PHD STUDENT · BIOINFORMATICS AND COMPUTATIONAL BIOLOG

Melbourne, VIC, Australia

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#### **Education**

Monash University

Melbourne, Australia

PHD STUDENT (BIOINFORMATICS)

2019 - 2022

· Project title: Identification of biological signals by combining distinct categories of biological data through machine learning.

**Australian National University** 

Canberra, Australia

BACHELOR OF SCIENCE (MOLECULAR BIOLOGY AND PLANT SCIENCE); HONS. (BIOINFORMATICS)

2010 - 2013

### Research Experience \_\_\_\_

## **Bioinformatics lab**, School of Biological Sciences, Faculty of Science, Monash University (Dr Sonika Tyagi)

Melbourne, Australia

PHD STUDENT (BIOINFORMATICS)

2019 - PRESENT

- Applying machine learning to integrate biological data across multiple modalities.
- · Large scale data (1-10 TB) analysis.
- · Contributed to writing and data analysis for peer-reviewed academic publications.
- Open source software development for biological data analysis.

# Stemformatics Project, Australian Institute for Bioengineering and Nanotechnology, University of Queensland; Centre for Stem Cell Systems, Department of Anatomy and Neuroscience, University of Melbourne (Prof. Christine Wells)

Brisbane and Melbourne, Australia

RESEARCH ASSISTANT (BIOINFORMATICS)

2014-2016; 2016 - 2019

- Curated and ran large quantities of microarray, high throughput sequencing and single cell data through pipelines on high performance compute clusters to contribute to a database of interest for both internal and external collaborators.
- Large scale data (1-10 TB) and metadata standardisation/management as well as retrieving and transmitting these in an accessible format to requesters.
- Contributed to data analysis for peer-reviewed academic publications.
- Open source software development for microarray and high throughput sequencing pipelines.

## John Curtin School of Medical Research, Australian National University (Prof. Gavin Huttley)

Canberra, Australia

HONOURS SCHOLAR (BIOINFORMATICS)

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- Honours project on the development of a computational motif discovery toolkit for identification of signals in biological data.
- Implemented an algorithm in python using information theory and numerical optimisation to discover contiguous and discontigous signals in biological sequence data.

#### Research School of Biology, Australian National University (Prof. Murray Badger)

Canberra, Australia

SUMMER RESEARCH STUDENT (PLANT BIOLOGY, MICROSCOPY)

2012 - 2013

- Carried out a systematic analysis of physiology of the land plant *Hirschfeldia incana* to identify the principles underlying its relatively high photosynthetic rate compared to closely related plants in its family.
- Performed plant growth under controlled conditions and conducted subsequent physiological measurements including microscopy as well as
  associated imaging and leaf gas exchange measurements.

#### Research School of Biology, Australian National University (Dr. Iain Searle)

Canberra, Australia

Undergraduate research student (Molecular Biology, Plant Biology)

2012

- · Investigated the possible effect of mutations in the small RNA silencing mechanism which might contribute to genomic instability.
- Performed sterile plant and bacterial growth, PCR, transformation and gel electrophoresis, antibiotic screens, RNAase digestion, DNA extraction and media preparation.

## Teaching Experience

Monash University

Melbourne, Australia

2019-PRESENT

DATA FLUENCY ASSOCIATE INSTRUCTOR

· Running software carpentry sessions in unix shell, github, python, R, high performance computing cluster usage and deep learning.

SEPTEMBER 23, 2020 TYRONE CHEN 1

#### College of Medicine, Biology & Environment, Australian National University

Canberra, Australia

2011

PEER MENTOR FOR BIOLOGY STUDENTS

· Facilitated learning among undergraduate biology students by running tutoring sessions and designing exercises.

### Other Experience \_\_\_\_\_

**TraffixGroup** Melbourne, Australia

SUPERVISOR (TRAM SURVEYOR)

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• Led a team to count and interview passengers across Melbourne's tram routes.

#### Research School of Biology, Australian National University

Canberra, Australia

LABORATORY TECHNICIAN

2012

• Maintained a molecular biology laboratory focused on non-coding RNA in plants.

#### Honours & Awards

#### **AUSTRALIA**

#### **Australian Government Research Training Program Scholarship**

2019-2022 Australian Government Research Training Program Scholarship to pursue full-time candidature in the Doctor *Canberra*, *Australia* of Philosophy (PhD) with the School of Biological Sciences, Monash University.

#### **Faculty of Science Dean's Postgraduate Research Scholarship**

2019-2022 Faculty of Science Dean's Postgraduate Research Scholarship to pursue full-time candidature in the Doctor of Philosophy (PhD) with the School of Biological Sciences, Monash University.

