# Pascal Raybaud

# List of Publications by Citations

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8,846 135 52 91 h-index g-index citations papers 6.16 9,613 6.5 140 avg, IF L-index ext. papers ext. citations

#	Paper	IF	Citations
135	Use of DFT to achieve a rational understanding of acidBasic properties of ⊞lumina surfaces. Journal of Catalysis, <b>2004</b> , 226, 54-68	7:3	721
134	Theoretical Study of the Dehydration Process of Boehmite to EAlumina. <i>Journal of Physical Chemistry B</i> , <b>2001</b> , 105, 5121-5130	3.4	362
133	Hydroxyl Groups on EAlumina Surfaces: A DFT Study. <i>Journal of Catalysis</i> , <b>2002</b> , 211, 1-5	7:3	295
132	Ab Initio Study of the H2H2S/MoS2 GasBolid Interface: The Nature of the Catalytically Active Sites. <i>Journal of Catalysis</i> , <b>2000</b> , 189, 129-146	7:3	292
131	Shape and Edge Sites Modifications of MoS2 Catalytic Nanoparticles Induced by Working Conditions: A Theoretical Study. <i>Journal of Catalysis</i> , <b>2002</b> , 207, 76-87	7.3	285
130	Effects of morphology on surface hydroxyl concentration: a DFT comparison of anataselliO2 and Ealumina catalytic supports. <i>Journal of Catalysis</i> , <b>2004</b> , 222, 152-166	7:3	266
129	Structure, Energetics, and Electronic Properties of the Surface of a Promoted MoS2 Catalyst: An ab Initio Local Density Functional Study. <i>Journal of Catalysis</i> , <b>2000</b> , 190, 128-143	7.3	266
128	Structure and Stability of Aluminum Hydroxides: A Theoretical Study. <i>Journal of Physical Chemistry B</i> , <b>2002</b> , 106, 5155-5162	3.4	209
127	Morphology and Surface Properties of Boehmite (FAlOOH): A Density Functional Theory Study. Journal of Catalysis, <b>2001</b> , 201, 236-246	7.3	183
126	Challenges on molecular aspects of dealumination and desilication of zeolites. <i>Microporous and Mesoporous Materials</i> , <b>2014</b> , 191, 82-96	5.3	173
125	Optimal promoter edge decoration of CoMoS catalysts: A combined theoretical and experimental study. <i>Catalysis Today</i> , <b>2008</b> , 130, 149-159	5.3	164
124	Periodic trends in hydrodesulfurization: in support of the Sabatier principle. <i>Applied Catalysis A: General</i> , <b>2002</b> , 227, 83-96	5.1	161
123	Kinetic interpretation of catalytic activity patterns based on theoretical chemical descriptors. <i>Journal of Catalysis</i> , <b>2003</b> , 216, 63-72	7-3	146
122	Promoter Sensitive Shapes of Co(Ni)MoS Nanocatalysts in Sulfo-Reductive Conditions. <i>Journal of Catalysis</i> , <b>2002</b> , 212, 33-38	7.3	144
121	H2-Induced Reconstruction of Supported Pt Clusters: MetalBupport Interaction versus Surface Hydride. <i>ChemCatChem</i> , <b>2011</b> , 3, 200-207	5.2	132
120	Mixed sites and promoter segregation: A DFT study of the manifestation of Le Chatelier's principle for the Co(Ni)MoS active phase in reaction conditions. <i>Catalysis Today</i> , <b>2008</b> , 130, 160-169	5.3	129
119	Adsorption of Unsaturated Hydrocarbons on Pd(111) and Pt(111): A DFT Study. <i>Journal of Physical Chemistry B</i> , <b>2003</b> , 107, 12287-12295	3.4	129

