

XIANG ZHANG

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Drug Discovery and Design Center, 647 Songtao Road, Shanghai, China

EDUCATION

Shanghai Institute of Materia Medica & NJUCM	Sept. 2021 - Jul. 2025
M.Pharm. in Drug design, GPA: 3.8/4 Advisor: Prof. Dr. Mingyue Zheng and Dr. Xutong Li	
Henan University	Sept. 2017 - Jul. 2021
B.S. in Pharmaceutics, GPA: 3.3/4	

RESEARCH INTEREST

AI for synthesis: Synthetic Accessibility Prediction & Retrosynthesis Planning.

Molecular property prediction: Drug's ADMET prediction, Uncertainty estimation.

Molecule generation: Multi-objective RL optimization, Multi-modal conditional generation

RESEARCH EXPERIENCE

Synthetic Accessibility Prediction	Jul. 2023 - Feb. 2025
<ul style="list-style-type: none">Developed SynFrag, a fragment assembly generation model for SA prediction, designed by simulating the stepwise assembly of building blocks in synthesis.Curated 9.2M pre-training and 800k fine-tuning dataset, contributing two application scenario test sets. SynFrag achieved SOTA while demonstrating chemical interpretability.Deployed SynFrag online service: sub-second & high-throughput & interpretable prediction support for in-silico to in-lab drug design.	
Molecular Property Prediction	Oct. 2024 - present
<ul style="list-style-type: none">Develop BioCLIP, a progressive multi-modal bootstrapping framework that address the scarcity of tri-modal data through hierarchical training, for diverse molecular property predictions. [Code]Integrate 1.3K tri-modal samples (Molecules, Gene Expression, Cell Painting) with 30K bi-modal pairs; using cross-modal synthesis to generate 6K high-confidence pseudo tri-modal samples.	
Organic Synthesis	Jan. 2022 - Sept. 2022
<ul style="list-style-type: none">Synthesis, Purification, and Analysis of PROTAC Intermediates in Prof. Chen's Lab.	

PUBLICATIONS

- Zhang, X.**, Liu, J., **Chen, K.** ([Academician](#)). SynFrag: Synthetic Accessibility via Fragment Assembly Generation *J. Chem. Inf. Model.*, peer review. [\[SynFrag\]](#) | [\[Code\]](#) | [\[PDF\]](#)
- Fan, Z., Yu, J., **Zhang, X.** Reducing overconfident errors in molecular property classification using Posterior Network. *Patterns*, 2024, **5**(6): 100991. [\[Code\]](#) | [\[PDF\]](#)

SKILLS

- Programming/Software: Python, HTML, CSS, L^AT_EX, Prism, ChemDraw, PyMOL, Matplotlib
- Cheminformatics: Pytorch, RDkit, DGL, deepchem, sklearn, TensorBoard, GNN, Transformer
- in Laboratory: Organic synthesis, purification & analysis
- Language: English (fluent, CET6: 520), Chinese (native), Cantonese (native), Hakka (native)

CONFERENCE

- Oral Presentation: The 13th Shanghai Symposium on Computer-Aided Drug Design | 2024
- Participation: World Artificial Intelligence Conference (WAIC), Shanghai | 2022, 2023, 2025

AWARDS & HONORS

- Academic Excellence Scholarship | 2021 - 2024
- Certificate of Honor for volunteer in COVID-19 Prevention | 2022