Garritt J Tucker

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#	Paper	IF	Citations
56	Spectral neighbor analysis method for automated generation of quantum-accurate interatomic potentials. <i>Journal of Computational Physics</i> , 2015 , 285, 316-330	4.1	330
55	Non-equilibrium grain boundary structure and inelastic deformation using atomistic simulations. <i>International Journal of Plasticity</i> , 2011 , 27, 841-857	7.6	100
54	Structure and free volume of <1 1 0> symmetric tilt grain boundaries with the E structural unit. <i>Acta Materialia</i> , 2007 , 55, 3959-3969	8.4	91
53	Atomistic simulations of tension moression asymmetry in dislocation nucleation for copper grain boundaries. <i>Computational Materials Science</i> , 2008 , 44, 351-362	3.2	79
52	Coarse-grained atomistic simulation of dislocations. <i>Journal of the Mechanics and Physics of Solids</i> , 2011 , 59, 160-177	5	77
51	Peierls potential of screw dislocations in bcc transition metals: Predictions from density functional theory. <i>Physical Review B</i> , 2013 , 87,	3.3	73
50	Evolution of structure and free volume in symmetric tilt grain boundaries during dislocation nucleation. <i>Acta Materialia</i> , 2010 , 58, 6464-6473	8.4	63
49	Quantifying the influence of twin boundaries on the deformation of nanocrystalline copper using atomistic simulations. <i>International Journal of Plasticity</i> , 2015 , 65, 191-205	7.6	58
48	A concurrent scheme for passing dislocations from atomistic to continuum domains. <i>Acta Materialia</i> , 2012 , 60, 899-913	8.4	58
47	Coarse-grained atomistic simulations of dislocations in Al, Ni and Cu crystals. <i>International Journal of Plasticity</i> , 2012 , 38, 86-101	7.6	56
46	Evidence for Bulk Ripplocations in Layered Solids. <i>Scientific Reports</i> , 2016 , 6, 33451	4.9	55
45	Stress-assisted grain growth in nanocrystalline metals: Grain boundary mediated mechanisms and stabilization through alloying. <i>Acta Materialia</i> , 2017 , 131, 39-47	8.4	49
44	Molecular dynamics simulations of rate-dependent grain growth during the surface indentation of nanocrystalline nickel. <i>Materials Science & Engineering A: Structural Materials: Properties, Microstructure and Processing,</i> 2013 , 571, 207-214	5.3	43
43	Investigating the deformation of nanocrystalline copper with microscale kinematic metrics and molecular dynamics. <i>Journal of the Mechanics and Physics of Solids</i> , 2012 , 60, 471-486	5	41
42	Spherical nanoindentation, modeling and transmission electron microscopy evidence for ripplocations in Ti3SiC2. <i>Acta Materialia</i> , 2017 , 131, 141-155	8.4	34
41	Synthesis and Surface Chemistry of 2D TiVC Solid-Solution MXenes. <i>ACS Applied Materials & Amp; Interfaces</i> , 2020 , 12, 20129-20137	9.5	34
40	Shear deformation kinematics of bicrystalline grain boundaries in atomistic simulations. <i>Modelling and Simulation in Materials Science and Engineering</i> , 2010 , 18, 015002	2	34

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39	Continuum metrics for deformation and microrotation from atomistic simulations: Application to grain boundaries. <i>International Journal of Engineering Science</i> , 2011 , 49, 1424-1434	5.7	33
38	Nanoindentation of monolayer Ti C T MXenes via atomistic simulations: The role of composition and defects on strength. <i>Computational Materials Science</i> , 2019 , 157, 168-174	3.2	29
37	Deformation of layered solids: Ripplocations not basal dislocations. <i>Scripta Materialia</i> , 2017 , 139, 166-1	73 .6	28
36	Achieving Radiation Tolerance through Non-Equilibrium Grain Boundary Structures. <i>Scientific Reports</i> , 2017 , 7, 12275	4.9	28
35	Interplay between Composition, Electronic Structure, Disorder, and Doping due to Dual Sublattice Mixing in Nonequilibrium Synthesis of ZnSnN:O. <i>Advanced Materials</i> , 2019 , 31, e1807406	24	27
34	Utilizing Site Disorder in the Development of New Energy-Relevant Semiconductors. <i>ACS Energy Letters</i> , 2020 , 5, 2027-2041	20.1	24
33	The mechanical behavior and deformation of bicrystalline nanowires. <i>Modelling and Simulation in Materials Science and Engineering</i> , 2013 , 21, 015004	2	22
32	Role of grain boundary character and its evolution on interfacial solute segregation behavior in nanocrystalline Ni-P. <i>Acta Materialia</i> , 2020 , 190, 113-123	8.4	20
31	Molecular dynamics studies of defect formation during heteroepitaxial growth of InGaN alloys on (0001) GaN surfaces. <i>Journal of Applied Physics</i> , 2017 , 121, 195301	2.5	19
30	Ripplocations: A universal deformation mechanism in layered solids. <i>Physical Review Materials</i> , 2019 , 3,	3.2	18
29	Mechanical properties of nanocrystalline nanoporous platinum. <i>Acta Materialia</i> , 2016 , 103, 624-632	8.4	16
28	Development of physically based atomistic microstructures: The effect on the mechanical response of polycrystals. <i>Computational Materials Science</i> , 2017 , 128, 29-36	3.2	16
27	Atomistic simulations of dislocation pinning points in pure face-centered-cubic nanopillars. <i>Modelling and Simulation in Materials Science and Engineering</i> , 2012 , 20, 075001	2	15
26	On the mechanistic origins of maximum strength in nanocrystalline metals. <i>Npj Computational Materials</i> , 2020 , 6,	10.9	11
25	Probing configurational disorder in ZnGeN2 using cluster-based Monte Carlo. <i>Physical Review Materials</i> , 2021 , 5,	3.2	10
24	Nucleation of ripplocations through atomistic modeling of surface nanoindentation in graphite. <i>Physical Review Materials</i> , 2018 , 2,	3.2	9
23	Quantifying grain boundary damage tolerance with atomistic simulations. <i>Modelling and Simulation in Materials Science and Engineering</i> , 2016 , 24, 075011	2	9
22	Perfect short-range ordered alloy with line-compound-like properties in the ZnSnN2:ZnO system. Npj Computational Materials, 2020, 6,	10.9	8

