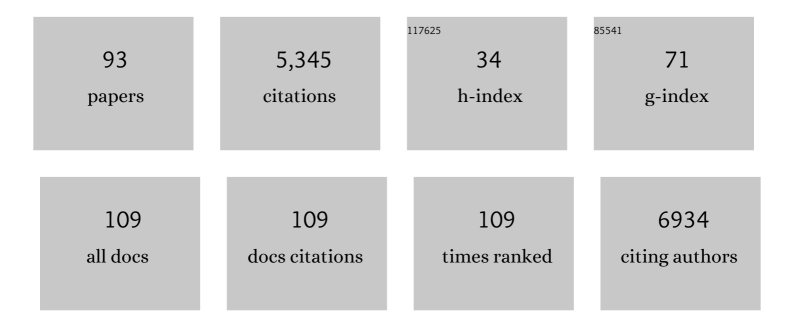
List of Publications by Year in descending order

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#	Article	lF	CITATIONS
1	How are transition states modeled in heterogeneous electrocatalysis?. Current Opinion in Electrochemistry, 2022, 33, 100940.	4.8	20
2	Autonomous high-throughput computations in catalysis. Chem Catalysis, 2022, 2, 940-956.	6.1	14
3	Genesis of MoS2 from model-Mo-oxide precursors supported on γ-alumina. Journal of Catalysis, 2022, 408, 303-315.	6.2	4
4	Mechanistic Investigation and Free Energies of the Reactive Adsorption of Ethanol at the Alumina/Water Interface. Journal of Physical Chemistry C, 2022, 126, 7446-7455.	3.1	8
5	How to Gain Atomistic Insights on Reactions at the Water/Solid Interface?. ACS Catalysis, 2022, 12, 6294-6301.	11.2	17
6	Modeling Electrochemical Processes with Grand Canonical Treatment of Many-Body Perturbation Theory. Journal of Physical Chemistry Letters, 2022, 13, 6079-6084.	4.6	8
7	Atomistic modeling of electrocatalysis: Are we there yet?. Wiley Interdisciplinary Reviews: Computational Molecular Science, 2021, 11, e1499.	14.6	79
8	The Impact of Water on Ru-Catalyzed Olefin Metathesis: Potent Deactivating Effects Even at Low Water Concentrations. ACS Catalysis, 2021, 11, 893-899.	11.2	25
9	Nature of High- and Low-Affinity Metal Surface Sites on Birnessite Nanosheets. ACS Earth and Space Chemistry, 2021, 5, 66-76.	2.7	11
10	(Dis)Similarities of adsorption of diverse functional groups over alumina and hematite depending on the surface state. Journal of Chemical Physics, 2021, 154, 084701.	3.0	11
11	Understanding electrified interfaces. Nature Reviews Materials, 2021, 6, 289-291.	48.7	38
12	Designing Active Sites for Structure-Sensitive Reactions via the Generalized Coordination Number: Application to Alcohol Dehydrogenation. Journal of Physical Chemistry C, 2021, 125, 10370-10377.	3.1	6
13	Efficient recursive least squares solver for rank-deficient matrices. Applied Mathematics and Computation, 2021, 399, 125996.	2.2	0
14	DockOnSurf: A Python Code for the High-Throughput Screening of Flexible Molecules Adsorbed on Surfaces. Journal of Chemical Information and Modeling, 2021, 61, 3386-3396.	5.4	13
15	How Stable Are 2H-MoS ₂ Edges under Hydrogen Evolution Reaction Conditions?. Journal of Physical Chemistry C, 2021, 125, 17058-17067.	3.1	25
16	Transferable Gaussian Attractive Potentials for Organic/Oxide Interfaces. Journal of Physical Chemistry B, 2021, 125, 10843-10853.	2.6	8
17	What does graphitic carbon nitride really look like?. Physical Chemistry Chemical Physics, 2021, 23, 2853-2859.	2.8	12
18	Same ligand, three first-row metals: comparing M-amido bifunctional reactivity (Mn, Fe, Co). Dalton Transactions, 2021, 50, 14542-14546.	3.3	5

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19	Adhesion of lubricant on aluminium through adsorption of additive head-groups on γ-alumina: A DFT study. Tribology International, 2020, 145, 106140.	5.9	15
20	Mononuclear Fe in N-doped carbon: computational elucidation of active sites for electrochemical oxygen reduction and oxygen evolution reactions. Catalysis Science and Technology, 2020, 10, 1006-1014.	4.1	34
21	Solvation Free Energies and Adsorption Energies at the Metal/Water Interface from Hybrid Quantum-Mechanical/Molecular Mechanics Simulations. Journal of Chemical Theory and Computation, 2020, 16, 6539-6549.	5.3	34
22	Water adlayers on noble metal surfaces: Insights from energy decomposition analysis. Journal of Chemical Physics, 2020, 153, 054703.	3.0	10
