L Senthilkumar

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26 78 897 15 h-index g-index citations papers 80 1,069 4.87 3.1 avg, IF L-index ext. papers ext. citations

#	Paper	IF	Citations
78	Adsorption behaviour of reduced graphene oxide towards cationic and anionic dyes: Co-action of electrostatic and Interactions. <i>Materials Chemistry and Physics</i> , 2017 , 194, 243-252	4.4	145
77	Calcium decorated and doped phosphorene for gas adsorption. <i>Applied Surface Science</i> , 2016 , 377, 317	1-362 3	69
76	Interaction studies of cysteine with Li+, Na+, K+, Be2+, Mg2+, and Ca2+ metal cation complexes. <i>Journal of Physical Organic Chemistry</i> , 2011 , 24, 553-567	2.1	46
75	Hydrogen bonding in substituted formic acid dimers. <i>Journal of Physical Chemistry A</i> , 2006 , 110, 12623	-82.8	44
74	Electron density and energy decomposition analysis in hydrogen-bonded complexes of azabenzenes with water, acetamide, and thioacetamide. <i>Journal of Physical Chemistry A</i> , 2005 , 109, 75	75 ² 82	41
73	Multiwalled Carbon Nanotube Oxygen Sensor: Enhanced Oxygen Sensitivity at Room Temperature and Mechanism of Sensing. <i>ACS Applied Materials & Amp; Interfaces</i> , 2015 , 7, 23857-65	9.5	28
72	Hydrogen-bonded complexes of serotonin with methanol and ethanol: a DFT study. <i>Structural Chemistry</i> , 2014 , 25, 139-157	1.8	22
71	Theoretical studies on hydrogen bonding in caffeinetheophylline complexes. <i>Computational and Theoretical Chemistry</i> , 2012 , 979, 54-63	2	22
70	Coordination and binding properties of zwitterionic glutathione with transition metal cations. <i>Inorganica Chimica Acta</i> , 2012 , 387, 125-136	2.7	22
69	Influence of in-plane StoneII hrower Wales defects and edge functionalisation on the adsorption of CO2 and H2O on graphene. <i>RSC Advances</i> , 2014 , 4, 39576	3.7	21
68	Degradation of methyl salicylate through Cl initiated atmospheric oxidation (b) theoretical study. <i>RSC Advances</i> , 2014 , 4, 23464	3.7	19
67	DFT study on X?[[HD])(n=1-10) (X=OH, NO[]NO[]CO][]anionic water cluster. <i>Journal of Molecular Graphics and Modelling</i> , 2014 , 54, 148-63	2.8	18
66	Interface energetics of [Emim] + [X] [and [Bmim] + [X] [X = BF 4 , Cl, PF 6 , TfO, Tf 2 N) based ionic liquids on graphene, defective graphene, and graphyne surfaces. <i>Journal of Molecular Liquids</i> , 2017 , 236, 124-134	6	17
65	Improved lithium adsorption in boron- and nitrogen-substituted graphene derivatives. <i>Journal of Materials Science</i> , 2017 , 52, 815-831	4.3	16
64	Theoretical investigations on the hydrogen bonding of nitrile isomers with H2O, HF, NH3 and H2S. <i>Molecular Simulation</i> , 2013 , 39, 908-921	2	15
63	Hydrogen-bond interactions in hydrated 6-selenoguanine tautomers: a theoretical study. <i>Structural Chemistry</i> , 2014 , 25, 197-213	1.8	15
62	Study of effective hardness and condensed Fukui functions using AIM, ab initio, and DFT methods. <i>Molecular Physics</i> , 2005 , 103, 547-556	1.7	14

(2014-2015)

Defect-Mediated Reduction in Barrier for Helium Tunneling through Functionalized Graphene Nanopores. <i>Journal of Physical Chemistry C</i> , 2015 , 119, 20940-20948	3.8	13	
Edge functionalised & Li-intercalated 555-777 defective bilayer graphene for the adsorption of CO 2 and H 2 O. <i>Applied Surface Science</i> , 2017 , 400, 375-390	6.7	12	
