Daniel B. Munro

Computational biologist at UC San Diego dmunro@health.ucsd.edu https://danmun.ro

EDUCATION

2019 **Ph.D.** Quantitative and Computational Biology, Princeton University

Thesis: Revealing disease-relevant alteration patterns through data

aggregation. Advisor: Mona Singh

2013 **B.S.** Biology, University of North Texas

RESEARCH EXPERIENCE

05/2023-Present Bioinformatics Programmer, UC San Diego and

Visiting Scientist, Seattle Children's Research Institute

Continuing postdoctoral research with Abraham Palmer and Pejman Mohammadi • Quantitative genetics and regulatory genomics • Integrating

genetic variation, multi-modal transcriptomics, and complex traits

03/2020-04/2023 Postdoctoral Fellow, Scripps Research and

Postdoctoral Scholar, UC San Diego

Advisors: Pejman Mohammadi & Abraham Palmer • Quantitative genetics and regulatory genomics • Integrating genetic variation, multi-modal

transcriptomics, and complex traits

03/2019–10/2019 **Postdoctoral Associate**, New York University

Advisor: Christine Vogel • Computational research in proteomics and

cancer genomics

01/2014-12/2018 Graduate Research Assistant, Princeton University

Advisor: Mona Singh • Computational research in cancer genomics, protein

variant impact, and histological image analysis

01/2011–05/2013 Undergraduate Research Assistant, University of North Texas

Advisor: Qunfeng Dong • Computational microbiome research and

genomics software development

PEER-REVIEWED PUBLICATIONS

D Munro, N Ehsan, SM Esmaeili-Fard, A Gusev, AA Palmer, P Mohammadi. Multimodal analysis of RNA sequencing data powers discovery of complex trait genetics. *Accepted at Nature Communications 2024-10-11*.

Preprint: https://www.biorxiv.org/content/10.1101/2024.05.14.594051v1

- F Okamoto, AS Chitre, TM Sanches, D Chen, **D Munro**, AT Aron, A Beeson, et al. Y and mitochondrial chromosomes in the heterogeneous stock rat population. *G3 GeneslGenomeslGenetics* (2024). ikae213.
- TV de Jong, Y Pan, P Rastas, **D Munro**, M Tutaj, H Akil, C Benner, et al. A revamped rat reference genome improves the discovery of genetic diversity in laboratory rats. *Cell Genomics* (2024). 4 (4): 100527.
- R Clifford, **D Munro**, D Dochtermann, P Devineni, S Pyarajan, Million Veteran Program, F Telese, et al. Genome-wide association study of chronic dizziness in the elderly identifies loci implicating *MLLT10*, *BPTF*, *LINC01224*, and *ROS1*. *JARO* (2023). 24: 575-591.
- JL Zhou, G de Guglielmo, AJ Ho, M Kallupi, N Pokhrel, H-R Li, AS Chitre, **D Munro**, et al. Single-nucleus genomics in outbred rats with divergent cocaine addiction-like behaviors reveals changes in amygdala GABAergic inhibition. *Nature Neuroscience* (2023). 26: 1868–1879.
- AS Chitre, O Polesskaya, **D Munro**, R Cheng, P Mohammadi, K Holl, J Gao, et al. Exponential increase in QTL detection with increased sample size. *GENETICS* (2023). 224 (2): iyad054.
- S Fowler, T Wang, **D Munro**, A Kumar, AS Chitre, TJ Hollingsworth, A Garcia Martinez, et al. Genome-wide association study finds multiple loci associated with intraocular pressure in HS rats. *Frontiers in Genetics* (2023). 13: 1029058.
- **D Munro**, T Wang, AS Chitre, O Polesskaya, N Ehsan, J Gao, A Gusev, LC Solberg Woods, LM Saba, H Chen, AA Palmer, P Mohammadi. The regulatory landscape of multiple brain regions in outbred heterogeneous stock rats. *Nucleic Acids Research* (2022). 50 (19): 10882–10895.
- JT Ash*, G Darnell*, **D Munro***, BE Engelhardt. Joint analysis of gene expression levels and histological images identifies genes associated with tissue morphology. *Nature Communications* (2021). 12 (1), 1–12.
- * These authors contributed equally
- **D Munro**, M Singh. DeMaSk: A deep mutational scanning substitution matrix and its use for variant impact prediction. *Bioinformatics* (2020). 36 (22–23): 5322–5329.
- GXL Li, **D Munro**, D Fermin, C Vogel, H Choi. A protein-centric approach for exome variant aggregation enables sensitive association analysis with clinical outcomes. *Human Mutation* (2020). 41 (5): 934–945.
- **D Munro**, D Ghersi, M Singh. Two critical positions in zinc finger domains are heavily mutated in three human cancer types. *PLoS Comput Biol* (2018). 14 (6): e1006290.
- C Cohen, E Toh, **D Munro**, Q Dong, H Hawlena. Similarities and seasonal variations in bacterial communities from the blood of rodents and from their flea vectors. *The ISME Journal* (2015). 2015-01-09.

