

# Joachim Sauer

## List of Publications by Year in descending order

Source: <https://exaly.com/author-pdf/860530/publications.pdf>

Version: 2024-02-01

351  
papers

26,550  
citations

3531  
90  
h-index

8630  
146  
g-index

389  
all docs

389  
docs citations

389  
times ranked

14809  
citing authors

#	ARTICLE	IF	CITATIONS
1	Oxygen vacancies in transition metal and rare earth oxides: Current state of understanding and remaining challenges. <i>Surface Science Reports</i> , 2007, 62, 219-270.	7.2	1,102
2	Oxygen Defects and Surface Chemistry of Ceria: Quantum Chemical Studies Compared to Experiment. <i>Chemical Reviews</i> , 2013, 113, 3949-3985.	47.7	849
3	Molecular models in ab initio studies of solids and surfaces: from ionic crystals and semiconductors to catalysts. <i>Chemical Reviews</i> , 1989, 89, 199-255.	47.7	726
4	Theoretical Study of van der Waals Complexes at Surface Sites in Comparison with the Experiment. <i>Chemical Reviews</i> , 1994, 94, 2095-2160.	47.7	704
5	Hybrid functionals applied to rare-earth oxides: The example of ceria. <i>Physical Review B</i> , 2007, 75, .	3.2	502
6	Density-Functional Calculations of the Structure of Near-Surface Oxygen Vacancies and Electron Localization on<math>\langle mml:math xmlns:mml="http://www.w3.org/1998/Math/MathML"> display="inline"><mml:msub><mml:mi>CeO</mml:mi><mml:mn>2</mml:mn></mml:msub><mml:mo stretchy="false">(</mml:mo><mml:mn>111</mml:mn><mml:mo> Tj ETQq0 0 0 rgBT /Overlock 10 Tf 50 527 Td (stretchy="false")</mml:mo>	7.8	501
7	Bridging hydroxyl groups in zeolitic catalysts: a computer simulation of their structure, vibrational properties and acidity in protonated faujasites ( $\text{H}-\text{Y}$ zeolites). <i>Chemical Physics Letters</i> , 1992, 188, 320-325.	2.6	370
8	Acidity Differences between Inorganic Solids Induced by Their Framework Structure. A Combined Quantum Mechanics/Molecular Mechanics ab Initio Study on Zeolites. <i>Journal of the American Chemical Society</i> , 1998, 120, 1556-1570.	13.7	359
9	Molecular mechanics potential for silica and zeolite catalysts based on ab initio calculations. 1. Dense and microporous silica. <i>The Journal of Physical Chemistry</i> , 1994, 98, 1238-1244.	2.9	317
10	Application of semiempirical long-range dispersion corrections to periodic systems in density functional theory. <i>Journal of Computational Chemistry</i> , 2008, 29, 2088-2097.	3.3	294
11	Quantum Chemical Modeling of Zeolite-Catalyzed Methylation Reactions: Toward Chemical Accuracy for Barriers. <i>Journal of the American Chemical Society</i> , 2009, 131, 816-825.	13.7	288
12	Evolution of water structures in metal-organic frameworks for improved atmospheric water harvesting. <i>Science</i> , 2021, 374, 454-459.	12.6	281
13	$^{23}\text{Na}$ NMR Spectroscopy of Solids: Interpretation of Quadrupole Interaction Parameters and Chemical Shifts. <i>The Journal of Physical Chemistry</i> , 1994, 98, 1544-1551.	2.9	277
14	Interaction of methanol with Brønsted acid sites of zeolite catalysts: an ab initio study. <i>Journal of the American Chemical Society</i> , 1995, 117, 3780-3789.	13.7	267
15	Vibrational spectra of alumina- and silica-supported vanadia revisited: An experimental and theoretical model catalyst study. <i>Journal of Catalysis</i> , 2004, 226, 88-100.	6.2	258
16	Thermal Activation of Methane by Tetranuclear $[\text{V}_4\text{O}_{10}]^+$ . <i>Angewandte Chemie - International Edition</i> , 2006, 45, 4681-4685.	13.8	236
17	Aluminum Siting in Silicon-Rich Zeolite Frameworks: A Combined High-Resolution $\langle \sup{27} \rangle_{\text{Al}} \text{...NMR}$ Spectroscopy and Quantum Mechanics/Molecular Mechanics Study of ZSM-5. <i>Angewandte Chemie - International Edition</i> , 2007, 46, 7286-7289.	13.8	234
18	Treating dispersion effects in extended systems by hybrid MP2:DFT calculations—protonation of isobutene in zeolite ferrierite. <i>Physical Chemistry Chemical Physics</i> , 2006, 8, 3955-3965.	2.8	232

#	ARTICLE	IF	CITATIONS
19	Predicting Absolute and Site Specific Acidities for Zeolite Catalysts by a Combined Quantum Mechanics/Interatomic Potential Function Approach. <i>Journal of Physical Chemistry B</i> , 1997, 101, 10035-10050.	2.6	227
20	Ab Initio Study of Hydrogen Adsorption in MOF-5. <i>Journal of the American Chemical Society</i> , 2009, 131, 4143-4150.	13.7	225
21	Combining quantum mechanics and interatomic potential functions in ab initio studies of extended systems. <i>Journal of Computational Chemistry</i> , 2000, 21, 1470-1493.	3.3	221
22	Coordination and siting of Cu <sup>+</sup> ions in ZSM-5: A combined quantum mechanics/interatomic potential function study. <i>Physical Chemistry Chemical Physics</i> , 1999, 1, 2019-2026.	2.8	215
23	The Atomic Structure of a Metal-Supported Vitreous Thin Silica Film. <i>Angewandte Chemie - International Edition</i> , 2012, 51, 404-407.	13.8	207
24	Molecular Mechanics Potential for Silica and Zeolite Catalysts Based on ab Initio Calculations. 2. Aluminosilicates. <i>The Journal of Physical Chemistry</i> , 1995, 99, 9536-9550.	2.9	203
25	Structure and reactivity of silica and zeolite catalysts by a combined quantum mechanics[-shell-model potential approach based on DFT]. <i>Faraday Discussions</i> , 1997, 106, 41-62.	3.2	203
26	Growth and Structure of Crystalline Silica Sheet on Ru(0001). <i>Physical Review Letters</i> , 2010, 105, 146104.	7.8	198
27	Aluminium siting in the ZSM-5 framework by combination of high resolution <sup>27</sup> Al NMR and DFT/MM calculations. <i>Physical Chemistry Chemical Physics</i> , 2009, 11, 1237-1247.	2.8	196
28	Mass-selective vibrational spectroscopy of vanadium oxide cluster ions. <i>Mass Spectrometry Reviews</i> , 2007, 26, 542-562.	5.4	192
29	Role of Ceria in Oxidative Dehydrogenation on Supported Vanadia Catalysts. <i>Journal of the American Chemical Society</i> , 2010, 132, 2345-2349.	13.7	191
30	Oxidation of Methanol to Formaldehyde on Supported Vanadium Oxide Catalysts Compared to Gas Phase Molecules. <i>Journal of the American Chemical Society</i> , 2005, 127, 10861-10868.	13.7	187
31	Interaction of Water with Brønsted Acidic Sites of Zeolite Catalysts. Ab Initio Study of 1:1 and 2:1 Surface Complexes. <i>The Journal of Physical Chemistry</i> , 1996, 100, 6199-6211.	2.9	183
32	Activation of Methane by Oligomeric (Al <sub>2</sub> O <sub>3</sub> ) <sub>x</sub> : The Role of Oxygen-Centered Radicals in Thermal Hydrogen-Atom Abstraction. <i>Angewandte Chemie - International Edition</i> , 2008, 47, 1946-1950.	13.8	183
33	Combining ab initio techniques with analytical potential functions for structure predictions of large systems: Method and application to crystalline silica polymorphs. <i>Journal of Computational Chemistry</i> , 1997, 18, 463-477.	3.3	181
34	Oxidative Dehydrogenation of Propane by Monomeric Vanadium Oxide Sites on Silica Support. <i>Journal of Physical Chemistry C</i> , 2007, 111, 6041-6050.	3.1	171
35	The Surface Structure of Sulfated Zirconia: Periodic ab Initio Study of Sulfuric Acid Adsorbed on ZrO <sub>2</sub> (101) and ZrO <sub>2</sub> (001). <i>Journal of the American Chemical Society</i> , 1998, 120, 13503-13512.	13.7	168
36	Electron Localization in Defective Ceria Films: A Study with Scanning-Tunneling Microscopy and Density-Functional Theory. <i>Physical Review Letters</i> , 2011, 106, 246801.	7.8	158

