Curriculum Vitae Updated Apr. 2025

# **Xin Dong**

#### Ph.D. in Bioinformatics

NO.1239 Siping Road, School of Life Sciences and Technology, Tongji University, Shanghai, China 

■ dongxin@tongji.edu.cn; xindongnov@gmail.com | Homepage | Scholar

## **EDUCATION**

Sep. 2018 – Jul. 2024 Ph.D. in Bioinformatics

School of Life Sciences and Technology, Tongji University, China

Advisors: Dr. Chenfei Wang & Dr. Xiaole Shirley Liu

Sep. 2014 – Jun. 2018 B.S. in Biotechnology

College of Life Sciences, Shandong Normal University, China

## **RESEARCH EXPERIENCES**

Graduate Student, Tongji University (2018-2024)

Developed and applied computational methods to analyze single-cell and spatial data
 Advisors: Dr. Chenfei Wang & Dr. Xiaole Shirley Liu

- o I developed SCRIP (Single-Cell Regulatory Network Inference using ChIP-seq and motif) (https://github.com/wanglabtongji/SCRIP), a method that integrates scATAC-seq and a large collection of bulk TR (Transcription Regulator) ChIP-seq to infer single-cell TR activity and targets. The method outperformed other motif-based approaches in assessing TR binding activity. Applying SCRIP to various biological systems shows its utility in lineage tracing, cell-type clustering, and inferring cell-type-specific GRNs. With the advances of single-cell multiomics techniques, we updated it to SCRIPro (https://github.com/wanglabtongji/SCRIPro), which suits both single-cell and spatial multiomics data.
- We developed TISCH (Tumor Immune Single Cell Hub; <a href="http://tisch.compbio.cn">http://tisch.compbio.cn</a>) and TISCH2, a large curated resource that integrates the single-cell transcriptome profiles of about 6 million cells from 190 high-quality tumor datasets across 50 cancer types. TISCH allows systematic comparison of gene expression across multiple datasets at the single-cell or cluster level for various cell types, patients, tissue origins, treatment and response groups, and cancer types, aiming to characterize the immune system heterogeneity in cancers and inform immunotherapy.
- Large-scale genomic data mining and integration analysis

Advisors: Dr. Xiaole Shirley Liu & Dr. Clifford Meyer

- To download the raw sequencing data of the chromatin occupancy experiments from the NCBI's GEO database and extract the metadata, we created a parser. The parser annotated and labeled the experiment type, tissue type, cell population, and other details of the experiments using the keywords-match strategy.
- We developed CHIPS (CHromatin enrIchment ProceSsor) (<a href="https://github.com/liulab-dfci/CHIPS">https://github.com/liulab-dfci/CHIPS</a>), a pipeline that processes the epigenome data to readable results with a single command line. It integrates states-of-the-art tools for cistrome NGS data and chooses the appropriate analysis parameters. It uses the Snakemake framework, which makes it easy to deploy on cloud or cluster servers.
- We developed and maintained the Cistrome Data Browser (<a href="http://cistrome.org/db/#/">http://cistrome.org/db/#/</a>), which hosts many public cistrome sequence datasets. It offers a user-friendly interface to access the data quality and the results of downstream analysis.

Undergraduate Student, Shandong Normal University (2014-2018)

- Rapid establishment of H7N9 dominant lineage by intragenic recombination in HA segment Undergraduate Dissertation Research; Advisor: Dr. Chengqiang He
- Screening of newcastle disease virus heat-resistant vaccine by homologous recombination Undergraduate Innovation Program (Presided); Advisor: Dr. Chengqiang He

Curriculum Vitae Updated Apr. 2025

• The origin, adaptive evolution and population dynamics of infectious Chicken Anemia Virus in China Undergraduate Innovation Program (Involved); Advisor: Dr. Chengqiang He

Diversity of fish in Nansi Lake
 Undergraduate Innovation Program (Involved); Advisors: Dr. Rongshu Fu & Dr. Mingsheng Miao

## **SKILLS**

- Bioinformatics: Single-cell/spatial RNA/ATAC-seq analysis; Epigenome data analysis (ChIP-seq, ATAC-seq, etc.); Machine learning (PyTorch, scikit-learn)
- Programming: Python; R; Shell; JavaScript; MySQL
- Tools & Frameworks: Git; Django; Vue.js
- Cloud & HPC: Snakemake; AWS/GCP computation
- Server operation and maintenance: Ubuntu; CentOS / Rocky Linux

#### **PUBLICATIONS**

Asterisk (\*) denotes the Co-first Authors and Hashtag (#) denotes the Corresponding Authors.

