Curriculum Vitae

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Contact

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- Address: Department of Biostatistics, Johns Hopkins Bloomberg School of Public Health, 615 N. Wolfe St. E3527, Baltimore, MD 21202, USA

Education and research experience

- 2016-present: **Postdoctoral Fellow** Department of Biostatistics, Johns Hopkins Bloomberg School of Public Health, Baltimore, USA.
 - Advisor: Assistant Professor Kasper Hansen
- 2011-2015: PhD (Statistics) Department of Mathematics and Statistics, The University of Melbourne, Australia.
 - Thesis: The statistical analysis of high-throughput assays for studying DNA methylation
 - Advisors: Professor Terry Speed and Professor Peter Hall
- 2009-2015: Research Assistant Bioinformatics Division, Walter and Eliza Hall Institute, Melbourne, Australia.
 - Advisor: Professor Melanie Bahlo
- 2008-2009: Undergraduate Research Opportunities Program scholar Bioinformatics Division, Walter and Eliza Hall Institute, Melbourne, Australia.
 - Advisor: Professor Melanie Bahlo
- 2006-2009: Bachelor of Science (Mathematics and Statistics) with First Class Honours The University of Melbourne, Australia.
 - Thesis: X chromosome association testing in genome-wide association studies
 - Advisors: Professor Melanie Bahlo and Professor Richard Huggins

Publications

*Indicates equal contributions

Journal articles

- [1] J. T. Hickey, P. F. Hickey, N. Maniar, R. G. Timmins, M. D. Williams, C. A. Pitcher and D. A. Opar. "A novel apparatus measuring knee flexor strength during various hamstring exercises: A reliability and retrospective study". In: *Journal of Orthopaedic & Sports Physical Therapy* in press ().
- [2] P. F. Hickey. "Representation and manipulation of genomic tuples in R". In: *The Journal of Open Source Software* 1.1 (May. 2016). DOI: 10.21105/joss.00020. URL: https://doi.org/10.21105/joss.00020.