### (2005-1999)

118	Sabatier with first principle calculations1This work has been undertaken within the AdR  Dynamique MolBulaire Quantique Applique Ila Catalyse Ia joint project of Centre National de la	5.3	128	
117	Recherche Scientifique, Technische Universitä Wien, and Institut Frantiis du Pfirole.1. Catalysis Modulation of Catalyst particle structure upon support hydroxylation: Ab initio insights into Pd13 and Pt13/EAl2O3. Journal of Catalysis, 2010, 274, 99-110	7.3	125	
116	Understanding and predicting improved sulfide catalysts: Insights from first principles modeling. <i>Applied Catalysis A: General</i> , <b>2007</b> , 322, 76-91	5.1	117	
115	Adsorption of Thiophene on the Catalytically Active Surface of MoS2: An Ab Initio Local-Density-Functional Study. <i>Physical Review Letters</i> , <b>1998</b> , 80, 1481-1484	7.4	114	
114	Ab initiodensity functional studies of transition-metal sulphides: I. Crystal structure and cohesive properties. <i>Journal of Physics Condensed Matter</i> , <b>1997</b> , 9, 11085-11106	1.8	106	
113	Free-energy profiles along reduction pathways of MoS2 M-edge and S-edge by dihydrogen: A first-principles study. <i>Journal of Catalysis</i> , <b>2011</b> , 280, 178-195	7.3	105	
112	Hemilabile Ligand Induced Selectivity: a DFT Study on Ethylene Trimerization Catalyzed by Titanium Complexes. <i>Organometallics</i> , <b>2003</b> , 22, 3404-3413	3.8	105	
111	Dealumination mechanisms of zeolites and extra-framework aluminum confinement. <i>Journal of Catalysis</i> , <b>2016</b> , 339, 242-255	7.3	105	
110	Atomic Description of the Interface between Silica and Alumina in Aluminosilicates through Dynamic Nuclear Polarization Surface-Enhanced NMR Spectroscopy and First-Principles Calculations. <i>Journal of the American Chemical Society</i> , <b>2015</b> , 137, 10710-9	16.4	104	
109	Pseudo-bridging silanols as versatile Brfisted acid sites of amorphous aluminosilicate surfaces. <i>Angewandte Chemie - International Edition</i> , <b>2009</b> , 48, 2891-3	16.4	104	
108	The role of the extra-framework cations in the adsorption of CO(2) on faujasite Y. <i>Physical Chemistry Chemical Physics</i> , <b>2010</b> , 12, 13534-46	3.6	97	
107	Hydrodeoxygenation pathways catalyzed by MoS2 and NiMoS active phases: A DFT study. <i>Journal of Catalysis</i> , <b>2011</b> , 279, 276-286	7.3	94	
106	Deoxygenation mechanisms on Ni-promoted MoS2 bulk catalysts: A combined experimental and theoretical study. <i>Journal of Catalysis</i> , <b>2012</b> , 286, 153-164	7.3	92	
105	Ab initiodensity functional studies of transition-metal sulphides: II. Electronic structure. <i>Journal of Physics Condensed Matter</i> , <b>1997</b> , 9, 11107-11140	1.8	92	
104	Structural and electronic properties of the MoS2(101 0) edge-surface. <i>Surface Science</i> , <b>1998</b> , 407, 237-2	2 <b>50</b> 8	92	
103	Brfisted acidity of amorphous silicallumina: The molecular rules of proton transfer. <i>Journal of Catalysis</i> , <b>2011</b> , 284, 215-229	7.3	87	
102	Origin of the Enhanced Visible-Light Absorption in N-Doped Bulk Anatase TiO2 from First-Principles Calculations. <i>Journal of Physical Chemistry C</i> , <b>2011</b> , 115, 19394-19404	3.8	86	
101	A density functional theory comparison of anatase (TiO2)- and EAl2O3-supported MoS2 catalysts.  Journal of Catalysis, <b>2005</b> , 232, 161-178	7.3	85	

