## Samuel Sledzieski

162 5th Ave, New York, NY 10010 ssledzieski@flatironinstitute.org • https://samsl.io

RESEARCH	Flatiron Institute	<ul> <li>New York, NY</li> <li>Sep 2024 – Present</li> <li>Princeton, NJ</li> <li>Sep 2024 – Present</li> <li>Cambridge, MA</li> <li>Feb 2020 – Aug 2024</li> <li>Monaco</li> <li>Jan 2024 – Feb 2024</li> <li>Cambridge, MA</li> <li>Feb 2023 – Dec 2023</li> <li>Redmond, WA</li> <li>May 2023 – Oct 2023</li> </ul>			
	Flatiron Research Fellow, Center for Computational Biology				
	Princeton University				
	Visiting Research Collaborator, Lewis-Sigler Institute for Integrative Genomic				
	Massachusetts Institute of Technology				
	Research Assistant, Computation and Biology Group				
	Centre Scientifique de Monaco				
	Visiting Researcher				
	Serinus Biosciences				
	Consultant				
	Microsoft Research				
	Research Intern, AI For Good Lab				
	Cellarity	Cambridge, MA			
	Machine Learning Intern, Perturbation Biology Group	May 2021 – Aug 2021 <b>Lexington, MA</b>			
	MIT Lincoln Laboratory				
	Summer Research Program, Advanced Lasercom Systems Group	May 2019 – Aug 2019			
	University of Connecticut	Storrs, CT			
	Undergraduate Research Assistant, Computational Biology Lab	Jan 2017 – May 2019 Aug 2018 – May 2019			
	Software Developer, Jackson Laboratory for Genomic Medicine				
	Undergraduate Research Assistant, Nelson Lab	Oct 2015 – Dec 2016			
EDUCATION	Massachusetts Institute of Technology	Cambridge, MA			
	PhD, Computer Science	2019 - 2024			
	<ul> <li>Minor in Biological Engineering</li> <li>Concentration: Protein language models, protein and drug interactions, protein structure</li> <li>Advisor: Dr. Bonnie Berger</li> </ul>				
	SM, Electrical Engineering and Computer Science	2019 – 2021			
	University of Connecticut	Storrs, CT			
	BS, Computer Science	2015 - 2019			
	<ul> <li>Minor in Molecular and Cellular Biology</li> <li>Concentration: Bioinformatics, Data Science</li> <li>Advisor: Dr. Mukul Bansal</li> <li>Magna Cum Laude, Honors Scholar</li> </ul>				
TEACHING	Massachusetts Institute of Technology	Cambridge, MA			
	Teaching Assistant, Machine Learning in Genomics (6.878)	Fall 2021			
	Teaching Assistant, Intro to Deep Learning (6.S191)	Winter 2021, 2022, 2023			
	University of Connecticut	Storrs, CT			
	Teaching Assistant, Theory of Computation	Spring 2018			