Y Gavish, H Kedem, I Messika, C Cohen, E Toh, **D Munro**, Q Dong, C Fuqua, K Clay, H Hawlena. Association of host and microbial species diversity across spatial scales in desert rodent communities. *PLoS ONE* (2014). 9: e109677.

JS Kuehn, PJ Gorden, **D Munro**, R Rong, Q Dong, PJ Plummer, C Wang, GJ Phillips. Bacterial community profiling of milk samples as a means to understand culture-negative bovine clinical mastitis. *PLoS ONE*. 8: e61959.

M Zhou, R Rong, **D Munro**, C Zhu, X Gao, Q Zhang, Q Dong (2013). Investigation of the effect of type 2 diabetes mellitus on subgingival plaque microbiota by high-throughput 16S rDNA pyrosequencing. *PLoS ONE* (2013). 8: e61516.

H Hawlena, E Rynkiewicz, E Toh, A Alfred, LA Durden, MW Hastriter, DE Nelson, R Rong, **D Munro**, Q Dong, C Fuqua, K Clay. The arthropod, but not the vertebrate host or its environment, dictates bacterial community composition of fleas and ticks. *The ISME Journal* (2013). 7: 221-223.

K Revanna, **D Munro**, A Gao, C Chiu, A Pathak, Q Dong. A web-based multi-Genome Synteny Viewer for customized data. *BMC Bioinformatics* (2012). 13: 190.

PREPRINTS

N Santhanam, S Sanchez-Roige, S Mi, Y Liang, AS Chitre, **D Munro**, D Chen, et al. RatXcan: A framework for cross-species integration of genome-wide association and gene expression data. Preprint: https://www.biorxiv.org/content/10.1101/2022.06.03.494719v5

EK Hebda-Bauer, MH Hagenauer, **DB Munro**, P Blandino Jr., F Meng, K Arakawa, AS Chitre, et al. Bioenergetic-Related Gene Expression in the Hippocampus Predicts Internalizing vs. Externalizing Behavior in a F_2 Cross of Selectively-Bred Rats.

Preprint: https://www.biorxiv.org/content/10.1101/2022.07.14.500129v4

HONORS & AWARDS

2019	Runner-up, Symbiosis Competition at Imagine Science Film Festival, New York, NY, in collaboration with Jin Angdoo
2013	National Science Foundation Graduate Research Fellowship
2013	Phi Kappa Phi National Fellowship (\$5000)

TEACHING EXPERIENCE

Spring 2016 Assistant in Instruction, "An Integrated, Quantitative Introduction to the Natural Sciences II", Princeton University

Fall 2015 Assistant in Instruction, "An Integrated, Quantitative Introduction to the Natural

Sciences I", Princeton University

PRESENTATIONS

2022 "The regulatory landscape of multiple brain regions in outbred heterogeneous

stock rats", talk at the Complex Trait Consortium-Rat Genomics 2022

meeting, September 29, 2022

2021 "Identification of regulatory landscape in multiple brain regions of outbred

heterogeneous stock rats", talk at the Complex Trait Consortium-Rat

Genomics 2021 meeting, September 1, 2021

2021, 2023 "Techniques for algorithmic graphics", guest lecture for Honors Colloquium on

Complex Systems, University of Nebraska Omaha, March 2, 2021 and March

28, 2023

2020 "Mapping eQTLs in five rat brain regions", talk at the 6th Annual Retreat for

P50 Center for GWAS in Outbred Rats, La Jolla, CA, November 2, 2020

PROFESSIONAL SERVICE & MEMBERSHIPS

Ad hoc peer reviewer for:

Developmental Biology (2019)

PLOS Computational Biology (2020)

PLOS ONE (2020)

BMC Bioinformatics (2021)

Journal of Computational Biology (2021)

Genome Biology (2023)

IEEE/ACM Transactions on Computational Biology and Bioinformatics (2023)

The American Journal of Human Genetics (2023)

Neuropsychopharmacology (2024)

Memberships:

American Society of Human Genetics (Since 2020)

International Society for Computational Biology (Since 2019)

CERTIFICATIONS

Initial Physiological Training Course – NASA (2024-11-07)

Open Water Diver – PADI (2024-07-28)

Amateur Radio License, General Class – FCC (2024-06-25)