100	The Origin of the C7-Hydroconversion Selectivities on Y, ∏ZSM-22, ZSM-23, and EU-1 Zeolites. <i>Journal of Catalysis</i> , <b>2001</b> , 197, 98-112	7.3	85
99	Influence of the hydroxylation of gamma-Al2O3 surfaces on the stability and diffusion of single Pd atoms: a DFT study. <i>Journal of Physical Chemistry B</i> , <b>2006</b> , 110, 1759-67	3.4	84
98	Edge wetting effects of FAl2O3 and anatase-TiO2 supports by MoS2 and CoMoS active phases: A DFT study. <i>Journal of Catalysis</i> , <b>2007</b> , 246, 325-343	7.3	80
97	Tuning the properties of visible-light-responsive tantalum (oxy)nitride photocatalysts by non-stoichiometric compositions: a first-principles viewpoint. <i>Physical Chemistry Chemical Physics</i> , <b>2014</b> , 16, 20548-60	3.6	77
96	Nucleation of Pdn (n=15) clusters and wetting of Pd particles on 🛭 203 surfaces: A density functional theory study. <i>Physical Review B</i> , <b>2007</b> , 75,	3.3	76
95	Interplay between molecular adsorption and metal@upport interaction for small supported metal clusters: CO and C2H4 adsorption on Pd4/FAl2O3. <i>Journal of Catalysis</i> , <b>2007</b> , 247, 339-355	7-3	73
94	Anionic or Cationic S-Doping in Bulk Anatase TiO2: Insights on Optical Absorption from First Principles Calculations. <i>Journal of Physical Chemistry C</i> , <b>2013</b> , 117, 8892-8902	3.8	70
93	Predictive approach for the design of improved HDT catalysts: EAlumina supported (Ni, Co) promoted Mo1\( \text{W}\) xS2 active phases. <i>Applied Catalysis A: General</i> , <b>2007</b> , 322, 92-97	5.1	70
92	Theoretical Unraveling of Selective 1-Butene Oligomerization Catalyzed by Iron <b>B</b> is(arylimino)pyridine. <i>Organometallics</i> , <b>2009</b> , 28, 5358-5367	3.8	68
91	Aging of Co(Ni)MoP/Al2O3 catalysts in working state. <i>Catalysis Today</i> , <b>2008</b> , 130, 97-108	5.3	67
90	Platinum Nanoclusters Stabilized on FAlumina by Chlorine Used As a Capping Surface Ligand: A Density Functional Theory Study. <i>ACS Catalysis</i> , <b>2012</b> , 2, 1346-1357	13.1	62
89	Effects of PH2O, PH2S, PH2 on the surface properties of anatase?TiO2 and ?-Al2O3: a DFT study. Journal of Catalysis, <b>2004</b> , 226, 260-272	7.3	62
88	A DFT study of the origin of the HDS/HydO selectivity on Co(Ni)MoS active phases. <i>Journal of Catalysis</i> , <b>2008</b> , 260, 276-287	7.3	60
87	Regioselectivity of Al® Bond Hydrolysis during Zeolites Dealumination Unified by Brlisted Evans Polanyi Relationship. ACS Catalysis, 2015, 5, 11-15	13.1	56
86	Quantitative Two-Dimensional (2D) MorphologyBelectivity Relationship of CoMoS Nanolayers: A Combined High-Resolution High-Angle Annular Dark Field Scanning Transmission Electron Microscopy (HR HAADF-STEM) and Density Functional Theory (DFT) Study. <i>ACS Catalysis</i> , <b>2016</b> , 6, 1081-	13.1 ·1092	56
85	DFT makes the morphologies of anatase-TiO2 nanoparticles visible to IR spectroscopy. <i>Journal of Catalysis</i> , <b>2005</b> , 236, 245-250	7.3	56
84	New insights into parameters controlling the selectivity in hydrocracking reactions. <i>Journal of Catalysis</i> , <b>2003</b> , 217, 376-387	7.3	54
83	From Ealumina to supported platinum nanoclusters in reforming conditions: 10 years of DFT modeling and beyond. <i>Journal of Catalysis</i> , <b>2013</b> , 308, 328-340	7.3	52

