ALEX MOREHEAD

result. Paper and Data.

Machine Learning & Computational Biology Researcher

	om Perkeley, Ca 00-0002-0586-6191	lifornia, USA	amorehead	
"Machine Learning & Computational Bioledge deep learning algorithms for geome ideas to life and implementing them using leading research teams, fostering collabo	tric (e.g., structure g Al-standard pro	al biology) data gramming lang	a. Track record o guages/framewo	of bringing new orks. Experienced
SKILLS				
Python PyTorch PyG PyTorch Lightn	ing W&B Hyd	Ira Slurm C	Git LaTeX Pa	ndas Java
Geometric deep learning Generative mode	eling Graph neura	al networks A	I4Science Com	npBio/Chem
EXPERIENCE				
Hopper Postdoctoral Fellow (Supervisor Lawrence Berkeley National Laboratory NER	SC			
☐ Jun 2025 - Ongoing		California, USA		JEDGG
Awarded Berkeley Lab's 2025 Hopper Postdo Research and develop deep generative mod 	els for science .			
Conduct large-scale deep learning HPC exp		contributions to	open-source cod	e along the way.
Deep learning Generative models Al4S	cience CompBio	/Chem		
Graduate Research Assistant (Supervis University of Missouri Bioinformatics & Mac Aug 2020 - May 2025	hine Learning Lab	neng) dissouri, USA		
Awarded two competitive first-year PhD fellow • Researched geometric and generative mode				
 Developed two state-of-the-art protein representations can be called the protein representation of the protein representation representation of the protein representation re	and flow model for	blind protein-liga	and docking. Publ	ished results at top-
Introduced the first deep learning benchman	k for broadly applic	able protein-liga	and docking.	
• Orchestrated several large-scale deep learn	ng HPC experiment	ts and contribute	ed to open-source	code.
Deep learning Geometric & graph represe	ntation learning	Generative mode	Structural b	ioinformatics
Research Intern (Supervisor: Dr. Jeffre Profluent Bio Al Research Lab	_	alifornia, USA		
 Created MMDiff, the first SE(3) diffusion mowhich achieved a 9% nucleic acid design such 	odel for joint sequer	nce-structure gei		
	Sequence generatio			ogle Cloud
Research Intern (Supervisor: Joshua M Absci Al Research Lab Jun 2022 - Apr 2023		lew York, USA		
 Collaboratively attained a 0.1% de novo anti 			ng deep learning, a	a first-of-its-kind

Protein design Generative Al Data science Model prototyping & benchmarking Kubernetes

MY LIFE PHILOSOPHY

"The cure to boredom is curiosity.
There is no cure for curiosity." - Dorothy Parker

ACHIEVEMENTS



Berkeley Lab's 2025 Hopper Postdoctoral Fellow

• Won a *competitive* postdoctoral fellowship for recent PhD graduates in computing sciences.



LoG Top-10 Reviewer

• Awarded large monetary prize for being a *top-3%* reviewer for the 2023 Learning on Graphs (LoG) conference.



Dean's Engineering Excellence & O'Neill Graduate Fellowships and Top-Ranked EECS/CS PhD Student Awards

Won *two* competitive graduate fellowships for first-year
 PhD students and the *outstanding EECS & CS PhD student* awards.



Public outreach

• Research featured in *two* public-facing venues including HPCwire and Marktechpost.

INVITED TALKS

- FlowDock: Geometric Flow Matching for Protein-Ligand Docking and Affinity Prediction, 3DSIG @ ISMB, Jul 2025, Liverpool, UK
- Geometric Deep Learning and Generative Models For Protein-Ligand Docking, Computational modeling of molecular structures course @ MU, Nov 2024, Columbia, Missouri, USA
- RNA-FrameFlow for de novo 3D RNA backbone design, SPIGM @ ICML, Jul 2024, Vienna, AT
- A Gated Graph Transformer for Protein Complex Structure Quality Assessment, 3DSIG @ ISMB, Jul 2023, Lyon, FR
- Geometry-Complete Perceptron Networks for 3D Molecular Graphs, Al2ASE @ AAAI, Feb 2023, Washington D.C., USA
- Neural Diffusion Models: Next-Generation Generative Deep Learning, Advanced topics in deep learning course @ MU, Nov 2022, Columbia, Missouri, USA

