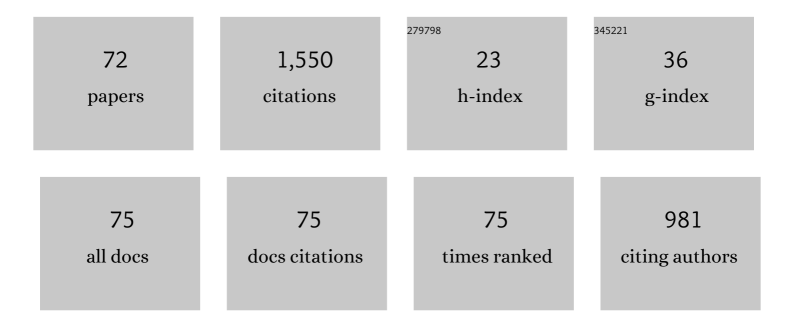
Carlos Cardenas

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/2705342/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	On the Prediction of Lattice Energy with the Fukui Potential: Some Supports on Hardness Maximization in Inorganic Solids. Journal of Physical Chemistry A, 2022, 126, 4507-4516.	2.5	13
2	Orbital energies and nuclear forces in <scp>DFT</scp> : Interpretation and validation. Journal of Computational Chemistry, 2021, 42, 334-343.	3.3	7
3	Designing boron and metal complexes for fluoride recognition: a computational perspective. Physical Chemistry Chemical Physics, 2021, 23, 22768-22778.	2.8	0
4	The change in the nature of bonding in the Li 2 dimer under confinement. International Journal of Quantum Chemistry, 2021, 121, e26644.	2.0	3
5	Coulomb Explosion of Multi-charged Atomic Alkaline Metal Clusters. Journal of Physical Chemistry A, 2021, 125, 2512-2517.	2.5	1
6	Understanding Topological Insulators in Real Space. Molecules, 2021, 26, 2965.	3.8	1
7	Links among the Fukui potential, the alchemical hardness and the local hardness of an atom in a molecule. Journal of Computational Chemistry, 2021, 42, 1681-1688.	3.3	10
8	Kick–Fukui: A Fukui Function-Guided Method for Molecular Structure Prediction. Journal of Chemical Information and Modeling, 2021, 61, 3955-3963.	5.4	14
9	On the nature of bonding in the photochemical addition of two ethylenes: C–C bond formation in the excited state?. Physical Chemistry Chemical Physics, 2021, 23, 20598-20606.	2.8	10
10	Electronegativity under Confinement. Molecules, 2021, 26, 6924.	3.8	2
11	Cluster Assembled Silicon-Lithium Nanostructures: A Nanowire Confined Inside a Carbon Nanotube. Frontiers in Chemistry, 2021, 9, 767421.	3.6	1
12	Electronic structure of first and second row atoms under harmonic confinement. International Journal of Quantum Chemistry, 2020, 120, e26132.	2.0	6
13	On the recognition of chloride, bromide and nitrate anions by anthracene–squaramide conjugated compounds: a computational perspective. New Journal of Chemistry, 2020, 44, 17831-17839.	2.8	9
14	On the chemical potential of the hydrogen atom. Theoretical Chemistry Accounts, 2020, 139, 1.	1.4	0
15	Computing the Fukui Function in Solid-State Chemistry: Application to Alkaline Earth Oxides Bulk and Surfaces. Journal of Physical Chemistry A, 2020, 124, 2826-2833.	2.5	20
16	Predicting Deprotonation Sites Using Alchemical Derivatives. Journal of Physical Chemistry A, 2020, 124, 3754-3760.	2.5	13
17	Magnon valley Hall effect in <mml:math xmlns:mml="http://www.w3.org/1998/Math/MathML"><mml:msub><mml:mrow><mml:mi>CrI</mml:mi>-based van der Waals heterostructures. Physical Review B, 2020, 101, .</mml:mrow></mml:msub></mml:math 	ຠ r ໝw> <mi< td=""><td>ന്മതന>3<!--ന</td--></td></mi<>	ന ്മത ന>3 ന</td

18 Interaction of Nitroxide Radicals with an Au₈ Nanostructure: Theoretical and Calorimetric Studies. Journal of Physical Chemistry C, 2019, 123, 21713-21720.