# (2013-2013)

82	Cobalt Catalyzed Fischer Tropsch Synthesis: Perspectives Opened by First Principles Calculations. <i>Catalysis Letters</i> , <b>2013</b> , 143, 1-17	2.8	52	
81	Growth of boehmite particles in the presence of xylitol: morphology oriented by the nest effect of hydrogen bonding. <i>Physical Chemistry Chemical Physics</i> , <b>2009</b> , 11, 11310-23	3.6	50	
80	Atomic scale insights on chlorinated gamma-alumina surfaces. <i>Journal of the American Chemical Society</i> , <b>2008</b> , 130, 11030-9	16.4	50	
79	Acidity of amorphous silica-alumina: from coordination promotion of Lewis sites to proton transfer. <i>ChemPhysChem</i> , <b>2010</b> , 11, 105-8	3.2	49	
78	Periodic trends in the selective hydrogenation of styrene over silica supported metal catalysts. Journal of Catalysis, <b>2013</b> , 307, 352-361	7.3	47	
77	Density functional theory simulations of complex catalytic materials in reactive environments: beyond the ideal surface at low coverage. <i>Catalysis Science and Technology</i> , <b>2014</b> , 4, 2797-2813	5.5	45	
76	Competitive adsorption of nitrogen and sulphur compounds on a multisite model of NiMoS catalyst: A theoretical study. <i>Journal of Catalysis</i> , <b>2016</b> , 333, 78-93	7.3	44	
75	Tuning the Magnetic Properties of MoS2 Single Nanolayers by 3d Metals Edge Doping. <i>Journal of Physical Chemistry C</i> , <b>2016</b> , 120, 10691-10697	3.8	43	
74	A QSPR Investigation of Thermal Stability of [Al(CH3)O]n Oligomers in Methylaluminoxane Solution: The Identification of a Geometry-Based Descriptor. <i>Organometallics</i> , <b>2012</b> , 31, 8312-8322	3.8	40	
73	Comment on "Examination of spinel and nonspinel structural models for gamma-Al2O3 by DFT and Rietveld refinement simulations". <i>Journal of Physical Chemistry B</i> , <b>2006</b> , 110, 20719-20; author reply 20	7246	39	
72	Potassium silanide (KSiH3): a reversible hydrogen storage material. <i>Chemistry - A European Journal</i> , <b>2011</b> , 17, 12302-9	4.8	38	
71	Revisiting carbenium chemistry on amorphous silica-alumina: Unraveling their milder acidity as compared to zeolites. <i>Journal of Catalysis</i> , <b>2015</b> , 325, 35-47	7.3	37	
70	Thermodynamic Stability of Buta-1,3-diene and But-1-ene on Pd(111) and (100) Surfaces under H2 Pressure: A DFT Study. <i>Journal of Physical Chemistry C</i> , <b>2011</b> , 115, 12135-12149	3.8	37	
69	Highly Active Nonpromoted Hydrotreating Catalysts through the Controlled Growth of a Supported Hexagonal WS2 Phase. <i>ACS Catalysis</i> , <b>2014</b> , 4, 4320-4331	13.1	35	
68	Evidence for the Iron(III) Oxidation State in Bis(imino)pyridine Catalysts. A Density Functional Theory Study. <i>Organometallics</i> , <b>2008</b> , 27, 3368-3377	3.8	35	
67	Microkinetic interpretation of HDS/HYDO selectivity of the transformation of a model FCC gasoline over transition metal sulfides. <i>Catalysis Today</i> , <b>2008</b> , 130, 221-230	5.3	34	
66	Transformation of a model FCC gasoline olefin over transition monometallic sulfide catalysts. <i>Journal of Catalysis</i> , <b>2007</b> , 248, 111-119	7.3	33	
65	A rational interpretation of improved catalytic performances of additive-impregnated dried CoMo hydrotreating catalysts: a combined theoretical and experimental study. <i>Catalysis Science and Technology</i> , <b>2013</b> , 3, 140-151	5.5	32	

