Steven Robert Kirk

List of Publications by Year in descending order

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236925 345221 1,965 108 25 36 citations h-index g-index papers 110 110 110 1486 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Next generation quantum theory of atoms in molecules for the design of emitters exhibiting thermally activated delayed fluorescence with laser irradiation. Journal of Computational Chemistry, 2022, 43, 206-214.	3.3	4
2	Chirality–helicity of cumulenes: A nonâ€scalar charge density derived perspective. International Journal of Quantum Chemistry, 2022, 122, .	2.0	9
3	Hydroxyl-assisted selective epoxidation of perillyl alcohol with hydrogen peroxide by vanadium-substituted phosphotungstic acid hinged on imidazolyl activated carbon. New Journal of Chemistry, 2022, 46, 6636-6645.	2.8	5
4	Beyond energetic and scalar measures: Next generation <scp>quantum theory of atoms in molecules</scp> . Wiley Interdisciplinary Reviews: Computational Molecular Science, 2022, 12, .	14.6	10
5	Chiral and steric effects in ethane: A next generation QTAIM interpretation. Chemical Physics Letters, 2022, 800, 139669.	2.6	8
6	Mixed chiral and achiral character in substituted ethane: A next generation QTAIM perspective. Chemical Physics Letters, 2022, 803, 139762.	2.6	7
7	Fatigue and fatigue resistance in S 1 excited state diarylethenes in electric fields. International Journal of Quantum Chemistry, 2021, 121, e26527.	2.0	8
8	Bond-path-rigidity and bond-path-flexibility of the ground state and first excited state of fulvene. Chemical Physics Letters, 2021, 766, 138339.	2.6	5
9	Photochemical ring-opening reactions of oxirane with the Ehrenfest force topology. Chemical Physics Letters, 2021, 769, 138432.	2.6	6
10	Selective Catalytic Isomerization of $\hat{1}^2\hat{a}\in P$ inene Oxide to Perillyl Alcohol Enhanced by Protic Tetraimidazolium Nitrate. ChemistryOpen, 2021, 10, 477-485.	1.9	2
11	Control of chirality, bond flexing and anharmonicity in an electric field. International Journal of Quantum Chemistry, 2021, 121, e26793.	2.0	8
12	Chirality without Stereoisomers: Insight fromÂthe Helical Response of Bond Electrons. ChemPhysChem, 2021, 22, 1989-1995.	2.1	8
13	Understanding chemical coupling in cyclic versus compact water clusters with the Ehrenfest Force. Chemical Physics Letters, 2021, 781, 138983.	2.6	3
14	Bond flexing, twisting, anharmonicity and responsivity for the infraredâ€active modes of benzene. International Journal of Quantum Chemistry, 2021, 121, e26584.	2.0	5
15	Nextâ€generation quantum theory of atoms in molecules for the S ₁ /S ₀ conical intersections in dynamics trajectories of a lightâ€driven rotary molecular motor. International Journal of Quantum Chemistry, 2020, 120, e26062.	2.0	12
16	Direct cyclohexanone oxime synthesis <i>via</i> oxidation–oximization of cyclohexane with ammonium acetate. Chemical Communications, 2020, 56, 1436-1439.	4.1	11
17	Nextâ€generation QTAIM for scoring molecular wires in Eâ€fields for molecular electronic devices. Journal of Computational Chemistry, 2020, 41, 913-921.	3.3	12
18	A comparison of QTAIM and the stress tensor for Chirality-Helicity equivalence in S and R stereoisomers. Chemical Physics Letters, 2020, 738, 136907.	2.6	12

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19	3-D bond-paths of QTAIM and the stress tensor in neutral lithium clusters, Li _m (<i>m</i> =) Tj ETQq1 22, 864-877.	1 0.78431 2.8	14 rgBT /0 16
20	Flip rearrangement in the water pentamer: Analysis of electronic structure. International Journal of Quantum Chemistry, 2020, 120, e26124.	2.0	6
21	The role of the natural transition orbital density in the SOÂ→ÂS1 and SOÂ→ÂS2 transitions of fulvene with next generation QTAIM. Chemical Physics Letters, 2020, 751, 137556.	2.6	5
