## Gilles A De Wijs

## 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.

109 4,512 37 64 g-index

117 4,894 4.8 5.28 ext. papers ext. citations avg, IF L-index

#	Paper	IF	Citations
109	Impact of F and S doping on (Mn,Fe)2(P,Si) giant magnetocaloric materials. <i>Acta Materialia</i> , <b>2022</b> , 234, 118057	8.4	1
108	Comparing GIPAW with numerically exact chemical shieldings: The role of two-center contributions to the induced current <i>Journal of Chemical Physics</i> , <b>2021</b> , 155, 234101	3.9	
107	Microscopic (Dis)order and Dynamics of Cations in Mixed FA/MA Lead Halide Perovskites. <i>Journal of Physical Chemistry C</i> , <b>2021</b> , 125, 1742-1753	3.8	14
106	Preactive Site in ZieglerNatta Catalysts. <i>Journal of Physical Chemistry C</i> , <b>2019</b> , 123, 14490-14500	3.8	6
105	The Rich Solid-State Phase Behavior of dl-Aminoheptanoic Acid: Five Polymorphic Forms and Their Phase Transitions. <i>Crystal Growth and Design</i> , <b>2018</b> , 18, 242-252	3.5	7
104	NMR shieldings from density functional perturbation theory: GIPAW versus all-electron calculations. <i>Journal of Chemical Physics</i> , <b>2017</b> , 146, 064115	3.9	12
103	Symmetry, Dynamics, and Defects in Methylammonium Lead Halide Perovskites. <i>Journal of Physical Chemistry Letters</i> , <b>2017</b> , 8, 61-66	6.4	63
102	The Role of Connectivity on Electronic Properties of Lead Iodide Perovskite-Derived Compounds. <i>Inorganic Chemistry</i> , <b>2017</b> , 56, 8408-8414	5.1	59
101	Theoretical models of Rashba spin splitting in asymmetric SrTiO3-based heterostructures. <i>Physical Review B</i> , <b>2017</b> , 95,	3.3	6
100	Band Offsets at the Interface between Crystalline and Amorphous Silicon from First Principles. <i>Physical Review Applied</i> , <b>2017</b> , 8,	4.3	7
99	Report on the sixth blind test of organic crystal structure prediction methods. <i>Acta Crystallographica Section B: Structural Science, Crystal Engineering and Materials</i> , <b>2016</b> , 72, 439-59	1.8	338
98	q-GRID: A New Method To Calculate Lattice and Interaction Energies for Molecular Crystals from Electron Densities. <i>Crystal Growth and Design</i> , <b>2016</b> , 16, 662-671	3.5	5
97	A multi-nuclear magnetic resonance and density functional theory investigation of epitaxially grown InGaP2. <i>Physical Chemistry Chemical Physics</i> , <b>2016</b> , 18, 21296-304	3.6	2
96	Structural Studies of Polyaramid Fibers: Solid-State NMR and First-Principles Modeling. <i>Macromolecules</i> , <b>2016</b> , 49, 5548-5560	5.5	9
95	Improved hydrogen storage in Ca-decorated boron heterofullerenes: a theoretical study. <i>Journal of Materials Chemistry A</i> , <b>2015</b> , 3, 7710-7714	13	23
94	Geometric, electronic, and magnetic structure of FexOy+ clusters. <i>Physical Review B</i> , <b>2015</b> , 92,	3.3	17
93	Carbon Support Effects on the Hydrogen Storage Properties of LiBH4 Nanoparticles: A First-Principles Study. <i>Journal of Physical Chemistry C</i> , <b>2014</b> , 118, 5102-5109	3.8	11

