# BRENDAN F. MILLER

Post-doctoral research fellow developing computational tools for analysis of spatial transcriptomics data. Molecular biologist with strong background in clinical diagnostics and cancer biology.



## **CURRENT POSITION**

Current 2020

## **Post-doctoral Research Fellow**

Johns Hopkins University Baltimore, MD Department of Biomedical Engineering JEFworks Lab

- Supervisor: Dr. Jean Fan
- Project: "Reference-free deconvolution of multi-cellular spatially resolved transcriptomics data"



## **EDUCATION**

2020 2014

#### Ph.D., Molecular Biology

Johns Hopkins University Paltimore, MD

- Supervisor: Dr. Laura Elnitski
- Dissertation: "Investigating Blood-based Biomarkers and Patterns of DNA Methylation in Tumors"
- Part of the National Institutes of Health and Johns Hopkins University Graduate Partnership Program

2012 2008

## B.S., Biochemistry (minor in Pharmacology)



## AWARDS AND HONORS

2020

National Human Genome Research Institute Intramural Research Award

National Institutes of Health 

Bethesda. MD

• Project: "Detection of Methylated Cancer Biomarkers in Cell-free DNA"

2020

#### National Institutes of Health Fellows Award for Research Excellence

National Institutes of Health 

Bethesda, MD

• Abstract: "A Methylation Density Binary Classifier for Predicting and Optimizing the Performance of Methylation Biomarkers in Clinical Samples"

2019

## National Institutes of Health Graduate Student Research Award

National Institutes of Health 

Bethesda, MD

- 15th Annual NIH Graduate Student Research Symposium
- Section: Pharmacology and. Clinical and Translational Science



## CONTACT

- https://github.com/bmill3r
- https://bmill3r.github.io/
- in brendan-f-miller

# **PROFICIENCIES**

## **Computational Biology**

Python, R, git, Unix

#### **Data Analysis**

Spatial transcriptomics Single-cell sequencing

# **Diagnostics and** Sequencing

Droplet-digital PCR Bisulfite sequencing Cell-free DNA

## **Molecular Biology**

DNA methylation Cell culture **DNA** cloning Nucleic acid purification siRNA knockdowns

2017	•	National Institutes of Health Fellows Award for Research Excellence
		National Institutes of Health ♥ Bethesda, MD
		<ul> <li>Abstract: "Digital Droplet PCR Liquid Biopsy Assay for Detecting Circulating Tumor DNA in Patient Plasma"</li> </ul>
2013		National Institutes of Health Post-Baccalaureate Outstanding Poster Award
		National Institutes of Health ♥ Bethesda, MD
2012		National Institutes of Health Post-Baccalaureate Training Award  National Institutes of Health    Bethesda, MD
2012		John Thanassi Research Award for Outstanding Achievements in Biochemistry
		University of Vermont ♥ Burlington, VT
	Д	RESEARCH EXPERIENCE
2020	•	Graduate Research Fellow
 2014		Translational and Functional Genomics Branch
2011		<ul> <li>Advisor: Dr. Laura Elnitski</li> <li>Project 1: "Detection of Methylated Cancer Biomarkers in Cell-free DNA"</li> <li>Project 2: "Elucidation of Molecular Commonalities in CpG Island Methylator Phenotype Tumors Across Cancer-types"</li> </ul>
2014		Post-Baccalaureate Research Fellow
 2012		Genomic Structure and Function Section ♥ Bethesda, MD
2012		Advisor: Dr. Anthony Furano
		Project: "Factors that Determine Strand Selection During Repair of T/G Mismatches"
2012	•	Undergraduate Researcher
 2011		Department of Pharmacology ♥ University of Vermont
2011		<ul> <li>Advisor: Dr. Wolfgang Dostmann</li> <li>Project: "Purification and Long-term Storage of Toxoplasma gondii PKGII Protein"</li> </ul>
2011		Undergraduate Researcher
		Department of Biochemistry • University of Vermont
		<ul> <li>Advisor: Dr. Anne B. Mason</li> <li>Project: "Identification of Key Residues of Human Transferrin and Transferrin Receptor"</li> </ul>
2007		Research Assistant  Department of Biological Sciences University of Massachusetts, Lowell, MA  • Advisor: Dr. Brian Bettencourt
		Project: "Variation inn Hsp70 Expression During Natural Thermal Stress Drives Differential Suppression of poly-Q Toxicity"