22	An explanation of the unusual strength of the <scp>hydrogen bond</scp> in small water clusters. International Journal of Quantum Chemistry, 2020, 120, e26361.	2.0	7
23	Intramolecular mode coupling of the isotopomers of water: a non-scalar charge density-derived perspective. Physical Chemistry Chemical Physics, 2020, 22, 2509-2520.	2.8	19
24	Stress tensor eigenvector following with nextâ€generation quantum theory of atoms in molecules. International Journal of Quantum Chemistry, 2019, 119, e25847.	2.0	23
25	Halogen and Hydrogen Bonding in Halogenabenzene/NH3 Complexes Compared Using Next-Generation QTAIM. Molecules, 2019, 24, 2875.	3.8	8
26	Stress Tensor Eigenvector Following with Next-Generation Quantum Theory of Atoms in Molecules: Excited State Photochemical Reaction Path from Benzene to Benzvalene. Journal of Physical Chemistry A, 2019, 123, 8254-8264.	2.5	10
27	Synergistic hydrogen atom transfer with the active role of solvent: Preferred one-step aerobic oxidation of cyclohexane to adipic acid by N-hydroxyphthalimide. Journal of Catalysis, 2019, 378, 256-269.	6.2	28
28	Imidazolyl activated carbon refluxed with ethanediamine as reusable heterogeneous catalysts for Michael addition. RSC Advances, 2019, 9, 185-191.	3.6	6
29	Next-generation quantum theory of atoms in molecules for the ground and excited state of DHCL. Chemical Physics Letters, 2019, 717, 91-98.	2.6	14
30	Explanation of the role of hydrogen bonding in the structural preferences of small molecule conformers. Chemical Physics Letters, 2019, 730, 206-212.	2.6	7
31	The directional bonding of [1.1.1]propellane with next generation QTAIM. Chemical Physics Letters, 2019, 730, 506-512.	2.6	15
32	Nano-Silica@PVC-Bonded <i>N</i> -Ethyl Sulfamic Acid as a Recyclable Solid Catalyst for the Hydroxyalkylation of Phenol with Formaldehyde to Bisphenol F. Bulletin of the Chemical Society of Japan, 2019, 92, 1394-1403.	3.2	1
33	Nextâ€generation quantum theory of atoms in molecules for the photochemical ringâ€opening reactions of oxirane. International Journal of Quantum Chemistry, 2019, 119, e25957.	2.0	12
34	A 3â€D bonding perspective of the factors influencing the relative stability of the S1/S0conical intersections of the pentaâ€2,4â€dieniminium cation (PSB3). International Journal of Quantum Chemistry, 2019, 119, e25903.	2.0	9
35	Next generation QTAIM for the design of quinone-based switches. Chemical Physics Letters, 2019, 722, 110-118.	2.6	9
36	Chirality–Helicity Equivalence in the S and R Stereoisomers: AÂTheoretical Insight. Journal of the American Chemical Society, 2019, 141, 5497-5503.	13.7	29

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37	The destabilization of hydrogen bonds in an external Eâ€field for improved switch performance. Journal of Computational Chemistry, 2019, 40, 1881-1891.	3.3	15
38	Nextâ€generation quantum theory of atoms in molecules for the ground and excited state of the ringâ€opening of cyclohexadiene. International Journal of Quantum Chemistry, 2019, 119, e25862.	2.0	18
39	Consequences of theory level choice evaluated with new tools from QTAIM and the stress tensor for a dipeptide conformer. Chemical Physics Letters, 2018, 696, 42-47.	2.6	1
40	Bamboo-derived porous biochar for efficient adsorption removal of dibenzothiophene from model fuel. Fuel, 2018, 211, 121-129.	6.4	73
41	The role of weak interactions in characterizing peptide folding preferences using a QTAIM interpretation of the Ramachandran plot (i•â€r). International Journal of Quantum Chemistry, 2018, 118, e25456.	2.0	7
42	Fatigue and photochromism $\langle i \rangle S \langle i \rangle \langle sub \rangle 1 \langle sub \rangle$ excited state reactivity of diarylethenes from QTAIM and the stress tensor. International Journal of Quantum Chemistry, 2018, 118, e25565.	2.0	23
