# Dr Keiran Rowell | Curriculum Vitæ

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# **Employment**

### **Bioinformatician**

Australian Proteomics Analysis Facility

2022-present

Working with R code for mass spectrometry proteomics analysis. Refactoring an existing experiment-specific codebase to be more modular, allowing easy addition of new analysis functions. Mentoring a final-year PhD candidate in refactoring code, while learning how to clean-up terse R scripts. Retiring legacy systems.

### **Research Computing Support Officer**

Research Technology Services, UNSW

2021–2022

HPC and research computing support, including day-to-day system administration. Installation and troubleshooting of scientific software. HPC outreach and uptake, including weekly 'Hacky Hour' sessions.

### Postdoctoral Research Associate

School of Chemistry, USYD

2020-2021

Continuation of my PhD research. Multiconfigurational quantum chemistry calculations (CASSCF & CASPT2). Mentoring PhD candidates to submission and employment. Preparing research papers.

### **Casual Science Teacher**

Matrix Education

2016-2020

Student experience questionnaire 2-year average: 4.7/5

Top quality high school curriculum tuition. Classroom lessons, quiz marking, workbook development.

### Postgraduate Teaching Fellow & Tutor

School of Chemistry, UNSW

2016–2018, 2019–2020

1<sup>st</sup> year tutorials, mentoring, 3<sup>rd</sup> year physical chemistry resource development, exam marking, outreach.

### **Laboratory Demonstrator**

School of Chemistry, UNSW

2014-2015, 2020

Laboratory teaching, supervision, and marking, for  $1^{st}$ ,  $2^{nd}$  and  $3^{rd}$  year chemistry students.

### **Education**

### PhD - Computational Chemistry

University of New South Wales

2016-2020

Thesis Title: Structure-Activity Relationships for Carbonyl Photolysis

Supervisors: Prof. Scott Kable, Prof. Meredith Jordan

Assessment: A,A — All categories 'Outstanding' — Nominated for a Dean's Award

Thesis summary: Structure-Activity relationships for carbonyl UV photochemistry were comprehensively elucidated for 38 molecules across seven structural classes, for ten unique reaction mechanisms, on all relevant ground  $(S_0)$  and excited  $(S_1 \& T_1)$  states. Theoretical protocols for photolysis thresholds and excitation energies were extensively validated to the available spectroscopic and kinetic data for  $\pm$  10 kJ/mol accuracy.

### B.Sc. (Adv.) - Chemistry - Honours Class 1 - Angyal Prize

University of New South Wales

2011-2014

Dissertation Title: Computational Studies on the Basis of 'Neighbour Exclusion' in a Series of Diacridine and Di-(terpy)Pt(II) Thiol Bisintercalators: combined MD & FMO approaches Supervisors: A/Prof. Graham Ball, A/Prof. Larry Wakelin, Dr Donald Thomas

### **Programming Experience**

#### Python

- Developed resources for Python and Jupyter Notebooks instruction in 3<sup>rd</sup> year chemistry
  - Received 'Rising Star' award for introduction of Jupyter Notebooks into UNSW's 3<sup>rd</sup> year physical chemistry course. Format is being adopted in some 1<sup>st</sup> and 2<sup>nd</sup> year chemistry courses

- Wrote 'Intro to Python and Plotting' lessons for physical science students
- o Molecular rotational constants calculator; exploratory data analysis with pandas and matplotlib
- o Developed code to process, search, and plot 1000s of pair-wise energies between DNA fragments

### High Performance Computing.

- o 5+ years experience in Linux and high performance computing environments
  - Bash scripting, environment modules, basic sys. admin., schedulers (PBS Pro)
  - Command line tools: vim, grep, awk, sed, regex, xargs, chmod, tar, etc.
  - Attended regular ResTech seminars on computational research methods and data management

# Undergraduate coursework

- Bioinformatics (BINF3020, HD)
  - String alignment (DNA and protein sequences), Statistical modelling (HMM, Bayes net)
  - Collaborated to develop a web app for saccharomyces cerevisiae regulatory networks
- Computer science (COMP1917, DN)
  - Memory allocation, alogrithms and data structures.
  - Object-oriented programming, classes, inheritance from COMP1927 recordings

# Fortran

• Worked with 'Grow' in-house Fortran code for potential energy surface interpolation

# **Teaching and Mentorship**

### Postgraduate Teaching Fellowship

o 3 year tenure, 75 hours commitment per year, developed physical chemistry resources.

