# Jan Janssen

Max-Planck-Institut für Eisenforschung

Max-Planck-Str. 1, 40237 Düsseldorf, Germany

## PROFESSIONAL APPOINTMENTS

2023 – present Group leader for Materials Informatics - Computational Materials Design

Research topic: machine learning for atomistic simulation Max-Planck-Institut für Eisenforschung, Düsseldorf, Germany

2022 – 2023 Director's Postdoctoral Fellowship - Theoretical Division (T-1)

Research topic: long-timescale simulations of materials with ab initio accuracy

Los Alamos National Laboratory, Los Alamos, NM, USA

2023 **Invited Postdoctoral Fellow** - Institute for Pure and Applied Mathematics (IPAM)

Research topic: new mathematics for the exascale: applications to materials science

Email: janssen@mpie.de

Website: jan-janssen.com

University of California, Los Angeles, CA, USA

2021 – 2022 Postdoctoral Research Associate - Theoretical Division (T-1)

Research topic: exascale atomistic capability for accuracy, length, and time

Los Alamos National Laboratory, Los Alamos, NM, USA

2017 Invited Fellow - Institute for Pure and Applied Mathematics (IPAM)

Research topic: complex high-dimensional energy landscapes

University of California, Los Angeles, CA, USA

2015 – 2021 PhD Candidate - Computational Materials Design

Research topic: computational phase studies and ab initio thermodynamics

Max-Planck-Institut für Eisenforschung, Düsseldorf, Germany

## **EDUCATION**

2015 – 2021 PhD in Theoretical Physics - Paderborn University, Germany

 $\bullet$  Thesis: pyiron - an integrated development environment for  $ab\ initio$  thermodynamics

• Advisor: Prof. Jörg Neugebauer

• Grade: summa cum laude

2009 – 2015 Advanced Degree in Physics - Technical University of Kaiserslautern, Germany

• Thesis: carbon in  $\alpha$ -iron-grainboundaries - an atomistic study of elastic properties

• Advisor: Prof. Herbert Urbassek

• Grade: thesis 1.0 (excellent) - total 1.6 (good)

## **FUNDED PROPOSALS**

2023 **Principal Investigator** of "workflows for machine learned interatomic potentials" as part of an internal call for funding for Postdoctoral Fellows at Los Alamos National Laboratory

• Computing Hardware Funding: \$13k

Co-Principal Investigator of "uncertainty propagation for multi-fidelity machine learned interatomic potentials" as part of the summer school for applied machine learning at Los Alamos National Laboratory

• Funding: \$25k

2022 **Co-Principal Investigator** of "development and deployment of a fully autonomous in *silico* processing and materials discovery platform" in collaboration with the department of Mechanical Engineering at Texas A&M University

• Funding: \$176k

Co-Principal Investigator of "transferability of interatomic machine learning potentials" as part of the summer school for applied machine learning at Los Alamos National Laboratory

• Funding: \$25k

**Principal Investigator** of "helping users build workflows with ontological typing" project funded by a NumFOCUS small development grant to hire an independent contractor

• Funding: \$6k

# **AWARDS & HONORS**

| 2023 | Invited postdoctoral fellow at the Institute for Pure and Applied Mathematics (IPAM)   |
|------|--|
| 2022 | Los Alamos National Laboratory (LANL) director's postdoctoral fellowship   |
| 2021 | PhD with highest distinction - summa cum laude   |
| 2019 | Runner-up for the Heinz Billing Award - a national biennial award for outstanding contributions to computational science by younger scientist without tenure |
| 2017 | Invited fellow at the Institute for Pure and Applied Mathematics (IPAM)  |
| 2015 | Scholarship of the Technical University of Kaiserslautern  |

# **PUBLICATIONS**

2021

Google Scholar, 08/23: Citations: 119

M.G. Taylor, D.J. Burrill, **J. Janssen**, E. Batista, D. Perez, and P. Yang. Architector: high-throughput cross-periodic table 3D complex builder. *Nature Communications*. doi:10.1038/s41467-023-38169-2. Citations: 0

A. Rohskopf, C. Sievers, N. Lubbers, M. A. Cusentino, J. Goff, **J. Janssen**, M. McCarthy, D. Montes de Oca Zapiain, S. Nikolov, K. Sargsyan, E. Sikorski, L. Williams, D. Sema, A. P. Thompson, and M. A. Wood FitSNAP: Atomistic machine learning in LAMMPS. *Journal of Open Source Software*. doi:10.21105/joss.05118. Citations: 3

• Code: FitSNAP/FitSNAP

L.F. Zhu, **J. Janssen**, S. Ishibashi, F. Körmann, B. Grabowski and J. Neugebauer. A fully automated approach to calculate the melting temperature of elemental crystals. *Computational Materials Science*. doi:10.1016/j.commatsci.2020.110065. Citations: 17

