## Hao-ran Gong

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Phase stability and mechanical property of W–Cu solid solutions from a newly derived W–Cu potential. Physica B: Condensed Matter, 2022, 624, 413436.	2.7	6
2	Atomistic modelling of the immiscible Fe–Bi system from a constructed bond order potential. Journal of Physics Condensed Matter, 2022, 34, 025901.	1.8	3
3	Heats of formation and stress-strain relationship of Fe-Cr solid solutions from a constructed Fe-Cr potential. Journal of Physics Condensed Matter, 2022, , .	1.8	0
4	Cohesive properties of PbBi/Fe <sub>3</sub> O <sub>4</sub> and PbBi/(Fe,Cr) <sub>3</sub> O <sub>4</sub> interfaces. Physical Chemistry Chemical Physics, 2022, 24, 6732-6741.	2.8	4
5	Fundamental effect of lead on mechanical properties of iron from a constructed iron-lead potential. Computational Materials Science, 2022, 212, 111587.	3.0	3

6 Addendum: Magnetic ground state of face-centered-cubic structure of iron (2020 J. Phys.: Condens.) Tj ETQq0 0 0 rgBT /Overlock 10 Tf

7	Phonon spectrum and thermoelectric properties of square/octagon structure of bismuth monolayer. RSC Advances, 2021, 11, 5107-5117.	3.6	2
8	Atomic structure, tensile property, and dislocation behavior of Fe–W interfaces from molecular dynamics simulation. Journal of Physics Condensed Matter, 2021, 33, 145901.	1.8	6
9	Stacking fault energy and electronic structure of molybdenum under solid solution softening/hardening. Journal of Central South University, 2021, 28, 39-47.	3.0	5
10	Hydrogen solubility and diffusivity at Σ3 grain boundary of PdCu. RSC Advances, 2021, 11, 13644-13652.	3.6	4
11	Magnetic ground state of face-centered-cubic structure of iron. Journal of Physics Condensed Matter, 2020, 32, 165806.	1.8	12
12	Band structure, phonon spectrum, and thermoelectric properties of β-BiAs and β-BiSb monolayers. Journal of Materials Chemistry C, 2020, 8, 581-590.	5.5	21
13	Liquid metal embrittlement of an Fe10Cr4Al ferritic alloy exposed to oxygen-depleted and -saturated lead-bismuth eutectic at 350°C. Corrosion Science, 2020, 165, 108364.	6.6	26
14	Construction of an n-body Fe–Cu potential and its application in atomistic modeling of Fe–Cu solid solutions. Journal of Applied Physics, 2020, 127, 045104.	2.5	6
15	Structural stability and mechanical property of Fe-W solid solutions from a constructed Fe-W potential. Journal of Applied Physics, 2019, 126, .	2.5	18
16	Band structure and thermoelectric performances of antimony under trigonal transformation. Journal of Applied Physics, 2019, 125, .	2.5	20
17	Strain-stress relationship and dislocation evolution of W–Cu bilayers from a constructed <i>n</i> -body W–Cu potential. Journal of Physics Condensed Matter, 2019, 31, 305002.	1.8	23
18	Effects of low dimensionality on electronic structure and thermoelectric properties of bismuth. RSC Advances, 2019, 9, 40670-40680.	3.6	10

Hao-ran Gong

#	Article	IF	CITATIONS
19	Influence of internal displacement on band structure, phase transition, and thermoelectric properties of bismuth. Journal of Materials Science, 2019, 54, 6347-6360.	3.7	17
20	8-Layer Shifted Hexagonal Perovskite Ba <sub>8</sub> MnNb <sub>6</sub> O <sub>24</sub> : Long-Range Ordering of High-Spin d <sup>5</sup> Mn <sup>2+</sup> Layers and Electronic Structure. Inorganic Chemistry, 2018, 57, 5732-5742.	4.0	10
21	Computationally predicted fundamental behaviors of embedded hydrogen at TiC/W interfaces. International Journal of Hydrogen Energy, 2018, 43, 16180-16186.	7.1	21
22	Effects of trigonal deformation on electronic structure and thermoelectric properties of bismuth. Journal of Physics Condensed Matter, 2018, 30, 285504.	1.8	17
23	Thermodynamic and mechanical properties of TiC from <i>ab initio</i> calculation. Journal of Applied Physics, 2014, 116, .	2.5	29
24	Structural, thermodynamic, mechanical, and magnetic properties of FeW system. Journal of Applied Physics, 2014, 116, .	2.5	25
25	Atomic structure, mechanical quality, and thermodynamic property of TiH <sub><i>x</i></sub> phases. Journal of Applied Physics, 2013, 114, 043510.	2.5	32
26	Interface structure and work function of W-Cu interfaces. Applied Physics Letters, 2013, 103, .	3.3	43
27	Comparative study of metastable phase formation in the immiscible Cu–W system byab initiocalculation andn-body potential. Journal of Physics Condensed Matter, 2004, 16, 5251-5258.	1.8	18
28	Metastability of an immiscible Cu-Mo system calculated from first-principles and a derivedn-body potential. Physical Review B, 2004, 69, .	3.2	18
29	Structural stability and magnetic properties of metastable Fe-Cu alloys studied byab initiocalculations and molecular dynamics simulations. Physical Review B, 2004, 69, .	3.2	17
30	Glass-forming ability determined by ann-body potential in a highly immiscible Cu-W system through molecular dynamics simulations. Physical Review B, 2003, 68, .	3.2	20
31	Atomistic modeling of solid-state amorphization in an immiscible Cu-Ta system. Physical Review B, 2002, 66, .	3.2	56