# University Subjects Taught

- $\circ$  3<sup>rd</sup> year physical chem demonstrator, coding facilitator satisfaction rose from **65% to 94%**
- o 2<sup>nd</sup> year physical chem laboratory demonstrator
- o 1st year chemistry tutor and laboratory demonstrator tutor feedback form average 4.6/5

### Academic Mentorship

- Mentored theoretical chemistry Honours students in performing quantum chemistry research
- o Mentored newer Teaching Fellows in delivering chemistry tutorials and pedagogical technique

# **Presentation Highlights**

**Group & 'Super-group' organiser**: Organised weekly group and monthly 'super-group' meetings; sourcing academic speakers, coordinating schedules. Ran computing & visualisation masterclasses

**APATCC 2019** (volunteer, poster)

USYD - Sydney Sep. 2019

Asia-Pacific Association of Theoretical and Computational Chemists

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**ISTCP-X** (poster)

UiT - Tromsø

10<sup>th</sup> Triennial Congress of the International Society of Theoretical Chemical Physics Jul. 2019

QUACCS 3.0 (talk)

ANU Campus - Kioloa

Computational chemistry workshops & student 'chalk 'n talk' presentations

Dec. 2018

Workshops attended: Global Optimisation, Stationary point search, Write your own HF & MP2, QM/MM

### ACCOMC 2018 (poster)

Atmospheric composition & modelling conference for Australasian region

CSIRO - Aspendale

Dec. 2018

2012

### **Awards**

Rising Star - PhD Casual Teacher School of Chemistry UNSW Awarded to two casual PhD tutors who have demonstrated educational excellence in a single year	2018
Poster presenter prize	
Association of Molecular Modellers of Australia Best Honours-level poster presenter	2015
Angyal prize	
School of Chemistry UNSW Top mark in Chemistry Honours	2014
Bosworth prize	
School of Chemistry UNSW	2013
Equal 1 <sup>st</sup> in third year Physical Chemistry	
Summer Vacation Research Scholarship	

UNSW Science competitive scholarships for undergraduates to gain early research experience

# **Computer Skills**

School of Chemistry UNSW

**Programming**: Python, Bash, C **Analysis**: Excel, Python, Pandas

**Documents**: Word, LATEX, Overleaf Figures: Matplotlib, Inkscape, GNU IMP

# **Publications** (refer to my Google Scholar page)

- 1. Rowell, K.; Kable, S.; Jordan, M., "An assessment of the tropospherically accessible photo-initiated ground state chemistry of carbonyls", doi:10.5194/acp-2021-424, [In Press]
- 2. Juan, C.Z.T; Syme, A-M.; Rowell, K.; et al., "Infrared Spectroscopy of Phosphorus-containing Molecules", Frontiers in Astronomy and Space Science, (2021), doi:10.3389/fspas.2021.639068
- **3. Rowell, K.**; Thomas, D.; Ball, G.; Wakelin, L., "Molecular Dynamic Simulations of Diacridine Binding to DNA: Indications that C6 Diacridine can Bisintercalate Spanning Two Base Pairs", *Biopolymers*, **(2020)**, doi:10.1002/bip.23409
- **4.** Rowell, K.; Kable, S; Jordan, M., "Substituent Effects on the Norrish Type I  $\alpha$ -bond Cleavage of Tropospheically Important Carbonyls", *The Journal of Physical Chemistry A*, **(2020)**, doi:10.1021/acs.jpca.9b05534
- **5.** Harrison, A.; Kharazmi, A.; Shaw, M.; Quinn, M.; Lee, K.; Klaas, N.; **Rowell, K.**; Jordan, M.; Kable, S., "Dynamics and Quantum Yields of  $H_2 + CH_2CO$  as a Primary Photolysis Channel in  $CH_3CHO$ ", *Physical Chemistry Chemical Physics*, **(2019)**, doi:10.1039/C8CP06412A

# Pre-prints.

**Rowell, K.**; Kable, S.; Jordan, M., "Predicting Carbonyl Excitation Energies Efficiently Using EOM-CC Trends", doi:10.26434/chemrxiv.12917369.v2

**Rowell, K.**; Kable, S.; Jordan, M., "Structural Causes of the Singlet/Triplet Preferences of Norrish Type II Reactions in Carbonyls", doi:10.26434/chemrxiv.12941702.v1