# Matthew G. Durrant, Ph.D.

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# **Education**

#### Stanford University - School of Medicine

Sep 2016 – Dec 2020

PhD in Genetics – National Science Foundation Graduate Research Fellow

- Doctoral Dissertation "Mobile genetic elements: mechanisms of microbial evolution and reservoirs of genome engineering tools"
- Advised by Dr. Ami Bhatt and Dr. Stephen Montgomery
- Courses of Study: Foundations in Experimental Biology, Frontiers in Biological Research, Advanced Genetics, Genomics, Statistical and Machine Learning Methods for Genomics, Deep Learning, Economics of Biotechnology

## **Brigham Young University**

Bachelor of Science – Summa cum laude – Provo, Utah

- Double Major: Bioinformatics and Neuroscience, Minor: Mathematics
- Cumulative GPA: 4.0
- Courses of Study: Computational Biology, Bioinformatics, Algorithm Design and Analysis, Advanced Programming, Statistics for Engineers and Scientists, Multivariable Calculus, Linear Algebra, Ordinary Differential Equations, Molecular Biology, Genetics, Advanced Neuroscience, Bioinnovation and Entrepreneurship

# Research Experience —

Arc Institute 7an 2022 – Present

Senior Scientist

- Working in the lab of Dr. Patrick Hsu
- Working with computational and experimental scientists to build a platform for the discovery of new genome engineering systems
- Discovered a family of programmable recombinases that encode a bridge RNA and was co-inventor on several related patents
- Directly mentoring graduate students attending UC Berkeley and Stanford.

**Stylus Medicine** 7an 2022 – Present

Co-founding Scientist

- Developed target product profiles, pitch decks, scientific resources
- Helped to recruit and build the computational biology team
- Implemented the first year of the computational biology research plan for the company
- Performed work that produced new intellectual property for the company
- Primarily using Google Cloud resources and cloud native tools such as Bigtable, BigQuery, Batch, Dataflow, Pub/Sub, etc.

# Pragma Bio (Formerly VastBiome)

Dec 2021 - July 2022

Computational Biology Consultant

- Worked on a team of computational biologists
- Developed data analysis pipelines and software packages for mining biological sequence data for new
- Operated through AWS batch for high-throughput computing

Sep 2016

# **UC Berkeley - Department of Bioengineering**

Jan 2021 – Dec 2021

Postdoctoral Fellow

- Worked in the lab of Dr. Patrick Hsu
- Led several projects to discover new molecular tools for genome engineering
- Gained extensive experience using cloud computing to store and analyze massive biological datasets

# **Bhatt & Montgomery Labs - Stanford School of Medicine**

June 2017 - Dec 2020

Genetics PhD Candidate

- Advisors: Dr. Ami Bhatt and Dr. Stephen Montgomery
- Research Focuses:
  - o Structural variation in microbial genomes
  - o Mining bacterial genomes to identify novel tools for genome editing
  - o Deep learning to annotate small open reading frames (smORFs) in bacterial genomes.
  - o Computational methods development
  - o Human RNA-seq, ATAC-seq, allele-specific expression, gene-environment interaction discovery
- Stanford SPARK Scholar A program to assist scholars as they develop new biotechnology products.
- Other interests:
  - Statistical genetics
  - Machine learning and Deep Learning in genomics

#### NorthShore Bio

Computational Biologist

July 2017 - Oct 2017

- Worked remotely on contract as a computational biologist
- Developed analytical software to accompany molecular analysis technology

# Yale University and the Rothberg Institute

7une 2015 – Dec 2015

Bioinformatics Research Intern – New Haven, CT

- Programmed an application to store millions of genetic variants and perform quality control analysis
- MongoDB, Python, R experience
- Wrote a manuscript documenting quality control procedure and drafting results of a rare variant analysis of an ADHD patient NGS dataset.
- Next Generation Sequencing (whole exome) data experience

Tute Genomics Nov 2014 - Feb 2015

Bioinformatics Research Intern - Provo, UT

- Programmed at Tute Genomics, a bioinformatics company which has developed a comprehensive cloud-based genome annotation platform for clinicians and researchers
- Advised by Bryce Daines, Ph.D. in Human Genetics
- Developed pharmacogenomics platform to be used for commercial purposes in team of 5 interns
- Project development, python programming, database curation, and MongoDB experience