64	CO adsorption on amorphous silica-alumina: electrostatic or Brfisted acidity probe?. <i>Chemical Communications</i> , <b>2012</b> , 48, 4076-8	5.8	32
63	A Rational Comparison of the Optimal Promoter Edge Decoration of HDT NiMoS vs CoMoS Catalysts. <i>Oil and Gas Science and Technology</i> , <b>2009</b> , 64, 719-730	1.9	32
62	Temperature-programed reduction of unpromoted MoS2-based hydrodesulfurization catalysts: Experiments and kinetic modeling from first principles. <i>Journal of Catalysis</i> , <b>2009</b> , 267, 67-77	7.3	32
61	THERMIDOR: A new model for combined simulation of operations and optimization of catalysts in residues hydroprocessing units. <i>Catalysis Today</i> , <b>2005</b> , 109, 135-153	5.3	32
60	Effect of confinement on the selectivity of hydrocracking. <i>Journal of Catalysis</i> , <b>2004</b> , 221, 500-509	7.3	31
59	Dehydrogenation mechanisms of methyl-cyclohexane on EAl2O3 supported Pt13: Impact of cluster ductility. <i>Journal of Catalysis</i> , <b>2019</b> , 370, 118-129	7-3	31
58	A DFT Study of CoMoS and NiMoS Catalysts: from Nano-Crystallite Morphology to Selective Hydrodesulfurization. <i>Oil and Gas Science and Technology</i> , <b>2009</b> , 64, 707-718	1.9	30
57	Structural, energetic, and electronic trends in low-dimensional late-transition-metal systems. <i>Physical Review B</i> , <b>2009</b> , 79,	3.3	29
56	Insights into the Geometry, Stability and Vibrational Properties of OH Groups on FAl2O3, TiO2-Anatase and MgO from DFT Calculations. <i>Topics in Catalysis</i> , <b>2009</b> , 52, 1005-1016	2.3	29
55	Effects of zeolite pore sizes on the mechanism and selectivity of xylene disproportionation DFT study. <i>Journal of Catalysis</i> , <b>2004</b> , 222, 323-337	7.3	29
54	Stability of Carbon on Cobalt Surfaces in Fischer Tropsch Reaction Conditions: A DFT Study. <i>Journal of Physical Chemistry C</i> , <b>2014</b> , 118, 22479-22490	3.8	28
53	An Atomistic Description of the EAlumina/Water Interface Revealed by Ab Initio Molecular Dynamics. <i>Journal of Physical Chemistry C</i> , <b>2017</b> , 121, 10351-10363	3.8	27
52	Understanding the role of aluminum-based activators in single site iron catalysts for ethylene oligomerization. <i>Journal of Catalysis</i> , <b>2014</b> , 317, 153-157	7.3	27
51	Tuning the metal-support interaction by structural recognition of cobalt-based catalyst precursors. <i>Angewandte Chemie - International Edition</i> , <b>2015</b> , 54, 6824-7	16.4	25
50	Ab Initio Simulation of the Acid Sites at the External Surface of Zeolite Beta. <i>ChemCatChem</i> , <b>2017</b> , 9, 2176-2185	5.2	24
49	Deep HDS of FCC gasoline over alumina supported CoMoS catalyst: Inhibiting effects of carbon monoxide and water. <i>Applied Catalysis B: Environmental</i> , <b>2016</b> , 183, 317-327	21.8	24
48	Enthalpy <b>E</b> ntropy Compensation Effect in Hydrogen Storage Materials: Striking Example of Alkali Silanides MSiH3 (M = K, Rb, Cs). <i>Journal of Physical Chemistry C</i> , <b>2014</b> , 118, 3409-3419	3.8	23
47	Quantum chemical and vibrational investigation of sodium exchanged gamma-alumina surfaces.  Physical Chemistry Chemical Physics, 2007, 9, 2577-82	3.6	23

