Kittusamy Senthilkumar

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/4033599/publications.pdf

Version: 2024-02-01

121 3,070 25
papers citations h-index

123 123 3372 all docs docs citations times ranked citing authors

51

g-index

| # | Article | IF | CITATIONS |
|----|--|-----|-----------|
| 1 | Coordination Behavior of Acylthiourea Ligands in Their Ru(II)–Benzene Complexes─Structures and Anticancer Activity. Organometallics, 2022, 41, 1621-1630. | 1.1 | 33 |
| 2 | The influence of the shape and configuration of sensitizer molecules on the efficiency of DSSCs: a theoretical insight. RSC Advances, 2021, 11, 5556-5567. | 1.7 | 7 |
| 3 | Investigation on surface interaction between graphene nanobuds and cerium(III) via fluorescence excimer, theoretical, real water sample, and bioimaging studies. Materials Chemistry and Physics, 2021, 264, 124453. | 2.0 | 13 |
| 4 | Crossover from static to dynamic Non-Condon effecton charge Transport in Organic Semiconductors. Journal of Physics: Conference Series, 2021, 1916, 012230. | 0.3 | 0 |
| 5 | Charge Transport and Optical Absorption Properties of Dibenzocoronene Tetracarboxdiimide Based Liquid Crystalline Molecules: A Theoretical Study. Journal of Physical Chemistry A, 2021, 125, 3852-3862. | 1.1 | 8 |
| 6 | Room temperature weakly ferromagnetic energy band opened graphene quantum dot coupled solid sheets – A possible carbon based dilute magnetic semiconductor. Applied Surface Science, 2021, 548, 149195. | 3.1 | 4 |
| 7 | Mechanism, Kinetics, and Ecotoxicity Assessment of \hat{A} ·OH-Initiated Oxidation Reactions of Sulfoxaflor. Journal of Physical Chemistry A, 2021, 125, 10052-10064. | 1.1 | 3 |
| 8 | Modified fullerenes as acceptors in bulk heterojunction organic solar cells – a theoretical study. Physical Chemistry Chemical Physics, 2021, 23, 27468-27476. | 1.3 | 3 |
| 9 | First principle studies on the atmospheric oxidation of HFC-C1436 initiated by the OH radical. New Journal of Chemistry, 2020, 44, 2070-2082. | 1.4 | 8 |
| 10 | A comprehensive quantum chemical study on the mechanism and kinetics of atmospheric reactions of 3-chloro-2-methyl-1-propene with OH radical. Theoretical Chemistry Accounts, 2020, 139, 1. | 0.5 | O |
| 11 | Aggregation induced emission behavior in oleylamine acetone system and its application to get improved photocurrent from In2S3 quantum dots. Scientific Reports, 2020, 10, 19712. | 1.6 | 11 |
| 12 | Unimolecular decomposition of acetyl peroxy radical: a potential source of tropospheric ketene. Physical Chemistry Chemical Physics, 2020, 22, 26819-26827. | 1.3 | 7 |
| 13 | Insight into the photophysics of strong dual emission (blue & green) producing graphene quantum dot clusters and their application towards selective and sensitive detection of trace level Fe ³⁺ and Cr ⁶⁺ ions. RSC Advances, 2020, 10, 26613-26630. | 1.7 | 11 |
| 14 | Adsorption of phenanthroline and its derivatives on Au $(1\hat{A}1\hat{A}1)$ surface $\hat{a}\in$ Influence of substitution on structure and electronic properties. Computational Materials Science, 2020, 182, 109778. | 1.4 | O |
| 15 | Theoretical investigation on the structure and antioxidant activity of (+) catechin and (\hat{a}) epicatechin \hat{a} \in " a comparative study. Molecular Physics, 2020, 118, e1745917. | 0.8 | 24 |
| 16 | Hydrolysis of HNSO ₂ : A potential route for atmospheric production of H ₂ SO ₄ and NH ₃ . International Journal of Quantum Chemistry, 2020, 120, e26182. | 1.0 | 5 |
| 17 | Mechanism and kinetics of diuron oxidation by hydroxyl radical addition reaction. Environmental Science and Pollution Research, 2020, 27, 12080-12095. | 2.7 | 14 |
| 18 | Atmospheric oxidation mechanism and kinetics of 2-bromo-4,6-dinitroaniline by OH radicals $\hat{a}\in$ a theoretical study. Physical Chemistry Chemical Physics, 2019, 21, 21109-21127. | 1.3 | 0 |

