Soledad Gutiérrez-Oliva

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Interacting Quantum Atoms Analysis of the Reaction Force: A Tool to Analyze Driving and Retarding Forces in Chemical Reactions. ChemPhysChem, 2021, 22, 1976-1988.	1.0	7
2	Spectral Decomposition of the Reaction Force Constant. Journal of Physical Chemistry A, 2020, 124, 2372-2379.	1.1	6
3	The effect of heteroatoms in carbonaceous surfaces: computational analysis of H chemisorption on to a PANH and Si-doped PAH. Monthly Notices of the Royal Astronomical Society, 2019, 490, 172-180.	1.6	7
4	Hydrogenation and hydration of carbon dioxide: a detailed characterization of the reaction mechanisms based on the reaction force and reaction electronic flux analyses. Journal of Molecular Modeling, 2019, 25, 16.	0.8	9
5	Reaction Electronic Flux Perspective on the Mechanism of the Zimmerman Di-ï€-methane Rearrangement. Journal of Organic Chemistry, 2018, 83, 5969-5974.	1.7	11
6	An extension of the Marcus equation: the Marcus potential energy function. Journal of Molecular Modeling, 2018, 24, 104.	0.8	10
7	Molecular hydrogen formation in the interstellar medium: the role of polycyclic aromatic hydrocarbons analysed by the reaction force and activation strain model. Monthly Notices of the Royal Astronomical Society, 2018, 481, 3052-3062.	1.6	13
8	Study of antiradical mechanisms with dihydroxybenzenes using reaction force and reaction electronic flux. Physical Chemistry Chemical Physics, 2017, 19, 14512-14519.	1.3	11
9	Why Low Valent Lead(II) Hydride Complex Would be a Better Catalyst for CO ₂ Activation than Its 14 Group Analogues?. Journal of Physical Chemistry C, 2017, 121, 12127-12135.	1.5	9
10	Double Gold Activation of 1â€Ethynylâ€2â€(Phenylethynyl)Benzene Toward 5â€ <i>exo</i> â€dig and 6â€ <i>endo</i> â€dig Cyclization Reactions. Chemistry - A European Journal, 2017, 23, 13360-13368.	1.7	21
11	The Role of Coâ€Activation and Ligand Functionalization in Neutral Methallyl Nickel(II) Catalysts for Ethylene Oligomerization and Polymerization. Chemistry - A European Journal, 2017, 23, 10167-10176.	1.7	13
12	The influence of the metal cations and microhydration on the reaction trajectory of the N3 ↔ O2 thymine proton transfer: Quantum mechanical study. Journal of Computational Chemistry, 2017, 38, 2680-2692.	1.5	4
13	ETSâ€NOCV Decomposition of the Reaction Force: The HCN/CNH Isomerization Reaction Assisted by Water. Journal of Computational Chemistry, 2017, 38, 2076-2087.	1.5	19
14	Insights into the Mechanism of Ground and Excited State Double Proton Transfer Reaction in Formic Acid Dimer. Journal of Physical Chemistry A, 2017, 121, 9531-9543.	1.1	16
15	Symmetry-adapted reaction electronic flux in cycloaddition reactions. Theoretical Chemistry Accounts, 2016, 135, 1.	0.5	8
16	Role of water in intramolecular proton transfer reactions of formamide and thioformamide. Theoretical Chemistry Accounts, 2016, 135, 1.	0.5	17
17	Catalytic Mechanism of H2Activation by a Carbenoid Aluminum Complex. Journal of Physical Chemistry C, 2015, 119, 26598-26604.	1.5	27
18	Effects of the ionization in the tautomerism of uracil: A reaction electronic flux perspective. Journal of Computational Chemistry, 2015, 36, 2135-2145.	1.5	6

