

# Phuti E Ngoepe

## List of Publications by Year in descending order

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58  
papers

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citations

361413

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docs citations

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times ranked

1369  
citing authors

#	ARTICLE	IF	CITATIONS
1	Structural characterisation and mechanical properties of nanosized spinel $\text{LiMn}_2\text{O}_4$ cathode investigated using atomistic simulation. Materials Research Bulletin, 2022, 146, 111611.	5.2	5
2	Adsorption mechanisms and effects of thiocarbamate collectors in the separation of chalcopyrite from pyrite minerals: DFT and experimental studies. Minerals Engineering, 2022, 176, 107318.	4.3	28
3	Oxidation behaviour of sperrylite and platarsite (100) surfaces: A DFT study. Materials Today Communications, 2022, 32, 103868.	1.9	1
4	Phase stability of $\text{TiPd}_{1-x}\text{Ru}_x$ and $\text{Ti}_{1-x}\text{PdR}_{1-x}$ shape memory alloys. Materials Today: Proceedings, 2021, 38, 1071-1076.	1.8	2
5	Intercalation voltages for spinel $\text{Li}_{x}\text{Mn}_2\text{O}_4$ ( $0 \leq x \leq 0.2$ ) cathode materials: Calibration of calculations with the ONETEP linear-scaling DFT code. Materials Today Communications, 2021, 27, 102380.	1.9	0
6	First Principles Study of Oxygen Adsorption on $\text{Li}-\text{MO}_2$ ( $\text{M} = \text{Mn, Ti and V}$ ) (110) Surface. Journal of the Electrochemical Society, 2021, 168, 070556.	2.9	5
7	The effects of process parameters on the properties of manganese-rich carbonate precursors: A study of co-precipitation synthesis using semi-batch reactors. Chemical Engineering Science, 2021, 241, 116694.	3.8	8
8	Density Functional Theory Study of Ethylene Carbonate Adsorption on the (0001) Surface of Aluminum Oxide $\text{Al}_2\text{O}_3$ . ACS Omega, 2021, 6, 29577-29587.	3.5	11
9	First-principles study: Effect of lithium and sodium intercalation in transition metal phosphates, $\text{MPO}_4$ ( $\text{M: Mn, Fe, Co}$ ). Computational Condensed Matter, 2020, 22, e00437.	2.1	10
10	Structural, thermodynamic, electronic and mechanical properties of spinel and phonon-harvested $\text{AMn}_2\text{O}_4$ ( $\text{A: Li, Na, Mg}$ ) systems: A First-Principles study. Materials Today Communications, 2020, 22, 100704.	1.9	6
11	Atomistic Simulation and Characterization of Spinel $\text{Li}_{1+x}\text{Mn}_2\text{O}_4$ ( $0 \leq x \leq 1$ ) Nanoparticles. ACS Applied Energy Materials, 2020, 3, 1429-1438.	5.1	12
12	Synthesis of the porous spinel $\text{Co-Al}_2\text{O}_4$ powder produced by ball milling and annealing. Advanced Powder Technology, 2020, 31, 2742-2748.	4.1	0
13	Simulated synthesis and structure of $\text{Li}_x\text{TiO}_2$ nanosheets as anode material for lithium ion batteries. Optical Materials, 2020, 102, 109831.	3.6	2
14	Ethylene carbonate adsorption on the major surfaces of lithium manganese oxide $\text{Li}_{1-x}\text{Mn}_2\text{O}_4$ spinel ( $0.000 \leq x \leq 0.375$ ): a DFT+U-D3 study. Physical Chemistry Chemical Physics, 2020, 22, 6763-6771.	2.8	18
15	Discharge induced structural variation of simulated bulk $\text{Li}_{1+x}\text{Mn}_2\text{O}_4$ ( $0 \leq x \leq 1$ ). Optical Materials, 2019, 92, 67-70.	3.6	2
16	Thermodynamic stability and pressure dependence of $\text{FePO}_4$ polymorphs. Computational and Theoretical Chemistry, 2019, 1155, 67-74.	2.5	12
17	<i>Ab initio</i> investigation of the thermodynamics of cation distribution and of the electronic and magnetic structures in the $\text{LiMn}_2\text{O}_4$ spinel. Physical Review B, 2018, 97, 104112.	2.8	18
18	Thermodynamically accessible titanium clusters $\text{Ti}_x\text{N}_y$ , $x\text{N}_y = 2$ –32. Physical Chemistry Chemical Physics, 2018, 20, 13962-13973.	2.8	18