92	Anionogenic mixed valency in KxBa1-xO2-\(\frac{1}{2}\)Inorganic Chemistry, <b>2014</b> , 53, 496-502	5.1	3
91	Phonons and electron-phonon coupling in graphene-h-BN heterostructures. <i>Annalen Der Physik</i> , <b>2014</b> , 526, 381-386	2.6	31
90	Interactions of adsorbed COIbn water ice at low temperatures. <i>Physical Chemistry Chemical Physics</i> , <b>2014</b> , 16, 15630-9	3.6	22
89	Stripline 75As NMR Study of Epitaxial III <b>V</b> Semiconductor Al0.5Ga0.5As. <i>Journal of Physical Chemistry C</i> , <b>2014</b> , 118, 13394-13405	3.8	4
88	Li intercalation in graphite: A van der Waals density-functional study. <i>Physical Review B</i> , <b>2014</b> , 90,	3.3	56
87	Quantum confinement and band offsets in amorphous silicon quantum wells. <i>Physical Review B</i> , <b>2014</b> , 90,	3.3	4
86	The electronic structure of organic-inorganic hybrid compounds: (NHIII աCl[I](CHINHIII աCl[I] nd (CHINHIII uCl[I] Journal of Physics Condensed Matter, <b>2013</b> , 25, 295502	1.8	33
85	Role of Magnetism in Catalysis: RuO2 (110) Surface. <i>Journal of Physical Chemistry C</i> , <b>2013</b> , 117, 6353-63	<b>5</b> ₹.8	64
84	First-Principles Study of Structural Prototypes for NaAlH4: Elevated Pressure Polymorph in Symmetry Fmm2 Leads to a Single-Step Decomposition Pathway. <i>Journal of Physical Chemistry C</i> , <b>2013</b> , 117, 8864-8870	3.8	3
83	Excess manganese as the origin of the low-temperature anomaly in NiMnSb. <i>Physical Review B</i> , <b>2013</b> , 88,	3.3	8
82	Finite-field implementation of NMR chemical shieldings for molecules: direct and converse gauge-including projector-augmented-wave methods. <i>Journal of Chemical Physics</i> , <b>2013</b> , 139, 014109	3.9	12
81	k[b subband structure of the LaAlO3/SrTiO3 interface. <i>Physical Review B</i> , <b>2013</b> , 88,	3.3	21
80	First-Principles Study of LiBH4 Nanoclusters and Their Hydrogen Storage Properties. <i>Journal of Physical Chemistry C</i> , <b>2012</b> , 116, 18038-18047	3.8	21
79	Switchable Fermi surface sheets in greigite. <i>Physical Review B</i> , <b>2012</b> , 86,	3.3	10
78	Hydrogen bonding and chemical shift assignments in carbazole functionalized isocyanides from solid-state NMR and first-principles calculations. <i>Physical Chemistry Chemical Physics</i> , <b>2011</b> , 13, 13082-9	5 <sup>3.6</sup>	24
77	Mixed Magnetism for Refrigeration and Energy Conversion. <i>Advanced Energy Materials</i> , <b>2011</b> , 1, 1215-1	<b>219</b> .8	192
76	Intrinsic defects and dopants in LiNH2: a first-principles study. <i>Physical Chemistry Chemical Physics</i> , <b>2011</b> , 13, 6043-52	3.6	14
75	Native Defects and the Dehydrogenation of NaBH4. <i>Journal of Physical Chemistry C</i> , <b>2011</b> , 115, 24429-2	4484	12