EDUCATION

M.S. & Ph.D. in Machine Learning & Computational Biology

University of Missouri | O'Neill and College of Engineering Dean's Graduate Fellow

🗖 Aug 2020 - May 2025

PhD Topic:

E Geometric Deep Learning & Generative Modeling of 3D Biomolecules

Collaborations:

 University of Cambridge, Research Labs of Prof. Pietro Lió & Prof. Sir Tom Blundell

Coursework:

Highest distinction (GPA=4.0/4.0)

B.S. in Computer Science

Missouri Western State University | General Studies and Outstanding Graduate Honors

🗖 Aug 2016 - May 2020

 Graduated top of class among all 2020 graduates in computer science, mathematics, and physics (GPA=4.0/4.0)

REFERENCES

Dr. Jianlin Cheng

- @ University of Missouri

Dr. Dong Xu

- Output
 <p
- xudong@missouri.edu

Dr. Xiaoqin Zou

- Output
 University of Missouri

REVIEWING

Conferences:

- Neural Information Processing Systems (NeurIPS), also AI4Sci, AI4D3, GenBio, & MLSB workshops [2021-...]
- International Conference on Learning Representations (ICLR), also GEM, Al4NA, and FPI workshops [2024-...]
- International Conference on Machine Learning (ICML), also Al4Sci, CB, AccMLBio, & SPIGM workshops [2024-...]
 - Also served on the ICML 2024 ML4LMS workshop's organizing committee [2024]
- Association for the Advancement of Artificial Intelligence (AAAI) [2025]
- Learning on Graphs Conference (LoG) of PMLR [2024-...], organizing comittee [2024-2025]
- ACM Conference on Bioinformatics, Computational Biology, & Health Informatics (ACM BCB) [2024-...]

Journals:

- Nature Machine Intelligence & Nature Methods [2023-...]
- Bioinformatics, Briefings in Bioinformatics, BMC Bioinformatics, JCIM, and CSB Journal [2024-...]
- IEEE Trans. on Neural Nets. & Learning Systems (TNNLS) & Emerging Topics in Comp. Intell. (TETCI) [2023-...]

PUBLICATIONS

Conference Proceedings

- [1] C. K. Joshi et al., "GRNAde: Geometric deep learning for 3d RNA inverse design," in *Thirteenth International Conference on Learning Representations (ICLR)*, Selected as an ICLR 2025 *spotlight*, 2025.
- [2] **A. Morehead** and J. Cheng, "Flowdock: Geometric flow matching for generative protein-ligand docking and affinity prediction," in *Intelligent Systems for Molecular Biology (ISMB)*, Presented as a CASP16 top-5 method, 2025.
- [3] A. R. Jamasb* et al., "Evaluating representation learning on the protein structure universe," in Twelth International Conference on Learning Representations (ICLR), also presented at NeurIPS 2023 MLSB (poster), 2024.
- [4] **A. Morehead**, N. Giri, J. Liu, and J. Cheng, "Deep learning for protein-ligand docking: Are we there yet?" In *ICML Al4Science Workshop*, presented as a *spotlight*, top 20% 30/159, 2024.
- [5] X. Chen*, A. Morehead*, J. Liu, and J. Cheng, "A gated graph transformer for protein complex structure quality assessment and its performance in casp15," in *Intelligent Systems for Molecular Biology (ISMB)*, 2023.
- [6] **A. Morehead**, A. Bhatnagar, J. A. Ruffolo, and A. Madani, "Towards joint sequence-structure generation of nucleic acid and protein complexes," in *NeurIPS Machine Learning in Structural Biology (MLSB) Workshop*, 2023.
- [7] **A. Morehead**, W. Chantapakul, and J. Cheng, "Semi-supervised graph learning meets dimensionality reduction," in *IEEE International Conference on Machine Learning and Applications*, 2023.
- [8] E. Soltanikazemi, R. S. Roy, F. Quadir, N. Giri, A. Morehead, and J. Cheng, "Drlcomplex: Reconstruction of protein quaternary structures using deep reinforcement learning," in *International Conference on Intelligent Biology and Medicine*, 2023.
- [9] **A. Morehead**, C. Chen, and J. Cheng, "Geometric transformers for protein interface contact prediction," in *Tenth International Conference on Learning Representations (ICLR)*, 2022.
- [10] M. Shoman, A. Aboah, A. Morehead, Y. Duan, A. Daud, and Y. Adu-Gyamfi, "A region-based deep learning approach to automated retail checkout," in *Proceedings of the IEEE/CVF CVPR Workshops*, 2022.
- [11] M. Gao et al., "High-performance deep learning toolbox for genome-scale prediction of protein structure and function," in IEEE/ACM Machine Learning with Graphs in High Performance Computing Environments (MLHPC) Workshop, 2021.
- [12] A. Morehead, L. Ogden, G. Magee, R. Hosler, B. White, and G. Mohler, "Low cost gunshot detection using deep learning on the raspberry pi," in *IEEE International Conference on Big Data*, 2019.