Molecular interaction study of formohydroxamic acid (FHA) with water. <i>Journal of Molecular Structure</i> , 2006 , 791, 149-157	3.4	12	
Post Hartree E ock and density functional theory studies on tautomerism of 6-thioxanthine in gas phase and in solution. <i>Computational and Theoretical Chemistry</i> , 2003 , 638, 69-78		12	
H, OH and COOH functionalized magnesium phthalocyanine as a catalyst for oxygen reduction reaction (ORR) [A DFT study. <i>International Journal of Hydrogen Energy</i> , 2020 , 45, 8540-8548	6.7	11	
Gas adsorption efficacy of graphene sheets functionalised with carboxyl, hydroxyl and epoxy groups in conjunction with Stone-Thrower-Wales (STW) and inverse Stone-Thrower-Wales (ISTW) defects. <i>Physical Chemistry Chemical Physics</i> , 2017 , 19, 30895-30913	3.6	11	
An experimental and theoretical study of the kinetics of the reaction between 3-hydroxy-3-methyl-2-butanone and OH radicals. <i>RSC Advances</i> , 2015 , 5, 26559-26568	3.7	11	
DFT study on abstraction reaction mechanism of oh radical with 2-methoxyphenol. <i>Journal of Physical Organic Chemistry</i> , 2017 , 30, e3713	2.1	10	
Glutathione functionalized copper nanoclusters as a fluorescence platform for specific biosensing of cysteine and application in cellular imaging. <i>Microchemical Journal</i> , 2020 , 158, 105253	4.8	10	
Theoretical study on the interaction of CO2 and H2O molecules with metal doped-fluorinated phthalocyanines. <i>New Journal of Chemistry</i> , 2018 , 42, 3465-3472	3.6	10	
Density functional theory investigation of cocaine water complexes. <i>Journal of Molecular Modeling</i> , 2013 , 19, 3411-25	2	10	
Elucidation of Binding Mechanism of Photodynamic Therapeutic Agent Toluidine Blue O with Chicken Egg White Lysozyme by Spectroscopic and Molecular Dynamics Studies. <i>Photochemistry and Photobiology</i> , 2017 , 93, 1043-1056	3.6	9	
Theoretical studies on interaction of anticancer drugs (dacarbazine, procarbazine and triethylenemelamine) with normal (AT and GC) and mismatch (GG, CC, AA and TT) base pairs. <i>Molecular Simulation</i> , 2015 , 41, 633-652	2	9	
Reaction of OH radical and ozone with methyl salicylate 🗈 DFT study. <i>Journal of Physical Organic Chemistry</i> , 2015 , 28, 542-553	2.1	9	
Cobalt phthalocyanine is a suitable scaffold for lithium polysulfide (Li2Sn n = $2B$). Chemical Physics Letters, 2020 , 739, 136942	2.5	9	
The influence of interfaces and intra-band transitions on the band gap of CdS/HgS and GaN/X (X=InN, In0.33Ga0.67N) core/shell/shell quantum dot quantum well IA theoretical study. <i>Physica E: Low-Dimensional Systems and Nanostructures</i> , 2015 , 74, 204-212	3	8	
Influence of metal ions (Zn2+, Cu2+, Ca2+, Mg2+ and Na+) on the water coordinated neutral and zwitterionic L-histidine dimer. <i>RSC Advances</i> , 2014 , 4, 49040-49052	3.7	8	
Structure and NLO properties of halogen (F, Cl) substituted formic acid dimers. <i>Spectrochimica Acta - Part A: Molecular and Biomolecular Spectroscopy</i> , 2014 , 132, 821-32	4.4	8	