74	Optical response of the sodium alanate system: GW0-BSE calculations and thin film measurements. <i>Physical Review B</i> , <b>2011</b> , 83,	3.3	19
73	First-principles modelling of magnesium titanium hydrides. <i>Journal of Physics Condensed Matter</i> , <b>2010</b> , 22, 074208	1.8	10
<del>72</del>	Tuning the Hydrogen Storage in Magnesium Alloys. <i>Journal of Physical Chemistry Letters</i> , <b>2010</b> , 1, 1982	-1 <i>0.</i> 846	20
71	A solid-state NMR and DFT study of compositional modulations in Al(x)Ga(1-x)As. <i>Physical Chemistry Chemical Physics</i> , <b>2010</b> , 12, 11517-35	3.6	25
70	Atomistic models of hydrogenated amorphous silicon nitride from first principles. <i>Physical Review B</i> , <b>2010</b> , 82,	3.3	20
69	Structural models of a-Si:H with a low defect concentration: A first-principles molecular dynamics study. <i>Physica Status Solidi (A) Applications and Materials Science</i> , <b>2010</b> , 207, 605-608	1.6	2
68	Publisher's Note: Modeling and analysis of the three-dimensional current density in sandwich-type single-carrier devices of disordered organic semiconductors [Phys. Rev. B 79, 085203 (2009)]. <i>Physical Review B</i> , <b>2009</b> , 79,	3.3	4
67	First-principles study of the optical properties of MgxTi1NH2. <i>Physical Review B</i> , <b>2009</b> , 79,	3.3	15
66	Amorphous semiconductors studied by first-principles simulations: structure and electronic properties. <i>Materials Research Society Symposia Proceedings</i> , <b>2009</b> , 1153, 1		1
65	First-principles study of hydrogenated amorphous silicon. <i>Physical Review B</i> , <b>2009</b> , 79,	3.3	49
64	Hydrogen Storage by Polylithiated Molecules and Nanostructures. <i>Journal of Physical Chemistry C</i> , <b>2009</b> , 113, 8997-9002	3.8	37
63	DFT Study of Planar Boron Sheets: A New Template for Hydrogen Storage. <i>Journal of Physical Chemistry C</i> , <b>2009</b> , 113, 18962-18967	3.8	101
62	Tunable hydrogen storage in magnesium <b>E</b> ransition metal compounds: First-principles calculations. <i>Physical Review B</i> , <b>2009</b> , 79,	3.3	45
61	Modeling and analysis of the three-dimensional current density in sandwich-type single-carrier devices of disordered organic semiconductors. <i>Physical Review B</i> , <b>2009</b> , 79,	3.3	98
60	Electronic band structure of tetracene-TCNQ and perylene-TCNQ compounds. <i>Journal of Physical Chemistry A</i> , <b>2008</b> , 112, 2497-502	2.8	41
59	First-principles calculations of the crystal structure, electronic structure, and thermodynamic stability of Be(BH4)2. <i>Physical Review B</i> , <b>2008</b> , 77,	3.3	28
58	A Density Functional Study of ⊞Mg(BH4)2. <i>Chemistry of Materials</i> , <b>2008</b> , 20, 4952-4956	9.6	70
57	Theoretical study of the stable radicals galvinoxyl, azagalvinoxyl and wursters blue perchlorate in the solid state. <i>Journal of Physical Chemistry A</i> , <b>2008</b> , 112, 7734-8	2.8	14

56	Spin tunneling in junctions with disordered ferromagnets. <i>Physical Review Letters</i> , <b>2008</b> , 100, 057205	7.4	28
55	Work function anisotropy and surface stability of half-metallic CrO2. <i>Physical Review B</i> , <b>2008</b> , 77,	3.3	17
54	Model for the Formation Energies of Alanates and Boranates. <i>Journal of Physical Chemistry C</i> , <b>2007</b> , 111, 9592-9594	3.8	9
53	Thermodynamic stability of boron: the role of defects and zero point motion. <i>Journal of the American Chemical Society</i> , <b>2007</b> , 129, 2458-65	16.4	147
52	Spintronic materials based on main-group elements. <i>Journal of Physics Condensed Matter</i> , <b>2007</b> , 19, 165	5208	21
51	Optimizing performance of half-metals at finite temperature. <i>Journal of Physics Condensed Matter</i> , <b>2007</b> , 19, 315212	1.8	18
50	Electronic structure and optical properties of lightweight metal hydrides. <i>Physical Review B</i> , <b>2007</b> , 75,	3.3	109
49	Ab initio study of the effects of transition metal doping of Mg2NiH4. <i>Physical Review B</i> , <b>2007</b> , 76,	3.3	53
48	Crystal Growth, Structure, and Electronic Band Structure of TetracenellCNQ. <i>Journal of Physical Chemistry C</i> , <b>2007</b> , 111, 3486-3489	3.8	34
47	Lattice and local-mode vibrations in anhydrous and protonized LiMn2O4 spinels from first-principles theory. <i>Journal of Materials Chemistry</i> , <b>2007</b> , 17, 4908		14
46	The continuing drama of the half-metal/semiconductor interface. <i>Journal Physics D: Applied Physics</i> , <b>2006</b> , 39, 793-796	3	39
45	Ab initio and work function and surface energy anisotropy of LaB6. <i>Journal of Physical Chemistry B</i> , <b>2006</b> , 110, 18459-65	3.4	69
44	Local Structure and Chemical Bonding of Protonated LixMn2O4 Spinels from First Principles. <i>Chemistry of Materials</i> , <b>2006</b> , 18, 1169-1173	9.6	17
43	Generalised coexistence of a low work function and a stable surface: CaAl4 and BaAuIn3. <i>Surface Science</i> , <b>2006</b> , 600, 2495-2500	1.8	7
42	Interrelation of Work Function and Surface Stability: The Case of BaAl4. <i>Chemistry of Materials</i> , <b>2005</b> , 17, 3879-3882	9.6	9
41	Anionogenic ferromagnets. <i>Journal of the American Chemical Society</i> , <b>2005</b> , 127, 16325-8	16.4	58
40	The role of the hydrogen bonding network for the shear modulus of PIPD. <i>Polymer</i> , <b>2005</b> , 46, 9144-915	43.9	18
39	Quantum mechanics calculations on the diastereomeric salts of cyclic phosphoric acids with ephedrine. <i>Computational and Theoretical Chemistry</i> , <b>2005</b> , 717, 205-214		

