Thomas Harrop

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I am a genomics scientist working with large-scale bioinformatic and genetic datasets.

Technical skills

I'm an experienced bioinformatician, geneticist and molecular biologist.

My main tools for programming and data analysis are Python and R. I use workflow managers, version control, containerisation and high-performance computing for bioinformatics. I'm also experienced with bash scripting and *nix system administration.

I am currently using my strategic development and user support skills in the context of a team working to provide national bioinformatics infrustructure, in collaboration with a distributed global team.

I am skilled at the entire wet-lab workflow for producing short and long read sequencing libraries, from nucleic acid extraction to library preparation and running instruments for 2^{nd} - and 3^{rd} -generation sequencing. I have experience with classical genetic approaches with live organisms, including transformation, crosses, microdissection and bioassays.

I enjoy communicating complex bioinformatics and genetic results to research partners, external stakeholders and students. I visualise and share my results with ggplot2, Markdown, Pandoc, TeX and a bit of html & css. I teach complex bioinformatics algorithms to postgraduate students in classroom and one-on-one settings.

Research

I'm interested in research related to agriculture and sustainable development, tools and techniques in bioinformatics and evolutionary and comparative biology.

Experience

Current positions

2023–Present Scientific Officer, CSIRO, Clayton, Australia.

I contribute to development of a genomics platform for natural resource management. This involves experiment design, method development and data analysis to address challenges in conservation genomics, species identification,

and biodiscovery in plants and insects.

2021-Present Academic Specialist, Melbourne Bioinformatics, University

of Melbourne, Australia, and Bioinformatician, Galaxy

Australia.

I provide training and support to researchers and projects who use bioinformatics in Melbourne's biosciences precinct. I am part of the team that develops and runs Galaxy Australia and global Galaxy infrastructure.

Previous research and University positions

2020–2021 Research Fellow, Biochemistry Department, University of

Otago, Dunedin, New Zealand.

I began developing an independent genomics and bioinformatics research program related to genomic selection and pest management problems in New Zealand.

2017–2020 Post-doctoral researcher, Laboratory for Evolution and

Development, Biochemistry Department, University of

Otago, Dunedin, New Zealand.

In this postdoc position I worked on a project to develop and apply tools to integrate genomic selection to breeding

programmes to improve honeybee performance.

2016–2017 Post-doctoral researcher, Laboratory for Evolution and

Development, Biochemistry Department, University of

Otago, Dunedin, New Zealand.

I worked on a bioprotection project investigating the

resistance of Argentine stem weevil to its biological control

agent.

2014–2016 Post-doctoral researcher, UMR DIADE, Institut de

Recherche pour le Développement (IRD), Montpellier,

France.

We used RNA sequencing to characterise molecular events that occur during development of inflorescence architecture in rice, which affects crop yield, and applied these results to study the effects of selection on gene expression during the parallel domestications of African and Asian rice.

2013–2014 Post-doctoral researcher, Department of Plant Breeding

and Genetics, Max Planck Institute for Plant Breeding

Research, Cologne, Germany.

At the Max Planck Institute I worked on a bioinformatic and molecular biology project to compare the transcriptional response of different land plants environmental signals.

Professional positions

2004–2007 Medical scientist, Network Pathology, the Austin Hospital,

Heidelberg, Australia.

2001–2004 Laboratory assistant, Network Pathology, the Austin

Hospital, Heidelberg, Australia.

Education

2006–2012 Doctor of Philosophy, Department of Genetics, the

University of Melbourne, Parkville, Australia.

Thesis: <u>The functions of cytochrome P450s in Drosophila</u>. I used classical genetics and modern molecular biology, including transgenic manipulation of gene expression and comparative studies of gene function, to investigate the functions of cytochrome P450 genes in the Drosophila

genus.

2002–2005 Bachelor of Science with Honours (first class), the

University of Melbourne, Parkville, Australia.