• Code: pyiron/pyiron meltingpoint

2020

T.D. Swinburne, J. Janssen, M. Todorova, G. Simpson, P. Plechac, M. Luskin and J. Neugebauer. Anharmonic free energy of lattice vibrations in fcc crystals from a mean field bond. Physical Review B. doi:10.1103/PhysRevB.102.100101. Citations: 4

• Code: O tomswinburne/BLaSA

2019

- J. Janssen, S. Surendralal, Y. Lysogorskiy, M. Todorova, T. Hickel, R. Drautz and J. Neugebauer. pviron: an integrated development environment for computational materials science. Computational Materials Science. doi:10.1016/j.commatsci.2018.07.043. Citations: 66
- Code: pyiron
- Y. Lysogorskiy, T. Hammerschmidt, J. Janssen, J. Neugebauer and R. Drautz. Transferability of interatomic potentials for molybdenum and silicon. and Simulation in Materials Science and Engineering. doi:10.1088/1361-651X/aafd13. Citations: 18

2016

J. Janssen, N. Gunkelmann and H. M. Urbassek. Influence of C concentration on elastic moduli of  $\alpha' - Fe_{1-x}C_x$  alloys. Philosophical Magazine. doi:10.1080/14786435.2016.1170224. Citations: 11

## PUBLICATIONS IN REVIEW

2021

- J. Janssen, E. Makarov, T. Hickel, A.V. Shapeev and J. Neugebauer. optimization of convergence parameters in plane wave density functional theory calculations via a tensor decomposition-based uncertainty quantification. npj Computational Materials. doi:10.48550/arXiv.2112.04081.
- Code: Opviron/pyiron-dft-uncertainty

## OPEN-SOURCE SOFTWARE

2015 - present pyiron - an integrated development environment for computational materials science

- Role: Lead developer for a team of eight core developers
- Code: Opyiron

2018 – present

Conda-forge - community-led software distribution for the conda package manager

- Role: Maintainer for materials science software
- Contribution: 600+ packages with a total of 145+ million downloads
- Code: conda-forge/staged-recipes

2022

Architector - high-throughput cross-periodic table 3D complex builder

- Contribution: Parallelization of the chemical complex building using mpi4py

2021

FitSNAP - Software for generating SNAP machine-learning interatomic potentials

- Contribution: Implemented a python library interface for the Exascale computing project
- Code: fitsnap/fitsnap

#### WORKSHOPS

2021 **Co-organizer** for the workshop on "workflows for atomistic simulation"

Max-Planck-Institut für Eisenforschung, Düsseldorf, Germany (online)

• Code: • pyiron/potentials-workshop-2021

2020 **Co-organizer** for the workshop on "software tools from atomistics to phase diagrams"

Pennsylvania State University, Pennsylvania, USA (online)

• Code: pyiron/phasediagram-workshop-2020

## **SUPERVISION**

2023 - present Ilgar Baghishov (intern): Multi-fidelity machine learned interatomic potentials

• Role: Primary Supervisor - acquiring the funding and leading the supervision of the intern

2023 – present Ahnaf Akif Alvi (PhD student): Autonomous in silico processing and materials discovery

• Role: Co-Supervisor - in collaboration with Prof. Arroyave at Texas A&M University

2022 – 2023 **Jason Blake Gibson** (intern): Transferability of interatomic machine learning potentials

• Role: Primary Supervisor - acquiring the funding and leading the supervision of the intern

2018 Ankita Biswas (student assistant): Calculation of vacancy formation energies with pyiron

• Role: Primary Supervisor - for a student research project

2017 Martin Böckmann (student assistant): Monte Carlo sampling with pyiron

• Role: Primary Supervisor - guiding the student as the first user of the pyiron software

### **PRESENTATIONS**

# INVITED TALKS

**J. Janssen**. Packaging Scientific Software with Conda-Forge, Workshop at the institute for pure and applied math (IPAM), Los Angeles, CA, USA.

- **J. Janssen**, D. Perez. Up-scaling atomistic simulation workflows with pyiron, Workshop at the institute for pure and applied math (IPAM), Los Angeles, CA, USA.
- **J. Janssen**, M. G. Taylor, P. Yang, D. Perez. Tutorial on high-throughput screening for chemical science with pyiron, 2nd International Workshop on Theory Frontiers in Actinide Science: Chemistry & Materials, Santa Fe, NM, USA.
- **J. Janssen**, T. Hickel, J. Neugebauer. Automated atomistic calculation of thermodynamic and thermophysical data, *DPG Fall Meeting*, Regensburg, Germany.
- **J. Janssen**. pyiron an integrated development environment (IDE) for scientific workflows at scale, *Department of Energy Python Exchange*, New York, NY, USA (online).
- **J. Janssen**. Up-scaling simulation protocols with pyiron, *Platform Material Digital Workflow Meeting 2022*, Berlin, Germany (online).
- **J. Janssen**. pyiron an integrated development environment (IDE) for materials science, Special Interest Group Data Infrastructure (SIGDIUS) Seminar, Stuttgart, Germany (online).

