Kousuke Nakano

List of Publications by Year in descending order

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516710 477307 49 973 16 29 citations g-index h-index papers 50 50 50 1089 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Valence Band Engineering of Layered Bismuth Oxyhalides toward Stable Visible-Light Water Splitting: Madelung Site Potential Analysis. Journal of the American Chemical Society, 2017, 139, 18725-18731.	13.7	144
2	Superconductivity in BaTi ₂ Sb ₂ O with a <i>d</i> < ¹ Square Lattice. Journal of the Physical Society of Japan, 2012, 81, 103706.	1.6	85
3	Uniform leader election protocols for radio networks. IEEE Transactions on Parallel and Distributed Systems, 2002, 13, 516-526.	5.6	84
4	Hierarchically Porous Monoliths Based on N-Doped Reduced Titanium Oxides and Their Electric and Electrochemical Properties. Chemistry of Materials, 2013, 25, 3504-3512.	6.7	52
5	Synthesis and Physical Properties of the New Oxybismuthides BaTi ₂ Bi ₂ O and (SrF) ₂ Ti ₂ Bi ₂ O with a <i>d¹Square Net. Journal of the Physical Society of Japan, 2013, 82, 013703.</i>	1.6	43
6	Electrical Properties of Epitaxial Thin Films of Oxyhydrides ATiO _{3–<i>x</i>} H _{<i>x</i>} (A = Ba and Sr). Chemistry of Materials, 2015, 27, 6354-6359.	6.7	40
7	Two Superconducting Phases in the Isovalent Solid Solutions BaTi ₂ <i>Pn</i> Sub>2O (<i>Pn</i> Sub, and Bi). Journal of the Physical Society of Japanl: aath, and Bi) are the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and Bi). Journal of the Physical Society of Japanl: aath, and aath, and all	1.6	39
8	display="inline"> <mml:mi>s</mml:mi> -wave superconductivity in superconducting BaTi <mml:math display="inline" xmlns:mml="http://www.w3.org/1998/Math/MathML"><mml:msub><mml:mrow></mml:mrow><mml:mn>2</mml:mn></mml:msub></mml:math> Sb <mml:math< td=""><td>3.2</td><td>39</td></mml:math<>	3.2	39
9	xmlns:mml="http://www.w3.org/1998/Math/MathML" display="inline"> <mml:msub><mml:mrow /><mml: <scp>TurboRVB</scp>: A many-body toolkit for <i>ab initio</i> electronic simulations by quantum Monte Carlo. Journal of Chemical Physics, 2020, 152, 204121.</mml: </mml:mrow </mml:msub>	3.0	37
10	A survey on leader election protocols for radio networks. , 0, , .		26
11	Highâ€Pressure Synthesis of A ₂ NiO ₂ Ag ₂ Se ₂ (A=Sr, Ba) with a Highâ€Spin Ni ²⁺ in Squareâ€Planar Coordination. Angewandte Chemie - International Edition, 2019, 58, 756-759.	13.8	25
12	LaPd2Sb2: A pnictide superconductor with CaBe2Ge2 type structure. Journal of Alloys and Compounds, 2014, 583, 151-154.	5 . 5	23
13	An image retrieval system using FPGAs., 0,,. Muon spin relaxation and electron/neutron diffraction studies of BaTi <mml:math display="inline" xmlns:mml="http://www.w3.org/1998/Math/MathML"><mml:msub><mml:mrow< td=""><td></td><td>22</td></mml:mrow<></mml:msub></mml:math>		22