| # | Article | IF | CITATIONS |
|----|--|-----|-----------|
| 19 | Mechanism and Kinetics of Diuron Oxidation Initiated by Hydroxyl Radical: Hydrogen and Chlorine Atom Abstraction Reactions. Journal of Physical Chemistry A, 2019, 123, 8954-8967. | 1.1 | 13 |
| 20 | Graphene Nanobuds: A New Second-Generation Phosgene Sensor with Ultralow Detection Limit in Aqueous Solution. ACS Applied Materials & Solution | 4.0 | 27 |
| 21 | Theoretical study on the gas phase reaction of methyl chavicol with hydroxyl radical. Computational and Theoretical Chemistry, 2019, 1151, 78-90. | 1.1 | 5 |
| 22 | Effect of site energy fluctuation on charge transport in disordered organic molecules. Journal of Chemical Physics, 2019, 151, 224301. | 1.2 | 9 |
| 23 | Adsorption of tetracyanoquinodimethane and tetrathiafulvalene on aluminium (100) surface $\hat{a} \in \hat{a}$ a first principle study of structural and electronic properties. Molecular Simulation, 2019, 45, 492-500. | 0.9 | 3 |
| 24 | Mechanism and kinetics of the oxidation of dimethyl carbonate by hydroxyl radical in the atmosphere. Environmental Science and Pollution Research, 2019, 26, 3357-3367. | 2.7 | 3 |
| 25 | Reaction mechanism and kinetics of the degradation of terbacil initiated by OH radical $\hat{a} \in A$ theoretical study. Chemical Physics, 2018, 501, 110-120. | 0.9 | 4 |
| 26 | Counter anion effect on structural, opto-electronic and charge transport properties of fused π-conjugated imidazolium compound. Molecular Physics, 2018, 116, 1145-1152. | 0.8 | 2 |
| 27 | Mechanism and kinetics of the reaction of methyl acetate with Cl atom $\hat{a}\in$ A theoretical study. Computational and Theoretical Chemistry, 2018, 1131, 40-50. | 1.1 | 4 |
| 28 | Exploring the mechanisms for the radical induced damage of 6â€thioguanine. International Journal of Quantum Chemistry, 2018, 118, e25544. | 1.0 | 1 |
| 29 | Theoretical Investigation on the Mechanism and Kinetics of Atmospheric Reaction of Methyldichloroacetate with Hydroxyl Radical. Journal of Physical Chemistry A, 2018, 122, 9316-9325. | 1.1 | 2 |
| 30 | Atmospheric Oxidation Mechanism and Kinetics of Hydrofluoroethers, CH ₃ OCF ₃ , CH ₃ OCHF ₂ , and CHF ₂ OCH ₂ CF ₃ , by OH Radical: A Theoretical Study. Journal of Physical Chemistry A, 2018, 122, 4972-4982. | 1.1 | 12 |
| 31 | Theoretical probe on modified organic dyes for high-performance dye-sensitised solar cell. Current Applied Physics, 2018, 18, 1071-1079. | 1.1 | 6 |
| 32 | Opto-electronic and interfacial charge transfer properties of azobenzene dyes on anatase TiO 2 (001) surface $\hat{a}\in$ The effect of anchoring group. Journal of Photochemistry and Photobiology A: Chemistry, 2017, 346, 372-381. | 2.0 | 10 |
| 33 | Conversion of toluene into benzyl radical on anatase TiO 2 (0 0 1) surface. Computational and Theoretical Chemistry, 2017, 1115, 13-21. | 1.1 | 1 |
| 34 | Mechanism and Kinetics of the Reaction of Nitrosamines with OH Radical: A Theoretical Study. International Journal of Chemical Kinetics, 2017, 49, 339-353. | 1.0 | 14 |
| 35 | The atmospheric oxidation mechanism and kinetics of 1,3,5-trimethylbenzene initiated by OH radicals $\hat{a}\in$ a theoretical study. New Journal of Chemistry, 2017, 41, 10259-10271. | 1.4 | 19 |
| 36 | Ab Initio QM/MM Modeling of the Rate-Limiting Proton Transfer Step in the Deamination of Tryptamine by Aromatic Amine Dehydrogenase. Journal of Physical Chemistry B, 2017, 121, 9785-9798. | 1.2 | 16 |

