Lando P Wolters

List of Publications by Year in descending order

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Version: 2024-02-01

471509 552781 1,351 25 17 26 citations h-index g-index papers 31 31 31 1457 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Câ^'X Bond Activation by Palladium: Steric Shielding versus Steric Attraction. Chemistry - A European Journal, 2022, 28, .	3.3	11
2	Cooperative Selfâ€Assembly in Linear Chains Based on Halogen Bonds. ChemPlusChem, 2021, 86, 812-819.	2.8	10
3	Nucleophilic Substitution in Solution: Activation Strain Analysis of Weak and Strong Solvent Effects. Chemistry - A European Journal, 2018, 24, 5927-5938.	3.3	53
4	Asymmetric identity SN2 transition states: Nucleophilic substitution at \hat{l}_{\pm} -substituted carbon and silicon centers. International Journal of Mass Spectrometry, 2017, 413, 85-91.	1.5	16
5	Mechanistic Insight into the Oxidation of Organic Phenylselenides by H ₂ O ₂ . Chemistry - A European Journal, 2017, 23, 2405-2422.	3.3	56
6	Enhanced Ï€â€Backâ€Donation as a Way to Higher Coordination Numbers in d ¹⁰ [M(NHC) _{<i>n</i>}] Complexes: A DFT Study. Chemistry - A European Journal, 2017, 23, 614-622.	3.3	17
7	Insights on selenium and tellurium diaryldichalcogenides: A benchmark DFT study. Journal of Computational Chemistry, 2016, 37, 1672-1680.	3.3	43
8	Addition–Elimination or Nucleophilic Substitution? Understanding the Energy Profiles for the Reaction of Chalcogenolates with Dichalcogenides. Journal of Chemical Theory and Computation, 2016, 12, 2752-2761.	5. 3	37
9	General AMBER Force Field Parameters for Diphenyl Diselenides and Diphenyl Ditellurides. Journal of Physical Chemistry A, 2016, 120, 4389-4400.	2.5	22
10	The activation strain model and molecular orbital theory. Wiley Interdisciplinary Reviews: Computational Molecular Science, 2015, 5, 324-343.	14.6	280
11	Selective Câ°'H and Câ°'C Bond Activation: Electronic Regimes as a Tool for Designing d ¹⁰ ML _{<i>n</i>} Catalysts. Chemistry - an Asian Journal, 2015, 10, 2272-2282.	3.3	13
12	Role of Steric Attraction and Bite-Angle Flexibility in Metal-Mediated C–H Bond Activation. ACS Catalysis, 2015, 5, 5766-5775.	11.2	58
13	Covalency in resonance-assisted halogen bonds demonstrated with cooperativity in N-halo-guanine quartets. Physical Chemistry Chemical Physics, 2015, 17, 1585-1592.	2.8	54
14	Peroxidase Activity of Organic Selenides: Mechanistic Insights from Quantum Chemistry. Current Organic Chemistry, 2015, 20, 189-197.	1.6	33
15	d10-ML2 Complexes: Structure, Bonding, and Catalytic Activity. Structure and Bonding, 2014, , 139-161.	1.0	1
16	New Concepts for Designing d ¹⁰ â€M(L) _{<i>n</i>} Catalysts: d Regime, s Regime and Intrinsic Biteâ€Angle Flexibility. Chemistry - A European Journal, 2014, 20, 11370-11381.	3.3	36
17	Understanding E2 versus S _N 2 Competition under Acidic and Basic Conditions. ChemistryOpen, 2014, 3, 29-36.	1.9	40
18	Controlling the oxidative addition of aryl halides to Au(I). Journal of Computational Chemistry, 2014, 35, 2140-2145.	3.3	65

#	Article	IF	CITATIONS
19	The many faces of halogen bonding: a review of theoretical models and methods. Wiley Interdisciplinary Reviews: Computational Molecular Science, 2014, 4, 523-540.	14.6	188
20	Nonlinear d ¹⁰ â€ML ₂ Transitionâ€Metal Complexes. ChemistryOpen, 2013, 2, 106-114.	1.9	52
21	In Silico Design of Heteroaromatic Halfâ€Sandwich Rh ^I Catalysts for Acetylene [2+2+2] Cyclotrimerization: Evidence of a Reverse Indenyl Effect. Chemistry - A European Journal, 2013, 19, 13337-13347.	3.3	27
22	Nonlinear d ¹⁰ â€ML ₂ Transitionâ€Metal Complexes. ChemistryOpen, 2013, 2, 78-78.	1.9	0
23	Halogen Bonding versus Hydrogen Bonding: A Molecular Orbital Perspective. ChemistryOpen, 2012, 1, 96-105.	1.9	185
24	Alkali-Metal-Supported Bismuth Polyhedraâ€"Principles and Theoretical Studies. Inorganic Chemistry, 2011, 50, 5755-5762.	4.0	8
25	Reaction Coordinates and the Transition-Vector Approximation to the IRC. Journal of Chemical Theory and Computation, 2008, 4, 920-928.	5.3	43