23	Demystifying the Atomistic Origin of the Electric Field Effect on Methane Oxidation. Journal of Physical Chemistry Letters, 2020, 11, 6976-6981.	4.6	16
24	Size-Dependent Structural, Energetic, and Spectroscopic Properties of MoS ₃ Polymorphs. Crystal Growth and Design, 2020, 20, 7750-7760.	3.0	9
25	Strong Affinity of Triazolium-Appended Dipyrromethenes (TADs) for BF4â^'. Molecules, 2020, 25, 4555.	3.8	2
26	Ten Facets, One Force Field: The GAL19 Force Field for Water–Noble Metal Interfaces. Journal of Chemical Theory and Computation, 2020, 16, 4565-4578.	5.3	26
27	Elucidating the role of electrochemical polarization on the selectivity of the CO2 hydrogenation reaction over Ru. Electrochimica Acta, 2020, 350, 136405.	5.2	20
28	Revisiting the Active Sites at the MoS ₂ /H ₂ O Interface via Grand-Canonical DFT: The Role of Water Dissociation. ACS Applied Materials & Interfaces, 2020, 12, 31401-31410.	8.0	36
29	Hydroxide-Induced Degradation of Olefin Metathesis Catalysts: A Challenge for Metathesis in Alkaline Media. ACS Catalysis, 2020, 10, 3838-3843.	11.2	15
30	Two-sites are better than one: revisiting the OER mechanism on CoOOH by DFT with electrode polarization. Physical Chemistry Chemical Physics, 2020, 22, 7031-7038.	2.8	45
31	The Mode of Incorporation of As(-I) and Se(-I) in Natural Pyrite Revisited. ACS Earth and Space Chemistry, 2020, 4, 379-390.	2.7	18
32	Parameter-free coordination numbers for solutions and interfaces. Journal of Chemical Physics, 2020, 152, 024124.	3.0	11
33	Revisiting the Atomistic Structures at the Interface of Au(111) Electrode–Sulfuric Acid Solution. Journal of the American Chemical Society, 2020, 142, 9439-9446.	13.7	35
34	Evaluating Thermal Corrections for Adsorption Processes at the Metal/Gas Interface. Journal of Physical Chemistry C, 2019, 123, 28828-28835.	3.1	17
35	The Pressure Gap for Thiols: Methanethiol Self-Assembly on Au(111) from Vacuum to 1 bar. Journal of Physical Chemistry C, 2019, 123, 12382-12389.	3.1	7
36	Theory and experiments join forces to characterize the electrocatalytic interface. Proceedings of the National Academy of Sciences of the United States of America, 2019, 116, 7611-7613.	7.1	5

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37	Theory-guided materials design: two-dimensional MXenes in electro- and photocatalysis. Nanoscale Horizons, 2019, 4, 809-827.	8.0	218
38	Theoretical insight into the origin of the electrochemical promotion of ethylene oxidation on ruthenium oxide. Catalysis Science and Technology, 2019, 9, 5915-5926.	4.1	26
39	Implicit self-consistent electrolyte model in plane-wave density-functional theory. Journal of Chemical Physics, 2019, 151, 234101.	3.0	561
40	Energy Decomposition Analysis for Metal Surface–Adsorbate Interactions by Block Localized Wave Functions. Journal of Chemical Theory and Computation, 2019, 15, 265-275.	5.3	13
41	Can microsolvation effects be estimated from vacuum computations? A case-study of alcohol decomposition at the H ₂ O/Pt(111) interface. Physical Chemistry Chemical Physics, 2019, 21, 5368-5377.	2.8	25
42	C6 Diacids from homocitric acid lactone using relay heterogeneous catalysis in water. Catalysis Today, 2019, 319, 191-196.	4.4	1
43	Computational screening for selective catalysts: Cleaving the C C bond during ethanol electro-oxidation reaction. Electrochimica Acta, 2018, 274, 274-278.	5.2	26
44	Evaluating the Risk of C–C Bond Formation during Selective Hydrogenation of Acetylene on Palladium. ACS Catalysis, 2018, 8, 1662-1671.	11.2	65
45	Force Field for Water over Pt(111): Development, Assessment, and Comparison. Journal of Chemical Theory and Computation, 2018, 14, 3238-3251.	5.3	38