- [3] A. Keniry, L. J. Gearing, N. Jansz, J. Liu, A. Z. Holik, **P. F. Hickey**, S. A. Kinkel, D. L. Moore, K. Breslin, K. Chen, R. Liu, C. Phillips, M. Pakusch, C. Biben, J. M. Sheridan, B. T. Kile, C. Carmichael, M. E. Ritchie, D. J. Hilton and M. E. Blewitt. "Setdb1-mediated H3K9 methylation is enriched on the inactive X and plays a role in its epigenetic silencing". In: *Epigenetics & Chromatin* 9 (May. 18, 2016), p. 16. ISSN: 1756-8935. DOI: 10.1186/s13072-016-0064-6. URL: http://dx.doi.org/10.1186/s13072-016-0064-6.
- [4] D. G. Phelan, D. J. Anderson, S. E. Howden, R. C. B. Wong, **P. F. Hickey**, K. Pope, G. R. Wilson, A. Pébay, A. M. Davis, S. Petrou, A. G. Elefanty, E. G. Stanley, P. A. James, I. Macciocca, M. Bahlo, M. M. Cheung, D. J. Amor, D. A. Elliott and P. J. Lockhart. "ALPK3-deficient cardiomyocytes generated from patient-derived induced pluripotent stem cells and mutant human embryonic stem cells display abnormal calcium handling and establish that ALPK3 deficiency underlies familial cardiomyopathy". In: *European Heart Journal* 37.33 (Sep. 01, 2016), pp. 2586-2590. ISSN: 0195-668X, 1522-9645. DOI: 10.1093/eurheartj/ehw160. URL: http://dx.doi.org/10.1093/eurheartj/ehw160.
- [5] D. Lacey, **P. F. Hickey**, B. D. Arhatari, L. A. O'Reilly, L. Rohrbeck, H. Kiriazis, X. Du and P. Bouillet. "Spontaneous retrotransposon insertion into TNF 3'UTR causes heart valve disease and chronic polyarthritis". In: *Proceedings of the National Academy of Sciences of the United States of America* 112.31 (Aug. 04, 2015), pp. 9698-9703. ISSN: 0027-8424, 1091-6490. DOI: 10.1073/pnas.1508399112. URL: http://dx.doi.org/10.1073/pnas.1508399112.
- [6] H. Oey, L. Isbel, **P. F. Hickey**, B. Ebaid and E. Whitelaw. "Genetic and epigenetic variation among inbred mouse littermates: identification of inter-individual differentially methylated regions". In: *Epigenetics & Chromatin* 8 (Dec. 12, 2015), p. 54. ISSN: 1756-8935. DOI: 10.1186/s13072-015-0047-z. URL: http://dx.doi.org/10.1186/s13072-015-0047-z.
- [7] P. F. Hickey and M. Bahlo. "X chromosome association testing in genome wide association studies". In: Genetic Epidemiology 35.7 (Nov. 2011), pp. 664-670. ISSN: 0741-0395, 1098-2272. DOI: 10.1002/gepi.20616. URL: http://dx.doi.org/10.1002/gepi.20616.
- [8] M. Bahlo, J. Stankovich, P. Danoy, **P. F. Hickey**, B. V. Taylor, S. R. Browning, Australian, ew Zealand Multiple Sclerosis Genetics Consortium (ANZgene), M. A. Brown and J. P. Rubio. "Saliva-derived DNA performs well in large-scale, high-density single-nucleotide polymorphism microarray studies". In: *Cancer Epidemiology, Biomarkers & Prevention* 19.3 (Mar. 2010), pp. 794-798. ISSN: 1055-9965, 1538-7755. DOI: 10.1158/1055-9965.EPI-09-0812. URL: http://dx.doi.org/10.1158/1055-9965.EPI-09-0812.
- [9] L. G. Riley, S. Cooper, **P. F. Hickey**, J. Rudinger-Thirion, M. McKenzie, A. Compton, S. C. Lim, D. Thorburn, M. T. Ryan, R. Giegé, M. Bahlo and J. Christodoulou. "Mutation of the mitochondrial tyrosyl-tRNA synthetase gene, YARS2, causes myopathy, lactic acidosis, and sideroblastic anemia-MLASA syndrome". In: *American Journal of Human Genetics* 87.1 (Jul. 09, 2010), pp. 52-59. ISSN: 0002-9297, 1537-6605. DOI: 10.1016/j.ajhg.2010.06.001. URL: http://dx.doi.org/10.1016/j.ajhg.2010.06.001.

Preprints

[1] L. F. Rizzardi*, **P. F. Hickey***, V. Rodriguez DiBlasi, R. Tryggvadóttir, C. M. Callahan, A. Idrizi, K. D. Hansen and A. P. Feinberg. "Neuronal brain region-specific DNA methylation and chromatin accessibility are associated with neuropsychiatric disease heritability". Mar. 24, 2017. DOI: 10.1101/120386. URL: http://biorxiv.org/content/early/2017/03/24/120386.

Theses

- [1] **P. F. Hickey**. "The statistical analysis of high-throughput assays for studying DNA methylation". PhD thesis. Department of Mathematics and Statistics, The University of Melbourne, 2015. URL: https://minerva-access.unimelb.edu.au/handle/11343/55699.
- [2] P. F. Hickey. "X chromosome association testing in genome-wide association studies". Honours Thesis. Department of Mathematics and Statistics, The University of Melbourne, Nov. 05, 2009.

