

# Francesca-Zhoufan Li



## AI for Science & Engineering

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- US permanent resident, Shanghainese American

## Goal

Determined to advance the intersection of ML and science with a protein engineering focus, a computational and experimental hybrid background, and an innovative interdisciplinary collaborative growth mindset

## Skills & Tools

- Python, Matlab, R, LaTeX, Git
- Bash, HPC, BLAST, ClustalW
- Pandas, ESM, CARP, Numpy
- Scikit-Learn, PyTorch, Biopython
- Data analysis, Machine learning
- Biotechnology, Protein engineering
- Analytical chemistry, Benchwork
- Research, Active Learning
- Leadership, Collaboration
- Proposal writing, Presentation
- Teaching, Mentoring, Outreach

## Talks & Posters

- SynBioBeta, 2023
- Caltech Bioscience Futures Day, 2023
- Seagate-Minnesota AI/ML Virtual Distinguished Speaker Series, 2023
- Google Research invited talk, 2022

## Other Experiences

- Code with Young Legends: led intro to coding workshop
- i-STEM: mentored under-invested high school students on Twitter terrorist indicators computational project
- Bioengineering Honor Society: mentored high school bioengineering research competitions, with one of the teams winning a 2nd place
- Biotech Connection Los Angeles: to grow the local biotech landscape
- Biology Scholars Program: to challenge who can do STEM

## Education

- 09/20-Present **Ph.D. in Bioengineering, GPA: 4.0** California Institute of Technology
  - NSF Graduate Research Fellowships Program • Amazon AI4Science Fellowship • Biotechnology Leadership Training Program
- 08/15-05/19 **B.S. in Bioengineering, GPA: 3.96** University of California, Berkeley
  - B.S. in Chemical Biology**
    - Highest Honors • Jack & Birthe Kirsch Prize • Tau Beta Pi Scholarship
    - John Gorton Davis Scholarship • T. Dale Stewart Scholarship
    - Genentech Outstanding Student Award Runner-Up

## Industry & Academic Experience

- 06/22-09/22 **BioML Research Intern** Microsoft Research
  - Developed protein transfer learning pipeline to systematically evaluate pre-trained transformer and convolutional models for 7 different downstream practical engineering and annotation tasks
  - Utilized and optimized computing resources for execution
  - Delivered talks and engaged in professional development
- 01/21-Present **Machine Learning for Proteins** Arnold Lab & Yue Group, Caltech
  - Developing multi-modal (ie. sequence, structure) representation learning pipeline for protein fitness (ie. binding, catalysis) prediction
  - Building easy-to-use protein single- to multi-mutant fitness prediction by incorporating mutation sequence context (coevolution, stability, and biochemical rules) on 11 diverse datasets
  - Led and facilitated 3 grant writing and cross-group collaborations
- 09/20-01/21 **Extremophile Genetic Component Discovery** Murray Lab, Caltech
  - Constructed an RNA-seq analysis pipeline in R to discover novel genetic circuit components in non-canonical cell-free extracts
  - Delivered results to groups at Caltech, the U.S. Army Chemical Biological Center, and the Imperial College London
- 06/20-08/20 **RNA-Seq Sample Preparation Pipeline Optimization** Zymergen
  - Developed a Python package to design DNA oligos for RNaseH-based ribosomal RNA depletion for 8 strains in 7 programs
  - Wrote R scripts to quality control and pre-process RNA extraction data from industry-standard electrophoresis instruments
  - Delivered talks, collaborated across and outside of the company
- 06/19-05/20 **Bioinformatics Tool Development** Koide Lab, NYU Langone Health
  - Developed Matlab software for SARS-CoV-2 mutation analysis from GISAID database covering 25k global sequence entries
  - Wrote easy-to-use Matlab scripts to identify monobody and antibody complementary-determined region mutations for protein engineering
  - Automated chromatogram visualization with user-chosen features
- 05/18-07/18 **Cell-Free Platform Streamline** Tierra Biosciences, QB3 Program
  - Optimized non-standard protein production in cell-free expression systems with Design Of Experiments methodology
- 01/16-05/19 **Independent Bioengineering Researcher** Dueber Lab, UC Berkeley
  - Automated time-course betaxanthin production analysis in Matlab
  - Engineered yeast to increase benzylisoquinoline alkaloids yield
  - Improved beta-glucosidase stability & activity in a basic solution for indigo bio-production in *E. coli* via error-prone PCR libraries
  - Honor thesis: A "Microbial Factory" Toolkit: Yeast Spheroplast Transformation Method Development for CRISPR-Cas9 Multiplexing

## Publications

- 2023 Yang J, Ducharme J, Johnston KE, Li F-Z, et al. DeCOIL: Optimization of Degenerate Codon Libraries for Machine Learning-Assisted Protein Engineering. *bioRxiv*.
- 2022 Li F-Z, et al. Pretrained protein language model transfer learning: is the final layer representation what we want? *NeurIPS 2022 Workshop*.
- 2021 Koide A, Panchenko T, Wang C, Thannickal SA, Romero LA, Teng KW, Li F-Z, et al. Two-dimensional multiplexed assay for rapid and DEEP SARS-COV-2 serology profiling and for machine learning prediction Of Neutralization capacity. *bioRxiv*.
- 2019 Savitskaya J, Protzko J, Li F-Z, et al. Iterative screening methodology enables isolation of strains with improved properties for a FACS-based screen and increased L-DOPA production. *Sci.Rep.* **9**