CURRICULUM VITAE

PERSONAL INFORMATION

Name: Yaqiang Cao

Organization: System Biology Center, National Heart, Lung, and Blood Institute

(NHLBI), National Institutes of Health (NIH)

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Links: Google Scholar | GitHub

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EDUCATION

2019 Ph.D. in Computational Biology

Chinese Academy of Sciences (CAS)- Max Planck Society (MPG) Partner Institute for Computational Biology, University of Chinese Academy of Sciences (UCAS), Shanghai, China.

Mentor: Prof. Jing-Dong Jackie Han

2012 B.S. in Bioengineering

East China University of Science and Technology, Shanghai, China.

RESEARCH EXPERIENCE

2024-Curr. Research Fellow

System Biology Center, National Heart, Lung, and Blood Institute (NHLBI).

National Institutes of Health (NIH), Bethesda, MD, USA.

Mentor: Dr. Keji Zhao

2019-2024 Visiting Fellow

NHLBI/NIH, Bethesda, MD, USA.

Mentor: Dr. Keji Zhao

2011-2018 Graduate Research Assistant

CAS-MPG Partner Institute for Computational Biology, Shanghai, China.

Mentor: Prof. Jing-Dong Jackie Han

RESEARCH SUMMARY

Focus: Chromatin regulation in immunity, leukemia, and aging.

Methods: Created computational tools (cLoops2, vince) and experimental techniques

(Hi-TrAC, iscMNase-seq) to map high-resolution enhancer-promoter

interactions and single-cell nucleosomes.

Discoveries: Chromatin loops/histone modifications/transcription factors driving T-cell

differentiation, acute myeloid leukemia, B-cell aging.

Collaborations: Cross-disciplinary partnerships bridging basic science and clinical

applications, involving biologists, physicians, and data scientists.

Impact:

User-friendly software for reproducible epigenomic and 3D genomic research; mechanism understandings for basic biology and clinical translations.

FIRST/CO-FIRST PUBLICATIONS (14 out of 31)

- * co-first author # correspondence.
- Tao Zhen *, **Yaqiang Cao** *, Tongyi Dou, Yun Chen, Guadalupe Lopez, Ana Catarina Menezes, Xufeng Wu, John A. Hammer, Jun Cheng, Lisa Garrett, Stacie Anderson, Martha Kirby, Stephen Wincovitch, Bayu Sisay, Abdel G. Elkahloun, Di Wu, Lucio Castilla, Wei Yang, Jiansen Jiang, Keji Zhao, Pu Paul Liu#: Enhancing RUNX1-DNA binding affinity is a critical step for leukemogenesis by CBFβ-SMMHC (Accepted, **JCI**)
- Ma, F. *, Cao, Y. *, Du, H., Braikia, F., Zong, L., Ollikainen, N., Bayer, M., Qiu, X., Park, B., Roy, R., Nandi, S., Sarantopoulou, D., Ziman, A., Bianchi, A., Beerman, I., Zhao, K., Grosschedl, R., Sen, R. *. (2024) Three-dimensional chromatin reorganization regulates B cell development during ageing. Nat Cell Biol. 1-12. Liu, S. *, Cao, Y. *, Cui, K*., Ren, G., Zhao, T., Wang, X., Wei, D., Chen, Z., Gurram, R.K., Liu, C. and Wu, C., Zhu, J., & Zhao, K. * (2024) Regulation of T helper cell differentiation by the interplay between histone modification and chromatin interaction. Immunity. 57(5), 987-1004.
- Cao, Y.*, Liu, S.*, Cui, K.*, Tang, Q., & Zhao, K.* (2023). Hi-TrAC detects active sub-TADs and reveals internal organizations of super-enhancers. Nucleic Acids Research, 51(12), 6172-6189.
 Cui, K.*, Chen, Z.*, Cao, Y.*, Liu, S.*, Ren, G., Hu, G., Fang, D., Wei, D., Liu, C., Zhu, J., Wu, C.*, & Zhao, K.* (2023). Restraint of IFN-γ expression through a distal silencer CNS–28 for tissue homeostasis. Immunity, 56(5), 944-958 (selected as cover, preview by Immunity)
- Cao, Y. *, Liu, S. *, Ren, G. *, Tang, Q., & Zhao, K. # (2022). cLoops2: a full-stack comprehensive analytical tool for chromatin interactions. <u>Nucleic Acids Research</u>, 50(1), 57-71.
 Liu, S. *, Cao, Y. *, Cui, K., Tang, Q., & Zhao, K. # (2022). Hi-TrAC reveals division of labor of transcription factors in organizing chromatin loops. <u>Nature communications</u>, 13(1), 1-17.
- 2020 Cao, Y. *, Chen, Z. *, Chen, X. *, Ai, D. *, Chen, G., McDermott, J., Huang, Y., Guo, X., & Han, J. # (2020). Accurate loop calling for 3D genomic data with cLoops.

