## Y Cheng

## List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/6353590/publications.pdf

Version: 2024-02-01

		1040056	1281871	
12	210	9	11	
papers	citations	h-index	g-index	
12	12	12	238	
all docs	docs citations	times ranked	citing authors	

#	Article	IF	CITATIONS
1	First-principles study of structural, elastic, electronic and thermodynamic properties of topological insulator Sb2Te3 under pressure. Computational Materials Science, 2015, 96, 342-347.	3.0	16
2	Structural, elastic, thermodynamic, electronic properties and phase transition in half-Heusler alloy NiVSb at high pressures. Computational Materials Science, 2015, 96, 72-80.	3.0	28
3	Thermodynamic and Electronic Properties of OsB2from First-Principles Calculations. Acta Physica Polonica A, 2014, 125, 1186-1190.	0.5	O
4	Elastic and electronic properties of MnTi2O4 under pressure: A first-principle study. Computational Materials Science, 2014, 84, 156-162.	3.0	9
5	Structural, electronic and elastic properties of AlFe 2 B 2 : First-principles study. Computational Materials Science, 2014, 92, 253-257.	3.0	32
6	Elastic, thermodynamic and electronic properties of LaF3 under pressure from first principles. Computational Materials Science, 2014, 89, 57-64.	3.0	9
7	Elastic and electronic properties of Ce2O3 from first principles. Journal of Alloys and Compounds, 2013, 551, 672-676.	5 <b>.</b> 5	13
8	First principles study of electronic, bonding, elastic properties and intrinsic hardness of CdSiP2. Computational Materials Science, 2013, 77, 114-119.	3.0	32
9	First principles investigations on elastic and electronic properties of BaHfN2 under pressure. Journal of Alloys and Compounds, 2012, 526, 74-78.	5 <b>.</b> 5	14
10	Elastic stability and electronic structure of tantalum boride investigated via first-principles density functional calculations. Journal of Physics and Chemistry of Solids, 2012, 73, 1197-1202.	4.0	11
11	Pressure effects on elastic and thermodynamic properties of Zr3Al intermetallic compound. Computational Materials Science, 2012, 58, 125-130.	3.0	31
12	Ab initio study for the intermolecular interaction potential surface of Ar-N2 complex. European Physical Journal D, 2005, 33, 43-48.	1.3	15