#	Article	IF	Citations
19	High- <i>T</i> _c Superconducting Hydrides Formed by LaH ₂₄ and YH ₂₄ Cage Structures as Basic Blocks. Chemistry of Materials, 2021, 33, 9501-9507.	6.7	16
20	Hypervalent Bismuthides La ₃ MBi ₅ (M = Ti, Zr, Hf) and Related Antimonides: Absence of Superconductivity. Inorganic Chemistry, 2017, 56, 5041-5045.	4.0	15
21	Highâ€Pressure Synthesis of A 2 NiO 2 Ag 2 Se 2 (A=Sr, Ba) with a Highâ€5pin Ni 2+ in Squareâ€Planar Coordination. Angewandte Chemie, 2019, 131, 766-769.	2.0	15
22	Estimation of maximum absorption wavelength of polymethine dyes in visible and near-infrared region based on time-dependent density functional theory. Chemical Physics, 2019, 518, 15-24.	1.9	15
23	All-Electron Quantum Monte Carlo with Jastrow Single Determinant Ansatz: Application to the Sodium Dimer. Journal of Chemical Theory and Computation, 2019, 15, 4044-4055.	5. 3	13
24	The Systematic Study on the Stability and Superconductivity of Yâ€Mgâ€H Compounds under High Pressure. Advanced Theory and Simulations, 2022, 5, .	2.8	13
25	High-pressure hydrogen by machine learning and quantum Monte Carlo. Physical Review B, 2022, 106, .	3.2	13
26	Superconductivity in LaPd2As2 with a collapsed 122 structure. Journal of Alloys and Compounds, 2014, 613, 370-374.	5. 5	12
27	Atomic forces by quantum Monte Carlo: Application to phonon dispersion calculations. Physical Review B, 2021, 103, .	3.2	12
28	General Correlated Geminal Ansatz for Electronic Structure Calculations: Exploiting Pfaffians in Place of Determinants. Journal of Chemical Theory and Computation, 2020, 16, 6114-6131.	5. 3	11
29	Speeding up <i>ab initio</i> diffusion Monte Carlo simulations by a smart lattice regularization. Physical Review B, 2020, 101, .	3.2	10
30	Sorting on single-channel wireless sensor networks. , 0, , .		8
31	Ab initio molecular dynamics simulation of structural and elastic properties of SiO ₂ â€"P ₂ O ₅ â€"Al ₂ O ₃ â€"Na ₂ O glass. Journal of the American Ceramic Society, 2022, 105, 6604-6615.	3.8	7
32	Diffusion Monte Carlo Study on Relative Stabilities of Boron Nitride Polymorphs. Journal of Physical Chemistry C, 2022, 126, 6000-6007.	3.1	6
33	<scp>Shry</scp> : Application of Canonical Augmentation to the Atomic Substitution Problem. Journal of Chemical Information and Modeling, 2022, 62, 2909-2915.	5.4	6
34	Investigation into Structural Phase Transitions in Layered Titanium-Oxypnictides by a Computational Phonon Analysis. Inorganic Chemistry, 2017, 56, 13732-13740.	4.0	5
35	High-Throughput Evaluation of Discharge Profiles of Nickel Substitution in LiNiO2 by Ab Initio Calculations. Journal of Physical Chemistry C, 2021, 125, 14517-14524.	3.1	5
36	Space-warp coordinate transformation for efficient ionic force calculations in quantum Monte Carlo. Journal of Chemical Physics, 2022, 156, 034101.	3.0	5

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37	Optimal algorithms for the multiple query problem on reconfigurable meshes, with applications. IEEE Transactions on Parallel and Distributed Systems, 2001, 12, 875-887.	5.6	4
38	Doubly-logarithmic energy-efficient initialization protocols for single-hop radio networks. , 2002, , .		4
39	Stochastic Estimations of the Total Number of Classes for a Clustering having Extremely Large Samples to be Included in the Clustering Engine. Advanced Theory and Simulations, 2021, 4, 2000301.	2.8	4
40	Pressure-Induced Collapse Transition in BaTi ₂ Pn ₂ O (Pn = As, Sb) with an Unusual Pn–Pn Bond Elongation. Inorganic Chemistry, 2021, 60, 2228-2233.	4.0	4
41	Superconducting properties of BaTi2Pn2O (Pn= Sb, Bi). Physica C: Superconductivity and Its Applications, 2014, 504, 36-38.	1.2	3
42	An optimal randomized ranking algorithm on the k-channel broadcast communication model. , 0, , .		2
43	Hydride-Reduced Eu2SrFe2O6: A T-to-T′ Conversion Enabling Fe2+ in Square-Planar Coordination. Inorganic Chemistry, 2020, 59, 12913-12919.	4.0	2
44	GaN bandgap bias caused by semi-core treatment in pseudopotentials analyzed by the diffusion Monte Carlo method. AIP Advances, 2021, 11, 025225.	1.3	2
45	A quantum annealing approach to ionic diffusion in solids. Scientific Reports, 2021, 11, 7261.	3.3	2
46	Computational Design to Suppress Thermal Runaway of Li-lon Batteries via Atomic Substitutions to Cathode Materials. ACS Applied Materials & Samp; Interfaces, 2022, , .	8.0	2
47	Workshop on advances in parallel and distributed computational models. , 2002, , .		O
48	Guest editors' introduction to special section on mobile computing and wireless networks. IEEE Transactions on Parallel and Distributed Systems, 2002, 13, 865-865.	5.6	0
49	Diffusion Monte Carlo evaluation of disiloxane linearisation barrier. Physical Chemistry Chemical Physics, 2022, , .	2.8	O