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19	Advances in Ti-Based Systems as High Temperature Shape Memory Alloys. <i>Key Engineering Materials</i> , 2018, 770, 230-238.	0.4	2
20	First Principle Study of $Ti_{50}Al_{50}$ Alloys. <i>Key Engineering Materials</i> , 2018, 770, 224-229.	0.4	0
21	Periodic modeling of zeolite Ti-LTA. <i>Journal of Chemical Physics</i> , 2017, 147, 074701.	3.0	11
22	â€˜Breathing-crystalsâ€™ the origin of electrochemical activity of mesoporous Liâ€“MnO <sub>2</sub> . <i>Journal of Materials Chemistry A</i> , 2016, 4, 6456-6464.	10.3	5
23	Computational Modelling as a Value Add in Energy Storage Materials. <i>Nanostructure Science and Technology</i> , 2016, , 481-513.	0.1	1
24	Structure of Surface Entrance Sites for Li Intercalation into TiO <sub>2</sub> Nanoparticles, Nanosheets, and Mesoporous Architectures with Application for Li-Ion Batteries. <i>Journal of Physical Chemistry C</i> , 2016, 120, 14001-14008.	3.1	6
25	Light scattering and computer simulation studies of superionic pure and La-doped BaF <sub>2</sub> . <i>Chemical Physics</i> , 2016, 467, 6-12.	1.9	2
26	Origin of electrochemical activity in nano-Li <sub>2</sub> MnO <sub>3</sub> ; stabilization via a â€˜point defect scaffoldâ€™. <i>Nanoscale</i> , 2015, 7, 1167-1180.	5.6	20
27	Martensitic Transformation Behaviour of $Ti_{50}Pt_{50-x}Co_x$ Shape Memory Alloys. <i>Advanced Materials Research</i> , 2014, 1019, 385-390.	0.3	0
28	Lithium and oxygen adsorption at the $\hat{\tau}^2\text{-MnO}_2$ (110) surface. <i>Journal of Materials Chemistry A</i> , 2013, 1, 14879.	10.3	58
29	Structural, elastic and electronic properties of equiatomic PtTi as potential high-temperature shape memory alloy. <i>Intermetallics</i> , 2013, 33, 27-32.	3.9	37
30	Phase transformation and crystal structure of IrTi. <i>Intermetallics</i> , 2012, 31, 26-33.	3.9	11
31	An alternative DFT-based model for calculating structural and elastic properties of random binary HCP, FCC and BCC alloys: Mgâ€“Li system as test case. <i>Intermetallics</i> , 2012, 21, 88-96.	3.9	15
32	Amorphization and recrystallization study of lithium insertion into manganese dioxide. <i>Physical Chemistry Chemical Physics</i> , 2011, 13, 1307-1313.	2.8	20
33	Solid-state transformation in nanocrystalline Ti induced by ball milling. <i>Materials Letters</i> , 2010, 64, 1215-1218.	2.6	41
34	Negative compressibility in platinum sulfide using density-functional theory. <i>Physical Review B</i> , 2010, 81, .	3.2	31
35	Electronic Structure and Redox Properties of the Ti-Doped Zirconia (111) Surface. <i>Journal of Physical Chemistry C</i> , 2010, 114, 15403-15409.	3.1	20
36	Theoretical investigation of the Pt <sub>3</sub> Al ground state. <i>Intermetallics</i> , 2010, 18, 417-421.	3.9	36