# Max Planck Institute for Chemical Ecology

May 2014 – Aug 2014

Bioinformatics Research Fellow – Jena, Germany

- Experience with Python, UNIX command line, R, and Adobe Illustrator
- Analyzed microarray and RNA-seq transcriptome data
- Published manuscript entitled Evidence of an evolutionary hourglass pattern in herbivory-induced transcriptomic responses in the journal New Phytologist

Undergraduate Research Assistant – Provo, Utah

- Researched under the guidance of Dr. David Busath and Dr. Mary Davis
- Published in BMC Genetics, *Investigation into a recent rise of dual-amantadine resistant mutations in the Influenza A M2 Sequence*. Presented article at The 11th Annual Biotechnology and Bioinformatics Symposium.
- Presented an abstract at the American Society of Human Genetics conference in October 2015 entitled Analysis of pathways associated with Body Mass Index in individuals with multiple sclerosis
- Contributed as a member of the YGen Consortium, a group investigating the relationship between human Y chromosome variation, complex traits, and clinical metadata.
- Molecular Dynamics Simulations experience with VMD, NAMD, CHARMM

# Other Work and Volunteer Experience -

# January, Inc.

June 2019 - Sep 2019

Machine Learning / Science Team

- Worked directly with both experimental biologists and machine learning engineers
- Collaborated with team members to develop features of a direct-to-consumer digital health product
- Clustered and typed users based on behavioral patterns collected from the company's app

# Stanford University - Department of Genetics

Jan. 2019 – Apr. 2019

Genomics Teaching Assistant – Stanford, CA

- Tutored for 2 hours per week
- Taught weekly Python programming lecture to graduate students.
- Evaluated students' code and project proposals.

# Software / Data Carpentry

Jan 2017 – Present

Volunteer Instructor – Stanford, CA

- Certified instructor trained in evidence-based teaching methods.
- Experience teaching graduate students to utilize Python, Jupyter Notebooks, R, and Unix in their research.
- Volunteering regularly as an instructor on a continual basis.

# **Vykon Technologies**

Jan 2015 – 2017

Co-founder - Provo, Utah

- Development of low-cost movement disorder monitoring software application that couples with commercially available 3D motion-capture cameras
- Developed the initial prototype of the leading product DextraSense
- Managed the development of the technology with co-founder Jacob Durrant, PhD
- Market survey development, deployment, and data analysis
- FDA medical device compliance experience

#### Publications -

- 1) \*Nguyen, E., \*Poli, M., \*Durrant, M., Thomas, A., Kang, B., Sullivan, J., Ng, M., Lewis, A., Patel, A., Lou, A., Ermon, A., Baccus, S., Hernandez-Boussard, T., Ré, C., Hsu, P., Hie, B. (2024), Sequence modeling and design from molecular to genome scale with Evo. *bioRxiv*, DOI: https://doi.org/10.1101/2024.02.27.582234. \*Authors contributed equally
- 2) \*Durrant, M., \*Perry, N., Pai, J., Jangid, A., Athukoralage, J., Hiraizumi, M., McSpedon, J., Pawluk, A., Nishimasu, H., Konermann, S., Hsu, P. (2024), Bridge RNAs direct modular and programmable recombination of target and donor DNA. bioRxiv, DOI: https://www.biorxiv.org/content/10.1101/2024.01.24.577089v1. \*Authors contributed equally

