# **Graduate Engineer · Computational Biologist**

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## **WORK EXPERIENCE**

# **Dr. Stephen Sansom** (University of Oxford)

Oxford, UK

POSTDOCTORAL RESEARCHER — COMPUTATIONAL BIOLOGIST

Sep. 2017 - PRESENT

- Molecular pathogenesis of inflammatory bowel disease, in collaboration with the group of Prof. Fiona Powrie
- Role of thymic epithelial cells in T-cell development, in collaboration with Prof. Georg Holländer and Prof. Chris Ponting.
- Development and maintainenance of software pipelines and programs for data analysis:
  - o 10x Genomics analysis pipeline ( sansomlab/tenx ).
  - o Configurable Shiny application for SummarizedExperiment objects (☑ iSEE).

## **Prof. David Mole** (University of Oxford)

Oxford, UK

POSTDOCTORAL RESEARCHER — COMPUTATIONAL BIOLOGIST

Dec. 2016 - Sep. 2017

- Management, quality control, and coordination of genomics data for integrative analyses
  of DNA targeted resequencing, ChIP-seq, RNA-seq in the context of oxygen sensing and
  renal cancer.
- Prototype of MeteorJS application to manage sequencing data (

   <u>kevinrue/Seqbook</u>).

#### **Prof. Martin Wilkins** (Imperial College London)

London, UK

RESEARCHER ASSOCIATE — BIOINFORMATICS / BIOSTATISTICS

Sep. 2015 - Dec. 2016

- Integration of genetic variation, proteomic and metabolomics data and associated deep phenotype data, in collaboration with the *Computational Genome Biology and Medicine* team led by Dr. Stefan Gräf (University of Cambridge).
- Functions for analysing genetic variants ( TVTB).

#### **Dr. Stewart Adcock** (MEDIT S.A.)

Palaiseau, France

SOFTWARE DEVELOPER (INTERN)

*Apr.* 2012 – Jul. 2012

• Implementation of structurally-constrained multiple alignment of protein sequences for the commercial software MED-SuMo.

**Dr. Nora Khaldi, Prof. Denis Shields** (University College Dublin)

Dublin, Ireland

RESEARCH ASSISTANT - BIOINFORMATICS (INTERN)

Jun. 2010 - Sep. 2010

Analysis of structural disorder in Saccharomyces cerevisiae interacting proteins.

**Dr. Henri De La Salle** (Établissement Français du Sang)

Strasbourg, France

RESEARCH ASSISTANT (INTERN)

Jul. 2009 – Aug. 2009

Recombinant protein expression in HeLa cells.

# **EDUCATION**

**Prof. David MacHugh, Prof. Stephen Gordon** (University College Dublin) Dublin, Ireland PhD CANDIDATE 2011 – 2015

- **Thesis**: Comparative systems biology analyses of the bovine transcriptional response to species of the *Mycobacterium* genus.
  - o Functions for Gene Ontology analysis of gene expression data (**Z** GOexpress).
- Rotation year projects:
  - In silico structure-based prediction of *Mycobacterium bovis* epitopes in cattle (Dr. Anthony Chubb).
  - Expression and purication of a Hepatitis B protein construct for experimental screening of interacting drug fragments (Dr. Neil Ferguson).
  - Evaluation of a gene candidate underlying Staphylococcus aureus antibiotic resistance by Sanger sequencing and biofilm assay (Prof. James O'Gara).

# École Polytechnique Universitaire Polytech Nice-Sophia

Sophia-Antipolis, France 2008 – 2011

- INGÉNIEUR DIPLÔMÉ
- Biological engineering; Pharmacology, Biotechnology, Toxicology, Human and Environmental Health.
- Major: Bioinformatics and Modelling for Biology.

# **FUNDING & AWARDS**

- Conference travel grant: Bioconductor 2019 conference (New York, USA).
- Winner of the 1st RStudio Shiny Contest: "Most technically impressive" app.
- Conference travel grant: Bioconductor 2017 conference (Boston, USA).
- Conference fee waiver: ISMB conference 2015 (Dublin, Ireland); Bioinformatics Open Source Conference (BOSC) Special Interest Group.
- Conference travel grant: FEMS Young Scientist Meeting Grant (YSMG); FEMS 2014 conference "Microbiology in the New Genomics Era" (Paris, France).
- **Best oral presentation**: Computational Biology and Innovation PhD Symposium 2014 (Dublin, Ireland).
- Wellcome Trust Four-year PhD Programme: Computational Infection Biology.

