List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Anion–π Interactions: Do They Exist?. Angewandte Chemie - International Edition, 2002, 41, 3389-3392.	13.8	690
2	Cooperativity in multiple unusual weak bonds. Theoretical Chemistry Accounts, 2010, 126, 1-14.	1.4	254
3	Halogen bonding versuschalcogen and pnicogen bonding: a combined Cambridge structural database and theoretical study. CrystEngComm, 2013, 15, 3137-3144.	2.6	206
4	A Topological Analysis of the Electron Density in Anion-Ï€ Interactions. ChemPhysChem, 2003, 4, 1344-1348.	2.1	190
5	A thorough anion–΀ interaction study in biomolecules: on the importance of cooperativity effects. Chemical Science, 2016, 7, 1038-1050.	7.4	188
6	Structure and Binding Energy of Anionâ^'Ï€ and Cationâ^'Ï€ Complexes:Â A Comparison of MP2, RI-MP2, DFT, and DF-DFT Methods. Journal of Physical Chemistry A, 2005, 109, 4632-4637.	2.5	186
7	Counterintuitive interaction of anions with benzene derivatives. Chemical Physics Letters, 2002, 359, 486-492.	2.6	178
8	Anion–π Interactions: Do They Exist?. Angewandte Chemie, 2002, 114, 3539-3542.	2.0	176
9	Cationâ~ï̃€ versus Anionâ~ï̃€ Interactions:Â Energetic, Charge Transfer, and Aromatic Aspects. Journal of Physical Chemistry A, 2004, 108, 9423-9427.	2.5	171
10	Anion-ï€ Interactions in Cyanuric Acids: A Combined Crystallographic and Computational Study. Chemistry - A European Journal, 2005, 11, 6560-6567.	3.3	167
11	Relevant Anion–π Interactions in Biological Systems: The Case of Urate Oxidase. Angewandte Chemie - International Edition, 2011, 50, 415-418.	13.8	164
12	Structural, Physicochemical, and Reactivity Properties of an All-Inorganic, Highly Active Tetraruthenium Homogeneous Catalyst for Water Oxidation. Journal of the American Chemical Society, 2009, 131, 17360-17370.	13.7	162
13	Cation‑ïl̃€ and anion‑'ïl̃€ interactions. Wiley Interdisciplinary Reviews: Computational Molecular Science, 2011, 1, 440-459.	14.6	156
14	MP2 study of cooperative effects between cation–π, anion–π and π–π interactions. New Journal of Chemistry, 2007, 31, 556-560.	2.8	151
15	Interplay Between Cation-ï€, Anion-ï€ and ï€-ï€ Interactions. ChemPhysChem, 2006, 7, 2487-2491.	2.1	145
16	Anion–π interactions: must the aromatic ring be electron deficient?. New Journal of Chemistry, 2003, 27, 211-214.	2.8	116
17	Pnicogen–π complexes: theoretical study and biological implications. Physical Chemistry Chemical Physics, 2012, 14, 14061.	2.8	113
18	Anionâ~ï€ Interactions in Bisadenine Derivatives:  A Combined Crystallographic and Theoretical Study. Inorganic Chemistry, 2007, 46, 10724-10735.	4.0	104

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19	Synthetic Prodiginine Obatoclax (GX15â€070) and Related Analogues: Anion Binding, Transmembrane Transport, and Cytotoxicity Properties. Chemistry - A European Journal, 2011, 17, 14074-14083.	3.3	102
20	Approximate Additivity of Anionâ^'Ï€ Interactions:  An Ab Initio Study on Anionâ^'Ï€, Anionâ^'Ï€2 and Anionâ^ Complexes. Journal of Physical Chemistry A, 2005, 109, 9341-9345.	Ϊ€3. <sub>5</sub>	101
21	Thermodynamic Characterization of Halideâ~ḯ€ Interactions in Solution Using "Two-Wall―Aryl Extended Calix[4]pyrroles as Model System. Journal of the American Chemical Society, 2014, 136, 3208-3218.	13.7	96
22	s-Tetrazine as a new binding unit in molecular recognition of anions. Chemical Physics Letters, 2003, 370, 7-13.	2.6	95
