



Qiao Fang

Machine Learning Scientist

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WORK EXPERIENCE

Machine Learning scientist, Fable Therapeutics

Jan 2025 - Present

- Managed and built a diffusion-based model for full-atoms antibody generation and built a side-chain packer that achieves state-of-the-art 0.35 Å sidechain RMSD.
- Developed an Equiformer-based confidence model to assess fully designed antibody-antigen complexes, proving to eliminate 80 % of low-quality candidates.
- Streamlined workflows by unifying datasets, modularizing code, and containerizing models—cutting setup time 40% and duplicate work.

Machine learning associate scientist, Fable Therapeutics

Oct 2022 - Jan 2025

- Built a diffusion-based model that folds complete antibody structures while simultaneously generating all CDR loops bound to specified targets achieving an average CDR RMSD of 2.83 Å.
- Built an H3-CDR co-design model that optimizes structure and sequence simultaneously, reaching state-of-the-art 1.37 Å RMSD and 56.7 % sequence recovery.
- Devised a comprehensive workflow for the systematic processing of protein and antibody datasets, encompassing data collection, curation, feature extraction, and integration, to provide a robust foundation for antibody predictive modeling

EDUCATION

M.Sc. in Department of Molecular Genetics

Sept 2020 - Sept 2022

University of Toronto

- Publications: [Kim J, McFee M, Fang Q, et al. Computational and artificial intelligence-based methods for antibody development\[J\]. *Trends in pharmacological sciences*, 2023, 44\(3\): 175-189.](#)

PhD candidate in Academy for Advanced Interdisciplinary Studies

Peking University

Sept 2017 - Aug 2020

- Publications:

[Zhang L, Li Z, Skrzypczynska K M, et al. Single-cell analyses inform mechanisms of myeloid-targeted therapies in colon cancer\[J\]. *Cell*, 2020, 181\(2\): 442-459. e29.](#)

[Zhang L, Yu X, Zheng L, et al. Lineage tracking reveals dynamic relationships of T cells in colorectal cancer\[J\]. *Nature*, 2018, 564\(7735\): 268-272.](#)

B.Sc. in School of Life Sciences

Sept 2013 - Sept 2017

Peking University

KEY SKILLS

- Python
- Bash
- Javascript
- Git
- AWS
- Docker
- Pytorch
- Tensorflow
- Deep learning
- Machine learning
- Statistical learning
- English and Mandarin