Elizete Ventura do Monte

List of Publications by Year in descending order

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567281 642732 55 651 15 23 citations h-index g-index papers 56 56 56 662 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	The Diels–Alder Reaction of Ethene and 1,3-Butadiene: An Extended Multireference ab initio Investigation. ChemPhysChem, 2004, 5, 1365-1371.	2.1	47
2	The valence-excited states T1–T4 and S1–S2 of acetylene: A high-level MR-CISD and MR-AQCC investigation of stationary points, potential energy surfaces, and surface crossings. Journal of Chemical Physics, 2003, 118, 1702-1713.	3.0	45
3	The generality of the GUGA MRCI approach in COLUMBUS for treating complex quantum chemistry. Journal of Chemical Physics, 2020, 152, 134110.	3.0	42
4	Determination of energy minima and saddle points using multireference configuration interaction methods in combination with reduced gradient following: The SO surface of H2CO and the T1 and T2 surfaces of acetylene. Journal of Computational Chemistry, 2002, 23, 576-583.	3.3	41
5	Topological twistons in crystalline polyethylene1This work is partially supported by the US Department of Energy (DOE) under cooperative research agreement DE-FC02-94ER40818, and by the Brazilian agencies CAPES and CNPq.1. Chemical Physics Letters, 1999, 303, 341-346.	2.6	36
6	Synthesis and conformational study of a new class of highly bioactive compounds. Chemical Physics Letters, 2007, 449, 336-340.	2.6	31
7	Theoretical study of cooperative effects in the homo- and heteromeric hydrogen bond chains (HCN)nHF withn = 1, 2, and 3. International Journal of Quantum Chemistry, 2006, 106, 2714-2722.	2.0	29
8	Cope Rearrangement of 1,5-Hexadiene:  Full Geometry Optimizations Using Analytic MR-CISD and MR-AQCC Gradient Methods. Journal of Physical Chemistry A, 2003, 107, 1175-1180.	2.5	27
9	Photochemistry of CH ₃ Cl: Dissociation and CH···Cl Hydrogen Bond Formation. Journal of the American Chemical Society, 2016, 138, 272-280.	13.7	27
10	An ab initio study of the C2H2HF, C2H(CH3) HF and C2(CH3)2HF hydrogen-bonded complexes. Spectrochimica Acta - Part A: Molecular and Biomolecular Spectroscopy, 2006, 63, 383-390.	3.9	23
11	Exact topological twistons in crystalline polyethylene. Chemical Physics Letters, 2000, 320, 587-593.	2.6	22
12	AM1 Sparkle modeling of Er(III) and Ce(III) coordination compounds. Journal of Organometallic Chemistry, 2006, 691, 2584-2588.	1.8	22
13	Challenges encountered during development of Mn porphyrin-based, potent redox-active drug and superoxide dismutase mimic, MnTnBuOE-2-PyP5+, and its alkoxyalkyl analogues. Journal of Inorganic Biochemistry, 2017, 169, 50-60.	3.5	18
14	Dissociation of ground and $n\ddot{l}f^*$ states of CF3Cl using multireference configuration interaction with singles and doubles and with multireference average quadratic coupled cluster extensivity corrections. Journal of Chemical Physics, 2007, 127, 164320.	3.0	17
15	Revisiting the concept of the (a)synchronicity of dielsa \in alder reactions based on the dynamics of quasiclassical trajectories. Journal of Computational Chemistry, 2016, 37, 701-711.	3.3	16
16	Photochemical Deactivation Process of HCFC-133a (C ₂ H ₂ F ₃ Cl): A Nonadiabatic Dynamics Study. Journal of Physical Chemistry A, 2014, 118, 12041-12049.	2.5	15
17	CASSCF and MR–CISD study of the nâ^'4s and nâ^'4pe Rydberg states of the CF3Cl. Chemical Physics Letters, 2012, 546, 30-33.	2.6	12
18	Dynamic Effects Dictate the Mechanism and Selectivity of Dehydration–Rearrangement Reactions of Protonated Alcohols [Me 2 (R)CCH(OH 2)Me] + (R=Me, Et, i Pr) in the Gas Phase. Chemistry - A European Journal, 2014, 20, 13742-13754.	3.3	12

