

Yury Basalaev

List of Publications by Year in descending order

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51

papers

256

citations

1163117

8

h-index

1125743

13

g-index

51

all docs

51

docs citations

51

times ranked

137

citing authors

#	ARTICLE	IF	CITATIONS
1	BAND STRUCTURE OF BePb ₂ PNICTIDES: AB INITIO CALCULATIONS. <i>Journal of Structural Chemistry</i> , 2022, 63, 588-592.	1.0	0
2	WANNIER FUNCTIONS AND CHEMICAL BONDING IN COMPOUNDS Be _{IV} P ₂ (IV=Al, Si, Ge, Sn) WITH CHALCOPYRITE STRUCTURE. <i>Journal of Structural Chemistry</i> , 2021, 62, 817-823.	1.0	1
3	ELECTRONIC AND VIBRATIONAL STRUCTURES OF BeC ₂ PNICTIDES. <i>Journal of Structural Chemistry</i> , 2021, 62, 985-994.	1.0	0
4	Electron Structure of Be _{IV} P ₂ Crystals with a Chalcopyrite Lattice. <i>Physics of the Solid State</i> , 2020, 62, 2016-2023.	0.6	6
5	A First-Principles Simulation of Electronic Structure of MCN ₂ Crystals (M = Be, Mg, Ca, Zn, Cd, Hg). <i>Journal of Structural Chemistry</i> , 2020, 61, 337-343.	1.0	9
6	SIMULATION OF ELECTRONIC STRUCTURES OF Hg _{IV} V ₂ COMPOUNDS. <i>Journal of Structural Chemistry</i> , 2020, 61, 1007-1016.	1.0	8
7	ELECTRONIC STRUCTURE OF PNICTIDES HgCX ₂ . <i>Journal of Structural Chemistry</i> , 2020, 61, 1839-1844.	1.0	3
8	Electronic Structure of CO ₂ and CS ₂ Crystals. <i>Journal of Structural Chemistry</i> , 2019, 60, 1019-1026.	1.0	2
9	Simulation of the Crystal Structure of GelSimCn Compounds and the Study of Their Electronic Structure. <i>Journal of Structural Chemistry</i> , 2019, 60, 518-523.	1.0	1
10	Modeling of Hypothetical BeXAs ₂ (X = Si, Ge, Sn) Crystal Structure and Phonon Spectra. <i>Russian Physics Journal</i> , 2019, 61, 1978-1984.	0.4	2
11	Electronic and Vibrational Properties of LiBO ₂ Crystals. <i>Russian Physics Journal</i> , 2019, 61, 1868-1875.	0.4	8
12	Electronic Structure and Elastic Properties of ZnCdSe ₂ Crystal with the Chalcopyrite Structure. <i>Journal of Structural Chemistry</i> , 2018, 59, 15-19.	1.0	2
13	Ab Initio Study of the ZnSnSb ₂ Semiconductor. <i>Semiconductors</i> , 2018, 52, 1715-1720.	0.5	2
14	Crystalline, Electronic, and Vibrational Structures of Zinc Cyanides. <i>Journal of Structural Chemistry</i> , 2018, 59, 1761-1767.	1.0	2
15	An Ab Initio Study of Electronic Structure of Lithium Metaborate. <i>Journal of Structural Chemistry</i> , 2018, 59, 1501-1506.	1.0	5
16	Electronic, Vibrational, and Elastic Properties of Zn ₂ SeTe Crystals with Antihalcopirite Structure. <i>Russian Physics Journal</i> , 2018, 61, 1425-1428.	0.4	1
17	Electronic structure of single-layer superlattices (GeC)1(SiC)1, (SnC)1(SiC)1, and (SnC)1(GeC)1. <i>Semiconductors</i> , 2017, 51, 617-620.	0.5	3
18	Ab initio study of the electronic and vibrational structures of tetragonal cadmium diarsenide. <i>Semiconductors</i> , 2017, 51, 783-788.	0.5	5

