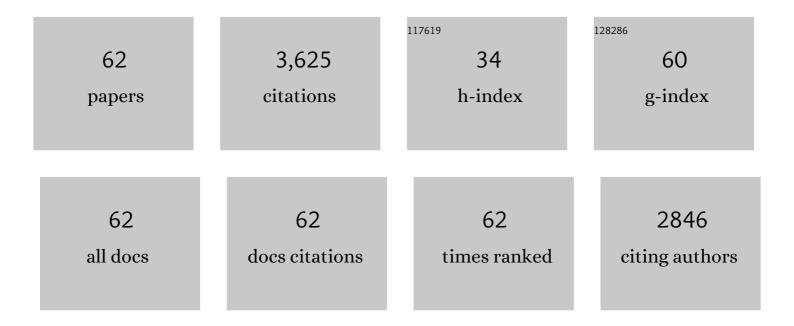
Jürgen Gräfenstein

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Interplay between Förster and Dexter Energy Transfer Rates in Isomeric Donor–Bridge–Acceptor Systems. Journal of Physical Chemistry A, 2020, 124, 7219-7227.	2.5	20
2	The Structure of the "Vibration Hole―around an Isotopic Substitution—Implications for the Calculation of Nuclear Magnetic Resonance (NMR) Isotopic Shifts. Molecules, 2020, 25, 2915.	3.8	3
3	Efficient calculation of NMR isotopic shifts: Difference-dedicated vibrational perturbation theory. Journal of Chemical Physics, 2019, 151, 244120.	3.0	7
4	Carbon's Three-Center, Four-Electron Tetrel Bond, Treated Experimentally. Journal of the American Chemical Society, 2018, 140, 17571-17579.	13.7	53
5	Halogen Bond Asymmetry in Solution. Journal of the American Chemical Society, 2018, 140, 13503-13513.	13.7	57
6	Photochemically Induced Aryl Azide Rearrangement: Solution NMR Spectroscopic Identification of the Rearrangement Product. Journal of Organic Chemistry, 2017, 82, 1812-1816.	3.2	3
7	The ¹⁵ N NMR chemical shift in the characterization of weak halogen bonding in solution. Faraday Discussions, 2017, 203, 333-346.	3.2	25
8	Halogen Bonding: A Powerful Tool for Modulation of Peptide Conformation. Biochemistry, 2017, 56, 3265-3272.	2.5	65
9	Naphthalene Derivatives from the Roots of <i>Pentas parvifolia</i> and <i>Pentas bussei</i> . Journal of Natural Products, 2016, 79, 2181-2187.	3.0	32
10	Substituent Effects on the [N–l–N] ⁺ Halogen Bond. Journal of the American Chemical Society, 2016, 138, 9853-9863.	13.7	89
11	Counterion influence on the N–I–N halogen bond. Chemical Science, 2015, 6, 3746-3756.	7.4	100
12	The nature of [N–Cl–N] ⁺ and [N–F–N] ⁺ halogen bonds in solution. Chemical Science, 2014, 5, 3226-3233.	7.4	66
13	Solvent effects on halogen bond symmetry. CrystEngComm, 2013, 15, 3087.	2.6	66
14	A computational study of the enantioselective addition of n-BuLi to benzaldehyde in the presence of a chiral lithium N,P amide. Organic and Biomolecular Chemistry, 2012, 10, 2807.	2.8	12
15	Symmetry of [N–X–N] ⁺ halogen bonds in solution. Chemical Communications, 2012, 48, 1458-1460.	4.1	76
16	Symmetric Halogen Bonding Is Preferred in Solution. Journal of the American Chemical Society, 2012, 134, 5706-5715.	13.7	159
17	Aggregation and Solvation of Chiral N,Pâ€Amide Ligands in Coordinating Solvents: A Computational and NMR Spectroscopic Study. ChemPlusChem, 2012, 77, 799-806.	2.8	11
18	An efficient algorithm for the density-functional theory treatment of dispersion interactions. Journal of Chemical Physics, 2009, 130, 124105.	3.0	98

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19	The self-interaction error and the description of non-dynamic electron correlation in density functional theory. Theoretical Chemistry Accounts, 2009, 123, 171-182.	1.4	51
20	Unusual long-range spin-spin coupling in fluorinated polyenes: A mechanistic analysis. Journal of Chemical Physics, 2007, 127, 174704.	3.0	10
21	Avoiding singularity problems associated with meta-GGA (generalized gradient approximation) exchange and correlation functionals containing the kinetic energy density. Journal of Chemical Physics, 2007, 127, 214103.	3.0	62
22	Efficient density-functional theory integrations by locally augmented radial grids. Journal of Chemical Physics, 2007, 127, 164113.	3.0	53
