Johannes Neugebauer

List of Publications by Year in descending order

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150 papers 6,472 citations

50170 46 h-index 74018 75 g-index

164 all docs

164 docs citations

164 times ranked 4210 citing authors

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| 1 | Quantum chemical calculation of vibrational spectra of large molecules?Raman and IR spectra for Buckminsterfullerene. Journal of Computational Chemistry, 2002, 23, 895-910. | 1.5 | 506 |
| 2 | Subsystem densityâ€functional theory. Wiley Interdisciplinary Reviews: Computational Molecular Science, 2014, 4, 325-362. | 6.2 | 282 |
| 3 | Fundamental vibrational frequencies of small polyatomic molecules from density-functional calculations and vibrational perturbation theory. Journal of Chemical Physics, 2003, 118, 7215. | 1.2 | 259 |
| 4 | Couplings between electronic transitions in a subsystem formulation of time-dependent density functional theory. Journal of Chemical Physics, 2007, 126, 134116. | 1.2 | 214 |
| 5 | The merits of the frozen-density embedding scheme to model solvatochromic shifts. Journal of Chemical Physics, 2005, 122, 094115. | 1.2 | 207 |
| 6 | Accurate frozen-density embedding potentials as a first step towards a subsystem description of covalent bonds. Journal of Chemical Physics, 2010, 132, 164101. | 1.2 | 172 |
| 7 | A flexible implementation of frozenâ€density embedding for use in multilevel simulations. Journal of Computational Chemistry, 2008, 29, 1011-1018. | 1.5 | 138 |
| 8 | An Explicit Quantum Chemical Method for Modeling Large Solvation Shells Applied to Aminocoumarin C151. Journal of Physical Chemistry A, 2005, 109, 7805-7814. | 1.1 | 130 |
| 9 | Assessment of a simple correction for the long-range charge-transfer problem in time-dependent density-functional theory. Journal of Chemical Physics, 2006, 124, 214102. | 1.2 | 126 |
| 10 | Chromophore-specific theoretical spectroscopy: From subsystem density functional theory to mode-specific vibrational spectroscopy. Physics Reports, 2010, 489, 1-87. | 10.3 | 118 |
| 11 | Decarboxylative Polymerization of 2,6-Naphthalenedicarboxylic Acid at Surfaces. Journal of the American Chemical Society, 2014, 136, 9658-9663. | 6.6 | 114 |
| 12 | Quantum Chemical Description of Absorption Properties and Excitedâ€State Processes in Photosynthetic Systems. ChemPhysChem, 2012, 13, 386-425. | 1.0 | 107 |
| 13 | A mode-selective quantum chemical method for tracking molecular vibrations applied to functionalized carbon nanotubes. Journal of Chemical Physics, 2003, 118, 1634-1641. | 1.2 | 106 |
| 14 | Properties of WAu12. Physical Chemistry Chemical Physics, 2004, 6, 11-22. | 1.3 | 97 |
| 15 | Subsystemâ€Based Theoretical Spectroscopy of Biomolecules and Biomolecular Assemblies. ChemPhysChem, 2009, 10, 3148-3173. | 1.0 | 91 |
| 16 | Photophysical Properties of Natural Light-Harvesting Complexes Studied by Subsystem Density Functional Theory. Journal of Physical Chemistry B, 2008, 112, 2207-2217. | 1.2 | 90 |
| 17 | Comparison of frozen-density embedding and discrete reaction field solvent models for molecular properties. Physical Chemistry Chemical Physics, 2006, 8, 2349. | 1.3 | 87 |
| 18 | A vibrational circular dichroism implementation within a Slater-type-orbital based density functional framework and its application to hexa- and hepta-helicenes. Theoretical Chemistry Accounts, 2008, 119, 245-263. | 0.5 | 87 |