21	Bond-order potentials for the Ti3AlC2 and Ti3SiC2 MAX phases. <i>Physical Review B</i> , 2019 , 100,	3.3	8
20	Simulated defect growth avalanches during deformation of nanocrystalline copper. <i>Philosophical Magazine</i> , 2013 , 93, 478-498	1.6	7
19	Molecular dynamics studies of InGaN growth on nonpolar (112[0) GaN surfaces. <i>Physical Review Materials</i> , 2018 , 2,	3.2	7
18	Characterization of ripplocation mobility in graphite. <i>Materials Research Letters</i> , 2020 , 8, 82-87	7.4	6
17	Mechanical properties of stabilized nanocrystalline FCC metals. <i>Journal of Applied Physics</i> , 2019 , 126, 110901	2.5	5
16	The effect of hydrostatic pressure on the shear deformation of Cu symmetric tilt interfaces. <i>International Journal of Plasticity</i> , 2019 , 118, 87-104	7.6	5
15	On the origin of kinking in layered crystalline solids. <i>Materials Today</i> , 2021 , 43, 45-52	21.8	5
14	Grain-Size-Dependent Grain Boundary Deformation during Yielding in Nanocrystalline Materials Using Atomistic Simulations. <i>Jom</i> , 2020 , 72, 1745-1754	2.1	4
13	The microstructural and stress evolution in sputter deposited Ni thin films. <i>Surface and Coatings Technology</i> , 2021 , 412, 126973	4.4	3
12	Continuum Metrics for Atomistic Simulation Analysis. Springer Series in Materials Science, 2016 , 297-315	0.9	3
11	Effects of grain boundary structure on lithium transport in graphite. <i>Molecular Simulation</i> , 2016 , 42, 135	5 6 -136	33
10	Correlating deformation mechanisms with X-ray diffraction phenomena in nanocrystalline metals using atomistic simulations. <i>Computational Materials Science</i> , 2018 , 154, 178-186	3.2	2
9	Manipulation of solute partitioning mechanisms for nanocrystalline stability. <i>Acta Materialia</i> , 2021 , 208, 116662	8.4	2
8	Bandgap analysis and carrier localization in cation-disordered ZnGeN2. APL Materials, 2022, 10, 011112	5.7	1
7	Basal dislocations in MAX phases: Core structure and mobility. <i>Materialia</i> , 2022 , 21, 101310	3.2	1
6	Threshold Determination and Engaging Materials Scientists in Ontology Design. <i>Communications in Computer and Information Science</i> , 2015 , 39-50	0.3	1
5	Introduction to Atomistic Simulation Methods. Springer Series in Materials Science, 2016, 1-52	0.9	1
4	A rapid preparation method for in situ nanomechanical TEM tensile specimens. <i>Journal of Materials Research</i> , 2021 , 36, 2315-2324	2.5	1

LIST OF PUBLICATIONS

3	Computational Materials Design: Interplay between Composition, Electronic Structure, Disorder, and Doping due to Dual Sublattice Mixing in Nonequilibrium Synthesis of ZnSnN2:O (Adv. Mater. 11/2019). <i>Advanced Materials</i> , 2019 , 31, 1970080	24	О
2	Improved computational method to generate properly equilibrated atomistic microstructures. <i>MethodsX</i> , 2021 , 8, 101217	1.9	О
1	Probing fundamental deformation mechanisms and trends during decohesion across random high angle grain boundaries. <i>Computational Materials Science</i> , 2021 , 186, 110063	3.2	