Saadi Berri

List of Publications by Year in descending order

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471061 476904 46 931 17 29 citations h-index g-index papers 46 46 46 391 all docs docs citations times ranked citing authors

#	Article	IF	CITATIONS
1	Probable thermoelectric materials for promising candidate of optoelectronics for Baâ€based complex perovskite compounds. International Journal of Energy Research, 2022, 46, 9968-9984.	2.2	8
2	First-principle calculations to investigate structural, electronic, optical, thermodynamic, and thermoelectric properties of ABO3 (A=Cs, Rb and B= Ta, Nb) compounds. Emergent Materials, 2022, 5, 1831-1847.	3.2	12
3	The magnetic and thermoelectric properties of Co- and Mn-doped CsPbI3. Emergent Materials, 2022, 5, 1859-1869.	3.2	5
4	First principle analysis of structural, electronic, optical, and thermoelectric characteristics of Ba3CaTa2O9 complex perovskite. Emergent Materials, 2022, 5, 1849-1857.	3.2	14
5	Study on quaternary diamond-like Li2CaGeO4 properties for optoelectronic applications. Computational Condensed Matter, 2022, 30, e00646.	0.9	1
6	Computational study of structural, electronic, half-metallic and thermodynamic properties of Sr3X2O7 (X = Fe, Mn) compounds. Indian Journal of Physics, 2021, 95, 2293-2301.	0.9	2
7	Electronic Structure and Thermoelectric Properties of Coâ€; Feâ€; Mnâ€; and Crâ€Doped Ba ₂ LuTaO ₆ from Spinâ€Polarized Calculations. Physica Status Solidi (B): Basic Research, 2021, 258, 2000402.	0.7	7
8	Influence of Cuâ€Doping on Linear and Nonlinear Optical Properties of Highâ€Quality ZnO Thin Films Obtained by Spinâ€Coating Technique. Physica Status Solidi (B): Basic Research, 2021, 258, 2000472.	0.7	5
9	Ab-initio calculations on structural, electronic, half-metallic and optical properties of Co-, Fe-, Mn- and Cr-doped $\$hbox \{Ba\}_{{mathbf \{2\}}}\box \{LuTaO\}_{{mathbf \{6\}}}$$. Pramana - Journal of Physics, 2021, 95, 1.	0.9	18
10	Band parameters and thermoelectric properties of chalcopyrite ternary compounds CdXP2 (X = Si, Ge) Tj ETQq0	0 0 rgBT /	Overlock 10 T
11	Half-metallic and thermoelectric properties of Sr2EuReO6. Computational Condensed Matter, 2021, 28, e00586.	0.9	40
12	Theoretical study of physical properties of Ba3B(Nb,Ta)2O9 (B = Mg, Ca, Sr, Cd, Hg, Zn, Fe, Mn, Ni, Co) perovskites. Computational Condensed Matter, 2021, 29, e00595.	0.9	31
13	First-principles investigation of the physical properties of XSb2O6 (XÂ= Ca, Sr, Ba) and YAs2O6 (YÂ= Mn,) Tj ETC	0.90.78	34314 rgBT /O
14	Theoretical analysis of the structural, electronic, optical and thermodynamic properties of trigonal and hexagonal Cs3Sb2I9 compound. European Physical Journal B, 2020, 93, 1.	0.6	21
15	Computational Study of Structural, Electronic, Elastic, Half-Metallic and Thermoelectric Properties of CoCrScZ (Z=Al, Si, Ge, and Ga) Quaternary Heusler Alloys. Journal of Superconductivity and Novel Magnetism, 2020, 33, 3809-3818.	0.8	22
16	First-principles studies of thermoelectric and thermodynamic properties of the complex perovskite Ba3MnNb2O9. Journal of Science: Advanced Materials and Devices, 2020, 5, 378-384.	1.5	6
17	First-Principles Predictions on Half-Metallic Results of RBaMn2O6- \hat{l} (R = Nd, Pr, La and \hat{l} = 0, 1) Double Perovskite Compounds. Journal of Superconductivity and Novel Magnetism, 2020, 33, 1737-1746.	0.8	18
18	First-principles investigation of structural, electronic, optical and thermodynamic properties of KAg2SbS4. Computational Condensed Matter, 2019, 19, e00365.	0.9	6