23	Calculation and analysis of NMR spin–spin coupling constants. Physical Chemistry Chemical Physics, 2007, 9, 2791-2816.	2.8	95
24	Development of a CAS-DFT method covering non-dynamical and dynamical electron correlation in a balanced way. Molecular Physics, 2005, 103, 279-308.	1.7	109
25	Analysis of long-range NMR spin–spin coupling in polyenes and the π-mechanism. Physical Chemistry Chemical Physics, 2005, 7, 452-462.	2.8	17
26	Elucidation of the Electronic Structure of Molecules with the Help of NMR Spinâ^'Spin Coupling Constants:Â The FH Molecule. Journal of Physical Chemistry A, 2005, 109, 2325-2339.	2.5	9
27	The impact of the self-interaction error on the density functional theory description of dissociating radical cations: Ionic and covalent dissociation limits. Journal of Chemical Physics, 2004, 120, 524-539.	3.0	141
28	Decomposition of nuclear magnetic resonance spin–spin coupling constants into active and passive orbital contributions. Journal of Chemical Physics, 2004, 120, 9952-9968.	3.0	16
29	Analysis of the paramagnetic spin–orbit transmission mechanism for NMR spin–spin coupling constants using the paramagnetic spin–orbit density distribution. Chemical Physics Letters, 2004, 383, 332-342.	2.6	33
30	Systematic strategy for decoding the NMR spin–spin coupling mechanism: the J-OC-PSP method. Magnetic Resonance in Chemistry, 2004, 42, S138-S157.	1.9	13
31	Analysis of the spin-dipole transmission mechanism for NMR spin–spin coupling constants using orbital contributions, spin polarization, and spin-dipole energy density distribution. Chemical Physics Letters, 2004, 387, 415-427.	2.6	24
32	Analysis of the NMR through-space coupling mechanism between 19F atoms. Chemical Physics Letters, 2004, 394, 5-13.	2.6	40
33	Effect of the self-interaction error for three-electron bonds: On the development of new exchange-correlation functionals. Physical Chemistry Chemical Physics, 2004, 6, 1096-1112.	2.8	107
34	Investigation of the Ï€ Character of a Câ^'C Bond with the Help of the Diamagnetic and Paramagnetic Spinâ^'Orbit Term of the NMR Spinâ^'Spin Coupling Constant. Journal of Physical Chemistry A, 2004, 108, 4520-4535.	2.5	18
35	Analysis of the NMR Spinâ^'Spin Coupling Mechanism Across a Hâ^'Bond:  Nature of the H-Bond in Proteins. Journal of Physical Chemistry B, 2004, 108, 1115-1129.	2.6	34
36	One-electron versus electron–electron interaction contributions to the spin–spin coupling mechanism in nuclear magnetic resonance spectroscopy: Analysis of basic electronic effects. Journal of Chemical Physics, 2004, 121, 12217.	3.0	9

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37	Long-range and short-range Coulomb correlation effects as simulated by Hartree-Fock, local density approximation, and generalized gradient approximation exchange functionals. Theoretical Chemistry Accounts, 2003, 109, 22-35.	1.4	81
38	Analysis of the Transmission Mechanism of NMR Spinâ~'Spin Coupling Constants Using Fermi Contact Spin Density Distribution, Partial Spin Polarization, and Orbital Currents:  XHn Molecules. Journal of Physical Chemistry A, 2003, 107, 7043-7056.	2.5	51
39	Can Unrestricted Density-Functional Theory Describe Open Shell Singlet Biradicals?. International Journal of Molecular Sciences, 2002, 3, 360-394.	4.1	182