JOURNAL	[9]	Kshirsagar, Meller, Humphreys, <b>Sledzieski</b> , Xu, Dodhia, Horvitz, Berger, Bowman, Lavista Ferres, Baker, Baek, "Rapid and accurate prediction of protein homo-oligomer symmetry," <i>Nature Communications</i> , 16, 2017 (2025)
	[8]	Singh, Im, Qiu, Macnkess, Gupta, Sorenson, <b>Sledzieski</b> , Erlach, Wendt, Nanfack, Bryson, Berger, "Learning the Language of Antibody Hypervariability," <i>Proceedings of the National Academy of</i> <i>Sciences</i> , 122.1 (2025): e2418918121
	[7]	<b>Sledzieski</b> , Kshirsagar, Baek, Berger, Dodhia, Lavista Ferres, "Democratizing Protein Language Models with Parameter-Efficient Fine-Tuning," <i>Proceedings of the National Academy of Sciences</i> , 121.26 (2024): e2405840121
	[6]	<b>Sledzieski*</b> , Devkota*, Singh, Cowen, Berger, "TT3D: Leveraging Pre-Computed Protein Sequence Models to Predict Protein-Protein Interactions," <i>Bioinformatics</i> , 2023: btad663
	[5]	Singh*, <b>Sledzieski*</b> , Bryson, Cowen, Berger, "Contrastive learning in protein language space predicts interactions between drugs and protein targets," <i>Proceedings of the National Academy of Sciences</i> , 120.24 (2023): e2220778120
	[4]	Kumar, Brenner, <b>Sledzieski</b> , Olaosebikan, Lynn-Goin, Putnam, Yang, Lewinski, Singh, Daniels, Cowen, Klein-Seetharaman, "Transfer of knowledge from model organisms to evolutionarily distant non-model organisms: The coral Pocillopora damicornis membrane signaling receptome," <i>Plos one</i> , 18.2 (2023). 10.1371/journal.pone.0270965
	[3]	Zaman*, <b>Sledzieski*</b> , Wu, Bansal, "virDTL: Viral recombination analysis through phylogenetic reconciliation and its application to sarbecoviruses and SARS-CoV-2," <i>J Comput Biol</i> , 2022 Sep 20. doi: 10.1089/cmb.2021.0507. Epub ahead of print. PMID: 36125448
	[2]	Singh*, Devkota*, <b>Sledzieski</b> , Berger, Cowen, "Topsy-Turvy: integrating a global view into sequence-based PPI prediction," <i>Bioinformatics</i> , 38.Supplement 1 (July 2022): i264–i272
	[1]	<b>Sledzieski*</b> , Singh*, Cowen, Berger, "D-SCRIPT translates genome to phenome with sequence-based, structure-aware, genome-scale predictions of protein-protein interactions," <i>Cell Systems</i> , 12.10 (2021): 969-982
CONFERENCE AND WORKSHOPS	[6]	<b>Sledzieski</b> , Versavel, Singh, Ocitti, Devkota, Kumar, Shhpilker, Roger, Yang, Lewinski, Putnam, Berger, Klein-Seetharaman, Cowen, "Decoding the Functional Interactome of Non-Model Organisms with PHILHARMONIC," <i>Conference on Research in Computational Molecular Biology</i> ( <i>RECOMB</i> ), 2025
	[5]	<b>Sledzieski</b> , Kshirsagar, Baek, Berger, Dodhia, Lavista Ferres, "Parameter-Efficient Fine-Tuning of Protein Language Models Improves Prediction of Protein-Protein Interactions," <i>Machine Learning for Structural Biology Workshop at NeurIPS</i> , 2023
	[4]	Sledzieski*, Singh*, Cowen, Berger, "Contrasting drugs from decoys," <i>Machine Learning for Structural Biology Workshop at NeurIPS</i> , 2022
	[3]	<b>Sledzieski*</b> , Singh*, Cowen, Berger, "Adapting Protein Language Models for Rapid Drug-Target Interaction Prediction," <i>Machine Learning for Structural Biology Workshop at NeurIPS</i> , 2021
	[2]	<b>Sledzieski*</b> , Singh*, Cowen, Berger, "Sequence-based prediction of protein-protein interactions: a structure-aware interpretable deep learning model," <i>Conference on Research in Computational Molecular Biology (RECOMB)</i> , 2021
	[1]	<b>Sledzieski*</b> , Zhang, Mandoiu, Bansal, "TreeFix-TP: Phylogenetic Error Correction for Accurate Reconstruction of Viral Transmission Networks," <i>Pacific Symposium on Biocomputing (PSB)</i> , 2021
PREPRINTS	[2]	Vizgaudis, Kumar, Olaosebikan, Roger, Brenner, <b>Sledzieski</b> , Yang, Lewinski, Singh, Daniels, Cowen, Klein-Seetharaman, "Insulin Signaling and Pharmacology in Corals," <i>Authorea Preprints</i> , posted 31 January 2024, 10.22541/au.170666200.07483513/v1
	[1]	Kousi, Boix, Park, Mathys, <b>Sledzieski</b> , Peng, Bennett, Tsai, Kellis, "Single-cell mosaicism analysis reveals cell-type-specific somatic mutational burden in Alzheimer's Dementia," <i>bioRxiv</i> , posted 22 April 2022, 10.1101/2022.04.21.489103

	Cold Spring Harbor Laboratory Meeting on Network B Machine Learning in Structural Biology (MLSB) Work		Mar 2021, Mar 2023 Dec 2021, Dec 2022, Dec 2023			
	Research on Computational Molecular Biology (RECO	Apr 2019, May 2022				
	Pacific Symposium on Biocomputing (PSB)	Jan 2021				
	IEEE ICCABS Workshop on Computational Advances for Next Generation Sequencing Oc					
	UConn Fall Frontiers in Undergraduate Research		Oct 2018			
	University of Connecticut Bioinformatics Seminar		Mar 2018, Oct 2018			
	Flatiron Postdoctoral Research Fellowship		2024 - 2026			
FELLOWSHIPS	National Science Foundation (NSF) Graduate Research	2021 - 2024				
	First Place, MIT Intro to Deep Learning Final Project C	2020				
	New England Scholar, University of Connecticut		2017 - 2019			
	Dean's List, College of Liberal Arts and Sciences, Scho	ool of Engineering	2015 - 2019			
	Academic Excellence Scholarship, University of Conne	ecticut	2015 - 2019			
	National Merit Scholarship Finalist		2014			
SOFTWARE	D-SCRIPT	https://github.com/samsledje/D-SCRIPT				
	36k+ PyPI downloads					
	ConPLex	https://github.com/samsledje/ConPLex				
	8k+ PyPI downloads					
	PHILHARMONIC	hub.com/samsledje/philharmonic				
			os://github.com/suz11001/virDTL			
	TreeFix-TP	https://github.com/samsledje/TreeFix-TP				
MENTORING	MIT Undergraduate Research Opportunities Program (	(IPOD) A divisor	2021 – 2024			
	MIT Research Summer Institute (RSI) Advisor	UKOF J AUVISOI	2021 - 2024 2022 - 2023			
	HackMIT Mentor		2022 - 2023			
	Nature, Nature Methods, Nature Communications, Current Opinions in Structural Biology, Bioinformatics, Journal of Computational Biology, PLOS Computational Biology, IEEE Transactions on Artificial Intelligence, NeurIPS, Machine Learning in Structural Biology (MLSB), RECOMB, ISMB					
	International Society for Computational Biology (ISCB	3)				
& ACTIVITIES	Institute of Electronics Engineers (IEEE)					
	Association for Computing Machinery (ACM)					
	Tau Beta Pi, Engineering Honor Society (TBII)					
	Eta Kappa Nu (IEEE-HKN)					
	Upsilon Pi Epsilon, Computer Science Honor Society (	UPE)				

[CV compiled on 2025-03-05]