Journal Articles

[13] R. Anand* et al., "Rna-frameflow for de novo 3d rna backbone design," *Transactions on Machine Learning Research*, 2025, presented at ICML 2024 SPIGM (*oral*) and AI4Science (*spotlight*) workshops.

- [14] H. La, A. Gupta, **A. Morehead**, J. Cheng, and M. Zhang, "Megafold: System-level optimizations for accelerating protein structure prediction models," *arXiv*, 2025.
- [15] A. Morehead, J. Liu, P. Neupane, N. Giri, and J. Cheng, "Protein-ligand structure and affinity prediction in casp16 using a geometric deep learning ensemble and flow matching," *Proteins: Structure, Function, and Bioinformatics*, 2025, Presented as a CASP16 top-5 method.
- [16] **A. Morehead** et al., "How to go with the flow: Flow matching in bioinformatics and computational biology," *Authorea Preprints*, 2025.
- [17] **A. Morehead** and J. Cheng, "Geometry-complete diffusion for 3d molecule generation," *Nature Communications Chemistry*, 2024, also presented at ICLR 2023 MLDD (poster).
- [18] **A. Morehead** and J. Cheng, "Geometry-complete perceptron networks for 3d molecular graphs," *Bioinformatics*, 2024, also presented at AAAI 2023 DLG (poster) and AI2ASE (*oral*).
- [19] C. Chen, X. Chen, A. Morehead, T. Wu, and J. Cheng, "3d-equivariant graph neural networks for protein model quality assessment," *Bioinformatics*, 2023.
- [20] M. F. Lensink et al., "Impact of alphafold on structure prediction of protein complexes: The casp15-capri experiment," *Proteins: Structure, Function, and Bioinformatics*, 2023.
- [21] S. Mahmud, A. Morehead, and J. Cheng, "Accurate prediction of protein tertiary structural changes induced by single-site mutations with equivariant graph neural networks," bioRxiv, 2023.
- [22] **A. Morehead**, C. Chen, A. Sedova, and J. Cheng, "Dips-plus: The enhanced database of interacting protein structures for interface prediction," *Nature Scientific Data*, 2023.

- [23] **A. Morehead** and J. Cheng, "Protein structure accuracy estimation using geometry-complete perceptron networks," *Protein Science*, 2023.
- [24] A. Shanehsazzadeh et al., "Unlocking de novo antibody design with generative artificial intelligence," *bioRxiv*, 2023, follow-up work presented at NeurIPS 2023 MLSB (poster).
- [25] **A. Morehead**, X. Chen, T. Wu, J. Liu, and J. Cheng, "Egr: Equivariant graph refinement and assessment of 3d protein complex structures," *arXiv*, 2022.
- [26] O. Kouckya et al., "Synthetic biology bicistronic designs support gene expression equally well in vitro and in vivo," AJUR, 2020.