Ilaria Ciofini

List of Publications by Year in Descending Order

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

10,226 50 91 234 h-index g-index citations papers 6.41 258 11,155 5.4 L-index avg, IF ext. papers ext. citations

#	Paper	IF	Citations
234	A global analysis of excited states: the global transition contribution grids. <i>Theoretical Chemistry Accounts</i> , 2021 , 140, 1	1.9	O
233	Assessing challenging intra- and inter-molecular charge-transfer excitations energies with double-hybrid density functionals. <i>Journal of Computational Chemistry</i> , 2021 , 42, 970-981	3.5	10
232	A combined Monte Carlo/DFT approach to simulate UV-vis spectra of molecules and aggregates: Merocyanine dyes as a case study. <i>Journal of Computational Chemistry</i> , 2021 , 42, 1054-1063	3.5	O
231	Role of dppf Monoxide in the Transmetalation Step of the SuzukiMiyaura Coupling Reaction. <i>Organometallics</i> , 2021 , 40, 1120-1128	3.8	2
230	Chasing unphysical TD-DFT excited states in transition metal complexes with a simple diagnostic tool. <i>Journal of Chemical Physics</i> , 2021 , 154, 204102	3.9	1
229	Modeling UV-Vis spectra of low dimensional materials using electrostatic embedding: The case of CdSe. <i>Journal of Computational Chemistry</i> , 2021 , 42, 1212-1224	3.5	О
228	Metal-Free Deoxygenation of Amine N-Oxides: Synthetic and Mechanistic Studies. <i>ChemPhysChem</i> , 2021 , 22, 1237-1242	3.2	1
227	Direct Synthesis of CFH-Substituted 2-Amidofurans via Copper-Catalyzed Addition of Difluorinated Diazoacetone to Ynamides. <i>Organic Letters</i> , 2021 , 23, 5528-5532	6.2	3
226	On the Interplay between Molecular Packing and Optical Response in Thin Films for Organic Photovoltaics. <i>Journal of Physical Chemistry C</i> , 2021 , 125, 16304-16315	3.8	
225	Electronic coupling in the reduced state lies at the origin of color changes of ommochromes. <i>Dyes and Pigments</i> , 2021 , 185, 108661	4.6	2
224	Mechanochromic LLDPE Films Doped with NIR Reflective Paliogen Black. <i>Macromolecular Rapid Communications</i> , 2021 , 42, e2000426	4.8	2
223	Copper-catalyzed transformation of alkyl nitriles to -arylacetamide using diaryliodonium salts <i>RSC Advances</i> , 2021 , 11, 15885-15889	3.7	2
222	Macrolactonization Reactions Driven by a Pentafluorobenzoyl Group*. <i>Angewandte Chemie - International Edition</i> , 2021 , 60, 19843-19851	16.4	1
221	Macrolactonization Reactions Driven by a Pentafluorobenzoyl Group**. <i>Angewandte Chemie</i> , 2021 , 133, 19996-20004	3.6	
220	Improving the heterointerface in hybrid organic-inorganic perovskite solar cells by surface engineering: Insights from periodic hybrid density functional theory calculations. <i>Journal of Computational Chemistry</i> , 2020 , 41, 1740-1747	3.5	5
219	General Density-Based Index to Analyze Charge Transfer Phenomena: From Models to Butterfly Molecules. <i>Journal of Chemical Theory and Computation</i> , 2020 , 16, 4543-4553	6.4	10
218	Electron Spin Densities and Density Functional Approximations: Open-Shell Polycyclic Aromatic Hydrocarbons as Case Study. <i>Journal of Chemical Theory and Computation</i> , 2020 , 16, 3567-3577	6.4	6

(2019-2020)

