# **BRIAN HIE**

brianhie@stanford.edu | github.com/brianhie | brianhie.com

Interests: Machine learning, computational biology, protein engineering and evolution, systems biology.

### **CURRENT POSITIONS**

Stanford University School of Medicine, Palo Alto, CA

Stanford Science Fellow

Machine learning at the host-pathogen interface.

Meta Platforms, Inc., Menlo Park, CA 2022-Present

2021-Present

Visiting Researcher, Meta Al

• Machine learning for protein biology.

### **EDUCATION**

Massachusetts Institute of Technology, Cambridge, MA

Electrical Engineering and Computer Science, Doctor of Philosophy 2017-2021

Electrical Engineering and Computer Science, Master of Science

• Areas of concentration: Computational biology, machine learning, statistics.

Stanford University, Palo Alto, CA

Computer Science, Bachelor of Science with Honors and Distinction 2012-2016

English Literature, Academic Minor

• Areas of concentration: Computational biology, computer systems, machine learning.

#### PROFESSIONAL EXPERIENCE

Massachusetts Institute of Technology, Cambridge, MA

### Graduate Researcher, Computer Science and Artificial Intelligence Laboratory 2017-2021

- Neural language modeling of viral evolution.
- Machine learning for biological discovery and design under uncertainty.
- Efficient geometric algorithms for single-cell biology.
- Cryptographically secure neural network training.

Google LLC, Mountain View, CA

# Artificial Intelligence/Machine Learning Resident, X – The Moonshot Factory 2019

• Machine learning for early-pipeline moonshots.

Illumina, Inc., San Diego, CA

### Machine Learning Intern, Bioinformatics 2018

Statistical signal processing for genomics-based health monitoring.

Salesforce.com, Inc., San Francisco, CA

### Software Engineer, Cloud Infrastructure 2016-2017

• Robust performance monitoring of globally distributed core application infrastructure.

Stanford University, Palo Alto, CA

# *Undergraduate Researcher, Biology* 2013-2016

Statistics and machine learning for computational genomics.

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Stantord	University.	Paio	AILO.	CA

## Undergraduate Researcher, Digital Humanities, Stanford Humanities Center

• Graph-theoretic analysis of the social network of early modern authors and publishers.

# Microsoft Corporation, Redmond, WA

### Software Engineering Intern, Azure Compute and Microsoft Research

2015

2016

Distributed scheduling algorithms and their impact on data center utilization and availability.

Synaptics, Inc., San Jose, CA

# Systems Architecture/Algorithms Intern

2014

• Algorithm design and implementation for embedded touchscreen firmware.

### **PUBLICATIONS**

\*Equal contribution. †Corresponding author.

### Journal articles

**B.** Hie<sup>†</sup>, D. Xu, V. Shanker, T. Bruun, P. Weidenbacher, S. Tang, and P.S. Kim<sup>†</sup>.

"Efficient evolution of human antibodies from general protein language models and sequence information alone."

bioRxiv (in press at Nature Biotechnology), DOI: 10.1101/2022.04.10.487811.

2022

Z. Lin\*, H. Akin\*, R. Rao\*, **B. Hie**\*, Z. Zhu, W. Lu, N. Smetanin, R. Verkuil, O. Kabeli, Y.

Shmueli, A. dos Santos Costa, M. Fazel-Zarandi, T. Sercu, S. Candido, and A. Rives.

"Evolutionary-scale prediction of atomic level protein structure with a language model." bioRxiv (in press at Science), DOI: 10.1101/2022.07.20.500902v2.

2022

**B.** Hie<sup>†</sup>, K.K. Yang, and P.S. Kim<sup>†</sup>.

"Evolutionary velocity with protein language models predicts evolutionary dynamics of diverse proteins."

Cell Systems, 13:4 (featured article).

2022

M.C. Maher, I. Bartha, S. Weaver, J. di Iulio, E. Ferri, L. Soriaga, F.A. Lempp, **B. Hie**, B. Bryson, B. Berger, D.L. Robertson, G. Snell, D. Corti, H.W. Virgin, S.L. Kosakovsky Pond, and A. Telenti.

"Predicting the mutational drivers of future SARS-CoV-2 variants of concern."

Science Translational Medicine, 14:633.

2022

B. Hie and K.K. Yang.

"Adaptive machine learning for protein engineering."

Current Opinion in Structural Biology, 72: February 2022.

2022

R. Singh\*, **B. Hie**\*, A. Narayan, and B. Berger.

"Schema: Metric learning enables interpretable synthesis of heterogeneous single-cell modalities."

Genome Biology, 22:131.

2021

**B.** Hie, E. Zhong, B. Berger, and B. Bryson.

"Learning the language of viral evolution and escape."

