### UNIVERSITY OF ROCHESTER SCHOOL OF MEDICINE & DENTISTRY CURRICULUM VITAE

# Matthew N. McCall, Ph.D.

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Date of Birth: 01/08/1982 Citizenship: USA

#### **CURRENT POSITIONS**

Associate Professor of Biostatistics & Computational Biology University of Rochester School of Medicine & Dentistry, Rochester, NY

### **EDUCATION**

2000 - 2004	B.S., Statistics, University of Michigan, Ann Arbor, MI
2009 - 2010	M.H.S., Bioinformatics, Johns Hopkins School of Public Health, Baltimore, MD
2005 - 2010	Ph.D., Biostatistics, Johns Hopkins School of Public Health, Baltimore, MD

# **POST-DEGREE TRAINING**

2010 - 2013	Postdoctoral Fellowship, University of Rochester School of Medicine & Dentistry,
	Rochester, NY
2013 - 2014	Postdoctoral Associate, University of Rochester School of Medicine & Dentistry,
	Rochester NY

#### FACULTY APPOINTMENTS

2014 - 2018	Assistant Professor, Department of Biostatistics & Computational Biology,
	University of Rochester School of Medicine & Dentistry, Rochester, NY
2014 - 2018	Assistant Professor, Department of Biomedical Genetics,
	University of Rochester School of Medicine & Dentistry, Rochester, NY
2019 – present	Associate Professor, Department of Biostatistics & Computational Biology,
	University of Rochester School of Medicine & Dentistry, Rochester, NY
2019 – present	Associate Professor, Department of Biomedical Genetics,
	University of Rochester School of Medicine & Dentistry, Rochester, NY
2015 – present	Affiliated Faculty, The Goergen Institute for Data Science,
	University of Rochester, Rochester, NY
2017 – present	Affiliated Faculty, Center for Biomedical Informatics,
	University of Rochester Medical Center, Rochester, NY
2018 – present	Member, Environmental Health Sciences Center,
	University of Rochester Medical Center, Rochester, NY

### **HONORS AND AWARDS**

2004	Honors College Concentration in Statistics with Highest Distinction,
	University of Michigan
2004	Phi Beta Kappa, University of Michigan
2004 - 2005	Intramural Research Training Award Fellow, NIH/NHGRI, Cancer Genetics Branch
2005 - 2007	Trainee, NIH Training Grant in the Epidemiology and Biostatistics of Aging,
	Johns Hopkins School of Public Health
2007 - 2010	Trainee, NIH Training Grant in Biostatistics for Genetics/Genomics,

2008 Travel Award, International Meeting of the Microarray and Gene Expression Data

Society

2009 & 2010 Teaching Assistant Recognition Award Nominee, Johns Hopkins School of Public Health Helen Abbey Award for Excellence in Teaching, Johns Hopkins School of Public Health

2010 – 2013 Trainee, NIH Training Grant in Experimental Therapy Models,

University of Rochester School of Medicine & Dentistry

2011 Conference Fellowship, q-bio Conference on Cellular Information Processing

2011 Conference Scholarship, Bioconductor Conference (BioC2011)

2015 Graduate Student Society Advocacy Award,

University of Rochester School of Medicine & Dentistry

# **ACADEMIC & PROFESSIONAL ORGANIZATIONS**

2009 – present Member, American Statistical Association (ASA)

2009 – present Member, The International Biometric Society, Eastern North American Region (ENAR)

2009 – present Member, ASA Biometrics Section
2014 – present Member, Rochester Chapter of the ASA

2015 – present Member, ASA Section on Statistics in Genomics and Genetics

#### COMMITTEES & OTHER ADMINISTRATIVE SERVICE

# UR Department of Biostatistics & Computational Biology

2015 - 2017	Chair, Biostatistics and Computational Biology Departmental Seminar Series
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2015 – present Member, Bioinformatics Concentration Curriculum Committee

2016 – present Member, Statistics Graduate Admissions Committee

2016 – present

2018 – present

Member, Biostatistics and Computational Biology Faculty Search Committee

Member, Biostatistics and Computational Biology Computing Committee

Member, Biostatistics and Computational Biology Curriculum Committee

2019 – present Member, Internal Advisory Board / Executive Committee for NIH T32 ES007271

2020 – present Associate Program Director, Statistics PhD Program

#### UR Extra-Departmental

2015 2016	$\alpha$ 1 .	D 1 C .	← 11	
2015 - 2016	Co-chair	Data Science	Colloc	illilim Series
2015 2010	Co onan,	Data Science	COHOC	aram Series

2015 – 2019 Judge, Graduate Student Society Poster Competition

2016 – 2018 Member, CTSI Informatics Education Cluster

2017, 2019, 2020 Reviewer, University Research Awards 2019 Reviewer, SAC Incubator Awards

2019 Reviewer, UNYTE Pipeline-to-Pilot Program

2017 – present Member, Center for Biomedical Informatics and Wilmot Cancer Institute Faculty Search

Committee

2018 – present Member, Biomedical Informatics Curriculum Committee

2019 – present Member, Bioinformatics Working Group Planning Committee

2019 – present Participant, Driving our Future: Culture, Climate and Leadership in the Sciences

