martina-yu.github.io

RESEARCH INTERESTS

Single cells & bulk genes, Crispr/Cas9 system, Gene editing, Spatial Transcriptomics, Cell-Cell interation

Graph Neural Networks & Machine Learning & Bioinformatics & Computational Biology

EDUCATION

• School of Medicine, Shanghai Jiao Tong University

Double Major: Preventive Medicine & Public Policy; GPA: 3.64(3.91/4 for last 3.5 years)

Shanghai, China

Sep 2020 - Jun 2025

Email: yuzimeng at gmail.com

Mobile: $+1\ 203-410-8140$

• KCL 2021 Summer School of Public Health

London, UK

• The Global Summer School 2022 of Tsinghua Vanke School of Public Health

Beijing, China

• A Courses: Over 43 of all 90 courses, including Statistics (Score: 91), R programming(93), Basic Calculus(96), Elementary Chemistry C(A-), Ogranci Chemyrsti C(A-), Machine Learning(89.1), Markov and Decision-trees(90)

RESEARCH EXPERIENCE

• Generate Chat-AI of Radiation Biology by Multi-model Team Member

Shanghai Jiao Tong University

Jun 2024 - Present

• Annotation: Label-Studio, configuring plantform for image labeling and annotation, which help doctors labeling radiology images manually.

• **Prompt Engineering**: Python and Markdown, designed and ensure exact and clear prompt for chat-bot to generate radiation biology text information according to input image

• Deep Learning for Detect Cell-Cell Communication

Yale University

 $Research\ Assistant$

Feb 2024 - Present

- Making Protocols and Strategies for this Project: Collect relative thesis and datas, then helping design some of the detailed experiments and prospectives of this project. Also helping other students who are interseted in it with their questions and problems.
- Preprocessing Datasets and Databases: Pytorch and R, preprocessing our raw datasets over 1k genes of single cells from collaborated biology laboratory and also public databases which includes MERFISH, STARmap and 10x Visium HD. Also handling on our referenced database NicheNet with their ligand-target datas.
- **Biological Hyperparameter Detecting**: Pytorch and Linux, figuring out the best biological hyperparameter of cells' diffusion diameter and activated communication with the least loss via adding model panelty and automated fine-tuning parameters of models.
- Predict Base Editor Editing Outcomes with Deep Learning Co-Leader

Shanghai Jiao Tong University Sep 2023 - Present

- Data Preprocessing: Linux, mapping designed library with raw mutated sequencing datas with tool bwe-mem2, calculating editing efficiency and outcomes proportion to generate processed data for training and testing.
- Data Analysis and Descriptive Statistics: Python and Matlab, describing and summarizing processed datasets including editing windows across 20bp protospacer region, mutation trends of single mutated position, effects of 1-bp and whole region context of targeted editing efficiency.
- Benchmark and Discuss Model Development: Jupyter Notebook, TensorFlow and Linux, processing and analysising raw datasets from 4 other works and configuring benchmark models in HPC for comparison, including AUC scores, Spearman and Pearson correlation.
- **Defining Class and Embedding**: Python, defining classes of somple, processing embedded inputs with embeddingbag method, then generating batchs.
- Building and Finetuning Model: Python, Linux, TensorFlow, building LSTM model mainly with additional attention layers, then use ray and cross-validation K-Fold and stratifiedkfold to finetune best parameters until the best performance of prediction.
- The First Global AI Drug Development Competition

Shanghai Jiao Tong University

May 2023 - Jul 2023

Team Member

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- Preprocessing: Processed and curated provided datasets to ensure high-quality inputs for predictive modeling.
- Benchmark Model: Collaborated with team members to analyze and compare model performance metrics.
- Competition Organization: Successfully uploaded and submitted competition results, achieving TOP 20%

• Cancer Integrative Analysis Via Multimodal Machine Learning

Shanghai Tong Ren Hospital

Student Researcher

Dec 2022 - May 2023

- Reproduce Research Paper: Utilized TCGA and cBioPortal databases for comprehensive genomic data retrieval and integration, reproduct the datasets in the research paper and get accurate and consistent results
- Visualization: Executed Whole Slide Image (WSI) analysis, contributing to the large-scale pathology datasets.

SKILLS

- Programming Languages: Python, SPSS, R, C, /LaTeX, Linux Operation System
- Languages: Chinese (Native), English (Professional), Japanese (Limited)
- Research Interests: Go to personal website

Honors

- 2022-2023 A Student Scholarship, School of Medicine (TOP 1 Student)
- 2023-2024 B Student Scholarship, School of Medicine (TOP 5 Student)