Jess Carrete

List of Publications by Year in Descending Order

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

123
papers5,747
citations40
h-index74
g-index128
ext. papers6,935
ext. citations5
avg, IF6.21
L-index

#	Paper	IF	Citations
123	Influence of Onion-like Carbonaceous Particles on the Aggregation Process of Hydrocarbons. <i>ACS Omega</i> , 2021 , 6, 27898-27904	3.9	1
122	Ultrahigh Thermal Conductivity of Phase Tantalum Nitride. <i>Physical Review Letters</i> , 2021 , 126, 115901	7.4	16
121	How dopants limit the ultrahigh thermal conductivity of boron arsenide: a first principles study. <i>Npj Computational Materials</i> , 2021 , 7,	10.9	5
120	Effects of doping substitutions on the thermal conductivity of half-Heusler compounds. <i>Physical Review B</i> , 2021 , 103,	3.3	1
119	Quantum Self-Consistent Ab-Initio Lattice Dynamics. <i>Computer Physics Communications</i> , 2021 , 263, 1079	945	1
118	Thermoelectric properties of the SnS monolayer: Fully ab initio and accelerated calculations. <i>Journal of Applied Physics</i> , 2021 , 130, 054301	2.5	1
117	Strain-tunable lattice thermal conductivity of the Janus PtSTe monolayer. <i>Journal of Physics Condensed Matter</i> , 2021 , 34,	1.8	1
116	Growth, charge and thermal transport of flowered graphene. Carbon, 2020, 161, 259-268	10.4	4
115	Anisotropic Thermal Conductivity in Few-Layer and Bulk Titanium Trisulphide from First Principles. <i>Nanomaterials</i> , 2020 , 10,	5.4	5
114	Anomalously large lattice thermal conductivity in metallic tungsten carbide and its origin in the electronic structure. <i>Materials Today Physics</i> , 2020 , 13, 100214	8	11
113	High-throughput study of the static dielectric constant at high temperatures in oxide and fluoride cubic perovskites. <i>Physical Review Materials</i> , 2020 , 4,	3.2	3
112	Machine-learning Prediction of Infrared Spectra of Interstellar Polycyclic Aromatic Hydrocarbons. <i>Astrophysical Journal</i> , 2020 , 902, 100	4.7	6
111	The AFLOW Fleet for Materials Discovery 2020 , 1785-1812		1
110	Optimisation of the thermoelectric efficiency of zirconium trisulphide monolayers through unixial and biaxial strain. <i>Nanoscale Advances</i> , 2020 , 2, 5352-5361	5.1	6
109	Combined treatment of phonon scattering by electrons and point defects explains the thermal conductivity reduction in highly-doped Si. <i>Journal of Materials Chemistry A</i> , 2020 , 8, 1273-1278	13	12
108	Localized dimers drive strong anharmonicity and low lattice thermal conductivity in ZnSe2. <i>Physical Review B</i> , 2020 , 102,	3.3	4
107	Shortcomings of meta-GGA functionals when describing magnetism. <i>Physical Review B</i> , 2020 , 102,	3.3	9

(2018-2020)

