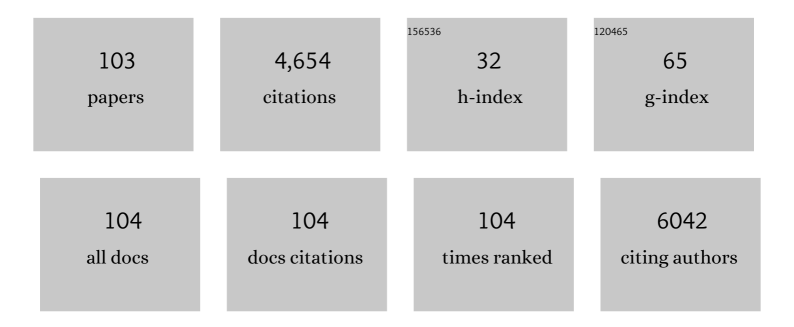
Jean-Sabin McEwen

List of Publications by Year in descending order

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#	Article	lF	CITATIONS
1	Nitrogen-doped char as a catalyst for wet oxidation of phenol-contaminated water. Biomass Conversion and Biorefinery, 2024, 14, 5861-5875.	2.9	4
2	Determining the Adsorption Energetics of 2,3-Butanediol on RuO2(110): Coupling First-Principles Calculations With Global Optimizers. Frontiers in Energy Research, 2022, 9, .	1.2	0
3	Elucidating the Role of B-Site Cations toward CO ₂ Reduction in Perovskite-Based Solid Oxide Electrolysis Cells. Journal of the Electrochemical Society, 2022, 169, 034532.	1.3	8
4	BrÃ,nsted acidity of H-adatoms at protic solvent-transition metal interfaces and its kinetic consequences in electrophilic addition reactions. Journal of Catalysis, 2022, 408, 179-195.	3.1	3
5	Chemisorption-Induced Formation of Biphenylene Dimer on Ag(111). Journal of the American Chemical Society, 2022, 144, 723-732.	6.6	20
6	Novel Amorphous Carbons for the Adsorption of Phosphate: Part I. Elucidation of Chemical Structure of N-Metal-Doped Chars. ACS Omega, 2022, 7, 14490-14504.	1.6	1
7	Elucidating CO Oxidation Pathways on Rh Atoms and Clusters on the "29―Cu ₂ O/Cu(111) Surface. Journal of Physical Chemistry C, 2022, 126, 11091-11102.	1.5	1
8	Catalytic consequences of hydrogen addition events and solvent-adsorbate interactions during guaiacol-H2 reactions at the H2O-Ru(0Â0Â0Â1) interface. Journal of Catalysis, 2021, 395, 467-482.	3.1	15
9	Developing single-site Pt catalysts for the preferential oxidation of CO: A surface science and first principles-guided approach. Applied Catalysis B: Environmental, 2021, 284, 119716.	10.8	19
10	Determining the hydration energetics on carbon-supported Ru catalysts: An adsorption calorimetry and density functional theory study. Catalysis Today, 2021, 365, 172-180.	2.2	3
11	Microscopic insights into long-range 1D ordering in a dense semi-disordered molecular overlayer. Chemical Communications, 2021, 57, 5937-5940.	2.2	1
12	Elucidating the Influence of Electric Fields toward CO2 Activation on YSZ (111). Catalysts, 2021, 11, 271.	1.6	5
13	Guiding the design of oxidation-resistant Fe-based single atom alloy catalysts with insights from configurational space. Journal of Chemical Physics, 2021, 154, 174709.	1.2	3
14	Elucidating the Cooperative Roles of Water and Lewis Acid–Base Pairs in Cascade C–C Coupling and Self-Deoxygenation Reactions. Jacs Au, 2021, 1, 1471-1487.	3.6	5
15	First-Principles Approach to Extracting Chemical Information from X-ray Absorption Near-Edge Spectra of Ga-Containing Materials. Journal of Physical Chemistry C, 2021, 125, 27901-27908.	1.5	5
16	Field-assisted suppression of coke in the methane steam reforming reaction. Applied Catalysis B: Environmental, 2020, 260, 118132.	10.8	20
17	Enhancing Chemical Interaction of Polysulfide and Carbon through Synergetic Nitrogen and Phosphorus Doping. ACS Sustainable Chemistry and Engineering, 2020, 8, 806-813.	3.2	11
18	Formulation of Multicomponent Lattice Gas Model Cluster Expansions Parameterized on Ab Initio Data: An Introduction to the Ab Initio Mean-Field Augmented Lattice Gas Modeling Code. Journal of Physical Chemistry C, 2020, 124, 2923-2938.	1.5	10

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19	Coverage-Dependent Adsorption of Phenol on Pt(111) from First Principles. Journal of Physical Chemistry C, 2020, 124, 356-362.	1.5	12
