Matthew Evans



energy storage materials • ab initio calculations crystal structure databases • open science & software

EDUCATION

2016- PhD Physics, Theory of Condensed Matter Group, University of Cambridge

2015–2016 MPhil Scientific Computing, University of Cambridge, Pass with distinction

2011–2015 MPhys Physics with Theoretical Physics, University of Manchester, First Class (Hons)

2021 Visiting Researcher

EXPERIENCE

with Prof Clare Grey (University of Cambridge)

 Data management, analysis software and open APIs for materials chemistry research. 2020- Researcher

with Prof Gian-Marco Rignanese (Université catholique de Louvain) Machine learning for small materials datasets (MODNet).

- High-throughput workflows for computational materials science.
- 2016–2020 PhD student: Crystal structure prediction for next-generation energy storage

with Dr Andrew Morris (University of Cambridge/University of Birmingham)

Computional materials discovery for conversion anodes for Li, Na and K-ion batteries

- Author of two open-source Python packages: database approaches for high-throughput
- ilustrado. - Active member of the OPTIMADE consortium for materials database interoperability and author of the optimade-python-tools package and odbx implementation.

calculations and materials design with matador and crystal structure prediction with

2019 Visiting Researcher

Department of Applied Physics, Aalto University

Enthought Inc., Cambridge

Received HPC-Europa funding to visit the group of Prof Adam Foster for 7 weeks, with

computational resources provided by the Finnish IT center for science (CSC). 2019 Scientific Software Developer (Intern)

Worked on the open source, Horizon 2020 FORCE project, adding functionality a workflow manager for multi-criteria optimisations. Helped develop Cython bindings for the ACADO toolkit.

2014, 2015 UG research: Interactions of quantised vortices in superfluid helium with Dr Paul Walmsley & Prof Andrei Golov (University of Manchester)

2014–2015 MPhys project: Electronic structure of defects in graphene superlattices with Prof Francisco Guinea (University of Manchester)

2013 UG research: Hard sphere packing of nanotube-encapsulated fullerenes with Dr Ho-Kei Chan & Prof Elena Besley (University of Nottingham)

COMPUTING

Languages Python, Fortran, Cython, C++

DFT **CASTEP**, Quantum Espresso, GPAW Packages NumPy, SciPy, matplotlib, scikit-

Exposure: **Daily**, Intermittent, *Occasional*.

learn, FastAPI, Tensorflow Tools git, vim, Docker Practices Test-driven development, CI

Databases MongoDB, SQL, Elasticsearch

2020 Co-chair of the Research Data Alliance (RDA) IG Materials Data, Infrastructure & Interoperability

(TEACHING + SERVICE)

2018– Reviewed manuscripts for npj. Computational Materials, J. Phys.: Cond. Mat., Scientific Reports

2021 Mentor at Acceleration Consortium Hackathon on Scientific Databases 2021 Developed and delivered a 2-day OPTIMADE tutorial for the NOMAD Virtual Tutorial Series.

2016–2020 Active member of TCM sysadmin team, Cavendish Laboratory

2019-2021 Demonstrator: Part II Computational Physics, Cavendish Laboratory

Demonstrated scientific Python to beginners in weekly labs (2019 only).

Conceptualised and delivered a tutorial on the basics of version control with Git (2019-2021).

Small group teaching, providing detailed feedback on assigned problems.

2016–2018 Supervisor: 2x Part IB Electromagnetism, Dynamics and Thermodynamics, Selwyn College

2016–2019 Demonstrator: 3x Part IB Introduction to Computing (C++), Cavendish Laboratory 2016–2019 Demonstrator: 4x at annual CASTEP workshop, University of Oxford

2017 Volunteer: 2nd Conference of Research Software Engineers, University of Manchester

2016–2017 Volunteer: Key Stage 2 Code Club, Ridgefield Primary School, Cambridge 2012–2015 Tutor: GCSE Maths & Key Stage 2 Programming for The Tutor Trust, Manchester

- Provided tuition to small groups and 'looked after children' across 15 schools. - Helped lead a successful pilot to teach primary school children programming using Scratch.

discovery for energy applications, M. L. Evans and A. J. Morris.

(AWARDS + HONOURS) 2021 PI for "Interoperable data management for fundamental battery research", BIG-MAP External

Stakeholder Initiative, total funding €150,000 (personal allocation €50,000). 2019 HPC-Europa 3 funding to visit Aalto University for 7 weeks and associated computing time.

Morris.

£12,000.

Meeting

2018 Tier-2 HPC Resource Allocation: PI on project awarded 2 MCPUh, Crystal structure prediction for next-generation solar absorbers, M. L. Evans, D. O. Scanlon and A. J. Morris.

HPC Midlands+ Substantial Project: awarded 1.3 MCPUh for High-throughput materials

2017 Tier-2 HPC Resource Allocation: Co-investigator on project awarded 4 MCPUh, Ab initio structure prediction for next-generation battery materials, B. Karasulu, M. L. Evans and A. J.