2019 – present Member, Wilmot Cancer Institute Scientific Leadership Committee

2019 – present Co-director, Wilmot Cancer Institute Biostatistics and Bioinformatics Shared Resource

2019 – present Co-chair, GIDS Working Group on Life Sciences and Biomedical Data Science

#### Local/Regional

2018 – 2019 Member, UP-STAT Organizing Committee
 2018 Member, UP-STAT Data Analytics Committee

2016, 2018, 2019 Session Chair, UP-STAT Conference

2018 - 2019	Judge, UP-STAT Conference Studen	nt Recearch Awards
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2012 - 2013	Judge, ASA/NCTM Statistical Project Competition
2016 – present	Member, Patient-Centered Outcomes Research Institute (PCORI) Reviewer Community
2017 - 2019	NIH Early Career Reviewer Program
2018 - 2019	Reviewer, CTSI Novel Biostatistical and Epidemiologic Methodology pilot program
2018	Reviewer, NIH Biodata Management and Analysis (BDMA) Study Section
2018 - 2019	Member, Bioconductor Conference (BioC2019) Organizing Committee
2019	Invited Session Organizer, Symposium on Data Science & Statistics
2019 - 2020	Member, Bioconductor Conference (BioC2020) Organizing Committee

# **EDUCATIONAL CONTRIBUTIONS**

# **Research Mentoring**

# Postdoctoral Fellows

2014 - 2015	Qingyan Lilly Xie, Ph.D.; Primary Mentor
2016 - 2017	Isar Nassiri, Ph.D.; Primary Mentor
2020 – present	Hongying Sun, Ph.D.; Primary Mentor

# **Graduate Students**

2015 - 2019	Valeriia Sherina	Statistics Ph.D.	Program;	Thesis Advisor (	co-advisor: Dr.	Tanzy Love)

2018 – present	David Burton, Statistics Ph.D. Program; Thesis Advisor
2018 – present	Zachary Brehm, Statistics Ph.D. Program; Thesis Advisor
2020 – present	Erik Vonkaenel, Statistics Ph.D. Program; Thesis Advisor

# **Graduate Research Assistants**

2015 - 2018	Yun Zhang, Statistics Ph.D. Program
2016 (fall)	Kejia Wang, Statistics M.A. Program
2017 (summer)	Jeremiah Jones, Statistics Ph.D. Program
2019 (summer)	Erik Vonkaenel, Statistics Ph.D. Program
2019 (fall)	Bokai Wang, Statistics Ph.D. Program
2020 (summer)	Sami Leon, Statistics Ph.D. Program

### **Rotation Students**

	2015 (fall)	Rokai Wang	Statistics Ph D	Program
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2018 (spring)	Mukta Palshikar, Biophysics, Structural & Computational Biology Ph.D. Program
2019 (spring)	Abhinav Mittal, Biophysics, Structural & Computational Biology Ph.D. Program

#### Undergraduate Students

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2015 (summer)	Lindsey Barden, University of North Carolina at Chapel Hill
2015 (summer)	Allison Maier, State University of New York at Geneseo
2015 (summer)	Scott Onestak, University of Rochester
2015 - 2017	Lauren Kemperman, University of Rochester
2015 - 2017	Jeffrey Hrebenach, University of Rochester
2016 (summer)	Alida Mooney, State University of New York at Geneseo
2016 - 2017	Winslow Powers, University of Rochester
2016 - 2018	Jonavelle Cuerdo, University of Rochester
2017 - 2019	Benjamin Hsu, University of Rochester
2018 (fall)	Oleg Demus, University of Rochester
2017 - 2019	Qidi Yang, University of Rochester

2018 - 2019	Kailey Ferger, University of Rochester
2019 (summer)	Joshua Dennis, University of Rochester
2019 (fall)	Daniel Munoz Huerta, University of Rochester
2019 – present	Yuewei Fei, University of Rochester
2019 – present	Michael Lansford, University of Rochester
2019 – present	Benyu Zhou, University of Rochester
2020 – present	Frank Gonzalez, University of Rochester

# High School Students

2017 (summer)	Fatima Umar, Pittsford Schools
2017 (summer)	Raymond Feng, Pittsford Schools
2019 (summer)	Lillian Stolberg, Pittsford Schools

# **Classroom Teaching**

2012	BST520, Current Topics in Bioinformatics (course co-designer and co-instructor)
2014	BST512, Advanced Bayesian Inference with an Emphasis on Computation (one lecture)
2014	BST432, Introduction to Bioinformatics (one lecture)
2014 - 2018	IND501, Ethics and Professional Integrity in Research (small group facilitator)
2015	GEN508, Genomics and Systems Biology (two lectures)
2015	GEN504, Genetics Seminar (one lecture)
2015	BST550, Genomic Data Analysis Using R/Bioconductor (course designer and instructor)
2016 – present	IND419, Introduction to Quantitative Biology (two to five lectures)
2016 - 2017	BST494/BST430, Statistical Computing (course designer and instructor)
2018 – present	BST487, Seminar in Statistical Literature (topic supervisor)
2019	BST494, Genomic Data Analysis (course designer and instructor)
2020	BST570, Statistical Analysis of Cell Mixtures (course designer and instructor)