	Edge functionalised & Li-intercalated 555-777 defective bilayer graphene for the adsorption of CO 2 and H 2 O. <i>Applied Surface Science</i> , 2017, 400, 375-390 Molecular interaction study of formohydroxamic acid (FHA) with water. <i>Journal of Molecular Structure</i> , 2006, 791, 149-157 Post Hartreeflock and density functional theory studies on tautomerism of 6-thioxanthine in gas phase and in solution. <i>Computational and Theoretical Chemistry</i> , 2003, 638, 69-78 H, OH and COOH functionalized magnesium phthalocyanine as a catalyst for oxygen reduction reaction (ORR) IA DFT study. <i>International Journal of Hydrogen Energy</i> , 2020, 45, 8540-8548 Gas adsorption efficacy of graphene sheets functionalised with carboxyl, hydroxyl and epoxy groups in conjunction with Stone-Thrower-Wales (STW) and inverse Stone-Thrower-Wales (ISTW) defects. <i>Physical Chemistry Chemical Physics</i> , 2017, 19, 30895-30913 An experimental and theoretical study of the kinetics of the reaction between 3-hydroxy-3-methyl-2-butanone and OH radicals. <i>RSC Advances</i> , 2015, 5, 26559-26568 DFT study on abstraction reaction mechanism of oh radical with 2-methoxyphenol. <i>Journal of Physical Organic Chemistry</i> , 2017, 30, e3713 Glutathione functionalized copper nanoclusters as a fluorescence platform for specific biosensing of cysteine and application in cellular imaging. <i>Microchemical Journal</i> , 2020, 158, 105253 Theoretical study on the interaction of CO2 and H2O molecules with metal doped-fluorinated phthalocyanines. <i>New Journal of Chemistry</i> , 2018, 42, 3465-3472 Density functional theory investigation of cocaine water complexes. <i>Journal of Molecular Modeling</i> , 2013, 19, 3411-25 Elucidation of Binding Mechanism of Photodynamic Therapeutic Agent Toluidine Blue O with Chicken Egg White Lysozyme by Spectroscopic and Molecular Dynamics Studies. <i>Photochemistry and Photobiology</i> , 2017, 93, 1043-1056 Theoretical studies on interaction of anticancer drugs (dacarbazine, procarbazine and triethylenemelamine) with normal (AT and GC) and mismatch (GC, CC, AA	Edge functionalised & Li-intercalated 555-777 defective bilayer graphene for the adsorption of CO 2 and H 2 O. Applied Surface Science, 2017, 400, 375-390 Molecular interaction study of formohydroxamic acid (FHA) with water. Journal of Molecular Structure, 2006, 791, 149-157 Post HartreeBock and density functional theory studies on tautomerism of 6-thioxanthine in gas phase and in solution. Computational and Theoretical Chemistry, 2003, 638, 69-78 H, OH and COOH functionalized magnesium phthalocyanine as a calayst for oxygen reduction reaction (ORR) (B DFT study. International Journal of Hydrogen Energy, 2020, 45, 8540-8548 Gas adsorption efficacy of graphene sheets functionalised with carboxyl, hydroxyl and epoxy groups in conjunction with Stone-Thrower-Wales (STW) and inverse stone-Thrower-Wales (ISTW) defects. Physical Chemistry, Chemical Physics, 2017, 19, 30895-30913 An experimental and theoretical study of the kinetics of the reaction between 3-hydroxy-3-methyl-2-butanone and OH radicals. RSC Advances, 2015, 5, 26559-26568 DFT study on abstraction reaction mechanism of oh radical with 2-methoxyphenol. Journal of Physical Organic Chemistry, 2017, 30, e3713 Glutathione functionalized copper nanoclusters as a fluorescence platform for specific biosensing of cysteine and application in cellular imaging. Microchemical Journal, 2020, 158, 105253 Theoretical study on the interaction of CO2 and H2O molecules with metal doped-fluorinated phthalocyanines. New Journal of Chemistry, 2018, 42, 3465-3472 Density functional theory investigation of cocaine water complexes. Journal of Molecular Modeling, 2013, 19, 3411-25 Elucidation