46	Tetrazine-Based Ligand Transformation Driving Metal–Metal Bond and Mixed-Valence Hg ^I /Hg ^{II} . ACS Omega, 2018, 3, 10273-10277.	3.5	3
47	Shining Light on Carbon Nitrides: Leveraging Temperature To Understand Optical Gap Variations. Chemistry of Materials, 2018, 30, 4253-4262.	6.7	28
48	Acetylene Adsorption on Pd–Ag Alloys: Evidence for Limited Island Formation and Strong Reverse Segregation from Monte Carlo Simulations. Journal of Physical Chemistry C, 2018, 122, 15456-15463.	3.1	35
49	Key Role of Anionic Doping for H ₂ Production from Formic Acid on Pd(111). ACS Catalysis, 2017, 7, 1955-1959.	11.2	72
50	Challenges in calculating the bandgap of triazine-based carbon nitride structures. Journal of Materials Chemistry A, 2017, 5, 5115-5122.	10.3	34
51	Group Additivity for Aqueous Phase Thermochemical Properties of Alcohols on Pt(111). Journal of Physical Chemistry C, 2017, 121, 21510-21519.	3.1	27
52	Molecular mechanics models for the image charge, a comment on "including image charge effects in the molecular dynamics simulations of molecules on metal surfaces― Journal of Computational Chemistry, 2017, 38, 2127-2129.	3.3	9
53	A machine learning approach to graph-theoretical cluster expansions of the energy of adsorbate layers. Journal of Chemical Physics, 2017, 147, 054106.	3.0	31
54	DFT Perspective on the Thermochemistry of Carbon Nitride Synthesis. Journal of Physical Chemistry C, 2016, 120, 24542-24550.	3.1	21

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55	C ₂ H ₂ -Induced Surface Restructuring of Pd–Ag Catalysts: Insights from Theoretical Modeling. Journal of Physical Chemistry C, 2016, 120, 26320-26327.	3.1	26
56	Solvation free energies for periodic surfaces: comparison of implicit and explicit solvation models. Physical Chemistry Chemical Physics, 2016, 18, 31850-31861.	2.8	80
57	Ethanol Electro-oxidation on Palladium Revisited Using Polarization Modulation Infrared Reflection Absorption Spectroscopy (PM-IRRAS) and Density Functional Theory (DFT): Why Is It Difficult To Break the C–C Bond?. ACS Catalysis, 2016, 6, 4894-4906.	11.2	109
58	Study of a novel hepta-coordinated FeIII bimetallic complex with an unusual 1,2,4,5-tetrazine-ring opening. Polyhedron, 2016, 108, 163-168.	2.2	13
59	Electro-carboxylation of butadiene and ethene over Pt and Ni catalysts. Journal of Catalysis, 2016, 343, 240-247.	6.2	31
60	Assessing a First-Principles Model of an Electrochemical Interface by Comparison with Experiment. Journal of Physical Chemistry C, 2016, 120, 5619-5623.	3.1	78
61	A fast chargeâ€Dependent atomâ€pairwise dispersion correction for DFTB3. International Journal of Quantum Chemistry, 2015, 115, 1265-1272.	2.0	16
62	Modeling the HCOOH/CO ₂ Electrocatalytic Reaction: When Details Are Key. ChemPhysChem, 2015, 16, 2307-2311.	2.1	44
63	Impacts of electrode potentials and solvents on the electroreduction of CO ₂ : a comparison of theoretical approaches. Physical Chemistry Chemical Physics, 2015, 17, 13949-13963.	2.8	90
64	Relationship between Carbon Nitride Structure and Exciton Binding Energies: A DFT Perspective. Journal of Physical Chemistry C, 2015, 119, 25188-25196.	3.1	104
65	Molecular adsorption at Pt(111). How accurate are DFT functionals?. Physical Chemistry Chemical Physics, 2015, 17, 28921-28930.	2.8	210
66	How important is self-consistency for the dDsC density dependent dispersion correction?. Journal of Chemical Physics, 2014, 140, 18A516.	3.0	24
67	Layer-Dependent Electrocatalysis of MoS ₂ for Hydrogen Evolution. Nano Letters, 2014, 14, 553-558.	9.1	667
68	Hierarchically Structured Microfibers of "Single Stack―Perylene Bisimide and Quaterthiophene Nanowires. ACS Nano, 2013, 7, 8498-8508.	14.6	88