3.1 4

CARLOS CARDENAS

#	Article	IF	CITATIONS
19	Electron Localization Function in Excited States: The Case of the Ultrafast Proton Transfer of the Salicylidene Methylamine. Journal of Chemical Theory and Computation, 2019, 15, 5532-5542.	5.3	14
20	Reactivity of Carbon Molecular Clusters from a Hückel-Type Model. Journal of Physical Chemistry A, 2019, 123, 8696-8701.	2.5	3
21	A chemical theory of topological insulators. Chemical Communications, 2019, 55, 12281-12287.	4.1	16
22	Formation of complex organic molecules in ice mantles: An ab initio molecular dynamics study. Astronomy and Astrophysics, 2019, 629, A28.	5.1	7
23	Hematene: a 2D magnetic material in van der Waals or non-van der Waals heterostructures. 2D Materials, 2019, 6, 045002.	4.4	24
24	The Pauli principle and the confinement of electron pairs in a double well: Aspects of electronic bonding under pressure. Journal of Chemical Physics, 2019, 150, 204304.	3.0	9
25	Theoretical analysis of the adsorption of ammonia–borane and their dehydrogenation products on the (001) surface of TiC and ZrC. Surface Science, 2019, 680, 95-106.	1.9	9
26	Molecular simulations of carbon allotropes in processes with creation and destruction of chemical bonds. Carbon, 2019, 144, 177-184.	10.3	8
27	A theoretical study of the photodynamics of salicylidene-2-anthrylamine in acetonitrile solution. Physical Chemistry Chemical Physics, 2018, 20, 29399-29411.	2.8	10
28	Local electrophilicity. Journal of Molecular Modeling, 2018, 24, 245.	1.8	21
29	Proposal of a simple and effective local reactivity descriptor through a topological analysis of an orbitalâ€weighted fukui function. Journal of Computational Chemistry, 2017, 38, 481-488.	3.3	58
30	How predictive could alchemical derivatives be?. Physical Chemistry Chemical Physics, 2017, 19, 16003-16012.	2.8	34
31	Negative Condensed-to-Atom Fukui Functions: A Signature of Oxidation-Induced Reduction of Functional Groups. , 2017, , 269-278.		7
32	On understanding the chemical origin of band gaps. Journal of Molecular Modeling, 2017, 23, 271.	1.8	9
33	The HSAB principle from a finite-temperature grand-canonical perspective. Theoretical Chemistry Accounts, 2017, 136, 1.	1.4	28
34	Ferromagnetic bond of Li10 cluster: An alternative approach in terms of effective ferromagnetic sites. Journal of Chemical Physics, 2016, 145, 094301.	3.0	3
35	Topological Analysis of the Fukui Function. Challenges and Advances in Computational Chemistry and Physics, 2016, , 227-241.	0.6	19
36	Benchmark values of chemical potential and chemical hardness for atoms and atomic ions (including) Tj ETQq	0 0 0 rgBT /0 2.8	Dverlock 10 1 51

^{18, 25721-25734.}

CARLOS CARDENAS

#	Article	IF	CITATIONS
37	An explicit approach to conceptual density functional theory descriptors of arbitrary order. Chemical Physics Letters, 2016, 660, 307-312.	2.6	46
38	A numerical study of the Lieb–Thirring kinetic energy lower bound. Molecular Physics, 2016, 114, 982-987.	1.7	0
39	A problematic issue for atoms in molecules: Impact of (quasi-)degenerate states on Quantum Theory Atoms in Molecules and Hirshfeld-I properties. Computational and Theoretical Chemistry, 2015, 1053, 106-111.	2.5	15
40	Reaction channels and spectroscopic constants of astrophysical relevant Silicon bearing molecules SiC3H,+ and SiC3H. Monthly Notices of the Royal Astronomical Society, 2014, 443, 3127-3133.	4.4	1
41	Density functional theory of chemical reactivity. Chemical Modelling, 2014, , 151-174.	0.4	30
42	Mechanism of fluorophore quenching in a pre-fluorescent nitroxide probe: A theoretical illustration. Chemical Physics Letters, 2014, 593, 89-92.	2.6	18
43	In pursuit of negative Fukui functions: molecules with very small band gaps. Journal of Molecular Modeling, 2014, 20, 2162.	1.8	22
44	How to Compute the Fukui Matrix and Function for Systems with (Quasi-)Degenerate States. Journal of Chemical Theory and Computation, 2014, 10, 202-210.	5.3	53
45	An information-theoretic resolution of the ambiguity in the local hardness. Physical Chemistry Chemical Physics, 2014, 16, 6019-6026.	2.8	34
46	<i>Ab Initio</i> Molecular Dynamics Study of Small Alkali Metal Clusters. Journal of Physical Chemistry A, 2014, 118, 1077-1083.	2.5	8
47	Regional Electrophilic and Nucleophilic Fukui Functions Efficiently Highlight the Lewis Acidic/Basic Regions in Ionic Liquids. Journal of Physical Chemistry B, 2014, 118, 3696-3701.	2.6	14
48	In pursuit of negative Fukui functions: examples where the highest occupied molecular orbital fails to dominate the chemical reactivity. Journal of Molecular Modeling, 2013, 19, 2779-2783.	1.8	43
49	How reliable is the hard–soft acid–base principle? An assessment from numerical simulations of electron transfer energies. Physical Chemistry Chemical Physics, 2013, 15, 13959.	2.8	44
50	On the exponential model for energy with respect to number of electrons. Journal of Molecular Modeling, 2013, 19, 2849-2853.	1.8	29
51	Performance of modified Lennard–Jones potential to seed ab initio calculations of small cadmium clusters. Computational and Theoretical Chemistry, 2013, 1021, 249-255.	2.5	8
52	Communication: A case where the hard/soft acid/base principle holds regardless of acid/base strength. Journal of Chemical Physics, 2013, 138, 181106.	3.0	31
53	Atomic Charges and the Electrostatic Potential Are Ill-Defined in Degenerate Ground States. Journal of Chemical Theory and Computation, 2013, 9, 4779-4788.	5.3	48
54	Ab Initio Molecular Dynamics Simulations of Ti ₂ on C ₂₀ Collisions and C ₂₀ Ti ₂ Configurations. Journal of Physical Chemistry C, 2013, 117, 4287-4291.	3.1	1