# **Short Courses & Workshops**

2018	Grad STEM for All Summer Program (instructor and program mentor)
2018	Genomic Data Analysis Workshop (course designer and instructor)

# **Thesis Committee Service**

2014 - 2017	Justin Komisarof (Genetics, Development, and Stem Cells Ph.D. Program)
2016 - 2018	Yun Zhang (Statistics Ph.D. Program)
2017 - 2020	Rohith Palli (Biophysics, Structural & Computational Biology Ph.D. Program)
2017 – present	Adam Cornwell (Genetics, Development, and Stem Cells Ph.D. Program)
2017 – present	Derek Crowe (Genetics, Development, and Stem Cells Ph.D. Program)
2017	Adan Becerra (Epidemiology Ph.D. Program); chair, Thesis Defense Committee
2018 - 2020	Matthew Turner (Genetics, Development, and Stem Cells Ph.D. Program)
2018	Kyle Berger (Biophysics, Structural & Computational Biology Ph.D. Program);
	chair, Thesis Defense Committee
2019 – present	Jiatong Sui (Statistics Ph.D. Program)

# **CONSULTATIONS**

2008 - 2009	University of Miami Miller School of Medicine, Miami, Florida, USA
2010 - 2011	GenomeDx Biosciences Inc, Vancouver, British Columbia, Canada
2011 - 2013	Janssen Research and Development, Beerse, Belgium
2012 - 2013	OpenAnalytics, Heist-op-den-Berg, Belgium
2013	Selventa Inc, Cambridge, Massachusetts, USA

#### EDITORIAL ASSIGNMENTS IN PROFESSIONAL JOURNALS

Ad hoc reviews for: Annals of Applied Statistics, APJ Renal Physiology, Bioinformatics, Biometrics, Biotechniques, biOverlay, BMC Bioinformatics, BMC Cancer, BMC Genomics, BMC Medical Genomics, Cancer Informatics, Database, Epigenomics, F1000Research, Gene, IEEE/ACM Transactions on Computational Biology and Bioinformatics, IMS Collections, iScience, Journal of the American Statistical Association, Journal of Postdoctoral Research, Microarrays, Nature Methods, Nucleic Acids Research, NAR Genomics and Bioinformatics, PLoS Computational Biology, PLoS Genetics, PLoS One, Proceedings of the National Academy of Sciences, Science, Scientific Reports, Statistical Applications in Genetics and Molecular Biology, Statistics in Biosciences

# **Editorial Assignments**

2012 – 2014 Editor, Journal of Postdoctoral Research 2015 – 2020 Review Editor, Frontiers in Genetics 2018 – present Associate Editor, Biostatistics

### **GRANTS & CONTRACTS**

# As Principal Investigator / Co-Principal Investigator

NIH R01 GM139928 (PI: McCall)

Statistical Methods for MicroRNA-seq Experiments

09/11/20-06/30/25; Total Award: \$1,975,065

The overall objective of this application is to improve the analysis of sRNA-seq data by developing statistical methods that account for challenges specific to sRNA-seq data.

### NIH R00 HG006853 (PI: McCall)

Statistical Methods for Estimation of Gene Regulatory Networks

09/01/14 - 08/31/18 (NCE); Total Award: \$726,419

The overall goals of the proposed research are: to improve the estimation of gene regulatory networks from perturbation experiments, by using methods that explicitly model and incorporate uncertainty in each step of the process, and to use these estimated networks to predict cellular response to intervention.

#### NIH K99 HG006853 (PI: McCall)

Statistical Methods for Estimation of Gene Regulatory Networks

09/10/13 - 08/31/14; Total Award: \$79,893

Responsibilities: This Award provided support to obtain the expertise required to address the proposed research aims and transition to an independent research career through a combination of coursework, mentorship, and research experience.

### UL1 TR002001 Pilot Studies Award (PI: McCall)

Development of qPCR Methodology for Clinical Testing

08/15/16 - 05/31/17; Total Award: \$35,000

The overall goals of the proposed research are: (1) to develop improved methodology to handle non-detects in qPCR data, (2) to develop a single sample version of our methodology for clinical biomarkers, and (3) to assess the applicability of our methods to microRNA transcriptome screening.

#### UL1 TR002001 Pilot Studies Award (PI: McCall)

Estimation of cell-type specific microRNA expression in complex tissue samples

06/01/17 - 05/31/18; Total Award: \$35,000

The overall goal of the proposed research is to develop statistical deconvolution methodology to estimate the cellular composition and cell-type specific microRNA expression of tissue samples.

UL1 TR002001 Pilot Studies Award (PI: McCall)

Statistical methods to quantify imaged microglia

06/01/19 - 05/31/20; Total Award: \$35,000

The overall goal of the proposed research is to improve the analysis of microglial morphology by developing statistical methods to quantify images of microglia.