 Bioinformatics, 36(3), 666-675.
 - Zhen, T. *, *Cao, Y.* *, Ren, G., Zhao, L., Hyde, R. K., Lopez, G., Feng, D., Alemu, L., Zhao, K., & Liu, P. P. # (2020). RUNX1 and CBFβ-SMMHC transactivate target genes together in abnormal myeloid progenitors for leukemia development. *Blood*, The Journal of the American Society of Hematology, 136(21), 2373-2385. (selected as Plenary Paper)
 - Han, M. *, Li, J. *, *Cao, Y.* *, Huang, Y., Li, W., Zhu, H., Zhao, Q., Han, J., Wu, Q., Li, J., Feng, J., & Wong, J. (2020). A role for LSH in facilitating DNA methylation by DNMT1

- through enhancing UHRF1 chromatin association. *Nucleic Acids Research*, 48(21), 12116-12134.
- Zeng, Y. *, *Cao, Y.* *, Halevy, R. S. *, Nguyen, P., Liu, D., Zhang, X., Ahituv, N., & Han, J. (2020). Characterization of functional transposable element enhancers in acute myeloid leukemia. *Science China Life Sciences*, 1-13.
- Huang, F. *, *Cao, Y.* *, Wu, G., Chen, J., Lin, W., Lan, R., Wu, B., Xie, X., Hong, J., & Fu, L. (2020). BMP2 signaling activation enhances bone metastases of non-small cell lung cancer. *Journal of Cellular and Molecular Medicine*, 24(18), 10768-10784.
- 2019 Cao, Y. *, Chen, G. *, Wu, G. *, Zhang, X. *, McDermott, J., Chen, X., Xu, C., Jiang, Q., Chen, Z., Zeng, Y., Ai, D., Huang, Y., &Han, J. # (2019). Widespread roles of enhancer-like transposable elements in cell identity and long-range genomic interactions. Genome research, 29(1), 40-52.
- Wang, L. *, Xu, X. *, Cao, Y. *, Li, Z., Cheng, H., Zhu, G., Duan, F., Na, J., Han, J., & Chen, Y. G. (2016). Activin/Smad2-induced H3K27me3 reduction is crucial to initiate mesendoderm differentiation of human embryonic stem cells. <u>Journal of Biological Chemistry</u>, jbc-M116.

OTHER PUBLICATIONS

- Roy, S., Ren, M., Li, P., Cui, K., *Cao, Y.*, Fisk, B., Markowitz, T. E., Redekar, N., Sakamoto, K., Nagao, K., Oh, J., Spolski, R., Liao, W., Dubois, S. P., Kelsall, B. L., Zhao, K., Phelan, J. D., & Leonard, W. J. (2025). BLIMP1 negatively regulates IL-2 signaling in T cells. *Science Advances*, 11(29), eadx8105.