23	Metalâ^'Peroxo versus Metalâ^'Oxo Oxidants in Non-Heme Iron-Catalyzed Olefin Oxidations:Â Computational and Experimental Studies on the Effect of Water. Journal of the American Chemical Society, 2005, 127, 6548-6549.	13.7	94
24	Substituent effects in halogen bonding complexes between aromatic donors and acceptors: a comprehensive ab initio study. Physical Chemistry Chemical Physics, 2011, 13, 20371.	2.8	92
25	Interplay between cation-ï€ and hydrogen bonding interactions. Chemical Physics Letters, 2008, 456, 257-261.	2.6	82
26	Quantification of Aromaticity in Oxocarbons: The Problem of the Fictitious "Nonaromatic―Reference System. Chemistry - A European Journal, 2002, 8, 433-438.	3.3	80
27	Very Longâ€Range Effects: Cooperativity between Anion‑'Ï€ and Hydrogenâ€Bonding Interactions. ChemPhysChem, 2009, 10, 2256-2264.	2.1	80
28	Interplay between anionâ€i€ and hydrogen bonding interactions. Journal of Computational Chemistry, 2009, 30, 75-82.	3.3	79
29	Theoretical Study on Cooperativity Effects between Anion–π and Halogenâ€Bonding Interactions. ChemPhysChem, 2011, 12, 2742-2750.	2.1	79
30	On the directionality of anion–π interactions. Physical Chemistry Chemical Physics, 2011, 13, 5696.	2.8	78
31	A theoretical study of aromaticity in squaramide and oxocarbons. Tetrahedron Letters, 2000, 41, 2001-2005.	1.4	74
32	Dual Binding Mode ofs-Triazine to Anions and Cations. Organic Letters, 2003, 5, 2227-2229.	4.6	74
33	Cation-Ï€ versus anion-Ï€ interactions: a comparative ab initio study based on energetic, electron charge density and aromatic features. Chemical Physics Letters, 2004, 392, 85-89.	2.6	74
34	A Theoretical ab initio Study of the Capacity of Several Binding Units for the Molecular Recognition of Anions. European Journal of Organic Chemistry, 2005, 2005, 179-183.	2.4	74
35	Coordination Complexes Exhibiting Anion···π Interactions: Synthesis, Structure, and Theoretical Studies. Inorganic Chemistry, 2008, 47, 5873-5881.	4.0	72
36	Ab Initio Study of [n.n]Paracyclophane (n= 2, 3) Complexes with Cations:Â Unprecedented Through-Space Substituent Effects. Journal of Physical Chemistry A, 2006, 110, 5144-5148.	2.5	71

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37	Rational Design, Synthesis, and Application of a New Receptor for the Molecular Recognition of Tricarboxylate Salts in Aqueous Media. Journal of Organic Chemistry, 2006, 71, 7185-7195.	3.2	66
38	Squaramide as a binding unit in molecular recognition. Chemical Physics Letters, 2000, 326, 247-254.	2.6	62
39	MP2 Study of synergistic effects between X–H/π (X = C,N,O) and π–π interactions. Theoretical Chemistry Accounts, 2008, 120, 385-393.	1.4	62
40	Experimental and theoretical study of uracil derivatives: the crucial role of weak fluorine–fluorine noncovalent interactions. CrystEngComm, 2010, 12, 3758.	2.6	60
41	A theoretical study of aromaticity in squaramide complexes with anions. Chemical Physics Letters, 2002, 351, 115-120.	2.6	57
42	Experimental and computational study of the interplay between C–H/π and anion–π interactions. Dalton Transactions, 2010, 39, 794-806.	3.3	57
43	Lithium diffusion in single-walled carbon nanotubes: a theoretical study. Chemical Physics Letters, 2003, 374, 548-555.	2.6	55
44	High‣evel Ab Initio Study of Anion–Ĩ€ Interactions in Pyridine and Pyrazine Rings Coordinated to Ag <sup>I</sup> . ChemPhysChem, 2008, 9, 397-399.	2.1	53
45	Simultaneous Interaction of Tetrafluoroethene with Anions and Hydrogen-Bond Donors: A Cooperativity Study. Journal of Chemical Theory and Computation, 2009, 5, 1186-1194.	5.3	52