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19	First-principles calculations to investigate structural, electronic, half-metallic and thermodynamic properties of hexagonal UX2O6 (X=Cr,V) compounds. Journal of Science: Advanced Materials and Devices, 2019, 4, 319-326.	1.5	5
20	Theoretical investigation of the structural, electronic and thermodynamic properties of cubic and orthorhombic XZrS3 (X = Ba,Sr,Ca) compounds. Journal of Computational Electronics, 2019, 18, 415-42	7 ^{1.3}	22
21	First-Principles Calculations to Investigate the Structural, Electronic, and Half-Metallic Properties of Ti2RhSn1-xSix, Ti2RhSn1-xGex, and Ti2RhGe1-xSix (x = 0, 0.25, 0.5, 0.75, and 1) Quaternary Heusler A Journal of Superconductivity and Novel Magnetism, 2019, 32, 2219-2228.	ulloys.	15
22	Ab initio study of fundamental properties of X AlO 3 (\times Â=ÂCs, Rb and K) compounds. Journal of Science: Advanced Materials and Devices, 2018, 3, 254-261.	1.5	21
23	Search for New Half-Metallic Ferromagnets in Quaternary Diamond-Like Compounds l–II2–III–VI4 and I2–II–VI4 (I = Cu; II = Mn, Fe, Co; III = In; IV = Ge, Sn; VI = S, Se, Te). Journal of Superconductivity and Novel Magnetism, 2018, 31, 1941-1947.	0.8	18
24	Density functional studies of magneto-optic properties of Sr2GdReO6. Modern Electronic Materials, 2018, 4, 13-19.	0.2	6
25	Electronic structure and half-metallicity of the new Heusler alloys PtZrTiAl, PdZrTiAl and Pt 0.5 Pd 0.5 ZrTiAl. Chinese Journal of Physics, 2017, 55, 195-202.	2.0	40
26	First-principles investigation for some physical properties of some fluoroperovskites compounds ABF3 (AÂ=ÂK, Na; BÂ=ÂMg, Zn). Indian Journal of Physics, 2017, 91, 1513-1523.	0.9	8
27	Magnetic and electronic properties of Li-, Be-, B- and F- doped C 3 N 4 : Ab initio calculations. Computational Condensed Matter, 2017, 12, 25-31.	0.9	2
28	Theoretical analysis of the structural, electronic and optical properties of tetragonal Sr2GaSbO6. Chinese Journal of Physics, 2017, 55, 2476-2483.	2.0	19
29	<i>Ab initio</i> study of electronic structure and magnetic properties of CoMnTaZ (Z = Si, Ge) quaternary Heusler compounds. Physica Status Solidi C: Current Topics in Solid State Physics, 2017, 14, 1700127.	0.8	3
30	Half-Metallic Ferromagnetism in Li6VCl8, Li6MnCl8, Li6CoCl8 and Li6FeCl8 from First Principles. Journal of Superconductivity and Novel Magnetism, 2016, 29, 2381-2386.	0.8	10
31	Electronic Structure and Magnetism of Ti2(Pd, Pt) (B, Al, Ga, In): A First-Principle Study. Journal of Superconductivity and Novel Magnetism, 2016, 29, 2189-2194.	0.8	4
32	Electronic structure and magnetic properties of Co2TaAl from ab initio calculations. Journal of Science: Advanced Materials and Devices, 2016, 1, 286-289.	1.5	11
33	First-principles Study on Half-metallic Properties of the CoMnCrSb Quaternary Heusler Compound. Journal of Superconductivity and Novel Magnetism, 2016, 29, 1309-1315.	0.8	57
34	The electronic structure and spin polarization of Co 2 Mn 0.75 (Gd, Eu) 0.25 Z (Z =Si, Ge, Ga, Al) quaternary Heusler alloys. Journal of Magnetism and Magnetic Materials, 2016, 401, 667-672.	1.0	33
35	First-principles study on half-metallic properties of the Sr 2 GdReO 6 double perovskite. Journal of Magnetism and Magnetic Materials, 2015, 385, 124-128.	1.0	75
36	Ab initio prediction of structural, electronic, magnetic and optical properties of Ba2GdSbO6. Materials Science in Semiconductor Processing, 2015, 40, 58-63.	1.9	9

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37	Ab initio study of electronic structure and magnetic properties in ferromagnetic Sr1â^'x(Mn, Cr)xO alloys. Materials Science in Semiconductor Processing, 2015, 38, 101-106.	1.9	19
38	Robust half-metallic ferromagnet of quaternary Heusler compounds ZrCoTiZ(ZÂ=ÂSi, Ge, Ga and Al). Computational Condensed Matter, 2014, 1, 26-31.	0.9	77
39	A first-principle study of half-metallic ferrimagnetism in the CoFeTiSb quaternary Heusler compound. Journal of Magnetism and Magnetic Materials, 2014, 354, 65-69.	1.0	62
40	First principles study of structural, electronic and magnetic properties of ZrFeTiAl, ZrFeTiSi, ZrFeTiGe and ZrNiTiAl. Journal of Magnetism and Magnetic Materials, 2014, 371, 106-111.	1.0	77
41	First principles study of structural, electronic and magnetic properties of Mn2CoAs. Journal of Magnetism and Magnetic Materials, 2014, 361, 132-136.	1.0	14
42	Electronic structure and magnetic properties of the perovskite cerium manganese oxide from ab initio calculations. Materials Science in Semiconductor Processing, 2014, 26, 199-204.	1.9	28
43	First principles study of structural, electronic and optical properties of AgSbS2. Materials Science in Semiconductor Processing, 2013, 16, 1439-1446.	1.9	18
44	Study of structural, electronic and magnetic properties of Rh2MnX (X=Al, Ge and Sn) Heusler alloys using GGA-WC and GGA+U approaches. Physica B: Condensed Matter, 2013, 418, 58-64.	1.3	18
45	Ab initio study of the structural, electronic and elastic properties of AgSbTe2, AgSbSe2, Pr3AlC, Ce3AlC, Ce3AlN, La3AlC and La3AlN compounds. Physica B: Condensed Matter, 2012, 407, 3320-3327.	1.3	13
46	Ab initio study of the structural, electronic, elastic and magnetic properties of Cu2GdIn, Ag2GdIn and Au2GdIn. Physica B: Condensed Matter, 2012, 407, 3328-3334.	1.3	8