Sarah A Gagliano Taliun

Sarah<dot>Gagliano<at>icm-mhi<dot>org; https://sgagliano.github.io

Research Focus

My research seeks to identify and better understand the genetic contributors to complex traits, using "big data" *in silico*, particularly disorders of the brain. My goal is to increase knowledge on the genetic aetiology of complex traits to allow for better diagnoses and more personalized treatment.

Academic Positions

Principal Investigator, Montréal Heart Institute, Montréal, Canada
 July 2020 – present

 Assistant Professor/professeure sous octroi adjointe,
 Departments of Medicine and Neurosciences, Université de Montréal, Montréal, Canada

 Honorary Senior Research Associate, UCL Institute of Neurology, Neurodegenerative Diseases

Education and Training

 Postdoctoral Research Fellow, School of Public Health, University of Michigan, Ann Arbor, USA
 Supervisors: Gonçalo Abecasis and Michael Boehnke

Weston Brain Institute International Fellowship in Neuroscience
 December 2015 – May 2016

 Supervisor: Michael Weale, King's College London

Project: In silico identification of genetic risk variants for Parkinson's disease

 PhD, Medical Science, Institute of Medical Science, University of Toronto, Canada Supervisors: James Kennedy and Jo Knight

Research conducted at Centre for Addiction and Mental Health (CAMH)

Dissertation: In silico prioritization of genetic risk variants using functional genomic information

• **Honours BSc**, Biochemistry & Human Biology with high distinction, University of June 2012 Toronto, Canada

Funding and Awards

Programme de Bourses de Chercheur-boursier Junior 1 en Intelligence Artificielle 2020-2024 et Santé Numérique, Fonds de la Recherche en Santé du Québec (FRQS)
 Project: Le développement des modèles prédictifs qui sont spécifiques au sexe pour la médecine personnalisée en santé cardiovasculaire

• Precision Health Scholars Award, University of Michigan 2019–20

Stellar Abstract Award Program in Quantitative Genomics (PQG), Harvard November 2018
 Postdoctoral Fellowship Award, Stanley Center for Psychiatric Research, Broad 2017–18

• Weston Brain Institute International Fellowship in Neuroscience December 2015 – May 2016

• "3 Minute Thesis" competition, University of Toronto finalist

April 2015

Trimate mess competition, of versity of forest manse

• McLaughlin Early Career Investigator Award October 2015

• Younger Family Foundation Award Fall 2015

• CIHR STAGE (Strategic Training for Advanced Genetic Epidemiology) trainee 2015–16