#	ARTICLE	IF	CITATIONS
37	Sites for Methane Activation on Lithium-Doped Magnesium Oxide Surfaces. <i>Angewandte Chemie - International Edition</i> , 2014, 53, 8774-8778.	13.8	152
38	Dealumination mechanisms of zeolites and extra-framework aluminum confinement. <i>Journal of Catalysis</i> , 2016, 339, 242-255.	6.2	149
39	Quantum Chemical Modeling of Benzene Ethylation over H-ZSM-5 Approaching Chemical Accuracy: A Hybrid MP2:DFT Study. <i>Journal of the American Chemical Society</i> , 2010, 132, 11525-11538.	13.7	144
40	Finding transition structures in extended systems: A strategy based on a combined quantum mechanics-empirical valence bond approach. <i>Journal of Chemical Physics</i> , 2000, 112, 6983-6996.	3.0	142
41	Resolving the Atomic Structure of Vanadia Monolayer Catalysts: Monomers, Trimers, and Oligomers on Ceria. <i>Angewandte Chemie - International Edition</i> , 2009, 48, 8006-8009.	13.8	138
42	Pyrazolate-Based Cobalt(II)-Containing Metal-Organic Frameworks in Heterogeneous Catalytic Oxidation Reactions: Elucidating the Role of Entatic States for Biomimetic Oxidation Processes. <i>Chemistry - A European Journal</i> , 2011, 17, 8671-8695.	3.3	138
43	Partial oxidation of ethanol on vanadia catalysts on supporting oxides with different redox properties compared to propane. <i>Journal of Catalysis</i> , 2012, 296, 120-131.	6.2	138
44	Potential Functions for Silica and Zeolite Catalysts Based on ab Initio Calculations. 3. A Shell Model Ion Pair Potential for Silica and Aluminosilicates. <i>The Journal of Physical Chemistry</i> , 1996, 100, 11043-11049.	2.9	135
45	(V2O5) <sub>n</sub> Gas-Phase Clusters (n = 1-12) Compared to V2O5 Crystal: DFT Calculations. <i>Journal of Physical Chemistry A</i> , 2001, 105, 8588-8598.	2.5	135
46	Siting of Al and bridging hydroxyl groups in ZSM-5: A computer simulation study. <i>Zeolites</i> , 1992, 12, 20-23.	0.5	132
47	Proton Mobility in Chabazite, Faujasite, and ZSM-5 Zeolite Catalysts. Comparison Based on ab Initio Calculations. <i>Journal of Physical Chemistry B</i> , 2001, 105, 1603-1613.	2.6	132
48	Models in Catalysis. <i>Catalysis Letters</i> , 2015, 145, 109-125.	2.6	130
49	A High-Resolution <sup>17</sup> O and <sup>29</sup> Si NMR Study of Zeolite Siliceous Ferrierite and ab Initio Calculations of NMR Parameters. <i>Journal of the American Chemical Society</i> , 2000, 122, 4948-4958.	13.7	129
50	Size-Dependent Catalytic Activity of Supported Vanadium Oxide Species: Oxidative Dehydrogenation of Propane. <i>Journal of the American Chemical Society</i> , 2014, 136, 7751-7761.	13.7	126
51	A hybrid MP2/planewave-DFT scheme for large chemical systems: proton jumps in zeolites. <i>Chemical Physics Letters</i> , 2004, 387, 388-394.	2.6	125
52	Proton NMR chemical shift and intrinsic acidity of hydroxyl groups. Ab initio calculations on catalytically active sites and gas-phase molecules. <i>Journal of the American Chemical Society</i> , 1993, 115, 7833-7838.	13.7	124
53	Coordination, Structure, and Vibrational Spectra of Titanium in Silicates and Zeolites in Comparison with Related Molecules. An ab Initio Study. <i>The Journal of Physical Chemistry</i> , 1996, 100, 5025-5034.	2.9	123
54	Accurate quantum chemical energies for the interaction of hydrocarbons with oxide surfaces: CH <sub>4</sub> /MgO(001). <i>Physical Chemistry Chemical Physics</i> , 2010, 12, 14330.	2.8	122

#	ARTICLE	IF	CITATIONS
55	Heats of Adsorption of CO and CO <sub>2</sub> in Metal-Organic Frameworks: Quantum Mechanical Study of CPO-27-M (M = Mg, Ni, Zn). <i>Journal of Physical Chemistry C</i> , 2011, 115, 21777-21784.	3.1	122
56	Effect of Al-Al and Al-Si-Al Pairs in the ZSM-5 Zeolite Framework on the <sup>27</sup> Al NMR Spectra. A Combined High-Resolution <sup>27</sup> Al NMR and DFT/MM Study. <i>Journal of Physical Chemistry C</i> , 2009, 113, 1447-1458.	3.1	121
57	Computational Elucidation of the Transition State Shape Selectivity Phenomenon. <i>Journal of the American Chemical Society</i> , 2004, 126, 936-947.	13.7	120
58	Accurate Adsorption Thermodynamics of Small Alkanes in Zeolites. Ab initio Theory and Experiment for H-Chabazite. <i>Journal of Physical Chemistry C</i> , 2015, 119, 6128-6137.	3.1	120
59	Toward an Understanding of Selective Alkyne Hydrogenation on Ceria: On the Impact of O Vacancies on H <sub>2</sub> Interaction with CeO <sub>2</sub> (111). <i>Journal of the American Chemical Society</i> , 2017, 139, 17608-17616.	13.7	120
60	Structure and reactivity of V <sub>2</sub> O <sub>5</sub> : bulk solid, nanosized clusters, species supported on silica and alumina, cluster cations and anions. <i>Dalton Transactions</i> , 2004, , 3116-3121.	3.3	119
61	The infrared spectrum of the O-H-O fragment of H[ <sub>5</sub> O] <sub>2</sub> [ <sup>+</sup> ]: Ab initio classical molecular dynamics and quantum 4D model calculations. <i>Journal of Chemical Physics</i> , 2001, 114, 240.	3.0	118
62	Polyhedral Vanadium Oxide Cages: Infrared Spectra of Cluster Anions and Size-Induced d Electron Localization. <i>Angewandte Chemie - International Edition</i> , 2005, 44, 3122-3125.	13.8	116
63	Surface Metal-Insulator Transition on a Vanadium Pentoxide (001) Single Crystal. <i>Physical Review Letters</i> , 2007, 99, 226103.	7.8	113
64	Unexpected Structures of Aluminum Oxide Clusters in the Gas Phase. <i>Angewandte Chemie - International Edition</i> , 2007, 46, 3372-3375.	13.8	113
65	Acidic sites in heterogeneous catalysis: structure, properties and activity. <i>Journal of Molecular Catalysis</i> , 1989, 54, 312-323.	1.2	112
66	Gas-Phase Vanadium Oxide Anions: Structure and Detachment Energies from Density Functional Calculations. <i>Journal of Physical Chemistry A</i> , 2000, 104, 10913-10922.	2.5	110
67	Gas phase infrared spectroscopy of mono- and divanadium oxide cluster cations. <i>Journal of Chemical Physics</i> , 2004, 120, 6461-6470.	3.0	110
68	Gas-Phase Oxidation of Propane and 1-Butene with [V <sub>3</sub> O <sub>7</sub> ] <sup>+</sup> : Experiment and Theory in Concert. <i>Angewandte Chemie - International Edition</i> , 2006, 45, 4677-4681.	13.8	110
69	Coordination of Cu+Ions to Zeolite Frameworks Strongly Enhances Their Ability To Bind NO <sub>2</sub> . An ab Initio Density Functional Study. <i>Journal of the American Chemical Society</i> , 1998, 120, 1545-1551.	13.7	109
70	Protonated Isobutene in Zeolites:tert-Butyl Cation or Alkoxide?. <i>Angewandte Chemie - International Edition</i> , 2005, 44, 4769-4771.	13.8	109
71	Probing the Electronic Structure of Early Transition-Metal Oxide Clusters: Polyhedral Cages of (V <sub>2</sub> O <sub>5</sub> ) <sub>n</sub> ( <i>n</i> = 2-4) and (Nb <sub>2</sub> O <sub>5</sub> ) <sub>n</sub> (Nb = Nb, Ta). <i>Journal of the American Chemical Society</i> , 2007, 129, 13270-13276.	13.7	109
72	Molecular structure of orthosilicic acid, silanol, and H <sub>3</sub> SiOH.cndot.AlH <sub>3</sub> complex: models of surface hydroxyls in silica and zeolites. <i>The Journal of Physical Chemistry</i> , 1987, 91, 2315-2319.	2.9	108

#	ARTICLE		IF	CITATIONS
73	1H NMR Chemical Shifts of Ammonia, Methanol, and Water Molecules Interacting with Brønsted Acid Sites of Zeolite Catalysts: Ab-Initio Calculations. <i>The Journal of Physical Chemistry</i> , 1994, 98, 3083-3085.	2.9	108	
74	Thin silica films on Ru(0001): monolayer, bilayer and three-dimensional networks of [SiO <sub>4</sub> ] tetrahedra. <i>Physical Chemistry Chemical Physics</i> , 2012, 14, 11344.	2.8	106	
75	O <sub>2</sub> Activation on Ceria Catalysts—The Importance of Substrate Crystallographic Orientation. <i>Angewandte Chemie - International Edition</i> , 2017, 56, 16399-16404.	13.8	106	
76	Heterogeneity of Brønsted Acidic Sites in Faujasite Type Zeolites due to Aluminum Content and Framework Structure. <i>Journal of Physical Chemistry B</i> , 1998, 102, 6397-6404.	2.6	105	
77	Coordination of Cu <sup>+</sup> and Cu <sup>2+</sup> ions in ZSM-5 in the vicinity of two framework Al atoms. <i>Physical Chemistry Chemical Physics</i> , 2001, 3, 1552-1559.	2.8	104	
78	Concentration of Vacancies at Metal-Oxide Surfaces: Case Study of MgO(100). <i>Physical Review Letters</i> , 2013, 111, 045502.	7.8	104	
79	Support Effect in Oxide Catalysis: Methanol Oxidation on Vanadia/Ceria. <i>Journal of the American Chemical Society</i> , 2014, 136, 14616-14625.	13.7	101	
80	Effect of Anharmonicity on Adsorption Thermodynamics. <i>Journal of Chemical Theory and Computation</i> , 2014, 10, 2479-2487.	5.3	101	
81	Ab initio molecular dynamics simulation of methanol adsorbed in chabazite. <i>Chemical Physics Letters</i> , 1997, 266, 397-402.	2.6	100	
82	Structure determination of neutral MgO clusters—hexagonal nanotubes and cages. <i>Physical Chemistry Chemical Physics</i> , 2012, 14, 2849.	2.8	100	
83	Ab initio Calculation of Rate Constants for Molecule-Surface Reactions with Chemical Accuracy. <i>Angewandte Chemie - International Edition</i> , 2016, 55, 5235-5237.	13.8	100	
84	Water in Interaction with Acid Sites in H-ZSM-5 Zeolite Does Not Form Hydroxonium Ions. A Comparison between Neutron Scattering Results and ab Initio Calculations. <i>The Journal of Physical Chemistry</i> , 1996, 100, 19545-19550.	2.9	99	
85	A High-Resolution <sup>17</sup> O NMR Study of Siliceous Zeolite Faujasite. <i>Journal of the American Chemical Society</i> , 1998, 120, 3510-3511.	13.7	97	
86	The <i>i</i> -tert-Butyl Cation in H-Zeolites: Deprotonation to Isobutene and Conversion into Surface Alkoxides. <i>Angewandte Chemie - International Edition</i> , 2010, 49, 4678-4680.	13.8	97	
87	Adsorption of NH <sub>3</sub> and H <sub>2</sub> O in Acidic Chabazite. Comparison of ONIOM Approach with Periodic Calculations. <i>Journal of Physical Chemistry B</i> , 2005, 109, 3539-3545.	2.6	96	
88	Modeling Zeolites with Metal-supported Two-dimensional Aluminosilicate Films. <i>Angewandte Chemie - International Edition</i> , 2012, 51, 6005-6008.	13.8	96	
89	Stable Mechanistically-Relevant Aromatic-Based Carbenium Ions in Zeolite Catalysts. <i>Journal of the American Chemical Society</i> , 2003, 125, 2136-2141.	13.7	95	
90	Stabilities of C <sub>3</sub> –C <sub>5</sub> alkoxide species inside H-FER zeolite: a hybrid QM/MM study. <i>Journal of Catalysis</i> , 2005, 231, 393-404.	6.2	91	