46	Energetic vs Synergetic Stability: A Theoretical Study. Journal of Physical Chemistry A, 2009, 113, 3266-3273.	2.5	52
47	Anionπ Interactions in Flavoproteins. Chemistry - an Asian Journal, 2011, 6, 2316-2318.	3.3	52
48	MP2 Study of Cationâ^'(Ï€)nâ^'Ï€ Interactions (n= 1â^'4). Journal of Physical Chemistry A, 2006, 110, 9307-9309.	2.5	49
49	Crystallographic and Theoretical Evidence of Anion–π and Hydrogenâ€Bonding Interactions in a Squaramide–Nitrate Salt. European Journal of Organic Chemistry, 2008, 2008, 1864-1868.	2.4	49
50	A Theoretical Study of Anion–Ĩ€ Interactions in Seven-Membered Rings. ChemPhysChem, 2007, 8, 1182-1187.	2.1	47
51	Cation–cation and anion–anion complexes stabilized by halogen bonds. Physical Chemistry Chemical Physics, 2016, 18, 27939-27950.	2.8	45
52	Counterintuitive Substituent Effect of the Ethynyl Group in Ionâ^'Ï€ Interactions. Journal of Physical Chemistry A, 2009, 113, 10367-10375.	2.5	43
53	Cation-π vs anion-π interactions: a complete π-orbital analysis. Chemical Physics Letters, 2004, 399, 220-225.	2.6	42
54	The Role of the Central Atom in Structure and Reactivity of Polyoxometalates with Adjacent d-Electron Metal Sites. Computational and Experimental Studies of Î <sup>3</sup> -[(Xn+O4)RullI2(OH)2(MFM)10O32](8-n)-for MFM= Mo and W, and X = AlIII, SiIV, PV, and SVI. Journal of Physical Chemistry B, 2006, 110, 170-173.	2.6	42

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55	Interplay between cationâ€"ï€ and hydrogen bonding interactions: Are non-additivity effects additive?. Chemical Physics Letters, 2009, 479, 316-320.	2.6	42
56	Anion-ï€ interactions in five-membered rings: a combined crystallographic and ab initio study. Chemical Physics Letters, 2003, 382, 534-540.	2.6	41
57	Unexpected chalcogen bonds in tetravalent sulfur compounds. Physical Chemistry Chemical Physics, 2019, 21, 11313-11319.	2.8	41
58	2-Aminopyrimidine Derivatives Exhibiting Anion-Ï€ Interactions: A Combined Crystallographic and Theoretical Study. Crystal Growth and Design, 2009, 9, 2363-2376.	3.0	39
59	Computational Studies of the Geometry and Electronic Structure of an All-Inorganic and Homogeneous Tetra-Ru-Polyoxotungstate Catalyst for Water Oxidation and Its Four Subsequent One-Electron Oxidized Forms. Journal of Physical Chemistry A, 2010, 114, 535-542.	2.5	39
60	Lone pair–π vs π–π interactions in 5-fluoro-1-hexyluracil and 1-hexyluracil: a combined crystallographic and computational study. CrystEngComm, 2010, 12, 362-365.	2.6	39
61	Anionâ^'Ï€ Interactions in Four-Membered Rings. Organic Letters, 2009, 11, 1987-1990.	4.6	38
62	Is the Use of Diffuse Functions Essential for the Properly Description of Noncovalent Interactions Involving Anions?. Journal of Physical Chemistry A, 2013, 117, 2651-2655.	2.5	38
63	Weak Câ^'H/Ï€ Interaction Participates in the Diastereoselectivity of a Hostâ 'Guest Complex in the Presence of Six Strong Hydrogen Bonds. Organic Letters, 2003, 5, 1135-1138.	4.6	37
64	Interaction of positively and negatively charged aromatic hydrocarbons with benzene and triphenylene: Towards a model of pure organic insulators. Chemical Physics Letters, 2008, 460, 406-410.	2.6	36
65	Preparation, Solid-State Characterization, and Computational Study of a Crown Ether Attached to a Squaramide. Organic Letters, 2005, 7, 1437-1440.	4.6	35
66	Does Dinitrogen Hydrogenation Follow Different Mechanisms for [(η5-C5Me4H)2Zr]2(μ2,η2,η2-N2) and {[PhP(CH2SiMe2NSiMe2CH2)PPh]Zr}2(μ2,η2,η2-N2) Complexes? A Computational Study. Journal of the American Chemical Society, 2006, 128, 11391-11403.	13.7	35