- 3) Wei, J., Lotfy, P.m Faizi, K., Baungaard, S., Gibson, E., Wang, E., Slabodkin, H., Kinnaman, E., Chandrasekaran, S., Kitano, H., <u>Durrant, M.</u>, Duffy, C., Pawluk, A., Hsu, P., Konermann, S. (2023), Deep learning and CRISPR-Cas13d ortholog discovery for optimized RNA targeting. *Cell Systems*, DOI: https://www.cell.com/cell-systems/pdf/S2405-4712(23)00329-0.pdf
- 4) \*Durrant, M., \*Fanton, A., \*Tycko, J., Hinks, M., Chandrasekaran, S., Perry, N., Schaepe, J., Du, P., Lotfy, P., Bassik, M., Bintu, L., Bhatt, A., Hsu, P. (2022), Systematic discovery of recombinases for efficient integration of large DNA sequences into the human genome. *Nature Biotechnology*, DOI: https://doi.org/10.1038/s41587-022-01494-w, \*Authors contributed equally
- 5) Smail, C., Ferraro, N., Hui, Q., <u>Durrant, M.</u>, Aguirre, M., Tanigawa, Y., Keever-Keigher, M., Rao, A., Justesen, J., Li, X., Gloudemans, M., Assimes, T., Koopergerg, C., Reiner, A., Huang, J., O'Donnel, C., Sun, Y., Million Veteran program, Rivas, M., Montgomery, S. (2022), Integration of rare expression outlier-associated variants improves polygenic risk prediction. *The American Journal of Human Genetics*, DOI: https://doi.org/10.1016/j.ajhg.2022.04.015
- 6) Gloudemans, M., Balliu, B., Nachun, D., Schnurr, T., <u>Durrant, M.</u>, Ingelsson, E., Wabitsch, M., Quertermous, T., Montgomery, S., Knowles, J., Carcamo-Orive I. (2022), Integration of genetic colocalizations with physiological and pharmacological perturbations identifies cardiometabolic disease genes. *Genome Medicine*, DOI: https://doi.org/10.1186/s13073-022-01036-8
- 7) Balliu, B., Carcamo-Orive, I., Gloudemans, M., Nachun, D., <u>Durrant, M.</u>, Gazal, S., Park, C., Knowles, D., Wabitsch, M., Quertermous, T., Knowles, J., Montgomery, S. (2021), An integrated approach to identify environmental modulators of genetic risk factors for complex traits. *The American Journal of Human Genetics*, DOI:https://doi.org/10.1016/j.ajhg.2021.08.014
- 8) \*Durrant, M., \*Fremin, B., Rao, A., Cribas, E., Montgomery, S., Bhatt, A. (preprint, 2021), Chromatin accessibility changes induced by the microbial metabolite butyrate reveal possible mechanisms of anticancer effects. *biorXiv*, DOI:https://doi.org/10.1101/2021.03.30.437582, \*Authors contributed equally
- 9) <u>Durrant, M.</u>, Bhatt, A. S. (2021), Automated Prediction and Annotation of Small Open Reading Frames in Microbial Genomes. *Cell Host & Microbe*, DOI:https://doi.org/10.1016/j.chom.2020.11.002
- 10) Contrepois, K., Wu, S., Moneghetti, K., Hornburg, D., Ahadi, S., Tsai, M., Metwally, A., Wei, E., Lee, B., Quijada, J., Chen, S., Christle, J., Ellenberger, M., Balliu, B., Taylor, S., <u>Durrant, M.</u>, Knowles, D., Choudhry, H., Ashland, M., Bahmani, A., Enslen, B., Amsallem, M., Kobayashi, Y., Avina, M., Perelman, D., Miryam, S., Zhou, W., Ashley, E., Montgomery, S., Chaib, H., Haddad, F., Snyder, M. (2020), Molecular Choreography of Acute Exercise. *Cell*, DOI:https://doi.org/10.1016/j.cell.2020.04.043
- 11) <u>Durrant, M.</u>, Li, M., Siranosian, B., Bhatt, A. S. (2020), A Bioinformatic Analysis of Integrative Mobile Genetic Elements Highlights Their Role in Bacterial Adaptation. *Cell Host & Microbe*, DOI:https://doiorg.stanford.idm.oclc.org/10.1016/j.chom.2019.10.022
- 12) Balliu, B., <u>Durrant, M.</u>, de Goede, O., Abell, N., Li, X., Liu, B., Gloudemans, M., Cook, N., Smith, K., Pala, M., Cucca, F., Schlessinger, D., Jaiswal, S., Sabatti, C., Lind, L., Ingelsson, E., Montgomery, S. B. (2019), Genetic dysregulation of gene expression and splicing during a ten-year period of human aging. *Genome Biology*. 519520; doi: 10.1186/s13059-019-1840-y.
- 13) <u>Durrant, M.</u>, Bhatt, A. S. (2019), Microbiome genome structure drives function. *Nature Microbiology*, News & Views. doi: 10.1038/s41564-019-0473-yx
- 14) <u>Durrant, M.</u>, Boyer, J., Zhou, W., Baldwin, I. T. and Xu, S. (2017), Evidence of an evolutionary hourglass pattern in herbivory-induced transcriptomic responses. *New Phytologist*. doi:10.1111/nph.14644

15) <u>Durrant, M.</u> , Eggett, D. L., & Busath, D. D. (2015). Investigation of a recent rise of dual amantadine-resistance mutations in the influenza A M2 sequence. <i>BMC genetics</i> , 16(2), S3.