3

| # | Article | IF | Citations |
|----|--|-----|-----------|
| 37 | Graphene Quantum Dot Solid Sheets: Strong blue-light-emitting & photocurrent-producing band-gap-opened nanostructures. Scientific Reports, 2017, 7, 10850. | 1.6 | 61 |
| 38 | Theoretical and experimental evaluation of a new organic proton transfer crystal aminoguanidinium p-nitrobenzoate monohydrate for optical limiting applications. Journal of Physics and Chemistry of Solids, 2017, 111, 82-94. | 1.9 | 31 |
| 39 | Theoretical Investigations on the Mechanism and Kinetics of OH Radical Initiated Reactions of Monochloroacetic Acid. Journal of Physical Chemistry A, 2017, 121, 6028-6035. | 1.1 | 8 |
| 40 | A theoretical investigation on the mechanism and kinetics of the gas-phase reaction of naphthalene with OH radical. Theoretical Chemistry Accounts, 2017, 136, 1. | 0.5 | 9 |
| 41 | Theoretical studies on adsorption of organic molecules on metal surface. , 2017, , 209-241. | | O |
| 42 | Structural diversity in aroylthiourea copper complexes – formation and biological evaluation of [Cu(<scp>i</scp>)(μ-S)SCl] ₂ , cis-Cu(<scp>ii</scp>)S ₂ O ₂ , trans-Cu(<scp>ii</scp>)S ₃ cores. New Journal of Chemistry, 2016, 40, 5401-5413. | 1.4 | 23 |
| 43 | Dissociation of N2O on anatase TiO2 (001) surface – The effect of oxygen vacancy and presence of Ag cluster. Applied Surface Science, 2016, 389, 1220-1232. | 3.1 | 13 |
| 44 | Atmospheric oxidation mechanism of OH-initiated reactions of diethyl ether – the fate of the 1-ethoxy ethoxy radical. RSC Advances, 2016, 6, 81354-81363. | 1.7 | 7 |
| 45 | Synthesis, structural characterization, DNA/protein binding and inÂvitro cytotoxicity of three structurally different organoruthenium metallates from single pot. Journal of Organometallic Chemistry, 2016, 825-826, 83-99. | 0.8 | 7 |
| 46 | N-heterocycles as corrosion inhibitors for mild steel in acid medium. Journal of Molecular Liquids, 2016, 216, 42-52. | 2.3 | 94 |
| 47 | Adsorption of proline, hydroxyproline and glycine on anatase (001) surface: a first-principle study. Theoretical Chemistry Accounts, 2016 , 135 , 1 . | 0.5 | 8 |
| 48 | Forth–back oscillated charge carrier motion in dynamically disordered hexathienocoronene molecules: a theoretical study. Physical Chemistry Chemical Physics, 2015, 17, 17729-17738. | 1.3 | 8 |
| 49 | Effect of dynamic disorder on charge carrier dynamics in Ph4DP and Ph4DTP molecules. RSC Advances, 2015, 5, 38722-38732. | 1.7 | 10 |
| 50 | Adsorption of RGD tripeptide on anatase (001) surface $\hat{a} \in \text{``A first principle study. Computational Materials Science, 2015, 104, 124-129.}$ | 1.4 | 16 |
| 51 | Ultrasonic, DFT and FT-IR studies on hydrogen bonding interactions in aqueous solutions of diethylene glycol. Journal of Molecular Liquids, 2015, 202, 115-124. | 2.3 | 38 |
| 52 | Effect of Structural Fluctuations on Charge Carrier Dynamics in Triazene Based Octupolar Molecules. Journal of Physical Chemistry C, 2014, 118, 27754-27762. | 1.5 | 14 |
| 53 | Structure and spectral properties of l-histidinium dipicrate dihydrate. Spectrochimica Acta - Part A: Molecular and Biomolecular Spectroscopy, 2014, 118, 102-111. | 2.0 | 17 |
| 54 | Structural, optical, and charge transport properties of cyclopentadithiophene derivatives: a theoretical study. Structural Chemistry, 2014, 25, 715-731. | 1.0 | 18 |