Science, 371:6526 (featured article).	2021
<b>B. Hie</b> , B. Bryson, and B. Berger.	
"Leveraging uncertainty in machine learning accelerates biological discovery and Cell Systems, 11:5.	design." 2020
<b>B. Hie</b> *, J. Peters*, S. Nyquist*, A. Shalek, B. Berger, and B. Bryson. "Computational methods for single-cell RNA sequencing."  Annual Review of Biomedical Data Science, 3:1.	2020
<b>B. Hie*</b> , H. Cho*, B. DeMeo, B. Bryson, and B. Berger.  "Geometric sketching compactly summarizes the single-cell transcriptomic landso Cell Systems, 8:6 (cover article).	<i>cape."</i> 2019
<b>B. Hie</b> , B. Bryson, and B. Berger.  "Efficient integration of heterogeneous single-cell transcriptomes using Scanorary	na "
Nature Biotechnology, 37:6.	2019
A. Tehranchi, <b>B. Hie</b> , M. Dacre, I. Kaplow, K. Pettie, P. Combs, and H. Fraser. <i>"Fine-mapping cis-regulatory variants in diverse human populations." eLife</i> , 8:e39595.	2019
<b>B. Hie*</b> , H. Cho*, and B. Berger.  "Realizing private and practical pharmacological collaboration."  Science, 362:6417.	2018
A. Tehranchi, M. Myrthil, T. Martin, <b>B. Hie</b> , D. Golan, and H. Fraser. "Pooled ChIP-seq links variation in transcription factor binding to complex diseas Cell, 165:3.	e risk." 2016
Conference papers	
C. Hsu, R. Verkuil, J. Liu, Z. Lin, <b>B. Hie</b> , T. Sercu, A. Lerer, and A. Rives. "Learning inverse folding from millions of predicted structures."  International Conference on Machine Learning (ICML).	2022
<b>B. Hie</b> , E. Zhong, B. Bryson, and B. Berger.  "Learning mutational semantics."	
Neural Information Processing Systems (NeurlPS).	2020
<b>B. Hie</b> , H. Cho, B. DeMeo, B. Bryson, and B. Berger.  "Geometric sketching of single-cell data preserves transcriptional structure."  Research in Computational Molecular Biology (RECOMB).	2019
Preprints	
<b>B. Hie*</b> , S. Candido*, Z. Lin, O. Kabeli, R. Rao, N. Smetanin, T. Sercu, and A. Rives "A high-level programming language for generative protein design." bioRxiv, DOI: 10.1101/2022.12.21.521526.	2022
C. Itoh, C. Gunnarson, G. Babunovic, A. Nibasumba, Ngomu A., M. Wadsworth III, Hughes II, S. Solomon, <b>B. Hie</b> , B. Berger, A. Shalek, S. Fortune, and B. Bryson.	Т.

"GM-CSF differentiation of human monocytes stabilizes macrophage state via oxidative signaling."

*bioRxiv,* DOI: 10.1101/2020.09.29.318352.

### **SOFTWARE**

Scanorama, Primary Developer

https://github.com/brianhie/scanorama, 90k+ PyPI downloads

Geosketch, Primary Developer

https://github.com/brianhie/geosketch, 63k+ PyPI downloads

**Evolocity, Primary Developer** 

https://github.com/brianhie/evolocity, 10k+ PyPI downloads

esm, Contributor

https://github.com/facebookresearch/esm

scverse, Contributor

https://github.com/scverse

### **TEACHING**

Massachusetts Institute of Technology, Cambridge, MA

Teaching Assistant, Algorithms for Inference (6.438)

2019

Graduate-level course on statistical inference with probabilistic graphical models. Responsible for preparing exams/assignments, leading discussion sections, and holding office hours.

#### ACADEMIC SERVICE

### Mentorship

Stanford University, Doctoral thesis research 2021-Present

University of Toronto, iGEM competition team 2021

Massachusetts Institute of Technology, Masters of Engineering thesis research 2020-2021

#### **Public Engagement and Science Communication**

Press interviews for Al Jazeera, Freethink, Ideas Roadshow, IEEE Spectrum, Inverse, MIT Technology Review, NSF The Discovery Files, Nautilus, Quanta Magazine, Swiss Radio, Wall Street Journal, and Wired

#### **Peer Review**

Contributed reviews to *Bioinformatics, BMC Bioinformatics, Cell, Cell Systems, Frontiers in Genetics, GigaScience, Intelligent Systems for Molecular Biology (ISMB), Journal of Molecular Biology, Nature, Nature Biotechnology, Nature Communications, Nature Methods, NeurIPS Workshop on Machine Learning in Structural Biology (MLSB), Nucleic Acids Research, PLOS Computational Biology, PLOS ONE, PNAS, PNAS Nexus, Research in Computational Molecular Biology (RECOMB), and Science Translational Medicine* 

### **PATENTS**

A. Thugabere Jagadeesh and **B. Hie**.

"Hyperspectral scanning to determine skin health."

US Patent No. 11,532,400.

2022
2021
2020
2020
2022
2021-Present
2019-2021
2019
2016
2015-Present
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2013
2013
2012