217	An electron density based analysis to establish the electronic adiabaticity of proton coupled electron transfer reactions. <i>Journal of Computational Chemistry</i> , 2020 , 41, 1835-1841	3.5	8
216	Rationally Designed Long-Wavelength Absorbing Ru(II) Polypyridyl Complexes as Photosensitizers for Photodynamic Therapy. <i>Journal of the American Chemical Society</i> , 2020 , 142, 6578-6587	16.4	74
215	Rationally designed ruthenium complexes for 1- and 2-photon photodynamic therapy. <i>Nature Communications</i> , 2020 , 11, 3262	17.4	83
214	Electron Storage System Based on a Two-Way Inversion of Redox Potentials. <i>Journal of the American Chemical Society</i> , 2020 , 142, 5162-5176	16.4	9
213	Double hybrids and time-dependent density functional theory: An implementation and benchmark on charge transfer excited states. <i>Journal of Computational Chemistry</i> , 2020 , 41, 1242-1251	3.5	26
212	Interfacial Engineering through Chloride-Functionalized Self-Assembled Monolayers for High-Performance Perovskite Solar Cells. <i>ACS Applied Materials & Discrete Solar Cells</i> , 12, 744-752	9.5	28
211	Silver(I) Oxide-/DBU-Promoted Synthesis of Dihydrofuran Units through Allenyl Silver Formation. <i>Chemistry - A European Journal</i> , 2020 , 26, 17455-17461	4.8	1
210	Theoretical insights on acceptor-donor dyads for organic photovoltaics. <i>Physical Chemistry Chemical Physics</i> , 2020 , 22, 27413-27424	3.6	O
209	Modeling the Electron Transfer Chain in an Artificial Photosynthetic Machine. <i>Journal of Physical Chemistry Letters</i> , 2020 , 11, 9738-9744	6.4	
208	A DFT Protocol for the Prediction of 31P NMR Chemical Shifts of Phosphine Ligands in First-Row Transition-Metal Complexes. <i>Organometallics</i> , 2020 , 39, 3121-3130	3.8	4
207	Computation of covalent and noncovalent structural parameters at low computational cost: Efficiency of the DH-SVPD method. <i>International Journal of Quantum Chemistry</i> , 2020 , 120, e26233	2.1	2
206	A Comparative Investigation of the Role of the Anchoring Group on Perylene Monoimide Dyes in NiO-Based Dye-Sensitized Solar Cells. <i>ChemSusChem</i> , 2020 , 13, 1844-1855	8.3	6
205	A Ru(II) polypyridyl complex bearing aldehyde functions as a versatile synthetic precursor for long-wavelength absorbing photodynamic therapy photosensitizers. <i>Bioorganic and Medicinal Chemistry</i> , 2019 , 27, 2666-2675	3.4	28
204	Atypical Lone PairInteraction with Quinone Methides in a Series of Imido-Ferrociphenol Anticancer Drug Candidates. <i>Angewandte Chemie</i> , 2019 , 131, 8509	3.6	
203	Excited state tracking during the relaxation of coordination compounds. <i>Journal of Computational Chemistry</i> , 2019 , 40, 1420-1428	3.5	8
202	Small Basis Set Allowing the Recovery of Dispersion Interactions with Double-Hybrid Functionals. Journal of Chemical Theory and Computation, 2019 , 15, 2944-2953	6.4	7
201	Atypical Lone Pair-Interaction with Quinone Methides in a Series of Imido-Ferrociphenol Anticancer Drug Candidates. <i>Angewandte Chemie - International Edition</i> , 2019 , 58, 8421-8425	16.4	21
200	Photophysical properties of fluorescent imaging biological probes of nucleic acids: SAC-CI and TD-DFT Study. <i>Journal of Computational Chemistry</i> , 2019 , 40, 127-134	3.5	3