| # | Article | IF | Citations |
|----|--|-----|-----------|
| 55 | A theoretical study on optical and charge transport properties of anthra-[1,2-b:4,3-b′:5,6-b″:8,7-bâ€]tetrathiophene molecules. Chemical Physics, 2014, 433, 48-59. | 0.9 | 14 |
| 56 | Adsorption of perfluoropentacene on aluminum (100) surface: Structural and electronic properties from first principle study. Computational Materials Science, 2014, 89, 216-223. | 1.4 | 2 |
| 57 | Charge transport and optical properties of cross-conjugated organic molecules: A theoretical study. Organic Electronics, 2014, 15, 1607-1623. | 1.4 | 15 |
| 58 | Triazolyl-donor–acceptor chromophore-decorated unnatural amino acids and peptides: FRET events in a β-turn conformation. Chemical Communications, 2014, 50, 433-435. | 2.2 | 17 |
| 59 | Theoretical studies on the quinoidal thiophene based dyes for dye sensitized solar cell and NLO applications. Physical Chemistry Chemical Physics, 2014, 16, 21496-21505. | 1.3 | 30 |
| 60 | Reaction mechanism and kinetics of the degradation of bromoxynil initiated by OH radical. RSC Advances, 2014, 4, 7749. | 1.7 | 7 |
| 61 | Oxidation and Nitration of Tyrosine by Ozone and Nitrogen Dioxide: Reaction Mechanisms and Biological and Atmospheric Implications. Journal of Physical Chemistry B, 2014, 118, 3479-3490. | 1.2 | 23 |
| 62 | Theoretical studies on charge transport and optical properties of tris(N-saclicylideneanilines). RSC Advances, 2014, 4, 25969. | 1.7 | 8 |
| 63 | Mechanism and kinetics of the atmospheric degradation of perfluoropolymethylisopropyl ether by OH radical: a theoretical study. Structural Chemistry, 2014, 25, 1773-1783. | 1.0 | 3 |
| 64 | A theoretical probe on the non-covalent interactions of sulfadoxine drug with pi-acceptors. Journal of Molecular Structure, 2014, 1074, 157-167. | 1.8 | 6 |
| 65 | Depletion of atmospheric ozone by nitrogen dioxide: a bifurcated reaction pathway. Theoretical Chemistry Accounts, 2013, 132, 1. | 0.5 | 7 |
| 66 | Structural properties and the effect of platinum drugs with DNA base pairs. Structural Chemistry, 2013, 24, 583-595. | 1.0 | 13 |
| 67 | Effect of structural fluctuations on charge carrier mobility in thiophene, thiazole and thiazolothiazole based oligomers. Physical Chemistry Chemical Physics, 2013, 15, 17947. | 1.3 | 32 |
| 68 | Theoretical and experimental studies on the structure and spectroscopic properties of Ni(II) complexes of the type [Ni(L)(PPh3)] [H2L=5-methyl-N-(2-mercaptophenyl)salicylideneimine and 5-chloro-N-(2-mercaptophenyl)salicylideneimine]. Journal of Molecular Structure, 2013, 1037, 367-375. | 1.8 | 8 |
| 69 | Wurtzite ZnSe quantum dots: synthesis, characterization and PL properties. Journal of Materials Science: Materials in Electronics, 2013, 24, 692-696. | 1.1 | 22 |
| 70 | Mechanism and Kinetics of the Atmospheric Oxidative Degradation of Dimethylphenol Isomers Initiated by OH Radical. Journal of Physical Chemistry A, 2013, 117, 4611-4626. | 1.1 | 26 |
| 71 | Opto-electronic properties of low band gap fused-ring thieno[3,4-b]pyrazine analogues – A theoretical study. Molecular Physics, 2013, 111, 3036-3046. | 0.8 | 5 |
| 72 | A theoretical study of structural and electronic properties of pentacene/Al(1 00) interface. Journal of Molecular Graphics and Modelling, 2012, 38, 334-341. | 1.3 | 10 |

