### Kim RUTHERFORD

Software Development and Bioinformatics

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### Profile

I am a software developer and bioinformatician. For the last 18 years I have been helping biologists to capture, analyse and interpret genomic data.

### Skills and Experience

- Software development: web, desktop and server
- Database administration and maintenance
- Genome annotation and analysis
- Processing, management and analysis of next generation sequencing data
- Systems administration including software and database installation and configuration

### Work experience

Sept 2012 - Scientific Officer Gemmell lab

Present Department of Anatomy, University of Otago

Tasks bioinformatics support for the group

genome and transcriptome assembly and analysis mentoring / training of group members in bioinformatics

☑ tuatara ☑ genome project

June 2010 – **Programmer** *S. pombe genome database (PomBase)*Present Department of Biochemistry, University of Cambridge

Tasks web and database programming

system administration for curation.pombase.org

Main project Canto - a community curation tool

Feb 2009 - June 
Computer Associate Baulcombe Group

2010 Department of Plant Sciences, University of Cambridge

Tasks bioinformatics support

creation of a short read analysis pipeline

Sept 2007 - Feb Programmer modENCODE Data Coordination Center

2009 Cambridge Systems Biology Centre

Tasks helped create the data warehouse for the data from modENCODE:

☑ modMine

Feb 2004 - Sept **Programmer** FlyMine/InterMine group

2007 Department of Genetics, University of Cambridge

Tasks | Java programming for the ☑ InterMine project

Sept 1998 - Feb Principal Computer Programmer Pathogen Sequencing Unit

2004 The Sanger Institute

Tasks programming and bioinformatics support

Major projects Artemis - a genome analysis and annotation tool

☑ ACT - a genome comparison viewer

### **Publications**

Feb 2000 The genome sequence of the food-borne pathogen Campylobacter jejuni reveals hypervariable sequences.

J. Parkhill et al.

Nature. 2000 Feb 10;403(6770):665-8.

# Mar 2000 Complete DNA sequence of a serogroup A strain of Neisseria meningitidis Z2491.

J. Parkhill et al.

Nature. 2000 Mar 30;404(6777):502-6.

#### Oct 2000 Artemis: sequence visualisation and annotation.

K. Rutherford, J. Parkhill, J. Crook, T. Horsnell, P. Rice, M-A. Rajandream and B. Barrell

Bioinformatics. 2000 Oct;16(10):944-5.

#### Feb 2001 Massive gene decay in the leprosy bacillus.

S. Cole et al.

Nature. 2001 Feb 22;409(6823):1007-11.

#### June 2001 A Re-annotation of the Saccharomyces cerevisiae Genome.

V. Wood, K. M. Rutherford, A. Ivens, M-Å Rajandream and B. Barrell Comp Funct Genomics. 2001 June; 2(3): 143-154.

### Oct 2001 Genome sequence of Yersinia pestis, the causative agent of plague.

I. Parkhill et al.

Nature. 2001 Oct 4;413(6855):523-7.

## Oct 2001 Complete genome sequence of a multiple drug resistant Salmonella enterica serovar Typhi CT18.

J. Parkhill et al.

Nature, 2001 Oct 25:413(6858):848-52.

#### Feb 2002 The genome sequence of Schizosaccharomyces pombe.

V. Wood et al.

Nature. 2002 Feb 21;415(6874):871-80.

## May 2002 Complete genome sequence of the model actinomycete Streptomyces coelicolor A3(2).

S. Bentley et al.

Nature. 2002 May 9;417(6885):141-7.

#### Oct 2002 Sequence of Plasmodium falciparum chromosomes 1, 3-9 and 13.

N. Hall et al.

Nature. 2002 Oct 3;419(6906):527-31.

## Oct 2002 **Genome sequence of the human malaria parasite** *Plasmodium falciparum*.

I. Malcolm et al.

Nature. 2002 Oct 3;419(6906):498-511.

### June 2003 Viewing and annotating sequence data with Artemis.

M. Berriman and K. Rutherford

Brief Bioinform. 2003 Jun;4(2):124-32.

### Aug 2003 The DNA sequence of chromosome I of an African trypanosome:

Gene content, chromosome organisation, recombination and polymorphism

N. Hall et al.

Nucl. Acids Res. (2003) 31 (16): 4864-4873.

## Nov 2003 The complete genome sequence and analysis of Corynebacterium diphtheriae NCTC13129.

A. Cerdeño-Tárraga et al.

Nucleic Acids Res. 2003 Nov 15;31(22):6516-23.

### Jan 2004 GeneDB: a resource for prokaryotic and eukaryotic organisms.

Hertz-Fowler C et al.

Nucleic Acids Res. 2004 Jan 1;32(Database issue):D339-43.

Complete	genomes o	of two clinica	I Staphyloco	occus aure	<i>eus</i> strains:
<b>Evidence</b>	for the rap	id evolution	of virulence	and drug	resistance.

M. Holden et al.

Proc Natl Acad Sci U S A. 2004 Jun 29;101(26):9786-91a

# Sep 2004 **Genomic plasticity of the causative agent of melioidosis, Burkholderia pseudomallei.**

M. Holden et al.

Proc Natl Acad Sci U S A. 2004 Sep 28;101(39):14240-5.

