

Prabhat K Sahu

List of Publications by Year in descending order

Source: <https://exaly.com/author-pdf/10867307/publications.pdf>

Version: 2024-02-01

18

papers

250

citations

1040056

9

h-index

940533

16

g-index

19

all docs

19

docs citations

19

times ranked

264

citing authors

#	ARTICLE	IF	CITATIONS
1	Many-body interaction in glycineâ€“(water)3 complex using density functional theory method. <i>Journal of Chemical Physics</i> , 2004, 120, 170-174.	3.0	71
2	Theoretical investigation for the hydrogen bond interaction in THFâ€“water complex. <i>Chemical Physics Letters</i> , 2004, 386, 351-355.	2.6	32
3	Hydrogen-bond interaction in 1:1 complexes of tetrahydrofuran with water, hydrogen fluoride, and ammonia: A theoretical study. <i>Journal of Chemical Physics</i> , 2005, 123, 044308.	3.0	20
4	Effect of microsolvation on zwitterionic glycine: an ab initio and density functional theory study. <i>Journal of Molecular Modeling</i> , 2008, 14, 385-392.	1.8	17
5	Quantum Mechanical Calculations for the Misincorporation of Nucleotides Opposite Mutagenic 3,⁴<i>N</i>-Ethenocytosine. <i>Journal of Physical Chemistry B</i> , 2012, 116, 11173-11179.	2.6	15
6	Hydrogen bonding interaction in sarcosine-water complex using ab initio and DFT method. <i>International Journal of Quantum Chemistry</i> , 2005, 101, 97-103.	2.0	14
7	A Density Functional Theory Study for the Hydrogen-Bonded Nucleic Acid Base Pair:Â Cytosine Dimer. <i>Journal of Physical Chemistry A</i> , 2005, 109, 2887-2893.	2.5	14
8	Net-sign identity information index: A novel approach towards numerical characterization of chemical signed graph theory. <i>Chemical Physics Letters</i> , 2008, 454, 133-138.	2.6	11
9	Interaction of Adenine Adducts with Thymine:Â A Computational Study. <i>Journal of Physical Chemistry B</i> , 2007, 111, 2991-2998.	2.6	9
10	Hydrogen-bond interactions in THFâ€“H2Oâ€“HF: A theoretical study. <i>International Journal of Quantum Chemistry</i> , 2007, 107, 2015-2023.	2.0	9
11	Model Calculations for the Misincorporation of Nucleotides Opposite Five-Membered Exocyclic DNA Adduct: ²<i>N</i>-,3-Ethenoguanine. <i>Journal of Physical Chemistry B</i> , 2011, 115, 10537-10546.	2.6	8
12	Density functional theory study of formaldehyde oligomers. <i>International Journal of Quantum Chemistry</i> , 2005, 101, 67-72.	2.0	6
13	Interaction of Serotonin and Fluoxetine: Toward Understanding the Importance of the Chirality of Fluoxetine (^S<i>S</i> form and ^R<i>R</i> form). <i>Journal of Physical Chemistry B</i> , 2009, 113, 14529-14535.	2.6	6
14	Many-body interactions of carbon monoxide cyclic oligomers: A computational study. <i>International Journal of Quantum Chemistry</i> , 2005, 103, 314-321.	2.0	5
15	Spectroscopic probe on Nâ€“Hâ€“N, Nâ€“Hâ€“O and controversial Câ€“Hâ€“O contact in Aâ€“T base pair: A DFT study. <i>Spectrochimica Acta - Part A: Molecular and Biomolecular Spectroscopy</i> , 2014, 120, 542-547.	3.9	5
16	Novel information theoretic topological index I_k for unsaturated hydrocarbons. <i>Chemical Physics Letters</i> , 2004, 396, 465-468.	2.6	4
17	Model calculations for the base-pairing specificity of mutagenic exocyclic DNA adduct 1,N-6-ethenoadenine. <i>Structural Chemistry</i> , 2014, 25, 561-573.	2.0	2
18	On the Development and Application of Net-Sign Graph Theory. , 2011, , 127-151.		0