#	ARTICLE	IF	CITATIONS
91	Water adsorption and O-defect formation on Fe <sub>2</sub> O <sub>3</sub> (0001) surfaces. <i>Physical Chemistry Chemical Physics</i> , 2016, 18, 25560-25568.	2.8	91
92	Ab Initio Prediction of Adsorption Isotherms for Small Molecules in Metal-Organic Frameworks: The Effect of Lateral Interactions for Methane/CPO-27-Mg. <i>Journal of the American Chemical Society</i> , 2012, 134, 18354-18365.	13.7	90
93	Oxidative dehydrogenation of propane: Differences between N <sub>2</sub> O and O <sub>2</sub> in the reoxidation of reduced vanadia sites and consequences for selectivity. <i>Journal of Catalysis</i> , 2008, 256, 84-94.	6.2	88
94	Point defects in CaF <sub>2</sub> and CeO <sub>2</sub> investigated by the periodic electrostatic embedded cluster method. <i>Journal of Chemical Physics</i> , 2009, 130, 174710.	3.0	88
95	Structures of the Ordered Water Monolayer on MgO(001). <i>Journal of Physical Chemistry C</i> , 2011, 115, 6764-6774.	3.1	88
96	<sup>29</sup> Si NMR Chemical Shifts of Silicate Species: A Ab Initio Study of Environment and Structure Effects. <i>Journal of the American Chemical Society</i> , 1996, 118, 13015-13020.	13.7	87
97	Understanding the Nature of Water Bound to Solid Acid Surfaces. Ab Initio Simulation on HSAPO-34. <i>Journal of the American Chemical Society</i> , 1998, 120, 8512-8516.	13.7	87
98	Coordination Change of Cu+ Sites in ZSM-5 on Excitation in the Triplet State: Understanding of the Photoluminescence Spectra. <i>Journal of Physical Chemistry B</i> , 2000, 104, 1738-1745.	2.6	87
99	Nature of the Cu+-NO Bond in the Gas Phase and at Different Types of Cu+ Sites in Zeolite Catalysts. <i>Journal of Physical Chemistry B</i> , 2004, 108, 13674-13682.	2.6	86
100	Oxidative Activation of Butane on Sulfated Zirconia. <i>Journal of the American Chemical Society</i> , 2005, 127, 16159-16166.	13.7	86
101	The effect of hydration on structure and location of Ti-sites in Ti-silicalite catalysts. A computational study. <i>Physical Chemistry Chemical Physics</i> , 2000, 2, 2195-2204.	2.8	85
102	Surface Structure of Hydroxylated and Sulfated Zirconia. A Periodic Density-Functional Study. <i>Physical Chemistry Chemical Physics</i> , 2004, 108, 14652-14662.	2.6	82
103	Gas phase vibrational spectroscopy of mass-selected vanadium oxide anions. <i>Physical Chemistry Chemical Physics</i> , 2008, 10, 3992.	2.8	81
104	Vanadium oxide surfaces and supported vanadium oxide nanoparticles. <i>Topics in Catalysis</i> , 2006, 38, 117-125.	2.8	80
105	Ab initio molecular dynamics simulation of methanol interacting with acidic zeolites of different framework structure. <i>Microporous and Mesoporous Materials</i> , 2000, 35-36, 379-385.	4.4	77
106	Periodic density functional study on structural and vibrational properties of vanadium oxide aggregates. <i>Physical Review B</i> , 2004, 69, .	3.2	76
107	Adsorption, Activation, and Dissociation of Oxygen on Doped Oxides. <i>Angewandte Chemie - International Edition</i> , 2013, 52, 11385-11387.	13.8	76
108	Degradation of Ionized OV(OCH <sub>3</sub> ) <sub>3</sub> in the Gas Phase. From the Neutral Compound All the Way down to the Quasi-terminal Fragments VO <sup>+</sup> and VOH <sup>+</sup> . <i>Inorganic Chemistry</i> , 2006, 45, 6235-6245.	4.0	75

#	ARTICLE	IF	CITATIONS
109	Regioselectivity of Al-O Bond Hydrolysis during Zeolites Dealumination Unified by Brønsted-Evans-Polanyi Relationship. <i>ACS Catalysis</i> , 2015, 5, 11-15.	11.2	73
110	Bonding ability of surface sites on silica and their effect on hydrogen bonds. A quantum-chemical and statistical thermodynamic treatment. <i>The Journal of Physical Chemistry</i> , 1981, 85, 4061-4067.	2.9	72
111	A comparison between plane wave and Gaussian-type orbital basis sets for hydrogen bonded systems: Formic acid as a test case. <i>Journal of Chemical Physics</i> , 2007, 127, 154102.	3.0	72
112	Minimal basis set MINI-1 ? powerful tool for calculating of molecular interactions. I. Neutral complexes. <i>Theoretica Chimica Acta</i> , 1984, 65, 279-290.	0.8	71
113	Translational proton motion in zeolite H-ZSM-5. Energy barriers and jump rates from DFT calculations. <i>Physical Chemistry Chemical Physics</i> , 2002, 4, 5207-5216.	2.8	71
114	Formaldehyde Formation on Vanadium Oxide Surfaces V <sub>2</sub> O <sub>3</sub> (0001) and V <sub>2</sub> O <sub>5</sub> (001): How does the Stable Methoxy Intermediate Form?. <i>Angewandte Chemie - International Edition</i> , 2009, 48, 3695-3698.	13.8	70
115	Ab Initio Calculations for Molecule-Surface Interactions with Chemical Accuracy. <i>Accounts of Chemical Research</i> , 2019, 52, 3502-3510.	15.6	70
116	The acidity of surface silanol groups. A theoretical estimate based on ab initio calculations on a model surface. <i>Chemical Physics Letters</i> , 1994, 218, 333-337.	2.6	69
117	Formation of hydronium ions on Brønsted sites in zeolitic catalysts: a quantum-chemical ab initio study. <i>Chemical Physics Letters</i> , 1990, 173, 26-32.	2.6	68
118	Gas-Phase Infrared Spectrum of the Protonated Water Dimer: Molecular Dynamics Simulation and Accuracy of the Potential Energy Surface. <i>ChemPhysChem</i> , 2005, 6, 1706-1710.	2.1	68
119	Gas phase acidities and molecular geometries of H <sub>3</sub> SiOH, H <sub>3</sub> COH, and H <sub>2</sub> O. <i>Journal of Chemical Physics</i> , 1990, 93, 2575-2583.	3.0	67
120	Structural Diversity and Flexibility of MgO Gas-Phase Clusters. <i>Angewandte Chemie - International Edition</i> , 2011, 50, 1716-1719.	13.8	67
121	Interaction of Probe Molecules with Bridging Hydroxyls of Two-Dimensional Zeolites: A Surface Science Approach. <i>Journal of Physical Chemistry C</i> , 2013, 117, 13547-13556.	3.1	67
122	Comparison of a combined quantum mechanics/interatomic potential function approach with its periodic quantum-mechanical limit: Proton siting and ammonia adsorption in zeolite chabazite. <i>Journal of Chemical Physics</i> , 1998, 109, 10379-10389.	3.0	66
123	Brønsted sites in zeolitic catalysts. An ab initio study of local geometries and of the barrier for proton jumps between neighbouring sites. <i>Chemical Physics Letters</i> , 1989, 164, 193-198.	2.6	64
124	Protonation of water clusters in the cavities of acidic zeolites: (H <sub>2</sub> O) <sub>n</sub> ·H-chabazite, n = 1-4. <i>Physical Chemistry Chemical Physics</i> , 2009, 11, 1702.	2.8	64
125	Accurate adsorption energies of small molecules on oxide surfaces: CO-MgO(001). <i>Physical Chemistry Chemical Physics</i> , 2013, 15, 16481.	2.8	64
126	The vibrational frequency of the donor OH group in the H-bonded dimers of water, methanol and silanol. Ab initio calculations including anharmonicities. <i>Chemical Physics Letters</i> , 1995, 238, 243-252.	2.6	63

