

PHD CANDIDATE - COMPUTATIONAL MATERIALS SCIENCE - EINDHOVEN UNIVERSITY OF TECHNOLOGY, THE NETHERLANDS

■ m.c.w.m.pols@tue.nl | **G** Google Scholar | **6** 0000-0002-1068-9599 | **1** mikepols | 5th of July, 1996

Leveraging extensive experience in training reactive and machine-learning force fields. I have investigated ion, phase, and lattice dynamics of halide perovskites. My goal is to advance machine-learning models beyond energy and force predictions to also cover charges, electronic structure, and spin properties, enabling applications in catalysis, optoelectronics, and spintronics.



Research Experience

Eindhoven University of Technology (TU/e)

PHD CANDIDATE IN MATERIALS SIMULATION & MODELLING (MSM)

Mar. 2021 - Feb. 2025

Computational modelling of halide perovskites using atomistic simulations, with a focus on their ion, phase, and lattice dynamics.

MSc in Applied Physics (Cum Laude), MSc in Chemical Engineering (Cum Laude)

May. 2018 - Feb. 2021

- Grade Point Average (GPA): 9.4/10
- Master thesis: A Reactive Force Field for Large Scale Simulations of Metal Halide Perovskites (9.5/10) [Pub. 7]
- Industrial internship at Software for Chemistry & Materials (SCM) on infrared and Raman spectroscopy in solids (9.5/10)
- Research assistant at TU/e investigating atom-thin 2D materials for perovskite solar cells using computational modelling [Pub. 18]

BSc in Applied Physics (Cum Laude), BSc in Chemical Engineering (Cum Laude)

Sep. 2014 - Apr. 2018

- Grade Point Average (GPA): 9.3/10
- Bachelor thesis: Triplet-triplet annihilation of phosphorescent host-guest materials for OLEDs (9.5/10)
- Research assistant at TU/e investigating triplet-triplet annihilation in phosphorescent host-guest systems for OLEDs [Pub. 19]

Teaching & Mentoring

Courses

Advanced Materials Modelling using Multiscale Methods (3MQ110)

- Design of the course materials, including programming exercises and introductory density functional theory simulations
- Teaching assistant for hands-on programming exercises for density functional theory
- General introduction to use of high-performance computing (HPC) systems
- Supervision of student projects using classical and quantum simulations of halide perovskites

COMPUTATIONAL MATERIALS SCIENCE (3MN200)

2019 - 2020

- Teaching assistant for hands-on density functional theory calculations with VASP
- General introduction to use of high-performance computing (HPC) systems

Student Mentoring

MENTORED A VARIETY OF STUDENTS AT VARIOUS LEVELS DURING THEIR THESIS AND INTERNSHIP PROJECTS:

- Master students (5) with projects covering ion migration, surface and grain boundary stability, and crystallization of halide perovskites
- · Bachelor student (1) with project focused at exploring the compositional effects on structural chirality in chiral perovskites

Honors & Awards

Jun. 2024 Participant of 73rd Lindau Meeting , Lindau Nobel Laureate Meetings	Lindau, Germany
Feb. 2023 HPC SYSTEMs Inc. Poster Award , APATCC-10	Quy Nhon, Vietnam
Nov. 2021 Best Poster Prize , ACOS conference 2021	Eindhoven, NL
Sep. 2021 Nominated: Best Student Assistant (Applied Physics, TU/e), STOOR Education Awards	Eindhoven, NL
Jun. 2020 Participant of 70th Lindau Meeting, Lindau Nobel Laureate Meetings	Lindau, Germany
May. 2018 ALSP Scholarship (top 1%), Eindhoven University of Technology (TU/e)	Eindhoven, NL
Jul. 2014 Bronze Medal (128th out of 291), 46th International Chemistry Olympiad	Hanoi, Vietnam
Jun. 2014 Gold Medal (4th out of 4090), Dutch National Chemistry Olympiad 2014	Amsterdam, NL



Languages Dutch: native, English: full professional proficiency, German: elementary

Programming Python, Matlab, Mathematica, Fortran, LaTeX **Software** VASP, AMS, Phonopy, OVITO, VESTA, LAMMPS

