Luis G Macdowell

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.

3,213 100 33 53 h-index g-index citations papers 103 3,512 4.5 5.33 avg, IF L-index ext. citations ext. papers

#	Paper	IF	Citations
100	Lifshitz theory of wetting films at three phase coexistence: The case of ice nucleation on Silver Iodide (AgI). <i>Journal of Colloid and Interface Science</i> , 2021 , 590, 527-538	9.3	2
99	Crystal growth of bcc titanium from the melt and interfacial properties: A molecular dynamics simulation study. <i>Journal of Chemical Physics</i> , 2021 , 154, 184704	3.9	0
98	How ice grows from premelting films and water droplets. <i>Nature Communications</i> , 2021 , 12, 239	17.4	8
97	Micelle-directed chiral seeded growth on anisotropic gold nanocrystals. <i>Science</i> , 2020 , 368, 1472-1477	33.3	78
96	Surface phase transitions and crystal habits of ice in the atmosphere. Science Advances, 2020, 6, eaay93	22 4.3	16
95	Rounded Layering Transitions on the Surface of Ice. <i>Physical Review Letters</i> , 2020 , 124, 065702	7.4	8
94	Premelting of ice adsorbed on a rock surface. <i>Physical Chemistry Chemical Physics</i> , 2020 , 22, 11362-1137	73 3.6	7
93	Structure and water attachment rates of ice in the atmosphere: role of nitrogen. <i>Physical Chemistry Chemical Physics</i> , 2019 , 21, 19594-19611	3.6	6
92	Structure and fluctuations of the premelted liquid film of ice at the triple point. <i>Molecular Physics</i> , 2019 , 117, 2846-2864	1.7	6
91	Disconnecting Symmetry Breaking from Seeded Growth for the Reproducible Synthesis of High Quality Gold Nanorods. <i>ACS Nano</i> , 2019 , 13, 4424-4435	16.7	59
90	Structural transitions and bilayer formation of CTAB aggregates. <i>Colloids and Surfaces A: Physicochemical and Engineering Aspects</i> , 2019 , 580, 123730	5.1	9
89	Surface van der Waals forces in a nutshell. <i>Journal of Chemical Physics</i> , 2019 , 150, 081101	3.9	10
88	Nanocapillarity and Liquid Bridge-Mediated Force between Colloidal Nanoparticles. <i>ACS Omega</i> , 2018 , 3, 112-123	3.9	8
87	The vicinity of an equilibrium three-phase contact line using density-functional theory: density profiles normal to the fluid interface. <i>Molecular Physics</i> , 2018 , 116, 2239-2243	1.7	2
86	Vapour-liquid interfacial properties of square-well chains from density functional theory and Monte Carlo simulation. <i>Physical Chemistry Chemical Physics</i> , 2017 , 19, 12296-12309	3.6	10
85	Femtosecond laser reshaping yields gold nanorods with ultranarrow surface plasmon resonances. <i>Science</i> , 2017 , 358, 640-644	33.3	176
84	Capillary wave theory of adsorbed liquid films and the structure of the liquid-vapor interface. <i>Physical Review E</i> , 2017 , 96, 022801	2.4	9