199	Low-Temperature Solution Synthesis of Au-Modified ZnO Nanowires for Highly Efficient Hydrogen Nanosensors. <i>ACS Applied Materials & Amp; Interfaces</i> , 2019 , 11, 32115-32126	9.5	29
198	Response Enhancement of Self-Powered Visible-Blind UV Photodetectors by Nanostructured Heterointerface Engineering. <i>Advanced Functional Materials</i> , 2019 , 29, 1903981	15.6	20
197	Following excited states in molecular systems using density-based indexes: A dual emissive system as a test case. <i>Journal of Photochemistry and Photobiology A: Chemistry</i> , 2019 , 383, 111978	4.7	1
196	Double-Hybrid Functionals and Tailored Basis Set: Fullerene (C) Dimer and Isomers as Test Cases. Journal of Physical Chemistry A, 2019 , 123, 10040-10046	2.8	4
195	Copper-Catalysed Hydroamination of N-Allenylsulfonamides: The Key Role of Ancillary Coordinating Groups. <i>Synthesis</i> , 2019 , 51, 1225-1234	2.9	10
194	Aggregation Effects on Pigment Coatings: Pigment Red 179 as a Case Study. ACS Omega, 2019 , 4, 2031.	5329032	.312
193	Quantifying partial hole-particle distance at the excited state: A revised version of the DCT index. <i>Chemical Physics Letters</i> , 2019 , 714, 81-86	2.5	11
192	Does the gradient-regulated connection improve the description of correlated metal bond properties?. <i>International Journal of Quantum Chemistry</i> , 2019 , 119, e25831	2.1	
191	Ligand exchange on CdSe nanoplatelets for the solar light sensitization of TiO2 and ZnO nanorod arrays. <i>Journal of Photochemistry and Photobiology A: Chemistry</i> , 2019 , 368, 182-189	4.7	6
190	Using density based indexes to characterize excited states evolution. <i>Journal of Computational Chemistry</i> , 2019 , 40, 650-656	3.5	4
189	Communication: Evaluating non-empirical double hybrid functionals for spin-state energetics in transition-metal complexes. <i>Journal of Chemical Physics</i> , 2018 , 148, 041103	3.9	16
188	Taming Nickel-Catalyzed Suzuki-Miyaura Coupling: A Mechanistic Focus on Boron-to-Nickel Transmetalation. <i>ACS Catalysis</i> , 2018 , 8, 4812-4823	13.1	49
187	Using Density Based Indexes and Wave Function Methods for the Description of Excited States: Excited State Proton Transfer Reactions as a Test Case. <i>Journal of Physical Chemistry A</i> , 2018 , 122, 375-3	382 ⁸	10
186	How are the charge transfer descriptors affected by the quality of the underpinning electronic density?. <i>Journal of Computational Chemistry</i> , 2018 , 39, 735-742	3.5	17
185	Pyrimidyl-substituted anthracene fluorophores: Syntheses, absorption spectra, and photophysical properties. <i>Dyes and Pigments</i> , 2018 , 159, 619-636	4.6	3
184	Evidence for a Cooperative Mechanism Involving Two Palladium(0) Centers in the Oxidative Addition of Iodoarenes. <i>Chemistry - A European Journal</i> , 2018 , 24, 2192-2199	4.8	6
183	Functionalized Pd/ZnO Nanowires for Nanosensors. <i>Physica Status Solidi - Rapid Research Letters</i> , 2018 , 12, 1700321	2.5	27
182	Ultra-sensitive and selective hydrogen nanosensor with fast response at room temperature based on a single Pd/ZnO nanowire. <i>Sensors and Actuators B: Chemical</i> , 2018 , 254, 1259-1270	8.5	91