 Khateb, M., Jung, R., Leibou, S., Hadley, P., Yu, Z., Dinerman, A. J., Dulemba, V., Gasmi, B., Levin, N., Kim, P., Bhasin, A., Bhat, D., Sindiri, S., Gartner, J. J., Prickett, T. D., Benzine, T., Farid, S. S., Parkhurst, M. R., Halas, H., *Cao, Y.,* Zhao, K., Yang, J. C., Robbins, P. F., Lowery, F., Krishna, S., Heller, T., McVicar, D., Rosenberg, S. A. & Klemen, N. D. (2025). Rapid enrichment of progenitor exhausted neoantigen-specific CD8 T cells from peripheral blood. *bioRxiv*, 2025-05.
- Zhu, X., Chen, X., *Cao, Y.*, Liu, C., Hu, G., Ganesan, S., Veres, T. Z., Fang, D., Liu, S., Chung, H., Germain, R. N., Schwartzberg, P. L., Zhao, K., & Zhu, J. (2024). Optimal CXCR5 Expression during Tfh Maturation Involves the Bhlhe40-Pou2af1 Axis. *bioRxiv*, 2024-05.
- Zhang, X., Jiang, Q., Li, J., Zhang, S., *Cao*, *Y*., Xia, X., Cai, D., Tan, J., Chen, J., & Han, J. D. J. (2022). KCNQ1OT1 promotes genome-wide transposon repression by guiding RNA–DNA triplexes and HP1 binding. *Nature Cell Biology*, 1-13. Lin, X., Liu, Y., Liu, S., Zhu, X., Wu, L., Zhu, Y., Zhao, D., Xu, X., Chemparathy, A., Wang, H., *Cao*, *Y*., Nakamura, M., Noordermeer, J., Russa, M., Wong, W., Zhao, K., & Qi, L. S. (2022). Nested epistasis enhancer networks for robust genome regulation. *Science*, 377(6610), 1077-1085.
 - Ren, G., Lai, B., Harly, C., Baek, S., Ding, Y., Zheng, M., *Cao, Y.*, Cui, K., Yang, Y., Zhu, J., Hager, G., Bhandoola, A., & Zhao, K. (2022). Transcription factors TCF-1 and GATA3 are key factors for the epigenetic priming of early innate lymphoid progenitors toward distinct cell fates. *Immunity*, 55(8), 1402-1413.
 - Pan, L., Ku, W. L., Tang, Q., *Cao*, *Y.*, & Zhao, K. (2022). scPCOR-seq enables coprofiling of chromatin occupancy and RNAs in single cells. *Communications Biology*, 5(1), 1-9.