106	The Effect of Janus Asymmetry on Thermal Transport in SnSSe. <i>Journal of Physical Chemistry C</i> , 2020 , 124, 17476-17484	3.8	9
105	Borophene vs. graphene interfaces: Tuning the electric double layer in ionic liquids. <i>Journal of Molecular Liquids</i> , 2020 , 303, 112647	6	7
104	Twisted bilayer graphene as a linear nanoactuator. <i>Physical Review B</i> , 2020 , 102,	3.3	2
103	Phonon Scattering by Dislocations in GaN. ACS Applied Materials & amp; Interfaces, 2019, 11, 8175-8181	9.5	18
102	Coupling of Spinons with Defects and Phonons in the Spin Chain Compound Ca_{2}CuO_{3}. <i>Physical Review Letters</i> , 2019 , 122, 185901	7.4	4
101	Tunable gap in stable arsenene nanoribbons opens the door to electronic applications <i>RSC Advances</i> , 2019 , 9, 11818-11823	3.7	2
100	The AFLOW Fleet for Materials Discovery 2019 , 1-28		
99	Thermal Resistance of GaN/AlN Graded Interfaces. <i>Physical Review Applied</i> , 2019 , 11,	4.3	15
98	Using nanotubes to study the phonon spectrum of two-dimensional materials. <i>Physical Chemistry Chemical Physics</i> , 2019 , 21, 5215-5223	3.6	2
97	Phonon transport across crystal-phase interfaces and twin boundaries in semiconducting nanowires. <i>Nanoscale</i> , 2019 , 11, 16007-16016	7.7	7
96	Parameter-free model to estimate thermal conductivity in nanostructured materials. <i>Physical Review B</i> , 2019 , 100,	3.3	3
95	Effect of local chemistry and structure on thermal transport in doped GaAs. <i>Physical Review Materials</i> , 2019 , 3,	3.2	5
94	An investigation of the structural properties of Li and Na fast ion conductors using high-throughput bond-valence calculations and machine learning. <i>Journal of Applied Crystallography</i> , 2019 , 52, 148-157	3.8	21
93	A theoretical model of the thermoelectric properties of SnSxSe1N and how to further enhance its thermoelectric performance. <i>Journal of Applied Physics</i> , 2019 , 126, 225105	2.5	12
92	Resonant phonon scattering in semiconductors. <i>Journal of Materials Chemistry C</i> , 2018 , 6, 4691-4697	7.1	13
91	Predictive Design and Experimental Realization of InAs/GaAs Superlattices with Tailored Thermal Conductivity. <i>Journal of Physical Chemistry C</i> , 2018 , 122, 4054-4062	3.8	12
90	Independently tuning the power factor and thermal conductivity of SnSe via Ag2S addition and nanostructuring. <i>Journal of Materials Chemistry A</i> , 2018 , 6, 7959-7966	13	17
89	Materials Screening for the Discovery of New Half-Heuslers: Machine Learning versus ab Initio Methods. <i>Journal of Physical Chemistry B</i> , 2018 , 122, 625-632	3.4	60

88	Influence of point defects on the thermal conductivity in FeSi. Physical Review B, 2018, 97,	3.3	15
87	Ab initio lattice thermal conductivity of bulk and thin-film AI2O3. MRS Communications, 2018, 8, 1119-1	12273	10
86	Phonon transport unveils the prevalent point defects in GaN. <i>Physical Review Materials</i> , 2018 , 2,	3.2	17
85	Thermal transport through Ge-rich Ge/Si superlattices grown on Ge(0 0 1). <i>Journal Physics D: Applied Physics</i> , 2018 , 51, 014001	3	19
84	The AFLOW Fleet for Materials Discovery 2018 , 1-28		9
83	First-principles quantitative prediction of the lattice thermal conductivity in random semiconductor alloys: The role of force-constant disorder. <i>Physical Review B</i> , 2018 , 98,	3.3	24
82	Vibrational Properties of Metastable Polymorph Structures by Machine Learning. <i>Journal of Chemical Information and Modeling</i> , 2018 , 58, 2460-2466	6.1	7
81	BoltzTraP2, a program for interpolating band structures and calculating semi-classical transport coefficients. <i>Computer Physics Communications</i> , 2018 , 231, 140-145	4.2	320
80	AFLOW-ML: A RESTful API for machine-learning predictions of materials properties. <i>Computational Materials Science</i> , 2018 , 152, 134-145	3.2	51
79	Density Functional Study of Charge Transfer at the Graphene/Ionic Liquid Interface. <i>Journal of Physical Chemistry C</i> , 2018 , 122, 15070-15077	3.8	9
78	Understanding Phonon Scattering by Nanoprecipitates in Potassium-Doped Lead Chalcogenides. <i>ACS Applied Materials & Doped Lead Chalcogenides</i> . 3686-3693	9.5	4
77	How Chemical Composition Alone Can Predict Vibrational Free Energies and Entropies of Solids. <i>Chemistry of Materials</i> , 2017 , 29, 6220-6227	9.6	69
76	Human Immune Protein C1q Selectively Disaggregates Carbon Nanotubes. <i>Nano Letters</i> , 2017 , 17, 3409	-34.55	12
75	Structural Complexity and Phonon Physics in 2D Arsenenes. <i>Journal of Physical Chemistry Letters</i> , 2017 , 8, 1375-1380	6.4	34
74	What will freestanding borophene nanoribbons look like? An analysis of their possible structures, magnetism and transport properties. <i>Physical Chemistry Chemical Physics</i> , 2017 , 19, 1054-1061	3.6	28
73	An efficient and accurate framework for calculating lattice thermal conductivity of solids: AFLOWAAPL Automatic Anharmonic Phonon Library. <i>Npj Computational Materials</i> , 2017 , 3,	10.9	46
72	Exceptionally Strong Phonon Scattering by B Substitution in Cubic SiC. <i>Physical Review Letters</i> , 2017 , 119, 075902	7.4	53
71	Influence of Antisite Defects on the Thermoelectric Properties of Fe2VAl. <i>Nanoscale and Microscale Thermophysical Engineering</i> , 2017 , 21, 237-246	3.7	13