#	ARTICLE	IF	CITATIONS
127	Quantum chemical studies on zeolites and silica. International Journal of Quantum Chemistry, 1984, 26, 793-822.	2.0	62
128	Water Interaction with Iron Oxides. Angewandte Chemie - International Edition, 2015, 54, 13942-13946.	13.8	62
129	Ab Initio Prediction of Adsorption Isotherms for Small Molecules in Metal-Organic Frameworks. Journal of the American Chemical Society, 2016, 138, 14047-14056.	13.7	62
130	Quantum chemical investigation of interaction sites in zeolites and silica. The Journal of Physical Chemistry, 1980, 84, 3318-3326.	2.9	61
131	Formation of the cerium orthovanadate CeVO <sub>4</sub> : DFT+U study. Physical Review B, 2007, 76, .	3.2	61
132	Oxidative conversion of C <sub>1</sub> -C <sub>3</sub> alkanes by vanadium oxide catalysts. DFT results and their accuracy. International Journal of Quantum Chemistry, 2008, 108, 2223-2229.	2.0	61
133	Periodic Density Functional Theory Study of VO <sub>n</sub> Species Supported on the CeO <sub>2</sub> (111) Surface. Journal of Physical Chemistry C, 2011, 115, 7399-7410.	3.1	61
134	Oligomeric Vanadium Oxide Species Supported on the CeO <sub>2</sub> (111) Surface: Structure and Reactivity Studied by Density Functional Theory. Journal of Physical Chemistry C, 2013, 117, 5274-5285.	3.1	60
135	Preferred stability of aluminum-oxygen-silicon-oxygen-aluminum linkages in high-silica zeolite catalysts: theoretical predictions contrary to Dempsey's rule. The Journal of Physical Chemistry, 1993, 97, 6579-6581.	2.9	59
136	Cooperative Formation of Long-Range Ordering in Water Adlayers on Fe <sub>3</sub> O <sub>4</sub> (111) Surfaces. Angewandte Chemie - International Edition, 2018, 57, 1409-1413.	13.8	59
137	Combining ab initio techniques with analytical potential functions. A study of zeolite-adsorbate interactions for NH <sub>3</sub> on H-faujasite. Journal of Molecular Catalysis A, 1997, 119, 19-33.	4.8	57
138	Isomorphous Substitution in Bimetallic Oxide Clusters. Physical Review Letters, 2006, 96, 233401.	7.8	57
139	Thickness-Dependent Hydroxylation of MgO(001) Thin Films. Journal of Physical Chemistry C, 2010, 114, 18207-18214.	3.1	57
140	The minimal basis set MINI-1?powerful tool for calculating intermolecular interactions. II. Ionic complexes. Theoretica Chimica Acta, 1984, 65, 291-302.	0.8	56
141	Low temperature adsorption of oxygen on reduced V <sub>2</sub> O <sub>3</sub> (0001) surfaces. Surface Science, 2006, 600, 1497-1503.	1.9	55
142	Physisorption and Chemisorption of Hydrocarbons in H-FAU Using QM-Pot(MP2//B3LYP) Calculations. Journal of Physical Chemistry C, 2008, 112, 11796-11812.	3.1	55
143	Oxidative Dehydrogenation of Hydrocarbons by V <sub>3</sub> O <sub>7</sub> + Compared to Other Vanadium Oxide Species. Journal of Physical Chemistry A, 2009, 113, 11586-11594.	2.5	55
144	Structural variability in transition metal oxide clusters: gas phase vibrational spectroscopy of V <sub>3</sub> O <sub>6</sub> 8+. Physical Chemistry Chemical Physics, 2012, 14, 9377.	2.8	55

#	ARTICLE of $\text{Ce}^{\text{3+}}$ in the $\text{CeO}_2$ lattice. <i>Topics in Catalysis</i> , 2008, 50, 106-115.	IF	CITATIONS
145	$\text{Ce}^{\text{3+}}$ in the $\text{CeO}_2$ lattice. <i>Topics in Catalysis</i> , 2008, 50, 106-115.	7.8	55
146	Selectivity in Methanol Oxidation as Studied on Model Systems Involving Vanadium Oxides. <i>Topics in Catalysis</i> , 2008, 50, 106-115.	2.8	53
147	Quantum Chemical Free Energies: Structure Optimization and Vibrational Frequencies in Normal Modes. <i>Journal of Chemical Theory and Computation</i> , 2013, 9, 5038-5045.	5.3	53
148	Ab initio calculations of the structure and properties of disiloxane. The effect of electron correlation and basis set extension. <i>Chemical Physics Letters</i> , 1994, 226, 405-412.	2.6	51
149	Environmental effects on vibrational proton dynamics in $\text{H}_5\text{O}_2^+$ : DFT study on crystalline $\text{H}_5\text{O}_2^+\text{ClO}_4^-$ . <i>Physical Chemistry Chemical Physics</i> , 2005, 7, 258-263.	2.8	51
150	Identification of Conical Structures in Small Aluminum Oxide Clusters: Infrared Spectroscopy of $(\text{Al}_2\text{O}_3)_n$ . <i>Journal of the American Chemical Society</i> , 2008, 130, 15143-15149.	13.7	51
151	Reactions of $\text{H}_2$ , $\text{CH}_4$ , $\text{C}_2\text{H}_6$ , and $\text{C}_3\text{H}_8$ with $[(\text{MgO})_n]^{+}$ Clusters Studied by Density Functional Theory. <i>ChemCatChem</i> , 2010, 2, 819-826.	3.7	51
152	Ultrathin Silica Films: The Atomic Structure of Two-dimensional Crystals and Glasses. <i>Chemistry - A European Journal</i> , 2014, 20, 9176-9183.	3.3	51
153	Structures and vibrational spectroscopy of partially reduced gas-phase cerium oxide clusters. <i>Physical Chemistry Chemical Physics</i> , 2011, 13, 19393.	2.8	50
154	Ab Initio Prediction of Proton Exchange Barriers for Alkanes at Brønsted Sites of Zeolite H-MFI. <i>Journal of the American Chemical Society</i> , 2018, 140, 18151-18161.	13.7	50
155	Electron hole formation in acidic zeolite catalysts. <i>Journal of Chemical Physics</i> , 2004, 121, 6034-6041.	3.0	49
156	Ab initio predictions of zeolite structures and $^{29}\text{Si}$ NMR chemical shifts. <i>Solid State Nuclear Magnetic Resonance</i> , 1997, 9, 155-164.	2.3	48
157	On the geometrical and electronic structure of an ultra-thin crystalline silica film grown on Mo(112). <i>Surface Science</i> , 2007, 601, 4849-4861.	1.9	48
158	Structure and thermochemistry of $\text{Fe}_2\text{S}_2\text{O}_4$ /gas phase clusters and their fragments. B3LYP calculations. <i>Physical Chemistry Chemical Physics</i> , 2002, 4, 5234-5243.	2.8	47
159	Electronic ground states of the $\text{V}_2\text{O}_4\text{O}_4$ species from multireference correlation and density functional studies. <i>Journal of Chemical Physics</i> , 2004, 120, 4207-4215.	3.0	47
160	Introduction: Surface Chemistry of Oxides. <i>Chemical Reviews</i> , 2013, 113, 3859-3862.	47.7	47
161	Initial stages of $\text{CO}_2$ adsorption on CaO: a combined experimental and computational study. <i>Physical Chemistry Chemical Physics</i> , 2017, 19, 4231-4242.	2.8	47
162	Bond Dissociation Energies and Structures of $\text{CuNO}_2^+$ and $\text{Cu}(\text{NO})_2^+$ . <i>Inorganic Chemistry</i> , 2002, 41, 5882-5890.	4.0	46