Awards 2006–2009, Australian Postgraduate Award

Personal information

Languages English (native)

German (<u>TELC Level</u> B2) French (<u>TELC Level</u> B1)

Nationalities Australian and British

Teaching

2021-current	BINF90002, Elements of Bioinformatics, guest lecturer.
2021-2023	COMP90014, Algorithms for Bioinformatics , Masters of Bioinformatics, The University of Melbourne. Subject coordinator and lecturer.
	BINF90002, Elements of Bioinformatics & COMP90016, Computational Genomics. Guest lecturer.
2020	GENE412 , Genetics Department, The University of Otago.
2017-2019	GENE312 project supervisor, Genetics Department, The University of Otago.
2007-2011	3rd Year Genetics , Department of Genetics, The University of Melbourne, Parkville, Australia.
2006-2009	High School Biology , Department of Genetics, The University of Melbourne, Parkville, Australia.
2006-2008	High School Biology , Gene Technology Access Centre, The University High School, Parkville, Australia.
2005-2008	1st Year Genetics , The University of Melbourne, Parkville, Australia.

Students (co-)supervised

Four graduate research students, Masters of Science (Bioinformatics), University of Melbourne.
Bachelor of Science with Honours , Genetics, University of Otago, Dunedin, New Zealand.
Bachelor of Science with Honours , Genetics, University of Otago, Dunedin, New Zealand.
Master of Science , Life Science Informatics, Universität Bonn, Germany.
Master of Science , Sciences et Technologies de l'Information et de la Communication pour la Santé, Université de Montpellier, France.

Service

Community group Founder and convenor of the popular *Bioinformatics Meetups*

group for people using bioinformatics in the local research

community, hosted by Melbourne Bioinformatics.

Genome Annotation Workshop development and lead trainer, Melbourne

Bioinformatics, 12 March 2024.

NextFlow Trainer, CSIRO pipeline development workshop, 23-24

August 2023.

Galaxy development Tool Development and Galaxy + R training workshops,

Galaxy Community Conference, 10–16 July 2023.

From reads to differentially expressed genes workshop for RNA-seq

Australian BioCommons, 23-24 February 2021.

Lead trainer, Melbourne Bioinformatics, 2021-2023.

Reproducible

Reproducible research workflows with containers webinar research

for New Zealand eScience Infrastructure (NeSI), 23 July

2020.

Sequencing Invited instructor for the Long-read, long-reach sequencing

workshop workshop. November 2019, ANU, Canberra, Australia.

MapNet Main organiser for the 2018 MapNet conference.

November 2018, Dunedin, New Zealand.

Main organiser for Genomics Aotearoa's long-read **Nanopore**

workshop sequencing workshop. April 2018, Dunedin, New Zealand.

Carpentries Assisting instructor for Carpentries workshops: Data

Carpentry, 2019; RNA-seq-workshop, 2020.

Publications

See Google scholar for the current list.

Journal articles

- Harrop, T.W.R.[§], Guhlin, J.[§], McLaughlin, G.M., Permina, E., Stockwell, P., Gilligan, J., Le Lec, M.F., Gruber, M.A.M., Quinn, O., Lovegrove, M., Duncan, E.J., Remnant E.J., Langford, K.L., Kronenberg, Z., Press, M.O., Eacker, S.M., Wilson-Rankin, E.E., Purcell, J., Lester, P.J., Dearden, P.K. (2020). High-quality assemblies for three invasive social wasps from the *Vespula* genus. G3: Genes, Genomes, Genetics 10 (10): 3479–3488. doi:10.1534/g3.120.401579.
- Harrop, T.W.R., Le Lec, M.F., Jauregui, R., Taylor, S.E., Inwood, S.N., van Stijn, T., Henry, H., Skelly, J., Ganesh, S., Ashby, R.L., Jacobs, J.M.E., Goldson, S.L., Dearden, P.K. (2020). Genetic Diversity in Invasive Populations of Argentine Stem Weevil Associated with Adaptation to Biocontrol. Insects 11 (7): 441. doi:10.3390/insects11070441.
- **Harrop, T.W.R.**, Mantegazza, O., Luong, A.M., Béthune, K., Lorieux, M., Jouannic, S., and Adam, H. (2019). A set of *AP2*-like genes is associated with inflorescence branching and architecture in domesticated rice. Journal of Experimental Botany 70, 5617–5629. doi:10.1093/jxb/erz340.
- Harrop, T.W.R.[§], Denecke, S.[§], Yang, Y.T., Chan, J., Daborn, P.J., Perry, T., Batterham, P., (2018). Evidence for activation of nitenpyram by a mitochondrial cytochrome P450 in *Drosophila melanogaster*. Pest Management Science 74, 1616–1622. doi:10.1002/ps.4852.
- Harrop, T.W.R.[§], Ud Din, I.[§], Gregis, V., Osnato, M., Jouannic, S., Adam, H. and Kater, M. (2016). Gene expression profiling of reproductive meristem types in early rice inflorescences by laser microdissection. The Plant Journal 86, 75–88. doi:10.1111/tpj.13147.
- Harrop, T.W.R.[§], Pearce, S.L.[§], Daborn, P.J. and Batterham, P. (2014). Whole-genome expression analysis in the third instar larval midgut of *Drosophila melanogaster*. G3 (Bethesda) 4, 2197–2205. doi:10.1534/g3.114.013870.
- **Harrop, T.W.R.**, Sztal, T., Lumb, C., Good, R.T., Daborn, P.J., Batterham, P. and Chung, H. (2014). Evolutionary changes in gene expression, coding sequence and copy-number at the *Cyp6g1* locus contribute to resistance to multiple insecticides in Drosophila. Plos One 9, e84879. doi:10.1371/journal.pone.0084879.