# (2008-2016)

46	Improved promoter effect in NiWS catalysts through a molecular approach and an optimized Ni edge decoration. <i>Journal of Catalysis</i> , <b>2016</b> , 340, 60-65	7.3	22	
45	A DFT Chemical Descriptor to Predict the Selectivity in Đlefins in the Catalytic Metallacyclic Oligomerization Reaction of Ethylene According to the (Hemi)labile Ligand Coordinating to Titanium. <i>Organometallics</i> , <b>2008</b> , 27, 4864-4872	3.8	22	
44	Dual Effect of H2S on Volcano Curves in Hydrotreating Sulfide Catalysis. <i>Oil and Gas Science and Technology</i> , <b>2006</b> , 61, 515-525	1.9	22	
43	Beyond EAl2O3 crystallite surfaces: The hidden features of edges revealed by solid-state 1H NMR and DFT calculations. <i>Journal of Catalysis</i> , <b>2019</b> , 378, 140-143	7.3	20	
42	On the origin of the difference between type A and type B skeletal isomerization of alkenes catalyzed by zeolites: The crucial input of ab initio molecular dynamics. <i>Journal of Catalysis</i> , <b>2019</b> , 373, 361-373	7.3	20	
41	Monitoring Morphology and Hydrogen Coverage of Nanometric Pt/EAl2O3 Particles by In Situ HERFDXANES and Quantum Simulations. <i>Angewandte Chemie</i> , <b>2014</b> , 126, 12634-12637	3.6	20	
40	Monitoring morphology and hydrogen coverage of nanometric Pt/EAl2 O3 particles by in situ HERFD-XANES and quantum simulations. <i>Angewandte Chemie - International Edition</i> , <b>2014</b> , 53, 12426-9	16.4	20	
39	Effect of Indium Doping of EAlumina on the Stabilization of PtSn Alloyed Clusters Prepared by Surface Organostannic Chemistry. <i>Journal of Physical Chemistry C</i> , <b>2012</b> , 116, 10073-10083	3.8	20	
38	On the understanding of the optoelectronic properties of S-doped MoO3 and O-doped MoS2 bulk systems: a DFT perspective. <i>Journal of Materials Chemistry C</i> , <b>2020</b> , 8, 9064-9074	7.1	19	
37	Temperature-programmed reduction of unpromoted MoS2-based hydrodesulfurization catalysts: First-principles kinetic Monte Carlo simulations and comparison with experiments. <i>Journal of Catalysis</i> , <b>2010</b> , 275, 117-128	7.3	19	
36	Competition of Secondary versus Tertiary Carbenium Routes for the Type B Isomerization of Alkenes over Acid Zeolites Quantified by Ab Initio Molecular Dynamics Simulations. <i>ACS Catalysis</i> , <b>2019</b> , 9, 9813-9828	13.1	18	
35	Hydrogenation properties of KSi and NaSi Zintl phases. <i>Physical Chemistry Chemical Physics</i> , <b>2012</b> , 14, 13319-24	3.6	18	
34	Pseudo-Bridging Silanols as Versatile Brilsted Acid Sites of Amorphous Aluminosilicate Surfaces. <i>Angewandte Chemie</i> , <b>2009</b> , 121, 2935-2937	3.6	18	
33	First principles surface thermodynamics of industrial supported catalysts in working conditions. Journal of Physics Condensed Matter, <b>2008</b> , 20, 064235	1.8	17	
32	Active sites speciation of supported CoMoS phase probed by NO molecule: A combined IR and DFT study. <i>Journal of Catalysis</i> , <b>2018</b> , 361, 62-72	7.3	16	
31	Magnifying the Morphology Change Induced by a Nickel Promoter in Tungsten(IV) Sulfide Industrial Hydrocracking Catalyst: A HAADF-STEM and DFT Study. <i>ChemCatChem</i> , <b>2014</b> , 6, 1594-1598	5.2	15	
30	Iron bis(arylimino)pyridine precursors activated to catalyze ethylene oligomerization as studied by DFT and QSAR approaches. <i>Computational and Theoretical Chemistry</i> , <b>2009</b> , 903, 100-107		15	
29	Thermodynamic Properties of Trialkali (Li, Na, K) Hexa-alanates: A Combined DFT and Experimental Study. <i>Journal of Physical Chemistry C</i> , <b>2008</b> , 112, 18598-18607	3.8	15	