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254, 1600443

Emissive Azobenzenes Delivered on a Silver Coordination Polymer. Inorganic Chemistry, 2018, 57, 15009-4.502210 181 Combined Computational and Experimental Study of CdSeS/ZnS Nanoplatelets: Structural, 180 Vibrational, and Electronic Aspects of Core-Shell Interface Formation. Langmuir, **2018**, 34, 13828-13836 4 Revealing the Origins of Mechanically Induced Fluorescence Changes in Organic Molecular Crystals. 58 179 24 Advanced Materials, 2018, 30, e1800817 Theoretical approaches for predicting the color of rigid dyes in solution. Journal of Computational 178 3.5 12 Chemistry, 2017, 38, 998-1004 Understanding Aggregation-Induced Emission in Molecular Crystals: Insights from Theory. Journal 3.8 177 37 of Physical Chemistry C, 2017, 121, 5747-5752 Comparison of structural dynamics and coherence of d-d and MLCT light-induced spin state 176 9.4 35 trapping. Chemical Science, 2017, 8, 4978-4986 Comparing the performance of TD-DFT and SAC-CI methods in the description of excited states potential energy surfaces: An excited state proton transfer reaction as case study. Journal of 175 3.5 13 Computational Chemistry, 2017, 38, 1084-1092 Copper-Catalyzed Hydroamination of Allenes: from Mechanistic Understanding to Methodology 13.1 174 35 Development. ACS Catalysis, 2017, 7, 4253-4264 Effects of Substituents on Transport Properties of Molecular Materials for Organic Solar Cells: A 28 9.6 173 Theoretical Investigation. Chemistry of Materials, 2017, 29, 673-681 Supramolecular Anchoring of NCN-Pincer Palladium Complexes into a Barrel Protein Host: 172 Molecular-Docking and Reactivity Insights. European Journal of Inorganic Chemistry, 2017, 2017, 3622-3634 Multiconfiguration Pair-Density Functional Theory Predicts Spin-State Ordering in Iron Complexes with the Same Accuracy as Complete Active Space Second-Order Perturbation Theory at a 171 6.4 43 Significantly Reduced Computational Cost. Journal of Physical Chemistry Letters, 2017, 8, 2026-2030 Stereoselective access to trisubstituted fluorinated alkenyl thioethers. Catalysis Science and 170 9 5.5 Technology, **2017**, 7, 1921-1927 Copper-Catalyzed Hydroamination of N-Allenylazoles: Access to Amino-Substituted N-Vinylazoles. 169 5.6 15 Advanced Synthesis and Catalysis, 2017, 359, 4388-4392 Direct Spirocyclization from Keto-sulfonamides: An Approach to Azaspiro Compounds. Organic 168 6.2 21 Letters, 2017, 19, 5042-5045 Metrics for Molecular Electronic Excitations: A Comparison between Orbital- and Density-Based 167 2.8 24 Descriptors. Journal of Physical Chemistry A, 2017, 121, 7543-7549 Facile One-Pot Synthesis of Ruthenium(II) Quaterpyridine-Based Photosensitizers for 166 5.1 22 Photocatalyzed Hydrogen Production. Inorganic Chemistry, 2017, 56, 9515-9524 Charge transfer excitations in TDDFT: A ghost-hunter index. Journal of Computational Chemistry, 165 3.5 30 2017, 38, 2151-2156

Ex situ and in situ sensitized quantum dot solar cells. Physica Status Solidi (B): Basic Research, 2017,