69	Bonding analysis of planar hypercoordinate atoms via the generalized BLW‣OL. Journal of Computational Chemistry, 2013, 34, 2242-2248.	3.3	10
70	Wave function methods for fractional electrons. Journal of Chemical Physics, 2013, 139, 074107.	3.0	19
71	Equivalence of particle-particle random phase approximation correlation energy and ladder-coupled-cluster doubles. Journal of Chemical Physics, 2013, 139, 104112.	3.0	51
72	Dynamical second-order Bethe-Salpeter equation kernel: A method for electronic excitation beyond the adiabatic approximation. Journal of Chemical Physics, 2013, 139, 154109.	3.0	23

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73	Benchmark tests and spin adaptation for the particle-particle random phase approximation. Journal of Chemical Physics, 2013, 139, 174110.	3.0	40
74	A ratiometric fluorescence sensor for caffeine. Organic and Biomolecular Chemistry, 2012, 10, 7487.	2.8	19
75	Role of π-Acceptor Effects in Controlling the Lability of Novel Monofunctional Pt(II) and Pd(II) Complexes: Crystal Structure of [Pt(tripyridinedimethane)Cl]Cl. Inorganic Chemistry, 2012, 51, 1516-1529.	4.0	48
76	π-Depletion as a criterion to predict π-stacking ability. Chemical Communications, 2012, 48, 9239.	4.1	68
77	Exploring the Limits of Density Functional Approximations for Interaction Energies of Molecular Precursors to Organic Electronics. Journal of Chemical Theory and Computation, 2012, 8, 4305-4316.	5.3	38
78	Quantification of "fuzzy―chemical concepts: a computational perspective. Chemical Society Reviews, 2012, 41, 4671.	38.1	108
79	How are small endohedral silicon clusters stabilized?. Physical Chemistry Chemical Physics, 2012, 14, 14842.	2.8	12
80	Why are the Interaction Energies of Charge-Transfer Complexes Challenging for DFT?. Journal of Chemical Theory and Computation, 2012, 8, 1629-1640.	5.3	153
81	Fluorescence sensing of caffeine in water with polysulfonated pyrenes. Chemical Communications, 2011, 47, 10584.	4.1	43
82	Dispersion-Corrected Energy Decomposition Analysis for Intermolecular Interactions Based on the BLW and dDXDM Methods. Journal of Physical Chemistry A, 2011, 115, 5467-5477.	2.5	43
83	The norbornene mystery revealed. Chemical Communications, 2011, 47, 227-229.	4.1	20
84	How do electron localization functions describe π-electron delocalization?. Physical Chemistry Chemical Physics, 2011, 13, 20584.	2.8	99
85	Comprehensive Benchmarking of a Density-Dependent Dispersion Correction. Journal of Chemical Theory and Computation, 2011, 7, 3567-3577.	5.3	400
86	A Density Dependent Dispersion Correction. Chimia, 2011, 65, 240.	0.6	40
87	A generalized-gradient approximation exchange hole model for dispersion coefficients. Journal of Chemical Physics, 2011, 134, 044117.	3.0	270
88	Overcoming systematic DFT errors for hydrocarbon reaction energies. Theoretical Chemistry Accounts, 2010, 127, 429-442.	1.4	51
89	Branched Alkanes Have Contrasting Stabilities. Organic Letters, 2010, 12, 3070-3073.	4.6	34
90	How Strained are Carbomeric-Cycloalkanes?. Journal of Physical Chemistry A, 2010, 114, 6705-6712.	2.5	22

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91	A System-Dependent Density-Based Dispersion Correction. Journal of Chemical Theory and Computation, 2010, 6, 1990-2001.	5.3	133
92	Direct Assessment of Electron Delocalization Using NMR Chemical Shifts. Angewandte Chemie - International Edition, 2009, 48, 9828-9833.	13.8	49
93	Unified Inter- and Intramolecular Dispersion Correction Formula for Generalized Gradient Approximation Density Functional Theory. Journal of Chemical Theory and Computation, 2009, 5, 2950-2958.	5.3	76