Institute of Medical Science Open Fellowship Award Fall 2014

• University of Toronto McLaughlin Centre Training Award Fall 2014

Peterborough K.M. Hunter Graduate Studentship
 Fall 2014

Institute of Medical Science Open Fellowship Award Fall 2013

School of Graduate Studies Conference Grants Fall 2012, 2013

• University of Toronto McLaughlin Centre Training Award September 2012

• Canadian Bioinformatics Workshop Registration Award September 2012

University of Toronto Fellowship – MSc
 September 2012

Publications

* Equal contribution † Co-senior author

Peer-reviewed

- 1. Z. Chen, D. Zhang, R.H. Reynolds, E. Gustavsson, K. D'Sa, S. García Ruiz, A. Fairbrother-Browne, J. Vandrovcova, J. Hardy, H. Houlden, **S.A. Gagliano Taliun**, J. Botía, M. Ryten. (2021) Human-lineage-specific genomic elements are enriched within genes implicated in neurodegenerative diseases. *Nature Communications*. [Accepted]
- 2. D. Taliun*, D.N. Harris*, M.D. Kessler*, J. Carlson*, Z.A. Szpiech*, R. Torres*, **S.A. Gagliano Taliun***, A. Corvelo*, [...] ~100 middle authors [...], C.C. Laurie, C.E. Jaquish, R.D. Hernandez, T.D. O'Connor, G.R. Abecasis. (2021) Sequencing of 53,831 diverse genomes from the NHLBI TOPMed Program. *Nature*. 590:290-299. PMID: 33568819
- 3. J. Nielsen, O. Rom, I. Surakka, S. Graham, W. Zhou, T. Roychowdhury, L. Fritsche, **S.A. Gagliano Taliun** [...] ~50 middle authors [...] M. Boehnke, S. Kathiresan, G.R. Abecasis, Y.E. Chen, C. Willer, K. Hveem. (2020) Loss-of-function genetic variants with impact on liver-related blood traits highlight potential therapeutic targets for cardiovascular disease. *Nature Communications*. 11:6417. PMID: 33339817
- 4. X. Zhao, D. Qiao, C. Yang, S. Kasela, W. Kim, N. Shrine, C. Batini, T. Sofer, **S.A. Gagliano Taliun** [...] \sim 60 middle authors [...] M. Cho, A. Manichaikul. (2020) Whole genome sequence analysis of pulmonary function and COPD in 19,996 multi-ethnic participants. *Nature Communications*. 11:5182. PMID: 33057025
- 5. Z. Chen, W. Yan Yau, Z. Jaunmuktane, A. Tucci, P. Sivakumar, **S.A. Gagliano Taliun** [...] ~20 middle authors [...] J. Hardy, M. Ryten, J. Vandrovcova, H. Houlden. (2020) Neuronal intranuclear inclusion disease is genetically heterogenous. *Annals of Clinical and Translational Neurology*. 10;7(9):1716-172. PMID: 32777174