67	Longâ€Range Effects in Anion–π Interactions: Their Crucial Role in the Inhibition Mechanism of <i>Mycobacterium Tuberculosis</i> Malate Synthase. Chemistry - A European Journal, 2014, 20, 6985-6990.	3.3	35
68	Ab initio investigations of lithium diffusion in single-walled carbon nanotubes. Chemical Physics, 2004, 297, 85-91.	1.9	34
69	Hydrogen Bond versus Halogen Bond in Cation–Cation Complexes: Effect of the Solvent. ChemPhysChem, 2017, 18, 3462-3468.	2.1	34
70	A novel fluoride selective optical chemosensor based on internal charge transfer signaling. Tetrahedron Letters, 2010, 51, 596-599.	1.4	33
71	Cations brought together by hydrogen bonds: the protonated pyridine–boronic acid dimer explained. Physical Chemistry Chemical Physics, 2019, 21, 5796-5802.	2.8	33
72	Feasibility of Single-Walled Carbon Nanotubes as Materials for CO <sub>2</sub> Adsorption: A DFT Study. Journal of Physical Chemistry C, 2012, 116, 21083-21092.	3.1	32

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73	Theoretical ab initio study of the interplay between hydrogen bonding, cation–π and π–π interactions. Theoretical Chemistry Accounts, 2009, 122, 325-332.	1.4	31
74	On the Importance of Anion–π Interactions in the Mechanism of Sulfide:Quinone Oxidoreductase. Chemistry - an Asian Journal, 2013, 8, 2708-2713.	3.3	31
75	Predicting Experimental Complexation-Induced Changes in1H NMR Chemical Shift for Complexes between Zinc-Porphyrins and Amines Using the ab Initio/GIAO-HF Methodology. Organic Letters, 2002, 4, 399-401.	4.6	30
76	Theoretical Studies of the Complex [(BPMEN)Fe(II)(NCCH3)2]2+, Precursor of Non-Heme Iron Catalysts for Olefin Epoxidation and Cis-Dihydroxylation. Inorganic Chemistry, 2003, 42, 8449-8455.	4.0	30
77	Dual Cation and Anion Acceptor Molecules. The Case of the (η6-C6H6)(η6C6F6)Cr(0) Complex. Journal of Physical Chemistry A, 2007, 111, 3137-3142.	2.5	29
78	A Combined Experimental and Theoretical Study of Anion–π Interactions in Bis(pyrÂɨmidine) Salts. European Journal of Organic Chemistry, 2007, 2007, 5821-5825.	2.4	29
79	MP2 study of anion–i̇̀€ complexes of trifluoro-s-triazine with tetrahedral and octahedral anions. Chemical Physics Letters, 2007, 438, 104-108.	2.6	29
80	New [2 × 2] Copper(I) Grids as Anion Receptors. Effect of Ligand Functionalization on the Ability to Host Counteranions by Hydrogen Bonds. Inorganic Chemistry, 2010, 49, 8828-8847.	4.0	28
81	Substituent effects in cationâ€"ï€ interactions revisited: a general approach based on intrinsic properties of the arenes. Physical Chemistry Chemical Physics, 2014, 16, 1322-1326.	2.8	28
82	Sigma-hole carbon-bonding interactions in carbon–carbon double bonds: an unnoticed contact. Physical Chemistry Chemical Physics, 2017, 19, 15530-15540.	2.8	28
83	Estimating ring strain energies in small carbocycles by means of the Bader's theory of â€~atoms-in-molecules'. Chemical Physics Letters, 2012, 536, 165-169.	2.6	27
84	Investigating Polyoxometalate–Protein Interactions at Chemically Distinct Binding Sites. Journal of Physical Chemistry B, 2018, 122, 7219-7232.	2.6	27
85	RI-MP2 and MPWB1K Study of π–Anionâ~π′ Complexes: MPWB1K Performance and Some Additivity Aspec Journal of Chemical Theory and Computation, 2011, 7, 3012-3018.	ts. 5.3	26
86	Ab initio investigations of lithium insertion in boron and nitrogen-doped single-walled carbon nanotubes. Chemical Physics Letters, 2005, 411, 256-261.	2.6	24
