Dilpuneet S Aidhy

List of Publications by Year in descending order

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39 1,205
papers citations

331670 377865
21 34
h-index g-index

40 40 all docs citations

40 times ranked 1475 citing authors

#	Article	IF	CITATIONS
1	Effect of different point-defect energetics in Ni80X20 (X=Fe, Pd) on contrasting vacancy cluster formation from atomistic simulations. Materialia, 2021, 15, 100974.	2.7	2
2	Breaking atomic-level ordering via biaxial strain in functional oxides: A DFT study. Journal of Applied Physics, 2021, 129, 095301.	2.5	8
3	A statistical approach for atomistic calculations of vacancy formation energy and chemical potentials in concentrated solid-solution alloys. Computational Materials Science, 2021, 190, 110308.	3.0	24
4	Coupling between interfacial strain and oxygen vacancies at complex-oxides interfaces. Journal of Applied Physics, 2021, 129, 171102.	2.5	9
5	Machine Learning Based Methodology to Predict Point Defect Energies in Multi-Principal Element Alloys. Frontiers in Materials, 2021, 8, .	2.4	19
6	Factors affecting the vacancy formation energy in Fe70Ni10Cr20 random concentrated alloy. Computational Materials Science, 2021, 198, 110669.	3.0	10
7	â~3 Twin Boundaries in Gd ₂ Ti ₂ O ₇ Pyrochlore: Pathways for Oxygen Migration. ACS Applied Materials & Samp; Interfaces, 2020, 12, 45558-45563.	8.0	9
8	Machine Learning Enabled Prediction of Stacking Fault Energies in Concentrated Alloys. Metals, 2020, 10, 1072.	2.3	23
9	Predicting vibrational entropy of fcc solids uniquely from bond chemistry using machine learning. Materialia, 2020, 12, 100804.	2.7	15
10	Effect of atomic order/disorder on vacancy clustering in concentrated NiFe alloys. Computational Materials Science, 2018, 147, 194-203.	3.0	9
11	Effect of atomic order/disorder on Cr segregation in Ni-Fe alloys. Journal of Applied Physics, 2018, 124,	2.5	4
12	Entropy contributions to phase stability in binary random solid solutions. Npj Computational Materials, 2018, 4, .	8.7	64
13	Segregation and binding energetics at grain boundaries in fluorite oxides. Journal of Materials Chemistry A, 2017, 5, 4026-4035.	10.3	24
14	Forging Fast Ion Conducting Nanochannels with Swift Heavy Ions: The Correlated Role of Local Electronic and Atomic Structure. Journal of Physical Chemistry C, 2017, 121, 975-981.	3.1	44
15	Cr-induced fast vacancy cluster formation and high Ni diffusion in concentrated Ni-Fe-Cr alloys. Journal of Alloys and Compounds, 2017, 725, 449-460.	5.5	23
16	Microstructure design for fast oxygen conduction. Journal of Materials Research, 2016, 31, 2-16.	2.6	22
17	Oxygen diffusion in ThO ₂ â€"CeO ₂ and ThO ₂ â€"UO ₂ solid solutions from atomistic calculations. Physical Chemistry Chemical Physics, 2016, 18, 15019-15024.	2.8	11
18	Influence of chemical disorder on energy dissipation and defect evolution in advanced alloys. Journal of Materials Research, 2016, 31, 2363-2375.	2.6	110

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19	Damage accumulation in ion-irradiated Ni-based concentrated solid-solution alloys. Acta Materialia, 2016, 109, 17-22.	7.9	114
20	Formation and growth of stacking fault tetrahedra in Ni via vacancy aggregation mechanism. Scripta Materialia, 2016, 114, 137-141.	5.2	42
21	Cation and vacancy disorder in U _{1â^'<i>y</i>} Nd _{<i>y</i>} 2.00â^' <i>x</i> alloys. Journal of Materials Research, 2015, 30, 3026-3040.	2.6	9
22	Fast ion conductivity in strained defect-fluorite structure created by ion tracks in Gd2Ti2O7. Scientific Reports, 2015, 5, 16297.	3.3	33
23	Chemical expansion affected oxygen vacancy stability in different oxide structures from first principles calculations. Computational Materials Science, 2015, 99, 298-305.	3.0	58
24	Point defect evolution in Ni, NiFe and NiCr alloys from atomistic simulations and irradiation experiments. Acta Materialia, 2015, 99, 69-76.	7.9	120
25	Radiation damage in cubic ZrO2 and yttria-stabilized zirconia from molecular dynamics simulations. Scripta Materialia, 2015, 98, 16-19.	5.2	20
26	Strain-Induced Phase and Oxygen-Vacancy Stability in Ionic Interfaces from First-Principles Calculations. Journal of Physical Chemistry C, 2014, 118, 30139-30144.	3.1	37
27	Impact of segregation energetics on oxygen conductivity at ionic grain boundaries. Journal of Materials Chemistry A, 2014, 2, 1704-1709.	10.3	31
28	The effect of electronic energy loss on irradiation-induced grain growth in nanocrystalline oxides. Physical Chemistry Chemical Physics, 2014, 16, 8051-8059.	2.8	62
29	Strained Ionic Interfaces: Effect on Oxygen Diffusivity from Atomistic Simulations. Journal of Physical Chemistry C, 2014, 118, 4207-4212.	3.1	27
30	(001) SrTiO ₃ (001) MgO Interface and Oxygen-Vacancy Stability from First-Principles Calculations. ACS Applied Materials & Samp; Interfaces, 2014, 6, 15536-15541.	8.0	9
31	The evolution mechanism of the dislocation loops in irradiated lanthanum doped cerium oxide. Journal of Nuclear Materials, 2014, 445, 209-217.	2.7	21
32	Investigation of oxygen point defects in cubic ZrO 2 by density functional theory. Computational Materials Science, 2014, 92, 22-27.	3.0	29
33	Stabilizing nanocrystalline grains in ceramic-oxides. Physical Chemistry Chemical Physics, 2013, 15, 18915.	2.8	11
34	On the Response of Ionic Crystals to Irradiation. Nuclear Technology, 2013, 182, 138-144.	1.2	2
35	Comparison of point-defect clustering in irradiated CeO2 and UO2: A unified view from molecular dynamics simulations and experiments. Scripta Materialia, 2011, 65, 867-870.	5.2	35
36	Effect of ionic polarizability on oxygen diffusion in Î-Bi2O3 from atomistic simulation. Ionics, 2010, 16, 297-303.	2.4	25

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37	Structure of Î-Bi2O3 from density functional theory: A systematic crystallographic analysis. Journal of Solid State Chemistry, 2009, 182, 1222-1228.	2.9	17
38	Grain-boundary source/sink behavior for point defects: An atomistic simulation study. International Journal of Materials Research, 2009, 100, 550-555.	0.3	28
39	Vacancyâ€Ordered Structure of Cubic Bismuth Oxide from Simulation and Crystallographic Analysis. Journal of the American Ceramic Society, 2008, 91, 2349-2356.	3.8	45