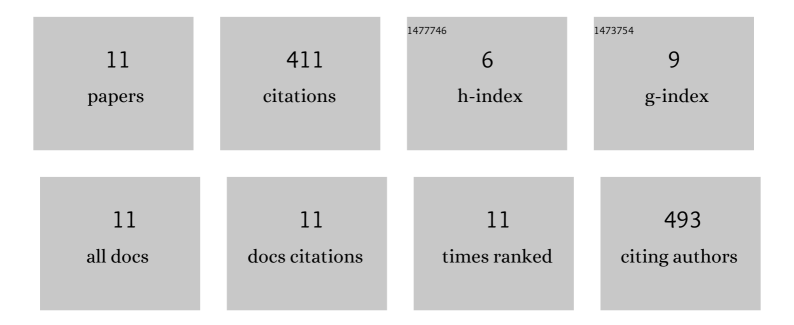
Preston B Moore

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Densely Packed Tethered Polymer Nanoislands: A Simulation Study. Polymers, 2021, 13, 2570.	2.0	0
2	Coordinated Responsive Arrays of Surface-Linked Polymer Islands—CORALs. ACS Applied Materials & Interfaces, 2018, 10, 7459-7468.	4.0	3
3	Computational Study of Intramolecular Heterocyclic Ring Formation with Cyclic Phosphazenes. International Journal of Engineering Research & Technology, 2014, 3, 1575-1582.	0.2	1
4	Raft registration across bilayers in a molecularly detailed model. Soft Matter, 2011, 7, 8182.	1.2	51
5	Quantitative Assessment of Force Fields on Both Low-Energy Conformational Basins and Transition-State Regions of the (أَهْمَ `أَ`) Space. Journal of Chemical Theory and Computation, 2011, 7, 402-419.	2.3	10
6	Nielsen, Moore, and Ensing Reply:. Physical Review Letters, 2011, 107, .	2.9	3
7	Calculating the surface tension between a flat solid and a liquid: a theoretical and computer simulation study of three topologically different methods. Journal of Mathematical Chemistry, 2009, 45, 161-174.	0.7	37
8	Energy Conservation in Adaptive Hybrid Atomistic/Coarse-Grain Molecular Dynamics. Journal of Chemical Theory and Computation, 2007, 3, 1100-1105.	2.3	151
9	Molecular Dynamics Investigations of Lipid Langmuir Monolayers Using a Coarse-Grain Model. Journal of Physical Chemistry B, 2003, 107, 13911-13917.	1.2	56
10	A combined time correlation function and instantaneous normal mode study of the sum frequency generation spectroscopy of the water/vapor interface. Journal of Chemical Physics, 2003, 118, 8411-8419.	1.2	98
11	Coarse Grained-to-Atomistic Mapping Algorithm: A Tool for Multiscale Simulations. , 0, , 73-88.		1