| # | Article | IF | CITATIONS |
|----|---|------------------|--------------------|
| 73 | Synthesis and characterization studies of ZnSe quantum dots. Journal of Materials Science: Materials in Electronics, 2012, 23, 2048-2052. | 1.1 | 25 |
| 74 | Understanding the absorption and emission spectra of borondipyrromethene dye and its substituted analogues. Molecular Physics, 2012, 110, 445-456. | 0.8 | 11 |
| 75 | Copper Ion Mediated Selective Cleavage of C–S Bond in Ferrocenylthiosemicarbazone Forming Mixed Geometrical [(PPh ₃)Cu(μ-S) ₂ Cu(PPh ₃) ₂] Having Cu ₂ S ₂ Core: Toward a New Avenue in Copper–Sulfur Chemistry. Inorganic Chemistry. 2012. 51. 3525-3532. | 1.9 | 29 |
| 76 | Reaction mechanism and kinetics of the atmospheric oxidation of 1,4-thioxane by NO ₃ Ââ€" A theoretical study. Canadian Journal of Chemistry, 2012, 90, 384-394. | 0.6 | 6 |
| 77 | Theoretical studies on the reaction mechanism and kinetics of the atmospheric reactions of 1,4-thioxane with OH radical. Structural Chemistry, 2012, 23, 1475-1488. | 1.0 | 25 |
| 78 | Low temperature method for synthesis of starch-capped ZnSe nanoparticles and its characterization studies. Journal of Applied Physics, 2012, 112, 114331. | 1.1 | 7 |
| 79 | Charge transport and optical properties of discotic liquid crystalline molecules THDDP and substituted THDP. International Journal of Quantum Chemistry, 2012, 112, 713-723. | 1.0 | 6 |
| 80 | Mechanism and kinetics of the reaction of 1,4-thioxane with O3 in the atmosphere – A theoretical study. Chemical Physics Letters, 2012, 525-526, 153-159. | 1.2 | 8 |
| 81 | Theoretical investigation of interaction between psoralen and altretamine with stacked DNA base pairs. Materials Science and Engineering C, 2012, 32, 423-431. | 3.8 | 25 |
| 82 | Structural and Spectral Properties of 4-Bromo-1-naphthyl Chalcones: A Quantum Chemical Study. Journal of Physical Chemistry A, 2011, 115, 6594-6602. | 1.1 | 36 |
| 83 | Optical Absorption and Emission Properties of Fluoranthene, Benzo[k]fluoranthene, and Their Derivatives. A DFT Study. Journal of Physical Chemistry A, 2011, 115, 14647-14656. | 1.1 | 41 |
| 84 | Hydrogen-bonding studies of amino acid side-chains with DNA base pairs. Molecular Physics, 2011, 109, 1995-2008. | 0.8 | 10 |
| 85 | Structural properties and the effect of interaction of alkali (Li+, Na+, K+) and alkaline earth (Be2+,) Tj ETQq1 1 0.57-65. | 784314 rg 1.1 | BT /Overlock 11 |
| 86 | Longâ€range charge transfer in donorâ€peptide bridgeâ€acceptor model systems—A theoretical study. International Journal of Quantum Chemistry, 2011, 111, 3904-3914. | 1.0 | 2 |
| 87 | Structural properties and the effect of 2,6â€diaminoanthraquinone on Gâ€tetrad, nonâ€Gâ€tetrads, and mixed tetradsâ€"A density functional theory study. International Journal of Quantum Chemistry, 2011, 111, 3239-3250. | 1.0 | 8 |
| 88 | A theoretical study on decomposition and rearrangement reaction mechanism of trichloroacetyl chloride (CCl ₃ COCl). International Journal of Quantum Chemistry, 2011, 111, 3482-3496. | 1.0 | 5 |
| 89 | Copper(I) hydrazone complexes: Synthesis, structure, DNA binding, radical scavenging and computational studies. Inorganic Chemistry Communication, 2011, 14, 1318-1322. | 1.8 | 53 |
| 90 | Hydrogen bond interactions in hydrated acetylsalicylic acid. Computational and Theoretical Chemistry, 2011, 966, 167-179. | 1.1 | 10 |

