Felix-Cosmin Mocanu

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• fcmocanu

ab-initio molecular dynamics • phase-change memory materials • thermoelectric materials • machine-learned interatomic potentials • structure searching

EDUCATION

2015- PhD Chemistry, Chemical Physics Group, University of Cambridge, Expected submission 2018.

2014–2015 MPhil Scientific Computing, University of Cambridge, Distinction.

2010–2014 MSci Natural Sciences, Department of Chemistry, University of Cambridge, Upper Second Class (Hons).

RESEARCH

PhD First and second principles simulations of Ge-Sb-Te phase-change memory materials

with Prof. Stephen R. Elliott and Prof. Gábor Csányi (University of Cambridge)

Monte-Carlo sampling techniques for the fitting of a transferable Ge-Sb-Te machine-learned interatomic potential. Non-equilibrium molecular dynamics and stochastic-boundary conditions methods for calculating thermal conductivity. Structure searching and prediction of novel compositions for phase-change memory and thermoelectric materials. PhD research undertaken as a member of the EPSRC CDT for Computational Methods in Materials Science.

MPhil Towards high-throughput simulation of Ge-Sb-Te phase change materials

with Prof. Stephen R. Elliott and Prof. Gábor Csányi (University of Cambridge)

Worked on a machine-learned Gaussian Approximation Potential for the $\rm Ge_2Sb_2Te_5$ phase-change memory material trained from *ab-initio* molecular dynamics trajectories and its application in large-scale and high-throughput simulations.

MSci Ab-inito Molecular Dynamics of Phase Change Materials - NAND gate simulation

with Dr. James Dixon and Prof. Stephen R. Elliott (University of Cambridge)

Worked on the *ab-initio* molecular-dynamics simulations of the atomistic effects of thermal pulse programming in the $Ge_2Sb_2Te_5$ phase-change memory material for applications in-memory logic and neuromorphic ("brain-like") computation.

UG Synthesis and characterisation of functional curcubituril-azobenzene host-guest assemblies summer student with Dr. Jesús del Barrio Lasheras and Prof. Oren Scherman (University of Cambridge)

INTERSHIPS

2012 Rotatives Engineering Intern

Cambridge Materials Placements for Undergraduates in Summer (CaMPUS) from the University of Cambridge Materials Science Department with Dr.-Ing. Hartmut Schlums (Rolls-Royce Deutschland) studying turbine blade lifetime in jet engines.

TEACHING

- 2016 Organiser of fortnightly simulation tutorials (Chemical Physics Group).
- 2017 Demonstrating Part IB Computational Practicals (Department of Chemistry).
- 2016 Demonstrating Part IA Labs (Department of Chemistry).
- 2016 Supervisor of Part II Theoretical Techniques course in Chemistry (Churchill College).
- 2015 Demonstrator IA C++ Computer Practicals (Engineering Laboratory).
- 2014–2016 Organiser of simulation sub-group meetings (Chemical Physics Group).

SIMULATION AND COMPUTING

DFT VASP, CP2K, CASTEP, GPAW MD LAMMPS, QUIP

Platforms **Linux** Packages **ASE**, quippy, pymatgen

Languages **Python**, Julia, Fortran, C++ Databases **MongoDB**, *SQL*

System administrator for the Elliott group Hathor computer cluster.

Tier-2 Resource Allocation (Project T2-CS020 "Thermoelectric Properties of Disordered Materials"): awarded 4 million CPU hours on the CSD3 supercomputer with Yuchen Hu and Prof. Stephen R. Elliott.

PUBLICATIONS

Status: in preparation, submitted, preprint, published.

- Konstantinou, K., Lee, T. H., Mocanu, Felix C. & Elliott, S. R. Origin of radiation tolerance in amorphous Ge₂Sb₂Te₅ phase-change random-access memory material. *Proceedings of the National Academy of Sciences of the United States of America* 115, 5353–5358. ISSN: 1091-6490 (May 2018).
- Mavračić, J., Mocanu, Felix C., Deringer, V. L., Csányi, G. & Elliott, S. R. Similarity Between Amorphous and Crystalline Phases: The Case of TiO₂. The Journal of Physical Chemistry Letters 9, 2985–2990. ISSN: 1948-7185 (June 2018).

CONFERENCE PRESENTATIONS

International Symposium on Doped Amorphous Chalcogenides and Devices

2018 Computer simulations of phase-change materials, Poster Presentation, Grasmere, United Kingdom - Runner-up Prize

European Phase-Change and Ovonics Symposium

2017 A machine-learned interatomic potential for Ge—Sb—Te phase-change materials, Poster Presentation, RWTH Aachen, Germany

Psi-k Conference on Electronic Structure

2015 A Gaussian Approximation Potential (GAP) for the Ge₂Sb₂Te₅ phase change material, Poster Presentation, San Sebastián, Spain

REFEREES

Prof. Stephen R. Elliott, University of Cambridge; sre1@cam.ac.uk Prof Gábor Csányi, University of Cambridge; gc121@cam.ac.uk