#### **CONFERENCE PRESENTATIONS**

- Short talk: Bioconductor 2019 conference (New York, USA).
- Poster: Genome Informatics 2018 (Cambridge, UK).
- Short talk: Bioconductor 2017 conference (Boston, USA).
- Talk: NGS-SIG Single-cell RNAseq (Oxford, UK).
- Poster: Quantitative Genomics 2015 (London, UK).
- Talk: Bioinformatics Open Source Conference (BOSC) 2015 (Dublin, Ireland).
- Poster: Wellcome Trust Final Year PhD Students' Meeting 2014 (London, UK).
- Poster: UCD Conway Festival of Research & Innovation 2014 (Dublin, Ireland).
- Talk: The Acid Fast Club, Summer Meeting 2014 (Berlin, Germany).
- Poster: EMBO conference 2014 (Paris, France).
- Talk: Virtual Institute of Bioinformatics & Evolution (VIBE) 2014 (Carlow, Ireland).
- Poster: Society for General Microbiology (SGM) 2014 (Dublin, Ireland).

# **CONFERENCE PRESENTATIONS (CONTINUED)**

- Poster: Dublin Academy of Pathogenomics & Infection Biology (DAPI) 2013 (Ireland).
- Poster: Computational Biology & Innovation Symposium 2013 (Dublin, Ireland).
- Talk: Virtual Institute of Bioinformatics and Evolution (VIBE) 2013 (Galway, Ireland)
- Poster: Animal Health Ireland Workshop & Conference 2013 (Cork, Ireland).
- Talk: Association of Veterinary Teachers and Research Workers (AVTRW) 2013 (Hillsborough, Ireland)

### **ORGANIZER & CHAIR**

- Co-organiser: Bioconductor 2019 Special Interest Group (New York, USA).
- Coordinator: Genomics Forum, Kennedy Institute of Rheumatology (2018-present).
- Co-organiser: Computational Biology and Innovation PhD Symposium (2012).

#### **TEACHING**

- Workshop: Bioconductor 2019 conference (New York, USA).
- Undergraduate module: Agricultural Microbiology (UCD, 2014).
- Undergraduate module: Molecular Genetics and Biotech (UCD, 2014).

#### **TRAINING & COURSES**

- Machine Learning (Coursera, Stanford University, 2016).
- Python for Genomic Data Science (Coursera, Johns Hopkins University, 2015).
- Command Line Tools for Genomic Data Science (Coursera, JHU, 2015).
- Bioconductor for Genomic Data Science (Coursera, Johns Hopkins University, 2015).
- Statistics for Genomic Data Science (Coursera, Johns Hopkins University, 2015).
- Teaching in Higher Education (2014).
- Network Analysis in Systems Biology (Coursera, 2013)
- Sequence Data Analysis Training (Wageningen Institute of Animal Sciences, 2013).
- Hot Topics in Food and Nutrition Research (University College Dublin, 2013)
- Genomics Principles and Practical Applications (University College Dublin, 2012).
- Bioinformatics Research Seminars (University College Dublin, 2012).
- Advances in Infection Biology (University College Dublin, 2012).
- Online Research Skills for Computational Biologists (University College Dublin, 2012).
- Python Programming for Computational Biologists (University College Dublin, 2012).
- Introductory Statistics using R for computational biologists (UCD, 2012).

#### PROFESSIONAL MEMBERSHIPS

• International Society for Computational Biology (ISCB): 2012-present.

#### PEER REVIEW

- OUP Bioinformatics: 1 article
- Journal of Open Source Software: 1 article

# SKILLS & EXPERIENCE

Programming (Alphabetical)	C#, Java, JavaScript, Matlab, PHP, Python, R, SQL, Unix, XHTML.
Software (Alphabetical)	Bioconductor, cgat-core, cgat-apps, cgat-flow, conda, Django, Eclipse, git, GitHub, MongoDB, MeteorJS, NodeJS, PyCharm, Rstudio, Ruffus, Subversion, Visual Studio.
<b>Creativity</b> (Release date)	Bioconductor packages (iSEE, TVTB, GOexpress).
Languages (Fluency)	French (native), English (fluent, TOEIC 970), Italian (working), Spanish (limited), German (limited).