Publications

First-author publications

- 1. M. Pols, E. Boom, G. Brocks, S. Calero, and S. Tao, 'Lattice Dynamics of Chiral 2D Perovskites: Pb vs. Sn', In preparation (2024).
- 2. M. Pols, G. Brocks, S. Calero, and S. Tao, 'Chiral Phonons in 2D Halide Perovskites', In preparation (2024), DOI: arXiv:2411.17225.
- 3. M. Pols, G. Brocks, S. Calero, and S. Tao, 'Temperature-Dependent Chirality in Halide Perovskites', *J. Phys. Chem. Lett.*, 15, 8057-8064 (2024), DOI: 10.1021/acs.jpclett.4c01629.
- 4. **M. Pols**, A.C.T. van Duin, S. Calero, and S. Tao, 'Mixing I and Br in Inorganic Perovskites: Atomistic Insights from Reactive Molecular Dynamics Simulations', *J. Phys. Chem. C*, 128, 4111-4118 (2024), DOI: 10.1021/acs.jpcc.4c00563.
- 5. **M. Pols**, V. Brouwers, S. Calero, and S. Tao, 'How Fast Do Defects Migrate in Halide Perovskites: Insights From On-the-Fly Machine-Learned Force Fields', *Chem. Commun.*, 59, 4660-4663 (2023), DOI: 10.1039/D3CC00953J.
- 6. **M. Pols**, T. Hilpert, I. Filot, A.C.T. van Duin, S. Calero, and S. Tao, 'What Happens at Surfaces and Grain Boundaries of Halide Perovskites: Insights from Reactive Molecular Dynamics Simulations of CsPbl₃', *ACS Appl. Mater. & Interfaces*, 14, 40841-40850 (2022), DOI: 10.1021/acsami.2c09239.
- 7. **M. Pols**, J.M. Vicent-Luna, I. Filot, A.C.T. van Duin, and S. Tao, 'Atomistic Insights Into the Degradation of Inorganic Halide Perovskite CsPbl₃: A Reactive Force Field Molecular Dynamics Study', *J. Phys. Chem. Lett.*, 12, 5519-5525 (2021), DOI: 10.1021/acs.jpclett. 1c01192.