- 6. **S.A. Gagliano Taliun***, P. VandeHaar*, A.P. Boughton, R.P. Welch, D. Taliun, E.M. Schmidt, W. Zhou, J.B. Nielsen, C.J. Willer, S. Lee, L.G. Fritsche, M. Boehnke, G.R. Abecasis (2020) Exploring and visualizing large-scale genetic associations using PheWeb. *Nature Genetics*. 52:550–552. PMID: 32504056
- 7. W. Zhou*, Z. Zhao*, J.B. Nielsen, L.G. Fritsche, J. LeFaive, **S.A. Gagliano Taliun**, W. Bi, M.J. Daly, B.M. Neale, K. Hveem, G.R. Abecasis, C.J. Willer, S. Lee. (2020) Scalable generalized linear mixed model for region-based association tests in large biobanks and cohorts. *Nature Genetics*. 52:634–639. PMID: 32424355
- 8. F. Zhang, M. Flickinger, **S.A. Gagliano Taliun**, InPSYght Psychiatric Genetics Consortium, G.R Abecasis, L.J. Scott, S.A. McCarroll, C.N. Pato, M. Boehnke, H.M. Kang (2020) Ancestry-agnostic estimation of DNA sample contamination from sequence reads. *Genome Research*. 30(2):185-194. PMID: 31980570
- S. Guelfi, K.D'Sa, J. Botía, J. Vandrovoca, R.H. Reynolds, D. Zhang, A. Ramasamy, D. Trabzuni, L. Collado-Torres, A. Thomason, P. Quijada Leyton, S.A. Gagliano Taliun, Mike A. Nalls, C. Smith, J. Hardy, M.E. Weale, K.S. Small, M. Ryten (2020) Regulatory sites for known and novel splicing in human basal ganglia are enriched for disease-relevant information. *Nature Communications*. 11:1041. PMID: 32098967
- 10. R.H. Reynolds, J. Hardy, M. Ryten[†], **S.A. Gagliano Taliun**[†] (2019) Informing disease modelling with brain-relevant functional genomic annotations. *Brain*. 0:1-19. PMID: 31603214
- D. Dutta, S.A. Gagliano Taliun, J. Weinstock, M. Zawistowski, C. Sidore, F. Cucca, D. Schlessinger, G. Abecasis, C. Brummett, S. Lee. (2019) Meta-MultiSKAT: Multiple phenotype meta-analysis for region-based association test. *Genetic Epidemiology*. 43(7):800-814. PMID: 31433078
- 12. L.G. Fritsche, L. J. Beesley, P. VandeHaar, R.B. Peng, M. Salvatore, M. Zawistowski, **S.A. Gagliano Taliun**, S. Das, J. LeFaive, E.O. Kaleba, T.T. Klumpner, S.E. Moser, V.M. Blanc, C.M. Brummett, S. Kheterpal, G. R. Abecasis, S.B. Gruber, B. Mukherjee (2019) Exploring various polygenic risk scores for skin cancer in the phenomes of the Michigan Genomics Initiative and the UK Biobank with a visual catalog: PRSWeb. *PLOS Genetics*. 15(6):e1008202. PMID: 31194742