of Binding Mechanism of Photodynamic Therapeutic Agent Toluidine Blue O with Chicken Egg White Lysozyme by Spectroscopic and Molecular Dynamics Studies. Photochemistry and Photobiology, 2017, 93, 1043-1056 Theoretical studies on interaction of anticancer drugs (dacarbazine, procarbazine and triethylenemelamine) with normal (AT and GC) and mismatch (GG, CC, AA and TT) base pairs. Mo	Edge functionalised & Li-intercalated 555-777 defective bilayer graphene for the adsorption of CO 2 and H2 O. Applied Surface Science, 2017, 400, 375-390 Molecular interaction study of Formohydroxamic acid (FHA) with water. Journal of Molecular Structure, 2006, 791, 149-157 Post HartreeBock and density functional theory studies on tautomerism of 6-thioxanthine in gas phase and in solution. Computational and Theoretical Chemistry, 2003, 638, 69-78 H, OH and COOH functionalized magnesium phthalocyanine as a catalyst for oxygen reduction reaction (ORR) In DFT study. International Journal of Hydrogen Energy, 2020, 45, 8540-8548 67 11 Gas adsorption efficacy of graphene sheets functionalised with carboxyl, hydroxyl and epoxy groups in conjunction with Stone-Thrower-Walas (STW) and inverse Stone-Thrower-Wales (ISTW) defects. Physical Chemistry Chemical Physics, 2017, 19, 30895-30913 An experimental and theoretical study of the kinetics of the reaction between 3-hydroxy-3-methyl-2-butanone and OH radicals. RSC Advances, 2015, 5, 26559-26568 37 11 Glutathione functionalized copper nanoclusters as a fluorescence platform for specific biosensing of cysteine and application in cellular imaging. Microchemical Journal, 2020, 158, 105253 Theoretical study on the interaction of CO2 and H2O molecules with metal doped-fluorinated phthalocyanines. New Journal of CO2 and H2O molecules with metal doped-fluorinated phthalocyanines. New Journal of Co2 and H2O molecules with metal doped-fluorinated phthalocyanines. New Journal of Co2 and H2O molecules with metal doped-fluorinated phthalocyanine water study on the interaction of CO2 and H2O molecules with metal doped-fluorinated phthalocyanines. New Journal of Co2 and H2O molecules with metal doped-fluorinated phthalocyanines in the process of the physical Organic Chemistry, 2017, 33, 1043-1056 Theoretical studies on interaction of anticancer drugs (dacarbazine, procarbazine and triethylenemelamine) with normal (AT and GC) and mismatch (GG, CC, AA and TT) base pairs. Molecu

43	Hydrogen-bonded complexes of nicotine with simple alcohols. <i>International Journal of Quantum Chemistry</i> , 2012 , 112, 2787-2793	2.1	8
42	Encapsulation of fluoroethanols in pristine and Stone Wales defect boron nitride nanotube A DFT study. <i>Applied Surface Science</i> , 2015 , 345, 369-378	6.7	7
41	Understanding molecular properties of halogenated cyclohexane IA DFT study. <i>Computational and Theoretical Chemistry</i> , 2014 , 1049, 55-61	2	7
40	Spectroscopic investigations and hydrogen bond interactions of 8-aza analogues of xanthine, theophylline and caffeine: a theoretical study. <i>Journal of Molecular Modeling</i> , 2013 , 19, 1835-51	2	7
39	Experimental and theoretical investigations of the kinetics and mechanism of the Cl + 4-hydroxy-4-methyl-2-pentanone reaction. <i>Atmospheric Environment</i> , 2017 , 166, 315-326	5.3	7
38	Atmospheric fate of diketones and OH radical linetics, reaction force, ETS-NOCV analysis. <i>Molecular Physics</i> , 2017 , 115, 839-859	1.7	6