[§] Equal contribution

Contributions to articles

- Inwood, S.N., Skelly, J., Guhlin, J.G., **Harrop, T.W.R.**, Goldson, S.L., and Dearden, P.K. (2023). Chromosome-level genome assemblies of two parasitoid biocontrol wasps reveal the parthenogenesis mechanism and an associated novel virus. BMC Genomics 24, 440 doi:10.1186/s12864-023-09538-4
- Inwood, S.N., **Harrop, T.W.R.**, and Dearden, P.K. (2023). The venom composition and parthenogenesis mechanism of the parasitoid wasp *Microctonus hyperodae*, a declining biocontrol agent. Insect Biochemistry and Molecular Biology 153. doi:10.1016/j.ibmb.2022.103897.
- McCulloch, G.A., Guhlin, J., Dutoit, L., **Harrop, T.W.R.**, Dearden P.K., and Waters, J.M. (2021). Genomic Signatures of Parallel Alpine Adaptation in Recently Evolved Flightless Insects. Molecular Ecology 30 (24): 6677–86. doi:10.1111/mec.16204.
- Dussex, N. *et al.* (2021). Population Genomics of the Critically Endangered Kākāpō. Cell Genomics 1 (1): 100002. doi:10.1016/j.xgen.2021.100002.
- McCulloch, G.A., Foster, B.J., Dutoit, L., **Harrop, T.W.R.**, Guhlin, J., Dearden, P.K., Waters, J.M. (2020). Genomics Reveals Widespread Ecological Speciation in Flightless Insects. Systematic Biology syaa094. doi:10.1093/sysbio/syaa094.
- Lamichhane, R., Munro, F., **Harrop, T.W.R.**, de la Harpe, S.M., Dearden, P.K., Vernall, A.J., McCall, J.L., Ussher, J.E. (2020). Human liver-derived MAIT cells differ from blood MAIT cells in their metabolism and response to TCR-independent activation. European Journal of Immunology 51: 879–892. doi:10.1002/eji.202048830.
- Morgan, B.M., Brown, A.N., Deo, N., **Harrop, T.W.R.**, Taiaroa, G., Mace, P.D., Wilbanks, S.M., Merriman, T.R., Williams, M.J.A., McCormick, S.P.A. (2020). Nonsynonymous SNPs in *LPA* homologous to plasminogen deficiency mutants represent novel null apo(a) alleles. Journal of Lipid Research 61(3), 432–44. doi:10.1194/jlr.M094540.
- Lamichhane, R., Schneider, M., de la Harpe, S.M., Harrop, T.W.R., Hannaway, R.F., Dearden, P.K., Kirman, J.R., Tyndall, J.D.A., Vernall, A.J., and Ussher, J.E. (2019). TCR- or Cytokine-Activated CD8+ Mucosal-Associated Invariant T Cells Are Rapid Polyfunctional Effectors That Can Coordinate Immune Responses. Cell Reports 28, 3061-3076.e5. doi:10.1016/j.celrep.2019.08.054.
- Guerrero, F.D., Bendele, K.G., Ghaffarim N., Guhlin, J., Gedye, K.R., Lawrence, K.E., Dearden, **Harrop, T.W.R.**, Heath, A.C.G, Lun, Y., Metz, R.P, Teel, P., Perez de Leon, A., Biggs, P.J., Pomroy, W.E., Johnson, C.D., Blood, P.D., Bellgard, S.E., Tompkins, D.M. (2019). The Pacific Biosciences *de novo* assembled genome dataset from a parthenogenetic New Zealand wild population of the longhorned tick, *Haemaphysalis longicornis* Neumann, 1901. Data in Brief 27, 104602. doi:10.1016/j.dib.2019.104602.