43	The 3-D bonding morphology of the infra-red active normal modes of benzene. Chemical Physics Letters, 2018, 710, 31-38.	2.6	9
44	A vectorâ€based representation of the chemical bond for predicting competitive and noncompetitive torquoselectivity of thermal ringâ€opening reactions. International Journal of Quantum Chemistry, 2018, 118, e25707.	2.0	6
45	QTAIM and stress tensor bond-path framework sets for the ground and excited states of fulvene. Chemical Physics Letters, 2018, 713, 125-131.	2.6	19
46	Non-nuclear attractors in small charged lithium clusters, Li _m ^q (<i>m</i> =) Tj ETQq0 (Physics, 2018, 20, 24695-24707.	0 0 rgBT /0 2.8	Overlock 10 Tf 24
47	Quinoneâ€based switches for candidate building blocks of molecular junctions with QTAIM and the stress tensor. International Journal of Quantum Chemistry, 2018, 118, e25676.	2.0	21
48	Nextâ€generation quantum theory of atoms in molecules for the ground and excited states of fulvene. International Journal of Quantum Chemistry, 2018, 118, e25768.	2.0	18
49	A vector-based representation of the chemical bond for the substituted torsion of biphenyl. Chemical Physics Letters, 2018, 702, 32-37.	2.6	6
50	A vectorâ€based representation of the chemical bond for the normal modes of benzene. International Journal of Quantum Chemistry, 2018, 118, e25698.	2.0	5
51	Preparation of Fe2O3 doped SBA-15 for vapor phase ortho-position C-alkylation of phenol with methanol. Catalysis Communications, 2017, 92, 90-94.	3.3	13
52	The normal modes of vibration of benzene from the trajectories of stress tensor eigenvector projection space. Chemical Physics Letters, 2017, 677, 156-161.	2.6	20
53	QTAIM and Stress Tensor Characterization of Intramolecular Interactions Along Dynamics Trajectories of a Light-Driven Rotary Molecular Motor. Journal of Physical Chemistry A, 2017, 121, 4778-4792.	2.5	17
54	Exploration of the forbidden regions of the Ramachandran plot (i•-i̇̀) with QTAIM. Physical Chemistry Chemical Physics, 2017, 19, 26423-26434.	2.8	13

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55	Insights into the all-metal [Sb3Au3Sb3]3â^ sandwich complex from a QTAIM and stress tensor analysis. Chemical Physics Letters, 2017, 685, 127-132.	2.6	8
56	Isomerization of the RPSB chromophore in the gas phase along the torsional pathways using QTAIM. Chemical Physics Letters, 2017, 685, 222-228.	2.6	3
57	Effective transformation of cellulose to 5-hydroxymethylfurfural catalyzed by fluorine anion-containing ionic liquid modified biochar sulfonic acids in water. Cellulose, 2017, 24, 95-106.	4.9	35
58	A stress tensor eigenvector projection space for the (H2O)5 potential energy surface. Chemical Physics Letters, 2017, 667, 25-31.	2.6	29
59	A QTAIM and stress tensor perspective of large-amplitude motions of the tetrasulfur tetranitride S ₄ N ₄ molecular graph. International Journal of Quantum Chemistry, 2016, 116, 1025-1039.	2.0	8
60	The substituent effects on the biphenyl H \hat{a} ⁻ H bonding interactions subjected to torsion. Chemical Physics Letters, 2016, 651, 251-256.	2.6	10
61	A QTAIM and stress tensor investigation of the torsion path of a light-driven fluorene molecular rotary motor. Journal of Computational Chemistry, 2016, 37, 2588-2596.	3.3	15
62	QTAIM and stress tensor interpretation of the (H ₂ O) ₅ potential energy surface. Journal of Computational Chemistry, 2016, 37, 2712-2721.	3.3	10
63	Distinguishing and quantifying the torquoselectivity in competitive ring-opening reactions using the stress tensor and QTAIM. Journal of Computational Chemistry, 2016, 37, 2722-2733.	3.3	38
