

# Rajendra R Zope

## List of Publications by Year in descending order

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81  
papers

1,616  
citations

257450  
24  
h-index

361022  
35  
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81  
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81  
docs citations

81  
times ranked

1190  
citing authors

#	ARTICLE	IF	CITATIONS
1	Snub boron nanostructures: Chiral fullerenes, nanotubes and planar sheet. <i>Chemical Physics Letters</i> , 2011, 501, 193-196.	2.6	69
2	Boron fullerenes: From $\text{B}_{80}$ to $\text{B}_{216}$ hole doped boron sheets. <i>Physical Review B</i> , 2009, 79, .	4.7	67
3	$\text{C}_{60}$ to $\text{C}_{216}$ characterized by an all-electron density functional theory. <i>Physical Review B</i> , 2008, 77, .	3.2	58
4	Self-interaction error overbinds water clusters but cancels in structural energy differences. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2020, 117, 11283-11288.	7.1	57
5	A step in the direction of resolving the paradox of Perdew-Zunger self-interaction correction. <i>Journal of Chemical Physics</i> , 2019, 151, 214108.	3.0	56
6	Electronic structure, vibrational stability, infra-red, and Raman spectra of $\text{B}_{24}\text{N}_{24}$ cages. <i>Chemical Physics Letters</i> , 2004, 393, 300-304.	2.6	50
7	Temperature Dependence of the Polarizability of Sodium Clusters. <i>Physical Review Letters</i> , 2000, 84, 4826-4829.	7.8	49
8	Efficient quantum-chemical geometry optimization and the structure of large icosahedral fullerenes. <i>Chemical Physics Letters</i> , 2006, 422, 451-454.	2.6	49
9	Are hemispherical caps of boron-nitride nanotubes possible?. <i>Chemical Physics Letters</i> , 2004, 386, 403-407.	2.6	47
10	Vibrational stability and electronic structure of a $\text{B}_{80}$ cage. <i>Physical Review B</i> , 2008, 78, .	3.2	47
11	Stretched or noded orbital densities and self-interaction correction in density functional theory. <i>Journal of Chemical Physics</i> , 2019, 150, 174102.	3.0	46
12	Electronic structure of fullerene-like cages and finite nanotubes of aluminum nitride. <i>Physical Review B</i> , 2005, 72, .	3.2	38
13	Fermi-Làwdin orbital self-interaction correction using the strongly constrained and appropriately normed meta-GGA functional. <i>Journal of Chemical Physics</i> , 2019, 151, 154105.	3.0	38
14	Charge transfer excited state energies by perturbative delta self-consistent field method. <i>Journal of Chemical Physics</i> , 2012, 137, 084316.	3.0	35
15	Stability of $\text{As}_n$ ( $n=4, 8, 20, 28, 32, 36, 60$ ) cage structures. <i>Chemical Physics Letters</i> , 2004, 387, 476-480.	2.6	33
16	Fermi-Làwdin orbital self-interaction correction to magnetic exchange couplings. <i>Journal of Chemical Physics</i> , 2018, 149, 164101.	3.0	33
17	Importance of self-interaction-error removal in density functional calculations on water cluster anions. <i>Physical Chemistry Chemical Physics</i> , 2020, 22, 3789-3799.	2.8	32
18	Shrinking Self-Interaction Errors with the Fermi-Làwdin Orbital Self-Interaction-Corrected Density Functional Approximation. <i>Journal of Physical Chemistry A</i> , 2018, 122, 9307-9315.	2.5	30

