk of disease progression. **Journal of Clinical Oncology**, 38(21), 2380 (2020). doi: 10.1200/JCO.20.00437
- [19] Taylor-Weiner, A., Aguet, F., Haradhvala, N. J., Gosai, S., **Anand, S.**, Kim, J. et al. Scaling computational genomics to millions of individuals with GPUs. **Genome Biology**, 20(1), 1-5 (2019). doi: 10.1186/s13059-019-1836-7
- [20] Lu, Y., **Anand, S.**, Shirley, W., Gedeck, P., Kelley, B. P., Skolnik, S. et al. Prediction of p K a using machine learning methods with rooted topological torsion fingerprints: application to aliphatic amines. **Journal of Chemical Information and Modeling**, 59(11), 4706-4719 (2019). doi: 10.1021/acs.jcim.9b00498
- [21] Anand, S., Kalesinskas, L., Smail, C., & Tanigawa, Y. SNPs2ChIP: Latent Factors of ChIP-seq to infer functions of non-coding SNPs. Proceedings of the Pacific Symposium (pp. 184-195) (2018). doi: 10.1142/9789813279827 0017

Peer-Reviewed Book Chapters

[1] **Anand, S.**, Verma, A. (2023). *Artificial Intelligence in Nephrology*. In C. Krittanawong (Ed.), Artificial Intelligence in Clinical Practice: How AI Technologies Impact Medical Research and Clinics (First Edition). New York City: Elsevier.

- [1] Roh, W., Geffen, Y., Miller, M., Anand, S., Kim, J., Heiman, D. et al. Identification and proteogenomic characterization of novel lung adenocarcinoma subtypes with therapeutic relevance. American Association of Cancer Research (AACR), 82(12_Supplement), 2151-2151 (2022). doi: 10.1158/1538-7445.AM2022-2151
- [2] Kubler, K., Nardone, A., Anand, S., Gorvich, D., Droog, M., Hermida-Prado, F. et al. Abstract GS2-09: Tamoxifen instigates uterine cancer development by activating PI3K signaling and supersedes PIK3CA driver mutations. San Antonio Breast Cancer Symposium (SABCS), 82(4_Supplement) (2021). doi: 10.1158/1538-7445.SABCS21-GS2-09
- [3] Sklavenitis-Pistofidis, R., Dutta, A. K., Ujwary, S., Redd, R. A., Savell, A., Fléchon, L. et al. Single-Cell RNA-Sequencing Identifies Immune Biomarkers of Response to Immunotherapy in Patients with High-Risk Smoldering Myeloma. **American Society of Hematology (ASH)**, 138, 330 (2021). doi: 10.1182/blood-2021-147623
- [4] **Anand, S.**[†], Bustoros, M.[†], Sklavenitis-Pistofidis, R., Redd, R. A., Boyle, E. M., Tai, Y. T. et al. Genomic Profiling of Smoldering Multiple Myeloma Classifies Molecular Groups with Distinct Pathogenic Phenotypes and Clinical Outcomes. **American Society of Hematology (ASH)**, 138, 723 (2021).
- [5] **Anand, S.** "Genomic profiling of smoldering myeloma identifies high- and low- risk patients | ASH 2021." **The Video Journal of Hematological Oncology**, Dec. 12, 2021. Interviews: Part 1. Part 2.
- [6] Geffen, Y.[†], **Anand, S.**[†], Akiyama, Y.[†], Yaron, T. M., Kerelsky, A., Johnson, J. L., ... & Oncogenic Drivers and Pathways Group, Clinical Proteomic Tumor Analysis Consortium (CPTAC). Patterns and regulation of post translational modifications in cancer. **American Association of Cancer Research (AACR)**, 81(13_Supplement), 16-16. doi: 10.1158/1538-7445.AM2021-16

Posters & Abstracts

- [1] **Anand, S.**, Staron, A., Mendelson, L. M., Joshi, T., Burke, N. J., Sanchorawala, V., Verma, A. Machine-Learning Based Subgroups of AL Amyloidosis and Cumulative Incidence of Mortality and End Stage Kidney Disease. **International Symposium on Amyloidosis**, (2024).
- [2] **Anand, S.**, Geffen, Y., Akiyama, Y., Aguet, F., Getz, G. *CLUMPS-PTM*: Spatial Clustering of Post-Translational Modifications Across Cancer Types. **American Association of Cancer Research** (**AACR**), 83(7_Supplement), 3132-3132 (2023). doi: 10.1158/1538-7445.AM2023-3132
- [3] Li, G. X., Hsiao, Y., Chen, L., Mannan, R., Zhang, Y., Petralia, F., et al. Comprehensive proteogenomic characterization of rare kidney tumors. American Association of Cancer Research (AACR), 83(7_Supplement), 3127-3127 (2023). doi: 10.1158/1538-7445.AM2023-3127
- [4] Boiarsky, R., Haradhvala, N., Sklavenitis-Pistofidis, R., Mouhieddine, T. H., Alberge, J. B., Zavidij, O. et al. Single Cell Characterization of Myeloma and its Precursor Conditions Reveals Transcriptional Signatures of Early Tumorigenesis. American Society of Hematology (ASH), 138, 2219 (2021). doi: 10.1182/blood-2021-150889
- [5] Akiyama, Y., Geffen, Y., Anand, S., Yaron, T., Johnson, J. L., Huntsman, E. et al. Pan-cancer proteogenomic analysis reveals functional mechanisms underlying DNA repair deficiencies. American Association of Cancer Research (AACR), 82(12_Supplement), 794-794 (2022). doi: 10.1158/1538-7445.AM2022-794
- [6] Anand, S., Bustoros, M., Aguet, F., Sklavenitis-Pistofidis, R., Redd, R., Zhitomirsky, B. et al. Genomic profiling of smoldering multiple myeloma classifies distinct molecular groups. American Association of Cancer Research (AACR), 81(13_Supplement), 2240-2240 (2021). doi: 10.1158/1538-7445.AM2021-2240
- [7] **Anand, S.**, Aguet, F., Taylor-Weiner, A., et al. Signature Analyzer: scaling Bayesian NMF to millions of individuals with GPUs. **Broad Institute of MIT & Harvard Retreat**, (2019). GitHub.
- HOBBIES **Rass/Garba** (**Indian**) **dance:** 5+ years of dancing and choreographing at cultural events (ex. Stanford Holi), national competitions, and family weddings.

Music: 15+ years of classical violin training, performed with Stanford Symphony Orchestra, chamber music groups, and musical theater pit orchestras; also enjoy guitar & ukulele.

Dog: My wife and I have a 4 year old Mudi dog, Hazel, who we do agility and frisbee catching with.

Ski Mountaineering: Ask me about the glacier we climbed and skied down in the Selkirk Mountains!

Rock Climbing: We love indoor & outdoor lead climbing + finding new routes along the Pacific Coast.