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163	Enzymatic oxidation of ansa-ferrocifen leads to strong and selective thioredoxin reductase inhibition in vitro. <i>Journal of Inorganic Biochemistry</i> , 2016 , 165, 146-151	4.2	12
162	Modelling photophysical properties of metal-organic frameworks: a density functional theory based approach. <i>Physical Chemistry Chemical Physics</i> , 2016 , 18, 25176-25182	3.6	20
161	Unveiling the Reactivity of a Synthetic Mimic of the Oxygen Evolving Complex. <i>Journal of Physical Chemistry Letters</i> , 2016 , 7, 5015-5021	6.4	15
160	Anchoring groups for dyes in p-DSSC application: insights from DFT. <i>Journal of Molecular Modeling</i> , 2016 , 22, 289	2	16
159	Antagonistic Effect of Acetates in CN Bond Formation with In Situ Generated Diazonium Salts: A Combined Theoretical and Experimental Study. <i>European Journal of Organic Chemistry</i> , 2016 , 2016, 588	37 ³ 5 ² 896	₅ 7
158	Towards the modeling of quantum-dot sensitized solar cells: from structural and vibrational features to electron injection through lattice-mismatched interfaces. <i>Journal of Materials Chemistry A</i> , 2016 , 4, 13081-13092	13	4
157	Gradient-regulated connection-based correction for the PBE exchange: the PBEtrans model. <i>Molecular Physics</i> , 2016 , 114, 1059-1065	1.7	3
156	Modeling emission features of salicylidene aniline molecular crystals: A QM/QMQ pproach. <i>Journal of Computational Chemistry</i> , 2016 , 37, 861-70	3.5	21
155	Excited-State Proton Transfer and Intramolecular Charge Transfer in 1,3-Diketone Molecules. <i>ChemPhysChem</i> , 2016 , 17, 1530-8	3.2	10
154	Electrostatic Embedding To Model the Impact of Environment on Photophysical Properties of Molecular Crystals: A Self-Consistent Charge Adjustment Procedure. <i>Journal of Chemical Theory and Computation</i> , 2016 , 12, 3316-24	6.4	19
153	Optical properties of the dibenzothiazolylphenol molecular crystals through ONIOM calculations: the effect of the electrostatic embedding scheme. <i>Theoretical Chemistry Accounts</i> , 2016 , 135, 1	1.9	13
152	Multiple Roles of Isocyanides in Palladium-Catalyzed Imidoylative Couplings: A Mechanistic Study. <i>Chemistry - A European Journal</i> , 2016 , 22, 15491-15500	4.8	17
151	Nonempirical Double-Hybrid Functionals: An Effective Tool for Chemists. <i>Accounts of Chemical Research</i> , 2016 , 49, 1503-13	24.3	75
150	A qualitative model to identify non-radiative decay channels: the spiropyran as case study. <i>Theoretical Chemistry Accounts</i> , 2016 , 135, 1	1.9	11
149	Morphological and charge transport properties of amorphous and crystalline P3HT and PBTTT: insights from theory. <i>Physical Chemistry Chemical Physics</i> , 2015 , 17, 18742-50	3.6	39
148	Exploring excited states using Time Dependent Density Functional Theory and density-based indexes. <i>Coordination Chemistry Reviews</i> , 2015 , 304-305, 166-178	23.2	94
147	Red-light-driven photocatalytic hydrogen evolution using a ruthenium quaterpyridine complex. <i>Chemical Communications</i> , 2015 , 51, 9261-4	5.8	39
146	Theoretical Investigation of Hole Transporter Materials for Energy Devices. <i>Journal of Physical Chemistry C</i> , 2015 , 119, 23890-23898	3.8	37

(2014-2015)