- 13. **S.A. Gagliano Taliun** (2019) Genetic determinants of low vitamin B12 levels in Alzheimer's disease risk. *Alzheimer's & Dementia: Diagnosis, Assessment & Disease Monitoring*. 11:430-434. PMID: 31206009
- 14. R.H. Reynolds, J.A. Botía, M.A. Nalls, International Parkinson's Disease Genomics Consortium (IPDGC), J. Hardy, **S.A. Gagliano Taliun**[†], M. Ryten[†] (2019) Moving beyond neurons: the role of cell type-specific gene regulation in Parkinson's disease heritability. 5:6. *NPJ Parkinsons Disease*. PMID: 31016231
- 15. **S.A. Gagliano**, S. Sengupta, C. Sidore, A. Maschio, F. Cucca, D. Schlessinger, G.R Abecasis (2018) Relative impact of indels versus SNPs on complex disease. *Genetic Epidemiology*. 43(1):112-117. PMID: 30565766
- C.D. Hughes, M.L. Choi, M. Ryten, L. Hopkins, A. Drews, J.A. Botía, M. Iljina, M. Rodrigues, S.A. Gagliano, S. Gandhi, C. Bryant, D. Klenerman (2018) Picomolar concentrations of oligomeric alpha synuclein sensitizes TLR4 to play an initiating role in PD pathogenesis. *Acta Neuropathologica*. 137(1):103-120. PMID: 30225556
- 17. W. Zhou, J.B. Nielsen, L.G. Fritsche, R. Dey, M.B. Elvestad, B.N. Wolford, J. LeFaive, P. VandeHaar, **S.A. Gagliano**, A. Gifford, L.A. Bastarache, W-Q. Wei, J.C. Denny, M. Lin, K. Hveem, H.M. Kang, G.R. Abecasis, C.J. Willer, S. Lee (2018) Efficiently controlling for case-control imbalance and sample relatedness in large-scale genetic association studies. *Nature Genetics*. 50(9):1335-1341. PMID: 30104761
- M. Hernandez-Fuentes, C. Franklin, I. Rebollo-Mesa, J. Mollon, F. Delaney, E. Perucha, C. Stapleton, R. Borrows, C. Byrne, G. Cavalleri, B. Clarke, M. Clatworthy, J. Feehally, S. Fuggle, S.A. Gagliano, S. Griffin, A. Hammad, R. Higgins, A. Jardine, M. Keogan, T. Leach, I. MacPhee, Patrick B. Mark, J. Marsh, P. Maxwell, W. McKane, A. McLean, C. Newstead, T. Augustine, P. Phelan, S. Powis, P. Rowe, N. Sheerin, E. Solomon, H. Stephens, R. Thuraisingham, R. Trembath, P. Topham, R. Vaughan, S.H. Sacks, P. Conlon, G. Opelz, N. Soranzo, M.E. Weale, G.M. Lord, United Kingdom and Ireland Renal Transplant Consortium (UKIRTC), Wellcome Trust Case Control Consortium (WTCCC)-3 (2018) Long- and short-term outcomes in renal cadaveric allografts: a large recipient and donor genome-wide association study. *American Journal of Transplantation*. 18(6):1370-1379. PMID: 29392897
- 19. **S.A. Gagliano** (2017) It's all in the brain: a review of available functional genomic annotations. *Biological Psychiatry*. 81(6):478-483. PMID: 27788914
- 20. **S.A. Gagliano**, J. G. Pouget, J. Hardy, J. Knight, M.R. Barnes, M. Ryten, M.E. Weale (2016) Genomics implicates adaptive and innate immunity in Alzheimer's and Parkinson's. *Annals of Clinical and Translational Neurology*. 3(12):924-933. PMID: 28097204
- 21. G. Zai, [...] **S.A. Gagliano** [...], ~40 alphabetized middle authors [...], J.L. Kennedy (2016) Rapporteur Summaries of Plenary, Symposia, and Oral sessions from the XXIIIrd World Congress of Psychiatric Genetics Meeting in Toronto, Canada, October 16-20, 2015. *Psychiatric Genetics*. 26(6):229-257. PMID: 27606929
- 22. **S.A. Gagliano**, C. Ptak, D.Y.F. Mak, M. Shamsi, G. Oh, J. Knight, P.C. Boutros, A. Petronis (2016) Allele-skewed DNA modification in the brain: relevance to schizophrenia GWAS. *American Journal of Human Genetics*. 98(5):956-962. PMID: 27087318
- 23. **S.A. Gagliano**, R. Ravji, M.R. Barnes, M.E. Weale, J. Knight (2015) Smoking gun or circumstantial evidence? comparison of statistical learning methods using functional annotations for prioritizing risk variants. *Scientific Reports*. 5:13373. PMID: 26300220
- 24. **S.A. Gagliano**, A.D. Paterson, M.E. Weale, J. Knight (2015) Assessing models for genetic prediction of complex traits: a comparison of visualization and quantitative methods. *BMC Genomics*. 16:405. PMID: 25997848
- 25. C.C. Zai, V. Gonçalves, A.K. Tiwari, **S.A. Gagliano**, G. Hosang, V. de Luca, S.A. Shaikh, N. King, Q. Chen, W. Xu, J. Strauss, G. Breen, C.M. Lewis, A.E. Farmer, P. McGuffin, J. Knight, J.B. Vincent, J.L. Kennedy (2014) A genome-wide association study of suicide severity scores in bipolar disorder. *Journal of Psychiatric Research*. 65:23-9. PMID: 25917933
- 26. **S.A. Gagliano**, A. K. Tiwari, N. Freeman, J.A. Lieberman, H.Y. Meltzer, J.L. Kennedy, J. Knight, D.J. Müller (2014) Protein kinase cAMP-dependent regulatory type II beta (*PRKAR2B*) gene variants in antipsychotic-induced weight gain. *Human Psychopharmacolgy*. 29(4):330-5. PMID: 24737441