40	Influence of the self-interaction error on the structure of the DFT exchange hole. Chemical Physics Letters, 2002, 352, 469-478.	2.6	65
41	On the diagnostic value of (Ŝ2) in Kohn-Sham density functional theory. Molecular Physics, 2001, 99, 981-989.	1.7	89
42	The ozone–acetylene reaction: concerted or non-concerted reaction mechanism? A quantum chemical investigation. Chemical Physics Letters, 2001, 347, 268-276.	2.6	25
43	4-Oxo-2,3,5,6-tetrafluorocyclohexa-2,5-dienylidene—A Highly Electrophilic Triplet Carbene. Chemistry - A European Journal, 2000, 6, 4567-4579.	3.3	43
44	The combination of density functional theory with multi-configuration methods – CAS-DFT. Chemical Physics Letters, 2000, 316, 569-577.	2.6	139
45	Can density functional theory describe multi-reference systems? Investigation of carbenes and organic biradicals. Physical Chemistry Chemical Physics, 2000, 2, 2091-2103.	2.8	135
46	What correlation effects are covered by density functional theory?. Molecular Physics, 2000, 98, 1639-1658.	1.7	55
47	An Accurate Description of the Bergman Reaction Using Restricted and Unrestricted DFT:Â Stability Test, Spin Density, and On-Top Pair Densityâ€. Journal of Physical Chemistry A, 2000, 104, 1748-1761.	2.5	250
48	Nuclear magnetic resonance spin–spin coupling constants from coupled perturbed density functional theory. Journal of Chemical Physics, 2000, 113, 3530-3547.	3.0	311
49	Density functional theory for open-shell singlet biradicals. Chemical Physics Letters, 1998, 288, 593-602.	2.6	103
50	α,3-Didehydro-5-methyl-6-hydroxytoluene: Matrix Isolation of a Diradical Related to the Neocarzinostatin Chromophore. Journal of the American Chemical Society, 1998, 120, 8480-8485.	13.7	33
51	Valence-band structure of group-IV semiconductors by means of local increments. Physical Review B, 1997, 55, 13588-13597.	3.2	43
52	Trimesitylsilylium cation verification of a free silylium cation in solution by NMR chemical shift calculations. Chemical Physics Letters, 1997, 279, 9-16.	2.6	16
53	Andersen's force theorem and the local stress field. Physical Review B, 1996, 53, 7143-7146.	3.2	75
54	Ab initioground-state correlation calculations for semiconductors with the local ansatz. Physical Review B, 1995, 51, 10556-10567.	3.2	17

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55	Computation of the valence band of diamond by means of local increments. Chemical Physics Letters, 1993, 215, 611-616.	2.6	37
56	Sum rule for planar jellium surfaces treated within the Kohn-Sham local-density approximation. Physical Review B, 1992, 46, 1715-1718.	3.2	6
57	An Alternative Approach to Calculate the Hydrostatic Pressure in Solids. Physica Status Solidi (B): Basic Research, 1990, 158, K133.	1.5	1
58	Comment on â€~â€~Stress field in quantum systems''. Physical Review B, 1990, 41, 3245-3247.	3.2	8
59	Calculated Fermi Surface Characteristics of the Noble Metals. Physica Status Solidi (B): Basic Research, 1988, 147, 575-582.	1.5	4
60	Local quantum mechanical stress in clusters and crystals. Physica Scripta, 1988, 37, 370-372.	2.5	10
61	Quantum-mechanical stress and a generalized virial theorem for clusters and solids. Physical Review B, 1988, 37, 8167-8178.	3.2	59
62	Anisotropic electron-impurity scattering rates of dilute noble-metal alloys: a comparison with surface state resonance data. Journal of Physics F: Metal Physics, 1988, 18, 731-743.	1.6	4