

Sadasivam Manogaran

List of Publications by Year in descending order

Source: <https://exaly.com/author-pdf/4807934/publications.pdf>

Version: 2024-02-01

11
papers

142
citations

1307594

7
h-index

1372567

10
g-index

11
all docs

11
docs citations

11
times ranked

157
citing authors

#	ARTICLE	IF	CITATIONS
1	Substituent effects on the aromaticity of benzene—An approach based on interaction coordinates. <i>Journal of Chemical Physics</i> , 2019, 150, 214108.	3.0	8
2	Quantification of Aromaticity of Heterocyclic Systems Using Interaction Coordinates. <i>Journal of Physical Chemistry A</i> , 2018, 122, 6953-6960.	2.5	22
3	Quantification of Hydrogen Bond Strength Based on Interaction Coordinates: A New Approach. <i>Journal of Physical Chemistry A</i> , 2017, 121, 6090-6103.	2.5	31
4	Quantification of Aromaticity Based on Interaction Coordinates: A New Proposal. <i>Journal of Physical Chemistry A</i> , 2016, 120, 2894-2901.	2.5	26
5	Intramolecular hydrogen bond: Can it be part of the basis set of valence internal coordinates in normal mode analysis?. <i>Journal of Chemical Sciences</i> , 2015, 127, 1127-1134.	1.5	8
6	Redundant internal coordinates, compliance constants and non-bonded interactions — some new insights. <i>Journal of Chemical Sciences</i> , 2013, 125, 9-15.	1.5	6
7	Vibrational spectra of triamantane X18H24, iso-tetramantane X22H28 and cyclohexamantane X26H30 (X=C, Si, Ge, Sn) — A theoretical study. <i>Computational and Theoretical Chemistry</i> , 2007, 816, 31-41.	1.5	10
8	Vibrational spectra of adamantanes X10H16 and diamantanes X14H20 (X=C, Si, Ge, Sn): A theoretical study. <i>Computational and Theoretical Chemistry</i> , 2006, 766, 125-135.	1.5	15
9	Electronic structure and vibrational spectra of X20H20 (X=C, Si, Ge, Sn): A theoretical study. <i>Computational and Theoretical Chemistry</i> , 2005, 730, 171-176.	1.5	5
10	Force field calculation of molecules with isotopomers of different symmetries in vibrational spectral analysis. <i>Computational and Theoretical Chemistry</i> , 2001, 574, 245-254.	1.5	11
11	Anharmonicity in compliance formalism: Potential constants and interaction coordinates. <i>International Journal of Quantum Chemistry</i> , 0, , e26841.	2.0	0