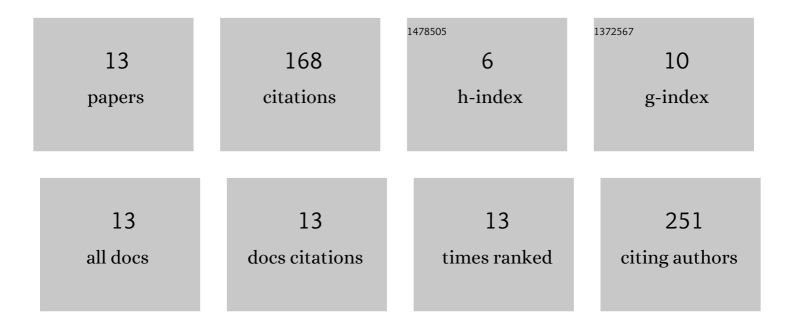
Haitao Yin

List of Publications by Year in descending order

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Ηλιτλο Υινι

#	Article	IF	CITATIONS
1	Electronic structure of III-V zinc-blende semiconductors from first principles. Physical Review B, 2013, 87, .	3.2	47
2	Composition dependent band offsets of ZnO and its ternary alloys. Scientific Reports, 2017, 7, 41567.	3.3	42
3	Energy storage performance of V _{n+1} C _n monolayer as electrode material studied by first-principles calculations. RSC Advances, 2016, 6, 54999-55006.	3.6	18
4	Spin transport through a junction entirely consisting of molecules from first principles. Applied Physics Letters, 2017, 111, .	3.3	18
5	Spin filtering in transition-metal phthalocyanine molecules from first principles. Frontiers of Physics, 2017, 12, 1.	5.0	13
6	Magnetic proximity, magnetoresistance and spin filtering effect in a binuclear ferric phthalocyanine from first principles. Journal Physics D: Applied Physics, 2020, 53, 035305.	2.8	8
7	Synthesis, photoluminescence and photocatalytic characteristics of Ag–ZnO sandwich structures. Journal of Physics and Chemistry of Solids, 2022, 165, 110697.	4.0	7
8	Theoretical limit of how small we can make MoS2 transistor channels. Journal Physics D: Applied Physics, 0, , .	2.8	5
9	Defect-Induced Different Band Alignment and Transport of All-Phosphorene Devices from First Principles. ACS Applied Electronic Materials, 2022, 4, 2070-2076.	4.3	4
10	Perfect dual spin filtering effect and large magnetoresistance in all-carbon devices based on C18 cyclo molecule from first principles. Physics Letters, Section A: General, Atomic and Solid State Physics, 2022, 441, 128166.	2.1	4
11	Predicting band offset of lattice matched ZnO and BeCdO heterojunction from first principles. Materials Research Letters, 2019, 7, 232-238.	8.7	2
12	Probing an Individual Electron Spin State in a Quantum Dot with Spin Bias. Journal of Nanoscience and Nanotechnology, 2018, 18, 2096-2099.	0.9	0
13	Electronic structures of Zn1â^'xGaxO1â^'xNx and band offsets of the ZnO/Zn1â^'xGaxO1â^'xNx heterojunction across the entire concentration range from first principles. Physical Chemistry Chemical Physics, 2021, 24, 375-381.	2.8	Ο