Collaborative publications

- 8. R. Nurdillayeva, R. Moral, **M. Pols**, D.-K. Lee, M.V. Altoe, C. Schwartz, S. Tao, and C.M. Sutter-Fella, 'Humidity Disrupts Structural and Chiroptical Properties of Chiral 2D Perovskites', Submitted (2024).
- 9. V. Tyagi, **M. Pols**, G. Brocks, and S. Tao, 'Tracing Ion Migration in Halide Perovskites with Machine Learned Force Fields', Under review (2024), DOI: arXiv:2409.16051.
- 10. S. Liu, **M. Pols**, X. Huang, Z. Huang, Z. Guo, L. Li, L. Wang, H. Dong, S. Tao, H. Zhou, L.-D. Sun, and C.-H. Yan, 'In-situ Polymerization Strategy for Blue Perovskite Light-Emitting Diodes with Ultra-high Efficiency Exceeding 20%', Under review (2024).
- 11. A.J. Chacón-García, H.G. Baldovi, **M. Pols**, S. Tao, S. Calero, S. Navalón, I.I. Vitorica-Yrezabal, A. Rodríguez-Diéguez, H. García, P. Horcajada, and Y. Pérez, 'Improving the Water Resistance of Bi-Based Perovskite-Inspired Materials for Vapor-Phase Photocatalytic Overall Water Splitting', *Solar RRL*, 8, 2400250 (2024), DOI: 10.1002/solr.202400250.
- 12. J. Liu, C. Zhu, **M. Pols**, Z. Zhang, F. Hu, L. Wang, C. Zhang, Z. Liu, S. Tao, M. Xiao, and X. Wang, 'Discrete Elemental Distributions Inside a Single Mixed-Halide Perovskite Nanocrystal for the Self-Assembly of Multiple Quantum-Light Sources', *Nano Lett.*, 23, 10089-10096 (2023), DOI: 10.1021/acs.nanolett.3c03761.
- 13. A.S. Mirza, **M. Pols**, W. Soltanpoor, S. Tao, G. Brocks, and M. Morales-Masis, 'The Role of Sulfur in Sulfur Doped Copper(I) Iodide p-type Transparent Conductors', *Matter*, 6, 1-15 (2023), DOI: 10.1016/j.matt.2023.10.003.
- 14. Z. Qin, **M. Pols**, M. Qin, J. Zhang, H. Yan, S. Tao, and X. Lu, 'Over-18%-Efficiency Quasi-2D Ruddlesden-Popper Pb-Sn Mixed Perovskite Solar Cells by Compositional Engineering', *ACS Energy Lett.*, 8, 3188-3195 (2023), DOI: 10.1021/acsenergylett.3c00853.
- 15. S. Raaijmakers, **M. Pols**, J.M. Vicent-Luna, and S. Tao, 'Refined GFN1-xTB Parameters for Engineering Phase-Stable CsPbX₃ Perovskites', *J. Phys. Chem. C*, 126, 9587-9596 (2022), DOI: 10.1021/acs.jpcc.2c02412.
- 16. C. Onwudinanti, **M. Pols**, G. Brocks, V. Koelman, A.C.T. van Duin, T. Morgan, and S. Tao, 'A ReaxFF Molecular Dynamics Study of Hydrogen Diffusion in Ruthenium–The Role of Grain Boundaries', *J. Phys. Chem. C*, 126, 5950-5959 (2022), DOI: 10.1021/acs.jpcc. 1c08776.
- 17. H. Xie, Z. Wang, Z. Chen, C. Pereyra, **M. Pols**, K. Gałkowski, M. Anaya, S. Fu, X. Jia, P. Tang, D.J. Kubicki, A. Agarwalla, H.-S. Kim, D. Prochowicz, X. Borrisé, M. Bonn, C. Bao, X. Sun, S.M. Zakeeruddin, L. Emsley, J. Arbiol, F. Gao, F. Fu, H.I. Wang, K.-J. Tielrooij, S. Stranks, S. Tao, M. Grätzel, A. Hagfeldt, and M. Lira-Cantu, 'Decoupling the Effects of Defects on Efficiency and Stability Through Phosphonates in Stable Halide Perovskite Solar Cells', *Joule*, 5, 1246-1266 (2021), DOI: 10.1016/j.joule.2021.04.003.
- 18. Y. Sun, Y. Yin, **M. Pols**, J. Zhong, Z. Huang, B. Liu, J. Liu, W. Wang, H. Xie, G. Zhan, Z. Zhou, W. Zhang, P. Wang, C. Zha, X. Jiang, Y. Ruan, C. Zhu, G. Brocks, X. Wang, L. Wang, J. Wang, S. Tao, and W. Huang, 'Engineering the Phases and Heterostructures of Ultrathin Hybrid Perovskite Nanosheets', *Adv. Mater.*, 32, 2002392 (2020), DOI: 10.1002/adma.202002392.

19. A. Ligthart, X. de Vries, L. Zhang, M. Pols, P.A. Bobbert, H. van Eersel, and R. Coehoorn, 'Effect of Triplet Confinement on Triplet-Triplet Annihilation in Organic Phosphorescent Host-Guest Systems', Adv. Func. Mater., 28, 1804618 (2018), DOI: 10.1002/adfm.2018 04618.

Other publications

20. M. Pols, 'Metal Halide Perovskites: Shimmering But Shaky', N! (TU/e Applied Physics Faculty Magazine), 57, 20-23 (2024).

Presentations

COMPUTATIONAL METHODS

NWO CHAINS 2024 TALK: STRUCTURAL AND DYNAMIC CHIRALITY IN 2D HALIDE PEROVSKITES Dec. 2024

AND DEGRADATION REACTIONS INDUCED BY THERMOCHEMICAL STRESS

Elementary School Tour at Applied Physics @ TU/e LECTURE: WHAT DOES A COMPUTATIONAL SCIENTIST DO AT TU/E? Jun. 2024

KNCV-CTC 2024

TALK: STABILITY OF METAL HALIDE PEROVSKITES: INSIGHTS FROM MULTISCALE REACTIVE MOLECULAR DYNAMICS Apr. 2024 SIMILI ATIONS