27. **S.A. Gagliano**, M.R. Barnes, M.E. Weale, J. Knight (2014) A Bayesian method to incorporate hundreds of functional characteristics with association evidence to improve variant prioritization. *PLoS ONE*. 9(5):e98122. PMID: 24844982

Under-review

- M. Riise Moksnes, H. Røsjø, A. Richmond, M. Nakrem Lyngbakken, S.E. Graham, A. Falkmo Hansen, B.N. Wolford, S.A. Gagliano Taliun, J. LeFaive, H. Rasheed, L.Thomas, W. Zhou, A. Campbell, D.J. Porteous, P. Welsh, N. Sattar, G. Davey Smith, L. Fritsche, J.B. Nielsen, B.O. Åsvold, K. Hveem, C. Hayward, C. Willer, B.M. Brumpton, T. Omland. Using human genetics to understand the causes and consequences of circulating cardiac troponin I in the general population.
- 2. T.N. Kelly*, X. Sun*, K.Y. Ye*, M.R. Brown*, **S.A. Gagliano Taliun***, J.N. Hellwge*, X. Mi*, J.A. Brody* [...] ~100 middle authors [...] T.L. Edwards, G.R. Abecasis, X. Zhu, D. Levy, D.K. Arnett, A.C. Morrison. Whole-genome sequencing study of blood pressure traits in the Trans-Omics for Precision Medicine and Centers for Common Disease Genomics programs.
- 3. K. D'Sa, R.H. Reynolds, S. Guelfi, D. Zhang, S.G. Ruiz, International Parkinson's Disease Genomics Consortium (IPDGC), System Genomics of Parkinson's Disease (SGPD), J.Hardy, **S.A. Gagliano Taliun**, K.S. Small, M. Ryten, J. Botía. ERASE: Extended Randomization for assessment of annotation enrichment in ASE datasets.

Scientific commentaries (edited, but not peer-reviewed)

- 1. **S.A. Gagliano Taliun** (2020) How to navigate academia as a female who is the first in her family with a PhD. Academic Matters. https://academicmatters.ca/how-to-navigate-academia-as-a-female-who-is-the-first-in-her-family-with-a-phd/
- 2. **S.A. Gagliano Taliun** (2019) Teaching at the university-level is not a hassle. *Nature*. 574(7777): 285. PMID: 31591538

Oral Presentations as an Invited Speaker

- Association canadienne-française pour l'avancement des sciences (ACFAS) meeting "L'imputation des génotypes permet la découverte de variations rares associées aux maladies" (Virtual meeting; May 7, 2021)
- Beyond Mendelian Genetics Symposium: Joint Genome Biology and Precision Medicine and Molecular Basis of Rare Diseases Section Symposium. "Leveraging dense genotype imputation for disease-associated rare variant discovery" (Virtual meeting; December 10, 2020)
- CAMH. "Using regulatory elements to prioritize genetic risk variants for neurodegeneration" (Toronto, Ontario; August 22, 2018)
- Michigan State University. "Prioritizing risk variants for neurodegeneration using functional genomics" (East Lansing, Michigan; June 15, 2018)
- Annual Canadian Human and Statistical Genetics meeting. "Understanding brain disorders through functional genomics" (Harrison Hot Springs, British Columbia; June 11, 2018)
- Genomics of Brain Disorders. "Neurodegeneration: Beyond the brain" (Hinxton, UK; April 24, 2018)

Oral Presentations

- Whole Genome Sequencing of Psychiatric Disorders (WGSPD) meeting. "InPSYght case-control and leveraging controls from TOPMed" (Cambridge, Massachusetts; January 24, 2019)
- Trans-omics for Precision Medicine (TOPMed) meeting "TOPMed imputed UK Biobank genetic data reveals disease-associated rare loss of function variation" (Tysons, Virginia; Dec. 5, 2018)
- Program in Quantitative Genomics (PQG) Conference. "Dense imputation of the UK Biobank genetic data reveals disease-associated rare loss of function variation" (Boston, Massachusetts; November 1, 2018)
- Whole Genome Sequencing of Psychiatric Disorders (WGSPD) meeting. "InPSYght case-control and controls from other sources" (Bethesda, Maryland; January 10, 2018)