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19	Structural and bonding properties of bcc-based $\text{B}_{80}$ . <i>Physical Review B</i> , 2008, 78, .	3.2	29
20	The effect of self-interaction error on electrostatic dipoles calculated using density functional theory. <i>Journal of Chemical Physics</i> , 2019, 151, 174106.	3.0	29
21	The $\text{B}_4$ -boron cages with four-member rings. <i>Europhysics Letters</i> , 2009, 85, 68005.	2.0	28
22	Self-interaction-free electric dipole polarizabilities for atoms and their ions using the Fermi-Lindgren self-interaction correction. <i>Physical Review A</i> , 2019, 100, .	2.5	27
23	Charge transfer excitations in cofacial fullerene-porphyrin complexes. <i>Journal of Chemical Physics</i> , 2012, 137, 084317.	3.0	26
24	$\text{Al}_{12}$ Superatom as Stable Building Block of Ionic Salts. <i>Journal of Physical Chemistry C</i> , 2015, 119, 5129-5137.	3.1	25
25	Theoretical infrared, Raman, and optical spectra of the $\text{B}_{36}\text{N}_{36}$ cage. <i>Physical Review A</i> , 2005, 71, .	2.5	24
26	Improvements in the orbitalwise scaling down of Perdew-Zunger self-interaction correction in many-electron regions. <i>Journal of Chemical Physics</i> , 2020, 152, 174112.	3.0	23
27	A step in the direction of resolving the paradox of Perdew-Zunger self-interaction correction. II. Gauge consistency of the energy density at three levels of approximation. <i>Journal of Chemical Physics</i> , 2020, 152, 214109.	3.0	23
28	On the Question of the Total Energy in the Fermi-Lindgren Orbital Self-Interaction Correction Method. <i>Journal of Chemical Theory and Computation</i> , 2018, 14, 4122-4128.	5.3	22
29	Study of self-interaction errors in density functional predictions of dipole polarizabilities and ionization energies of water clusters using Perdew-Zunger and locally scaled self-interaction corrected methods. <i>Journal of Chemical Physics</i> , 2020, 153, 164304.	3.0	21
30	Density functional study of structural and electronic properties of $\text{Na}_{n}\text{Mg}_{(1-n)/2}$ clusters. <i>Journal of Chemical Physics</i> , 2001, 115, 2109-2116.	3.0	19
31	Slater's Exchange Parameters $\alpha_s$ for Analytic and Variational X $\alpha$ Calculations. <i>Journal of Chemical Theory and Computation</i> , 2005, 1, 1193-1200.	5.3	19
32	The limitations of Slater's element-dependent exchange functional from analytic density-functional theory. <i>Journal of Chemical Physics</i> , 2006, 124, 044107.	3.0	18
33	The static dipole polarizability of $\text{C}_{70}$ fullerene. <i>Journal of Physics B: Atomic, Molecular and Optical Physics</i> , 2007, 40, 3491-3496.	1.5	18
34	Low-lying planar isomers of neutral and charged $\text{B}_{22}$ clusters. <i>Journal of Physics B: Atomic, Molecular and Optical Physics</i> , 2012, 45, 225101.	1.5	17
35	The electronic structure and charge transfer excited states of the endohedral trimetallic nitride $\text{C}_{80}(\text{Li}_h)$ fullerenes-Zn-tetraphenyl porphyrin dyads. <i>Physical Chemistry Chemical Physics</i> , 2015, 17, 5832-5839.	2.8	17
36	Full-potential LAPW calculation of electron momentum density and related properties of Li. <i>Physical Review B</i> , 1999, 60, 10770-10775.	3.2	16