Melbourne, Australia

#### **Data Fluency Digital Toolkit Poster Competition 2019 prize**

Awarded a prize for an oral presentation and associated poster presented as part of the Data Fluency Digital Toolkit Poster Competition 2019 held in Monash University.

Melbourne, Australia

#### **USA**

2013-2015 Sponsored Membership

Washington, DC,

American Association for the Advancement of Science (Science Program for Excellence in Science)

USA

#### Skills

#### COMPUTATIONAL

- · Machine learning: pytorch and tensorflow
- Languages and markup: python, R, bash, pSQL, LaTeX, restructured text
- · Version control: git, mercurial
- · Virtual environment management: conda, virtualenv, pip
- $\bullet \ \ \text{Usage of high performance compute clusters and associated job submissions: PBS, SGE, slurm$
- · Efficient management, organisation, archival, retrieval and standardisation of large quantities of data and metadata (1-10TB) per dataset
- Reproducible research practices and documentation: jupyter notebooks, Rmarkdown
- Workflow languages: Snakemake and Nextflow
- Graphic design for posters and scientific articles: Adobe inDesign, Adobe Illustrator example poster available online

#### BIOINFORMATICS, MATHEMATICS AND STATISTICS

- Data visualisation: PCA, interactive plots, genome browsers
- Differential gene expression analyses: ANOVA, pairwise t-tests
- Information theory: mutual information, relative entropy
- Signal processing: Fourier transform, autocorrelation
- Numerical optimisation: simulated annealing
- Motif enrichment analyses: ISMARA, MEME suite
- Batch effect correction: limma, ComBAT, supervised normalisation

#### BIOLOGY

- Microscopy: light microscopy and associated sample preparation
- · Plant physiology: Leaf gas exchange measurements, Membrane inlet mass spectrometry, fieldwork, plant growth
- Molecular biology: PCR, Molecular cloning, bacterial transformation, antibiotic screens, gel electrophoresis, RNAase digestion, DNA extraction, sterile plant growth, media preparation

#### OTHER

- Teaching and training: usage of microarray and high throughput sequencing pipelines, introduction to programming (python, R, unix shell, HPC), github, deep learning, peer mentoring for undergraduate biology students
- Project management and task organisation
- Awareness of cultural and professional diversity and clear communication
- · Social media editor [Stemformatics Project twitter], [Monash Computational Biology lab twitter]

### **Professional Development**

#### SOCIETY MEMBERSHIPS

- Software carpentries instructor 2020-present.
- Professional membership with the Australasian Genomic Technologies Association (AGTA) 2014-2015.
- · Professional membership with the Australian Bioinformatics And Computational Biology Society (ABACBS) 2014-present.
- Sponsored membership with the American Association for the Advancement of Science (Science Program for Excellence in Science) 2013-2015.
- Professional membership with the Institute of Electrical and Electronics Engineers (IEEE) 2013-present.

#### VOLUNTEER EXPERIENCE

- · Workshop assistance at the Introduction to Python workshop, 28th March, 2019, Monash University, Clayton, Australia.
- Community outreach with visiting students at the University of Melbourne (2016), University of Queensland (2014) and Australian National University (2013)

#### CONTRIBUTIONS TO OPEN SOURCE SOFTWARE LIBRARIES

- MONET multi-omics clustering algorithm. git repository
- Modules for use with the s4m shell framework used in the Stemformatics project, git repository
- The extra tools needed to run the Stemformatics project and add datasets into it. git repository
- Software carpentries lesson material. git repository

#### CONFERENCE AND WORKSHOP ATTENDANCE

- Conference presentation at the 28th International Conference on Intelligent Systems for Molecular Biology (ISMB), 13-16th July, 2020\*
- Conference presentation at the Bioinformatics Open Source Conference (BOSC 2020), 20-23rd July, 2020
- Conference presentation at the Australian Bioinformatics and Computational Biology Conference, 9-11th December, 2019, University of Sydney, Sydney, Australia.
- Conference presentation at the Australian Bioinformatics and Computational Biology Conference, 26-29th November, 2018, University of Melbourne, Melbourne, Australia.
- Conference presentation at the Single Cell Genomics Conference, 14-16th September, 2016, Wellcome Genome Campus Conference Centre, Cambridge, United Kingdom.
- Workshop attendance at the QFAB RNA-Sequencing workshop, 19-20th November, 2014, Queensland Biosciences Precinct, University of Oueensland, Australia.
- Conference presentation at the Australasian Genomic Technologies Association 14th Annual Conference (AGTA), 13-15th October, 2014, Crown Promenade Hotel, Melbourne, Australia.
- \* virtual participation due to the COVID-19 pandemic of 2020