NIH R01 HL137811 (PI: Halushka; Subcontract-PI: McCall)

Fine Dissection of Atherosclerosis Microenvironment RNA Expression

09/01/17 – 08/30/22; Annual Direct Costs to McCall Lab: \$65,866/year

This project will identify genes, long noncoding RNAs and microRNAs that are involved in the development of coronary artery atherosclerosis.

NIH R01 GM083084 (PI: Irizarry; Subcontract-PI: McCall)

Preprocessing and Analysis Tools for High-Throughput Technologies

09/01/16 - 06/30/20; Annual Direct Costs to McCall Lab: \$25,000/year

The goal of the proposed research is to leverage the public data repositories to gain insights into the stochastic nature of measurement and develop methods to model application-specific systematic bias and unwanted variability. The proposed research requires the use of a vast amount of publicly available data across several high-throughput platforms.

#### Other Roles

Invidumed GmbH (PI: Land)

Global Cancer Database Project

01/01/18 - 12/31/19; Percent Effort: 20%

The URMC and INDIVUMED bioinformatics collaboration will focus on the development of a multi-dimensional analytics platform (Portal Solution) linked to a global biospecimen and clinical data resource controlled for molecular integrity, comprehensive data annotation and diversity of patient population. The Global Cancer Data Base Solution (GCDS) will result from the convergence of the Portal Solution and a global biospecimen resource, allowing for an investigator, regardless of their analytical bias (e.g. genomics, proteomics, metabolomics, immune-profiling, etc.,) to interrogate research hypothesis. The GCDS, when developed should represent the most comprehensive analytical tool to interpret and utilize cancer phenomic data linked to a dynamic and growing pool of molecular and phenomic data curated from patients on a global scale.

### NIH R35 CA197562 (PI: Land)

Mediators of Cancer Cell Homeostasis: Intervention Targets Common to Diverse Types of Cancer 09/01/15 - 08/31/22; Percent Effort: 10%

The goals of this research program are to develop a rational path towards cancer interventions independent of the tumors' mutational status and with lower rates of associated disease recurrence that can be delivered to a large fraction of patients.

NIH HHSN272201200005C (PI: Topham)

NIAID Respiratory Pathogens Research Center (RPRC) at the University of Rochester

12/13/11 – 12/12/18; Percent Effort: 10%

This center's purpose is to establish, direct, and maintain the UR-RPRC, a multidisciplinary, collaborative, integrated, and iterative program focused on the conduct of research activities to understand the etiology and impact of respiratory pathogens and to develop and evaluate strategies for their control, treatment and prevention. Specifically, we will provide and maintain a multidisciplinary team of experienced investigators and technical staff with expertise in clinical research and laboratory methods, and experience working with respiratory pathogens and the diseases they cause.

NIH R01 HL127891 (PI: Brookes)

The Role of the Mitochondrial UPR in Ischemic Protection

06/01/15 - 02/28/19; Percent Effort: 5%

This proposal aims to identify mechanisms through which the UPRmt protects C. elegans and to translate these findings to a mammalian cardiac model. Our approach will include defining mechanistic crosstalk with other signaling pathways that are also protective and will result in the identification of functional orthologs that perform similarly in mammals as ATFS-1 and HAF-1.

NIH U54 CA228110 (PI: Rahman)

WNY Center for Research on Flavored Tobacco Products

09/01/2018 - 06/30/2023; Precent Effect: 5%

We propose a Center for Research on Flavored Tobacco Products (CRoFT), the goal of which is to develop a novel framework and approaches for assessing the impact of tobacco product flavors and flavorings on consumer behavior, exposures, and health.

NIH P30 ES001247 (PI: Lawrence)

Environmental Agents as Modulators of Disease Processes

04/01/2019 - 03/31/2021; Percent Effort: 5%

The mission of the EHSC at Rochester is to improve public health through the generation of fundamental knowledge and elaboration of mechanisms by which chemical exposures, alone or through interaction with other modifying factors, contribute to cumulative health risk across the lifespan.

NIH R01 NS114480 (PI: Majewska)

Mechanisms that regulate microglial dynamics in the context of plasticity

01/01/2020 - 12/31/2024; Percent Effort: 5%

The goals of this research program are to investigate microglial mechanisms that govern changes in neuronal networks. This will yield information with broad implications for understanding and treating a large spectrum of human neurological disorders.

NIH T32 ES007271 (PI: Thurston)

Training in Environmental Health Biostatistics

09/30/15 - 06/30/20

I am a research mentor on this training grant; no salary support.

