## Sijia S Dong

List of Publications by Year in descending order

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SULA S DONG

#	Article	IF	CITATIONS
1	Mechanism of halide exchange in reactions of CpRu(PPh <sub>3</sub> ) <sub>2</sub> Cl with haloalkanes. New Journal of Chemistry, 2022, 46, 6603-6608.	2.8	1
2	Targeting oncogenic KRAS with molecular brush-conjugated antisense oligonucleotides. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119, .	7.1	14
3	Machine learning dielectric screening for the simulation of excited state properties of molecules and materials. Chemical Science, 2021, 12, 4970-4980.	7.4	16
4	Nature of the 1 <sup>1</sup> B <sub>u</sub> and 2 <sup>1</sup> A <sub>g</sub> Excited States of Butadiene and the Goldilocks Principle of Basis Set Diffuseness. Journal of Chemical Theory and Computation, 2019, 15, 4591-4601.	5.3	15
5	OpenMolcas: From Source Code to Insight. Journal of Chemical Theory and Computation, 2019, 15, 5925-5964.	5.3	661
6	State-Interaction Pair-Density Functional Theory Can Accurately Describe a Spiro Mixed Valence Compound. Journal of Physical Chemistry A, 2019, 123, 2100-2106.	2.5	12
7	Assessment of MC-PDFT Excitation Energies for a Set of QM/MM Models of Rhodopsins. Journal of Chemical Theory and Computation, 2019, 15, 1915-1923.	5.3	16
8	Automatic Selection of an Active Space for Calculating Electronic Excitation Spectra by MS-CASPT2 or MC-PDFT. Journal of Chemical Theory and Computation, 2018, 14, 2017-2025.	5.3	45
9	Excitation spectra of retinal by multiconfiguration pair-density functional theory. Physical Chemistry Chemical Physics, 2018, 20, 7265-7276.	2.8	13
10	Extended Hamiltonian molecular dynamics: semiclassical trajectories with improved maintenance of zero point energy. Physical Chemistry Chemical Physics, 2018, 20, 30209-30218.	2.8	6
11	Identifying multiple active conformations in the G protein-coupled receptor activation landscape using computational methods. Methods in Cell Biology, 2017, 142, 173-186.	1.1	3
12	Conformational and Thermodynamic Landscape of GPCR Activation from Theory and Computation. Biophysical Journal, 2016, 110, 2618-2629.	0.5	18
13	The Predicted Ensemble of Lowâ€Energy Conformations of Human Somatostatin Receptor Subtypeâ€5 and the Binding of Antagonists. ChemMedChem, 2015, 10, 650-661.	3.2	6
14	Computational predictions of corroles as a class of Hsp90 inhibitors. Molecular BioSystems, 2015, 11, 2907-2914.	2.9	11
15	Computational Prediction of the Class a GPCR Active State Conformations. Biophysical Journal, 2014, 106, 308a.	0.5	0
16	Electronic Structures of Group 9 Metallocorroles with Axial Ammines. Inorganic Chemistry, 2011, 50, 764-770.	4.0	18
17	Highly Efficient and Diastereoselective Gold(I) atalyzed Synthesis of Tertiary Amines from Secondary Amines and Alkynes: Substrate Scope and Mechanistic Insights. Chemistry - A European Journal, 2011, 17, 12932-12945.	3.3	42