- Chang, Z.\*, Xu, Y.\*, <u>Dong, X.\*</u>, Gao, Y. & Wang, C.# Single-cell and spatial multiomic inference of gene regulatory networks using SCRIPro. *Bioinformatics* 40, btae466 (2024).
- Sun, F.\*, Li, H.\*, Sun, D.\*, Fu, S.\*, Gu, L.\*, Shao, X.\*, Wang, Q.\*, <u>Dong, X.\*</u>, Duan, B.\*, Xing, F.\*, Wu, J.\*, Xiao, M.#, Zhao, F.#, Han, J.-D. J.#, Liu, Q.#, Fan, X.#, Li, C.#, Wang, C.# & Shi, T.# Single-cell omics: experimental workflow, data analyses and applications. SCIENCE CHINA Life Sciences (2024).
- Ren, P.\*, Shi, X.\*, Yu, Z., <u>Dong, X.</u>, Ding, X., Wang, J., Sun, L., Yan, Y., Hu, J., Zhang, P., Chen, Q., Zhang, J., Li, T. & Wang, C.# Single-cell assignment using multiple-adversarial domain adaptation network with large-scale references. *Cell Reports Methods*, 3, 100577 (2023).
- Han, Y.\*, Wang, Y.\*, <u>Dong, X.\*</u>, Sun, D., Liu, Z., Yue, J., Wang, H., Li, T.# & Wang, C.# TISCH2: expanded datasets and new tools for single-cell transcriptome analyses of the tumor microenvironment. *Nucleic Acids Research*, 51, D1029-D1037 (2022).
- Shi, X.\*, Yu, Z.\*, Ren, P., <u>Dong, X.</u>, Ding, X., Song, J., Zhang, J., Li, T.# & Wang, C.# HUSCH: an integrated single-cell transcriptome atlas for human tissue gene expression visualization and analyses. *Nucleic Acids Research*, 51, D1425-D1431 (2022).
- <u>Dong, X.\*</u>, Tang, K.\*, Xu, Y., Wei, H., Han, T. & Wang, C. # Single-cell gene regulation network inference by large-scale data integration. *Nucleic Acids Research*, 50, e126 (2022).
- Xu, R.\*, Li, S.\*, Wu, Q.\*, Li, C.\*, Jiang, M.\*, Guo, L., Chen, M., Yang, L., <u>Dong, X.</u>, Wang, H., Wang, C.#, Liu, X.#, Ou, X.# & Gao, S.# Stage-specific H3K9me3 occupancy ensures retrotransposon silencing in human pre-implantation embryos. *Cell Stem Cell*, 29, 1051-1066.e8 (2022).
- Sun, D.\*, Wang, J.\*, Han, Y.\*, <u>Dong, X.</u>, Ge, J., Zheng, R., Shi, X., Wang, B., Li, Z., Ren, P., Sun, L., Yan, Y., Zhang, P., Zhang, F.#, Li, T.# & Wang, C.# TISCH: a comprehensive web resource enabling interactive single-cell transcriptome visualization of tumor microenvironment. *Nucleic Acids Research*, 49, D1420-D1430 (2021).
- Zheng, R.\*, <u>Dong, X.\*</u>, Wan, C., Shi, X., Zhang, X.# & Meyer, C.A.# Cistrome Data Browser and Toolkit: analyzing human and mouse genomic data using compendia of ChIP-seq and chromatin accessibility data. *Quantitative Biology*, 8, 267-276 (2020).
- Chen, C.-H., Zheng, R., Tokheim, C., <u>Dong, X.</u>, Fan, J., Wan, C., Tang, Q., Brown, M., Liu, J.S., Meyer, C.A. # & Liu, X.S.# Determinants of transcription factor regulatory range. *Nature Communications*, 11, 2472 (2020).
- Li, S.\*, Wan, C.\*, Zheng, R., Fan, J., **Dong, X.**, Meyer, C.A.# & Liu, X.S.# Cistrome-GO: a web server for functional enrichment analysis of transcription factor ChIP-seq peaks. *Nucleic Acids Research*, 47, W206–W211 (2019).

#### **CONFERENCES**

Curriculum Vitae Updated Apr. 2025

- 2024 Poster Speaking (Awarded), The 2<sup>nd</sup> Bioinformatics Conference of Yangtze River Delta, Shanghai, China
- 2023 Poster Speaking, Systems biology of gene regulation and genome editing, Cold Spring Harbor Conferences Asia, Suzhou, China
- 2023 Poster Speaking, The WLA Research Conference on Cells and Genes, Shanghai, China
- 2023 Poster Speaking, The 11st National Conference on Bioinformatics and Systems Biology, Guangzhou, China
- 2023 Poster Speaking, Human Cell Atlas Asia 2021 Meeting, online
- 2021 Poster Speaking, The 10th National Conference on Bioinformatics and Systems Biology, Chengdu, China
- 2019 Attendee, 14th International Bioinformatics Workshop, Beijing, China

#### **TEACHING**

- 2023 Teaching assistant, Summer Deep Learning Course, School of Life Sciences and Technology, Tongji University, Shanghai, China
- 2022 Teaching assistant, Summer Bioinformatics Course, School of Life Sciences and Technology, Tongji University, Shanghai, China
- 2021 Teaching assistant, Summer Bioinformatics Course, School of Life Sciences and Technology, Tongji University, Shanghai, China
- 2020 Teaching assistant, Summer Bioinformatics Course, School of Life Sciences and Technology, Tongji University, Shanghai, China
- 2019 Teaching assistant, Dragon Star Course, The 14th International Bioinformatics Workshop, Beijing, China

### AWARDS AND DISTINCTIONS

- 2024 Outstanding Graduates Awards of Shanghai, Shanghai Municipal Education Commission
- 2023 The 17th Academic Pioneer (10 of the University), Tongji University
- 2023 National Scholarship, Ministry of Education of the People's Republic of China
- 2022 Outstanding Student Leader, Tongji University
- 2019 Outstanding Graduate Student Scholarship, Tongji University
- 2018 Outstanding Undergraduate Thesis for Bachelor's Degree, Shandong Normal University
- 2017 Outstanding Graduates Awards of Shandong, Human Resources and Social Security Department of Shandong Province
- 2017 The First Prize of the 15<sup>th</sup> Challenge Cup Competition in Shandong, Shandong Association for Science & Technology
- 2017 Outstanding Student Leader of Shandong, Shandong Provincial Education Department
- 2016 The Grand Prize of Shandong College Students' Biochemistry Experiment Skills Competition, Shandong Association of Science & Technology