#### **PRESENTATIONS**

### **National & International Conferences**

May 2005	A Novel Algorithm for Microarray Time Series Data Analysis
	IEEE International Workshop on Genomic Signal Processing and Statistics, Newport, RI
Sept 2008	An Improved Gene Expression Barcode based on a Hierarchical Mixture Model
	International Meeting of MGED, Riva del Garda, Italy
July 2009	From a Single CEL File to a Gene Expression Barcode
	Bioconductor Conference (BioC2009), Seattle, WA
Mar 2010	Gene Expression Barcodes
	ENAR Spring Meeting, New Orleans, LA
July 2011	Thawing Frozen Robust Multi-array Analysis (fRMA)
	Bioconductor Conference (BioC2011), Seattle, WA
Aug 2011	Assessing Microarray Quality
	Joint Statistical Meeting, Miami Beach, FL
Aug 2011	A Computational Bayesian Approach for the Inference of Boolean Networks

Q-Bio Conference in Cellular Information Processing, Santa Fe, NM

July 2012	Batch effect reduction using the frma package
•	Bioconductor Conference (BioC2012), Seattle, WA
Aug 2013	Addressing Within-Subject Genomic Heterogeneity
	Joint Statistical Meeting, Montreal, Canada
Feb 2015	A Gene Regulatory Network of Cooperation Response Genes Governs the Cancer Phenotype
	Winter q-bio Meeting, Maui, HI
June 2015	Estimation of a Regulatory Network of Cooperation Response Genes in a Model of Cancer
	Malignancy, Statistical Learning of Biological Systems from Perturbations, Ascona, Switzerland
May 2016	Estimation of a Regulatory Network of Cooperation Response Genes in a Model of Cancer
	Malignancy, Great Lakes Bioinformatics Conference, Toronto, Canada
July 2016	Complex Sources of Variation in Tissue Expression Data: Analysis of the Genotype-Tissue
	Expression (GTEx) Lung Transcriptome, GTEx Project Community Meeting, Stanford, CA

# **Invited Presentations: Local**

Apr 2008	Cross-platform comparisons using microarray spike-in data The Expressionist Seminar Series, Baltimore, MD
Oct 2009	Barcodes for Beginners
	The Expressionist Seminar Series, Baltimore, MD
Mar 2011	What can I do with a single microarray?
	Transcriptomics and Integrated Genomics Meeting, Rochester, NY
May 2012	Harnassing heterogeneity to improve genomic biomarkers
	Transcriptomics and Integrated Genomics Meeting, Rochester, NY
Sept 2012	Batch effects in genomic data
	Transcriptomics and Integrated Genomics Meeting, Rochester, NY
Dec 2013	Gene Regulatory Network Estimation
	Transcriptomics and Integrated Genomics Meeting, Rochester, NY
Sept 2014	On Non-Detects in qPCR Data
	Transcriptomics and Integrated Genomics Meeting, Rochester, NY
Mar 2015	Overview of McCall Group Research
	Transcriptomics and Integrated Genomics Meeting, Rochester, NY
Apr 2015	Incorporating Uncertainty in the Estimation of Gene Regulatory Networks
	Center for Integrated Research Computing Symposium Series, Rochester, NY
Apr 2015	Challenges in Translational Genomics
	Workshop sponsored by the Graduate Student Society, Rochester, NY
Dec 2015	Mechanics of RNA-Seq Data Analysis Using R
	Workshop sponsored by the Center for Professional Development, Rochester, NY
Oct 2016	Sources of Variation in GTEx Lung Tissue Expression Data
	Lung Biology Research Seminar Series, Rochester, NY
Jan 2017	Incorporating Uncertainty in the Estimation of Gene Regulatory Networks
	Bioinformatics Cluster, Rochester, NY
Mar 2017	Sources and Effects of Transcriptomic Heterogeneity
	Center for Biomedical Informatics, Rochester, NY
Dec 2017	What can we learn from co-expression analyses?
F 1 2010	Transcriptomics and Integrated Genomics Meeting, Rochester, NY
Feb 2019	Co-expression analysis of complex tissues
	PhD Applicant Weekend, Rochester, NY

# **Invited Presentations: National & International**

Sept 2008 EMERALD Workshop on Array Quality Assessment Methods International Meeting of MGED, Riva del Garda, Italy

Feb 2011	Frozen Robust Multi-array Analysis (fRMA)
	La Calestienne Meeting, Nismes, Belgium
Oct 2011	Microarray Preprocessing and Quality Control
	La Calestienne Meeting, Nismes, Belgium
Oct 2011	MicroRNA Microarray Performance
	Janssen Research and Development, Beerse, Belgium
Apr 2012	Batch Effects and Splice Variants
	La Calestienne Meeting, Nismes, Belgium
Aug 2012	Summarization of Affymetrix GeneChip Probe Level Data: from RMA to fRMA
	International Biometric Conference, Kobe, Japan
Oct 2012	fRMA for Exon / Gene ST Arrays
	La Calestienne Meeting, Nismes, Belgium
July 2013	A Computational Bayesian Approach to Gene Regulatory Network Estimation
	Bioconductor Conference (BioC2013), Seattle, WA
June 2015	Modeling qPCR Non-Detects as Missing Data
	qPCR and Digital PCR Congress: USA, San Diego, CA
Dec 2016	Incorporating Uncertainty in the Estimation of Gene Regulatory Networks
	University of Buffalo Biostatistics Seminar Series, Buffalo, NY
Aug 2017	It's all about the residuals: estimating gene networks by leveraging residual structure in multi-
	tissue transcriptomic data
	Joint Statistical Meeting, Baltimore, MD
Mar 2018	How Amazing Mentors Think About Mentoring
	Webinar organized by the Center for the Integration of Research, Teaching and Learning
May 2019	Case Studies in Interoperability: From Generic Classes To Specific Functions
	Symposium on Data Science & Statistics, Bellevue, WA
Aug 2020	Co-Expression Analysis of Complex Tissues
	Joint Statistical Meeting, Virtual Conference

