

# Fangfei Li

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## SUMMARY & SKILLS

- **Ph.D. in Applied Mathematics & Statistics** | 2 yrs postdoc + 2 yrs biotech startup + 3 yrs wet-lab (synthetic barcoding-based *in vivo* lineage tracing)
- **Skills:** Bayesian optimization, machine learning (deep generative model), data analysis & visualization, Python (NumPy, pandas, Matplotlib, Flet), Git, SQL
- Strong analytical and critical thinking; detail-oriented, highly organized, goal-driven

## INDUSTRIAL & ACADEMIC EXPERIENCE

**AI Training with Certificates** 9/2025-current

*Online*

- Artificial Intelligence Professional Program (Stanford): XCS236 - Deep Generative Models (Certificate)

**Staff Scientist (Computational Biology)** 11/2023-8/2025

BacStitch DNA, South San Francisco, CA

- Developed an automated end-to-end DNA assembly workflow (**Python, web/GUI**) that scores sequences, designs optimal overlapping blocks, checks synthesizability via supplier APIs, and generates automated high-throughput experimental plans.
- Built a **Python tool for optimal DNA block design** that segments sequences into overlapping blocks under customizable constraints (block count, block/overlap length, and specific regions as blocks/overlaps) while minimizing structural features in overlaps.
- Created **Python visualization tools** to (i) plot BLAST results and structural scores ([Example](#)), (ii) generate .ab1 files encoding structural scores as ATGC traces at single-nucleotide resolution, and (iii) produce publication-quality figures for publications and internal use.

**Postdoctoral Research Fellow** 4/2021-10/2023

Department of Genetics, Stanford University, CA

- Developed a Bayesian optimization algorithm and Python pipeline ([FitMut2](#)) to identify genome-wide adaptive lineages in experimental evolution from fitness phenotype (without WGS), and estimate their fitness effects and occurrence times.
- Developed an optimization algorithm to segment DNA sequences into overlapping fragments that minimize overlap-region scores (e.g., undesirable DNA structural features) under fixed or ranged constraints on fragment count, and fragment/overlap length.

**Artificial Intelligence Fellow** 9/2019-10/2019

Insight Data Science, San Francisco, CA

- Developed a deep learning pipeline for portrait editing (a 6-week project), providing continuous adjustment of facial features.

**Graduate Researcher** 8/2014-12/2020

Laufer Center for Physical and Quantitative Biology, Stony Brook University, NY

- Developed a fitness estimation tool ([Fit-Seq](#)) that can estimate unbiased fitness for complex cell pools, using a maximum likelihood algorithm.
- Designed and conducted *in vivo* bar-seq experimental yeast evolution (yeast genetics, cloning, barcoded library construction, lineage-traced evolution, and NGS prep). Per-

formed computational analysis (NGS processing, mathematical modeling to infer evolutionary parameters from lineage dynamics, and evolutionary simulations for validation) to explore the epistasis of adaptive mutations in yeast evolution.

**Graduate Researcher** ————— **9/2010-6/2013**

Chinese Academy of Sciences, Academy of Mathematics and Systems Science, China

- Established (1) a consensus criterion for second-order multi-agent systems (MASs) with switching topology and communication delays, and (2) a cluster consensus criterion for Boolean MASs with fixed or switching topologies.

**EDUCATION**

**Ph.D., Applied Mathematics and Statistics (Computational Biology)** — **8/2013-12/2020**

Stony Brook University, Stony Brook, NY

**M.S., Systems Theory** ————— **9/2010-7/2013**

University of Chinese Academy of Sciences, Beijing, China

**B.S., Mathematics and Applied Mathematics** ————— **9/2006-7/2010**

Beijing Normal University, Beijing, China

**PRESENTATIONS**

**Plenary Talk: Fitness Estimation of Pooled Amplicon Sequencing Studies** ————— **8/2018**

2018 Yeast Genetics Meeting, Stanford CA, 8/22/2018-8/26/2018

**PUBLICATIONS**

- [1] T Matsui, P-H Hung, H Mei, X Liu, **F Li**, J Collins, W Li, D Miller, N Wilson, E Toro, G J Taghon, G Sherlock, S Levy. **High-throughput DNA engineering by mating bacteria**. Preprint (2024).
- [2] **F Li**, A Mahadevan, G Sherlock. **An improved algorithm for inferring mutational parameters from Bar-seq evolution experiments**. *BMC Genomics* 24, 246 (2023).
- [3] **F Li\***, J Tarkington\*, G Sherlock. **Fit-Seq-2.0: an improved software for high throughput fitness measurements using pooled competition assays**. *Journal of Molecular Evolution* 91, 334–344 (2023).
- [4] G Avecilla, J N Chuong, **F Li**, G Sherlock, D Gresham, Y Ram. **Neural networks enable efficient and accurate simulation-based inference of evolutionary parameters from adaptation dynamics**. *PLoS Biology* 20, e3001633 (2022).
- [5] Z Liu, D Miller, **F Li**, X Liu, S Levy. **A large accessory protein interactome is rewired across environments**. *eLife* 9, e62365 (2020).
- [6] X Liu, Z Liu, A K Dziulko, **F Li**, D Miller, R D Morabito, D Francois, S Levy. **iSeq 2.0: a modular and interchangeable toolkit for interaction screening in yeast**. *Cell Systems* 8, 338–344 (2019).
- [7] **F Li**, M L Salit, S Levy. **Unbiased fitness estimation of pooled barcode or amplicon sequencing studies**. *Cell Systems* 7, 521–525 (2018).
- [8] I Frumkin, D Schirman, A Rotman, **F Li**, L Zahavi, E Mordret, O Asraf, S Wu, S Levy, Y Pilpel. **Gene architectures that minimize cost of gene expression**. *Molecular Cell* 65, 142–153 (2017).
- [9] Y Chen, W Xiong, **F Li**. **Convergence of infinite products of stochastic matrices: a graphical decomposition criterion**. *IEEE Transactions on Automatic Control* 61, 3599–3605 (2016).
- [10] Y Chen, **F Li**, B Hou, S Tan, H Zhu. **Convergence analysis of discrete-time consensus algorithm with both self and transmission delays**. *Journal of the Franklin Institute* 353, 2467–2481 (2016).
- [11] **F Li**, Y Chen, J Lü, D Hill. **Cluster consensus of Boolean multi-agent systems**. *2013 9th Asian Control Conference (ASCC)* (2013).
- [12] Y Chen, W Yu, **F Li**, S Feng. **Synchronization of complex networks with impulsive control and disconnected topology**. *IEEE Transactions on Circuits and Systems II: Express Briefs* 60, 292–296 (2013).