## Publications . JOURNAL ARTICLES

Journa	AL ARTICLES	
2020	Paul W Angel, Nadia Rajab, Yidi Deng, Chris M Pacheco, <b>Tyrone Chen</b> , Kim-Anh Lê Cao, Jarny Choi, Christine A Wells. A simple, scalable approach to building a cross-platform transcriptome atlas <i>PLOS Computational Biology. doi: 10.1101/2020.03.09.984468</i>	Link
2020	<b>Tyrone Chen</b> , Sonika Tyagi. Integrative computational epigenomics to build data-driven gene regulation hypotheses <i>GigaScience. doi: 10.1093/gigascience/giaa064</i>	Link
2020	Ali Motazedian, Freya F. Bruveris, Santhosh V. Kumar, Jacqueline V. Schiesser, <b>Tyrone Chen</b> , Elizabeth S. Ng, Ann P. Chidgey, Christine A. Wells, Andrew G. Elefanty, Edouard G. Stanley. Multipotent RAG1+ progenitors emerge directly from haemogenic endothelium in human pluripotent stem cell-derived haematopoietic organoids. <i>Nature Cell Biology. doi: 10.1038/s41556-019-0445-8</i>	Link
2019	[Acknowledged] Nadia Rajab, Paul W Angel, Mariola Kurowska-Stolarska, Simon Milling, Chris M Pacheco, Matt Rutar, Jarny Choi, Christine A Wells. iMAC: An interactive atlas to explore phenotypic differences between in vivo, ex vivo and in vitro-derived myeloid cells in the Stemformatics platform. bioRxiv. doi: 10.1101/719237	Link
2018	[Acknowledged] Shahzya Chaudhury, Caitríona O'Connor, Ana Cañete, Joana Bittencourt-Silvestre, Evgenia Sarrou, Áine Prendergast, Jarny Choi, Pamela Johnston, Christine A. Wells, Brenda Gibson, Karen Keeshan. Age-specific biological and molecular profiling distinguishes paediatric from adult acute myeloid leukaemias. Nature communications. doi: 10.1038/s41467-018-07584-1	Link
2018	Jarny Choi, Chris M. Pacheco, Rowland Mosbergen, Othmar Korn, <b>Tyrone Chen</b> , Isha Nagpal, Steve Englart, Paul W. Angel, Christine A. Wells. Stemformatics: visualise and download curated stem cell data. <i>Nucleic Acids Research. doi: 10.1093/nar/gky1064</i>	Link
2018	[Acknowledged] Suzanne K. Butcher, Christine E. O'Carroll, Christine A. Wells, Ruaidhrí J. Carmody. Toll-like receptors drive specific patterns of tolerance and training on restimulation of macrophages. Frontiers in immunology. 2018;9.	Link
2016	Florian Rohart, Elizabeth A. Mason, Nicholas Matigian, Rowland Mosbergen, Othmar Korn, <b>Tyrone Chen</b> , Suzanne Butcher, Jatin Patel, Kerry Atkinson, Kiarash Khosrotehrani, Nicholas M. Fisk, Kim-Anh Lê Cao, Christine A. Wells (2016). A molecular classification of human mesenchymal stromal cells. <i>PeerJ</i> , <i>4</i> , <i>e1845</i> .	Link
2016	Kenneth Sinclair, Stephanie Yerkovich, <b>Tyrone Chen</b> , Jonathan McQualter, Peter Hopkins, Christine Wells, Daniel Chambers (2016). Mesenchymal stromal cells are readily recoverable from lung tissue, but not the alveolar space, in healthy humans. <i>Stem Cells. doi: 10.1002/stem.2419.</i>	Link
Confer	ENCE PROCEEDINGS AND POSTER PRESENTATIONS	
2020	<b>Tyrone Chen</b> , Kim-Anh Le Cao, Sonika Tyagi. Multi-omics data integration for the discovery of COVID-19 drug targets [version 1; not peer reviewed]. F1000Research 2020, 9 (ISCB Comm J):698 (poster) (https://doi.org/10.7490/f1000research.1118023.1). Presentation at the International Society for Computational Biology Conference 2020, Montreal, Canada (virtual due to COVID-19), Bioinformatics Community Conference 2020, Toronto, Canada (virtual due to COVID-19)	Link
2019	<b>Tyrone Chen</b> , Jason D. Rigby, Matthew D. McGee, Sonika Tyagi. Deep learning has potential for harmonising multi-omics data to discover weak regulatory features [version 1; not peer reviewed]. F1000Research 2019, 8:2068 (poster) (https://doi.org/10.7490/f1000research.1117749.1). Presentation at the Australian Bioinformatics and Computational Biology Society Annual Conference 2019, Sydney, New South Wales, Australia.	Link
2019	Paul W Angel, Matthew Rutar, Nadia Rajab, <b>Tyrone Chen</b> , Isha Nagpal, Jarny Choi, Christine A Wells. Mapping the blood cell landscape in stemformatics [version 1; not peer reviewed]. F1000Research 2019, 8:206 (poster) (https://doi.org/10.7490/f1000research.1116449.1). Presentation at the 40th Annual Lorne Genome Conference 2019, Lorne, Victoria, Australia.	Link