- University of Michigan Biostatistics Grand Rounds "PheWAS of >100 traits in the SardiNIA study: Insights" (Ann Arbor, Michigan; November 2, 2017)
- Whole Genome Sequencing of Psychiatric Disorders (WGSPD) meeting. "Joint analysis of sequenced InPSYght African American schizophrenia and bipolar cases and controls with TOPMed external controls" (Cambridge, Massachusetts; June 5, 2017)
- Center for Biomedicine, European Academy of Bolzano (EURAC). "PheWAS of >100 traits in SardiNIA" (Bolzano, Italy; Nov 22, 2016)
- European Mathematical Genetics Meeting (EMGM). "In silico identification of genetic risk variants for Parkinson's disease" (Newcastle, UK; May 12, 2016)
- World Congress of Psychiatric Genetics (WCPG) 2015 annual meeting. "In silico prioritization of genetic risk variants for psychiatric disorders using functional genomic information" (Toronto, Ontario; October 20, 2015)
- Harvey Stancer Research Day 2014. "Assessing models for genetic prediction of complex traits: visualization and quantitative methods" (Toronto, Ontario; June 19, 2014)
- Statistical Society of Canada annual meeting. "Investigation of Predictive Accuracy Measures for Genetic Models" (Toronto, Ontario; May 26, 2014)
- Toronto Bioinformatics User Group. "A method to incorporate hundreds of functional characteristics with association evidence to improve SNP prioritization" (Toronto, Ontario; November 27, 2013) https://www.youtube.com/watch?v=xpDYFfAfPZE
- Harvey Stancer Research Day 2013. "Protein kinase cAMP-dependent regulatory type II beta (PRKAR2B) gene variants in antipsychotic-induced weight gain" (Toronto, Ontario; June 13, 2013)
- Neuroscience Research Exchange Day. "Distribution of epiSNPs in sub-threshold variants from genome-wide association studies for psychiatric disorders" (Toronto, Ontario; April 12, 2013)
- Genetic Analysis Workshop 18 Conference. "GAW18: Gene-based tests" (Stevenson,
 Washington; October 16, 2012); Co-presenters: Heather Cordell and Indranil Mukhopadhyay

Poster presentations as first author

- S.A. Gagliano Taliun, Y. Li, D. Ray, P. Yajnik, NIMH InPSYght Consortium and NHLBI TOPMed Program, S. Lee, L.J. Scott, S.A. McCarroll, C.N. Pato, G.R. Abecasis, M. Boehnke, H.M. Kang. "High-specificity variant filter enables joint analysis of whole genome sequence data from multiple studies and sequencing centres" American Society of Human Genetics (ASHG) Meeting. (Houston, Texas; October 17, 2019)
- **S.A. Gagliano**, W. Zhou, J.B. Nielson, J. LeFaive, S. Das, D. Taliun, R. Dey, G.R. Abecasis. "Analysis of densely imputed UK Biobank genetic data reveals disease-associated rare loss of function variation" American Society of Human Genetics (ASHG) Meeting. (San Diego, California; October 17, 2018) *Top 10% of submissions*.
- **S.A. Gagliano**, W. Zhou, J.B. Nielson, J. LeFaive, S. Das, D.Taliun, R. Dey, G.R. Abecasis. "Analysis of densely imputed UK Biobank genetic data reveals disease-associated rare loss of function variation previously only implicated in family studies" UK Biobank Conference Early-Career Researcher of the Year (London, UK; June 21, 2018) *Top 20% of submissions*.
- S.A. Gagliano, S. Sengupta, C. Sidore, A. Maschio, F. Cucca, D. Schlessinger, G.R. Abecasis.
 "Pinpointing GWAS signals: Indels vs. SNPs" American Society of Human Genetics (ASHG) Meeting. (Orlando, Florida; October 19, 2017)
- S.A. Gagliano, J.G. Pouget, J. Hardy, J. Knight, M.R. Barnes, M. Ryten, M.E. Weale. "Genetic variability in both the adaptive and innate immune systems contribute to Alzheimer's and Parkinson's disease risk" International Genetic Epidemiology Society (IGES) Meeting (Toronto, Ontario; October 25, 2016)
- S.A. Gagliano, C. Ptak, D.Y.F. Mak, M. Shamsi, G. Oh, J. Knight, P.C. Boutros, A. Petronis. "Allelespecific DNA modification: relevance to GWAS of complex traits" World Congress of Psychiatric Genetics (WCPG). (Toronto, Ontario; October 17, 2015)
- S.A. Gagliano, A.D. Paterson, M.E. Weale, J. Knight. "Assessing models for genetic prediction of

