

# Badis Bennecer

## List of Publications by Year in descending order

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Version: 2024-02-01

33

papers

497

citations

687363

13

h-index

713466

21

g-index

34

all docs

34

docs citations

34

times ranked

476

citing authors

#	ARTICLE	IF	CITATIONS
1	Ab initio predictions of structures and physical properties of the KCuX (X = Se and Te) phases under pressure. Computational Condensed Matter, 2022, 30, e00616.	2.1	2
2	Theoretical study of the structural, elastic, vibrational and thermal properties of perovskite halides $\text{Cs}_{1-x}\text{TiBr}_x$ : Ab initio calculations. Computational Condensed Matter, 2021, 29, e00587.	2.1	6
3	Pressure induced phase transition, electronic and optical properties of LiBeX (X = As, Sb and Bi). Journal of Physics Condensed Matter, 2020, 32, 325503.	1.8	4
4	Phase transitions and lattice dynamics in perovskite-type hydride $\text{Li}_{1-x}\text{Na}_{x}\text{MgH}_3$ . Journal of Physics Condensed Matter, 2019, 31, 505402.	1.8	2
5	Magneto-optical Kerr effect in ZnTMO <sub>2</sub> (TM=Cr, Mn, Fe, Co and Ni). Journal of Magnetism and Magnetic Materials, 2017, 424, 327-338.	2.3	8
6	Structural, electronic and optical properties of LiBeP in its normal and high pressure phases. Journal of Physics and Chemistry of Solids, 2014, 75, 838-848.	4.0	4
7	Elastic and lattice dynamical properties of ternary strontium chalcogenide alloys. Materials Science in Semiconductor Processing, 2014, 26, 267-275.	4.0	7
8	Electronic and optical properties of the orthorhombic compounds FeX <sub>2</sub> (X=P, As and Sb). Materials Science and Engineering B: Solid-State Materials for Advanced Technology, 2013, 178, 1249-1256.	3.5	10
9	Elastic, vibrational and thermodynamic properties of based group IV semiconductors and GeC under pressure. Journal of Physics and Chemistry of Solids, 2013, 74, 1615-1625.	4.0	16
10	Pressure effect on the electronic and optical properties of the FeP <sub>2</sub> and FeAs <sub>2</sub> compounds. Journal of Physics and Chemistry of Solids, 2013, 74, 1336-1340.	4.0	4
11	Ab initio lattice dynamics and thermodynamic properties of SrO under pressure. Journal of Physics and Chemistry of Solids, 2012, 73, 129-135.	4.0	25
12	Ab initio calculation of vibrational and thermodynamic properties of SrX (S, Se, Te) in the B1 (NaCl) and B2 (CsCl) structures. Computational Materials Science, 2011, 50, 1701-1710.	3.0	36
13	Elastic properties and lattice dynamics of alkali chalcogenide compounds Na <sub>2</sub> S, Na <sub>2</sub> Se and Na <sub>2</sub> Te. Computational Materials Science, 2011, 50, 1806-1810.	3.0	17
14	Elastic and electronic properties of the alkali pnictide compounds Li <sub>3</sub> Sb, Li <sub>3</sub> Bi, Li <sub>2</sub> NaSb and Li <sub>2</sub> NaBi. Computational Materials Science, 2011, 50, 2880-2885.	3.0	24
15	Optical properties of the alkali antimonide semiconductors $\text{Na}_x\text{Sb}_y$ , $\text{Na}_x\text{Bi}_y$ and $\text{Na}_x\text{Bi}_y\text{Sb}_z$ . Journal of Physics and Chemistry of Solids, 2010, 71, 314-322.	4.0	39
16	Pressure effect on the electronic and optical properties of the alkali antimonide semiconductors Cs <sub>3</sub> Sb, KCs <sub>2</sub> Sb, CsK <sub>2</sub> Sb and K <sub>3</sub> Sb: Ab initio study. Journal of Physics and Chemistry of Solids, 2010, 71, 1732-1741.	4.0	20
17	Structural and electronic properties of the pseudo-binary compounds ( $\text{Na}_x\text{Sb}_y$ , S and Se). Journal of Physics and Chemistry of Solids, 2010, 71, 42-46.	4.0	22
18	Elastic and electronic properties of Li <sub>2</sub> ZnGe. Computational Materials Science, 2010, 47, 869-874.	3.0	12

#	ARTICLE	IF	CITATIONS
19	Electronic and optical properties of the orthorhombic compounds PdPX (X=S and Se). Computational Materials Science, 2010, 48, 115-123.	3.0	16
20	Calculation of the vibrational properties of LiMgAs. Journal of Physics Condensed Matter, 2009, 21, 305402.	1.8	4
21	Optical Properties of the Strained Layer Superlattices (GaAs) <sub>6</sub> /(GaP) <sub>m</sub> (001) (m=2, 4 and 6). Materials Science Forum, 2009, 609, 41-48.	0.3	2
22	Structural and elastic properties of the half-Heusler compounds IrMnZ (, Sn and Sb). Materials Chemistry and Physics, 2009, 114, 732-735.	4.0	15
23	Electronic and optical properties of LiMgN, LiMgP and LiMgAs under hydrostatic pressure. Journal of Physics and Chemistry of Solids, 2009, 70, 26-31.	4.0	7
24	Vibrational properties of the filled tetrahedral compounds LiCdP and LiCdAs. Computational Materials Science, 2009, 44, 876-880.	3.0	5
25	Electronic structure of Li <sub>3</sub> GaN <sub>2</sub> . Physics Letters, Section A: General, Atomic and Solid State Physics, 2008, 372, 1324-1326.	2.1	4
26	Electronic and optical properties of the antifluorite semiconductors Be <sub>2</sub> C and Mg <sub>2</sub> X (, Si, Ge) under hydrostatic pressure. Journal of Physics and Chemistry of Solids, 2008, 69, 1775-1781.	4.0	40
27	First principles calculations of the structural and elastic properties of the filled tetrahedral compounds LiCdX (X=N,P,As). Computational Materials Science, 2008, 42, 579-583.	3.0	12
28	Electronic structure of the filled tetrahedral compound LiCdP and zinc-blende InP: Application of the interstitial insertion rule. Computational Materials Science, 2008, 43, 791-795.	3.0	15
29	Pressure effect on the optical properties of the filled tetrahedral semiconductors LiZnX. Journal of Physics and Chemistry of Solids, 2007, 68, 2286-2292.	4.0	13
30	Optical properties of the filled tetrahedral semiconductors LiMgX (X = N, P and As). Journal of Physics Condensed Matter, 2006, 18, 7237-7247.	1.8	27
31	Structural and elastic properties of the filled tetrahedral semiconductors LiZnX (X=N, P, and As). Journal of Physics and Chemistry of Solids, 2006, 67, 846-850.	4.0	39
32	Optical properties of the filled tetrahedral semiconductors LiZnX (X=N, P, and As). Journal of Physics and Chemistry of Solids, 2006, 67, 1850-1857.	4.0	39
33	Calculated acoustic plasmon spectra in SnTe: effect on anisotropy. Semiconductor Science and Technology, 1992, 7, 822-827.	2.0	1