#	ARTICLE	IF	CITATIONS
163	Surface Termination of $\text{Fe}_{3\text{O}_4}$ (111) Films Studied by CO Adsorption Revisited. Journal of Physical Chemistry B, 2018, 122, 527-533.	2.6	46
164	Confirmation of $\hat{\Sigma}$ and $\hat{\Xi}$ ground states of $\text{Fe}_2$ and $\text{Fe}_2\hat{\alpha}$ by CASSCF/MRCI. Chemical Physics Letters, 2002, 358, 442-448.	2.6	45
165	Synthesis and Structure of Ultrathin Aluminosilicate Films. Angewandte Chemie - International Edition, 2006, 45, 7636-7639.	13.8	45
166	Selective oxidation of propene by vanadium oxide monomers supported on silica. Journal of Catalysis, 2014, 317, 75-82.	6.2	45
167	Molecular and electronic structure of disiloxane, an ab initio MO study. Chemical Physics Letters, 1979, 65, 587-591.	2.6	44
168	The electronic states of $\text{Fe}_2\text{S}_2\hat{\alpha}/0+/2+$ . Journal of Chemical Physics, 2002, 116, 617-628.	3.0	44
169	Catalytic dehydrogenation of ethane over mononuclear Cr(III) surface sites on silica. part I. H activation by $\hat{\sigma}$ -bond metathesis. Journal of Physical Organic Chemistry, 2004, 17, 990-1006.	1.9	44
170	Partial oxidation of methanol on well-ordered V <sub>2</sub> O <sub>5</sub> (001)/Au(111) thin films. Physical Chemistry Chemical Physics, 2009, 11, 3290.	2.8	44
171	Hydrogen Spillover to Copper Clusters on Hydroxylated $\hat{\beta}\text{-Al}_2\text{O}_3$ . Journal of Physical Chemistry C, 2018, 122, 18445-18455.	3.1	44
172	Hydrogen adsorbed in a metal organic framework-5: Coupled translation-rotation eigenstates from quantum five-dimensional calculations. Journal of Chemical Physics, 2012, 137, 014701.	3.0	43
173	Comparing the Acidities of Microporous Aluminosilicate and Silico-Aluminophosphate Catalysts: A Combined Quantum Mechanics-Interatomic Potential Function Study. Collection of Czechoslovak Chemical Communications, 1998, 63, 1394-1408.	1.0	42
174	Vanadium Oxides on Aluminum Oxide Supports. 1. Surface Termination and Reducibility of Vanadia Films on $\hat{\pm}\text{-Al}_2\text{O}_3$ (0001). Journal of Physical Chemistry B, 2005, 109, 23523-23531.	2.6	42
175	Electron Distribution in Partially Reduced Mixed Metal Oxide Systems: Infrared Spectroscopy of $\text{Ce}_{m\text{V}_n}\text{O}_o$ Gas-Phase Clusters. Journal of Physical Chemistry A, 2011, 115, 11187-11192.	2.5	42
176	Interaction of $\text{C}_3\text{C}_5\text{Alkenes}$ with Zeolitic Brønsted Sites: $\ell$ -Complexes, Alkoxides, and Carbenium Ions in H-FER. Journal of Physical Chemistry C, 2020, 124, 10067-10078.	3.1	41
177	Structure and reactivity of zeolite catalysts: Atomistic modelling using ab initio techniques. Studies in Surface Science and Catalysis, 1994, , 2039-2057.	1.5	40
178	Trapping Aluminum Hydroxide Clusters with Trisilanols during Speciation in Aluminum(III)-Water Systems: Reproducible, Large Scale Access to Molecular Aluminate Models. Angewandte Chemie - International Edition, 2016, 55, 12325-12329.	13.8	40
179	Crystal Structure and Vibrational Spectra of $\text{AlVO}_4$ . A DFT Study. Journal of Physical Chemistry B, 2005, 109, 394-400.	2.6	39
180	Fast atom diffraction during grazing scattering from a MgO(001) surface. Surface Science, 2012, 606, 161-173.	1.9	39

#	ARTICLE	IF	CITATIONS
181	Preferential Activation of Primary C <sub>i</sub> H Bonds in the Reactions of Small Alkanes with the Diatomic MgO <sup>+</sup> Cation. <i>Chemistry - A European Journal</i> , 2010, 16, 4110-4119.	3.3	38
182	Oxidation of Methanol to Formaldehyde on Silica-Supported Molybdena: Density Functional Theory Study on Models of Mononuclear Sites. <i>Journal of Physical Chemistry C</i> , 2010, 114, 2967-2979.	3.1	38
183	Quantum chemical <i>ab initio</i> prediction of proton exchange barriers between CH <sub>4</sub> and different H-zeolites. <i>Journal of Chemical Physics</i> , 2015, 143, 102810.	3.0	38
184	Dissociative Water Adsorption by Al <sub>3</sub> O <sub>4</sub> <sup>+</sup> in the Gas Phase. <i>Journal of Physical Chemistry Letters</i> , 2017, 8, 1272-1277.	4.6	38
185	A CASSCF/ACPF study of spectroscopic properties of FeS and FeS <sup>+</sup> and the photoelectron spectrum of FeS <sup>+</sup> . <i>Chemical Physics Letters</i> , 1998, 294, 37-44.	2.6	37
186	Oxygen adsorption on Mo(112) surface studied by ab initio genetic algorithm and experiment. <i>Journal of Chemical Physics</i> , 2007, 126, 234710.	3.0	37
187	Theoretical studies of Cu(i) sites in faujasite and their interaction with carbon monoxide. <i>Physical Chemistry Chemical Physics</i> , 2007, 9, 5446.	2.8	37
188	Vibrational properties of surface hydroxyls: Nonempirical model calculations including anharmonicities. <i>Collection of Czechoslovak Chemical Communications</i> , 1988, 53, 2191-2202.	1.0	36
189	Vibrational spectra of the methanol tetramer in the OH stretch region. Two cyclic isomers and concerted proton tunneling. <i>Journal of Chemical Physics</i> , 2001, 114, 2623-2628.	3.0	36
190	Interaction of ethene, 2-methylpropene and benzene with the Na <sup>+</sup> ion. II. Quantum chemical study of sorption complexes in faujasites. <i>Zeolites</i> , 1982, 2, 114-120.	0.5	35
191	Molecular structure of orthosilicic acid and importance of (p-d) $\pi$ bonding. An ab initio molecular orbital study. <i>Chemical Physics Letters</i> , 1983, 97, 275-278.	2.6	35
192	Methane activation by silica-supported Zr(iv) hydrides: the dihydride [(SiO) <sub>2</sub> ZrH <sub>2</sub> ] is much faster than the monohydride [(SiO) <sub>3</sub> ZrH]. <i>Chemical Communications</i> , 2004, , 1729-1731.	4.1	35
193	Atomic Structure of an Ultrathin Fe-Silicate Film Grown on a Metal: A Monolayer of Clay?. <i>Journal of the American Chemical Society</i> , 2013, 135, 19222-19228.	13.7	35
194	Relative propensity of methanol and silanol towards hydrogen bond formation. <i>Chemical Physics Letters</i> , 1992, 191, 537-547.	2.6	34
195	Quantum anharmonic frequencies of the O <sup>-</sup> H <sup>-</sup> O fragment of the H <sub>5</sub> O <sub>2</sub> <sup>+</sup> ion: a model three-dimensional study. <i>Chemical Physics Letters</i> , 1999, 312, 591-597.	2.6	34
196	Vibrations of Silica Supported Vanadia: Variation with Particle Size and Local Surface Structure. <i>Journal of Physical Chemistry C</i> , 2009, 113, 12454-12464.	3.1	34
197	Water on the MgO(001) Surface: Surface Reconstruction and Ion Solvation. <i>Journal of Physical Chemistry Letters</i> , 2015, 6, 2310-2314.	4.6	34
198	Methanol adsorption on monocrystalline ceria surfaces. <i>Journal of Catalysis</i> , 2016, 336, 116-125.	6.2	34

#	ARTICLE	IF	CITATIONS
199	On the Existence of CuI Pairs in ZSM-5—A Computational Study. <i>Chemistry - A European Journal</i> , 2002, 8, 2099.	3.3	33
200	Electron stimulated hydroxylation of a metal supported silicate film. <i>Physical Chemistry Chemical Physics</i> , 2016, 18, 3755-3764.	2.8	33
201	Interactions of Water with the (111) and (100) Surfaces of Ceria. <i>Journal of Physical Chemistry C</i> , 2017, 121, 21571-21578.	3.1	33
202	Ab Initio Adsorption Isotherms for Molecules with Lateral Interactions: CO <sub>2</sub> in Metal-Organic Frameworks. <i>Journal of Physical Chemistry C</i> , 2017, 121, 12789-12799.	3.1	33
203	Probing Catalysts with Water. <i>Science</i> , 1996, 271, 774-775.	12.6	32
204	Acidity of two-dimensional zeolites. <i>Physical Chemistry Chemical Physics</i> , 2015, 17, 27873-27882.	2.8	32
205	Transferable analytical potential based on nonempirical quantum chemical calculations (QOPEN) for water-silica interactions. <i>The Journal of Physical Chemistry</i> , 1984, 88, 6375-6383.	2.9	31
206	Ion Chemistry of OV(OCH <sub>3</sub> ) <sub>3</sub> in the Gas Phase: Molecular Cations and Anions and Their Primary Fragmentations. <i>Inorganic Chemistry</i> , 2004, 43, 1976-1985.	4.0	31
207	Combined Quantum Mechanics: A Interatomic Potential Function Investigation of rac-meso Configurational Stability and Rotational Transition in Zirconocene-Based Ziegler-Natta Catalysts. <i>Journal of Physical Chemistry A</i> , 2000, 104, 10932-10938.	2.5	30
208	Adsorption and Diffusion of Fructose in Zeolite HZSM-5: Selection of Models and Methods for Computational Studies. <i>Journal of Physical Chemistry C</i> , 2011, 115, 21785-21790.	3.1	30
209	Surface Structure of V <sub>2</sub> O <sub>3</sub> (0001) Revisited. <i>Physical Review Letters</i> , 2015, 114, 216101.	7.8	30
210	Brønsted activity of two-dimensional zeolites compared to bulk materials. <i>Faraday Discussions</i> , 2016, 188, 227-234.	3.2	30
211	Including dispersion in density functional theory for adsorption on flat oxide surfaces, in metal-organic frameworks and in acidic zeolites. <i>Physical Chemistry Chemical Physics</i> , 2020, 22, 7577-7585.	2.8	30
212	Formic acid tetramers: structure isomers in the gas phase. <i>Chemical Physics Letters</i> , 1997, 267, 111-115.	2.6	29
213	Infrared spectroscopic characterization of the oxidative dehydrogenation of propane by V <sub>4</sub> O <sub>10</sub> <sup>+</sup> . <i>International Journal of Mass Spectrometry</i> , 2010, 297, 102-106.	1.5	29
214	Thermal Methane Activation by a Binary Nb Transition-Metal Oxide Cluster Cation: A Further Example for the Crucial Role of Oxygen-Centered Radicals. <i>Chemistry - A European Journal</i> , 2013, 19, 11496-11501.	3.3	29
215	CO adsorption on a silica bilayer supported on Ru(0001). <i>Surface Science</i> , 2016, 648, 2-9.	1.9	29
216	<i>&lt; i&gt;Ab initio&lt;/i&gt;</i> study of methanol and ethanol adsorption on Brønsted sites in zeolite H-MFI. <i>Physical Chemistry Chemical Physics</i> , 2018, 20, 19964-19970.	2.8	29