38	Geometry of {001} Surfaces of Spinel (MgAl2O4): First-Principles Simulations and Experimental Measurements. <i>Journal of the American Ceramic Society</i> , <b>2005</b> , 88, 1544-1548	3.8	30
37	Tunable spin transport in CrAs: Role of correlation effects. <i>Physical Review B</i> , <b>2005</b> , 71,	3.3	38
36	Ab initio study of Mg(AlH4)2. <i>Physical Review B</i> , <b>2005</b> , 72,	3.3	46
35	Lattice vibrations and thermal properties of carbon nitride with defect ZnS structure from first-principles calculations. <i>Journal of Physics Condensed Matter</i> , <b>2004</b> , 16, 3027-3034	1.8	6
34	Low work function of the (1000) Ca2N surface. <i>Journal of Applied Physics</i> , <b>2004</b> , 96, 1751-1753	2.5	17
33	Defects in half-metals and finite temperature. <i>Journal of Physics Condensed Matter</i> , <b>2004</b> , 16, S5517-S5	5248	33
32	Local structure and electronic properties of BaTaO2N with perovskite-type structure. <i>Journal of Physics and Chemistry of Solids</i> , <b>2003</b> , 64, 281-286	3.9	59
31	Weakening of a Polyethylene Chain by Methyl Side Groups. <i>Soft Materials</i> , <b>2003</b> , 1, 223-233	1.7	3
30	Anisotropy of the mobility of pentacene from frustration. Synthetic Metals, 2003, 139, 109-114	3.6	123
29	Phonon spectrum and thermal properties of cubic Si3N4 from first-principles calculations. <i>Journal of Applied Physics</i> , <b>2003</b> , 93, 5175-5180	2.5	44
28	Modeling the polymorphism of pentacene. <i>Journal of the American Chemical Society</i> , <b>2003</b> , 125, 6323-3	016.4	198
27	Spin-polarization in half-metals (invited). <i>Journal of Applied Physics</i> , <b>2002</b> , 91, 8340	2.5	173
26	First-principles calculation of the phonon spectrum of MgAl2O4 spinel. <i>Physical Review B</i> , <b>2002</b> , 65,	3.3	56
25	Phonon spectrum of ZnAl2O4 spinel from inelastic neutron scattering and first-principles calculations. <i>Physical Review B</i> , <b>2002</b> , 66,	3.3	49
24	Amorphous WO3: a first-principles approach. <i>Electrochimica Acta</i> , <b>2001</b> , 46, 1989-1993	6.7	46
23	Patterning of Si(001) with halogens: Surface structure as a function of the halogen chemical potential. <i>Physical Review B</i> , <b>2001</b> , 64,	3.3	22
22	Towards 100% spin-polarized charge-injection: The half-metallic NiMnSb/CdS interface. <i>Physical Review B</i> , <b>2001</b> , 64,	3.3	166
21	The electronic structure of tantalum (oxy)nitridesTaON and Ta3N5. <i>Journal of Materials Chemistry</i> , <b>2001</b> , 11, 1248-1252		164

