Rees B Rankin

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/7982512/publications.pdf

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34 1,797 18 32 g-index

43 43 43 2658 all docs docs citations times ranked citing authors

#	Article	IF	CITATIONS
1	Progress, Opportunities, and Challenges for Applying Atomically Detailed Modeling to Molecular Adsorption and Transport in Metalâ 'Organic Framework Materials. Industrial & Engineering Chemistry Research, 2009, 48, 2355-2371.	3.7	283
2	Unique Electrochemical Adsorption Properties of Ptâ€Skin Surfaces. Angewandte Chemie - International Edition, 2012, 51, 3139-3142.	13.8	264
3	Trends in Selective Hydrogen Peroxide Production on Transition Metal Surfaces from First Principles. ACS Catalysis, 2012, 2, 2664-2672.	11.2	137
4	Rational Development of Ternary Alloy Electrocatalysts. Journal of Physical Chemistry Letters, 2012, 3, 1668-1673.	4.6	130
5	Adsorption and Diffusion of Light Gases in ZIF-68 and ZIF-70: A Simulation Study. Journal of Physical Chemistry C, 2009, 113, 16906-16914.	3.1	126
6	Elucidation of Pathways for NO Electroreduction on $Pt(111)$ from First Principles. Angewandte Chemie - International Edition, 2015, 54, 8255-8258.	13.8	111
7	Structures of Glycine, Enantiopure Alanine, and Racemic Alanine Adlayers on Cu(110) and Cu(100) Surfaces. Journal of Physical Chemistry B, 2005, 109, 16764-16773.	2.6	101
8	Assessment of heterochiral and homochiral glycine adlayers on Cu(110) using density functional theory. Surface Science, 2004, 548, 301-308.	1.9	99
9	Structure of enantiopure and racemic alanine adlayers on Cu(110). Surface Science, 2005, 574, L1-L8.	1.9	99
10	First-principles studies of chiral step reconstructions of Cu(100) by adsorbed glycine and alanine. Journal of Chemical Physics, 2006, 124, 074703.	3.0	44
11	The importance of charge–quadrupole interactions for H2adsorption and diffusion in CuBTC. Molecular Simulation, 2009, 35, 60-69.	2.0	39
12	Structures of Dense Glycine and Alanine Adlayers on Chiral Cu(3,1,17) Surfaces. Langmuir, 2006, 22, 8096-8103.	3.5	38
13	Trends in methanol decomposition on transition metal alloy clusters from scaling and Brønsted–Evans–Polanyi relationships. Physical Chemistry Chemical Physics, 2012, 14, 8644.	2.8	30
14	Exploring Computational Design of Size-Specific Subnanometer Clusters Catalysts. Topics in Catalysis, 2012, 55, 353-365.	2.8	30
15	Elucidation of Pathways for NO Electroreduction on Pt(111) from First Principles. Angewandte Chemie, 2015, 127, 8373-8376.	2.0	22
16	Computational Screening for Developing Optimal Intermetallic Transition Metal Pt-Based ORR Catalysts at the Predictive Volcano Peak. Journal of Physical Chemistry C, 2019, 123, 13236-13245.	3.1	21
17	Tracking Amino Acids in Chiral Quantum Corrals. Journal of Physical Chemistry C, 2013, 117, 11757-11763.	3.1	19
18	Characterization of Bulk Structure in Zinc Orthotitanate: A Density Functional Theory and EXAFS Investigation. Journal of the American Ceramic Society, 2008, 91, 584-590.	3.8	17

#	Article	IF	CITATIONS
19	Chiral "Pinwheel―Heterojunctions Self-Assembled from C ₆₀ and Pentacene. ACS Nano, 2013, 7, 3086-3094.	14.6	16
20	Computational predictive design for metal-decorated-graphene size-specific subnanometer to nanometer ORR catalysts. Catalysis Today, 2018, 312, 105-117.	4.4	13
21	Graphene Supported Tungsten Carbide as Catalyst for Electrochemical Reduction of CO2. Catalysts, 2019, 9, 604.	3.5	12
22	Surface reactions of AsH3, H2Se, and H2S on the Zn2TiO4(010) surface. Surface Science, 2011, 605, 818-823.	1.9	10
23	Amino Acid Immobilization of Copper Surface Diffusion on Cu(111). Advanced Materials Interfaces, 2019, 6, 1900021.	3.7	7
24	Towards quaternary alloy Au–Pd catalysts for direct synthesis of hydrogen peroxide. Materials Today Energy, 2020, 16, 100399.	4.7	5
25	Influence of Surface Reactions on Complex Hydride Reversibility. Journal of Physical Chemistry C, 2008, 112, 18270-18279.	3.1	4
26	Adsorption Energy Shifts for Oxygen and Hydroxyl on 4-atom Metal-Decorated Graphene Catalysts Via Solvation, pH, and Substrate Dopants: Effects on ORR Activity. Metals, 2019, 9, 227.	2.3	4
27	Density functional theory calculations of the surface structure of the inverse spinel zinc orthotitanate. Journal of Physics Condensed Matter, 2008, 20, 095001.	1.8	3
28	Size, Composition, and Support-Doping Effects on Oxygen Reduction Activity of Platinum-Alloy and on Non-platinum Metal-Decorated-Graphene Nanocatalysts. Frontiers in Chemistry, 2019, 7, 610.	3.6	3
29	Effects of surface structure and halogen substitution on electronic charge transfer in adlayers of (BETS)2-GaCl4 on silver surfaces. Surface Science, 2019, 687, 34-40.	1.9	3
30	DFT characterization of adsorption and diffusion mechanisms of H, As, S, and Se on the zinc orthotitanate (010) surface. Surface Science, 2008, 602, 1877-1882.	1.9	2
31	Computational screening of transition metal/pâ€block hybrid electrocatalysts for CO ₂ reduction. Journal of Computational Chemistry, 2020, 41, 1384-1394.	3.3	2
32	Innenr $\tilde{A}^{1}\!\!/\!$ cktitelbild: Elucidation of Pathways for NO Electroreduction on Pt(111) from First Principles (Angew. Chem. 28/2015). Angewandte Chemie, 2015, 127, 8417-8417.	2.0	0
33	Similarities and differences for atomic and diatomic molecule adsorption on the B-5 type sites of the HCP(101ì6) surfaces of Co, Os, and Ru from DFT calculations. Heliyon, 2019, 5, e01924.	3.2	O

134 Immobilized Cu Adatoms: Amino Acid Immobilization of Copper Surface Diffusion on Cu(111) (Adv.) Tj ETQq0 0 0 0 rgBT /Overlock 10 Tf