Timothy (Tim) Barry

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EMPLOYMENT

Harvard University Postdoctoral researcher, Department of Biostatistics Advisor: Xihong Lin	2024 –
University of Pennsylvania Postdoctoral researcher, Department of Statistics Advisor: Eugene Katsevich	- December 2023
EDUCATION	
Carnegie Mellon University (CMU) PhD in Statistics Advisors: Kathryn Roeder (CMU), Eugene Katsevich (University of Pennsylvania)	2018 – 2023
University of Maryland, College Park BS in Mathematics with high honors Minor in Computer Science	2014 – 2018
AWARDS	
Howard Hughes Medical Institute Fellowship	2017
 Maryland Summer Scholars Research Grant Banneker-Key Scholarship, University of Maryland's highest academic scholarship 	2016 2014
PAPERS	
• T Barry, K Mason, E Katsevich, K Roeder. "Robust differential expression testing for single-cell CRISPR screens at low multiplicity of infection." <i>Genome Biology</i> . Link. (Mihaela Serban Memorial Award, American Statistical Association, Pittsburgh chapter)	2024
• T Barry , K Roeder, E Katsevich. "Exponential family measurement error models for single-cell CRISPR screens." <i>Biostatistics</i> . Link.	2024
• J Morris, C Caragine, Z Daniloski, J Domingo, T Barry , L Lu, K Davis, M Ziosi, D Glinos, S Hao, E Mimitou, P Smibert, K Roeder, E Katsevich, T Lappalainen, N Sanjana. "Discovery of target genes and pathways at GWAS loci by pooled single-cell CRISPR screens." <i>Science</i> . Link.	2023
• T Barry , X Wang, J Morris, K Roeder, E Katsevich. "SCEPTRE improves calibration and sensitivity in single-cell CRISPR screen analysis." <i>Genome Biology</i> . Link. (Reviewers' choice, American Society of Human Genetics conference)	2021
• T Barry *, E Gurarie*, F Cheraghi, I Kajola, W Fagan. "Does dispersal make the heart grow bolder? Avoidance of anthropogenic habitat elements across wolf life history." <i>Animal Behaviour</i> 166. * <i>Joint first authorship</i> . Link.	2020
• T Barry . "Collections in R: Review and Proposal." The R Journal 10.1. Link.	2018

T Barry, J Deutsch, E Katsevich. "Hands-on single-cell CRISPR screen analysis." e-book preprint. Link

SOFTWARE PACKAGES

- sceptre: statistically rigorous and massively scalable single-cell CRISPR screen analysis. Link.
 sceptre is the first package for single-cell CRISPR screen analysis endorsed by 10x Genomics, the main commercial supplier of single-cell CRISPR screen kits.
- ondisc: out-of-core and cluster-scale computing on single-cell data. Link.

PROFESSIONAL SERVICE

- Reviewer, Annals of Applied Statistics, Biometrika, Nature Biotechnology
- Judge, New England Statistical Society student paper competition

MENTORING

• Songcheng Dai (Computational Biology Masters student at CMU). Topic: algorithms, data structures, and software for large-scale single-cell data.

TALKS

• "Let it SNO: massive-scale perturb-seq analysis using sceptre, Nextflow, and ondisc (SNO)." RECOMB-seq conference. Boston, MA. Contributed poster.	2024
• "Let it SNO: massive-scale perturb-seq analysis using sceptre, Nextflow, and ondisc (SNO)." IGVF CAMP working group. Invited.	2024
• "Robust inference by resampling score statistics, with application to single-cell CRISPR screens." IGVF conference. St Louis, MO. Contributed poster.	2023
• "Robust inference by resampling score statistics, with application to single-cell CRISPR screens." Bioconductor Conference. Boston, MA. Contributed poster.	2023
• "Robust inference by resampling score statistics, with application to single-cell CRISPR screens." ASA Pittsburgh chapter spring banquet. Pittsburgh, PA. Contributed poster.	2023
• "Robust differential expression analysis for single-cell CRISPR screens." Hicks and Hansen Labs, Department of Biostatistics, Johns Hopkins University. Baltimore, MD. Invited.	2023
• "Exponential family measurement error models for single-cell CRISPR screens." Joint Statistics Meetings. Washington, DC. Contributed.	2022
• "Conditional resampling improves calibration and sensitivity in single-cell CRISPR screen analysis." RECOMB-Seq. Held virtually. Contributed.	2021
• "Conditional resampling improves calibration and sensitivity in single-cell CRISPR screen analysis." He Lab, Department of Human Genetics, University of Chicago. Held virtually. Invited.	2021

TEACHING ASSISTANTSHIPS (CMU)

 Statistics 36-350: 	Statistical comp	puting	Fall 2018
 Statistics 36-469: 	Statistical genor	omics and high-dimensional inference	Spring 2020

COMPUTING

 \bullet Languages: R, Python, C/C++, Nextflow

• Clusters and clouds: SLURM, Sun Grid Engine, Azure

• Database systems: HDF5

Updated May 2024.