28	Compensation effect and volcano curve in toluene hydrogenation catalyzed by transition metal sulfides. <i>Dalton Transactions</i> , <b>2010</b> , 39, 8420-2	4.3	14
27	Atomic Scale Insight into the Formation, Size, and Location of Platinum Nanoparticles Supported on FAlumina. <i>ACS Catalysis</i> , <b>2020</b> , 10, 4193-4204	13.1	12
26	Impact of CO on the transformation of a model FCC gasoline over CoMoS/Al2O3 catalysts: A combined kinetic and DFT approach. <i>Applied Catalysis B: Environmental</i> , <b>2010</b> , 97, 323-332	21.8	11
25	Adsorption of NO on Pd-Exchanged Mordenite: Ab Initio DFT Modeling. <i>Journal of Physical Chemistry C</i> , <b>2008</b> , 112, 12349-12362	3.8	11
24	Dynamic Features of Transition States for Escission Reactions of Alkenes over Acid Zeolites Revealed by AIMD Simulations. <i>Angewandte Chemie - International Edition</i> , <b>2020</b> , 59, 18938-18942	16.4	10
23	Topological Analysis of the Interactions between Organic Molecules and Co(Ni)MoS Catalytic Active Phases. <i>Journal of Chemical Theory and Computation</i> , <b>2009</b> , 5, 580-93	6.4	10
22	Atomistic Models for Highly-Dispersed PtSn/EAl2O3 Catalysts: Ductility and Dilution Affect the Affinity for Hydrogen. <i>ChemCatChem</i> , <b>2019</b> , 11, 3941-3951	5.2	9
21	In Silico Prediction of Catalytic Oligomerization Degrees. <i>Organometallics</i> , <b>2011</b> , 30, 3911-3914	3.8	9
20	Interplay of the adsorption of light and heavy paraffins in hydroisomerization over H-beta zeolite. <i>Catalysis Science and Technology</i> , <b>2019</b> , 9, 5368-5382	5.5	8
19	Theoretical Insights into the Interaction of Oxygenated Organic Molecules and Cobalt(II) Precursor with EAl2O3 Surfaces. <i>Journal of Physical Chemistry C</i> , <b>2018</b> , 122, 19560-19574	3.8	7
18	Comment on "Electronic properties and charge transfer phenomena in Pt nanoparticles on EAl2O3: size, shape, support, and adsorbate effects" by F. Behafarid et al., Phys. Chem. Chem. Phys., 2012, 14, 11766-11779. <i>Physical Chemistry Chemical Physics</i> , <b>2012</b> , 14, 16773-4	3.6	7
17	Catalytic Reforming: Methodology and Process Development for a Constant Optimisation and Performance Enhancement. <i>Oil and Gas Science and Technology</i> , <b>2016</b> , 71, 41	1.9	7
16	Combined Experimental and Theoretical Molecular Approach of the Catalytically Active Hydrotreating MoS2 Phases Promoted by 3d Transition Metals. <i>Journal of Physical Chemistry C</i> , <b>2019</b> , 123, 24659-24669	3.8	6
15	Prediction of optimal catalysts for a given chemical reaction. <i>Catalysis Science and Technology</i> , <b>2020</b> , 10, 2069-2081	5.5	6
14	2D MoOS/MoS van der Waals Assembly: A Tunable Heterojunction with Attractive Properties for Photocatalysis. <i>ACS Applied Materials &amp; Amp; Interfaces</i> , <b>2021</b> , 13, 36465-36474	9.5	6
13	Competitive Deposition of C and O Species on Cobalt Surface in Fischer Tropsch Synthesis Conditions: A Plausible Origin of Deactivation. <i>Journal of Physical Chemistry C</i> , <b>2015</b> , 119, 23515-23526	3.8	5
12	Surface speciation of Co based Fischer-Tropsch catalyst under reaction conditions: Deactivation by coke or by oxidation?. <i>Applied Catalysis A: General</i> , <b>2020</b> , 590, 117332	5.1	5
11	Computational chemistry approaches for the preparation of supported catalysts: Progress and challenges. <i>Journal of Catalysis</i> , <b>2020</b> , 391, 539-547	7.3	5

#### LIST OF PUBLICATIONS

10	Electronic structures of the MoS/TiO (anatase) heterojunction: influence of physical and chemical modifications at the 2D- or 1D-interfaces <i>Physical Chemistry Chemical Physics</i> , <b>2022</b> , 24, 2646-2655	3.6	2
9	Multiscale Modeling as a Tool for the Prediction of Catalytic Performances: The Case of n-Heptane Hydroconversion in a Large-Pore Zeolite. <i>ACS Catalysis</i> ,1068-1081	13.1	2
8	Dynamic Features of Transition States for Escission Reactions of Alkenes over Acid Zeolites Revealed by AIMD Simulations. <i>Angewandte Chemie</i> , <b>2020</b> , 132, 19100-19104	3.6	2
7	Size-Dependent Structural, Energetic, and Spectroscopic Properties of MoS3 Polymorphs. <i>Crystal Growth and Design</i> , <b>2020</b> , 20, 7750-7760	3.5	2
6	Structural Characterization of Phosphate Species Adsorbed on EAlumina by Combining DNP Surface Enhanced NMR Spectroscopy and DFT Calculations. <i>ACS Catalysis</i> , <b>2021</b> , 11, 11278-11292	13.1	2
5	Thermokinetic and Spectroscopic Mapping of Carbon Monoxide Adsorption on Highly Dispersed Pt/EAl2O3. <i>ACS Catalysis</i> ,13280-13293	13.1	1
4	Tuning the MetalBupport Interaction by Structural Recognition of Cobalt-Based Catalyst Precursors. <i>Angewandte Chemie</i> , <b>2015</b> , 127, 6928-6931	3.6	О
3	Genesis of MoS2 from model-Mo-oxide precursors supported on 🗟 lumina. <i>Journal of Catalysis</i> , <b>2022</b> , 408, 303-315	7-3	O
2	Evidence for H2-Induced Ductility in a Pt/Al2O3 Catalyst. ACS Catalysis,5979-5989	13.1	О
1	Hydrogenolysis and Blimination mechanisms for C S bond scission of dibenzothiophene on CoMoS edge sites. <i>Journal of Catalysis</i> , <b>2021</b> , 403, 32-32	7.3	