2015 Tesella Prize for Software, University of Manchester, for the most effective use of software in a

final year physics project. 2013, 2014 Undergraduate research bursary for two summers as an undergraduate, totalling £4200. 2011–2015 Means-tested and merit based scholarship to study at the University of Manchester, worth

2021 Invited panel discussions: International Materials Data: Joint Meeting and Metadata for Data Indexing and Discovery in Materials Science, Research Data Alliance (RDA) 18th Virtual Plenary

Invited talk: The OPTIMADE Ecosystem, DoE Battery Genome Initiative

CECAM Workshop, Open Databases Integration for Materials Design 2020

Invited panel discussion: Delivery platforms for open marketplaces, Research Data Alliance (RDA) 17th Virtual Plenary Meeting

prediction, EMRS Spring 2019, Nice, France

Seminar Series, University of Cambridge

SELECTED PRESENTATIONS

Invited talk: The OPTIMADE Specification, Research Data Alliance (RDA) 16th Virtual Plenary Meeting: Data Infrastructure for Collaborations in Materials Research Invited talk and workshop demonstration: odbx & OPTIMADE and optimade-python-tools,

2019 Contributed talk: Phosphorus anodes for potassium-ion batteries: insights from crystal structure

Invited talk: matador & OPTIMADE, CECAM Workshop, Open Databases Integration for Materials Design 2019, EPFL, Switzerland

prediction, SMARTER6 Conference, Ljubljana, Slovenia Invited talk: matador: databases and crystal structure prediction (slides), CECAM Workshop, Open Databases Integration for Materials Design 2018, EPFL, Switzerland

2018 Contributed talk: Sn-P anodes for potassium-ion batteries: insights from crystal structure

Poster Presentation: 13th RSC Conference in Materials Chemistry (poster), University of Liverpool

2016 Poster Presentation: SMARTER5 Conference, University of Bayreuth, Germany

2017 Invited talk: Crystal structure prediction for next-generation battery anodes (slides), Solid State

PUBLICATIONS Status: preprint, published. Underline indicates (joint) first authorship or lead theory authorship.

1361-648x/ac1280. arXiv:2102.02263.

optimade-python-tools: a Python library for serving and consuming materials data via OPTIMADE APIs. Journal of Open Source Software 6, 3458, (2021). DOI:10.21105/joss.03458.

8. Evans, M. L., Andersen, C. W., Dwaraknath, S., Scheidgen, M., Fekete, A. & Winston, D.

- 7. Andersen, C., Armiento, R., Blokhin, E., Conduit, G., Dwaraknath, S., Evans, M. L., Fekete, A., Gopakumar, A., Grazulis, S., Merkys, A., Mohamed, F., Oses, C., Pizzi, G., Rignanese, G., Scheidgen, M., Talirz, L., Toher, C., Winston, D., et al. OPTIMADE, an API for exchanging materials data. Sci Data 8, 217, (2021). DOI:10.1038/ s41597-021-00974-z. arXiv:2103.02068.
- Evans, M. L. & Morris, A. J. matador: a Python library for analysing, curating and performing high-throughput density-functional theory calculations. *Journal of Open Source Software* **5**, 2563, (2020). DOI:10.21105/joss. 02563.

6. De Breuck, P., **Evans, M. L.** & Rignanese, G. Robust model benchmarking and bias-imbalance in data-driven materials science: a case study on MODNet. J. Phys.: Condens. Matter 33, 404002, (2021). DOI:10.1088/

- 4. Harper, A. F., Evans, M. L. & Morris, A. J. Phases of Copper Phosphides from Computational Structure Searches. Chem. Mater. 32, (2020). DOI:10.1021/acs.chemmater.0c02054. arXiv:2005.05375. 3. Harper, A. F., Evans, M. L., Darby, J. P., Karasulu, B., Koçer, C. P., Nelson, J. R. & Morris, A. J. Ab initio structure
- prediction methods for battery materials: A review of recent computational efforts to predict the atomic level structure and bonding in materials for rechargeable batteries. Johnson Matthey Technology Review 64, 103-118, (2020). DOI:10/ggrmgf. 2. Marbella, L. E., Evans, M. L., Groh, M. F., Nelson, J., Griffith, K. J., Morris, A. J. & Grey, C. P. (De)Sodiation via Helical Phosphorus Intermediates in High Capacity Anodes for Sodium-ion Batteries. J. Am. Chem. Soc.
- **140**, 7994–8004, (2018). DOI:10.1021/jacs.8b04183. 1. Zhu, T., Evans, M. L., Brown, R. A., Walmsley, P. M. & Golov, A. I. Interactions between unidirectional quantized vortex rings. Phys. Rev. Fluids 1, 044502, (2016). DOI:10.1103/PhysRevFluids.1.044502. arXiv:1603. 04313.