NWO Physics 2024 TALK: STRUCTURAL DESCRIPTORS FOR CHIRAL HALIDE PEROVSKITES AT FINITE TEMPERATURES Jan. 2024

IUPAC | CHAINS 2023 TALK: REACTIVE MOLECULAR DYNAMICS SIMULATIONS OF METAL HALIDE PEROVSKITES Aug. 2023

Elementary School Tour at Applied Physics @ TU/e LECTURE: WHAT DOES A COMPUTATIONAL SCIENTIST DO AT TU/E? Jun. 2023

CCER Seminar (3rd Year) TALK: MULTISCALE MODELLING OF DEFECTS IN HALIDE PEROVSKITES: INSIGHTS FROM A VARIETY OF Apr. 2023

Asia Pacific Conference of Theoretical and Computational Chemistry (APATCC-10) TALK: REACTIVE MOLECULAR DYNAMICS SIMULATIONS OF DEFECT-INDUCED DEGRADATION IN HALIDE PEROVSKITES: Mar. 2023 INSIGHTS FROM MACHINE LEARNED FORCE FIELDS AND REAXFF

2022 MRS Fall Meeting Talk: Multiscale Modelling of Defects in Halide Perovskites — Implications for the Phase Stability Nov. 2022

Computational Methods in Nanothermodynamics TALK: REACTIVE MOLECULAR DYNAMICS SIMULATIONS OF METAL HALIDE PEROVSKITES Apr. 2022

Green Week Lunch Lecture

Talk: What can I do as a physicist in the field of sustainability within Materials Simulations & Mar. 2022 MODELLING (MSM)?

Han-sur-Lesse Winterschool 2021 LECTURE: A REACTIVE FORCE FIELD FOR LARGE SCALE SIMULATIONS OF METAL HALIDE PEROVSKITES Dec. 2021

CCER Seminar (First year) TALK: A REACTIVE FORCE FIELD FOR LARGE SCALE SIMULATIONS OF METAL HALIDE PEROVSKITES Apr. 2021

Posters

DuComS Day 2023 POSTER: ON-THE-FLY MACHINE LEARNED FORCE FIELDS FOR HALIDE PEROVSKITES: HOW FAST DO DEFECTS MOVE? Nov. 2023

NWO Physics 2023 POSTER: ON-THE-FLY MACHINE LEARNED FORCE FIELDS FOR HALIDE PEROVSKITES: HOW FAST DO DEFECTS MOVE? Apr. 2023

Asia Pacific Conference of Theoretical and Computational Chemistry (APATCC-10) POSTER: WHAT HAPPENS AT SURFACES AND GRAIN BOUNDARIES OF HALIDE PEROVSKITES: INSIGHTS FROM Feb. 2023 REACTIVE MOLECULAR DYNAMICS SIMULATIONS OF CSPBI₃ (HPC SYSTEMS INC. POSTER AWARD)

Next-Generation V+ PV Materials 2022 POSTER: ATOMISTIC INSIGHTS INTO THE STABILITY OF INORGANIC CSPBI₃ FROM REACTIVE MOLECULAR DYNAMICS:

Jul. 2022 THE IMPACT OF DEFECTS, SURFACES AND GRAIN BOUNDARIES

Physics@Veldhoven 2022 POSTER: ATOMISTIC INSIGHTS INTO DEGRADATION PATHWAYS OF INORGANIC METAL HALIDE PEROVSKITES Jan. 2022

Applied Computational Science (ACOS) conference 2021 POSTER: ATOMISTIC INSIGHTS INTO THE DEGRADATION OF INORGANIC HALIDE PEROVSKITE CSPBI₃ (BEST POSTER PRIZE) 2020 Virtual MRS Spring/Fall Meeting & Exhibit POSTER: A REACTIVE FORCE FIELD FOR LARGE SCALE SIMULATIONS OF METAL HALIDE PEROVSKITES Online POSTER: A REACTIVE FORCE FIELD FOR LARGE SCALE SIMULATIONS OF METAL HALIDE PEROVSKITES Online POSTER: A REACTIVE FORCE FIELD FOR LARGE SCALE SIMULATIONS OF METAL HALIDE PEROVSKITES Sep. 2020