Byung Chul Yeo

List of Publications by Year in descending order

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	623734 940533		940533
17	689	14	16
papers	citations	h-index	g-index
17	17	17	1335
all docs	docs citations	times ranked	citing authors

#	Article	IF	CITATIONS
1	High-throughput Screening Computation for Discovery of Porous Zeolites for Hydrogen Storage. Journal of Korean Institute of Metals and Materials, 2022, 60, 537-544.	1.0	O
2	Accelerated mapping of electronic density of states patterns of metallic nanoparticles via machine-learning. Scientific Reports, 2021, 11, 11604.	3.3	11
3	High-throughput computational-experimental screening protocol for the discovery of bimetallic catalysts. Npj Computational Materials, 2021, 7, .	8.7	20
4	Unlocking the Potential of Nanoparticles Composed of Immiscible Elements for Direct H2O2 Synthesis. ACS Catalysis, 2019, 9, 8702-8711.	11.2	32
5	Pattern Learning Electronic Density of States. Scientific Reports, 2019, 9, 5879.	3.3	38
6	Atomistic Sodiation Mechanism of a Phosphorene/Graphene Heterostructure for Sodium-Ion Batteries Determined by First-Principles Calculations. Journal of Physical Chemistry C, 2018, 122, 20653-20660.	3.1	35
7	Dissimilar anisotropy of electron versus hole bulk transport in anatase <mml:math xmlns:mml="http://www.w3.org/1998/Math/MathML"><mml:msub><mml:mi>TiO</mml:mi><mml:mn>2<td>നനു.മ/നനി</td><td>l:m28b></td></mml:mn></mml:msub></mml:math>	നന ു. മ/നനി	l:m 28 b>
8	Simulation Protocol for Prediction of a Solid-Electrolyte Interphase on the Silicon-based Anodes of a Lithium-Ion Battery: ReaxFF Reactive Force Field. Journal of Physical Chemistry Letters, 2017, 8, 2812-2818.	4.6	51
9	Atomistic Simulation Protocol for Improved Design of Si–O–C Hybrid Nanostructures as Li-Ion Battery Anodes: ReaxFF Reactive Force Field. Journal of Physical Chemistry C, 2017, 121, 23268-23275.	3.1	14
10	A comparative first-principles study of the lithiation, sodiation, and magnesiation of black phosphorus for Li-, Na-, and Mg-ion batteries. Physical Chemistry Chemical Physics, 2016, 18, 21391-21397.	2.8	73
11	Atomistics of the lithiation of oxidized silicon (SiO _x) nanowires in reactive molecular dynamics simulations. Physical Chemistry Chemical Physics, 2016, 18, 32078-32086.	2.8	41
12	High-Throughput Screening to Investigate the Relationship between the Selectivity and Working Capacity of Porous Materials for Propylene/Propane Adsorptive Separation. Journal of Physical Chemistry C, 2016, 120, 24224-24230.	3.1	37
13	Simultaneously Controllable Doping Sites and the Activity of a W–N Codoped TiO ₂ Photocatalyst. ACS Catalysis, 2016, 6, 2745-2753.	11.2	84
14	Development of the ReaxFF _{CBN} reactive force field for the improved design of liquid CBN hydrogen storage materials. Physical Chemistry Chemical Physics, 2016, 18, 1818-1827.	2.8	27
15	Atomistic Observation of the Lithiation and Delithiation Behaviors of Silicon Nanowires Using Reactive Molecular Dynamics Simulations. Journal of Physical Chemistry C, 2015, 119, 3447-3455.	3.1	56
16	Unraveling the Atomistic Sodiation Mechanism of Black Phosphorus for Sodium Ion Batteries by First-Principles Calculations. Journal of Physical Chemistry C, 2015, 119, 15041-15046.	3.1	135
17	Energy Efficient Scalable Video Coding Based Cooperative Multicast Scheme with Selective Layer Forwarding. IEEE Communications Letters, 2013, 17, 1116-1119.	4.1	12