20	Unravelling the reaction mechanism of gas-phase formic acid decomposition on highly dispersed Mo2C nanoparticles supported on graphene flakes. Applied Catalysis B: Environmental, 2020, 264, 118478.	10.8	23
21	The Role of Protons and Hydrides in the Catalytic Hydrogenolysis of Guaiacol at the Ruthenium Nanoparticle–Water Interface. ACS Catalysis, 2020, 10, 12310-12332.	5.5	29
22	Identifying Trends in the Field Ionization of Diatomic Molecules over Adsorbate Covered Pd(331) Surfaces. Topics in Catalysis, 2020, 63, 1510-1521.	1.3	0
23	Promoting the Cleavage of C–O Bonds at the Interface between a Metal Oxide Cluster and a Co(0001) Support. ACS Catalysis, 2020, 10, 14722-14731.	5.5	8
24	Controlling the Oxidation State of Fe-Based Catalysts through Nitrogen Doping toward the Hydrodeoxygenation of <i>m</i> -Cresol. ACS Catalysis, 2020, 10, 7884-7893.	5.5	32
25	Templated Growth of a Homochiral Thin Film Oxide. ACS Nano, 2020, 14, 4682-4688.	7.3	5
26	Accelerated Cu ₂ O Reduction by Single Pt Atoms at the Metal-Oxide Interface. ACS Catalysis, 2020, 10, 4215-4226.	5.5	34
27	Probing Active-Site Relocation in Cu/SSZ-13 SCR Catalysts during Hydrothermal Aging by In Situ EPR Spectroscopy, Kinetics Studies, and DFT Calculations. ACS Catalysis, 2020, 10, 9410-9419.	5.5	64
28	Coverage-Dependent Adsorption of Hydrogen on Fe(100): Determining Catalytically Relevant Surface Structures via Lattice Gas Models. Journal of Physical Chemistry C, 2020, 124, 7254-7266.	1.5	15
29	Deconvoluting the XPS spectra for nitrogen-doped chars: An analysis from first principles. Carbon, 2020, 162, 528-544.	5.4	323
30	Enhancing the partial oxidation of gasoline with Mo-doped Ni catalysts for SOFC applications: An integrated experimental and DFT study. Applied Catalysis B: Environmental, 2020, 266, 118626.	10.8	24
31	Microstructural analysis of nitrogen-doped char by Raman spectroscopy: Raman shift analysis from first principles. Carbon, 2020, 167, 559-574.	5.4	52
32	Visualizing the origin of rotational entropy effects in coadsorbed systems. Physical Review Research, 2020, 2, .	1.3	4
33	Controlling Molecular Switching via Chemical Functionality: Ethyl vs Methoxy Rotors. Journal of Physical Chemistry C, 2019, 123, 23738-23746.	1.5	9
34	Thermodynamic stability of nitrogen functionalities and defects in graphene and graphene name name of the second nanoribbons from first principles. Carbon, 2019, 152, 715-726.	5.4	22
35	Benchmarking the accuracy of coverage-dependent models: adsorption and desorption of benzene on Pt (1â∈¯1 1) and Pt3Sn (1 1 1) from first principles. Progress in Surface Science, 2019, 94, 100538.	3.8	10
36	Understanding Enantioselective Interactions by Pulling Apart Molecular Rotor Complexes. ACS Nano, 2019, 13, 5939-5946.	7.3	8

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37	Improving the electrochemical oxidation of formic acid by tuning the electronic properties of Pd-based bimetallic nanoparticles. Applied Catalysis B: Environmental, 2019, 254, 685-692.	10.8	73
38	The partial reduction of clean and doped α-Fe2O3(0001) from first principles. Applied Catalysis A: General, 2019, 582, 116989.	2.2	6
39	Surface-Templated Assembly of Molecular Methanol on the Thin Film "29―Cu(111) Surface Oxide. Journal of Physical Chemistry C, 2019, 123, 2911-2921.	1.5	9
40	An atomic-scale view of single-site Pt catalysis for low-temperature CO oxidation. Nature Catalysis, 2018, 1, 192-198.	16.1	292