2018	Jarny Choi, <b>Tyrone Chen</b> , Chris M. Pacheco, Rowland Mosbergen, Othmar Korn, Isha Nagpal, Steve Englart, Paul W. Angel, Christine A. Wells. Stemformatics – visualise and download curated stem cell data [version 1; not peer reviewed]. F1000Research 2018, 7:1843 (poster) (https://doi.org/10.7490/f1000research.1116311.1). Presentation at the Australian Bioinformatics and Computational Biology Society Annual Conference 2018, Melbourne, Victoria, Australia. Joint presentation between the first two authors	Link
2018	Paul W Angel, <b>Tyrone Chen</b> , Jarny Choi, Kim-Anh Le Cao, Elizabeth Mason, Christine A Wells. Mapping the blood cell landscape in stemformatics [version 1; not peer reviewed]. F1000Research 2018, 7:1872 (poster) (https://doi.org/10.7490/f1000research.1116323.1). Presentation at the Australian Bioinformatics and Computational Biology Society Annual Conference 2018, Melbourne, Victoria, Australia.	Link
2018	Paul W Angel, <b>Tyrone Chen</b> , Jarny Choi, Othmar Korn, Rowland Mosbergen, Isha Nagpal, Chris Pacheco Rivera, Christine A. Wells. Stemformatics: Easy visualisation platform for well-curated stem cell data [version 1; not peer reviewed]. F1000Research 2018, 7:1007 (poster) (doi: 10.7490/f1000research.1115766.1). Presentation at the 16th International Society for Stem Cell Research Annual Meeting 2018, Melbourne, Victoria, Australia.	Link
2017	Chris Pacheco Rivera, Rowland Mosbergen, Othmar Korn, <b>Tyrone Chen</b> , Isha Nagpal, Christine A. Wells. Ontology challenges for the stem cell community: towards integrative data mining in the Stemformatics atlas. <i>ICBO 2017, 2137(1), 1 (Conference paper). Proceedings of the 8th International Conference on Biomedical Ontology 2017, Newcastle-upon-Tyne, United Kingdom.</i>	Link
2016	<b>Tyrone Chen</b> , Rowland Mosbergen, Othmar Korn and Christine Wells. Stemformatics: a visualisation platform for well-curated biological sequence data [version 1; not peer reviewed]. F1000Research 2016, 5:2352 (poster) (doi: 10.7490/f1000research.1113160.1). Presentation at the Single Cell Genomics Conference 2016, Cambridge, United Kingdom.	Link
2016	Jackson, L., Hopcroft, L.E.M., Rogers, S., Jorgensen, H., Pellicano, F., Wells, C., Mosbergen, R., <b>Chen, T.</b> , Vetrie, D. and Holyoake, T.L., 2016, January. Identifying genes and pathways deregulated in chronic myeloid leukaemia stem cells through meta-analysis of transcriptomic data. <i>In British Journal of Haematology (Vol. 173, pp. 122-122). Wiley-Blackwell.</i>	Link not available
2016	L Jackson, LEM Hopcroft, R Mosbergen, <b>T Chen</b> , S Kelly, O Khan, S Rogers, HG Jorgensen, D Vetrie, M Copland, CA Wells, TL Holyoake (2016). LEUKomics: Bringing high throughput stem cell data to the leukaemia research community. <i>Presentation at the 18th Annual John Goldman Conference on Chronic Myeloid Leukemia: Biology and Therapy 2016, Houston, USA.</i>	Link not available
2015	Suzanne Butcher, Edward Huang, Ashley Waardenberg, Othmar Korn, Rowland Mosbergen, <b>Tyrone Chen</b> , Kelly Hitchens, Dipti Vijayan, Anthony Beckhouse, Mark Walker, Tim Barnett, James Frazer, Antje Blumenthal, Matt Sweet, The FANTOM5 consortium, Christine Wells (2015). Diversification of innate immune responses by transcriptional mechanisms. <i>Presentation at the Lorne Infection and Immunity Conference 2015, Lorne, Australia.</i>	Link not available
2014	<b>Tyrone Chen</b> , Rowland Mosbergen, Othmar Korn, Christine A Wells (2014). Stemformatics: a user-friendly database of well-curated biological data. <i>Presentation at the 14th Annual Conference of the Australasian Genomic Technologies Association (AGTA), Melbourne, Victoria, Australia.</i>	Link
2013	<b>Tyrone Chen</b> , Yicheng Zhu, Gavin A Huttley (2013). Do DNA sequences influence recruitment of histone variants? ASMR New Investigator's Conference, Canberra. Australian Capital Territory, Australia.	Link not available