- complex traits: a comparison of visualization and quantitative methods" American Society of Human Genetics (ASHG) Meeting. (Baltimore, Maryland; October 7, 2015)
- S.A. Gagliano, R. Ravji, M.R. Barnes, M.E. Weale, J. Knight. "Comparing statistical learning methods for genetic variant prioritization" International Genetic Epidemiology Society (IGES) Meeting. (Baltimore, Maryland; October 4, 2015)
- **S.A. Gagliano**, R. Ravji, M.R. Barnes, M.E. Weale, J. Knight. "Comparison of machine-learning methodologies to prioritize genetic variants based on functional data" American Society of Human Genetics (ASHG) Meeting (San Diego, California; October 21, 2014)
- S.A. Gagliano, M.R. Barnes, M.E. Weale, J. Knight. "A Bayesian method to incorporate hundreds
 of functional characteristics with association evidence to improve variant prioritization" Institute
 of Medical Science Scientific Day (Toronto, Ontario; May 22, 2014)
- S.A. Gagliano, M.R. Barnes, M.E. Weale, J. Knight. "Enrichment of functional information (543 annotation tracks) in GWAS hits" American Society of Human Genetics (ASHG) Meeting (Boston, Massachusetts; October 24, 2013)
- S.A. Gagliano, D.Y.F. Mak, C. Ptak, P.C. Boutros, A. Petronis, J. Knight. "Distribution of epiSNPs in Sub-threshold Variants from Genome-wide Association Studies for Psychiatric Disorders" World Congress of Psychiatric Genetics (Boston, Massachusetts; October 18, 2013)
- S.A. Gagliano, A.K. Tiwari, N. Freeman, J.A. Lieberman, H.Y. Meltzer, D.J. Mueller, J.L. Kennedy, J. Knight "Protein kinase cAMP-dependent regulatory type II beta (PRKAR2B) gene variants in antipsychotic-induced weight gain" Canadian College of Neuropsychopharmacology annual meeting (Toronto, Ontario; May 30, 2013)
- S.A. Gagliano, D.Y.F. Mak, C. Ptak, P.C. Boutros, A. Petronis, J. Knight. "Distribution of epiSNPs in sub-threshold variants from genome-wide association studies for psychiatric disorders" Institute of Medical Science Scientific Day (Toronto, Ontario; May 28, 2013)
- **S.A. Gagliano**, K. Benke, J. Knight. Genetic Analysis Workshop 18 Conference, "Functional Annotation of Rare Variants in GAW18 Data" (Stevenson, Washington; October 16, 2012)

Teaching and Mentoring

Graduate level

- Guest lecturer, MSO 6018: Introduction to Genetic Epidemiology Université de Montréal
- Winter 2021
- Prepared and taught a 3h class on Mendelian Randomization for this graduate-level course of 12 students.
- **Co-instructor**, Human Genetics 542: Molecular Basis of Human Genetic Winter 2019, 2020 Disease, University of Michigan
 - Prepared and delivered four classes on complex traits and gene-environment interactions for this graduate-level course of ~30 students. Created and graded assignments and exam.
- **Guest lecturer**, Epidemiology 516: Genomics in Epidemiology, University Spring 2018, 2019 of Michigan
 - Co-prepared and co-taught a lecture and a hands-on practical on Biobanks and Electronic Health Records in Genetics for this graduate-level course of ~24 students.
- **Co-instructor**, Human Genetics 544: Basic Concepts in Population and Statistical Genetics, University of Michigan
 Prepared and led four 1.5h white-board-format lectures and two in-class discussions for ~20
- graduate students. Met regularly with the co-instructors for planning and grading.

 Practical Sessions Assistant at a five-day Advanced GWAS Course

 June 2013

 Tested tutorials provided by the lecturers, and aided students with computer issues.
- **Co-leader hands-on tutorial series** in statistical genetics at CAMH 2013-15 Topics included methods for genome-wide association and sequencing studies.

