# Konrad Rudolph

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# Wellcome Trust/Cancer Research UK Gurdon Institute Tennis Court Road Cambridge CB2 1QN, UK

2008

2008-2011

### DEGREES

### UNIVERSITY OF CAMBRIDGE

PhD 2016 **Bioinformatics** 

### FREIE UNIVERSITÄT BERLIN

Master of Science 2011 Bioinformatics

Bachelor of Science 2008 Major: bioinformatics

Minor: physics

### EMBL PhD fellowship

Awarded 4 year funded PhD fellowship

### **LANGUAGES**

Fluency in written & spoken English, German, and

### **ANALYSIS**

Data exploration · data visualisation · statistical modelling · RNA-seq · ChIP-seq · motif analysis

### **PROGRAMMING**

 $C++\cdot C\cdot R\cdot Python\cdot Perl\cdot Bash\cdot Make\cdot C\#\cdot$ Java · PHP · VB · JavaScript · Ruby · ...

### TOOLS

Version control · unit testing · reproducible research · literate programming · lexical/semantic analysis · Unix · Windows · macOS

# MANAGEMENT

Software development team lead · EMBL PhD Symposium organising committee · EMBL Bioinformatics Workshop coordination

### **INVITED TALKS**

Cambridge Epigenetics Symposium 2014

### **GITHUB**

klmr

### ORCID

0000-0002-9866-7051

## STACK OVERFLOW

Konrad Rudolph

# **FXPERIENCE**

### UNIVERSITY OF CAMBRIDGE

Postdoctoral research associate May 2016-

Gene regulation by non-coding RNA · Nuclear RNAi · RNA modifications · transposable elements · transgenerational epigenetic inheritance

Group leader: Prof Dr Eric Miska

### **FMRI-FRI**

Postdoctoral fellow

Oct 2015-Mar 2016 Group leader: Dr John Marioni

Predoctoral fellow

Oct 2011-Sep 2015

Link between tRNA and mRNA abundance in mammals via codon-anticodon pairing Group leader: Dr John Marioni

# INDEPENDENT CONSULTANCY

Developer Jun 2011-Aug 2011

Integration of FPGA kernel with C++ library

# ILLUMINA INC.

Research associate (intern) Oct 2008-Jan 2009

Implementation of short-read mapping on GPGPUs with Nvidia CUDA

# FREIE UNIVERSITÄT BERLIN

Master project student 2010-2011

Thesis: Generic parallelisation of a sequence analysis library

Group leader: Prof Dr Knut Reinert

Tutor 2008-2011

Bachelor project student

Thesis: Implementation of a read mapping tool based on the pigeon-hole principle

Group leader: Prof Dr Knut Reinert

### **ITOSA**

Full stack developer Jan 2007-Dec 2007

C# Windows application; PHP/JavaScript/HTML web application & database frontend

# UNIVERSITY OF CAMBRIDGE

smallRNA sequencing 2017 NST Part II BBS Bioinformatics minor 2017 Next generation sequencing (focus on RNA-seg & ChIP-seg) 2013-2015

**EMBL-EBI** 

Bioinformatics workshop 2011-2015 Bash scripting · Unix · Git · R · LATEX

FREIE UNIVERSITÄT BERLIN

Algorithms 101 · Algorithms 102 · Algorithms in bioinformatics · Database systems · C++

Katharina Gapp, Gretchen van Steenwyck, Wayo Matsushima, Konrad L M Rudolph, Francesca Manuella, Grégoire Vernaz, Tanay Gosh, Pawel Pelzcar, Isabelle M Mansuy, Eric A Miska. "A sperm long RNA fraction transmits effects of early life traumatic stress." (submitted).

Alper Akay, Tomás Di Domenico, Kin M Suen, Amena Nabih, Guillermo E Parada, Mark Larance, Ragini Medhi, Ahmet C Berkyurek, Xinlian Zhang, Christopher J Wedeles, Konrad L M Rudolph, Jan Engelhardt, Martin Hemberg, Ping Ma, Angus I Lamond, Julie M Claycomb, Eric A Miska. "Small RNA pathways require the helicase Aquarius/EMB0-4 to heritably silence transcription." Developmental Cell (accepted).

Pieter van Delft, Alper Akay, Sabrina M Huber, Christoph Bueschl, Konrad L M Rudolph, Tomás Di Domenico, Rainer Schuhmacher, Eric A Miska, Shankar Balasubramanian. "The profile and dynamics of RNA modifications in animals." ChemBioChem (Apr 2017), DOI: 10.1002/cbic.201700093.

Konrad L M Rudolph\*, Bianca M Schmitt\*, Diego Villar, Robert J White, John C Marioni, Claudia Kutter, Duncan T Odom. "Codon-driven translational efficiency is stable across diverse mammalian cell states." PLOS Genetics (May 2016), DOI: 10.1371/journal.pgen.1006024.

Bianca M Schmitt\*, Konrad L M Rudolph\*, Panagiota Karagianni, Nuno A Fonseca, Robert J White, Iannis Talianidis, Duncan T Odom, John C Marioni, Claudia Kutter. "High-resolution mapping of transcriptional dynamics across tissue development reveals a stable mRNA-tRNA interface." Genome Research (Aug 2014), DOI: 10.1101/gr.176784.114.

Anja Thormann, Konrad L M Rudolph, Ina Koch. "TInA (T-Invariant Analysis): a tool box for exploring pathways in biochemical systems at steady state." Abstract book of CGB 2009 (Sep 2009).