## Martina Kieninger

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	SVECV â€f12: Benchmark of a composite scheme for accurate and cost effective evaluation of reaction barriers. International Journal of Quantum Chemistry, 2021, 121, e26745.	2.0	18
2	A reinvestigation of the deceptively simple reaction of toluene with OH, and the fate of the benzyl radical: a combined thermodynamic and kinetic study on the competition between OH-addition and H-abstraction reactions. Theoretical Chemistry Accounts, 2020, 139, 1.	1.4	5
3	Reinvestigation of the Deceptively Simple Reaction of Toluene with OH and the Fate of the Benzyl Radical: The "Hidden―Routes to Cresols and Benzaldehyde. Journal of Physical Chemistry A, 2020, 124, 5917-5930.	2.5	18
4	Enthalpies of formation of the benzyloxyl, benzylperoxyl, hydroxyphenyl radicals and related species on the potential energy surface for the reaction of toluene with the hydroxyl radical. Theoretical Chemistry Accounts, 2019, 138, 1.	1.4	3
5	Basis Set Effects in the Description of the Cl-O Bond in ClO and XClO/ClOX Isomers (X = H, O, and Cl) Using DFT and CCSD(T) Methods. Journal of Chemistry, 2019, 2019, 1-23.	1.9	5
6	Using density functional theory to increase the accuracy of experimental crystal structures: The case of potassium peroxocarbonate. Journal of Molecular Structure, 2017, 1146, 1-4.	3.6	1
7	Atmospheric reactivity of HCî€,CCH2OH (2-propyn-1-ol) toward OH radicals: experimental determination and theoretical comparison with its alkyne analogue. RSC Advances, 2015, 5, 106668-106679.	3.6	0
8	A Festschrift in honor of Sándor Suhai's 65th birthday. Theoretical Chemistry Accounts, 2010, 125, 101-105.	1.4	0
9	On the structure, infrared and Raman spectra of the 2:1 cysteine–Zn complex. Theoretical Chemistry Accounts, 2010, 125, 279-291.	1.4	6
10	On the experimental structure of monoperoxocarbonic acid and the enthalpy of formation of carbonic acid, peroxyformic acid and monoperoxocarbonic acid in gas phase. Chemical Physics Letters, 2009, 480, 52-56.	2.6	3
11	Tautomeric Forms of 2-Thiobarbituric Acid As Studied in the Solid, in Polar Solutions, and on Gold Nanoparticles. Journal of Physical Chemistry C, 2007, 111, 3369-3383.	3.1	59
12	Comparison of large basis set DFT and MP2 calculations in the study of the barrier for internal rotation of 2,3,5,6-tetrafluoroanisole. International Journal of Quantum Chemistry, 2007, 107, 403-417.	2.0	3
13	A New Perspective in the Lewis Acid Catalyzed Ring Opening of Epoxides. Theoretical Study of Some Complexes of Methanol, Acetic Acid, Dimethyl Ether, Diethyl Ether, and Ethylene Oxide with Boron Trifluoride. Journal of Physical Chemistry A, 2006, 110, 11734-11751.	2.5	12
14	Molecular Structure and Internal Rotation in 2,3,5,6-Tetrafluoroanisole as Studied by Gas-Phase Electron Diffraction and Quantum Chemical Calculations. Journal of Physical Chemistry A, 2005, 109, 394-399.	2.5	11
15	A comparative density functional study of the torsional potential of 4-fluoro (trifluoromethoxy)benzene and related species. Chemical Physics Letters, 2004, 389, 405-412.	2.6	7
16	Density Functional Computational Thermochemistry:  Determination of the Enthalpy of Formation of Methanethial-S,S-dioxide (Sulfene). Journal of Physical Chemistry A, 2003, 107, 518-521.	2.5	11
17	Computational determination of the enthalpy of formation of alkylthial S-oxides and alkylthione S-oxides: a study of (Z)-propanethial-S-oxide, the lachrymatory factor of the onion (Allium cepa). Physical Chemistry Chemical Physics, 2002, 4, 4328-4333.	2.8	3
18	Density functional computational thermochemistry: solving the discrepancy between MO and DFT calculations on the enthalpy of formation of sulfine, CH2r̃Sr̃O. Chemical Physics Letters, 2002, 355, 207-213.	2.6	19

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19	Complete basis set and density functional determination of the enthalpy of formation of the controversial HO3 radical: a discrepancy between theory and experiment. Chemical Physics Letters, 2002, 365, 440-449.	2.6	39
20	Density Functional Computational Thermochemistry:Â Isomerization of Sulfine and Its Enthalpy of Formation. Journal of Physical Chemistry A, 2001, 105, 9912-9916.	2.5	17
21	Density functional computational thermochemistry: determination of the enthalpy of formation of sulfine, CH2ĩSĩO, at room temperature. Chemical Physics Letters, 2000, 329, 145-153.	2.6	17
22	Density functional and coupled-cluster calculations of isodesmic reactions involving fluorine oxides. Chemical Physics Letters, 1999, 301, 331-335.	2.6	20
23	Density Functional Theory Is More Accurate Than Coupled-Cluster Theory in the Study of the Thermochemistry of Species Containing the Fâ^'O Bond. Journal of Physical Chemistry A, 1999, 103, 147-151.	2.5	32
24	A discrepancy between experimental and theoretical thermochemical characterization of some oxygen fluorides. Chemical Physics Letters, 1998, 287, 597-600.	2.6	28
25	Density functional and ab initio study of the free radical MgNC. Computational and Theoretical Chemistry, 1998, 422, 133-141.	1.5	10
26	Glycine conformations: gradient-corrected DFT-studies. Computational and Theoretical Chemistry, 1998, 433, 193-201.	1.5	6
27	Density functional investigations of carboxyl free radicals: Formyloxyl, acetyloxyl, and benzoyloxyl radicals. International Journal of Quantum Chemistry, 1998, 70, 253-267.	2.0	34
28	Computational chemistry as an analytical tool: thermochemical examples in atmospheric chemistry. Pure and Applied Chemistry, 1998, 70, 2301-2307.	1.9	5
29	Equilibrium structure of the carbon dioxide-water complex in the gas phase: an ab initio and density functional study. Computational and Theoretical Chemistry, 1997, 390, 157-167.	1.5	21
30	Title is missing!. Molecular Engineering, 1997, 7, 317-348.	0.2	2
31	Density functional study of isomerization of fluoro- and chloroformaldehyde radical cations. Journal of Computational Chemistry, 1996, 17, 1309-1317.	3.3	10
32	Conformational and energetic properties of the ammonia dimer?comparison of post-Hartree?Fock and density functional methods. Journal of Computational Chemistry, 1996, 17, 1508-1519.	3.3	17
33	Density functional studies of internal rotation: formamide as a prototype of the peptide bond. Journal of Molecular Structure, 1996, 375, 181-188.	3.6	9
34	The FO2 radical: a new success of density functional theory. Chemical Physics Letters, 1995, 245, 488-497.	2.6	50
35	Density functional studies on hydrogen-bonded complexes. International Journal of Quantum Chemistry, 1994, 52, 465-478.	2.0	46
36	The chemical Hamiltonian approach in density functional theory. Chemical Physics Letters, 1994, 230, 485-490.	2.6	22