Undergraduate level

• Faculty mentor, Genomics, Big Data Summer Institute, University of Michigan Summer 2019
Prepared Mendelian randomization (MR) practical for ~15 students and supervised four of the

students on a project using MR on real data from large-scale cohorts

 Guest lecturer, Biostatistics 666: Statistical Methods in Human Genetics, University of Michigan

Prepared and gave Functional Genomics lecture for this graduate-level course of ~30 students.

• Mentor for high school summer student

Supervised the student through the completion of a genome-wide association study

Secondary school level

- Mentor for students Winter 2015, Fall 2016, Winter 2017, Fall 2017, Winter 2019, Fall 2019 for the Ontario On-Line Research Co-op program

 Created/graded assignments, met regularly with student/group of students, assisted with writing a final research paper, and provided feedback to the co-op teacher.
- **DNA Day Ambassador**, Skyline High School, Ann Arbor, Michigan April 2018 Co-led a lesson on genetics and inheritance for Grade 9 students.

Co-mentoring of undergraduate/graduate students

- Honorary Senior Research Associate, UCL Institute of Neurology, Neurodegenerative Diseases
 - Affiliation for my role as mentor for Regina H. Reynolds (supervisor: Mina Ryten)
- **Teaching Assistant**, HMB265H- General and Human Genetics, University of *May June 2015* Toronto
 - Prepared and led weekly tutorials, graded, helped with exam invigilation.
- **Co-mentor** (with one of my PhD supervisors) for an undergraduate research student from Vellore Institute of Technology, India
- Co-mentor (with one of my PhD supervisors) for an undergraduate exchange student from University College London, UK
 Supervised methods comparison of statistical learning models for genetic risk prediction.

Mentorship

Bioinformatics Masters Research Internship Supervisor, Université de Montréal

Oumaima Hamza
 Hester Faou
 May 2021 – Oct 2021
 Jan 2021 – June 2021

<u>Doctoral Supervisor, Université de Montréal</u>

Qiang Ye
 Jan 2021 –

Graduate Student Mentor ("Parrain"), Université de Montréal

• Justin Pelletier (Masters) 2020 – 21

<u>Thesis Committee Member, Université de Mo</u>ntréal

• Alex Richard-St-Hilaire (Masters) 2021

Marc-André Legault (Doctorate)
 2021

Certifications

Training for Diversity & Inclusive Teaching (University of Michigan)
 THE500H: Teaching in Higher Education (University of Toronto)
 Scientific Computing & High Performance Computing (SciNet)
 Advanced University Teaching Preparation (University of Toronto)
 May 2014

Professional Service

Co-Moderator for ASHG platform session "A Deep Dive into Deep Learning"
 ASHG abstract reviewer, Statistical Genetics and Genetic Epidemiology
 Co-Moderator for ASHG platform session "Biases of Polygenic Risk Scores"
 Writer, ASHG newsletter, The Nascent Transcript
 Proposer/Moderator for ASHG invited session
 October 20, 2017
 October 20, 2017

"Using controls from external studies: issues, methods & successes"

IGES Webmaster (Communications Committee)
 July 2016 – present

Post announcements to the website, updated to new website

IGES Young Investigators Committee member
 Co-organized Young Investigators Mixer at the 2016 Meeting in Toronto

2015-18

• ASHG Genetics Education Outreach Network member

January 2015 – present

 Poster co-judge at the Summer Undergraduate Research Day for the Institute of Medical Science, University of Toronto August 2014

• DNA Day Essay Contest Judge for ASHG

2014-17

• **Reviewer** for Neurobiology of Aging, Annals of Oncology, Annals of Medicine, BMC Psychiatry, Current Drug Targets, Frontiers in Immunology, Genomics, International Journal of Psychiatry in Clinical Practice, Molecular Biology Reports, PeerJ, PLoS Genetics, PLoS ONE, Psychiatric Genetics, Psychiatry Research, Scientific Reports