# Dave Ting Pong Tang

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### Summary

I am currently a post-doctoral researcher at the University of Western Australian working on single cell transcriptomics. Prior to this position, I was working on the analysis of whole exome sequencing in patients with rare genetic disorders at the Telethon Kids Institute. During my PhD I was a Marie Curie Early Stage Researcher in the lab of Piero Carninci in RIKEN Yokohama and was working primarily on the analysis of high-throughput transcriptome sequencing data sets. I have also worked as a bioinformatician at the University of Queensland and the Commonwealth Scientific and Industrial Research Organisation.

## Education

2010-2015	PhD, Vrije University, the Netherlands. PhD thesis: High-throughput sequencing and transcriptomics: methods development and data analysis of large ex-
2001-2005	pression data sets.  BSc (Honours) in biochemistry and microbiology, University of Queensland, Australia. Honours thesis: Using a supertree approach to detect laterally trans-
	ferred genes within Staphylococcus.

### Past Scientific Positions

2015 - 2017	Post-doctoral researcher in the lab of Timo Lassmann at the Telethon Kids
	Institute, Australia
2010 – 2015	Research Associate in the lab of Piero Carninci at RIKEN Yokohama, Japan
2008 – 2010	Research Assistant in the lab of Sean Grimmond at the University of Queens-
	land, Australia
2006 – 2008	Research Assistant in the lab of Brian Dalrymple at the Commonwealth Sci-
	entific and Industrial Research Organisation, Australia
2005 - 2006	Research Assistant in the lab of Mark Ragan at the University of Queensland,
	Australia

### Grants

2017	Dissecting the leukaemia microenvironment - Cancer Council WA Collaborative
	Cancer Grant Scheme CIC AUD 43,395
2016	Targeting four-stranded DNA conformations to modulate gene expression in
	breast cancer - Cancer Council WA Collaborative Cancer Grant Scheme CIC
	AUD 47,669
2015	SeqNextGen: Translating NextGen Sequencing for the Diagnosis of Develop-
	mental Anomalies and Rare Diseases Telethon - Perth Children's Hospital Re-
	search Fund AI4 AUD 192,505

#### Honours and Awards

2010	CSIRO Chairman's Medal
2008	CSIRO Partnership Excellence Award

## Academic and Administrative Experience

2017	Presenter at the Australian Bioinformatics and Computational Biology Society
	(ABACBS) WA meeting
2015	MODHEP workshop presenter on analysing CAGE data
2013 – 2014	Organiser of the Chat with Guest sessions at RIKEN CLST DGT
2013 – 2014	Organiser of the Student Journal Club at RIKEN CLST DGT
2013	Organising committee for the BrainTrain conference
2013	Organiser of the BrainTrain workshop: Identifying regulatory elements in the
	genome
2012	Session chair for the Patients and Medicines forum
2012	Presenter at the RIKEN OSC bioinformatics course

### Workshop and Course Attendances

2016	AGHA Workshop: reducing morbidity and mortality from genetic disease
	through screening
2016	Consumer and Community Involvement in Research workshop
2015	UQ winter school in mathematical and computational biology
2014	RIKEN/KI doctoral course: Employing Genome-wide Technologies for Func-
	tional Regulation in Development and Disease
2013	AMATA conference ECR workshop
2013	Coursera data analysis course from Johns Hopkins University
2013	BrainTrain courses and workshops
2013	RIKEN/KI doctoral course: Epigenomics: Methods and Applications to Dis-
	ease and Development
2012	SISSA summer school on dopaminergic neurons
2012	RIKEN/KI doctoral course: Functional Regulation in Development and Dis-
	ease
2011	UQ winter school in mathematical and computational biology
2011	RIKEN/EBI bioinformatics roadshow

#### **Editorial Activities**

Referee/Reviewer (number of grant proposals/manuscripts reviewed in parenthesis) for: National Health and Medical Research Council (1), BMC Genomics (1), PeerJ (1), and Scientific Reports (2).

#### **Bioinformatic Skills**

- Data analysis of high-throughput sequencing data including DNA-seq, RNA-seq, CAGE-seq, sRNA-seq, ChIP-seq, and scRNA-seq.
- Knowledge and the ability to use various bioinformatic databases, APIs, repositories, and tools.
- Ability to implement bioinformatic pipelines using pipelining tools such as Bpipe and Snakemake.