87	Synthesis, X-ray structure analysis and computational studies of novel bis(thiocarbamoyl) disulfides with non-covalent Sâ< N and Sâ< S interactions. Chemical Physics Letters, 2006, 422, 234-239.	2.6	24
88	Interplay between Edge-to-Face Aromatic and Hydrogen-Bonding Interactions. Journal of Physical Chemistry A, 2008, 112, 6017-6022.	2.5	24
89	Unexpected Nonadditivity Effects in Anionâ°'Ï€ Complexes. Journal of Physical Chemistry A, 2011, 115, 7849-7857.	2.5	23
90	A methodological analysis for the assessment of non-covalent π interactions. Chemical Physics Letters, 2011, 508, 144-148.	2.6	23

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91	Substituent Effects in Ionâ^"í€ Interactions: Fine-Tuning via the Ethynyl Group. Journal of Physical Chemistry A, 2010, 114, 1926-1930.	2.5	22
92	Theoretical and Crystallographic Study of the Dual σ/π Anion Binding Affinity of Quinolizinylium Cation. Journal of Chemical Theory and Computation, 2008, 4, 1981-1989.	5.3	21
93	Theoretical and crystallographic study of edge-to-face aromatic interactions between pyridine moieties and benzene. Chemical Physics Letters, 2009, 468, 280-285.	2.6	21
94	Induced-Polarization Energy Map:  A Helpful Tool for Predicting Geometric Features of Anion-π Complexes. Journal of Chemical Theory and Computation, 2007, 3, 2098-2107.	5.3	20
95	Erroneous behaviour of the widely used MP2(full)/aug-cc-pVXZ (X=D,T) level of theory for evaluating the BSSE in ion–Ĩ€ complexes. Chemical Physics Letters, 2010, 489, 254-258.	2.6	20
96	Tuning of the anion–݀ interaction. Theoretical Chemistry Accounts, 2012, 131, 1.	1.4	20
97	Structural and energetic features of single-walled carbon nanotube junctions: a theoretical ab initio study. Chemical Physics, 2004, 303, 265-270.	1.9	19
98	A Combined Experimental and Theoretical Study of Anion–΀ Interactions in <i>N</i> <sup>6</sup> ― and <i>N</i> <sup>9</sup> â€Đecyladenine Salts. European Journal of Organic Chemistry, 2010, 2010, 5171-5180.	2.4	19
99	Synthetic Tripodal Squaramidoâ€Based Receptors for the Complexation of Antineoplastic Folates in Water. European Journal of Organic Chemistry, 2011, 2011, 6187-6194.	2.4	19
100	Self-assembly hexanuclear metallacontainer hosting halogenated guest species and sustaining structure of 3D coordination framework. Chemical Communications, 2011, 47, 1764-1766.	4.1	18
101	Highly efficient coordination of Hg <sup>2+</sup> and Pb <sup>2+</sup> metals in water with squaramide-coated Fe <sub>3</sub> O <sub>4</sub> nanoparticles. Journal of Materials Chemistry A, 2014, 2, 8796-8803.	10.3	18
102	Theoretical ab initio study of anion–π interactions in inorganic rings. Chemical Physics Letters, 2012, 530, 145-150.	2.6	17
103	Anion–π interactions in [S4N3]+ rings. New Journal of Chemistry, 2013, 37, 2636.	2.8	17
104	Hydrolysis of chemically distinct sites of human serum albumin by polyoxometalate: A hybrid QM/MM (ONIOM) study. Journal of Computational Chemistry, 2019, 40, 51-61.	3.3	17
105	Metastable Dianions and Dications. ChemPhysChem, 2020, 21, 1597-1607.	2.1	16
106	The resonance model in amides: a combined crystallographic and ab initio investigation. New Journal of Chemistry, 2001, 25, 259-261.	2.8	15
107	Conformational Analysis of a Model Synthetic Prodiginine. Journal of Organic Chemistry, 2012, 77, 6538-6544.	3.2	15
108	Ab initio calculations on zinc porphyrins complexed to amines: geometrical details and NMR chemical shifts. Computational and Theoretical Chemistry, 2000, 531, 381-386.	1.5	14