- Fang, D., Cui, K., *Cao, Y.*, Zheng, M., Kawabe, T., Hu, G., Khillan, J., Li, D., Zhong, C., Jankovic, D., Sher, A., Zhao, K., & Zhu, J. (2022). Differential regulation of transcription factor T-bet induction during NK cell development and T helper-1 cell differentiation. *Immunity*, 55(4), 639-655.
- Ku, W. L., Pan, L., Cao, Y., Gao, W., & Zhao, K. (2021). Profiling single-cell histone modifications using indexing chromatin immunocleavage sequencing. Genome Research, 31(10), 1831-1842.
 Huang, F., Cao, Y., Wang, C., Lan, R., Wu, B., Xie, X., Hong, J., Fu, L., & Wu, G. (2021). PNMA5 promotes bone metastasis of non-small-cell lung cancer as a target of BMP2 signaling. Frontiers in Cell and Developmental Biology, 1400.
- Qiu, X., Ma, F., Zhao, M., *Cao, Y.*, Shipp, L., Liu, A., Dutta, A., Singh, A., Braikia, F.Z., De, S., Wood, W. H., Becker, K.G., Zhou, W., Ji, H., Zhao, K., Atchison, M.L., & Sen, R. (2020). Altered 3D chromatin structure permits inversional recombination at the IgH locus. *Science Advances*, 6(33), eaaz8850.
 Xia, X., Chen, X., Wu, G., Li, F., Wang, Y., Chen, Y., Chen, M., Wang, X., Chen, W., Xian, B., Chen, W., *Cao, Y.*, Xu, C., Gong, W., Chen, G., Cai, D., Wei, W., Yan, Y., Liu, K., Qiao, N., Zhao, X., Jia, J., Wang, W., Kennedy, B., Zhang, K., Cannistraci, C., Zhou, Y., & Han, J. (2020). Three-dimensional facial-image analysis to predict heterogeneity of the human ageing rate and the impact of lifestyle. *Nature Metabolism*, 2(9), 946-957.
- 2019 Chen, X., Xu, C., Hong, S., Xia, X., *Cao, Y.*, McDermott, J., Mu, Y., & Han, J. (2019). Immune cell types and secreted factors contributing to inflammation-to-cancer transition and immune therapy response. *Cell Reports*, 26(7), 1965-1977.
- 2018 Xu, C., Ai, D., Shi, D., Suo, S., Chen, X., Yan, Y., *Cao, Y.*, Zhang, R., Sun, N., Chen, W., McDermott, J., Zhang, S., Zeng, Y. & Han, J. (2018). Accurate drug repositioning through non-tissue-specific core signatures from cancer transcriptomes. *Cell Reports*, 25(2), 523-535.
- Chen, W., Qian, W., Wu, G., Chen, W., Xian, B., Chen, X., Cao, Y., Green, C.D., Zhao, F., Tang, K., & Han, J. (2015). Three-dimensional human facial morphologies as robust aging markers. Cell Research, 25(5), 574-587.
 Huang, Y., Yu, X., Sun, N., Qiao, N., Cao, Y., Boyd-Kirkup, J. D., Shen, Q., & Han, J. (2015). Single-cell-level spatial gene expression in the embryonic neural differentiation niche. Genome Research, 25(4), 570-581.
- 2014 Hong, S., Huang, Y., *Cao, Y.*, Chen, X., & Han, J. D. J. (2014). Approaches to uncovering cancer diagnostic and prognostic molecular signatures. *Molecular & Cellular Oncology*, 1(2), e957981.

SOFTWARES

2022-Curr. astroBoy

An automated, Al-assisted platform for large-scale next-generation sequencing data analysis, supporting both in-house and publicly deposited datasets on a

single-node server. Deployed at NHLBI/NIH, it has efficiently processed over 300 billion reads from more than 8,000 samples across 600+ analysis jobs since December 2022.

2023-Curr glitter

A custom-built, in-house web application for the analysis and visualization of single-cell RNA-seq data, developed to support collaborative research.

2023-Curr VINCE

A de novo Python package designed for nucleosome data analysis from single-cell MNase-seq, currently under active development and optimization.

2025 Ryder

A Python package for epigenome data normalization with internal reference and variable feature detection (manuscript submitting).

2021 <u>cLoops2</u>

A Python package for full stack analysis tool for chromatin interaction data such as Hi-TrAC, HiChIP and Hi-C (*Nucleic Acids Research*, 2022).

2019 TOWN

A Python package built on Keras for high-throughput, deep learning—augmented bright-field microscopy to automatically count live *C. elegans* worms. Developed for in-house use in drug screening to identify compounds that extend worm survival curves.

2017 cLoops

A Python package for accurate and flexible loops calling tool for 3D genomic data (*Bioinformatics*, 2020).

UNDER REVIEW/REVISE

2025 1. Guangzhe G

- 1. Guangzhe Ge *, **Yaqiang Cao** *, Danping Wei *, Mary Attaway, Barbara L Kee, Jinfang Zhu, and Keji Zhao *: Deciphering chromatin priming of T cell lineage by single-cell micrococcal nuclease sequencing
- 2. **Yaqiang Cao** **, Guangzhe Ge *, and Keji Zhao *: Ryder: Epigenome Normalization and Variable Feature Identification

AWARDS and HONORS

2024 NHLBI Director's Award for Innovation

Discovery of a novel regulatory element for IFN-y expression (Immunity, 2023).

MENTORING

2024,2023 Hannah Zhu

University of Maryland; NIH Summer Internship Program.