| # | Article | IF | CITATIONS |
|-----|---|-----|-----------|
| 91 | First and second coordination spheres in 8-azaxanthinato salts of divalent metal aquacomplexes – Ab initio and DFT study. Polyhedron, 2011, 30, 1431-1445. | 1.0 | 8 |
| 92 | Reaction mechanism of cysteine proteases model compound HSH with diketone inhibitor PhCOCOCH _{$3\hat{a}^2$(i>n} x (sub> <i>n</i> , (X = F, Cl, <i>n</i> = 0, 1, 2). International Journal of Quantum Chemistry, 2010, 110, 1660-1674. | 1.0 | 12 |
| 93 | FOBEZ-256 (A hashing function using Bezier curve)., 2010,,. | | O |
| 94 | Tautomerization and solvent effects on the absorption and emission properties of the Schiff base <i>N</i> , <i>N</i> ,6>N,6>N,6>N,6>N,6>N,6>N,6>N,6>N,6>N,6>N,6>N,7 <n< i="">,7>N,7<n< i="">,7<n< td=""><td>0.8</td><td>18</td></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<></n<> | 0.8 | 18 |
| 95 | Absorption and emission properties of phenylene ethynylene oligomers: effect of substitution and ï€-conjugation length. Molecular Physics, 2009, 107, 1629-1639. | 0.8 | 17 |
| 96 | Calculation of ionization potential and chemical hardness: A comparative study of different methods. International Journal of Quantum Chemistry, 2009, 109, 764-771. | 1.0 | 59 |
| 97 | Effect of conformational degrees of freedom on the charge transfer in model tripeptide. Journal of Molecular Graphics and Modelling, 2009, 27, 784-791. | 1.3 | 6 |
| 98 | Lennardâ^'Jones Parameters for B3LYP/CHARMM27 QM/MM Modeling of Nucleic Acid Bases. Journal of Chemical Theory and Computation, 2009, 5, 396-410. | 2.3 | 17 |
| 99 | Interactions of anticancer drugs with usual and mismatch base pairs â€" Density functional theory studies. Biophysical Chemistry, 2008, 136, 50-58. | 1.5 | 26 |
| 100 | Analysis of polarization in QM/MM modelling of biologically relevant hydrogen bonds. Journal of the Royal Society Interface, 2008, 5, 207-216. | 1.5 | 49 |
| 101 | Post Hartree–Fock and density functional theory studies on Di-Protonated Allopurinol2+. Computational and Theoretical Chemistry, 2007, 810, 25-30. | 1.5 | 2 |
| 102 | Theoretical investigation on intramolecular electron transfer in polypeptides. Chemical Physics Letters, 2007, 440, 302-307. | 1.2 | 7 |
| 103 | Charge Transfer in Polypeptides:  Effect of Secondary Structures on Charge-Transfer Integral and Site Energies. Journal of Physical Chemistry A, 2006, 110, 11551-11556. | 1.1 | 15 |
| 104 | Two-Dimensional Charge Delocalization in X-Shaped Phenylenevinylene Oligomers. Chemistry of Materials, 2006, 18, 2118-2129. | 3.2 | 23 |
| 105 | Effect of substitution of electron-donating and -withdrawing groups on the stability of flavin–diaminepyridine complexes—a density functional theory study. Computational and Theoretical Chemistry, 2006, 758, 107-112. | 1.5 | 12 |
| 106 | Absolute Rates of Hole Transfer in DNA. Journal of the American Chemical Society, 2005, 127, 14894-14903. | 6.6 | 325 |
| 107 | Charge Transport in Self-Organized π-Stacks ofp-Phenylene Vinylene Oligomers. Journal of Physical Chemistry B, 2005, 109, 18267-18274. | 1.2 | 90 |
| 108 | Charge Transport Properties in Discotic Liquid Crystals:  A Quantum-Chemical Insight into Structureâ^'Property Relationships. Journal of the American Chemical Society, 2004, 126, 3271-3279. | 6.6 | 464 |