# **♣** TEACHING EXPERIENCE **Lecturer - Research Tools for Studying Disease** 2016 National Institutes of Health Bethesda, MD 2015 • Developed syllabus, overall course structure, and administered grades for "Proteins I" and "Computational Biology" lectures **Teaching Assistant - General Biology** 2015 Managed laboratory section, which included experimental setup, development of instructional lectures, and adminstration of assignments and exams **Program Leader** 2014 Health Education Outreach Program Pethesda, MD 2012 • Taught topics in medicine and healthcare to underpriviledged and underrepresented **Tutor - General Chemistry and Organic Chemistry** 2012 University of Vermont ◆ Burlington, VT 2010 • Established and oversaw individual and group tutoring sessions INVITED SPEAKER Reference-free cell-type deconvolution of multi-cellular pixel resolution 2022 spatially resolved transcriptomics data CNS Research Showcase at Indiana University Virtual Reference-free cell-type deconvolution of multi-cellular pixel resolution 2021 spatially resolved transcriptomics data University of Sydney Bioinformatics Seminar Series Virtual A methylation density binary classifier for predicting and optimizing the 2019 performance of methylation biomarkers in clinical samples Circulating Nucleic Acids/Liquid Biopsy Interest Group Virtual Advancement in ovarian carcinoma detection using circulating cell-free 2018 DNA from patient plasma samples National Human Genome Research Institute Symposium Bethesda, MD Detecting DNA methylation patterns in patient plasma to improve cancer 2018 diagnostics NIEHS Inflammation Faculty Workshop Pasearch Triangle Park, NC Detecting DNA methylation patterns in patient plasma to improve cancer 2018 diagnostics

3rd Annual Liquid Biopsy Summit San Francisco, CA



# 📮 SELECTED POSTER PRESENTATIONS

2020

A methylation density binary classifier for predicting and optimizing the performance of methylation biomarkers in clinical samples

Advances in Genome Biology and Technology 

Marco Island, FL

2017

Detecting DNA methylation in blood for cancer diagnostics

2017

**Detecting DNA methylation in blood for cancer diagnostics** 

EMBO Chromatin and Epigenetics Meeting 

Heidelberg, Germany

Factors that determine strand selection during repair of T/G mismatches 2013

13th Annual Postbac Poster Day Pethesda, MD



## PREPRINT PUBLICATIONS

2015

Building genomic analysis pipelines in a hackathon setting with bioinformatician teams: DNA-Seq, Epigenomics, Metagenomics, and RNA-Seq

bioRxiv 018085

Ben Busby, Allissa Dillman, Claire L. Simpson, Ian Fingerman, Sijung Yun, David M. Kristensen, Lisa Federer, Naisha Shah, Matthew C. LaFave, Laura Jimenez-Barron, Manjusha Pande, Wen Luo, Brendan Miller, Cem Mayden, Dhruva Chandramohan, Kipper Fletez-Brant, Paul W. Bible, Sergej Nowoshilow, Alfred Chan, Eric JC Galvez, Jeremy Chignell, Joseph N. Paulson, Manoj Kandpal, Suhyeon Yoon, Esther Asaki, Abhinav Nellore, Adam Stine, Robert Sanders, Jesse Becker, Matt Lesko, Mordechai Abzug, Eugene Yaschenko



# SELECTED PUBLICATIONS

2022

Reference-free celltype deconvolution of multi-cellular pixel-resolution spatially resolved transcriptomics data

Nat Commun 13, 2339 (2022)

Brendan F. Miller, Feiyang Huang, Lyla Atta, Arpan Sahoo, Jean Fan

Software: STdeconvolve

2021

Characterizing spatial gene expression heterogeneity in spatially resolved single-cell transcriptomics data with nonuniform cellular densities

Genome Res. 2021. 31: 1843-1855

Brendan F. Miller, Dhananjay Bambah-Mukku, Catherine Dulac, Xiaowei Zhuang, Jean

Software: MERINGUE

Assessing ZNF154 methylation in patient plasma as a multicancer marker in liquid biopsies from colon, liver, ovarian and pancreatic cancer patients