(2015-2017)

70	Phonon thermal transport in 2H, 4H and 6H silicon carbide from first principles. <i>Materials Today Physics</i> , 2017 , 1, 31-38	8	29
69	almaBTE: A solver of the spacelime dependent Boltzmann transport equation for phonons in structured materials. <i>Computer Physics Communications</i> , 2017 , 220, 351-362	4.2	124
68	Ab initio phonon scattering by dislocations. <i>Physical Review B</i> , 2017 , 95,	3.3	43
67	Temperature and Thickness Dependences of the Anisotropic In-Plane Thermal Conductivity of Black Phosphorus. <i>Advanced Materials</i> , 2017 , 29, 1603756	24	75
66	Glass-like thermal conductivity in nanostructures of a complex anisotropic crystal. <i>Physical Review B</i> , 2017 , 96,	3.3	5
65	Influence of the optical-acoustic phonon hybridization on phonon scattering and thermal conductivity. <i>Physical Review B</i> , 2016 , 93,	3.3	33
64	Optimizing phonon scattering by nanoprecipitates in lead chalcogenides. <i>Applied Physics Letters</i> , 2016 , 108, 113901	3.4	4
63	Structural and electronic properties of zigzag InP nanoribbons with Stone-Wales type defects. Journal of Physics Condensed Matter, 2016 , 28, 065503	1.8	O
62	Physically founded phonon dispersions of few-layer materials and the case of borophene. <i>Materials Research Letters</i> , 2016 , 4, 204-211	7.4	158
61	Cross-plane heat conduction in thin films with ab-initio phonon dispersions and scattering rates. <i>Applied Physics Letters</i> , 2016 , 108, 193104	3.4	36
60	High-Throughput Computation of Thermal Conductivity of High-Temperature Solid Phases: The Case of Oxide and Fluoride Perovskites. <i>Physical Review X</i> , 2016 , 6,	9.1	41
59	Unraveling the dominant phonon scattering mechanism in the thermoelectric compound ZrNiSn. <i>Journal of Materials Chemistry A</i> , 2016 , 4, 15940-15944	13	36
58	Ultralow lattice thermal conductivity in topological insulator TlBiSe2. <i>Applied Physics Letters</i> , 2016 , 108, 233902	3.4	19
57	Basal-plane thermal conductivity of nanocrystalline and amorphized thin germanane. <i>Applied Physics Letters</i> , 2016 , 109, 131907	3.4	9
56	Anomalous thermal conductivity and suppression of negative thermal expansion in ScF3. <i>Physical Review B</i> , 2016 , 94,	3.3	41
55	Ab initio study of the effect of vacancies on the thermal conductivity of boron arsenide. <i>Physical Review B</i> , 2016 , 94,	3.3	53
54	Microstructure investigations and thermoelectrical properties of an N-type magnesium lilicon lin alloy sintered from a gas-phase atomized powder. <i>Acta Materialia</i> , 2015 , 96, 437-451	8.4	14
53	Twisting phonons in complex crystals with quasi-one-dimensional substructures. <i>Nature Communications</i> , 2015 , 6, 6723	17.4	52

