# **Matthew Evans**



decentralized data management • open science & software materials discovery • ab initio calculations

#### RESEARCH INTERESTS

My background in computational materials science has left me with an overarching interest in the application of machine learning, open source software & infrastructure, and data management practices to accelerate and enhance scientific workflows for discovery in the chemical and materials sciences.

#### **EDUCATION**

2016-2023	PhD Physics, (submitted July 2023), 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)

## SELECTED EXPERIENCE

#### 2020— Research Assistant then BEWARE Research Fellow (2022 onwards)

Université catholique de Louvain and Matgenix, with Prof Gian-Marco Rignanese

- Co-creator and architect of datalab, open source data management software for sample tracking and characterisation, lab management, and machine learning, deployed at several labs internationally.
- High-throughput machine-learning accelerated workflows for materials discovery and design.
- Leading development of the OPTIMADE API specification and associated software.

## 2021 Visiting Researcher: Data management platforms for materials chemistry research University of Cambridge, with Prof Clare Grey FRS

- Developing bespoke data management platforms for materials chemistry and battery research.
- Supervising contributions from a full-time software developer and providing user training.

## 2024- Scientific Software Consultant and Director (part-time)

datalab industries ltd.

- Supporting the open source development of OPTIMADE and datalab via consultancy services.
- Customisation and deployment of dαtαlαb for industrial R&D and academic labs.

#### 2016–2020 PhD student: Crystal structure prediction for next-generation energy storage

University of Cambridge, with Dr Andrew Morris

- 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 calculations and materials design with matador and crystal structure prediction with ilustrado.

### COMPUTING

Languages **Python**, Javascript, Vue.js, Fortran, C++ Tools git, vim, Docker, Ansible, Terraform Practices **Test-driven** development, Expertise Web APIs, databases, machine learning, CI/CD, **Cloud Automation. HPC** high-throughput workflows

#### OTHER EXPERIENCE

- 2022 **Postdoctoral Researcher: Recommender systems for crystal structure search**Cambridge Crystallographic Data Centre, Cambridge, UK
- 2019 Visiting Researcher: Machine learning for materials discovery Aalto University, Finland, with Profs Adam Foster & Patrick Rinke
- 2019 Scientific Software Developer (Intern): Multi-objective optimisation Enthought Inc., Cambridge
- 2014, 2015 **UG research: Interactions of quantised vortices in superfluid helium**University of Manchester, with Dr Paul Walmsley & Prof Andrei Golov
  - 2013 **UG research: Hard sphere packing of nanotube-encapsulated fullerenes**University of Nottingham, with Dr Ho-Kei Chan & Prof Elena Besley

## SELECTED (AWARDS + HONOURS)

- 2022 BEWARE2 Fellowship from the Wallonia-Brussels Federation (approx. €300,000).
- 2021 PI for "Interoperable data management for fundamental battery research", BIG-MAP External Stakeholder Initiative (approx. €150,000).

## SELECTED (TEACHING + SERVICE)

- 2018– Reviewed manuscripts and data for JOSS (x6), Digital Discovery (x5), J. Phys.: Cond. Mat. (x4), Mach. Learn.: Sci. Technol. (x2), npj. Comp. Mater. (x1), Sci Data (x1), Sci. Rep. (x1)
- 2022–2024 Proposed and co-lead a MaRDA working group on metadata extractors for materials science.
- 2022–2024 Initiator and organiser of the CECAM Workshop series Machine-actionable Data Interoperability for Chemical Sciences (MADICES, February 2022 and April 2024)
- 2019-2021 Demonstrator: Part II Computational Physics, 3x Part IB Intro to Computing, Cavendish Laboratory
- 2016–2018 Supervisor: 2x Part IB Electromagnetism, Dynamics and Thermodynamics, Selwyn College
- 2012–2015 Tutor: GCSE Maths & Key Stage 2 Programming for The Tutor Trust, Manchester

#### SELECTED RECENT PRESENTATIONS

- 2025 Invited seminar: Decentralized materials research data management, curation and dissemination for accelerated discovery, Computational Chemistry Seminar Series, University of Warwick
- 2024 Invited talk: Decentralized materials research data management, curation and dissemination for accelerated discovery, Democratizing AI in Materials Science A Pathway to Broaden the Impact of Materials Research, MRS Fall Meeting, Boston, USA

Contributed talk/paper: Optical materials discovery and design with federated databases and machine learning, Faraday Discussions, University of Oxford, United Kingdom.

Invited talk: Federated, interoperable databases for accelerated materials discovery and design, CECAM Flagship Workshop on MLIPs and Accessible Databases, Grenoble, France.

Invited talk: Open Databases Integration for Materials Design at the CECAM Flagship Workshop for FAIR and TRUE Soft Matter Simulations, Max Planck Institute for Polymer Research, Germany.