CARLOS CARDENAS

#	Article	IF	CITATIONS
55	Symmetric Nonlocal Weighted Density Approximations from the Exchange-Correlation Hole of the Uniform Electron Gas. Journal of Chemical Theory and Computation, 2012, 8, 4081-4093.	5.3	22
56	Understanding chemical binding using the Berlin function and the reaction force. Chemical Physics Letters, 2012, 539-540, 168-171.	2.6	11
57	A new isomer of C20 and a way to a new C240. Physical Chemistry Chemical Physics, 2012, 14, 14810.	2.8	6
58	Reactivity indicators for degenerate states in the density-functional theoretic chemical reactivity theory. Journal of Chemical Physics, 2011, 134, 174103.	3.0	74
59	Assembling Small Silicon Clusters Using Criteria of Maximum Matching of the Fukui Functions. Journal of Chemical Theory and Computation, 2011, 7, 3995-4001.	5.3	20
60	Should negative electron affinities be used for evaluating the chemical hardness?. Physical Chemistry Chemical Physics, 2011, 13, 2285-2293.	2.8	69
61	The Fukui potential is a measure of the chemical hardness. Chemical Physics Letters, 2011, 513, 127-129.	2.6	34
62	The Fukui Potential and the Capacity of Charge and the Global Hardness of Atoms. Journal of Physical Chemistry A, 2011, 115, 2325-2331.	2.5	56
63	Relationships between the third-order reactivity indicators in chemical density-functional theory. Journal of Chemical Physics, 2009, 130, 244105.	3.0	35
64	Chemical Reactivity Descriptors for Ambiphilic Reagents: Dual Descriptor, Local Hypersoftness, and Electrostatic Potential. Journal of Physical Chemistry A, 2009, 113, 8660-8667.	2.5	166
65	Theoretical study of the surface reactivity of alkaline earth oxides: Local density of states evaluation of the local softness. Journal of Chemical Physics, 2008, 128, 034708.	3.0	31
66	Theoretical Study on CDK2 Inhibitors Using a Global Softness Obtained from the Density of States. Journal of Physical Chemistry B, 2007, 111, 3293-3297.	2.6	12
67	Further exploration of the Fukui function, hardness, and other reactivity indices and its relationships within the Kohn–Sham scheme. International Journal of Quantum Chemistry, 2007, 107, 37-45.	2.0	27
68	Nuclear Fukui functions from nonintegral electron number calculations. International Journal of Quantum Chemistry, 2007, 107, 807-815.	2.0	11
69	Nuclear reactivity indices in the context of spin polarized density functional theory. Chemical Physics, 2006, 322, 303-310.	1.9	16
70	Condensation of the highest occupied molecular orbital within the electron localization function domains. Journal of Chemical Sciences, 2005, 117, 419-424.	1.5	13
71	Nature of Bonding in the Cyclization Reactions of (2-Ethynylphenyl)triazene and 2-Ethynylstyrene. Journal of Physical Chemistry A, 2005, 109, 4352-4358.	2.5	23
72	Molecular Interactions From the Density Functional Theory for Chemical Reactivity: The Interaction Energy Between Two-Reagents. Frontiers in Chemistry, 0, 10, .	3.6	6