41	Modeling the adsorbate coverage distribution over a multi-faceted catalytic grain in the presence of an electric field: O/Fe from first principles. Catalysis Today, 2018, 312, 92-104.	2.2	4
42	Local Environment Sensitivity of the Cu K-Edge XANES Features in Cu-SSZ-13: Analysis from First-Principles. Journal of Physical Chemistry Letters, 2018, 9, 3035-3042.	2.1	34
43	Mechanistic understanding of methanol carbonylation: Interfacing homogeneous and heterogeneous catalysis via carbon supported Ir La. Journal of Catalysis, 2018, 361, 414-422.	3.1	28
44	Identifying the Thermal Decomposition Mechanism of Guaiacol on Pt(111): An Integrated X-ray Photoelectron Spectroscopy and Density Functional Theory Study. Journal of Physical Chemistry C, 2018, 122, 4261-4273.	1.5	5
45	Mechanistic Effects of Water on the Fe-Catalyzed Hydrodeoxygenation of Phenol. The Role of BrÃ,nsted Acid Sites. ACS Catalysis, 2018, 8, 2200-2208.	5.5	50
46	Characterization of CoCu- and CoMn-Based Catalysts for the Fischer–Tropsch Reaction Toward Chain-Lengthened Oxygenates. Topics in Catalysis, 2018, 61, 1016-1023.	1.3	10
47	Elucidating the Roles of Electric Fields in Catalysis: A Perspective. ACS Catalysis, 2018, 8, 5153-5174.	5.5	215
48	Dissolution of CoCu catalyst step defects by Co subcarbonyl formation. Journal of Catalysis, 2018, 368, 31-37.	3.1	11
49	Modeling the Adsorption of NO and NH ₃ on Fe-SSZ-13 from First-Principles: A DFT Study. Industrial & Engineering Chemistry Research, 2018, 57, 13396-13405.	1.8	16
50	Computational Catalysis at NAM25. Catalysis Today, 2018, 312, 1.	2.2	0
51	Predicting the Electric Field Effect on the Lateral Interactions Between Adsorbates: O/Fe(100) from First Principles. Topics in Catalysis, 2018, 61, 763-775.	1.3	11
52	Water activation by single Pt atoms supported on a Cu2O thin film. Journal of Catalysis, 2018, 364, 166-173.	3.1	18
53	Hydrogen Oxidation and Water Dissociation over an Oxygen-Enriched Ni/YSZ Electrode in the Presence of an Electric Field: A First-Principles-Based Microkinetic Model. Industrial & Engineering Chemistry Research, 2017, 56, 1201-1213.	1.8	13
54	DFT-Based Method for More Accurate Adsorption Energies: An Adaptive Sum of Energies from RPBE and vdW Density Functionals. Journal of Physical Chemistry C, 2017, 121, 4937-4945.	1.5	80

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55	Innenrücktitelbild: Catalytic Reaction Rates Controlled by Metal Oxidation State: Câ^'H Bond Cleavage in Methane over Nickelâ€Based Catalysts (Angew. Chem. 13/2017). Angewandte Chemie, 2017, 129, 3777-3777.	1.6	0
56	Catalytic Reaction Rates Controlled by Metal Oxidation State: Câ^'H Bond Cleavage in Methane over Nickelâ€Based Catalysts. Angewandte Chemie - International Edition, 2017, 56, 3557-3561.	7.2	39
57	Catalytic Reaction Rates Controlled by Metal Oxidation State: Câ^'H Bond Cleavage in Methane over Nickelâ€Based Catalysts. Angewandte Chemie, 2017, 129, 3611-3615.	1.6	16
58	Effect of pyrolysis temperature on aromatic cluster size of cellulose char by quantitative multi cross-polarization 13C NMR with long range dipolar dephasing. Carbon, 2017, 116, 210-222.	5.4	22
59	Role of Carbon Monoxide in Catalyst Reconstruction for CO Hydrogenation: First-Principles Study of the Composition, Structure, and Stability of Cu/Co(10112) as a Function of CO Pressure. Journal of Physical Chemistry C, 2017, 121, 2181-2191.	1.5	20