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19	A detailed analysis of the mechanism of a carbocationic triple shift rearrangement. Physical Chemistry Chemical Physics, 2015, 17, 9771-9779.	1.3	24
20	Insights into the chemical meanings of the reaction electronic flux. Theoretical Chemistry Accounts, 2015, 134, 1.	0.5	20
21	The performance of methallyl nickel complexes and boron adducts in the catalytic activation of ethylene: a conceptual DFT perspective. Journal of Molecular Modeling, 2015, 21, 227.	0.8	14
22	The mechanism of chemisorption of hydrogen atom on graphene: Insights from the reaction force and reaction electronic flux. Journal of Chemical Physics, 2014, 141, 134701.	1.2	27
23	Revisiting the seemingly straightforward hydrogen cyanide/hydrogen isocyanide isomerisation. Molecular Physics, 2014, 112, 349-354.	0.8	17
24	Perspectives on the Reaction Force. Advances in Quantum Chemistry, 2012, 64, 189-209.	0.4	36
25	A Relation between Different Scales of Electrophilicity: Are the Scales Consistent Along a Chemical Reaction?. Journal of Physical Chemistry A, 2012, 116, 7074-7081.	1.1	9
26	Electropolymerization of 3′,4′-disubstituted 2,2′:5′,2″-terthiophene derivatives. A theoretical and photovoltaic characterization. Journal of Molecular Modeling, 2011, 17, 81-88.	0.8	4
27	Theoretical study of the hydrogen abstraction from vitamin-E analogues. The usefulness of DFT descriptors. Journal of Molecular Modeling, 2011, 17, 593-598.	0.8	6
28	The reaction electronic flux in chemical reactions. Science China Chemistry, 2011, 54, 1982-1988.	4.2	52
29	Enhanced reactivity of Lys182 explains the limited efficacy of biogenic amines in preventing the inactivation of glucose-6-phosphate dehydrogenase by methylglyoxal. Bioorganic and Medicinal Chemistry, 2011, 19, 1613-1622.	1.4	6
30	The reaction electronic flux: A new descriptor of the electronic activity taking place during a chemical reaction. Application to the characterization of the mechanism of the Schiff's base formation in the Maillard reaction. Computational and Theoretical Chemistry, 2010, 943, 121-126.	1.5	46
31	Insights into the Maillard reaction. The mechanism of Schiff's base formation from the reaction force perspective. Molecular Physics, 2009, 107, 1587-1596.	0.8	6
32	The mechanism of double proton transfer in dimers of uracil and 2â€ŧhiouracil—The reaction force perspective. Journal of Computational Chemistry, 2009, 30, 389-398.	1.5	26
33	The reaction force and the transition region of a reaction. Journal of Molecular Modeling, 2009, 15, 707-710.	0.8	101
34	Reversibility from DFT-Based Reactivity Indices: Intramolecular Side Reactions in the Polymerization of Poly(vinyl chloride). Journal of Physical Chemistry A, 2009, 113, 7899-7908.	1.1	11
35	Reaction Force Analysis of Solvent Effects in the Addition of HCl to Propene. Journal of Physical Chemistry A, 2009, 113, 6500-6503.	1.1	32
36	Theoretical Study on a Multicenter Model Based on Different Metal Oxidation States for the Bis(imino)pyridine Iron Catalysts in Ethylene Polymerization. Organometallics, 2009, 28, 5889-5895.	1.1	43

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37	Rationalization of Diels–Alder reactions through the use of the dual reactivity descriptor Δf(r). Physical Chemistry Chemical Physics, 2008, 10, 7239.	1.3	94
38	On the Nature of the Active Site in bis(imino)Pyridyl Iron, a Catalyst for Olefin Polymerization. Journal of Physical Chemistry C, 2008, 112, 5023-5028.	1.5	34
39	Chapter 7 Using the reactivity-selectivity descriptor Δ f(r) in organic chemistry. Theoretical and Computational Chemistry, 2007, , 101-117.	0.2	8
40	Reaction Force Decomposition of Activation Barriers To Elucidate Solvent Effects. Journal of Physical Chemistry A, 2007, 111, 2455-2457.	1.1	71
41	Bridging the Gap between the Topological and Orbital Description of Hydrogen Bonding:Â The Case of the Formic Acid Dimer and Its Sulfur Derivatives. Journal of Physical Chemistry A, 2006, 110, 5102-5107.	1.1	13
42	The reaction force: Three key points along an intrinsic reaction coordinate. Journal of Chemical Sciences, 2005, 117, 467-472.	0.7	122
43	On the Mechanism of Hydrogen Transfer in the HSCH(O) ⇌ (S)CHOH and HSNO ⇌ SNOH Reactions. Journal of Physical Chemistry A, 2005, 109, 1748-1751.	1.1	93
44	The torsional problem of oxalyl chloride: a challenge for theoretical methods. Chemical Physics Letters, 2004, 383, 435-440.	1.2	17
45	Analysis of two intramolecular proton transfer processes in terms of the reaction force. Journal of Chemical Physics, 2004, 121, 4570-4576.	1.2	78
46	Towards understanding the molecular internal rotations and vibrations and chemical reactions through the profiles of reactivity and selectivity indices: an ab initio SCF and DFT study. Molecular Physics, 2003, 101, 2841-2853.	0.8	42
47	Using Sanderson's Principle to Estimate Global Electronic Properties and Bond Energies of Hydrogen-Bonded Complexes. Journal of Physical Chemistry A, 2000, 104, 8955-8964.	1.1	57
48	Energy, chemical potential and hardness profiles for the rotational isomerization of HOOH, HSOH and HSSH. Molecular Physics, 1999, 96, 61-70.	0.8	47
49	Relations between Potential Energy, Electronic Chemical Potential, and Hardness Profiles. Journal of Physical Chemistry A, 1997, 101, 4621-4627.	1.1	45