Christian S. Ahart

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Professional summary

Following the completion of my PhD at University College London with Prof. Jochen Blumberger I have joined the group of Dr. Clotilde Cucinotta at Imperial College London, developing methodologies to enable the dynamical modelling of electrochemical systems under applied potential.

Research interests

- · Charge transport
- Ab initio molecular dynamics
- Electronic structure of solids and interfaces
- Method development

Publications

- 1. Hannah C Nerl, **Christian S. Ahart**, Alberto Eljarrat, Christoph T Koch, Clotilde S Cucinotta, Milivoj Plodinec. Transitional surface Pt carbide formation during carbon nanotube growth. *submitted to Nat. Catal.*, 2023.
- 2. **Christian S. Ahart**, Kevin M. Rosso and Jochen Blumberger. Implementation and Validation of Constrained Density Functional Theory Forces in the CP2K Package. *J. Chem. Theory Comput.* 18, 4438–4446, 2022.
- 3. **Christian S. Ahart**, Kevin M. Rosso and Jochen Blumberger. Electron and Hole Mobilities in Bulk Hematite from Spin-Constrained Density Functional Theory. *J. Am. Chem. Soc.* 144, 4623–4632, 2022.
- 4. **Christian S. Ahart**, Jochen Blumberger and Kevin M. Rosso. Polaronic structure of excess electrons and holes for a series of bulk iron oxides. *Phys. Chem. Chem. Phys.* 22, 10699–10709, 2020.

Conferences and talks

- 1. Materials and Molecular Modelling Hub. UK, 2023. Poster presentation.
- 2. Psi-k. Switzerland, 2022. Poster presentation.
- 3. Supercomputer modelling of advanced materials at the Royal Society. UK, 2022. Poster presentation.
- 4. Research Group Seminar at Chicheley Hall. UK, 2022. Oral Presentation.
- 5. Computational Molecular Science. UK, 2019. Poster presentation.

Research Experience

2022-2024 Imperial College London, UK

Research Associate

- Developing methodologies to enable the dynamical modelling of electrochemical systems under applied potential.
- Supporting and supervising PhD and Master's students.
- Giving lectures and leading demonstrating sessions in computer labs.
- Collaborating with experimental scientists, contributing theoretical and computational expertise.

Education

2018 - 2022 University College London, UK

PhD Condensed Matter and Materials Physics Research group of Prof. Jochen Blumberger

Thesis: Charge transport in bulk hematite and at the hematite/water interface

The mobility for excess electrons and electron holes in bulk hematite was calculated using spin-constrained and gap-optimised hybrid density functional theory, with comparison to calculations of charge transport at the hematite/water interface.

2014 - 2018 University of Nottingham, UK

MSc. Chemistry and Molecular Physics (First class Honours)

Modules include:

- Scientific Computing
- Quantum Dynamics
- · Solids, Interfaces and Surfaces
- · Advanced Physical Chemistry

Master's project: Quantum mechanics of rotating electron nuclear spin systems

This project involved research into, and application of, theoretical and computational techniques to model nuclear magnetic resonance with dynamic nuclear polarisation.

2007 - 2014 William Howard School, Brampton, UK

A Levels: Mathematics (A), Physics (A), Chemistry (A), Biology AS (A)

GCSEs: 9 including Maths and English (A*-B)

Other Skills

- Programming: Fortran including MPI and OpenMP, Python, MATLAB.
- IT: Microsoft Office Suite, Adobe Creative Suite.

Interests

- Homebrew: member of the London Amateur Brewers, participate in homebrew competitions.
- eSports: captain of a 5-member team within the Nottingham Gaming Society competing in National tournaments.
- Rock climbing, badminton.

Reference contacts

Dr. Clotilde S. Cucinotta Imperial College London, UK c.cucinotta@imperial.ac.uk Relationship: Supervisor

Dr. Sergey Chulkov University of Lincoln, UK schulkov@lincoln.ac.uk Relationship: Research collaborator

Prof. Jochen Blumberger University College London, UK j.blumberger@ucl.ac.uk Relationship: PhD supervisor