37	Reaction of NO3 radical with benzyl alcohol - A DFT study. <i>Computational and Theoretical Chemistry</i> , 2017 , 1102, 51-59	2	6
36	DFT study on the tautomerism of organic linker 1H-imidazole-4,5-tetrazole (HIT). <i>Computational and Theoretical Chemistry</i> , 2015 , 1068, 149-159	2	6
35	Theoretical investigations on hydrated 6,8-dithioguanine tautomers. Structural Chemistry, 2012, 23, 120	3:.821	86
34	Impact of heterogeneous passivation of trimethylphosphine oxide and di-methylphosphine oxide surface ligands on the electronic structure of CdnSen (n=6, 15) quantum dots: A DFT study. <i>Physica E: Low-Dimensional Systems and Nanostructures</i> , 2016 , 83, 284-296	3	6
33	Addition and abstraction reaction mechanism of 2,4,5-trimethylphenol with OH radical IA first principle study. <i>Computational and Theoretical Chemistry</i> , 2016 , 1092, 90-107	2	6
32	Impact of functional groups substitution on the molecular properties of magnesium and scandium phthalocyanines. <i>Inorganica Chimica Acta</i> , 2018 , 483, 203-210	2.7	5
31	Photoactive amorphous molecular materials based on bisquinoline diamines and their synthesis by Friedlider condensation reaction. <i>Journal of Photochemistry and Photobiology A: Chemistry</i> , 2014 , 283, 45-55	4.7	5
30	First Experimental and Theoretical Kinetic Study of the Reaction of 4-Hydroxy-4-methyl 2-pentanone as a Function of Temperature. <i>International Journal of Chemical Kinetics</i> , 2016 , 48, 584-600	1.4	4
29	Metal-interacted histidine dimer: an ETS-NOCV and XANES study. <i>RSC Advances</i> , 2016 , 6, 38919-38930	3.7	4
28	Influence of dopants Cu, Ga, In, Hg on the electronic structure of CdnSn (n = 6, 15) clusters a DFT study. <i>RSC Advances</i> , 2016 , 6, 93056-93067	3.7	4
27	CHIIIIO interaction between cation and anion in amino acid-based ionic liquids DFT study in gas and solvent phase. <i>Structural Chemistry</i> , 2019 , 30, 185-194	1.8	4
26	Synthesis of metal-free nitrogen-enriched porous carbon and its electrochemical sensing behavior for the highly sensitive detection of dopamine: Both experimental and theoretical investigation. <i>Materials Chemistry and Physics</i> , 2021 , 260, 124094	4.4	4

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25	Chemical Properties of Lithium Cluster (Lix, x = 28) on Stone Wales Defect Graphene Sheet: A DFT Study. <i>Journal of Physical Chemistry C</i> , 2020 , 124, 7229-7237	3.8	3
24	Water confined (H2O) n=1110 amino acid-based ionic liquids IA DFT study on the bonding, energetics and IR spectra. <i>Journal of Molecular Liquids</i> , 2020 , 304, 112720	6	3
23	Reaction of Pentanol isomers with OH radical IA theoretical perspective. <i>Molecular Physics</i> , 2018 , 116, 1153-1165	1.7	3
22	The study of performance of DFT functional for van der Waals interactions. <i>Computational and Theoretical Chemistry</i> , 2013 , 1004, 56-60	2	3
21	Facile Hydrothermal Synthesis and First Principle Computational Studies of NiSb2O4 and Its Electrochemical Properties with Ni3(Fe(CN)6)2(H2O) for Hybrid Supercapacitors. <i>ChemistrySelect</i> , 2017 , 2, 6823-6832	1.8	3
20	Importance of hydrogen bonding for second harmonic generation effect: X-ray diffraction and DFT study on S-benzyl isothiouronium chloride. <i>Journal of Physics and Chemistry of Solids</i> , 2009 , 70, 322-325	3.9	3