Publications: 18 h-index: 11 Total citations: 540

# Manuscripts awaiting publication

Robert A. Amezquita, Vince J. Carey, Lindsay N. Carpp, Ludwig Geistlinger, Aaron T. L. Lun, Federico Marini, **Kevin Rue-Albrecht**, Davide Risso, Charlotte Soneson, Levi Waldron, Hervé Pagès, Mike Smith, Wolfgang Huber, Martin Morgan, Raphael Gottardo, Stephanie C. Hicks (2019). "Orchestrating Single-Cell Analysis with Bioconductor". <u>Nature Methods</u>

2019

Schulthess, J., S. Pandey, M. Capitani, **K. C. Rue-Albrecht**, I. Arnold, F. Franchini, A. Chomka, N. E. Ilott, D. G. W. Johnston, E. Pires, J. McCullagh, S. N. Sansom, C. V. Arancibia-Carcamo, H. H. Uhlig and F. Powrie (2019). "The Short Chain Fatty Acid Butyrate Imprints an Antimicrobial Program in Macrophages." <a href="mailto:linearing-in-ma

#### 2018

Aulicino, A., **K. C. Rue-Albrecht**, L. Preciado-Llanes, G. Napolitani, N. Ashley, A. Cribbs, J. Koth, B. C. Lagerholm, T. Ambrose, M. A. Gordon, D. Sims and A. Simmons (2018). "Invasive Salmonella exploits divergent immune evasion strategies in infected and bystander dendritic cell subsets." <u>Nat Commun</u> **9**(1): 4883.

O'Doherty, A. M., **K. C. Rue-Albrecht**, D. A. Magee, S. Ahting, R. E. Irwin, T. J. Hall, J. A. Browne, N. C. Nalpas, C. P. Walsh, S. V. Gordon, M. W. Wojewodzic and D. E. MacHugh (2019). "The bovine alveolar macrophage DNA methylome is resilient to infection with Mycobacterium bovis." <u>Sci Rep</u> **9**(1): 1510.

Malone, K. M., **K. Rue-Albrecht**, D. A. Magee, K. Conlon, O. T. Schubert, N. C. Nalpas, J. A. Browne, A. Smyth, E. Gormley, R. Aebersold, D. E. MacHugh and S. V. Gordon (2018). "Comparative 'omics analyses differentiate Mycobacterium tuberculosis and Mycobacterium bovis and reveal distinct macrophage responses to infection with the human and bovine tubercle bacilli." <u>Microb Genom</u>.

Villarreal-Ramos, B., S. Berg, A. Whelan, S. Holbert, F. Carreras, F. J. Salguero, B. L. Khatri, K. Malone, **K. Rue-Albrecht**, R. Shaughnessy, A. Smyth, G. Ameni, A. Aseffa, P. Sarradin, N. Winter, M. Vordermeier and S. V. Gordon (2018). "Experimental infection of cattle with Mycobacterium tuberculosis isolates shows the attenuation of the human tubercle bacillus for cattle." <u>Sci Rep</u> **8**(1): 894.

**Rue-Albrecht, K.**, F. Marini, C. Soneson and A. T. L. Lun (2018). "iSEE: Interactive SummarizedExperiment Explorer." <u>F1000Res</u> **7**: 741.

Correia, C. N., K. E. McLoughlin, N. C. Nalpas, D. A. Magee, J. A. Browne, **K. Rue-Albrecht**, S. V. Gordon and D. E. MacHugh (2018). "RNA Sequencing (RNA-Seq) Reveals Extremely Low Levels of Reticulocyte-Derived Globin Gene Transcripts in Peripheral Blood From Horses (Equus caballus) and Cattle (Bos taurus)." <u>Front Genet</u> **9**: 278.