60	Reducing Reaction Temperature, Steam Requirements, and Coke Formation During Methane Steam Reforming Using Electric Fields: A Microkinetic Modeling and Experimental Study. ACS Catalysis, 2017, 7, 6957-6968.	5.5	30
61	Chemical Sensitivity of Valence-to-Core X-ray Emission Spectroscopy Due to the Ligand and the Oxidation State: A Computational Study on Cu-SSZ-13 with Multiple H ₂ O and NH ₃ Adsorption. Journal of Physical Chemistry C, 2017, 121, 25759-25767.	1.5	12
62	Improving Ni Catalysts Using Electric Fields: A DFT and Experimental Study of the Methane Steam Reforming Reaction. ACS Catalysis, 2017, 7, 551-562.	5.5	73
63	Characterizing the geometric and electronic structure of defects in the "29―copper surface oxide. Journal of Chemical Physics, 2017, 147, 224706.	1.2	9
64	Perspective on Catalytic Hydrodeoxygenation of Biomass Pyrolysis Oils: Essential Roles of Fe-Based Catalysts. Catalysis Letters, 2016, 146, 1621-1633.	1.4	42
65	Structurally Accurate Model for the "29―Structure of Cu _{<i>x</i>} O/Cu(111): A DFT and STM Study. Journal of Physical Chemistry C, 2016, 120, 10879-10886.	1.5	45
66	Improving the deconvolution and interpretation of XPS spectra from chars by ab initio calculations. Carbon, 2016, 110, 155-171.	5.4	351
67	CO Adsorption on the "29―Cu _{<i>x</i>} O/Cu(111) Surface: An Integrated DFT, STM, and TPD Study. Journal of Physical Chemistry C, 2016, 120, 25387-25394.	1.5	24
68	Elucidating the Role of the Electric Field at the Ni/YSZ Electrode: A DFT Study. Journal of Physical Chemistry C, 2016, 120, 14608-14620.	1.5	12
69	Elucidating the field influence on the energetics of the methane steam reforming reaction: A density functional theory study. Applied Catalysis B: Environmental, 2016, 195, 77-89.	10.8	45
70	Ab initio X-ray absorption modeling of Cu-SAPO-34: Characterization of Cu exchange sites under different conditions. Catalysis Today, 2016, 267, 28-40.	2.2	10
71	Structural analysis of char by Raman spectroscopy: Improving band assignments through computational calculations from first principles. Carbon, 2016, 100, 678-692.	5.4	288
72	Adsorption of guaiacol on Fe (110) and Pd (111) from first principles. Surface Science, 2016, 648, 227-235.	0.8	36

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73	The interaction of reactants, intermediates and products with Cu ions in Cu-SSZ-13 NH ₃ SCR catalysts: an energetic and ab initio X-ray absorption modeling study. Catalysis Science and Technology, 2016, 6, 5812-5829.	2.1	23
74	CO-induced inversion of the layer sequence of a model CoCu catalyst. Surface Science, 2016, 648, 74-83.	0.8	30
75	Catalytic water dehydrogenation and formation on nickel: Dual path mechanism in high electric fields. Journal of Catalysis, 2015, 332, 187-200.	3.1	52
76	Adsorption of aromatics on the (111) surface of PtM and PtM ₃ (M = Fe, Ni) alloys. RSC Advances, 2015, 5, 85705-85719.	1.7	14
77	Phenol Deoxygenation Mechanisms on Fe(110) and Pd(111). ACS Catalysis, 2015, 5, 523-536.	5.5	116
78	First-principles reaction site model for coverage-sensitive surface reactions: Pt(111)–O temperature programmed desorption. Surface Science, 2014, 622, L1-L6.	0.8	24
79	Density functional theory studies of methyl dissociation on a Ni(111) surface in the presence of an external electric field. Physical Chemistry Chemical Physics, 2014, 16, 2399-2410.	1.3	31
80	Decomposition of methyl species on a Ni(211) surface: investigations of the electric field influence. Catalysis Science and Technology, 2014, 4, 4020-4035.	2.1	29
