



Daniil Poletaev

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Personal information

Date of Birth October 24, 1990
Sex Male
Family status Married, two children
Citizenship Russia
Languages English (intermediate), Russian
(native speaker)

Education

09.2012–02.2017 **Candidate of Sciences in Physics and Mathematics (PhD)**, Laboratory of Theoretical investigations and Computer simulation, Belgorod State University, Belgorod, Russia.
Speciality: Condensed Matter Physics
Thesis: *Prediction of hydrogen and silicon solubility in an α -Ti within the density functional theory*

09.2007–06.2012 **Specialist in the field of nanomaterials**, Belgorod State University, Belgorod, Russia.
Department of Physics and Engineering
Thesis: *Methods of molecular dynamics and density functional theory in the studying of the phase transitions in metallic systems on the example of Ti and Cu-Nb composite*

Working experience

2022–present **Research Scientist**, *Materials Discovery Laboratory, Skolkovo Institute of Science and Technology, Moscow, Russia.*
Ab initio calculations of superconducting critical temperature in the predicted materials with account of anharmonic contributions, Supervisor - Oganov A.R.

2022–present **Research Scientist**, *Frank Laboratory of Neutron Physics, Joint Institute for Nuclear Research, Dubna, Russia.*
Ab initio investigations of coherent cluster ordering in magnetostrictive materials of Fe-Ga-Al system, Supervisor - Balagurov A.M.

- 2019–2021 **Research Scientist**, *Laboratory of Theoretical investigations and Computer simulation, Belgorod State University, Belgorod, Russia.*
 – Russian Science Foundation, Grant No. 19-73-00313, "Development of an approach to constructing of phase diagrams of materials with metallic and covalent bond types within the CALPHAD on the basis of experimental data and results of atomistic modeling using the example of the Ti-Si-H system. ", Senior researcher - Poletaev D.O.
- 2011–2019 **Programmer**, *Belgorod virtual branch of the Russian Museum, Belgorod State Institute of Arts and Culture, Belgorod, Russia.*
- 2014–2016 **Junior Scientist**, *Laboratory of Theoretical investigations and Computer simulation, Belgorod State University, Belgorod, Russia.*
 – Ministry of Education of Russian Federation, Grant No. 3.1282.2014/K, "Development of an universal model of interatomic interactions in systems with metallic and covalent types of bonding for the computer design of new materials with predetermined properties within the molecular dynamics, and approbation of the model using the vanadium-iron-titanium system as an example", Senior researcher - Lipnitskii A.G.
- 2012–2015 **Junior Scientist**, *Laboratory of Theoretical investigations and Computer simulation, Belgorod State University, Belgorod, Russia.*
 – Ministry of Education of Russian Federation, Contract No. 02.G25.31.0103, "Development of the technology for the creation of porous bioactive nanostructured coatings on the surface of elements of hip and knee endoprostheses made from titanium alloys that have controlled pore space, specified relief parameters and biochemical activity", Senior researcher - Kolobov Yu.R.

Research interests

With practical background, Computational search for new inorganic materials. Refinement and extension of phase diagrams of inorganic materials with metallic and covalent types of bonding by calculation of the thermodynamic properties of solid phases within the density functional theory, atomistic simulations with semiempirical potentials, and CALPHAD approach. Investigation of grain boundary segregation and diffusion characteristics of impurities in metals within the density functional theory. Construction of semiempirical interatomic potentials.

Without practical background, Development of automated or semi-automated pipelines for computational materials design from crystal structure prediction to practical guidelines for materials manufacturing.

Computer skills

Scientific packages	ABINIT, VASP, Wien2k, LAMMPS, Thermocalc, Pymatgen, Siman
Programming	Python, Fortran, C++, Bash, HTML
General	Experienced user of Windows, Linux OS and \LaTeX
Github	https://github.com/Kurufinve

List of the most important publications

- [1] **D.O. Poletaev**, A.G. Lipnitskii, V.N. Maksimenko, Yu.R. Kolobov, A.G. Beresnev, and M.S. Gusakov. The N-body interatomic potentials for molecular dynamics simulations of diffusion in C15 Cr₂Ta Laves phase. *Computational Materials Science*, 216(June 2022):111841, jan 2023.
- [2] M. Ait Boukideur, N. Selhaoui, F.Z. Chrifi Alaoui, **D.O. Poletaev**, H. Bouchta, K. Achgar, and A. Aharoune. Thermodynamic assessment of the Ga–Lu system by the combination of ab-initio calculations and the CALPHAD approach. *Calphad*, 79(March):102464, dec 2022.
- [3] V. N. Maksimenko, A. G. Lipnitskii, A. I. Kartamyshev, **D. O. Poletaev**, and Yu R. Kolobov. The N-body interatomic potential for molecular dynamics simulations of diffusion in tungsten. *Computational Materials Science*, 202(October 2021):110962, 2022.
- [4] **D.O. Poletaev**, D.A. Aksyonov, and A.G. Lipnitskii. Evolutionary search for new compounds in the Ti–Si system. *Calphad*, 71:102201, dec 2020.
- [5] Anton Boev, **Daniel Poletaev**, Andrey Kartamyshev, Mariya Boeva, and Tatiana Vershinina. Influence of the Mo₁₀Ni₃C₃B phase on the hardness and fracture toughness of Mo–Ni–C–B cermet: experimental and theoretical study. *Letters on Materials*, 10(4):387–391, 2020.
- [6] A.I. Kartamyshev, **D.O. Poletaev**, and A.G. Lipnitskii. The influence of lattice vibrations and electronic free energy on phase stability of titanium silicides and Si solubility in hcp titanium: A DFT study. *Calphad*, 65(February):194–203, jun 2019.
- [7] A.I. Kartamyshev, A.G. Lipnitskii, V.N. Saveliev, V.N. Maksimenko, I.V. Nelasov, and **D.O. Poletaev**. Development of an interatomic potential for titanium with high predictive accuracy of thermal properties up to melting point. *Computational Materials Science*, 160:30–41, apr 2019.
- [8] **D.O. Poletaev**, D.A. Aksyonov, Dat Duy Vo, and A.G. Lipnitskii. Hydrogen solubility in hcp titanium with the account of vacancy complexes and hydrides: A DFT study. *Computational Materials Science*, 114:199–208, 2016.
- [9] **D.O. Poletaev**, A.G. Lipnitskii, A.I. Kartamyshev, D.A. Aksyonov, E.S. Tkachev, S.S. Manokhin, M.B. Ivanov, and Yu. R. Kolobov. Ab initio-based prediction and TEM study of silicide precipitation in titanium. *Computational Materials Science*, 95:456–463, 2014.

Conferences

1. Inaugural Symposium for “Computational Materials Program of Excellence”, 04 – 06 September 2019, Skoltech, Russia (<https://cmp.skoltech.ru/>) - poster presentation
2. International Conference on Computer Coupling of Phase Diagrams and Thermochemistry "CALPHAD GLOBAL", June 2021, World Wide Web (<https://calphad.org/calphad-global-home>) - online poster presentation