#### **PUBLICATIONS**

 $^*$  denotes trainee / supervisee;  $^\circ$  denotes senior statistician;  $^\dagger$  denotes corresponding author(s)

# **Pre-prints**

- 1. **Sherina V**\*, McMurray HR, **Powers W**\*, Land H, Love TMT, and **McCall MN**†° (2017). Statistical Approaches to Decreasing the Discrepancy of Non-detects in qPCR Data. bioRxiv. doi: <a href="http://dx.doi.org/10.1101/231621">http://dx.doi.org/10.1101/231621</a>
  [ENAR Distinguished Student Paper Award to V. Sherina]
- 2. **Kemperman L**\* and **McCall MN**†° (2017). *miRcomp-Shiny*: Interactive assessment of qPCR-based microRNA quantification and quality control algorithms. F1000Research. 6:2046.
- 3. Wang L, Chu CY, **McCall MN**, Slaunwhite C, Holden-Wiltse J, Corbett A, Falsey AR, Topham DJ, Caserta MT, Mariani TJ, Walsh EE, and Qiu X (2019). Airway Gene-Expression Classifiers for Respiratory Syncytial Virus (RSV) Disease Severity in Infants. bioRxiv. doi: <a href="https://doi.org/10.1101/628701">https://doi.org/10.1101/628701</a>
- 5. **Hsu B\***, **Sherina V\***, **McCall MN**<sup>†</sup>° (2019). Auto-regressive modeling and diagnostics for qPCR amplification. bioRxiv. doi: <a href="https://doi.org/10.1101/665596">https://doi.org/10.1101/665596</a>
- 6. **Sherina V**\*, **McCall MN**, Love TMT<sup>†°</sup> (2019). Fully Bayesian imputation model for non-random missing data in qPCR. arXiv:1910.13936.
- 7. Fomchenko KM, Verma RX, Kannan S, Lin BL, Yang X, Nieuwenhuis TO, Patil AH, Fox-Talbot K, **McCall MN**, Kwon C, Kass DA, Rosenberg AZ, Halushka MK<sup>†</sup> (2020). Proteogenomic single cell analysis of skeletal muscle myocytes. bioRxiv. doi: https://doi.org/10.1101/2020.01.23.916791

8. Mendes MS, Atlas J, **Brehm Z**\*, Ladron-de-Guevara A, **McCall MN**°, Majewska AK† (2020). In vivo imaging of the kinetics of microglial self-renewal and maturation in the adult visual cortex. bioRxiv. doi: https://doi.org/10.1101/2020.03.05.977553

