

Jan Janssen

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PROFESSIONAL APPOINTMENTS

- 2021 – present **Postdoctoral Research Associate**
Theoretical Division (T-1)
Los Alamos National Laboratory
Los Alamos, NM, USA
- 2015 – 2021 **PhD Candidate**
Computational Materials Design
Max-Planck-Institut für Eisenforschung
Düsseldorf, Germany
- 2017 **Invited Fellow**
Institute for Pure and Applied Mathematics
University of California
Los Angeles, CA, USA



EDUCATION


- 2015 – 2021 **PhD in Theoretical Physics**, Paderborn University, Germany
- 2009 – 2015 **Advanced degree in Theoretical Physics (Master's equivalent)**, Technical University of Kaiserslautern, Germany

AWARDS & HONORS



- 2019 Runner-up for the Heinz Billing Award of 2019

PUBLICATIONS



- 2021 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](https://doi.org/10.1016/j.commatsci.2020.110065).
• Code:  [pyiron/pyiron_meltingpoint](https://github.com/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](https://doi.org/10.1103/PhysRevB.102.100101).
• Code:  [tomswinburne/BLaSA](https://github.com/tomswinburne/BLaSA)

- 2019 **J. Janssen**, S. Surendralal, Y. Lysogorskiy, M. Todorova, T. Hickel, R. Drautz and J. Neugebauer. pyiron: an integrated development environment for computational materials science. *Computational Materials Science*. doi:[10.1016/j.commatsci.2018.07.043](https://doi.org/10.1016/j.commatsci.2018.07.043).
 • Code:  [pyiron](https://github.com/pyiron/pyiron)
- Y. Lysogorskiy, T. Hammerschmidt, **J. Janssen**, J. Neugebauer and R. Drautz. Transferability of interatomic potentials for molybdenum and silicon. *Modelling and Simulation in Materials Science and Engineering*. doi:[10.1088/1361-651X/aafd13](https://doi.org/10.1088/1361-651X/aafd13)
- 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](https://doi.org/10.1080/14786435.2016.1170224)

OPEN-SOURCE SOFTWARE

- 2015 – present **pyiron**
 An integrated development environment for computational materials science
 • Role: Project Lead Developer
 • Code:  [pyiron](https://github.com/pyiron/pyiron)
 • Website: pyiron.org
- 2018 – present **Conda-Forge**
 A community-led collection of recipes, build infrastructure and distributions for the conda package manager.
 • Role: Maintainer for 400+ Materials Science Software Packages
 • Code:  [conda-forge](https://github.com/conda-forge)
 • Website: conda-forge.org

WORKSHOPS

- 2021 Workflows for Atomistic Simulation
Max-Planck-Institut für Eisenforschung, Düsseldorf, Germany (online).
 • Code:  [pyiron/potentials-workshop-2021](https://github.com/pyiron/potentials-workshop-2021)
- 2020 Software Tools from Atomistics to Phase Diagrams
Max-Planck-Institut für Eisenforschung, Düsseldorf, Germany (online).
 • Code:  [pyiron/phasediagram-workshop-2020](https://github.com/pyiron/phasediagram-workshop-2020)

PRESENTATIONS

INVITED TALKS

- 2022 **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).
- 2021 **J. Janssen**, T. Hickel, J. Neugebauer. pyiron - an integrated development environment for materials science, *CECAM Workshop: Simulation Workflows in Materials Modelling (SWiMM)*, Lausanne, Switzerland (online).

- 2020 **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*, Gaitersburg, MD, USA (online).
- 2019 **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

- 2019 **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.
- 2018 **J. Janssen**, T. Hickel, J. Neugebauer. Generation of ab initio datasets with predefined precision using uncertainty quantification, *DPG Spring Meeting*, Berlin, Germany.
- 2017 **J. Janssen**, T. Hickel, J. Neugebauer. Towards an uncertainty quantification for ab initio thermodynamics, *MRS Fall Meeting*, Boston, MA, USA.
- 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.
- 2016 **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.