64	A QTAIM exploration of the competition between hydrogen and halogen bonding in halogenated 1-methyluracil: Water systems. Chemical Physics Letters, 2016, 662, 67-72.	2.6	10
65	Highly efficient and recyclable alkylammonium hydrosulfate catalyst for formation of bisphenol F by condensation of phenol with formaldehyde. RSC Advances, 2016, 6, 92716-92722.	3.6	4
66	Ï€â€Extended "Earring―Porphyrins with Multiple Cavities and Nearâ€Infrared Absorption. Angewandte Chemie - International Edition, 2016, 55, 6438-6442.	13.8	47
67	11-cis retinal torsion: A QTAIM and stress tensor analysis of the S1 excited state. Chemical Physics Letters, 2016, 652, 112-116.	2.6	19
68	Visible light-triggered vanadium-substituted molybdophosphoric acids to catalyze liquid phase oxygenation of cyclohexane to KA oil by nitrous oxide. Applied Catalysis B: Environmental, 2016, 182, 392-404.	20.2	37
69	Azobenzeneâ€Bridged Porphyrin Nanorings: Syntheses, Structures, and Photophysical Properties. Chemistry - A European Journal, 2015, 21, 15328-15338.	3.3	20
70	Biphenyl: A stress tensor and vector-based perspective explored within the quantum theory of atoms in molecules. International Journal of Quantum Chemistry, 2015, 115, 1678-1690.	2.0	46
71	Vanadiumâ€Substituted Tungstophosphoric Acids as Efficient Catalysts for Visibleâ€Lightâ€Driven Oxygenation of Cyclohexane by Dioxygen. ChemCatChem, 2015, 7, 2637-2645.	3.7	26
72	Hybrid QTAIM and electrostatic potential-based quantum topology phase diagrams for water clusters. Physical Chemistry Chemical Physics, 2015, 17, 15258-15273.	2.8	9

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73	A non-nitric acid method of adipic acid synthesis: organic solvent- and promoter-free oxidation of cyclohexanone with oxygen over hollow-structured Mn/TS-1 catalysts. Green Chemistry, 2015, 17, 1884-1892.	9.0	36
74	Solvent-free selective oxidation of toluene by oxygen over MnOx/SBA-15 catalysts: Relationship between catalytic behavior and surface structure. Chemical Engineering Journal, 2015, 280, 737-747.	12.7	46
75	Quantum topological resolution of catalyst proficiency. International Journal of Quantum Chemistry, 2015, 115, 875-883.	2.0	3
76	Consideration of roles of commercial TiO2 pigments in aromatic polyurethane coating via the photodegradation of dimethyl toluene-2,4-dicarbamate in non-aqueous solution. Research on Chemical Intermediates, 2015, 41, 7785-7797.	2.7	8
77	Oxidation of cyclohexane to adipic acid catalyzed by Mn-doped titanosilicate with hollow structure. Catalysis Communications, 2015, 58, 46-52.	3.3	36
78	Biochar sulfonic acid immobilized chlorozincate ionic liquid: an efficiently biomimetic and reusable catalyst for hydrolysis of cellulose and bamboo under microwave irradiation. Cellulose, 2014, 21, 1227-1237.	4.9	36
79	Light-triggered oxy-chlorination of cyclohexane by metal chlorides. Applied Catalysis A: General, 2014, 469, 483-489.	4.3	21
80	A QTAIM perspective of the Si6Li6 potential energy surface using quantum topology phase diagrams. Chemical Physics Letters, 2014, 609, 117-122.	2.6	5
81	New non-metallic mesoporous SBA-15 catalyst with high selectivity for the gas-phase oxidation of cyclohexylamine to cyclohexanone oxime. Catalysis Communications, 2014, 56, 148-152.	3.3	10
82	Allylic oxidation of α-isophorone to keto-isophorone with molecular oxygen catalyzed by copper chloride in acetylacetone. Applied Catalysis A: General, 2014, 486, 193-200.	4.3	8
83	Quantum topology phase diagrams for the <i>cis</i> ê•and <i>trans</i> êisomers of the cyclic contryphanâ€6m peptide. International Journal of Quantum Chemistry, 2014, 114, 1697-1706.	2.0	10
84	The response of the electronic structure to electronic excitation and double bond torsion in fulvene: a combined QTAIM, stress tensor and MO perspective. Physical Chemistry Chemical Physics, 2014, 16, 7115-7126.	2.8	58