81	Synergistic Catalysis between Pd and Fe in Gas Phase Hydrodeoxygenation of <i>m</i> -Cresol. ACS Catalysis, 2014, 4, 3335-3345.	5.5	173
82	Enhanced Fe ₂ O ₃ Reducibility via Surface Modification with Pd: Characterizing the Synergy within Pd/Fe Catalysts for Hydrodeoxygenation Reactions. ACS Catalysis, 2014, 4, 3381-3392.	5.5	114
83	Adsorption of phenol on Fe (110) and Pd (111) from first principles. Surface Science, 2014, 630, 244-253.	0.8	52
84	NO Chemisorption on Cu/SSZ-13: A Comparative Study from Infrared Spectroscopy and DFT Calculations. ACS Catalysis, 2014, 4, 4093-4105.	5.5	139
85	Carbon-supported bimetallic Pd–Fe catalysts for vapor-phase hydrodeoxygenation of guaiacol. Journal of Catalysis, 2013, 306, 47-57.	3.1	384
86	Tailoring the Adsorption of Benzene on PdFe Surfaces: A Density Functional Theory Study. Journal of Physical Chemistry C, 2013, 117, 24317-24328.	1.5	45
87	Integrated X-ray photoelectron spectroscopy and DFT characterization of benzene adsorption on Pt(111), Pt(355) and Pt(322) surfaces. Physical Chemistry Chemical Physics, 2013, 15, 20662.	1.3	25
88	How low can you go? Minimum energy pathways for O2 dissociation on Pt(111). Physical Chemistry Chemical Physics, 2012, 14, 16677.	1.3	50
89	Integrated operando X-ray absorption and DFT characterization of Cu–SSZ-13 exchange sites during the selective catalytic reduction of NO with NH3. Catalysis Today, 2012, 184, 129-144.	2.2	212
90	Electric field induced oscillations in the catalytic water production on rhodium: A theoretical analysis. Surface Science, 2010, 604, 1353-1368.	0.8	22

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91	Non-equilibrium surface pattern formation during catalytic reactions with nanoscale resolution: Investigations of the electric field influence. Catalysis Today, 2010, 154, 75-84.	2.2	14
92	Catalytic Reduction of NO ₂ with Hydrogen on Pt Field Emitter Tips: Kinetic Instabilities on the Nanoscale. Langmuir, 2010, 26, 16381-16391.	1.6	20
93	Nanometric chemical clocks. Proceedings of the National Academy of Sciences of the United States of America, 2009, 106, 3006-3010.	3.3	50
94	Oscillations and Bistability in the Catalytic Formation of Water on Rhodium in High Electric Fields. Journal of Physical Chemistry C, 2009, 113, 17045-17058.	1.5	45
95	Field-assisted oxidation of rhodium. Chemical Physics Letters, 2008, 452, 133-138.	1.2	26
96	Hopping kinetics on a finite 1D chain: An exact analysis. International Journal of Quantum Chemistry, 2006, 106, 2889-2903.	1.0	5
97	Lateral interactions and nonequilibrium in adsorption and desorption. Part 2. A kinetic lattice gas model for (2×2)-(3O+NO)/Ru(001). Surface Science, 2006, 600, 4660-4669.	0.8	8
98	CO oxidation on electrically charged gold nanotips. Journal of Chemical Physics, 2006, 125, 214707.	1.2	11
99	Two-Dimensional Roughening of Adsorbate Islands in Thermodynamic Equilibrium. Physical Review Letters, 2006, 96, 166102.	2.9	7
100	Adsorption and desorption of CO on Ru(0001): A comprehensive analysis. Surface Science, 2005, 594, 240-262.	0.8	52
101	Adsorption and desorption of CO on Pt(111): a comprehensive analysis. Surface Science, 2003, 545, 47-69.	0.8	99
102	Phase diagram of O/Ru(0001) from first principles. Chemical Physics Letters, 2002, 361, 317-320.	1.2	23
103	Quantifying Errors in Effective Cluster Interactions of Lattice Gas Cluster Expansions. Journal of Physical Chemistry C, 0, , .	1.5	2