- Ta, K. N., Adam. H., Staedler, Y.M., Schönenberger, J., Harrop. T.W.R., Tregear, J., Do, N.V., Gantet, P., Ghesquière, A., and Jouannic, S. (2017). Differences in meristem size and expression of branching genes are associated with variation in panicle phenotype in wild and domesticated African rice. EvoDevo 8: 2. doi:10.1186/s13227-017-0065-y.
- Daborn, P.J., Lumb, C., **Harrop, T.W.R.**, Blasetti, A., Pasricha, S., Morin, S., Mitchell, S.N., Donnelly, M.J., Müller, P. and Batterham, P. (2012). Using *Drosophila melanogaster* to validate metabolism-based insecticide resistance from insect pests. Insect Biochemistry and Molecular Biology 42, 918–924. doi:10.1016/j.ibmb.2012.09.003.

Book chapters

Cridge, A., **Harrop, T.W.R.**, Lovegrove, M., Remnant, E. & Dearden, P. (2017). *Nutrition and Epigenetic Change in Insects: Evidence and Implications*, in Advances in Insect Physiology (ed. Verlinden, H.) 53, 31–54 (Academic Press). doi:10.1016/bs.aiip.2017.06.001

Selected scientific presentations

- Genetic diversity and biocontrol in Argentine stem weevils. Genomics Aotearoa seminar series, New Zealand, 2021.
- The genomes of three invasive social wasps. **Oral presentation** at the Annual Conference of the Genetics Society of Australasia (GSA), Melbourne, Australia, 2019. **Symposium chair**: Genomics & bioinformatics.
- Comparative genomics of sexual and asexual parasitoid wasps from the Microctonus genus.

 Poster presentation at the 7th meeting of the European Society for Evolutionary

 Developmental Biology, Galway, Ireland, 2018.
- Genomics of a weevil pest and its parasitoid biocontrol agent. **Poster presentation** at the Annual Conference of the Genetics Society of Australasia (GSA) with the New Zealand Society for Biochemistry and Molecular Biology (NZSBMB), Dunedin, New Zealand, 2017.
- De novo sequencing of Argentine stem weevil and its parasitoid wasp biocontrol agent. **Poster presentation** at the 10th Annual Arthropod Genomics Symposium, Notre Dame University, South Bend, U.S.A., 2017.
- Sequencing and community annotation of the Vespula vulgaris genome. **Oral presentation** at the Entomological Society of NZ Conference, Wellington, New Zealand, 2017.
- Genes controlling transition of reproductive meristem identity during inflorescence development. **Oral presentation** at the 14th International Symposium on Rice Functional Genomics, Montpellier, France, 2016.

Mechanisms underlying the parallel evolution of inflorescence phenotype during independent domestication of African and Asian rice. **Contributed oral presentation** at the 6th meeting of the European Society for Evolutionary Developmental Biology, Uppsala, Sweden, 2016. **Joint symposium chair**: Branching across the tree of life.

Gene expression during O. sativa panicle development. **Invited seminar** at the Agricultural Genetics Institute, Hanoi, Vietnam, 2015.

Molecular mechanisms underlying the phenotypic convergence of inflorescence architecture in domesticated rice species. **Poster presentation** at the Society for Molecular Biology and Evolution annual meeting, Vienna, Austria, 2015.

Transcriptomic analysis of early developmental stages of the rice panicle. **Poster presentation** at the Workshop in Molecular Mechanisms Controlling Flowering, Aiguablava, Spain, 2015

RNA interference of the Cytochrome P450 redox partners Cpr and dare uncovers novel P450 functions in Drosophila melanogaster. **Poster presentation** at the 50th Annual Drosophila Research Conference, Chicago, USA, 2009.

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