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109	Counterintuitive affinity of [2.2]paracyclophane to cations. Chemical Physics Letters, 2005, 408, 59-64.	2.6	14
110	Can lone pair-Ï€ and cation-Ï€ interactions coexist? A theoretical study. Open Chemistry, 2011, 9, 25-34.	1.9	14
111	Adsorption and Quantification of Volatile Organic Compounds (VOCs) by using Hybrid Magnetic Nanoparticles. Chemistry - A European Journal, 2018, 24, 12820-12826.	3.3	14
112	On the importance of the inclusion of the basis set superposition error counterpoise correction during optimization of ion-Ï€ complexes. Chemical Physics Letters, 2008, 455, 325-330.	2.6	13
113	Computational insights to the mechanism of alkene epoxidation by manganese-based catalysts in the presence of bicarbonate. Computational and Theoretical Chemistry, 2009, 903, 115-122.	1.5	13
114	Cooperativity effects between non-covalent interactions: Are they important for Z-DNA stability?. Chemical Physics Letters, 2010, 485, 221-225.	2.6	13
115	Radical cation (CE™+–Ĩ€) and radical anion (AE™â^'–Ĩ€) interactions with aromatic rings: energetic, orbitalic and spin density considerations. Physical Chemistry Chemical Physics, 2011, 13, 16698.	2.8	13
116	OPLS all-atom force field for squaramides and squaric acid. Chemical Physics Letters, 2001, 350, 331-338.	2.6	12
117	A theoretical ab initio study of [n.n]paracyclophane complexes with cations. Chemical Physics Letters, 2006, 417, 371-377.	2.6	12
118	MP2 Study of the Dual σ/Ï€â~'Anion-Binding Affinity of Fluorinated Phthallic Acid Anhydrides. Journal of Physical Chemistry A, 2008, 112, 1622-1626.	2.5	12
119	Interplay between ion–݀ and Ar/݀ Van der Waals interactions. Computational and Theoretical Chemistry, 2012, 998, 51-56.	2.5	12
120	Hydrogen Bond versus Halogen Bond in HXOn (X = F, Cl, Br, and I) Complexes with Lewis Bases. Inorganics, 2019, 7, 9.	2.7	12
121	Anion–΀ Interactions Involving [MX <sub><i>n</i></sub> ] <sup><i>m</i>â^`</sup> Anions: A Comprehensive Theoretical Study. ChemPhysChem, 2013, 14, 145-154.	2.1	11
122	Substituent Effects in Multivalent Halogen Bonding Complexes: A Combined Theoretical and Crystallographic Study. Molecules, 2018, 23, 18.	3.8	11
123	A density functional study of geometry and electronic structures of [(SiO4)(MIII)2(OH)2W10O32]4â^', M=Mo, Ru and Rh. Journal of Molecular Catalysis A, 2007, 262, 227-235.	4.8	10
124	New Chlorido(dimethyl sulfoxide)iridium(III) Complexes with N6-Substituted Adenines - Kinetic N(7) versus Thermodynamic N(9) Coordinated Adenine Isomers. European Journal of Inorganic Chemistry, 2010, 2010, 5617-5628.	2.0	10
125	Anion Recognition by Pyrylium Cations and Thio-, Seleno- and Telluro- Analogues: A Combined Theoretical and Cambridge Structural Database Study. Molecules, 2015, 20, 11632-11659.	3.8	10
126	Reconciling Experiment and Theory in the Use of Aryl-Extended Calix[4]pyrrole Receptors for the Experimental Quantification of Chloride–i€ Interactions in Solution. International Journal of Molecular Sciences, 2015, 16, 8934-8948.	4.1	10

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127	Influence of the aromatic surface on the capacity of adsorption of VOCs by magnetite supported organic–inorganic hybrids. RSC Advances, 2019, 9, 24184-24191.	3.6	10
128	Squaramido-based receptors: applicability of molecular interaction potential to molecular recognition of polyalkylammonium compounds. Theoretical Chemistry Accounts, 2000, 104, 50-66.	1.4	9