Collaborated on STARR-seq and Hi-TrAC data analysis to study enhancer—enhancer interactions.

2024 Emily Yu

The Bryn Mawr School; High School Senior Project.

Currently Yale University; mentored on single-cell RNA-seg analysis.

2019-2020 Jonathan Perrie

NIH Postbac Program. Co-mentored in bioinformatics.

Successfully enrolled into the bioinformatics Ph.D. program at UCLA.

2015-18 Yingying Zeng

Partner Institute for Computational Biology; Junior master's student. Co-mentored in bioinformatics and collaborated on research investigating the epigenetic features of MIR retrotransposons. Co-first-authored a publication and successfully enrolled into the Ph.D. program at Nanyang Technological

University (Singapore).

2016-18 Zhaoxiong Chen

Partner Institute for Computational Biology; Junior Ph.D. student. Co-mentored in bioinformatics and collaborated on research optimizing the

cDBSCAN algorithm and cLoops. Co-first-authored a publication.

2016-18 Daosheng Ai

Partner Institute for Computational Biology; intern. Co-mentored in

bioinformatics.

Co-first-authored a publication. Successfully enrolled into the Ph.D. program at

Peking University (China).

PEER REVIEWER

Bioinformatics, Communications Biology, BMC Genomics, F1000Research, Frontiers in Immunology, Frontiers in Bioinformatics, Genes, Cells etc.

TALKS

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- 2024 Bioinformatics and Scientific Programming Core, NICHD, Bethesda, USA
- 2024 Invited by Dr. Gordon L. Hager, NCI, Bethesda, USA
- 2023 Invited by Prof. Lei Hou, Boston University, Boston, USA
- 2023 DECODE Seminar Series, NIH, Bethesda, USA
- 2017 Epigenetics Retreat, Ningbo, China

CONFERENCE & POSTERS

- 2025 NIH/FDA Immunology Interest Group Retreat, DC, USA
- 2024 NCI Histone Modifications and Chromatin Structure Symposium, Bethesda, USA
- 2024 Epigenetics & Chromatin, Cold Spring Harbor Laboratory, New York, USA
- 2024 NHLBI Fellow Festival, NIH, Bethesda, USA

- 2024 NIH/FDA Immunology Interest Group Retreat, DC, USA
- 2023 Genome Architecture in Cell Fate and Disease, Ventura, California, USA
- 2023 NHLBI Fellow Festival, Bethesda, USA
- 2022 SCBA DC-Baltimore Chapter Annual Scientific Symposium, University of Maryland, USA
- 2019 4D Nucleosome Annual Meeting, DC, USA
- 2018 Chromatin, Epigenetics & Transcription, Suzhou, China
- 2017 First Chromatin Biology Conference, Shenzhen, China
- 2016 3rd International Symposium on 3D Genomics, Huazhong Agricultural University, Wuhan, China
- 2015 Otto Warburg Summer School, Max Planck Institute for Molecular Genetics, Berlin, Germany

TECHNICAL SKILLS

- Epigenomic, 3D Genome, and single-cell data analysis (Hi-C, ATAC-seq, ChIP-seq, MNase-seq, Hi-TrAC, RNA-seq, Micro-C, 10x single-cell RNA-seq, STARR-seq), with expertise in customized and sophisticated data mining approaches
- Proficient in Python programming and package development
- Experienced CPU/GPU servers setup and maintenance
- Web-application development
- Application and development of deep-learning and machine-learning algorithms

REFERENCE

Dr. Keji Zhao	Distinguished Investigator, NHLBI/NIH	<u>zhaok@nhlbi.nih.gov</u>
Dr. Paul P. Liu	Senior Investigator, NHGRI/NIH	pliu@nhgri.nih.gov
Dr. Chuan Wu	Senior Investigator, NCI/NIH	chuan.wu@nih.gov
Dr. Jingfang Zhu	Senior Investigator, NIAID/NIH	jfzhu@niaid.nih.gov