52	Influence of the addition of Half-Heusler nanoparticles on the thermoelectrical properties of an N-type magnesium ilicon in alloy. <i>Scripta Materialia</i> , 2015 , 104, 5-8	5.6	5
51	Solvation of molecular cosolvents and inorganic salts in ionic liquids: A review of molecular dynamics simulations. <i>Journal of Molecular Liquids</i> , 2015 , 210, 178-188	6	70
50	Anomalous pressure dependence of thermal conductivities of large mass ratio compounds. <i>Physical Review B</i> , 2015 , 91,	3.3	44
49	Role of force-constant difference in phonon scattering by nano-precipitates in PbTe. <i>Journal of Applied Physics</i> , 2015 , 118, 085701	2.5	12
48	Structural, magnetic, and vibrational properties of stoichiometric clusters of CrN. <i>International Journal of Quantum Chemistry</i> , 2015 , 115, 523-528	2.1	2
47	How does lithium nitrate dissolve in a protic ionic liquid?. <i>Journal of Molecular Liquids</i> , 2015 , 205, 16-21	6	50
46	Superdiffusive heat conduction in semiconductor alloys. I. Theoretical foundations. <i>Physical Review B</i> , 2015 , 91,	3.3	59
45	Ti-decorated zigzag graphene nanoribbons for hydrogen storage. A van der Waals-corrected density-functional study. <i>International Journal of Hydrogen Energy</i> , 2015 , 40, 4960-4968	6.7	48
44	Nanostructure of mixtures of protic ionic liquids and lithium salts: effect of alkyl chain length. <i>Physical Chemistry Chemical Physics</i> , 2015 , 17, 5298-307	3.6	34
43	Complex Network Analysis in Socioeconomic Models. <i>Dynamic Modeling and Econometrics in Economics and Finance</i> , 2015 , 209-245		7
42	Solvation of lithium salts in protic ionic liquids: a molecular dynamics study. <i>Journal of Physical Chemistry B</i> , 2014 , 118, 761-70	3.4	74
41	Effect of nitrogen and vacancy defects on the thermal conductivity of diamond: An ab initio Greenß function approach. <i>Physical Review B</i> , 2014 , 90,	3.3	72
40	Low thermal conductivity and triaxial phononic anisotropy of SnSe. <i>Applied Physics Letters</i> , 2014 , 105, 101907	3.4	200
39	Unexpected high-temperature stability of EZn4Sb3 opens the door to enhanced thermoelectric performance. <i>Journal of the American Chemical Society</i> , 2014 , 136, 1497-504	16.4	97
38	Molecular dynamics simulations of the structure of the graphene-ionic liquid/alkali salt mixtures interface. <i>Physical Chemistry Chemical Physics</i> , 2014 , 16, 13271-8	3.6	54
37	Nanograined Half-Heusler Semiconductors as Advanced Thermoelectrics: An Ab Initio High-Throughput Statistical Study. <i>Advanced Functional Materials</i> , 2014 , 24, 7427-7432	15.6	97
36	Phonon thermal transport in strained and unstrained graphene from first principles. <i>Physical Review B</i> , 2014 , 89,	3.3	270
35	ShengBTE: A solver of the Boltzmann transport equation for phonons. <i>Computer Physics Communications</i> , 2014 , 185, 1747-1758	4.2	1243

(2011-2014)

34	Strong enhancement of phonon scattering through nanoscale grains in lead sulfide thermoelectrics. NPG Asia Materials, 2014 , 6, e108-e108	10.3	119
33	Mixtures of protic ionic liquids and molecular cosolvents: a molecular dynamics simulation. <i>Journal of Chemical Physics</i> , 2014 , 140, 214502	3.9	69
32	Finding Unprecedentedly Low-Thermal-Conductivity Half-Heusler Semiconductors via High-Throughput Materials Modeling. <i>Physical Review X</i> , 2014 , 4,	9.1	168
31	Electronic properties of pure and p-type doped hexagonal sheets and zigzag nanoribbons of InP. <i>Journal of Physics Condensed Matter</i> , 2013 , 25, 085506	1.8	2
30	Molecular hydrogen uptake by zigzag graphene nanoribbons doped with early 3d transition-metal atoms. <i>International Journal of Hydrogen Energy</i> , 2013 , 38, 8872-8880	6.7	17
29	Thermal conductivity and phonon linewidths of monolayer MoS2 from first principles. <i>Applied Physics Letters</i> , 2013 , 103, 253103	3.4	236
28	Atomistic origin of glass-like Zn4Sb3 thermal conductivity. <i>Applied Physics Letters</i> , 2013 , 103, 103902	3.4	14
27	MD simulations of the formation of stable clusters in mixtures of alkaline salts and imidazolium-based ionic liquids. <i>Journal of Physical Chemistry B</i> , 2013 , 117, 3207-20	3.4	77
26	Thermal Conductivity of Ionic Liquids: A Pseudolattice Approach. <i>Journal of Physical Chemistry C</i> , 2012 , 116, 1265-1273	3.8	14
25	Investigation of the local structure of mixtures of an ionic liquid with polar molecular species through molecular dynamics: cluster formation and angular distributions. <i>Journal of Physical Chemistry B</i> , 2012 , 116, 5941-50	3.4	22
24	Effect of temperature and cationic chain length on the physical properties of ammonium nitrate-based protic ionic liquids. <i>Journal of Physical Chemistry B</i> , 2012 , 116, 11302-12	3.4	77
23	Thermoelectric Properties of Hybrid OrganicIhorganic Superlattices. <i>Journal of Physical Chemistry C</i> , 2012 , 116, 10881-10886	3.8	23
22	Liquid Bolid Dquid phase transition hysteresis loops in the ionic conductivity of ten imidazolium-based ionic liquids. <i>Fluid Phase Equilibria</i> , 2012 , 320, 1-10	2.5	54
21	Al enhances the H2 storage capacity of graphene at nanoribbon borders but not at central sites: A study using nonlocal van der Waals density functionals. <i>Physical Review B</i> , 2012 , 85,	3.3	24
20	Molecular dynamics simulations of the structural and thermodynamic properties of imidazolium-based ionic liquid mixtures. <i>Journal of Physical Chemistry B</i> , 2011 , 115, 11170-82	3.4	51
19	Molecular dynamics simulation of the structure and dynamics of water-1-alkyl-3-methylimidazolium ionic liquid mixtures. <i>Journal of Physical Chemistry B</i> , 2011 , 115, 6995-7008	3.4	111
18	Magnetism of substitutional Fe impurities in graphene nanoribbons. <i>Journal of Chemical Physics</i> , 2011 , 134, 024704	3.9	26
17	Prediction of phonon thermal transport in thin GaAs, InAs and InP nanowires by molecular dynamics simulations: influence of the interatomic potential. <i>Nanotechnology</i> , 2011 , 22, 185704	3.4	10