#	ARTICLE	IF	CITATIONS
217	Adsorption and cracking of propane by zeolites of different pore size. <i>Journal of Catalysis</i> , 2021, 395, 117-128.	6.2	29
218	Ab initio molecular dynamics simulation of H <sub>5</sub> O+2 and H <sub>7</sub> O+3 gas phase clusters based on density functional theory. <i>Molecular Physics</i> , 1997, 91, 963-975.	1.7	29
219	Structural Analysis of Silica-Supported Molybdena Based on X-ray Spectroscopy: Quantum Theory and Experiment. <i>Journal of Physical Chemistry C</i> , 2011, 115, 15449-15458.	3.1	28
220	Kinetic study of the reaction of vanadium and vanadium-titanium oxide cluster anions with SO <sub>2</sub> . <i>Physical Chemistry Chemical Physics</i> , 2012, 14, 14344.	2.8	28
221	Oxidative dehydrogenation of methanol at ceria-supported vanadia oligomers. <i>Journal of Catalysis</i> , 2017, 352, 382-387.	6.2	28
222	Nonempirical direct SCF calculations on sodalite and double six-ring models of SiO <sub>2</sub> and AlPO <sub>4</sub> minerals: H <sub>2</sub> 4Si24O60, H <sub>12</sub> Si12O30, H <sub>12</sub> Al6P6O30. <i>Chemical Physics Letters</i> , 1989, 164, 199-204.	2.6	27
223	Absolute acidities and site specific properties of zeolite catalysts modelled by advanced computational chemistry technology. <i>Chemical Physics Letters</i> , 1999, 308, 147-154.	2.6	27
224	Acidic Catalysis by Zeolites: Ab Initio Modeling of Transition Structures. <i>ACS Symposium Series</i> , 1999, , 358-367.	0.5	27
225	Vanadium Oxides on Aluminum Oxide Supports. 2. Structure, Vibrational Properties, and Reducibility of V <sub>2</sub> O <sub>5</sub> Clusters on $\hat{\gamma}$ -Al <sub>2</sub> O <sub>3</sub> (0001). <i>Journal of Physical Chemistry B</i> , 2005, 109, 23532-23542.	2.6	27
226	Interplay between theory and experiment in the quest for silica with reduced dimensionality grown on a Mo(112) surface. <i>Chemical Physics Letters</i> , 2006, 424, 115-119.	2.6	27
227	Vanadium Oxides on Aluminum Oxide Supports. 3. Metastable $\hat{\gamma}$ -Al <sub>2</sub> O <sub>3</sub> (001) Compared to $\hat{\gamma}$ -Al <sub>2</sub> O <sub>3</sub> (0001). <i>Journal of Physical Chemistry C</i> , 2007, 111, 5141-5153.	3.1	27
228	Gas-phase Infrared Photodissociation Spectroscopy of Tetravanadiumoxo and Oxo-Methoxo Cluster Anions. <i>ChemPhysChem</i> , 2007, 8, 1640-1647.	2.1	27
229	The [(Al <sub>2</sub> O <sub>3</sub> ) <sub>2</sub> ] <sup>2+</sup> Anion Cluster: Electron Localization-Delocalization Isomerism. <i>ChemPhysChem</i> , 2009, 10, 2410-2413.	2.1	27
230	Applicability of the supermolecule MP2 approach to intermolecular interactions: He <sub>2</sub> and Ne <sub>2</sub> . <i>Chemical Physics Letters</i> , 1987, 134, 553-559.	2.6	26
231	Internal silanols in zeolites - inferences from quantum chemical calculations. <i>Catalysis Today</i> , 1988, 3, 485-492.	4.4	26
232	V <sub>2</sub> O <sub>5</sub> /SiO <sub>2</sub> surface inspired, silsesquioxane-derived oxovanadium complexes and their properties. <i>Dalton Transactions</i> , 2008, , 326-331.	3.3	26
233	Aluminum Siting in the ZSM-22 and Theta-1 Zeolites Revisited: A QM/MM Study. <i>Collection of Czechoslovak Chemical Communications</i> , 2008, 73, 909-920.	1.0	26
234	Chemically accurate adsorption energies for methane and ethane monolayers on the MgO(001) surface. <i>Physical Chemistry Chemical Physics</i> , 2018, 20, 9760-9769.	2.8	26

#	ARTICLE	IF	CITATIONS
235	Water adsorption on the Fe <sub>3</sub> O <sub>4</sub> (111) surface: dissociation and network formation. <i>Physical Chemistry Chemical Physics</i> , 2018, 20, 15764-15774.	2.8	26
236	Interaction of Water Molecules with the $\hat{t}$ -Fe <sub>2</sub> O <sub>3</sub> (0001) Surface: A Combined Experimental and Computational Study. <i>Journal of Physical Chemistry C</i> , 2019, 123, 8324-8335.	3.1	26
237	Activation and isomerization of n-butane on sulfated zirconia model systems—“an integrated study across the materials and pressure gaps. <i>Physical Chemistry Chemical Physics</i> , 2007, 9, 3600-3618.	2.8	25
238	Stability and migration barriers of small vanadium oxide clusters on the CeO <sub>2</sub> (111) surface studied by density functional theory. <i>Faraday Discussions</i> , 2013, 162, 233.	3.2	25
239	Hydrogen Atom or Proton Coupled Electron Transfer? H Bond Activation by Transition-Metal Oxides. <i>Journal of the American Chemical Society</i> , 2019, 141, 14603-14611.	13.7	25
240	Formation of one-dimensional molybdenum oxide on Mo(112). <i>Surface Science</i> , 2008, 602, 3338-3342.	1.9	23
241	Chemically Accurate Adsorption Energies: CO and H <sub>2</sub> O on the MgO(001) Surface. <i>Journal of Chemical Theory and Computation</i> , 2019, 15, 1329-1344.	5.3	23
242	THE ELECTRONIC STRUCTURE AND PHYSICAL PROPERTIES OF THIONITROSO COMPOUNDS—“A QUANTUM CHEMICAL STUDY. <i>Phosphorous and Sulfur and the Related Elements</i> , 1981, 11, 325-334.	0.2	22
243	Insights into Reaction Kinetics in Confined Space: Real Time Observation of Water Formation under a Silica Cover. <i>Journal of the American Chemical Society</i> , 2021, 143, 8780-8790.	13.7	22
244	Starting from first principles. <i>Nature</i> , 1993, 363, 493-494.	27.8	21
245	Hydration Structures of MgO, CaO, and SrO (001) Surfaces. <i>Journal of Physical Chemistry C</i> , 2016, 120, 24762-24769.	3.1	21
246	Ab Initio Prediction of Adsorption Isotherms for Gas Mixtures by Grand Canonical Monte Carlo Simulations on a Lattice of Sites. <i>Journal of Physical Chemistry Letters</i> , 2017, 8, 2713-2718.	4.6	21
247	O <sub>2</sub> Aktivierung an Cerdioxid-Katalysatoren — Zur Bedeutung der kristallographischen Orientierung des Substrats. <i>Angewandte Chemie</i> , 2017, 129, 16618-16623.	2.0	21
248	Inhibition of Interleukin-2-Mediated Lymphocyte Activation in Patients with Cushing’s Syndrome: A Comparison with Hypocortisolemic Patients. <i>Neuroendocrinology</i> , 1994, 59, 144-151.	2.5	20
249	Vanadia Aggregates on an Ultrathin Aluminum Oxide Film on NiAl(110). <i>Journal of Physical Chemistry C</i> , 2010, 114, 4983-4994.	3.1	20
250	The vibrational spectrum of FeO <sub>2+</sub> isomers—“Theoretical benchmark and experiment. <i>Journal of Chemical Physics</i> , 2014, 140, 204315.	3.0	20
251	Catalytic Performance of Vanadium MIL-47 and Linker-Substituted Variants in the Oxidation of Cyclohexene: A Combined Theoretical and Experimental Approach. <i>ChemPlusChem</i> , 2014, 79, 1183-1197.	2.8	20
252	Accurate adsorption energies for small molecules on oxide surfaces: CH <sub>4</sub> /MgO(001) and C <sub>2</sub> H <sub>6</sub> /MgO(001). <i>Journal of Computational Chemistry</i> , 2016, 37, 2374-2385.	3.3	20

