

# 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

- Designed and implemented the company's DNA assembly computational workflow, a modular Python pipeline (web/GUI) to: (1) score sequences by structural features, (2) segment sequences into overlapping blocks that minimizes unfavorable structural features in overlaps, (3) query supplier **APIs** for block synthesizability and order processing, (4) generate streamlined experimental plans for automated high-throughput assembly.
- Developed a Python pipeline to segment DNA sequences into optimal overlapping blocks that minimizes structural score of the overlaps, under customizable constraints on block count/length, overlap length, and inclusion of specific regions as block or overlap.
- Created Python tools for visualizations, including: (1) Generate figures visualizing BLAST results and DNA structural scores ([Example](#)). (2) Generate .ab1 files encoding up to four structural scores (as ATGC traces) at single-nucleotide resolution. (3) Make publication-quality figures for research papers and internal documentation.

### 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 multi-dimensional data (e.g. DNA sequences) into optimal overlapping fragments by minimizing total scores (e.g. undesirable DNA structural features) in overlap regions, under fixed or ranged constraints on fragment count, fragment length, and 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). Performed 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 Research** ————— 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**<sup>\*</sup>, J Tarkington<sup>\*</sup>, 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).