Sci Rep 11, 221 (2021)

2020

2019

2016

2012

Brendan F. Miller, Hanna M. Petrykowska, Laura Elnitski

Leveraging locus-specific epigenetic heterogeneity to improve the performance of blood-based DNA methylation biomarkers

Clin Epigenet 12, 154 (2020)

**Brendan F. Miller**, Thomas R. Pisanic II, Gennady Margolin, Hanna M. Petrykowska, Pornpat Athamanolap, Akosua Osei-Tutu, Tza-Huei Wang, Christina Annunziata, Laura Elnitski

Software: EpiClass

Identification of human silencers by correlating cross-tissue epigenetic profiles and gene expression

Genome Research. 2019 March; 29(3):1-11

Di Huang, Hanna M. Petrykowska, **Brendan F. Miller**, Laura Elnitski, Ivan Ovcharenko

2018 Transient reduction of DNA methylation at the onset of meiosis in male mice

Epigenetics and Chromatin. 2018 April; 11:15

Valeriya Gaysinskaya, **Brendan F. Miller**, Godfried W. van der Heijden, Kasper D. Hansen, Alex Bortvin

The emergence of pan-cancer CIMP and its elusive interpretation

Biomolecules. 2016 Nov; 6(4):45

Brendan F. Miller, Francisco Sanchez-Vega, Laura Elnitski

Page 2014 Repair of naturally occurring mispairs can induce mutations in flanking DNA

eLife 2014;3:e02001

Jia Chen, Brendan F. Miller, Anthony V. Furano

 Structure-based Mutagenesis Reveals Critical Residues in the Transferrin Receptor Participating in the Mechanism of pH-induced Release of Iron from Human Serum Transferrin

Biochemistry. 2012 Feb; 51(10):2113-21

Ashley N. Steere, N. Dennis Chasteen, **Brendan F. Miller**, Valerie C. Smith, Ross T. A. MacGillivray, Anne B. Mason

lonic Residues of Human Serum Transferrin that Affect Binding to the Transferrin Receptor and Iron Release from the Complex

Biochemistry. 2012 Dec; 51(2):686-94

Ashley N. Steere, **Brendan F. Miller**, Samantha E. Roberts, Shaina L. Byrne, N. Dennis Chasteen, Valerie C. Smith, Ross T. A. MacGillivray, Anne B. Mason.

# SOFTWARE

2021

#### **STdeconvolve**

https://jef.works/STdeconvolve/

• Unsupervised machine learning approach to deconvolve multi-cellular pixel resolution spatial transcriptomics datasets in order to recover the putative transcriptomic profiles of cell-types and their proportional representation within spatially resolved pixels without reliance on external single-cell transcriptomics references.

2021

#### **MERINGUE**

https://jef.works/MERINGUE/

• MERINGUE characterizes spatial gene expression heterogeneity in spatially resolved single-cell transcriptomics data with non-uniform cellular densities.

2020

#### **EpiClass**

https://pypi.org/project/EpiClass/

• Optimizing and predicting performance of DNA methylation biomarkers using sequence methylation density information.



# SERVICE

Current 2019

## Circulating Nucleic Acids/Liquid Biopsy Interest Group

Research Webinar Committee Co-Leader

• The aims of the circulating nucleic acid/liquid biopsy interest group are (1) working towards establishing standard practices for CNA studies; (2) sharing and optimization of DNA techniques; (3) communication of CAN research

2021

#### **Journal Reviewer - Nature Protocols**

Prioritization of cell-types responsive to biological perturbations in single-cell data with Augur. Nat Protoc 16, 3836-3873 (2021)

2019

#### **Journal Reviewer - Epigenomics**



## REFERENCES

## Dr. Jean Fan

Johns Hopkins University Department of Biomedical Engineering jeanfan@jhu.edu

### Dr. Laura Elniski

National Institutes of Health National Human Genome Research Institute elnitski@nih.gov

## Dr. Thomas Pisanic II

Johns Hopkins University Department of Mechanical Engineering tpisani1@jhu.edu