### **Peer-Reviewed Journal Articles**

- 1. Chu CY, Qiu X, **McCall MN**, Wang L, Corbett A, Holden-Wiltse J, Slaunwhite C, Grier A, Gill SR, Pryhuber GS, Falsey AR, Topham DJ, Caserta MT, Walsh EE, Mariani TJ (2020). Airway gene expression correlates of RSV disease severity and microbiome composition in infants, *The Journal of Infectious Diseases*, jiaa576, doi: https://doi.org/10.1093/infdis/jiaa576
- 2. Nieuwenhuis TO, Yang S, Verma RX, Pillalamarri V, Arking D, Rosenberg AZ, **McCall MN**°, Halushka MK<sup>†</sup> (2020). Consistent RNA sequencing contamination in GTEx and other data sets. *Nat Communications*, 11:1933.
- 3. **Zhang Y\***, **Cuerdo J\***, Halushka MK, **McCall MN**<sup>†</sup>° (2019). The effect of tissue composition on gene co-expression. *Briefings in Bioinformatics*, bbz135, doi: https://doi.org/10.1093/bib/bbz135
- 4. Wang YT, Lim Y, **McCall MN**°, Huang KT, Haynes CM, Nehrke KW, Brookes PS<sup>†</sup> (2019). Cardioprotection by the mitochondrial unfolded protein response requires ATF5. *American Journal of Physiology-Heart and Circulatory Physiology*, 317(2):H472-H478.
- 5. Walsh EE<sup>†</sup>, Mariani TJ, Chu C, Grier A, Gill SR, Qiu X, Wang L, Jeanne HW, Corbett A, Thakar J, Benoodt L, **McCall MN**, Topham DJ, Falsey AR, Caserta MT (2019). Aims, study design and enrollment results from the Assessing Predictors of Infant Respiratory Syncytial Virus Effects and Severity (AsPIRES) study. *JMIR Research Protocols*, 8(6):e12907.
- 6. Spinelli P, Latchney SE, Reed JM, Fields A, Baier BS, Lu X\*, McCall MN°, Murphy SP, Mak W, Susiarjo M† (2018). Identification of the novel Ido1 imprinted locus and its potential epigenetic role in pregnancy loss. *Human Molecular Genetics*, 28(4):662-674.
- 7. **Nassiri I**\* and **McCall MN**†° (2018). Systematic exploration of cell morphological phenotypes associated with a transcriptomic query. *Nucleic Acids Research*, 46(19):e116.
- 8. Rosenberg AZ, Wright C, Fox-Talbot K, Rajpurohit A, Williams C, Porter C, Kovbasnjuk O, **McCall MN**°, Shin JH, Halushka MK<sup>†</sup> (2018). xMD-miRNA-seq to generate near in vivo miRNA expression estimates in colon epithelial cells. *Scientific Reports*, 8(1):9783.
- 9. Hill MA, Alexander WB, Guo B, Kato Y, Patra KC, O'Dell MR, McCall MN°, Whitney-Miller CL, Bardeesy N, and Hezel AF† (2018). Kras and Tp53 mutations cause cholangiocyte- and hepatocyte-derived cholangiocarcinoma. *Cancer Research*, 78(16):4445-4451.
- 10. Halushka MK<sup>†</sup>, Fromm B, Peterson KJ, **McCall MN°** (2018). Big Strides in Cellular MicroRNA Expression. *Trends in Genetics*, 34(3):165-167.
- 11. **McCall MN°**, Kim MS, Adil M, Patil AH, Lu Y, Mitchell CJ, Leal-Rojas P, Xu J, Kumar M, Dawson VL, Dawson TM, Baras AS, Rosenberg AZ, Arking DE, Burns KH, Pandey A, Halushka MK<sup>†</sup> (2017). Toward the human cellular microRNAome. *Genome Research*, 27(10):1769-1781.
- 12. Li N, **McCall MN**, Wu Z<sup>†°</sup> (2017). Establishing Informative Prior for Gene Expression Variance from Public Databases. *Statistics in Biosciences*, 9(1):160-177.
- 13. Komisarof J, **McCall MN**°, Newman L, Bshara W, Mohler JL, Morrison C, Land H<sup>†</sup> (2017). A Four Gene Signature Predictive of Recurrent Prostate Cancer. *Oncotarget*. 8(2):3430-3440.
- 14. **McCall MN**<sup>†</sup>°, Illei PB, Halushka MK<sup>†</sup> (2016). Complex Sources of Variation in Tissue Expression Data: Analysis of the GTEx Lung Transcriptome. *The American Journal of Human Genetics*, 99(3):624-635.
- 15. **Xie QY**\*, Almudevar A, Whitney-Miller CL, Barry CT, **McCall MN**<sup>†°</sup> (2016). A microRNA biomarker of hepatocellular carcinoma recurrence following liver transplantation accounting for within-patient heterogeneity. *BMC Medical Genomics*, 9(1):18.