# Jan 2005 A comprehensive survey of the *Plasmodium* life cycle by genomic, transcriptomic, and proteomic analyses

N. Hall

Science 07 Jan 2005: Vol. 307, Issue 5706, pp. 82-86

# June 2005 A Human-Curated Annotation of the Candida albicans Genome Braun et al.

PLoS Genet. 2005 Jul;1(1):36-57. Epub 2005 Jun 17

#### Aug 2005 ACT: the Artemis comparison tool.

T. Carver, K. Rutherford, M. Berriman, M-A. Rajandream, B.. Barrell and J. Parkhill

Bioinformatics. 2005 Aug 15;21(16):3422-3

### Sep 2005 WebACT - An online companion for the Artemis Comparison Tool

J. Abbott, D. Aanensen, K. Rutherford, S. Butcher and B. Spratt Bioinformatics. 2005 Sep 15;21(18):3665-6. Epub 2005 Aug 2

## July 2007 FlyMine: an integrated database for *Drosophila* and *Anopheles* genomics.

R. Lyne et al.

Genome Biol. 2007;8(7):R129.

### Dec 2010 Integrative Analysis of the *C. elegans* Genome by the modENCODE Project.

The modENCODE Consortium Science. 2010 Dec 24;330(6012):1775-87.

## Dec 2010 Identification of Functional Elements and Regulatory Circuits by Drosophila modENCODE.

The modENCODE Consortium *Science. 2010 Dec 24;330(6012):1787-97.* 

#### Nov 2011 The Gene Ontology: enhancements for 2011.

Gene Ontology Consortium. *Nucleic Acids Res. 2012 Jan;40(Database issue):D559-64. Epub 2011 Nov 18.* 

#### Jan 2012 PomBase: a comprehensive online resource for fission yeast.

V. Wood, M.A. Harris, M.D. McDowall, K. Rutherford, B.W. Vaughan, D.M. Staines, M. Aslett, A. Lock, J. Bähler, P.J. Kersey, S.G. Oliver *Nucleic Acids Res. 2012 Jan;40(Database issue):D695-9.* 

### Jan 2012 modMine: flexible access to modENCODE data.

S. Contrino, R.N. Smith, D. Butano, A. Carr, F. Hu, R. Lyne, K. Rutherford, A. Kalderimis, J. Sullivan, S. Carbon, E.T. Kephart, P. Lloyd, E.O. Stinson, N.L. Washington, M.D. Perry, P. Ruzanov, Z. Zha, S.E. Lewis, L.D. Stein, G. Micklem

Nucleic Acids Res. 2012 Jan; 40 (Database issue): D1082-8.

## Sep 2012 InterMine: a flexible data warehouse system for the integration and analysis of heterogeneous biological data

R.N. Smith, J. Aleksic, D. Butano, A. Carr, S. Contrino, F. Hu, M. Lyne, R. Lyne, A. Kalderimis, K. Rutherford, R. Stepan, J. Sullivan, M. Wakeling, X. Watkins, G. Micklem

Bioinformatics. (2012) 28 (23):3163-3165.

- Gene ontology annotations and resources. Jan 2013 Gene Ontology Consortium Nucleic Acids Res. 2013 Jan;41(Database issue):D530-5. doi: 10.1093/nar/gks1050 Feb 2014 Canto: An online tool for community literature curation K. Rutherford, M. A. Harris, A. Lock, S. G. Oliver and V. Wood Bioinformatics (2014) doi: 10.1093/bioinformatics/btu103 lune 2014 Improving functional annotation for industrial microbes: A case study with Pichia pastoris D. Dikicioglu, V. Wood, K. Rutherford, M. McDowall, S. Oliver Trends in Biotechnology (2014) 10.1016/j.tibtech.2014.05.003 Oct 2014 PomBase 2015: updates to the fission yeast database M. McDowall, M. Harris, A. Lock, K. Rutherford, D. Staines, J. Bähler, P. Kersey, S. Oliver and V. Wood Nucl. Acids Res. (2014) doi: 10.1093/nar/gku1040 Dec 2014 Molecular evolution of Dmrt1 accompanies change of sexdetermining mechanisms in reptilia D. E. Janes et al. Biol Lett. 2014 Dec; 10(12): 20140809. Jan 2015 Gene Ontology Consortium: going forward The Gene Ontology Consortium Nucl. Acids Res. (28 January 2015) 43 (D1): D1049-D1056. Nov 2015 Large-scale transcriptome sequencing reveals novel expression patterns for key sex-related genes in a sex-changing fish H. Liu, M. Lamm, K. Rutherford, M. Black, J. Godwin and N. Gemmell Biology of Sex Differences (2015) 6:26 Feb 2016 Uncovering the pathways underlying whole body regeneration in a chordate model, Botrylloides leachi using de novo transcriptome analysis L. Zondag, K. Rutherford, N. Gemmell and M. Wilson BMC Genomics (2016) 17:114 DOI 10.1186/s12864-016-2435-6 PHI-base: A new interface and further additions for the multi-Jan 2017 species pathogen-host interactions database.
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Nucleic Acids Research (Jan 2017database issue) accepted.

Urban, M., Alayne Cuzick, A., Rutherford, K., Irvine, A., Pedro, H., Pant, R., Sadanadan, V., Khamari, L., Billal, S., Mohanty S., and. Hammond-Kosack,