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19	Revisiting the stationary points on the potential energy surface of tetramethylene at the MR-AQCC level using analytic gradients. Journal of Chemical Physics, 2003, 118, 10963-10972.	3.0	11
20	A theoretical prediction of stability in hydrogen-bonded complexes formed between oxirane and oxetane rings with HX (X=F and Cl). Spectrochimica Acta - Part A: Molecular and Biomolecular Spectroscopy, 2006, 64, 156-160.	3.9	11
21	Assessment of density-functionals for describing the $X\hat{a}^{\circ}$ + CH3ONO2 gas-phase reactions with X = F, OH, CH2CN. Physical Chemistry Chemical Physics, 2014, 16, 26769-26778.	2.8	11
22	Ab initio and DFT conformational study on nitrosamine (H2N–N=O) and N-Nitrosodimethylamine [(CH3)2N–N=O]. Structural Chemistry, 2011, 22, 497-507.	2.0	10
23	Matrix isolation infrared spectroscopic and theoretical study of 1,1,1-trifluoro-2-chloroethane (HCFC-133a). Journal of Chemical Physics, 2013, 139, 204302.	3.0	10
24	Valence and Rydberg states of CH ₃ Cl: a MR-CISD study. RSC Advances, 2014, 4, 64085-64092.	3.6	10
25	Spinâ€Forbidden Branching in the Mechanism of the Intrinsic Haber–Weiss Reaction. ChemistryOpen, 2017, 6, 360-363.	1.9	10
26	Photochemistry of CF ₃ Cl: Quenching of Charged Fragments Is Caused by Nonadiabatic Effects. Journal of Chemical Theory and Computation, 2018, 14, 4844-4855.	5. 3	10
27	Photoinduced Formation of H-Bonded Ion Pair in HCFC-133a. Journal of Physical Chemistry A, 2019, 123, 1953-1961.	2.5	10
28	A theoretical study of hydrogen complexes of the XH-ï€ type between propyne and HF, HCL or HCN. Spectrochimica Acta - Part A: Molecular and Biomolecular Spectroscopy, 2006, 64, 412-417.	3.9	6
29		3.3	6
30	Effects of Ï€ bond type, backbone size, and halogen on structural and spectroscopic properties of hydrogen-bonded complexes of the XH … Ï€ type between alkenes or alkynes and haloacids (HF and) Tj ETÇ	190 20 00 rgE	BT © verlock 1
31	Matrix Isolation FTIR Spectroscopic and Theoretical Study of 3,3-Dichloro-1,1,1-Trifluoropropane (HCFC-243). Journal of Physical Chemistry A, 2008, 112, 11641-11648.	2.5	5
32	CASSCF and multireference CI with singles and doubles study of lowâ€lying valence and Rydberg states of 2Hâ€tetrazole. Journal of Computational Chemistry, 2009, 30, 1075-1081.	3.3	5
33	Hydrogen-bonded contact ion pair in gaseous chloroethane: a multi-reference configuration interaction with singles and doubles (MR-CISD) study including extensivity corrections. Theoretical Chemistry Accounts, 2020, 139, 1.	1.4	5
34	Accurate calculation of the ionization energies of the chlorine lone pairs in 1,1,1-trifluoro-2-chloroethane (HCFC-133a). Journal of Molecular Modeling, 2014, 20, 2393.	1.8	4
35	Can a gas phase contact ion pair containing a hydrocarbon carbocation be formed in the ground state?. RSC Advances, 2021, 11, 4221-4230.	3.6	4
36	Crystallographic and theoretical investigations of the preferred conformations of three isomeric N-acetyl-dihydro-oxadiazoles. Journal of Molecular Structure, 2001, 561, 29-41.	3.6	3