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37	Dipole polarizability of isovalent carbon and boron cages and fullerenes. Physical Review B, 2009, 80, .	3.2	16
38	Analytic atomic gradients in the fermi-LÀ¶wdin orbital selfâ€‘interaction correction. Journal of Computational Chemistry, 2019, 40, 820-825.	3.3	16
39	Accurate molecular energies by extrapolation of atomic energies using an analytic quantum mechanical model. Physical Review B, 2005, 71, .	3.2	14
40	Comparative study of unscreened and screened molecular static linear polarizability in the Hartree-Fock, hybridâ€‘density functional, and density functional models. International Journal of Quantum Chemistry, 2008, 108, 307-317.	2.0	14
41	Optical excitation energies, Stokes shift, and spin-splitting of C <sub>24</sub> H <sub>72</sub> Si <sub>14</sub> . Journal of Chemical Physics, 2010, 133, 034301.	3.0	14
42	Crystalline Alloys of Organic Donors and Acceptors Based on TIPS-Pentacene. Journal of Physical Chemistry C, 2015, 119, 20823-20832.	3.1	14
43	Local self-interaction correction method with a simple scaling factor. Physical Chemistry Chemical Physics, 2021, 23, 2406-2418.	2.8	14
44	Fermi-LÀ¶wdin-orbital self-interaction correction using the optimized-effective-potential method within the Krieger-Li-lafrate approximation. Physical Review A, 2021, 103, .	2.5	14
45	How well do self-interaction corrections repair the overestimation of static polarizabilities in density functional calculations?. Physical Chemistry Chemical Physics, 2021, 23, 18678-18685.	2.8	14
46	Geometry and electronic structure of neutral and charged B <sub>21</sub> clusters. Chemical Physics Letters, 2013, 557, 15-18.	2.6	13
47	Atomic Compton profiles within different exchange-only theories. European Physical Journal D, 1999, 7, 151-155.	1.3	12
48	Static dipole polarizabilities of polyacenes using self-interaction-corrected density functional approximations. Journal of Chemical Physics, 2021, 154, 114305.	3.0	12
49	Exploring and enhancing the accuracy of interior-scaled Perdewâ€‘Zunger self-interaction correction. Journal of Chemical Physics, 2021, 154, 094105.	3.0	12
50	Study of self-interaction-errors in barrier heights using locally scaled and Perdewâ€‘Zunger self-interaction methods. Journal of Chemical Physics, 2022, 156, 014306.	3.0	12
51	Momentum-space properties of atoms: Application of the generalized-gradient approximation. Physical Review A, 2000, 62, .	2.5	11
52	Calcium coated B <sub>80</sub> fullerene: A study on various coating configurations of B <sub>80</sub> . Chemical Physics Letters, 2011, 514, 66-69.	2.6	11
53	Site specific atomic polarizabilities in endohedral fullerenes and carbon onions. Journal of Chemical Physics, 2015, 143, 084306.	3.0	11
54	Electronic and Structural Properties of C <sub>60</sub> and Sc <sub>3</sub> N@C <sub>80</sub> Supported on Graphene Nanoflakes. Journal of Physical Chemistry C, 2016, 120, 26083-26092.	3.1	11