16	Dynamical properties of alcohol + 1-hexyl-3-methylimidazolium ionic liquid mixtures: a computer simulation study. <i>Journal of Physical Chemistry B</i> , 2011 , 115, 15313-22	3.4	31
15	Pseudolattice Theory of Ionic Liquids 2011 ,		1
14	Theoretical model for moisture adsorption on ionic liquids: A modified Brunauer Emmet Teller isotherm approach. <i>Fluid Phase Equilibria</i> , 2011 , 301, 118-122	2.5	13
13	Re-entrant phase behavior for systems with competition between phase separation and self-assembly. <i>Journal of Chemical Physics</i> , 2011 , 134, 104905	3.9	30
12	Surfactant self-assembly nanostructures in protic ionic liquids. <i>Journal of Physical Chemistry B</i> , 2011 , 115, 8145-54	3.4	40
11	Ab initio study of 3d, 4d, and 5d transition metal adatoms and dimers adsorbed on hydrogen-passivated zigzag graphene nanoribbons. <i>Physical Review B</i> , 2011 , 83,	3.3	45
10	Surface roughness and thermal conductivity of semiconductor nanowires: Going below the Casimir limit. <i>Physical Review B</i> , 2011 , 84,	3.3	70
9	Structural, magnetic, and electronic properties of Nin and Fen nanostructures (n=1월) adsorbed on zigzag graphene nanoribbons. <i>Physical Review B</i> , 2010 , 81,	3.3	41
8	Pseudolattice theory of charge transport in ionic solutions: Corresponding states law for the electric conductivity. <i>Fluid Phase Equilibria</i> , 2010 , 298, 280-286	2.5	43
7	A density-functional study of the vertical ionization potentials of the cluster Mn13. <i>Journal of Chemical Physics</i> , 2009 , 131, 046101	3.9	5
6	A density-functional study of the structures and electronic properties of neutral, anionic, and endohedrally doped In(x)P(x) clusters. <i>Journal of Chemical Physics</i> , 2009 , 131, 074504	3.9	7
5	Pseudolattice theory of the surface tension of ionic liquid-water mixtures. <i>Journal of Physical Chemistry B</i> , 2009 , 113, 12500-5	3.4	27
4	Molecular-dynamics prediction of the thermal conductivity of thin InP nanowires: Similarities to Si. <i>Physical Review B</i> , 2009 , 80,	3.3	12
3	Nonequilibrium nanothermodynamics. <i>Physical Review E</i> , 2008 , 77, 022102	2.4	5
2	Microcanonical equations obtained from the Tsallis entropy. <i>Physica A: Statistical Mechanics and Its Applications</i> , 2008 , 387, 6752-6758	3.3	4
1	Nonextensive statistical mechanics of ionic solutions. <i>Physics Letters, Section A: General, Atomic and Solid State Physics</i> , 2007 , 370, 405-412	2.3	13