- 16. **McCall MN**<sup>†°</sup>, Baras AS, Crits-Christoph A, Ingersoll R, McAlexander MA, Witwer KW, Halushka MK (2016). A benchmark for microRNA quantification algorithms using the OpenArray platform. *BMC Bioinformatics*, 17(1):138.
- 17. Verbist BM, Verheyen GR, Vervoort L, Crabbe M, Beerens D, Bosmans C, Jaensch S, Osselaer S, Talloen W, Van den Wyngaert I, Van Hecke G, Wuyts D, **Qstar Consortium**, Van Goethem F, Gohlmann HW (2015). Integrating High-Dimensional Transcriptomics and Image Analysis Tools into Early Safety Screening: Proof of Concept for a New Early Drug Development Strategy, *Chemical Research in Toxicology*, 28(10):1914-25.
- 18. Verbist B, Klambauer G, Vervoort L, Talloen W, **Qstar Consortium**, Shkedy Z, Thas O, Bender A, Gohlmann HWH, Hochreiter S (2015). Using transcriptomics to guide lead optimization in drug discovery projects: Lessons learned from the QSTAR project, *Drug Discovery Today*, Volume 20, Issue 5, Pages 505-513.
- 19. Ravindranath AC, Perualila-Tan N, Kasim A, Drakakis G, Liggi S, Brewerton SC, Mason D, Bodkin MJ, Evans DA, Bhagwat A, Talloen W, Gohlmann HW, **Qstar Consortium**, Shkedy Z, Bender A (2015). Connecting gene expression data from connectivity map and in silico target predictions for small molecule mechanism-of-action analysis, *Molecular Biosystems*, 11(1):86-96.
- 20. Kent O, **McCall MN**°, Cornish T, Halushka MK<sup>†</sup> (2014). Lessons from miR-143/145: The importance of cell-type localization of miRNAs, *Nucleic Acids Research*, 42(12):7528-7538.
- 21. **McCall MN**<sup>†</sup>, McMurray H, Land H, Almudevar A° (2014). On Non-Detects in qPCR Data, *Bioinformatics*, 30(16):2310-2316.
- 22. Haider B, Baras AS, **McCall MN**°, Hertel JA, Cornish TC, Halushka MK<sup>†</sup> (2014). A critical evaluation of microRNA biomarkers in non-neoplastic disease, *PLoS ONE*, 9(2): e89565.
- 23. **McCall MN**°, Jaffee HA, Zelisko SJ, Sinha N, Hooiveld G, Irizarry RA, Zilliox MJ<sup>†</sup> (2014). The Gene Expression Barcode 3.0: improved data processing and mining tools, *Nucleic Acids Research*, 42(D1):D938-D943.
- 24. Tran V, **McCall MN**<sup>†</sup>, McMurray H, Almudevar A° (2013). On the underlying assumptions of threshold Boolean networks as a model for genetic regulatory network behavior, *Frontiers in Genetics*, 4:263.
- 25. Wu G, Yustein JT, **McCall MN**, Zilliox M, Irizarry RA, Zeller K, Dang CV, Ji H<sup>†</sup>° (2013). ChIP-PED enhances the analysis of ChIP-seq and ChIP-chip data, *Bioinformatics*, 29(9):1182-1189.
- 26. **McCall MN**<sup>†°</sup> (2013). Estimation of Gene Regulatory Networks, *Journal of Postdoctoral Research*, 1(1):60-69.
- 27. Bravo HC<sup>†</sup>°, Pihur V, **McCall MN**, Irizarry RA, Leek JT (2012). Gene expression anti-profiles as a basis for accurate universal cancer signatures, *BMC Bioinformatics*, 13:272.
- 28. **McCall MN**<sup>†</sup>, Jaffee HA, Irizarry RA° (2012). fRMA ST: Frozen robust multiarray analysis for Affymetrix Exon and Gene ST arrays, *Bioinformatics*, 28 (23): 3153-3154.
- 29. **McCall MN**<sup>†</sup> and Almudevar A° (2012). Affymetrix GeneChip microarray preprocessing for multivariate analyses, *Briefings in Bioinformatics*, 13(5):536-546.
- 30. Barry CT<sup>†</sup>, D'Souza M, **McCall MN**, Safadjou S, Ryan C, Kashyap R, Marroquin C, Orloff M, Almudevar A°, Godfrey TE (2012). Micro RNA expression profiles as adjunctive data to assess the risk of hepatocellular carcinoma recurrence after liver transplantation, *American Journal of Transplantation*, 12(2):428-437.
- 31. **McCall MN°**, Kent OA, Yu J, Fox-Talbot K, Zaiman AL, Halushka MK<sup>†</sup> (2011). MicroRNA profiling of diverse endothelial cell types, *BMC Medical Genomics*, 4:78.
- 32. Almudevar A<sup>†°</sup>, **McCall MN**, McMurray H, Land H (2011). Fitting Boolean Networks from Steady State Perturbation Data, *Statistical Applications in Genetics and Molecular Biology*, 10(1): Article 47.
- 33. **McCall MN** and Irizarry RA<sup>†o</sup> (2011). Thawing Frozen Robust Multi-array Analysis (fRMA), *BMC Bioinformatics*, 12:369.

- 34. **McCall MN**, Murakami PN, Lukk M, Huber W, Irizarry RA<sup>†°</sup> (2011). Assessing Affymetrix GeneChip Microarray Quality, *BMC Bioinformatics*, 12:137.
- 35. **McCall MN**, Uppal K, Jaffee HA, Zilliox MJ<sup>†</sup>, and Irizarry RA<sup>†o</sup> (2011). The Gene Expression Barcode: leveraging public data repositories to begin cataloging the human and murine transcriptomes, *Nucleic Acids Research*, 39:D1011-5.
- 36. Shi L<sup>†</sup>, Campbell G, Jones WD, ..., **McCall MN**, ..., Scherf U, Tong W, Wolfinger RD; MAQC Consortium (2010). The MicroArray Quality Control (MAQC)-II study of common practices for the development and validation of microarray-based predictive models, *Nature Biotechnology*, 28(8):827-38.
- 37. **McCall MN**, Bolstad BM, and Irizarry RA<sup>†°</sup> (2010). Frozen Robust Multi-Array Analysis (fRMA), *Biostatistics*, 11(2):242-253.
- 38. Sah S, **McCall MN**, Eveleigh D, Wilson M, and Irizarry RA<sup>†o</sup> (2010). Performance Evaluation of Commercial miRNA Expression Array Platforms, *BMC Research Notes*, 3:80.
- 39. **McCall MN** and Irizarry RA<sup>†o</sup> (2008). Consolidated strategy for the analysis of microarray spike-in data, *Nucleic Acids Research*, 36(17), e108.

### Peer-Reviewed R/Bioconductor Software Packages

- 1. frma: Single microarray preprocessing and analysis
- 2. frmaTools: Extension and customization of the frma package
- 3. miRcomp: Tools to assess and compare miRNA expression estimation methods
- 4. nondetects: Methods to model and impute non-detects in qPCR data
- 5. spkTools: Methods for microarray spike-in data
- 6. ternarynet: Computational Bayesian approach to ternary network estimation

# Peer-Reviewed R/Bioconductor Data Packages

- 1. microRNAome: This package provides a SummarizedExperiment object of read counts for microRNAs across tissues, cell-types, and cancer cell-lines.
- 2. miRcompData: Raw amplification data from a large microRNA mixture / dilution study. These data are used by the miRcomp package to assess the performance of methods that estimate expression from the amplification curves.
- 3-12. <platform>frmavecs: data packages containing the frozen parameter vectors used by the frma package for 10 different microarray platforms.