| # | Article | IF | CITATIONS |
|-----|--|-----|-----------|
| 109 | A quantative study of the charge-transfer between conjugated thiophene rings in vibrationally excited states. Physica B: Condensed Matter, 2004, 350, 220-223. | 1.3 | 7 |
| 110 | Structure, conformation and NMR studies on 1,2-dioxane and halogen substituted 1,2-dioxane molecules. Computational Biology and Chemistry, 2003, 27, 173-183. | 1.1 | 13 |
| 111 | Charge transport in columnar stacked triphenylenes: Effects of conformational fluctuations on charge transfer integrals and site energies. Journal of Chemical Physics, 2003, 119, 9809-9817. | 1.2 | 395 |
| 112 | Mapping the Sites for Selective Oxidation of Guanines in DNA. Journal of the American Chemical Society, 2003, 125, 13658-13659. | 6.6 | 97 |
| 113 | Hartree-Fock and density functional theory studies on ionization and fragmentation of halomethane molecules by positron impact. Molecular Physics, 2002, 100, 3817-3822. | 0.8 | 12 |
| 114 | Molecular structure, conformational stability and cis effect of 1,4-dichlorobutadiene $\hat{a}\in$ " a quantum chemical study. Computational and Theoretical Chemistry, 2002, 577, 69-79. | 1.5 | 1 |
| 115 | Origin of the cis effect—nonbonded intramolecular interactions: quantum chemical studies on 1,2-dihaloethylene molecules. Computational and Theoretical Chemistry, 2002, 589-590, 95-102. | 1.5 | 18 |
| 116 | Post Hartreeâ€"Fock and density functional theory studies on structure and conformational stability of nitrosoethylene and substituted compounds of nitrosoethylene. Computers & Chemistry, 2002, 26, 207-221. | 1.2 | 20 |
| 117 | Quantum chemical studies on tautomerism of barbituric acid in gas phase and in solution. Journal of Computer-Aided Molecular Design, 2002, 16, 263-272. | 1.3 | 47 |
| 118 | Studies of chemical hardness and Fukui function using the exact solution of the density functional theory. International Journal of Quantum Chemistry, 2001, 81, 4-10. | 1.0 | 5 |
| 119 | Ab initio and DFT studies on structure and stability of aliphatic aldoxime molecules. Computational and Theoretical Chemistry, 2001, 535, 61-70. | 1.5 | 33 |
| 120 | Study of chemical bonding in H2 and HF molecules: Wave function and density functional theory (DFT) parameters approach. International Journal of Quantum Chemistry, 2000, 76, 662-669. | 1.0 | 6 |
| 121 | Simultaneous Reduction of NOx and Soot Using Early Post Injection. , 0, , . | | 8 |