85	Chlorocuprate Ionic Liquid Functionalized Biochar Sulfonic Acid as an Efficiently Biomimetic Catalyst for Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Microwave Irradiation. Industrial & Direct Hydrolysis of Bamboo under Hydrolysis of Bambo	3.7	15
86	The Pt site reactivity of the molecular graphs of Au6Pt isomers. Chemical Physics Letters, 2013, 590, 41-45.	2.6	11
87	The Ehrenfest force topology: a physically intuitive approach for analyzing chemical interactions. Physical Chemistry Chemical Physics, 2013, 15, 17823.	2.8	38
88	A bond, ring and cage resolved Poincaré–Hopf relationship for isomerisation reaction pathways. Molecular Physics, 2013, 111, 3104-3116.	1.7	17
89	The <i>cis</i> -effect using the topology of the electronic charge density. Molecular Physics, 2013, 111, 793-805.	1.7	14
90	Origin of the cis -Effect: a Density Functional Theory Study of Doubly Substituted Ethylenes. Wuli Huaxue Xuebao/ Acta Physico - Chimica Sinica, 2013, 29, 43-54.	4.9	25

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91	Structures, energies and bonding in neutral and charged Li microclusters. Journal of Molecular Modeling, 2012, 18, 4171-4189.	1.8	35
92	Exploring hydrogen bond in the excited state leading toward intramolecular proton transfer: detailed analysis of the structure and charge density topology along the reaction path using QTAIM. Journal of Molecular Modeling, 2012, 18, 4225-4237.	1.8	26
93	Molecular dynamics simulations of the aggregation of nanocolloidal amorphous silica monomers and dimers. Procedia Engineering, 2011, 18, 188-193.	1.2	5
94	Spanning Set of Silica Cluster Isomer Topologies from QTAIM. Journal of Physical Chemistry A, 2011, 115, 12503-12511.	2.5	28
95	Electronic Stress as a Guiding Force for Chemical Bonding. Topics in Current Chemistry, 2011, 351, 103-124.	4.0	10
96	Pointing the way to the products? Comparison of the stress tensor and the second-derivative tensor of the electron density. Journal of Chemical Physics, 2011, 134, 234106.	3.0	52
97	Spanning QTAIM topology phase diagrams of water isomers W4, W5 and W6. Physical Chemistry Chemical Physics, 2011, 13, 11644.	2.8	48
98	The mechanics of charge-shift bonds: A perspective from the electronic stress tensor. Chemical Physics Letters, 2011, 510, 18-20.	2.6	31
99	Molecular dynamics simulation of nanocolloidal amorphous silica particles: Part III. Journal of Chemical Physics, 2009, 130, 134702.	3.0	11
100	The role of hydrogen bonding in nanocolloidal amorphous silica particles in electrolyte solutions. Journal of Colloid and Interface Science, 2009, 339, 351-361.	9.4	12
101	Bond metallicity of materials from real space charge density distributions. Chemical Physics Letters, 2009, 471, 174-177.	2.6	51
102	Tiling for Performance Tuning on Different Models of GPUs. , 2009, , .		12
103	Molecular dynamics simulation of nanocolloidal amorphous silica particles: Part II. Journal of Chemical Physics, 2008, 128, 164711.	3.0	20
104	Molecular dynamics simulation of nanocolloidal amorphous silica particles: Part I. Journal of Chemical Physics, 2007, 127, 224711.	3.0	29
105	Software architecture graphs as complex networks: A novel partitioning scheme to measure stability and evolution. Information Sciences, 2007, 177, 2587-2601.	6.9	85
106	Information theory-based software metrics and obfuscation. Journal of Systems and Software, 2004, 72, 179-186.	4.5	26
107	Dependence of the normal modes on the electronic structure of various phases of ice as calculated by ab initio methods. Canadian Journal of Physics, 2003, 81, 225-231.	1.1	42
108	The chirality of isotopomers of glycine compared using nextâ€generation <scp>QTAIM</scp> . International Journal of Quantum Chemistry, 0, , .	2.0	3