145	Transition-Metal-Free Arylation of Enolizable Aryl Ketones and Mechanistic Evidence for a Radical Process. <i>Angewandte Chemie - International Edition</i> , 2015 , 54, 10587-91	16.4	101
144	Kinetics of Multielectron Transfers and Redox-Induced Structural Changes in N-Aryl-Expanded Pyridiniums: Establishing Their Unusual, Versatile Electrophoric Activity. <i>Journal of the American Chemical Society</i> , 2015 , 137, 11349-64	16.4	22
143	Non-parametrized functionals with empirical dispersion corrections: A happy match?. <i>Theoretical Chemistry Accounts</i> , 2015 , 134, 1	1.9	16
142	Describing excited state intramolecular proton transfer in dual emissive systems: a density functional theory based analysis. <i>Journal of Physical Chemistry B</i> , 2015 , 119, 2459-66	3.4	37
141	Computational Insights into Excited-State Proton-Transfer Reactions in Azo and Azomethine Dyes. <i>ChemPhysChem</i> , 2015 , 16, 3966-73	3.2	18
140	Low-Temperature Preparation of Ag-Doped ZnO Nanowire Arrays, DFT Study, and Application to Light-Emitting Diode. <i>ACS Applied Materials & Diodes amp; Interfaces</i> , 2015 , 7, 11871-80	9.5	53
139	Solvent tuned single molecule dual emission in protic solvents: effect of polarity and H-bonding. <i>Faraday Discussions</i> , 2015 , 185, 285-97	3.6	15
138	Solvent-tuned dual emission: a structural and electronic interplay highlighting a novel planar ICT (OPICT). <i>Physical Chemistry Chemical Physics</i> , 2015 , 17, 7639-42	3.6	14
137	A family of Ru(II) complexes built on a novel sexipyridine building block: synthesis, photophysical properties and the rare structural characterization of a triruthenium species. <i>Dalton Transactions</i> , 2015 , 44, 11551-61	4.3	11
136	Piano-stool d 6 -rhodium(III) complexes of chelating pyridine-based ligands and their papain bioconjugates for the catalysis of transfer hydrogenation of aryl ketones in aqueous medium. <i>Journal of Molecular Catalysis B: Enzymatic</i> , 2015 , 122, 314-322		8
135	Intrinsic and dynamical reaction pathways of an excited state proton transfer. <i>Journal of Physical Chemistry B</i> , 2015 , 119, 2650-7	3.4	30
134	The nature of vertical excited states of dyes containing metals for DSSC applications: insights from TD-DFT and density based indexes. <i>Physical Chemistry Chemical Physics</i> , 2014 , 16, 14435-44	3.6	51
133	Two-electron versus one-electron reduction of chalcogens by uranium(III): synthesis of a terminal U(V) persulfide complex. <i>Chemical Science</i> , 2014 , 5, 841-846	9.4	54
132	Toward a fast evaluation of g-tensor of Cu containing systems: A DFT parametrized approach. <i>Chemical Physics Letters</i> , 2014 , 614, 226-233	2.5	2
131	Ligand-stabilized CdSe nanoplatelet hybrid structures with tailored geometric and electronic properties. New insights from theory. <i>RSC Advances</i> , 2014 , 4, 55980-55989	3.7	16
130	Intermolecular proton shuttling in excited state proton transfer reactions: insights from theory. <i>Physical Chemistry Chemical Physics</i> , 2014 , 16, 8661-6	3.6	33
129	Non-radiative decay paths in rhodamines: new theoretical insights. <i>Physical Chemistry Chemical Physics</i> , 2014 , 16, 20681-8	3.6	35
128	Investigation of the bulk and surface properties of CdSe: insights from theory. <i>Physical Chemistry Chemical Physics</i> , 2014 , 16, 23251-9	3.6	17