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55	Donor-fullerene dyads for energy cascade organic solar cells. <i>Inorganica Chimica Acta</i> , 2017, 468, 192-202.	2.4	10
56	A DFT analysis of the ground and charge-transfer excited states of Sc <sub>3</sub> N@I <sub>h</sub> -C <sub>80</sub> fullerene coupled with metal-free and zinc-phthalocyanine. <i>Physical Chemistry Chemical Physics</i> , 2018, 20, 25841-25848.	2.8	10
57	Positron binding: A positron-density viewpoint. <i>Physical Review A</i> , 1994, 50, 2191-2196.	2.5	9
58	Electronic structure, vibrational stability, and predicted infrared-Raman spectra of the As <sub>20</sub> , As@Ni <sub>12</sub> , and As@Ni <sub>12</sub> @As <sub>20</sub> clusters. <i>Journal of Chemical Physics</i> , 2004, 121, 11007.	3.0	9
59	Momentum-space properties from coordinate-space electron density. <i>Journal of Chemical Physics</i> , 2005, 122, 204110.	3.0	9
60	The effect of structural changes on charge transfer states in a light-harvesting carotenoid-diaryl-porphyrin-C <sub>60</sub> molecular triad. <i>Journal of Chemical Physics</i> , 2014, 140, 204309.	3.0	8
61	Density-related properties from self-interaction corrected density functional theory calculations. <i>Journal of Chemical Physics</i> , 2021, 154, 024102.	3.0	8
62	Self-interaction-corrected Kohn-Sham effective potentials using the density-consistent effective potential method. <i>Journal of Chemical Physics</i> , 2021, 155, 064109.	3.0	8
63	On the optimal value of $\hat{t} \pm$ for the Hartree-Fock-Slater method. <i>Chemical Physics Letters</i> , 2004, 399, 417-421.	2.6	7
64	Density functional study of the electronic structure of dye-functionalized fullerenes and their model donor-acceptor complexes containing P3HT. <i>Journal of Chemical Physics</i> , 2016, 144, 144304.	3.0	7
65	Electronic and Optical Properties of VSc <sub>2</sub> N@C <sub>68</sub> Fullerene. <i>Journal of Physical Chemistry C</i> , 2016, 120, 27813-27819.	3.1	7
66	Implementation of Perdew-Zunger self-interaction correction in real space using Fermi-Wedin orbitals. <i>Journal of Chemical Physics</i> , 2021, 154, 084112.	3.0	7
67	Electronic and Structural Study of Zn <sub>i</sub> x <sub>i</sub> S <sub>i</sub> [i = 12, 16, 24, 28, 36, 48, 96, and 108] Cage Structures. <i>Journal of Physical Chemistry A</i> , 2017, 121, 3486-3493.	2.5	6
68	Assessing the effect of regularization on the molecular properties predicted by SCAN and self-interaction corrected SCAN meta-GGA. <i>Physical Chemistry Chemical Physics</i> , 2020, 22, 18060-18070.	2.8	6
69	Study of Self-Interaction Errors in Density Functional Calculations of Magnetic Exchange Coupling Constants Using Three Self-Interaction Correction Methods. <i>Journal of Physical Chemistry A</i> , 2022, 126, 1923-1935.	2.5	6
70	Total atomic energies using indirect-path methods. <i>Physical Review A</i> , 1996, 53, 3652-3655.	2.5	5
71	Full-potential LAPW calculation of magnetic Compton profiles of Ni. <i>Physical Review B</i> , 2000, 62, 16435-16441.	3.2	5
72	Smooth scaling of valence electronic properties in fullerenes: From one carbon atom, to C <sub>60</sub> , to graphene. <i>Physical Review A</i> , 2013, 87, .	2.5	5

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73	Diels-Alder addition to H <sub>2</sub> O@C <sub>60</sub> an electronic and structural study. <i>Chemical Physics Letters</i> , 2017, 685, 198-204.	2.6	4
74	Dipole moments from atomic-number-dependent potentials in analytic density-functional theory. <i>Journal of Chemical Physics</i> , 2006, 125, 214104.	3.0	3
75	Magnetic structure, excitations and short-range order in honeycomb Na <sub>2</sub> Ni <sub>2</sub> TeO <sub>6</sub> . <i>Journal of Physics Condensed Matter</i> , 2021, 33, 375803.	1.8	3
76	Leading corrections to the compton profiles beyond the impulse approximation: second-order correction. <i>Chemical Physics Letters</i> , 1995, 242, 555-559.	2.6	2
77	Electronic structure calculation of vanadium- and scandium-based endohedral fullerenes VSc <sub>2</sub> N@C <sub>2n</sub> (2<math>n</math> = 70, 76, 78, 80). <i>International Journal of Quantum Chemistry</i> , 2018, 118, e25785.	2.0	2
78	Positron and positronium affinities in the work-formalism Hartree-Fock approximation. <i>Physical Review A</i> , 1999, 60, 218-223.	2.5	1
79	Fully Analytic Implementation of Density Functional Theory for Efficient Calculations on Large Molecules. , 0, , 157-168.		1
80	Excited Electronic States of Porphyrin-Based Assemblies Using Density Functional Theory. , 2016, , 233-289.		1
81	Polarizabilities of intermediate sized lithium clusters from density-functional theory. <i>Journal of Computational Methods in Sciences and Engineering</i> , 2008, 7, 495-505.	0.2	0