2022

- J. Janssen, J. Neugebauer. pyiron an integrated development environment for materials science, CECAM Workshop: Simulation Workflows in Materials Modelling (SWiMM), Lausanne, Switzerland (online).
- J. Janssen, T. Hickel, J. Neugebauer. Uncertainty quantification for *ab initio* thermodynamics, *Group Seminar Professor R. Drautz*, Bochum, Germany (online).
  - **J.** Janssen, T. Hickel, J. Neugebauer. Automated *ab initio* determination of materials properties at finite temperatures with pyiron, *NIST Workshop: Atomistic simulations for industrial needs*, Gaitgersburg, MD, USA (online).
- J. Janssen, T. Hickel, J. Neugebauer. Automated *ab initio* determination of materials properties at finite temperatures with pyiron, *CLNS Seminar Los Alamos National Laboratory*, Los Alamos, NM, USA.
  - **J. Janssen**, T. Hickel, J. Neugebauer. pyiron an integrated development environment for computational materials science, *Group Seminar Professor G. Kresse*, Vienna, Austria.

#### TALKS AT INTERNATIONAL CONFERENCES

2022

2019

- J. Janssen, D. Perez. Parameter studies for interatomic potentials using LAMMPS and pyiron, *LAMMPS Virtual Workshop and Symposium 2023*, Philadelphia, PA, USA (online).
  - **J. Janssen**, D. Perez. Enabling long timescale molecular dynamics simulation with ab initio precision, *MRS Spring Meeting*, San Francisco, CA, USA.
  - **J. Janssen**, D. Perez. pyiron an integrated development environment for the development and assessment of interatomic models, 11th Annual Mach Conference, Baltimore, MD, USA.
  - **J. Janssen**, D. Perez. Enabling long timescale molecular dynamics simulation with ab initio precision, *TMS Spring Meeting*, San Diego, CA, USA.
  - **J. Janssen**, T. Hickel, J. Neugebauer. Predicting melting temperatures from bulk properties with pyiron, *Multiscale Materials Modelling (MMM) conference*, Baltimore, MD, USA.
    - **J. Janssen**, M. G. Taylor, P. Yang, D. Perez. Screening of ligand-metal-complexes for separation science with pyiron, *ACS Fall Meeting*, Chicago, IL, USA.
    - **J. Janssen**. pyiron an integrated development environment (IDE) for scientific workflows, *Scientific Computing with Python (SciPy)*, Austin, TX, USA.
    - **J. Janssen**. Predicting melting temperatures from bulk properties with pyiron, *Artificial Intelligence for Materials Science (AIMS)*, Gaithersburg, MD, USA (online).
    - **J. Janssen**, T. Hickel, J. Neugebauer. Automated uncertainty quantification for *ab initio* thermodynamics, *MRS Fall Meeting*, Boston, MA, USA.
    - **J. Janssen**, T. Hickel, J. Neugebauer. Automated sensitivity analysis for high-throughput ab initio calculations, *IPAM Workshop*, Los Angeles, CA, USA.
    - **J. Janssen**, T. Hickel, J. Neugebauer. Automated error analysis and control for *ab initio* calculations, *DPG Spring Meeting*, Regensburg, Germany.
    - **J. Janssen**, T. Hickel, J. Neugebauer. Automated sensitivity analysis for high-throughput ab initio calculations, *TMS Spring Meeting*, San Antonio, TX, USA.

- J. Janssen, T. Hickel, J. Neugebauer. Generation of *ab initio* datasets with predefined precision using uncertainty quantification, *DPG Spring Meeting*, Berlin, Germany.
- **J. Janssen**, T. Hickel, J. Neugebauer. Towards an uncertainty quantification for *ab initio* thermodynamics, *MRS Fall Meeting*, Boston, MA, USA.

2016

- **J. Janssen**, T. Hickel, J. Neugebauer. Sensitivity analysis for large sets of density functional theory calculations, *DPG Spring Meeting*, Dresden, Germany.
- **J. Janssen**, T. Hickel, J. Neugebauer. Automated convergence and error analyses for high-precision DFT calculations, *TMS Spring Meeting*, San Diego, CA, USA.
- **J. Janssen**, T. Hickel, J. Neugebauer. Automated convergence checks with the python based library pyiron, *DPG Spring Meeting*, Regensburg, Germany.
  - **J. Janssen**, T. Hickel, J. Neugebauer. Automated convergence checks with the python based workbench pyiron, *TMS Spring Meeting*, Nashville, TN, USA.