127	Modeling of charge transfer processes to understand photophysical signatures: The case of Rhodamine 110. <i>Chemical Physics Letters</i> , 2014 , 610-611, 148-152	2.5	16
126	Benchmark Study on the Triplet Excited-State Geometries and Phosphorescence Energies of Heterocyclic Compounds: Comparison Between TD-PBE0 and SAC-CI. <i>Journal of Chemical Theory and Computation</i> , 2014 , 10, 3969-79	6.4	32
125	In silico assessment of the HPLCDV response coefficients. <i>Computational and Theoretical Chemistry</i> , 2014 , 1040-1041, 1-5	2	4
124	Copper-amyloid-Itomplex may catalyze peroxynitrite production in brain: evidence from molecular modeling. <i>Physical Chemistry Chemical Physics</i> , 2014 , 16, 10169-74	3.6	14
123	Simulations of UVIIisible spectra for analytical applications: phenothiazines as a case study. <i>Molecular Simulation</i> , 2014 , 40, 169-175	2	3
122	Computational Protocol for Modeling Thermochromic Molecular Crystals: Salicylidene Aniline As a Case Study. <i>Journal of Chemical Theory and Computation</i> , 2014 , 10, 5577-85	6.4	39
121	Chemically intuitive indices for charge-transfer excitation based on SAC-CI and TD-DFT calculations. Journal of Computational Chemistry, 2013 , 34, 2498-501	3.5	24
120	Understanding the redox properties of dinuclear ruthenium(II) complexes by a joint experimental and theoretical analysis. <i>Dalton Transactions</i> , 2013 , 42, 5281-91	4.3	16
119	Ethylene dimerization catalyzed by mixed phosphine-iminophosphorane nickel(II) complexes: a DFT investigation. <i>Journal of Molecular Modeling</i> , 2013 , 19, 2107-18	2	13
118	Evaluating push-pull dye efficiency using TD-DFT and charge transfer indices. <i>Physical Chemistry Chemical Physics</i> , 2013 , 15, 20210-9	3.6	53
117	Molecular dyads of ruthenium(II)- or osmium(II)-bis(terpyridine) chromophores and expanded pyridinium acceptors: equilibration between MLCT and charge-separated excited states. <i>Inorganic Chemistry</i> , 2013 , 52, 11944-55	5.1	25
116	Assessing the performances of some recently proposed density functionals for the description of organometallic structures. <i>Theoretical Chemistry Accounts</i> , 2013 , 132, 1	1.9	12
115	Discriminating role of bases in diketonate copper(I)-catalyzed C-O couplings: phenol versus diarylether. <i>Dalton Transactions</i> , 2013 , 42, 5348-54	4.3	32
114	Modeling Dye-Sensitized Solar Cells: From Theory to Experiment. <i>Journal of Physical Chemistry Letters</i> , 2013 , 4, 1044-50	6.4	93
113	Excited-State Geometries of Heteroaromatic Compounds: A Comparative TD-DFT and SAC-CI Study. Journal of Chemical Theory and Computation, 2013 , 9, 2368-79	6.4	54
112	Photophysical properties of NIR-emitting fluorescence probes: insights from TD-DFT. <i>Physical Chemistry Chemical Physics</i> , 2013 , 15, 10019-27	3.6	29
111	Spectral signature of a Ru(II, III, IV) complex: a combined experimental and theoretical investigation. <i>Dalton Transactions</i> , 2013 , 42, 7943-51	4.3	6
110	Confinement Effects on UV-Visible Absorption Spectra: ECarotene Inside Carbon Nanotube as a Test Case. <i>Journal of Physical Chemistry Letters</i> , 2013 , 4, 1239-43	6.4	22

(2012-2013)