83	Premelting-Induced Smoothening of the Ice-Vapor Interface. <i>Physical Review Letters</i> , 2016 , 117, 09610	1 7.4	29
82	Analytic perturbative FMSA equation of state and thermodynamic properties from Monte Carlo simulation of the Kihara potential with a spherical core. <i>Molecular Physics</i> , 2015 , 113, 1076-1090	1.7	6
81	Interfacial free energy of the NaCl crystal-melt interface from capillary wave fluctuations. <i>Journal of Chemical Physics</i> , 2015 , 142, 134706	3.9	8
80	Mesoscopic Hamiltonian for the fluctuations of adsorbed Lennard-Jones liquid films. <i>Physical Review E</i> , 2015 , 91, 062404	2.4	4
79	A study of the ice-water interface using the TIP4P/2005 water model. <i>Physical Chemistry Chemical Physics</i> , 2014 , 16, 22159-66	3.6	36
78	Disjoining Pressure, Healing Distance, and Film Height Dependent Surface Tension of Thin Wetting Films. <i>Journal of Physical Chemistry C</i> , 2014 , 118, 22079-22089	3.8	14
77	Disjoining pressure and the film-height-dependent surface tension of thin liquid films: new insight from capillary wave fluctuations. <i>Advances in Colloid and Interface Science</i> , 2014 , 206, 150-71	14.3	33
76	Effect of molecular flexibility of Lennard-Jones chains on vapor-liquid interfacial properties. Journal of Chemical Physics, 2014 , 140, 114705	3.9	11
75	Computer simulation study of surface wave dynamics at the crystal-melt interface. <i>Journal of Chemical Physics</i> , 2014 , 141, 034701	3.9	22
74	Damped reaction field method and the accelerated convergence of the real space Ewald summation. <i>Journal of Chemical Physics</i> , 2014 , 141, 164108	3.9	4
73	Capillary fluctuations and film-height-dependent surface tension of an adsorbed liquid film.		
73	Physical Review Letters, 2013 , 111, 047802	7.4	25
72		7·4 3·9	19
	Physical Review Letters, 2013, 111, 047802 Vapor-liquid interfacial properties of rigid-linear Lennard-Jones chains. Journal of Chemical Physics,		
72	Physical Review Letters, 2013, 111, 047802 Vapor-liquid interfacial properties of rigid-linear Lennard-Jones chains. Journal of Chemical Physics, 2012, 137, 084706 Universal scaling behaviour of surface tension of molecular chains. Journal of Chemical Physics,	3.9	19
72 71	Physical Review Letters, 2013, 111, 047802 Vapor-liquid interfacial properties of rigid-linear Lennard-Jones chains. Journal of Chemical Physics, 2012, 137, 084706 Universal scaling behaviour of surface tension of molecular chains. Journal of Chemical Physics, 2012, 137, 024702 Semi-infinite boundary conditions for the simulation of interfaces: the Ar/CO2(s) model revisited.	3.9	19
7 ² 7 ¹	 Physical Review Letters, 2013, 111, 047802 Vapor-liquid interfacial properties of rigid-linear Lennard-Jones chains. Journal of Chemical Physics, 2012, 137, 084706 Universal scaling behaviour of surface tension of molecular chains. Journal of Chemical Physics, 2012, 137, 024702 Semi-infinite boundary conditions for the simulation of interfaces: the Ar/CO2(s) model revisited. Journal of Chemical Physics, 2012, 136, 104703 Computer simulation of interface potentials: Towards a first principle description of complex 	3.9 3.9 3.9	19 17 19
7 ² 7 ¹ 7 ⁰ 69	Vapor-liquid interfacial properties of rigid-linear Lennard-Jones chains. <i>Journal of Chemical Physics</i> , 2012, 137, 084706 Universal scaling behaviour of surface tension of molecular chains. <i>Journal of Chemical Physics</i> , 2012, 137, 024702 Semi-infinite boundary conditions for the simulation of interfaces: the Ar/CO2(s) model revisited. <i>Journal of Chemical Physics</i> , 2012, 136, 104703 Computer simulation of interface potentials: Towards a first principle description of complex interfaces?. <i>European Physical Journal: Special Topics</i> , 2011, 197, 131-145 Discussion notes on Disjoining pressure of planar adsorbed films[by J.R. Henderson. <i>European</i>	3.9 3.9 3.9	19 17 19 20

65	Solvation effects for polymers at an interface: a hybrid self-consistent field-density functional theory approach. <i>Journal of Chemical Physics</i> , 2011 , 135, 204901	3.9	12
64	Phase diagram of water under an applied electric field. <i>Physical Review Letters</i> , 2011 , 107, 155702	7.4	41
63	Liquid Vapor Phase Equilibria and Surface Tension of Ethane As Predicted by the TraPPE and OPLS Models (Journal of Chemical & Engineering Data, 2010, 55, 5465-5470)	2.8	15
62	Dielectric constant of ice Ih and ice V: a computer simulation study. <i>Journal of Physical Chemistry B</i> , 2010 , 114, 6089-98	3.4	29
61	Coarse-grained models for fluids and their mixtures: Comparison of Monte Carlo studies of their phase behavior with perturbation theory and experiment. <i>Journal of Chemical Physics</i> , 2009 , 130, 04410	13 .9	49
60	Coarse-graining dipolar interactions in simple fluids and polymer solutions: Monte Carlo studies of the phase behavior. <i>Physical Chemistry Chemical Physics</i> , 2009 , 11, 1923-33	3.6	10
59	Surface tension of fully flexible Lennard-Jones chains: role of long-range corrections. <i>Journal of Chemical Physics</i> , 2009 , 131, 074705	3.9	41
58	Towards the Quantitative Prediction of the Phase Behavior of Polymer Solutions by Computer Simulation. <i>Macromolecular Symposia</i> , 2009 , 278, 1-9	0.8	5
57	Efficient prediction of thermodynamic properties of quadrupolar fluids from simulation of a coarse-grained model: the case of carbon dioxide. <i>Journal of Chemical Physics</i> , 2008 , 128, 104501	3.9	44
56	Self-consistent field/density functional study of conformational properties of polymers at interfaces: role of intramolecular interactions. <i>Journal of Chemical Physics</i> , 2008 , 129, 104901	3.9	18
55	Vapor-liquid interfacial properties of fully flexible Lennard-Jones chains. <i>Journal of Chemical Physics</i> , 2008 , 129, 144703	3.9	69
54	Ice: A fruitful source of information about liquid water. <i>Journal of Molecular Liquids</i> , 2007 , 136, 214-220	6	14
53	Direct calculation of interfacial tensions from computer simulation: results for freely jointed tangent hard sphere chains. <i>Physical Review E</i> , 2007 , 75, 061609	2.4	45
52	Adsorption of polymers on a brush: Tuning the order of the wetting phase transition. <i>Journal of Chemical Physics</i> , 2006 , 124, 084907	3.9	58
51	Computer simulation of two new solid phases of water: Ice XIII and ice XIV. <i>Journal of Chemical Physics</i> , 2006 , 125, 116101	3.9	15
50	Nucleation and cavitation of spherical, cylindrical, and slablike droplets and bubbles in small systems. <i>Journal of Chemical Physics</i> , 2006 , 125, 34705	3.9	116
49	Can simple models describe the phase diagram of water?. <i>Journal of Physics Condensed Matter</i> , 2005 , 17, S3283-S3288	1.8	62
48	The range of meta stability of ice-water melting for two simple models of water. <i>Molecular Physics</i> , 2005 , 103, 1-5	1.7	50