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37	Link between structural and mechanical stability of fcc- and bcc-based ordered Mg-Li alloys. <i>Intermetallics</i> , 2010, 18, 2083-2089.	3.9	59
38	Generating structural distributions of atomistic models of Li <sub>2</sub> O nanoparticles using simulated crystallisation. <i>Journal of Materials Chemistry</i> , 2010, 20, 10452.	6.7	8
39	Simulating Mechanical Deformation in Nanomaterials with Application for Energy Storage in Nanoporous Architectures. <i>ACS Nano</i> , 2009, 3, 3308-3314.	14.6	23
40	Atomistic simulation of the surface structure of electrolytic manganese dioxide. <i>Surface Science</i> , 2009, 603, 3184-3190.	1.9	32
41	Predicting the Electrochemical Properties of MnO <sub>2</sub> Nanomaterials Used in Rechargeable Li Batteries: Simulating Nanostructure at the Atomistic Level. <i>Journal of the American Chemical Society</i> , 2009, 131, 6161-6173.	13.7	74
42	The Displacive Phase Transition of Vanadium Dioxide and the Effect of Doping with Tungsten. <i>Chemistry of Materials</i> , 2008, 20, 1764-1772.	6.7	68
43	Evolving microstructure in MnO <sub>2</sub> using amorphisation and recrystallisation. <i>Journal of Crystal Growth</i> , 2006, 294, 118-129.	1.5	13
44	A computer modelling study of the interaction of organic adsorbates with fluorapatite surfaces. <i>Physics and Chemistry of Minerals</i> , 2006, 33, 314-331.	0.8	25
45	Nanoscale coatings of AuAl <sub>x</sub> and PtAl <sub>x</sub> and their mesoporous elemental derivatives. <i>Current Applied Physics</i> , 2006, 6, 440-443.	2.4	20
46	Generating MnO <sub>2</sub> Nanoparticles Using Simulated Amorphization and Recrystallization. <i>Journal of the American Chemical Society</i> , 2005, 127, 12828-12837.	13.7	69
47	Light scattering and computational simulations of the superionic behaviour of CaF <sub>2</sub> doped with lanthanide ions. <i>Physica Status Solidi C: Current Topics in Solid State Physics</i> , 2005, 2, 490-494.	0.8	1
48	Brillouin scattering studies and computational simulations of the elastic properties of pyrite (FeS <sub>2</sub> ) at high temperatures. <i>Physica Status Solidi C: Current Topics in Solid State Physics</i> , 2004, 1, 3073-3076.	0.8	4
49	Atomistic simulation of the structure and elastic properties of pyrite (FeS <sub>2</sub> ) as a function of pressure. <i>Physics and Chemistry of Minerals</i> , 2003, 30, 615-619.	0.8	27
50	Ab initio calculation of the voltage profile for LiC <sub>6</sub> . <i>Solid State Ionics</i> , 2003, 159, 21-23.	2.7	24
51	Modeling the Surface Structure and Reactivity of Pyrite: Introducing a Potential Model for FeS <sub>2</sub> . <i>Journal of Physical Chemistry B</i> , 2000, 104, 7969-7976.	2.6	72
52	Calculated cell discharge curve for lithium batteries with a V <sub>2</sub> O <sub>5</sub> cathode. <i>Journal of Materials Chemistry</i> , 2000, 10, 239-240.	6.7	20
53	Behavior of elastic constants, refractive index, and lattice parameter of cubic zirconia at high temperatures. <i>Journal of Applied Physics</i> , 1993, 73, 7268-7274.	2.5	44
54	High-temperature elastic constants of yttrium aluminum garnet. <i>Journal of Applied Physics</i> , 1993, 73, 7298-7301.	2.5	39

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55	Brillouin-scattering and computer-simulation studies of fast-ion conductors: a review. Journal of the Chemical Society, Faraday Transactions, 1990, 86, 1183.	1.7	27
56	Brillouin scattering investigation of the rare-earth fluorides at high temperatures. Journal of the Less Common Metals, 1989, 148, 375-380.	0.8	5
57	Temperature Dependence and Martensitic Transformation of Ti<sub>50</sub>Pt<sub>50</sub> Shape Memory Alloys. Advanced Materials Research, 0, 1019, 379-384.	0.3	0
58	Unravelling the Catalytic Activity of MnO<sub>2</sub>, TiO<sub>2</sub>, and VO<sub>2</sub> (110) Surfaces by Oxygen Coadsorption on Sodium-Adsorbed MO<sub>2</sub> {M = Mn, Ti, V}. ACS Omega, 0, ..	3.5	5