Invited talk: Open Databases Integration for Materials Design at the Actively Learning Materials Science (AL4MS2023) workshop, Aalto University, Finland.

## PUBLICATIONS

Underline indicates (joint) first authorship (reordered where appropriate). Full list with OA links available online (https://ml-evs.science/papers, ORCiD, Google Scholar).

- 18. Mroz, A. M., **Evans, M. L.**, Basford, A. R., et al. Cross-disciplinary perspectives on the potential for artificial intelligence across chemistry (submitted). 2025.
- 17. **Evans, M. L.**, Trinquet, V., *et al.* Optical materials discovery and design with federated databases and machine learning. *Faraday Discussions* **256**, 459–482, (2025). DOI:10.1039/D4FD00092G.
- 16. Zimmermann, Y., Bazgir, A., **Evans, M. L.**, et al. Reflections from the 2024 Large Language Model (LLM) Hackathon for Applications in Materials Science and Chemistry (submitted). 2024. DOI:10.48550/arXiv.2411.15221.
- 15. **Evans, M. L.**, Rignanese, G.-M., Elbert, D. & Kraus, P. Metadata, automation, and registries for extractor interoperability in the chemical and materials sciences (submitted). 2024. DOI:10.48550/arXiv.2410.18839.
- 14. **Evans, M. L.**, Bergsma, J., Merkys, A., Andersen, C. W., *et al.* Developments and applications of the OPTIMADE API for materials data exchange and discovery. *Digital Discovery* **3**, (2024). DOI:10.1039/D4DD00039K.
- 13. Rosen, A. S., Gallant, M., George, J., et al. Jobflow: Computational Workflows Made Simple. *Journal of Open Source Software* **9**, 5995, (2024) ISSN: 2475-9066. DOI:10.21105/joss.05995. (2024).
- 12. Wang, Z., Gong, Y., **Evans, M. L.**, *et αl.* Machine learning-accelerated discovery of A<sub>2</sub>BC<sub>2</sub> ternary electrides with diverse anionic electron densities. *J. Amer. Chem. Soc.* **145**, 26412–26424, (2023). DOI:10.1021/jacs.3c10538.
- 11. Lertkiattrakul, M., **Evans, M. L.** & Cliffe, M. J. PASCal Python: A Principal Axis Strain Calculator. *Journal of Open Source Software* **8,** 5556, (2023). DOI:10.21105/joss.05556.
- 10. Jablonka, K. M., Ai, Q., Al-Feghali, A., et al. 14 Examples of How LLMs Can Transform Materials Science and Chemistry: A Reflection on a Large Language Model Hackathon. *Digital Discovery*, (2023). DOI:10/gswbnx.
- 9. Ells, A. W., **Evans, M. L.**, Groh, M., Morris, A. J. & Marbella, L. E. Phase transformations and phase segregation during potassiation of Sn<sub>x</sub>P<sub>ν</sub> anodes. *Chemistry of Materials*, (2022). DOI:10/h69d.
- 8. Evans, M. L., Andersen, C. W., et αl. 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/gn3w9f.
- 7. **Evans, M. L.**, Andersen, C. W., Armiento, R., *et al.* OPTIMADE, an API for exchanging materials data. *Scientific Data* 8, 217, (2021). DOI:10/gmnrxj.
- 6. Breuck, P.-P. D., **Evans, M. L.** & Rignanese, G.-M. Robust model benchmarking and bias-imbalance in data-driven materials science: a case study on MODNet. *J. Phys.: Cond. Mat.* **33**, 404002, (2021). DOI:10/gpw93d.
- 5. **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/gmf4mv.
- 4. Harper, A. F., **Evans, M. L.** & Morris, A. J. Computational Investigation of Copper Phosphides as Conversion Anodes for Lithium-Ion Batteries. *Chemistry of Materials*, (2020). DOI:10/gg5sx3.
- 3. Harper, A. F., **Evans, M. L.**, Darby, J. P., *et al.* Ab initio Structure Prediction Methods for Battery Materials. *Johnson Matthey Technology Review* **64**, 103–118, (2020). DOI:10/ggrmgf.
- 2. Marbella, L. E., **Evans, M. L.**, Groh, M. F., *et αl.* Sodiation and Desodiation via Helical Phosphorus Intermediates in High-Capacity Anodes for Sodium-Ion Batteries. *J. Amer. Chem. Soc.* **140**, 7994–8004, (2018). DOI:10/gdq6h4.
- 1. Zhu, T., **Evans, M. L.**, Brown, R. A., Walmsley, P. M. & Golov, A. I. Interactions between unidirectional quantized vortex rings. *Physical Review Fluids* **1**, 044502, (2016). DOI:10/gf2529.