#	ARTICLE		IF	CITATIONS
19	Energy Band Structure of Beâ€“(C, Si, Ge, Sn)â€“N ₂ Crystals. Russian Physics Journal, 2017, 60, 900-907.		0.4	12
20	Ab initio and phenomenological simulation of the phonon spectra BeMN ₂ (M = C, Si, Ge, Sn) crystals. Journal of Structural Chemistry, 2017, 58, 1588-1596.		1.0	6
21	Modeling of half-Heusler crystals with the chalcopyrite structure: Li ₂ MgZnX ₂ (X = N, P, As, Sb). Journal of Structural Chemistry, 2016, 57, 1499-1504.		1.0	1
22	Sublattice effect on the formation of the band structure of crystals with the chalcopyrite lattice: B ₂ CN, BC ₂ N, BCN ₂ . Journal of Structural Chemistry, 2016, 57, 8-13.		1.0	6
23	Modeling of Electronic Structure of a Monolayer Superlattice SiC/GeC. Russian Physics Journal, 2016, 59, 1111-1114.		0.4	1
24	Features of the valence electron charge distribution in LiB ₁₁ X ₅ crystals. Journal of Structural Chemistry, 2015, 56, 1036-1042.		1.0	1
25	Modeling of Electronic Structure of Cu ₂ BrCl Crystals with Chalcopyrite Lattice. Russian Physics Journal, 2015, 58, 1201-1204.		0.4	2
26	Ab Initio and Phenomenological Modeling of the Phonon Spectrum of Superhard cp-BC ₂ N. Russian Physics Journal, 2015, 58, 978-986.		0.4	5
27	Electronic structure and lattice dynamics of the $\hat{I}\pm$ -ZnCl ₂ crystal. Journal of Structural Chemistry, 2015, 56, 823-828.		1.0	2
28	New Diamond-Like Compounds with Anti-chalcopyrite Structure. Russian Physics Journal, 2014, 57, 558-560.		0.4	10
29	Features of the band structure of (CuInSe ₂) _{1-x} (MeSe) _x alloys (Me = Mn, Fe). Semiconductors, 2014, 48, 417-422.		0.5	3
30	Electronic structure of the CuBS ₂ crystal. Physics of the Solid State, 2012, 54, 1764-1767.		0.6	4
31	Influence of the sublattice symmetry on the band structure of a MgSiN ₂ crystal. Russian Physics Journal, 2012, 54, 1145-1151.		0.4	4
32	Role of sublattices in the formation of the electronic structure and chemical bonding in a Zn ₂ SiO ₄ crystal with a defect chalcopyrite lattice. Journal of Structural Chemistry, 2012, 53, 35-38.		1.0	4
33	Zone structure and its genesis from the states of sublattices in orthorhombic MgGeN ₂ . Moscow University Physics Bulletin (English Translation of Vestnik Moskovskogo Universiteta, Fizika), 2011, 66, 39-44.		0.4	6
34	Special features of the electronic structure of Ga ₂ AsSb crystals. Russian Physics Journal, 2011, 53, 1216-1218.		0.4	4
35	Energy band genesis from sublattice states in MgSiN ₂ and MgGeN ₂ crystals. Journal of Structural Chemistry, 2010, 51, 1191-1194.		1.0	14
36	Electron structure of hypothetical IVâ€“IVâ€“IV ₂ -type crystals having a chalcopyrite-type lattice. Russian Physics Journal, 2009, 52, 992-994.		0.4	7

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37	Role of sublattices in the formation of the chemical bond in ion-covalent crystals. Journal of Structural Chemistry, 2009, 50, 1177-1180.	1.0	11
38	Lattice dynamics of chalcopyrite semiconductors LiAlTe_{2} , LiGaTe_{2} and LiInTe_{2} . Physica Status Solidi (B): Basic Research, 2009, 246, 364-371.	1.5	46
39	Specific features of electronic and vibrational properties of I-V-V2 crystals. Semiconductors, 2009, 43, 735-739.	0.5	2
40	Optical functions of ternary Li-containing tellurides. Russian Physics Journal, 2007, 50, 1232-1236.	0.4	0
41	LiPN_2 and NaPN_2 crystals: Structural features and chemical bonding. Journal of Structural Chemistry, 2007, 48, 996-1000.	1.0	5
42	Chemical bonding in isostructural Li-containing ternary chalcogenides. Journal of Structural Chemistry, 2007, 48, 1001-1005.	1.0	11
43	Optical Properties of ZnGeP_2 in the UV Spectral Region. Semiconductors, 2005, 39, 1004.	0.5	3
44	Electronic Structure of Triple Phosphides MgSiP_2 , ZnSiP_2 , and CdSiP_2 . Russian Physics Journal, 2005, 48, 78-83.	0.4	9
45	The genesis of energy bands formed by sublattice states in alkaline-earth metal oxides and sulfides. Physics of the Solid State, 2004, 46, 848-852.	0.6	3
46	Title is missing!. Theoretical and Experimental Chemistry, 2003, 39, 81-84.	0.8	0
47	Electron Density Calculations for Crystals with a NaCl Lattice. Journal of Structural Chemistry, 2001, 42, 172-176.	1.0	3
48	Electronic Structure of Alkali Metal Oxides and Sulfides. Russian Physics Journal, 2001, 44, 398-403.	0.4	10
49	Structure effects in the electronic energy spectrum of sodium nitrate. Journal of Structural Chemistry, 1996, 37, 508-510.	1.0	0
50	Bond charges and structural features in $\text{A}_2\text{B}_4\text{C}_2\text{S}_5$ crystals. Journal of Structural Chemistry, 1992, 32, 534-537.	1.0	0
51	The electronic structure and chemical bonding in MgPbP_2 . Physica Status Solidi (B): Basic Research, 0, ..	1.5	1