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| 19 | Modelling charge transfer reactions with the frozen density embedding formalism. Journal of Chemical Physics, 2011, 135, 234103. | 1.2 | 87 |
| 20 | Gas-Phase Câ^'H and Nâ^'H Bond Activation by a High Valent Nitrido-Iron Dication and ã€^NH〉-Transfer to Activated Olefins. Journal of the American Chemical Society, 2008, 130, 4285-4294. | 6.6 | 85 |
| 21 | Resonance Raman spectra of uracil based on Kramers–Kronig relations using time-dependent density functional calculations and multireference perturbation theory. Journal of Chemical Physics, 2004, 120, 11564-11577. | 1.2 | 81 |
| 22 | Vibronic coupling and double excitations in linear response time-dependent density functional calculations: Dipole-allowed states of N2. Journal of Chemical Physics, 2004, 121, 6155-6166. | 1.2 | 80 |
| 23 | A Subsystem TDDFT Approach for Solvent Screening Effects on Excitation Energy Transfer Couplings. Journal of Chemical Theory and Computation, 2010, 6, 1843-1851. | 2.3 | 77 |
| 24 | The electronic structure of the primary electron donor of reaction centers of purple bacteria at atomic resolution as observed by photo-CIDNP ¹³ C NMR. Proceedings of the National Academy of Sciences of the United States of America, 2009, 106, 22281-22286. | 3.3 | 74 |
| 25 | Effects of Complex Formation on Vibrational Circular Dichroism Spectra. Journal of Physical Chemistry A, 2008, 112, 6978-6991. | 1.1 | 73 |
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| 28 | First-principles calculation of electronic spectra of light-harvesting complex II. Physical Chemistry Chemical Physics, 2011, 13, 10475. | 1.3 | 67 |
| 29 | An accurate and linear-scaling method for calculating charge-transfer excitation energies and diabatic couplings. Journal of Chemical Physics, 2013, 138, 054101. | 1.2 | 67 |
| 30 | Finding a needle in a haystack: direct determination of vibrational signatures in complex systems. New Journal of Chemistry, 2007, 31, 818. | 1.4 | 66 |
| 31 | Analysis of electron density distributions from subsystem density functional theory applied to coordination bonds. Chemical Physics Letters, 2008, 461, 353-359. | 1.2 | 66 |
| 32 | Vibronic Structure of the Permanganate Absorption Spectrum from Time-Dependent Density Functional Calculations. Journal of Physical Chemistry A, 2005, 109, 1168-1179. | 1.1 | 65 |
| 33 | Modeling solvent effects on electron-spin-resonance hyperfine couplings by frozen-density embedding. Journal of Chemical Physics, 2005, 123, 114101. | 1.2 | 64 |
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| 36 | M <scp>O</scp> V <scp>I</scp> P <scp>AC</scp> : Vibrational spectroscopy with a robust metaâ€program for massively parallel standard and inverse calculations. Journal of Computational Chemistry, 2012, 33, 2186-2198. | 1.5 | 59 |

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| 42 | Phytochrome as Molecular Machine: Revealing Chromophore Action during the Pfr â†' Pr Photoconversion by Magic-Angle Spinning NMR Spectroscopy. Journal of the American Chemical Society, 2010, 132, 4431-4437. | 6.6 | 55 |
| 43 | Importance of vibronic effects on the circular dichroism spectrum of dimethyloxirane. Journal of Chemical Physics, 2005, 122, 234305. | 1.2 | 54 |
| 44 | Design of Ru(II)-NHC-Diamine Precatalysts Directed by Ligand Cooperation: Applications and Mechanistic Investigations for Asymmetric Hydrogenation. Journal of the American Chemical Society, 2020, 142, 7100-7107. | 6.6 | 53 |
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| 51 | Direct determination of exciton couplings from subsystem time-dependent density-functional theory within the Tamm–Dancoff approximation. Journal of Chemical Physics, 2013, 138, 034104. | 1.2 | 41 |
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| 86 | Photochemically induced dynamic nuclear polarization NMR on photosystem II: donor cofactor observed in entire plant. Scientific Reports, 2018, 8, 17853. | 1.6 | 16 |
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