Dave Ting Pong Tang

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Education

2010 – 2015	PhD Candidate, Vrije University, the Netherlands
2001 – 2005	BSc (Honours) in biochemistry and microbiology, University of Queens-
	land, Australia

Past Scientific Positions

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
	Queensland, Australia
2006 – 2008	Research Assistant in the lab of Brian Dalrymple at the Commonwealth
	Scientific and Industrial Research Organisation, Australia
2005 - 2006	Research Assistant in the lab of Mark Ragan at the University of Queens-
	land, Australia

Research Interests

- Genomics and transcriptomics; in particular the study of non-coding RNAs and transposable elements expression
- Open science and reproducible research
- The application of bioinformatics, in particular machine learning and data mining methods, to biological problems

Bioinformatic Skills

- Data analysis of high-throughput sequencing data from various *Seq protocols
- Knowledge and the ability to use various bioinformatic databases, repositories, and tools
- Biostatistics for the analysis of high-throughput sequencing data

Computer Skills

- Programming/scripting languages: Perl, R, SQL, bash, PHP, JavaScript, Python, and C
- Operating systems: Linux/Unix, OS X, and Windows
- Pre Web 2.0: CGI, PHP, HTML, MySQL, Apache2, and JavaScript
- Open science and reproducible research tools: git, GitHub, WordPress, Jekyll, and R Markdown/Markdown

Honours and Awards

2010	CSIRO Chairman's Medal
2008	CSIRO Partnership Excellence Award

Academic and Administrative Experience

2015	MODHEP Workshop presenter on analysing CAGE data using R
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

Workshop and Course Attendances

KI/RIKEN Doctoral Course: Employing Genome-wide Technologies for
Functional Regulation in Development and Disease
AMATA Conference ECR workshop
Courseras Data Analysis from Johns Hopkins University
BrainTrain courses and workshops
RIKEN/KI Doctoral Course: Epigenomics: Methods and Applications
to Disease and Development
SISSA Summer School on Dopaminergic Neurons
RIKEN/KI Doctoral Course: Functional regulation in development and
disease
UQ Winter School in Mathematical and Computational Biology
RIKEN/EBI Bioinformatics Roadshow

Hobbies and Interests

Sports (especially basketball), cycling, blogging, self study, and computer games.

Publications

[1] **Tang, Dave** and Piero Carninci. The regulated expression of repetitive elements across human cell types and tissues. *In preparation*, 2015.

- [2] **Tang, Dave**, Ana Maria Suzuki, Raffaella Calligaris, Stefano Gustincich, and Piero Carninci. Deep transcriptome sequencing of whole blood samples from Parkinson's disease patients. *In preparation*, 2015.
- [3] Y. Hasegawa, **Tang**, **D.**, 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.
- [4] Tang, D. T., C. Plessy, M. Salimullah, A. M. Suzuki, R. Calligaris, S. Gustincich, and P. Carninci. Suppression of artifacts and barcode bias in high-throughput transcriptome analyses utilizing template switching. *Nucleic Acids Res.*, 41(3):e44, Feb 2013. URL: http://www.ncbi.nlm.nih.gov/pubmed/23180801.
- [5] A. Saxena, **Tang, D.**, and P. Carninci. piRNAs warrant investigation in Rett Syndrome: an omics perspective. *Dis. Markers*, 33(5):261–275, 2012. URL: http://www.ncbi.nlm.nih.gov/pubmed/22976001.
- [6] S. Francia, F. Michelini, A. Saxena, Tang, D., M. de Hoon, V. Anelli, M. Mione, P. Carninci, and F. d'Adda di Fagagna. Site-specific DICER and DROSHA RNA products control the DNA-damage response. *Nature*, 488(7410):231–235, Aug 2012. URL: http://www.ncbi.nlm.nih.gov/pubmed/22722852.
- [7] R. D. Thiagarajan, N. Cloonan, B. B. Gardiner, T. R. Mercer, G. Kolle, E. Nourbakhsh, S. Wani, **Tang, D.**, K. Krishnan, K. M. Georgas, B. A. Rumballe, H. S. Chiu, J. A. Steen, J. S. Mattick, M. H. Little, and S. M. Grimmond. Refining transcriptional programs in kidney development by integration of deep RNA-sequencing and array-based spatial profiling. *BMC Genomics*, 12:441, 2011. URL: http://www.ncbi.nlm.nih.gov/pubmed/21888672.
- [8] R. D. Thiagarajan, K. M. Georgas, B. A. Rumballe, E. Lesieur, H. S. Chiu, D. Taylor, Tang, D. T., S. M. Grimmond, and M. H. Little. Identification of anchor genes during kidney development defines ontological relationships, molecular subcompartments and regulatory pathways. *PLoS ONE*, 6(2):e17286, 2011. URL: http://www.ncbi.nlm.nih.gov/pubmed/21386911.
- [9] H. S. Chiu, J. C. Szucsik, K. M. Georgas, J. L. Jones, B. A. Rumballe, **Tang, D.**, S. M. Grimmond, A. G. Lewis, B. J. Aronow, J. L. Lessard, and M. H. Little. Comparative gene expression analysis of genital tubercle development reveals a putative appendicular Wnt7 network for the epidermal differentiation. *Dev. Biol.*, 344(2):1071–1087, Aug 2010. URL: http://www.ncbi.nlm.nih.gov/pubmed/20510229.
- [10] N. Cloonan, Q. Xu, G. J. Faulkner, D. F. Taylor, **Tang, D. T.**, G. Kolle, and S. M. Grimmond. RNA-MATE: a recursive mapping strategy for high-throughput RNA-sequencing data. *Bioinformatics*, 25(19):2615–2616, Oct 2009. URL: http://www.ncbi.nlm.nih.gov/pubmed/19648138.
- [11] K. Georgas, B. Rumballe, M. T. Valerius, H. S. Chiu, R. D. Thiagarajan, E. Lesieur, B. J. Aronow, E. W. Brunskill, A. N. Combes, Tang, D., D. Taylor, S. M. Grimmond, S. S. Potter, A. P. McMahon, and M. H. Little. Analysis of early nephron patterning reveals a role for distal RV proliferation in fusion to the ureteric tip via a cap mesenchyme-derived connecting segment. Dev. Biol., 332(2):273–286, Aug 2009. URL: http://www.ncbi.nlm.nih.gov/pubmed/19501082.

- [12] Tang, D. T., E. A. Glazov, S. M. McWilliam, W. C. Barris, and B. P. Dalrymple. Analysis of the complement and molecular evolution of tRNA genes in cow. *BMC Genomics*, 10:188, 2009. URL: http://www.ncbi.nlm.nih.gov/pubmed/19393063.
- [13] J. W. Kijas, D. Townley, B. P. Dalrymple, M. P. Heaton, J. F. Maddox, A. McGrath, P. Wilson, R. G. Ingersoll, R. McCulloch, S. McWilliam, **Tang, D.**, J. McEwan, N. Cockett, V. H. Oddy, F. W. Nicholas, and H. Raadsma. A genome wide survey of SNP variation reveals the genetic structure of sheep breeds. *PLoS ONE*, 4(3):e4668, 2009. URL: http://www.ncbi.nlm.nih.gov/pubmed/19270757.