109	Is There Still Room for Parameter Free Double Hybrids? Performances of PBE0-DH and B2PLYP over Extended Benchmark Sets. <i>Journal of Chemical Theory and Computation</i> , 2013 , 9, 3444-52	6.4	33	
108	Exploring the metric of excited state proton transfer reactions. <i>Journal of Physical Chemistry B</i> , 2013 , 117, 16165-73	3.4	50	
107	Probing the performances of HISS functionals for the description of excited states of molecular systems. <i>Theoretical Chemistry Accounts</i> , 2013 , 132, 1	1.9	1	
106	Ruthenium(II) complexes with new large-surface ligands based on electron-accepting expanded pyridiniums: insights from density functional theory. <i>Theoretical Chemistry Accounts</i> , 2012 , 131, 1	1.9	1	
105	Contribution to the Mechanism of Copper-Catalyzed CN and CD Bond Formation. <i>Organometallics</i> , 2012 , 31, 7694-7707	3.8	101	
104	Verdict: Time-Dependent Density Functional Theory "Not Guilty" of Large Errors for Cyanines. Journal of Chemical Theory and Computation, 2012 , 8, 1255-9	6.4	110	
103	Photoinduced electron transfer in Os(terpyridine)-biphenylene-(bi)pyridinium assemblies. <i>Inorganic Chemistry</i> , 2012 , 51, 5342-52	5.1	24	
102	Impact of Vibronic Couplings on Perceived Colors: Two Anthraquinones as a Working Example. <i>Journal of Physical Chemistry Letters</i> , 2012 , 3, 468-71	6.4	69	
101	Through-Space Charge Transfer in Rod-Like Molecules: Lessons from Theory. <i>Journal of Physical Chemistry C</i> , 2012 , 116, 11946-11955	3.8	193	
100	A DFT study of magnetic interactions in photoswitchable systems. <i>Chemical Physics Letters</i> , 2012 , 542, 13-18	2.5	11	
99	Revisiting the importance of dye binding mode in dye-sensitized solar cells: a periodic viewpoint. Journal of Materials Chemistry, 2012 , 22, 12205		22	
98	Amphiphilic acids as co-adsorbents of metal-free organic dyes for the efficient sensitization of nanostructured photoelectrode. <i>RSC Advances</i> , 2012 , 2, 11836	3.7	23	
97	Toward blue emission in ZnO based LED 2012 ,		2	
96	Tictoid expanded pyridiniums: assessing structural, electrochemical, electronic, and photophysical features. <i>Journal of Physical Chemistry A</i> , 2012 , 116, 7880-91	2.8	15	
95	Promising anchoring groups for ZnO-based hybrid materials: A periodic density functional theory investigation. <i>International Journal of Quantum Chemistry</i> , 2012 , 112, 2062-2071	2.1	9	
94	First-principles modeling of dye-sensitized solar cells: challenges and perspectives. <i>Accounts of Chemical Research</i> , 2012 , 45, 1268-77	24.3	182	
93	Influence of the Formation of the Halogen Bond ArX N on the Mechanism of Diketonate Ligated Copper-Catalyzed Amination of Aromatic Halides. <i>Organometallics</i> , 2012 , 31, 914-920	3.8	41	
92	Single-step versus stepwise two-electron reduction of polyarylpyridiniums: insights from the steric switching of redox potential compression. <i>Journal of the American Chemical Society</i> , 2012 , 134, 2691-70)5 ^{16.4}	24	

91	What is the "best" atomic charge model to describe through-space charge-transfer excitations?. <i>Physical Chemistry Chemical Physics</i> , 2012 , 14, 5383-8	3.6	242
90	Generalized gradient exchange functionals based on the gradient-regulated connection: a new member of the TCA family. <i>Theoretical Chemistry Accounts</i> , 2012 , 131, 1	1.9	12
89	TD-DFT Vibronic Couplings in Anthraquinones: From Basis Set and Functional Benchmarks to Applications for Industrial Dyes. <i>Journal of Chemical Theory and Computation</i> , 2011 , 7, 1882-92	6.4	96
88	High Aspect Ratio Ternary Zn1⊠CdxO Nanowires by Electrodeposition for Light-Emitting Diode Applications. <i>Journal of Physical Chemistry C</i> , 2011 , 115, 14548-14558	3.8	61
87	Phosphorescent binuclear iridium complexes based on terpyridine-carboxylate: an experimental and theoretical study. <i>Inorganic Chemistry</i> , 2011 , 50, 8197-206	5.1	42
86	A Qualitative Index of Spatial Extent in Charge-Transfer Excitations. <i>Journal of Chemical Theory and Computation</i> , 2011 , 7, 2498-506	6.4	657
85	Mechanistic Insights into C?N Coupling Catalyzed by 1,3-Diketonate-Ligated Copper: Unprecedented Activation of Aryl Iodide. <i>ChemCatChem</i> , 2011 , 3, 305-309	5.2	26
84	Theoretical procedure for optimizing dye-sensitized solar cells: from electronic structure to photovoltaic efficiency. <i>Journal of the American Chemical Society</i> , 2011 , 133, 8005-13	16.4	80
83	Insights into Working Principles of Ruthenium Polypyridyl Dye-Sensitized Solar Cells from First Principles Modeling. <i>Journal of Physical Chemistry C</i> , 2011 , 115, 4297-4306	3.8	64
82	Assessment of the B 97 family for excited-state calculations. <i>Theoretical Chemistry Accounts</i> , 2011 , 128, 127-136	1.9	115
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