(2003-2005)

47	Polymer + Solvent Systems: Phase Diagrams, Interface Free Energies, and Nucleation. <i>Advances in Polymer Science</i> , 2005 , 1-110	1.3	61
46	Observation of autophobic dewetting on polymer brushes from computer simulation. <i>Journal of Physics Condensed Matter</i> , 2005 , 17, S3523-S3528	1.8	24
45	Tracing the phase diagram of the four-site water potential (TIP4P). <i>Journal of Chemical Physics</i> , 2004 , 121, 1165-6	3.9	68
44	Combinatorial entropy and phase diagram of partially ordered ice phases. <i>Journal of Chemical Physics</i> , 2004 , 121, 10145-58	3.9	50
43	Phase behavior of n-alkanes in supercritical solution: a Monte Carlo study. <i>Journal of Chemical Physics</i> , 2004 , 121, 2169-79	3.9	88
42	Molecular modeling of flexible molecules. Vaporliquid and fluidBolid equilibria. <i>Journal of Molecular Liquids</i> , 2004 , 113, 37-51	6	11
41	Computer simulation study of the global phase behavior of linear rigid Lennard-Jones chain molecules: comparison with flexible models. <i>Journal of Chemical Physics</i> , 2004 , 120, 3957-68	3.9	25
40	Phase diagram of water from computer simulation. <i>Physical Review Letters</i> , 2004 , 92, 255701	7.4	245
39	The evaporation/condensation transition of liquid droplets. <i>Journal of Chemical Physics</i> , 2004 , 120, 529	3-33-98	133
38	Phase separation kinetics in compressible polymer solutions: computer simulation of the early stages. <i>New Journal of Physics</i> , 2004 , 6, 7-7	2.9	23
37	The Droplet Evaporation/Condensation Transition in a Finite Volume. <i>Springer Proceedings in Physics</i> , 2004 , 129-133	0.2	
36	Wetting of polymer liquids: Monte Carlo simulations and self-consistent field calculations. <i>Journal of Physics Condensed Matter</i> , 2003 , 15, R609-R653	1.8	49
35	Third virial coefficients and critical properties of quadrupolar two center Lennard-Jones models. <i>Physical Chemistry Chemical Physics</i> , 2003 , 5, 2851-2857	3.6	17
34	Critical properties of molecular fluids from the virial series. <i>Journal of Chemical Physics</i> , 2003 , 119, 1136	57 5 .1 ₉ 13	73 7
33	Short chains at surfaces and interfaces: A quantitative comparison between density-functional theories and Monte Carlo simulations. <i>Journal of Chemical Physics</i> , 2003 , 118, 2929	3.9	96
32	Formal study of nucleation as described by fluctuation theory. <i>Journal of Chemical Physics</i> , 2003 , 119, 453-463	3.9	19
31	Bonded hard-sphere theory and computer simulation of the equation of state of linear fused ard-sphere fluids. <i>Journal of Chemical Physics</i> , 2003 , 119, 9633-9639	3.9	8
30	How Do Droplets Depend on the System Size? Droplet Condensation and Nucleation in Small Simulation Cells 2003 , 125-135		