Technical reports, conference papers and published abstracts

- [1] I. Scheffer, K. Williams, C. Green, K. Pereira, N. Brown, **P. F. Hickey**, V. Lukic, G. Gillies, M. Delatycki, P. J. Lockhart, M. Bahlo and S. Wilson. "The Victorian Collaborative Autism Study: A family and community study of the genetics of autism spectrum disorder". In: *Journal of Intellectual Disability Research*. Ed. by J. of Intellectual Disability Research. Vol. 60. 7. 2016, p. 732. DOI: 10.1111/jir.12305.
- [2] D. G. Phelan, G. R. Wilson, K. Pope, G. Gillies, J. Sim, M. Bahlo, **P. F. Hickey**, C. Bromhead, P. A. James, D. du Sart, M. Delatyki, R. Leventer, D. J. Amor and P. J. Lockhart. "Identification and characterisation of a novel hypertrophic cardiomyopathy gene". In: *Pathology: The Journal of the Royal College of Pathologists of Australasia*. Ed. by P. T. J. of the Royal College. Vol. 46, 2014, pp. S91-S92.
- [3] D. G. Phelan, G. R. Wilson, J. Sim, M. Bahlo, **P. F. Hickey**, P. A. James, D. du Sart, M. Delatyki, D. J. Amor and P. J. Lockhart. "Identification and characterisation of a novel hypertrophic cardiomyopathy gene". In: *Global Heart*. Ed. by G. Heart. Vol. 9. 1. Elsevier, 2014, p. e316. DOI: http://dx.doi.org/10.1016/j.gheart.2014.03.2365.

Commentaries and meeting reports

[1] **P. F. Hickey** and M. D. Robinson. *Genomics by the beach*. Apr. 14, 2014. DOI: 10.1186/gb4171. URL: http://dx.doi.org/10.1186/gb4171.

Computer skills

Very proficient with statistical computing, particularly R, and standard tools for analysing genomics data such as Bioconductor, SAMtools, and BEDtools. Familiar with C++, C, Python, Unix shell tools, Sun Grid Engine, Slurm. Developed pipelines for analysing genomics data including whole-genome and whole-exome sequencing, ATAC-seq, RNA-seq, and whole-genome bisulfite-sequencing.

Software packages

Developer and contributor to several R packages through the Bioconductor project. Also developed and published Python software. Please see https://github.com/PeteHaitch for additional projects.

R packages

Download statistics from http://bioconductor.org/packages/stats/bioc for years since I have been a substantial author/contributor to the package.

Author

- Genomic Tuples: Representation and Manipulation of Genomic Tuples
 - Number of downloads (2014-2017): 497, 2426, 2022, 870
- bsseq: Analyze, Manage and Store Bisulfite Sequencing Data
 - Number of downloads (2014-2017): 5049, 6631, 8419, 5069

Contributor

- minfi: Analysing Illumina Infinium DNA Methylation Arrays
 - Number of downloads (2017): 18539

Python packages

Author

• methtuple: A caller for DNA methylation events that co-occur on the same DNA fragment from high-throughput bisulfite sequencing data, such as methylC-seq

Funding

Grants

- 2015: Bioconductor Travel Grant
 - To attend the Bioconductor meeting in Seattle, USA
- 2013: EMBL Australia Travel Grant
 - To attend the EMBL PhD Symposium in Heidelberg, Germany

Scholarships

- 2013: Victorian Life Sciences Computation Initiative PhD Top-Up Scholarship.
- 2012: Statistical Society of Australia (Victoria Branch) scholarship to attend the Young Statisticians Conference (2013).
- 2011-2015: Australian Postgraduate Award.
- 2009: Maurice Belz scholarship, The University of Melbourne
 - A competitive scholarship awarded to complete Honours degree in statistics, stochastic processes or operations research.
- 2009: Alan W Harris Honours scholarship, The Walter and Eliza Hall Institute of Medical Research

Awards

- 2015: Edith Moffat Travel Award
- To interview for international for postdoctoral positions and attend the European Bioconductor meeting
- 2013: Prize for best lightning talk, Australian Epigenetics Conference 2013.
- 2013: Prize for third best oral presentation, Young Statisticians Conference 2013.
- 2010: Best presentation (Statistics), 2010 Victorian Mathematics and Students' Conference.