# Computer Skills

- Operating systems: Linux/Unix (RHEL/CentOS and Ubuntu), OS X, and Windows.
- Programming/scripting languages: Perl, R, C, Python, Bash, SQL, and PHP.
- Reproducible research tools: git, cloud computing (AWS), WordPress, Jekyll, Docker, Markdown, R Markdown, and bookdown.

### **Hobbies and Interests**

Sports (especially basketball), cycling, blogging, data science, reading, and movies.

### Academic References

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Institute of Agriculture,
The University of Western Australia
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#### **Publications**

- [1] D. Tang, M. Fakiola, G. Syn, D. Anderson, H. J. Cordell, E. S. H. Scaman, E. Davis, S. J. Miles, T. McLeay, S. E. Jamieson, T. Lassmann, and J. M. Blackwell. Arylsulphatase A Pseudodeficiency (ARSA-PD), hypertension and chronic renal disease in Aboriginal Australians. *Sci Rep*, 8(1):10912, Jul 2018. URL: http://www.ncbi.nlm.nih.gov/pubmed/30026549.
- [2] K. M. Rothacker, K. L. Ayers, D. Tang, K. Joshi, J. A. van den Bergen, G. Robevska, N. Samnakay, L. Nagarajan, K. Francis, A. H. Sinclair, and C. S. Choong. A novel, homozygous mutation in desert hedgehog (DHH) in a 46, XY patient with dysgenetic testes presenting with primary amenorrhoea: a case report. *Int J Pediatr Endocrinol*, 2018:2, 2018. URL: https://www.ncbi.nlm.nih.gov/pubmed/29507583.
- [3] F. Roudnicky, L. C. Dieterich, C. Poyet, L. Buser, P. Wild, D. Tang, P. Camenzind, C. Hsien Ho, V. I. Otto, and M. Detmar. High expression of insulin receptor on tumor-associated blood vessels in invasive bladder cancer predicts poor overall and progression-free survival. J. Pathol., Mar 2017. URL: http://www.ncbi.nlm.nih.gov/pubmed/28295307.
- [4] C. C. Hon, J. A. Ramilowski, J. Harshbarger, N. Bertin, O. J. Rackham, J. Gough, E. Denisenko, S. Schmeier, T. M. Poulsen, J. Severin, M. Lizio, H. Kawaji, T. Kasukawa, M. Itoh, A. M. Burroughs, S. Noma, S. Djebali, T. Alam, Y. A. Medvedeva, A. C. Testa, L. Lipovich, C. W. Yip, I. Abugessaisa, M. Mendez, A. Hasegawa, D. Tang, et al. An atlas of human long non-coding RNAs with accurate 5' ends. Nature, Mar 2017. URL: http://www.ncbi.nlm.nih.gov/pubmed/28241135.
- [5] M. B. Abraham, D. Li, D. Tang, S. M. O'Connell, F. McKenzie, E. M. Lim, H. Hakonarson, M. A. Levine, and C. S. Choong. Short stature and hypoparathyroidism in a child with Kenny-Caffey syndrome type 2 due to a novel mutation in FAM111A gene. *Int J Pediatr Endocrinol*, 2017:1, 2017. URL: http://www.ncbi.nlm.nih.gov/pubmed/28138333.
- [6] G. Baynam, N. Pachter, F. McKenzie, S. Townshend, J. Slee, C. Kiraly-Borri, A. Vasudevan, A. Hawkins, S. Broley, L. Schofield, H. Verhoef, C. E. Walker, C. Molster, J. M. Blackwell, S. Jamieson, D. Tang, T. Lassmann, K. Mina, J. Beilby, M. Davis, N. Laing, L. Murphy, T. Weeramanthri, H. Dawkins, and J. Goldblatt. The rare and undiagnosed diseases diagnostic service application of massively parallel sequencing in a state-wide clinical service. Orphanet J Rare Dis, 11(1):77, 2016. URL: http://www.ncbi.nlm.nih.gov/pubmed/27287197.
- [7] D. Tang, D. Anderson, R. W. Francis, G. Syn, S. E. Jamieson, T. Lassmann, and J. M. Blackwell. Reference genotype and exome data from an Australian Aboriginal population for health-based research. *Sci Data*, 3:160023, 2016. URL: http://www.ncbi.nlm.nih.gov/pubmed/27070114.
- [8] Y. Hasegawa, D. Tang, N. Takahashi, Y. Hayashizaki, A. R. Forrest, et al. CCL2 enhances pluripotency of human induced pluripotent stem cells by activating hypoxia related genes. *Sci Rep*, 4:5228, 2014. URL: http://www.ncbi.nlm.nih.gov/pubmed/24957798.

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