29	How do droplets on a surface depend on the system size?. <i>Colloids and Surfaces A: Physicochemical and Engineering Aspects</i> , 2002 , 206, 277-291	5.1	33
28	Phase diagrams of hexadecane IO2 mixtures from histogram-reweighting Monte Carlo. <i>Computer Physics Communications</i> , 2002 , 147, 378-381	4.2	24
27	Interface properties and bubble nucleation in compressible mixtures containing polymers. <i>Journal of Chemical Physics</i> , 2002 , 117, 5480-5496	3.9	60
26	A computer simulation study of racemic mixtures. <i>Molecular Physics</i> , 2002 , 100, 2397-2415	1.7	10
25	Critical lines and phase coexistence of polymer solutions: A quantitative comparison between Wertheim thermodynamic perturbation theory and computer simulations. <i>Journal of Chemical Physics</i> , 2002 , 117, 6360-6371	3.9	31
24	The effect of flexibility on the phase diagram of simple molecular models. <i>Physical Chemistry Chemical Physics</i> , 2002 , 4, 853-862	3.6	18
23	Wetting of a short chain liquid on a brush: First-order and critical wetting transitions. <i>Europhysics Letters</i> , 2001 , 55, 221-227	1.6	36
22	Isotropic-nematic phase transition: Influence of intramolecular flexibility using a fused hard sphere model. <i>Physical Review E</i> , 2001 , 64, 011703	2.4	44
21	Extending Wertheim perturbation theory to the solid phase: The freezing of the pearl-necklace model. <i>Journal of Chemical Physics</i> , 2001 , 114, 10411-10418	3.9	44
20	Liquid crystal phase formation for the linear tangent hard sphere model from Monte Carlo simulations. <i>Journal of Chemical Physics</i> , 2001 , 115, 4203-4211	3.9	45
19	Nano-dewetting: Interplay between van der Waals- and short-ranged interactions. <i>Journal of Chemical Physics</i> , 2001 , 115, 9960-9969	3.9	67
18	Equation of state of model branched alkanes: Theoretical predictions and configurational bias Monte Carlo simulations. <i>Journal of Chemical Physics</i> , 2001 , 115, 6220-6235	3.9	14
17	Critical temperature of infinitely long chains from Wertheim's perturbation theory. <i>Molecular Physics</i> , 2000 , 98, 1295-1308	1.7	28
16	The virial coefficients of the pearl-necklace model. <i>Journal of Chemical Physics</i> , 2000 , 113, 10398-10409	3.9	23
15	Equation of state and critical behavior of polymer models: A quantitative comparison between Wertheim thermodynamic perturbation theory and computer simulations. <i>Journal of Chemical Physics</i> , 2000 , 113, 419-433	3.9	72
14	Interface and Surface Properties of Short Polymers in Solution: Monte Carlo Simulations and Self-Consistent Field Theory. <i>Macromolecules</i> , 2000 , 33, 3902-3923	5.5	127
13	Excess properties of mixtures of n-alkanes from perturbation theory. <i>Journal of Chemical Physics</i> , 1999 , 111, 3192-3202	3.9	7
12	On the calculation of the frequency sum rules of the heat flux correlation function. <i>Molecular Physics</i> , 1999 , 96, 881-884	1.7	1

LIST OF PUBLICATIONS

11	Critical properties of mixtures of alkanes from perturbation theory. <i>Journal of Chemical Physics</i> , 1999 , 111, 3183-3191	3.9	8	
10	RESEARCH NOTE On the calculation of the frequency sum rules of the heat flux correlation function. <i>Molecular Physics</i> , 1999 , 96, 881-884	1.7	2	
9	Vaporliquid equilibria of linear and branched alkanes from perturbation theory. <i>Journal of Chemical Physics</i> , 1998 , 109, 5681-5690	3.9	14	
8	The second virial coefficient of hard alkane models. <i>Journal of Chemical Physics</i> , 1998 , 109, 5670-5680	3.9	20	
7	Nonequilibrium properties of linear polar Kihara fluids from molecular dynamics. Results for models and for liquid acetonitrile. <i>Journal of Chemical Physics</i> , 1997 , 107, 2034-2045	3.9	12	
6	Dynamical properties and transport coefficients of Kihara linear fluids. <i>Journal of Chemical Physics</i> , 1997 , 106, 4753-4767	3.9	12	
5	The vapour - liquid equilibrium ofn-alkanes. <i>Journal of Physics Condensed Matter</i> , 1996 , 8, 9643-9648	1.8	3	
4	Equation of state for hard n-alkane models: Long chains. <i>Journal of Chemical Physics</i> , 1996 , 104, 701-713	33.9	26	
3	Understanding the critical properties of chain molecules. <i>Molecular Physics</i> , 1996 , 88, 1575-1602	1.7	20	
2	Vapour-liquid equilibria of propane and n-alkane conformers. <i>Molecular Physics</i> , 1995 , 85, 679-699	1.7	6	
1	Understanding the critical properties of chain molecules		1	