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37	Ab-initio study of the electronic factor (Î") conformational dependence in [CH2â€"(phenylene)nâ€"CH2]â-" electron transfer model systems. Chemical Physics, 2005, 313, 311-323.	1.9	3
38	Theoretical calculations of the substituent effect on molecular properties of the RCNâcHF hydrogen-bonded complexes with R=NH2, CH3O, CH3, OH, SH, H, Cl, F, CF3, CN and NO2. Vibrational Spectroscopy, 2009, 49, 133-141.	2.2	3
39	Ab initio and DFT conformational study on N-nitrosodiethylamine, (C2H5)2N-N=O. Journal of Molecular Modeling, 2012, 18, 339-350.	1.8	3
40	Photochemistry of Monohydrated Chloromethane: Formation of Free and Hydrated Cl [–] and CH ₃ ⁺ lons from a Solvent-Shared Semi-Ion-Pair. Journal of Physical Chemistry A, 2021, 125, 8603-8614.	2.5	3
41	Modeling the heating and cooling of a chromophore after photoexcitation. Physical Chemistry Chemical Physics, 2022, 24, 9403-9410.	2.8	3
42	Diastereoselective Epoxidation of Allylic Diols Derived from Baylis-Hillman Adducts. Synthesis, 2005, 2005, 2297-2306.	2.3	2
43	Effects of wave function modifications on calculated carbon–carbontriple bond lengths. Computational and Theoretical Chemistry, 2006, 758, 253-258.	1.5	2
44	Control of ionic properties of N-nitrosodimethylamine through hydrogen substitution by fluorine atoms. Structural Chemistry, 2012, 23, 1193-1201.	2.0	2
45	Electronic properties of the low-lying spin states of dimethylnitrosamine coordinated to Fe(III) heme models: An ab initio study. International Journal of Quantum Chemistry, 2014, 114, 508-520.	2.0	2
46	A multi-reference CI study of the low-lying valence and Rydberg states of CF ₃ radical. Molecular Physics, 2018, 116, 2187-2195.	1.7	2
47	Increasing the kinetic stability of a gasâ€phase contact ionâ€pair through enhancement of the carbocation stability. International Journal of Quantum Chemistry, 2022, 122, .	2.0	2
48	Effects of wave function modifications on calculated carbon–hydrogen bond lengths. Computational and Theoretical Chemistry, 2006, 760, 21-27.	1.5	1
49	O ensino de reações orgânicas usando quÃmica computacional: I. reações de adição eletrofÃłica a alquenos. Quimica Nova, 2008, 31, 1243-1249.	0.3	1
50	Effect of methylation on relative energies of tautomers and on the intramolecular proton transfer barriers of protonated nitrosamine: A <scp>MR</scp> â€ <scp>CISD</scp> study. Journal of Computational Chemistry, 2015, 36, 2027-2036.	3.3	1
51	Solvent effect on the tautomers' stabilities of protonated <i>N</i> , <i>N</i> ,ê€dimethylnitrosamine: The role of hydrogen bonds network. International Journal of Quantum Chemistry, 2017, 117, e25311.	2.0	1
52	Competition between electron transfer and base-induced elimination mechanisms in the gas-phase reactions of superoxide with alkyl hydroperoxides. Physical Chemistry Chemical Physics, 2021, 23, 5583-5595.	2.8	1
53	A multireference configuration interaction study with singles and doubles of some mesoionic rings: reaction and activation free energies for the ringâ€opening reaction. International Journal of Quantum Chemistry, 2020, 120, e26391.	2.0	0
54	A comparative multiâ€reference configuration interaction study of the lowâ€lying states of two thione isomers of thiophenol. International Journal of Quantum Chemistry, 2020, 120, e26263.	2.0	0

#	Article	IF	CITATIONS
55	A importância do método de Hartree no ensino de quÃmica quântica. Quimica Nova, 2011, 34, 527-534.	0.3	0