Presentations

Talks

Conferences

• 2017: Developing statistical methods for large epigenomic studies in the human brain. Contributed talk, ENAR 2017 Spring Meeting, Washington D.C., USA (03/13)

- 2016: New features in bsseq for analysing large whole genome bisulfite-sequencing datasets. Lightning talk, BioC 2016, San Franciso, USA (24/06)
- 2016: GenomicTuples. Lightning talk, BioC 2015 Developers' Day, Seattle, USA (20/7)
- 2015: Genomic tuples and DNA methylation patterns. Contributed talk, European Bioconductor Developers' Meeting, Heidelberg, Germany (12/01) [slides]
- 2014: Making sense of DNA methylation data. PhD completion seminar, Melbourne, Australia (15/09) [video | pdf]
- 2014: Simulating whole-genome DNA methylation data. Contributed talk, Australian Statistical Conference/International Mathematical Statistics Annual Meeting, Sydney, Australia (10/07) [slides]
- 2013: Exploiting local dependencies in genome-wide studies of DNA methylation. Contributed talk, Young Statisticians Conference, Melbourne, Australia (07/02)
- 2012: Spatial dependence of CpG-methylation from whole genome bisulfite sequencing. Contributed talk, Epigenomics of Common Diseases Meeting, Baltimore, USA (15/10)
- 2012: Spatial dependence of DNA methylation. Contributed talk, Australian Statistical Conference, Adelaide, Australia (15/07).
- 2010: Bioinformatics Applied statistics in modern molecular biology. Contributed talk (with Davis McCarthy), 2010 Victorian Mathematics and Statistics Students' Conference, Melbourne, Australia (02/07).

Other

- 2013: Bioinformatics for bisulfite sequencing. Invited talk, La Trobe University Sequencing Users Group, Melbourne, Australia (28/08)
- 2010: X chromosome association testing in genome wide association studies. Invited talk, Statistical Society of Australia Victorian branch meeting, Melbourne, Australia (24/08).

Posters

- 2014: Simulating whole-genome bisulfite-sequencing data. Poster, Lorne Genome, Lorne, Australia (17/01) [pdf]
- 2013: Simulating whole-genome bisulfite-sequencing data. Poster, Epigenetics 2013, Shoal Bay, Australia (03/12) [pdf]
- 2011: Analysis of mouse exome sequencing: filtering institute specific single nucleotide variants (SNVs). Poster, GeneMappers - 8th Australian Human Gene Mapping Conference, Hobart, Australia (04/04).
- 2010: X chromosome association testing in genome wide association studies. Poster, The International Genetic Epidemiology Society Conference, Boston, USA (12/10) 2010: X chromosome association testing in genome wide association studies. Poster, The Australasian Microarray and Associated Technologies Association Conference, Hobart, Australia (16/09) 2009: Homozygosity by state analysis in highly inbred pedigrees. Poster, GeneMappers 7th Australian Human Gene Mapping Conference, Sydney, Australia (15/06)

Teaching experience

Workshops

• 2016: Analysing DNA methylation data with Bioconductor. BioC 2016, Palo Alto, USA (26/06)

Tutoring

- 2006-2014: Private tutoring
 - One-on-one and small-group tutoring for students studying Masters-level biostatistics, third-year
 university level statistics and first-year university level engineering mathematics. Tutoring senior
 high school students in elementary, intermediate and advanced mathematics subjects and physics.
- 2011: Classroom assistant
 - Assisting in mathematics classes for final year high school students

Professional activities and service

Reviewing

Referee for Nature Methods, PLoS Genetics, Genome Biology, Bioinformatics, PLoS Computational Biology, Genetic Epidemiology, Heredity, and F1000.

Professional society memberships

- Member, Statistical Society of Australia
- Member, International Biometric Society (ENAR)

References

Professor Terry Speed

Bioinformatics Division

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