#	ARTICLE	IF	CITATIONS
253	Trapping Aluminum Hydroxide Clusters with Trisilanols during Speciation in Aluminum(III)-Water Systems: Reproducible, Large Scale Access to Molecular Aluminate Models. <i>Angewandte Chemie</i> , 2016, 128, 12513-12517.	2.0	20
254	Reduction of the (001) Surface of $\beta$ -V2O5 Compared to $\gamma$ -V2O5. <i>Journal of Physical Chemistry B</i> , 2005, 109, 374-380.	2.6	19
255	Formation of one-dimensional crystalline silica on a metal substrate. <i>Surface Science</i> , 2006, 600, L164-L168.	1.9	19
256	Activation of C-H bonds mediated by Mo-Mo moieties in heterobimetallic Zn/O/Moclusters. <i>Dalton Transactions</i> , 2010, 39, 103-106.	3.3	19
257	Vanadium Oxide Oligomers and Ordered Monolayers Supported on CeO <sub>2</sub> (111): Structure and Stability Studied by Density Functional Theory. <i>Journal of Physical Chemistry C</i> , 2018, 122, 9101-9110.	3.1	19
258	Predicting adsorption selectivities from pure gas isotherms for gas mixtures in metal-organic frameworks. <i>Chemical Science</i> , 2020, 11, 643-655.	7.4	19
259	Methanol Adsorption on V2O3(0001). <i>Topics in Catalysis</i> , 2011, 54, 669-684.	2.8	18
260	Embedded and DFT Calculations on the Crystal Structures of Small Alkanes, Notably Propane. <i>Crystal Growth and Design</i> , 2017, 17, 1636-1646.	3.0	18
261	Acid strength of zeolitic Brønsted sites—Dependence on dielectric properties. <i>Catalysis Today</i> , 2019, 323, 86-93.	4.4	18
262	Interaction of ethene, 2-methylpropene, and benzene with the sodium(1+) ion. 1. Quantum chemical study of gas-phase complexes. <i>The Journal of Physical Chemistry</i> , 1982, 86, 1327-1332.	2.9	17
263	Optimized molecular integration schemes for density functional theory ab initio molecular dynamics simulations. <i>Chemical Physics Letters</i> , 1996, 255, 187-194.	2.6	17
264	The Th1 and Th2 cytokines IFN- $\beta$ and IL-4 antagonize the inhibition of monocyte IL-1 receptor antagonist by glucocorticoids: involvement of IL-1. <i>European Journal of Immunology</i> , 1998, 28, 2075-2085.	2.9	17
265	Influence of Ti Substitution on the $^{29}\text{Si}$ NMR Spectra of Silicalite. A Computational Study. <i>Zeitschrift Fur Physikalische Chemie</i> , 1999, 209, 21-32.	2.8	17
266	Vanadium Oxides Supported on a Thin Silica Film Grown on Mo(112): Insights from Density Functional Theory. <i>Journal of Physical Chemistry C</i> , 2009, 113, 8336-8342.	3.1	17
267	Vanadia and Water Coadsorption on Tetragonal Zirconia Surfaces. <i>Journal of Physical Chemistry C</i> , 2009, 113, 18191-18203.	3.1	17
268	Ultrathin Ti-Silicate Film on a Ru(0001) Surface. <i>Journal of Physical Chemistry C</i> , 2015, 119, 15443-15448.	3.1	17
269	Applications of Quantum Chemical Methods in Zeolite Science. <i>Studies in Surface Science and Catalysis</i> , 2007, , 701-XXI.	1.5	16
270	Stabilizing Gold Adatoms by Thiophenyl Derivatives: A Possible Route toward Metal Redispersion. <i>Journal of the American Chemical Society</i> , 2012, 134, 11161-11167.	13.7	16

#	ARTICLE	IF	CITATIONS
271	The structure of epitaxial V <sub>2</sub> O <sub>3</sub> films and their surfaces: A medium energy ion scattering study. <i>Surface Science</i> , 2012, 606, 1716-1727.	1.9	16
272	Gas-Phase Vibrational Spectroscopy of the Aluminum Oxide Anions (Al <sub>2</sub> O <sub>3</sub> ) <sub>n</sub> <sup>6-</sup> . <i>ChemPhysChem</i> , 2017, 18, 868-872.	2.1	16
273	Structure and Reactivity of Al <sup>3+</sup> O(H) <sup>2+</sup> Al Moieties in Siloxide Frameworks: Solution and Gas-Phase Model Studies. <i>Angewandte Chemie - International Edition</i> , 2019, 58, 902-906.	13.8	16
274	Elucidating Surface Structure with Action Spectroscopy. <i>Journal of the American Chemical Society</i> , 2020, 142, 2665-2671.	13.7	16
275	Isomerization and Selective Hydrogenation of Propyne: Screening of Metal-Organic Frameworks Modified by Atomic Layer Deposition. <i>Journal of the American Chemical Society</i> , 2020, 142, 20380-20389.	13.7	15
276	INDO/S calculations on simple silicon compounds and some silicon organic molecules. <i>Collection of Czechoslovak Chemical Communications</i> , 1982, 47, 1149-1168.	1.0	14
277	Charge differences between silicon atoms in aluminosilicates and their relation to <sup>29</sup> Si NMR chemical shifts. A quantum-chemical study. <i>Computational and Theoretical Chemistry</i> , 1986, 139, 113-124.	1.5	14
278	A Two-Dimensional "Zigzag" Silica Polymorph on a Metal Support. <i>Journal of the American Chemical Society</i> , 2018, 140, 6164-6168.	13.7	14
279	Valence and Structure Isomerism of Al <sub>2</sub> FeO <sub>4</sub> <sup>+</sup> : Synergy of Spectroscopy and Quantum Chemistry. <i>Journal of the American Chemical Society</i> , 2020, 142, 18050-18059.	13.7	14
280	Structure and Reactivity of Single-Site Vanadium Catalysts Supported on Metal-Organic Frameworks. <i>ACS Catalysis</i> , 2020, 10, 10051-10059.	11.2	14
281	Konsequenzen des Koopmanschen Theorems in den Restricted Hartree Fock Methoden für open-shell-Systeme. <i>Theoretica Chimica Acta</i> , 1975, 40, 129-141.	0.8	13
282	Sodium doped sodium sodalite: magnetic coupling between F centers and hyperfine interactions with framework atoms. <i>Physical Chemistry Chemical Physics</i> , 1999, 1, 4505.	2.8	13
283	On the interpretation of the experimental Raman spectrum of $\tilde{\ell}^2$ -eucryptite LiAlSiO <sub>4</sub> from atomistic computer modeling. <i>Journal of Non-Crystalline Solids</i> , 2000, 274, 264-270.	3.1	13
284	Structure and properties of bimetallic titanium and vanadium oxide clusters. <i>Physical Chemistry Chemical Physics</i> , 2014, 16, 8441.	2.8	13
285	Gas phase structures and charge localization in small aluminum oxide anions: Infrared photodissociation spectroscopy and electronic structure calculations. <i>Journal of Chemical Physics</i> , 2016, 144, 244305.	3.0	13
286	Ab initio Calculation of Rate Constants for Molecule-Surface Reactions with Chemical Accuracy. <i>Angewandte Chemie</i> , 2016, 128, 5321-5323.	2.0	13
287	Reaction dynamics of metal/oxide catalysts: Methanol oxidation at vanadium oxide films on Rh(111) from UHV to 10 <sup>-2</sup> mbar. <i>Journal of Catalysis</i> , 2020, 385, 255-264.	6.2	13
288	Chemically Accurate Vibrational Free Energies of Adsorption from Density Functional Theory Molecular Dynamics: Alkanes in Zeolites. <i>Journal of Chemical Theory and Computation</i> , 2021, 17, 5849-5862.	5.3	13

#	ARTICLE	IF	CITATIONS
289	Relative Stability of $\text{Al}^{\text{III}}\text{O}^{\text{IV}}\text{Al}$ Linkages in Zeolites. A Nonempirical Molecular Orbital Study. <i>Zeitschrift Fur Naturforschung - Section A Journal of Physical Sciences</i> , 1982, 37, 277-279.	1.5	12
290	The HF-AlF <sub>3</sub> gas-phase complex: An ab initio molecular orbital study. <i>Chemical Physics Letters</i> , 1989, 156, 125-128.	2.6	12
291	Interaction of SO <sub>3</sub> with c-ZrO <sub>2</sub> (111) films on Pt(111). <i>Physical Chemistry Chemical Physics</i> , 2006, 8, 1593.	2.8	12
292	Modeling environmental effects on charge density distributions in polar organometallics: Validation of embedded cluster models for the methyl lithium crystal. <i>Journal of Computational Chemistry</i> , 2010, 31, 2568-2576.	3.3	12
293	Oxygen Scrambling of CO <sub>2</sub> Adsorbed on CaO(001). <i>Journal of Physical Chemistry C</i> , 2017, 121, 18625-18634.	3.1	12
294	Syndnon- $\text{\AA}$ thylene. III. Chemische Reaktionen elektronisch angeregter 4-Styryl-syndnone. <i>Journal F\AAr Praktische Chemie</i> , 1978, 320, 81-90.	0.2	11
295	Orbital energies in open shell systems. <i>Journal of Chemical Physics</i> , 1978, 69, 495.	3.0	11
296	Opening of 3-membered rings to 4 <i>\pi</i> -electron systems. <i>Tetrahedron</i> , 1979, 35, 2109-2112.	1.9	11
297	Electronic structure and magnetic coupling in sodium electro sodalite: All-electron density functional calculations. <i>Journal of Chemical Physics</i> , 2000, 113, 5466.	3.0	11
298	Interaction Between Gold Atoms and Thio-Aryl Ligands on the Au(111) Surface. <i>Journal of Physical Chemistry C</i> , 2011, 115, 24871-24879.	3.1	11
299	Catalytically Active Vanadia Species on Silica: Effect of Oxygen and Water. <i>Journal of Physical Chemistry C</i> , 2014, 118, 29159-29163.	3.1	11
300	Dimerization of Linear Butenes and Pentenes in an Acidic Zeolite (H-MFI). <i>Angewandte Chemie - International Edition</i> , 2021, 60, 3529-3533.	13.8	11
301	Restricted Hartree-Fock-Berechnungen von open-shell-Systemen mittels halbempirischer MO-LCAO-SCF-Verfahren. <i>Collection of Czechoslovak Chemical Communications</i> , 1974, 39, 1235-1245.	1.0	11
302	Transferability test of open/2-type potential functions based on quantum-chemical interaction energies (QPEN). <i>Chemical Physics Letters</i> , 1984, 107, 530-534.	2.6	10
303	Adsorption of CH <sub>4</sub> on the Pt(111) surface: Random phase approximation compared to density functional theory. <i>Journal of Chemical Physics</i> , 2021, 155, 174702.	3.0	10
304	Charge-transfer-Anregungsenergien von Tetracyan- $\text{\AA}$ thylene-Molek\AA{l}komplexen phenylsubstituierter $\text{\AA}$ thylene und substituierter trans-Stilbene. <i>Journal F\AAr Praktische Chemie</i> , 1976, 318, 618-626.	0.2	9
305	Ab initio calculation of nuclear motion corrections to the geometries of water, methanol and silanol. <i>Molecular Physics</i> , 1991, 73, 335-348.	1.7	9
306	Ab initio molecular dynamics simulation of H <sub>5</sub> O <sub>2</sub> <sup>+</sup> and H <sub>7</sub> O <sub>3</sub> <sup>+</sup> gas phase clusters based on density functional theory. <i>Molecular Physics</i> , 1997, 91, 963-975.	1.7	9