19	Adsorption and sensing properties of non-planar Burfaces towards high energy molecules: A density functional theory study. <i>Journal of Physics and Chemistry of Solids</i> , 2020 , 138, 109198	3.9	3
18	OH initiated oxidation mechanism of monoterpene (linalool) IA first comprehensive theoretical study. <i>Atmospheric Environment</i> , 2018 , 189, 235-243	5.3	3
17	DFT study of chemical reactivity parameters of lithium polysulfide molecules Li2Sn(1BB) in gas and solvent phase. <i>Computational and Theoretical Chemistry</i> , 2021 , 1202, 113323	2	3
16	An experimental and theoretical study on the kinetics of the reaction between 4-hydroxy-3-hexanone CH3CH2C(O)CH(OH)CH2CH3 and OH radicals. <i>International Journal of Chemical Kinetics</i> , 2018 , 50, 556-567	1.4	2
15	Effect of alkyl chain on the NLO property of nonylphenol isomers: a DFT study. <i>Monatshefte Fill Chemie</i> , 2015 , 146, 1983-1994	1.4	2
14	Pyrrolic, pyridinic, and graphitic sumanene as metal-free catalyst for oxygen reduction reaction IA density functional theory study. <i>Fuel Cells</i> , 2021 , 21, 490	2.9	2
13	Cation-Anion Interactions, Stability, and IR Spectra of Dicationic Amino Acid-Based Ionic Liquids Probed Using Density Functional Theory. <i>Journal of Molecular Modeling</i> , 2021 , 27, 180	2	2
12	Hydrogen bonds in Zif268 proteins - a theoretical perspective. <i>Journal of Biomolecular Structure and Dynamics</i> , 2016 , 34, 1607-24	3.6	1
11	Molecular properties of metal difluorides and their interactions with CO and HO molecules: a DFT investigation. <i>Journal of Molecular Modeling</i> , 2017 , 23, 345	2	1
10	ZnO and TiO2 clusters as catalyst in the addition and abstraction reaction of acrylic acid with the OH radical. <i>International Journal of Chemical Kinetics</i> , 2020 , 52, 3-15	1.4	1
9	Nitrogen-Doped Buckybowls as Potential Scaffold Material for Lithium-Sulfur Battery: A DFT Study. <i>Electrocatalysis</i> , 2021 , 12, 678	2.7	1
8	Effect of metal substitution (Mg, Sc) and functionalization (H, F, NH2, CH3, OH, CHO, and COOH) on the absorption properties of phthalocyanines [A TDDFT study. <i>Polyhedron</i> , 2020 , 176, 114244	2.7	O

7	Theoretical perspective on the interaction of CO2 and H2O molecules with functionalized magnesium and scandium phthalocyanines. <i>Theoretical Chemistry Accounts</i> , 2021 , 140, 1	1.9	О
6	Adsorption properties of amino acid-based ionic liquids (AAILs) on edge fluorinated graphene surface 🗈 DFT study. <i>Molecular Simulation</i> , 2021 , 47, 1066-1077	2	0
5	Complexes of criegee intermediate CH2OO with CO, CO2, H2O, SO2, NO2, CH3OH, HCOOH and CH3CH3CO molecules IA DFT study on bonding, energetics and spectra. <i>Computational and Theoretical Chemistry</i> , 2021 , 1203, 113341	2	0
4	Interaction between arginine conformers and Hofmeister halide anions. <i>Computational and Theoretical Chemistry</i> , 2016 , 1095, 93-103	2	
3	Experimental and Theoretical Studies of Trans-2-Pentenal Atmospheric Ozonolysis. <i>Atmosphere</i> , 2022 , 13, 291	2.7	
2	The first-principles study of CoSb2O4 and its electrochemical properties for supercapacitors. <i>Electrochimica Acta</i> , 2018 , 283, 949-958	6.7	
1	Enhancement of electrochemical performances of Li-S batteries using PPESK and Nelumbo nucifera derived porous carbon modified separator. <i>Materials Letters</i> , 2022 , 315, 131935	3.3	