129	A topological analysis of charge density in complexes between derivatives of squaric acid and ammonium cation. Chemical Physics Letters, 2001, 339, 369-374.	2.6	9
130	Applicability of the 1H NMR chemical shifts computed by the ab initio/GIAO-HF methodology to the study of geometrical features of Zn-porphyrin dimers. Tetrahedron Letters, 2004, 45, 9387-9391.	1.4	8
131	Weak interactions within nitryl halide heterodimers. New Journal of Chemistry, 2016, 40, 9060-9072.	2.8	8
132	Theoretical ab initio study of substituted benzene trimer: Interplay between hydrogen bonding and π–π interactions. Computational and Theoretical Chemistry, 2011, 975, 106-110.	2.5	7
133	Theoretical ab initio study of lone pair and anion–π interactions in fluorinated tropolones. Computational and Theoretical Chemistry, 2012, 998, 20-25.	2.5	7
134	Predicting experimental complexation-induced changes in NMR chemical shift for complexes between metalloporphyrins and ligands using the Ab initio/GIAO-HF methodology. Chemical Physics Letters, 2002, 360, 72-78.	2.6	6
135	Theoretical Study of the Structure and Properties of [(η5-C5Me4H)2Zr]2(μ2,η2,η2-N2). Journal of Chemical Theory and Computation, 2006, 2, 336-341.	5.3	6
136	Quadrupole moment versus Molecular Electrostatic Potential: Strange behavior of ethynyl-substituted benzenes. Chemical Physics Letters, 2013, 567, 60-65.	2.6	5
137	Affinity of ferrocene and (1,1′)(3,3′)[3,3]ferrocenophane to cations. Chemical Physics Letters, 2006, 424, 204-208.	2.6	4
138	New 1,8-naphthyridine-based probes for the selective fluorescence signalling of toxic cadmium: synthesis, photophysical studies and molecular modelling. Supramolecular Chemistry, 2010, 22, 524-531.	1.2	4
139	Anion–π Interactions: Do They Exist?. Angewandte Chemie - International Edition, 2004, 43, 141-141.	13.8	3
140	The Role of the Ethynyl Substituent on the π–π Stacking Affinity of Benzene: A Theoretical Study. ChemPhysChem, 2011, 12, 283-288.	2.1	3
141	Internal rotation in squaramide and related compounds. A theoretical ab initio study. Theoretical Chemistry Accounts, 2002, 108, 157-167.	1.4	2
142	Kinetics and mechanism of the oxidation of hydroxylamine by a {Mn3O4}4+ core in aqueous acidic media. Dalton Transactions, 2011, 40, 9571.	3.3	2
143	Molecular Interaction Potential with Polarization (MIPp) Study of the Interplay Between Ion-Ï $\in$ and Hydrogen Bonding Interactions. The Open Chemical Physics Journal, 2008, 1, 36-41.	0.7	2
144	Synthesis and structure of <i>cis</i> -[RuCl(bpzm)(î <sup>e</sup> <sup>1</sup> - <i>P</i> -dpim)(î <sup>e</sup> <sup>2</sup> - <i>P,N</i> -dpim)]Cl·(CHCl <sub>3<td>ub&gt;)<sub 1<b>.2</b></sub </td><td>&gt;5<sub>4</sub>/sub&gt;.</td></sub>	ub>) <sub 1<b>.2</b></sub 	>5 <sub>4</sub> /sub>.

Stability of [Cl(HCCl<sub>3</sub><i>n</i></sub>]<sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<sub>3</sub>]<su

#	Article	IF	CITATIONS
145	A Squaramide-Based Citrate Receptor. Synfacts, 2006, 2006, 1225-1225.	0.0	0
146	Frontispiece: Adsorption and Quantification of Volatile Organic Compounds (VOCs) by using Hybrid Magnetic Nanoparticles. Chemistry - A European Journal, 2018, 24, .	3.3	0
147	Anionπ, lone pairπ, and FF interactions in nucleobase derivatives. Acta Crystallographica Section A: Foundations and Advances, 2011, 67, C600-C601.	0.3	0
148	Quantification of Aromaticity in Oxocarbons: The Problem of the Fictitious "Nonaromatic―Reference System. Chemistry - A European Journal, 2002, 8, 433.	3.3	0