#	ARTICLE	IF	CITATIONS
307	Interaction of CO with Electron-Rich Defects on MgO(100). <i>Journal of Physical Chemistry C</i> , 2013, 117, 8365-8373.	3.1	9
308	Theoretical interpretation of $^{29}\text{Si}$ NMR chemical shifts of aluminosilicates. <i>Computational and Theoretical Chemistry</i> , 1989, 183, 223-232.	1.5	8
309	The beryllium atomâ€”water molecule interaction A many-body perturbation theory study. <i>Chemical Physics Letters</i> , 1990, 174, 19-24.	2.6	8
310	â€œElektronenberechnungen an substituierten Sydnonen. <i>Zeitschrift fÃ¼r Chemie</i> , 1973, 13, 434-435.	0.0	8
311	Characterization of Phonon Vibrations of Silica Bilayer Films. <i>Journal of Physical Chemistry C</i> , 2019, 123, 7110-7117.	3.1	8
312	Hybrid Quantum Mechanics/ Molecular Mechanics Methods and their Application. , 2005, , 241-258.		8
313	Kooperative Bildung einer langreichweitig geordneten Wasserschicht auf der $\text{Fe}_{\langle \text{sub} \rangle 3} \text{O}_{\langle \text{sub} \rangle 4} \langle \text{sub} \rangle (111)$ â€”OberflÄche. <i>Angewandte Chemie</i> , 2018, 130, 1423-1428.	2.0	7
314	Thermochemistry of $\text{FeO}_{\langle \text{i} \rangle} \langle \text{sub} \rangle \text{m} \langle / \text{sub} \rangle \langle / \text{i} \rangle \text{H}_{\langle \text{i} \rangle} \langle \text{sub} \rangle \text{n} \langle / \text{sub} \rangle \langle \text{sup} \rangle \text{z} \langle / \text{sup} \rangle \langle / \text{i} \rangle$ Species: Assessment of Some DFT Functionals. <i>Journal of Chemical Theory and Computation</i> , 2020, 16, 2430-2435.	5.3	7
315	Sydnon-Äthylene. II. Elektronische Wechselwirkungen in 4-Styryl-sydnonen. <i>Journal fÃ¼r Praktische Chemie</i> , 1978, 320, 71-80.	0.2	6
316	Struktur und ReaktivitÄt der Alâ€O(H)â€Alâ€Einheiten in SiloxidgerÃ¼stverbindungen â€“ Modellstudien in LÄ¶sung und in Isolation. <i>Angewandte Chemie</i> , 2019, 131, 912-917.	2.0	6
317	Ab Initio Simulation of Cu-Species in Zeolites: Siting, Coordination, UV-Vis Spectra and Reactivity. , 2001, , 221-234.		6
318	Gasâ€Phase Mechanism of $\text{O}^{\langle \text{sup} \rangle \sim} \langle / \text{sup} \rangle \text{Ni}^{\langle \text{sup} \rangle 2+} \langle / \text{sup} \rangle$ Mediated Methane Conversion to Formaldehyde. <i>Angewandte Chemie - International Edition</i> , 2022, 61, .	13.8	6
319	Quantum Chemical Studies of Zeolite Acidity. <i>Studies in Surface Science and Catalysis</i> , 1989, 52, 73-90.	1.5	5
320	Nonuniform temperature dependence of the reactivity of disordered $\text{VOx}/\text{Al}_2\text{O}_3(001)$ surfaces: A density functional theory based Monte Carlo study. <i>Journal of Chemical Physics</i> , 2008, 129, 224710.	3.0	5
321	Aluminum siting in the framework of silicon rich zeolites. A ZSM-5 study. <i>Studies in Surface Science and Catalysis</i> , 2008, , 781-786.	1.5	4
322	Imaging of individual adatoms on oxide surfaces by dynamic force microscopy. <i>Physical Review B</i> , 2010, 81, .	3.2	4
323	Designing new catalysts: synthesis of new active structures: general discussion. <i>Faraday Discussions</i> , 2016, 188, 131-159.	3.2	4
324	Electronic States of $\text{Fe}_2\text{S}_-/0/+$ . <i>Collection of Czechoslovak Chemical Communications</i> , 2003, 68, 405-422.	1.0	4

#	ARTICLE	IF	CITATIONS
325	Effective semi-empirical excited state calculations using morokuma's EHP method. <i>Chemical Physics Letters</i> , 1978, 55, 119-124.	2.6	3
326	The H <sub>2</sub> O—Mg van der waals complex — a theoretical study. <i>Chemical Physics</i> , 1987, 113, 201-209.	1.9	3
327	Relative stability of alkoxides and carbocations in zeolites. QM/MM embedding and QM calculations applying periodic boundary conditions. <i>Studies in Surface Science and Catalysis</i> , 2002, 142, 643-649.	1.5	3
328	<i>Zeitschrift fÃ¼r Chemie</i> , 1977, 17, 30-30.	0.0	3
329	BasizitÃ¤t von Disiloxan: Nichtempirische Berechnung der Protonierungsenergie. <i>Zeitschrift fÃ¼r Chemie</i> , 1982, 22, 60-61.	0.0	3
330	Bridging model and real catalysts: general discussion. <i>Faraday Discussions</i> , 2016, 188, 565-589.	3.2	3
331	PROTON TRANSFER FROM ACIDIC SITES TO WATER, METHANOL AND AMMONIA. A COMPARATIVE AB INITIO STUDY. , 1993, , 679-686.		3
332	The Chemical Nature of Ti <sub>4</sub> O <sub>10</sub> : Vibrational Predissociation Spectroscopy Combined with Global Structure Optimization. <i>Journal of Physical Chemistry A</i> , 2021, 125, 9571-9577.	2.5	3
333	Structure and Reactivity of Solid Catalysts — Quantum Chemical Approach. <i>Studies in Surface Science and Catalysis</i> , 2007, , 19-26.	1.5	2
334	Catalyst design from theory to practice: general discussion. <i>Faraday Discussions</i> , 2016, 188, 279-307.	3.2	2
335	Wechselwirkung von Ethen mit Na <sup>+</sup> -Clonen in der Gasphase und in Zeolithen: Ab initio-Berechnung des Schwingungsspektrums. <i>Zeitschrift fÃ¼r Chemie</i> , 1985, 25, 254-255.	0.0	1
336	Publications of Joachim Sauer. <i>Journal of Physical Chemistry C</i> , 2019, 123, 7531-7539.	3.1	1
337	Chapter 25. Zeolite Modelling: Active Sites in Different Framework Structures and in Different Crystallographic Positions. , 2007, , 441-456.		1
338	Dimerization of Linear Butenes and Pentenes in an Acidic Zeolite (H-MFI). <i>Angewandte Chemie</i> , 2021, 133, 3571-3575.	2.0	1
339	Interactions of open-shell systems, RHF molecular potential equations. <i>Chemical Physics Letters</i> , 1977, 46, 438-441.	2.6	0
340	Isomers of weak C-C-complexes predicted by CT energy calculations. TCNE complexes with stilbene and analogs. <i>Advances in Molecular Relaxation and Interaction Processes</i> , 1978, 12, 1-11.	0.5	0
341	Ab initio study of the interaction of methanol with BrÃ¶nsted acid sites of zeolites. <i>Studies in Surface Science and Catalysis</i> , 1995, 98, 242-243.	1.5	0
342	HISTORY OF SCIENCE:Before the Fall of the Wall. <i>Science</i> , 2000, 287, 1210-1211.	12.6	0

#	ARTICLE	IF	CITATIONS
343	Bond Dissociation Energies and Structures of CuNO+ and Cu(NO)2+. ChemInform, 2003, 34, no.	0.0	0
344	Electronic States of Fe2S-/0/+.. ChemInform, 2003, 34, no.	0.0	0
345	Environmental Effects on Vibrational Proton Dynamics in H5O2+: DFT Study on Crystalline H5O2+ClO4-. ChemInform, 2005, 36, no.	0.0	0
346	Local geometry of AlO4â˜” and SiO4 tetrahedra in the silicone rich chabazite. A combined high resolution NMR and QM/MM study. Studies in Surface Science and Catalysis, 2008, , 729-732.	1.5	0
347	Structure and Thermochemistry of Fe<sub>2</sub>S<sub>2</sub><sup>â€¢0/+</sup> Gas Phase Clusters and Their Fragments. B3LYP Calculations.. ChemInform, 2002, 33, 2-2.	0.0	0
348	Students, Postdoctoral Associates, and Collaborators of Joachim Sauer. Journal of Physical Chemistry C, 2019, 123, 7527-7528.	3.1	0
349	Curriculum Vitae of Joachim Sauer. Journal of Physical Chemistry C, 2019, 123, 7529-7530.	3.1	0
350	Autobiography of Joachim Sauer. Journal of Physical Chemistry C, 2019, 123, 7521-7526.	3.1	0
351	Gas Phase Mechanism of Oâ€¢â˜”/Ni2+â€¢mediated Methane Conversion to Formaldehyde. Angewandte Chemie, 0, , .	2.0	0