20	Lithium trapping by excess oxygen in WO3: A first-principles study. <i>Physical Review B</i> , <b>2000</b> , 62, 1508-15	51313	6
19	Bond Scission in a Perfect Polyethylene Chain and the Consequences for the Ultimate Strength. <i>Macromolecules</i> , <b>2000</b> , 33, 9098-9108	5.5	25
18	O/N Ordering in Y2Si3O3N4 with the Melilite-type Structure from First-Principles Calculations. <i>Chemistry of Materials</i> , <b>2000</b> , 12, 1071-1075	9.6	27
17	Bulk and Surface Electronic Structure of the Layered Sub-Nitrides Ca2N and Sr2N. <i>Chemistry of Materials</i> , <b>2000</b> , 12, 1847-1852	9.6	22
16	Structure and electronic properties of amorphous WO3. <i>Physical Review B</i> , <b>1999</b> , 60, 16463-16474	3.3	79
15	Transport coefficients of liquids from first principles. <i>Journal of Non-Crystalline Solids</i> , <b>1999</b> , 250-252, 82-90	3.9	11
14	The viscosity of liquid iron at the physical conditions of the Earth's core. <i>Nature</i> , <b>1998</b> , 392, 805-807	50.4	232
13	First-order phase transitions by first-principles free-energy calculations: The melting of Al. <i>Physical Review B</i> , <b>1998</b> , 57, 8223-8234	3.3	115
12	First-principles study of chlorine adsorption and reactions on Si(100). <i>Physical Review B</i> , <b>1998</b> , 57, 1002	1-3.902	944
11	Reversed spin polarization at the Co(001)-HfO2(001) interface. <i>Physical Review B</i> , <b>1998</b> , 58, 15422-1542	253.3	13
10	Mechanism for SiCl2 Formation and Desorption and the Growth of Pits in the Etching of Si(100) with Chlorine. <i>Physical Review Letters</i> , <b>1997</b> , 78, 4877-4880	7.4	49
9	First principles calculations on crystalline and liquid iron at Earth\$ core conditions. <i>Faraday Discussions</i> , <b>1997</b> , 106, 205-218	3.6	94
8	Chlorine on Si(001)-(2 x 1): Bridge versus Terminal Bonding. <i>Physical Review Letters</i> , <b>1996</b> , 77, 881-884	7.4	21
7	Electron-ion correlation in liquid metals from first principles: Liquid Mg and liquid Bi. <i>Physical Review Letters</i> , <b>1995</b> , 75, 4480-4483	7.4	31
6	First-principles molecular-dynamics simulation of liquid CsPb. <i>Journal of Chemical Physics</i> , <b>1995</b> , 103, 5031-5040	3.9	36
5	Clusters in liquid K-Tl and Cs-Tl alloys. <i>Journal of Physics Condensed Matter</i> , <b>1994</b> , 6, A255-A260	1.8	12
4	Ab initio molecular dynamics study of liquid Li12Si7. <i>Journal of Non-Crystalline Solids</i> , <b>1993</b> , 156-158, 961-964	3.9	1
3	Structure of liquid caesium-lead alloys. <i>Journal of Non-Crystalline Solids</i> , <b>1993</b> , 156-158, 34-37	3.9	7

Nanometre superstructure in liquid alkali-thallium alloys. *Journal of Physics Condensed Matter*, **1993** , 5, 9253-9260

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First-principles molecular-dynamics simulation of liquid Li12Si7. *Physical Review B*, **1993**, 48, 13459-1346**8**.3

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