# Abstracts, Posters, and Presentations -

- 1) \*Durrant, M., \*Perry, N., Pai, J., Jangid, A., Athukoralage, J., Hiraizumi, M., McSpedon, J., Pawluk, A., Nishimasu, H., Konermann, S., Hsu, P. (2024), Bridge RNAs direct modular and programmable recombination of target and donor DNA. Poster presented at *Keystone Symposia Precision Genome Engineering*. \*Authors contributed equally.
- 2) \*Durrant, M., \*Fanton, A., \*Tycko, J., Hinks, M., Chandrasekaran, S., Perry, N., Schaepe, J., Du, P., Lotfy, P., Bassik, M., Bintu, L., Bhatt, A., Hsu, P (2022), Systematic discovery of recombinases for efficient integration of large DNA sequences into the human genome. Talk and poster presented at Keystone Symposia Precision Genome Engineering. \*Authors contributed equally.
- 3) \*<u>Durrant MG</u>, \*Tycko J, Bintu L, Bassik M, Bhatt A. (2019), Mining bacterial genomes to identify thousands of novel recombinases for genome editing. Poster abstract presented at *Stanford Medicine Genetics Retreat*. \*Authors contributed equally.
- 4) \*<u>Durrant MG</u>, \*Fremin B, Montgomery S, Bhatt A. (2018), Microbial metabolite butyrate affects allele-specific expression and disrupts chromatin architecture in a colon cancer cell line. Reviewers' choice poster abstract presented at *American Society of Human Genetics Annual Meeting*. \*Authors contributed equally.
- 5) \*<u>Durrant MG</u>, \*Fremin B, Greenside P, Kundaje A, Montgomery S, Bhatt A. (2017) The impact of butyrate, a microbial metabolite, on allele-specific expression and chromatin accessibility in colon cancer cells. Poster abstract presented at *Stanford Medicine Genetics Retreat*. \*Authors contributed equally.
- 6) <u>Durrant MG</u>, Denny JC, Davis MF. (2015) Analysis of pathways associated with Body Mass Index in individuals with multiple sclerosis. Poster abstract presented at *American Society of Human Genetics Annual Meeting*.
- 7) <u>Durrant MG</u>, Eggett DL, Busath DD. (2014) Investigation of a recent rise of dual amantadine-resistance mutations in the influenza A M2 sequence. Article presented at the 11th Annual Biotechnology and Bioinformatics Symposium.

# Intellectual Property —

 $\underline{Durrant,\,M.,\,}$  Hsu, P., Fanton, A., Moon, C. (2022), Integration of large nucleic acids into genomes (WO2023177424A1). Patent application.

<u>Durrant, M.</u>, Bhatt, A., Tycko, J., Hsu, P., Fanton, A., Bassik, M., Bintu, L. (2022), Serine recombinases (WO2023081762A2). Patent application.

### **Code Portfolio** -

- 1) Systematic Discovery Recombinases 2022 Code to accompany Durrant, Fanton & Tycko et al. (2022). Includes a custom pipeline for identifying engineered recombinase plasmid insertions in the human genome.
- 2) GenomeSearch A command line tool to quickly identify closely related genomes using a marker-gene based approach. https://github.com/bhattlab/GenomeSearch
- 3) *SmORFinder* A command line tool to identify and annotate small proteins in genomes and metagenomes. https://github.com/bhattlab/SmORFinder
- 4) *MGEFinder* A toolbox for identifying mobile genetic element (MGE) insertions from short-read sequencing data of bacterial isolates. https://github.com/bhattlab/MGEfinder
- 5) Github: <a href="https://github.com/durrantmm">https://github.com/durrantmm</a>

#### **Awards and Distinctions**

- National Science Foundation Graduate Research Fellow
- Stanford SPARK Scholar
- College of Life Sciences Dean's List 2010, 2011, 2013, 2014, 2015. (Deferred enrollment 2012)
- Brigham Young University Full Ride Scholarship
- Regent's Scholar Utah System of Higher Education
- LFSCI 399R Internship Grant Helped cover the cost of my internship at the Max Planck Institute
- \$15,000 Miller New Venture Challenge Prize Winner Cofounder of Winning Company Vykon Technologies

# Other Skills and Accomplishments

- Proficient in Bash, Python, and R
- Extensive experience with cloud computing, SQL, snakemake workflows, anaconda, shiny web applications
- Fluent in Slovak Language
- Eagle Scout