#### 2017

Rhodes, C. J., P. Ghataorhe, J. Wharton, **K. C. Rue-Albrecht**, C. Hadinnapola, G. Watson, M. Bleda, M. Haimel, G. Coghlan, P. A. Corris, L. S. Howard, D. G. Kiely, A. J. Peacock, J. Pepke-Zaba, M. R. Toshner, S. J. Wort, J. S. Gibbs, A. Lawrie, S. Graf, N. W. Morrell and M. R. Wilkins (2017). "Plasma Metabolomics Implicates Modified Transfer RNAs and Altered Bioenergetics in the Outcomes of Pulmonary Arterial Hypertension." <u>Circulation</u> **135**(5): 460-475.

#### 2016

**Rue-Albrecht, K.**, P. A. McGettigan, B. Hernandez, N. C. Nalpas, D. A. Magee, A. C. Parnell, S. V. Gordon and D. E. MacHugh (2016). "GOexpress: an R/Bioconductor package for the identification and visualisation of robust gene ontology signatures through supervised learning of gene expression data." <u>BMC Bioinformatics</u> **17**: 126.

Farrell, D., G. Jones, C. Pirson, K. Malone, **K. Rue-Albrecht**, A. J. Chubb, M. Vordermeier and S. V. Gordon (2016). "Integrated computational prediction and experimental validation identifies promiscuous T cell epitopes in the proteome of Mycobacterium bovis." <u>Microb Genom</u> **2**(8): e000071.

#### 2015

Nalpas, N. C., D. A. Magee, K. M. Conlon, J. A. Browne, C. Healy, K. E. McLoughlin, **K. Rue-Albrecht**, P. A. McGettigan, K. E. Killick, E. Gormley, S. V. Gordon and D. E. MacHugh (2015). "RNA sequencing provides exquisite insight into the manipulation of the alveolar macrophage by tubercle bacilli." <u>Sci Rep</u> **5**: 13629.

Park, S. D., D. A. Magee, P. A. McGettigan, M. D. Teasdale, C. J. Edwards, A. J. Lohan, A. Murphy, M. Braud, M. T. Donoghue, Y. Liu, A. T. Chamberlain, **K. Rue-Albrecht**, S. Schroeder, C. Spillane, S. Tai, D. G. Bradley, T. S. Sonstegard, B. J. Loftus and D. E. MacHugh (2015). "Genome sequencing of the extinct Eurasian wild aurochs, Bos primigenius, illuminates the phylogeography and evolution of cattle." Genome Biol **16**: 234.

Aulicino, A., A. M. Dinan, A. A. Miranda-CasoLuengo, J. A. Browne, **K. Rue-Albrecht**, D. E. MacHugh and B. J. Loftus (2015). "High-throughput transcriptomics reveals common and strain-specific responses of human macrophages to infection with Mycobacterium abscessus Smooth and Rough variants." <u>BMC Genomics</u> **16**: 1046.

#### 2014

**Rue-Albrecht, K.**, D. A. Magee, K. E. Killick, N. C. Nalpas, S. V. Gordon and D. E. MacHugh (2014). "Comparative functional genomics and the bovine macrophage response to strains of the mycobacterium genus." <u>Front Immunol</u> **5**: 536.

McLoughlin, K. E., N. C. Nalpas, **K. Rue-Albrecht**, J. A. Browne, D. A. Magee, K. E. Killick, S. D. Park, K. Hokamp, K. G. Meade, C. O'Farrelly, E. Gormley, S. V. Gordon and D. E. MacHugh (2014). "RNA-seq Transcriptional Profiling of Peripheral Blood Leukocytes from Cattle Infected with Mycobacterium bovis." <u>Front Immunol</u> **5**: 396.

### 2013

Nalpas, N. C., S. D. Park, D. A. Magee, M. Taraktsoglou, J. A. Browne, K. M. Conlon, **K. Rue-Albrecht**, K. E. Killick, K. Hokamp, A. J. Lohan, B. J. Loftus, E. Gormley, S. V. Gordon and D. E. MacHugh (2013). "Whole-transcriptome, high-throughput RNA sequence analysis of the bovine macrophage response to Mycobacterium bovis infection in vitro." <u>BMC Genomics</u> **14**: 230.

#### 2012

**Rue-Albrecht, K.**, D. C. Shields and N. Khaldi (2012). "Correlation of disorder between S. cerevisiae interacting proteins." Mol Biosyst 8(1): 417-425.