

# OÄuz GÃ¼lseren

## List of Publications by Year in descending order

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107  
papers

5,128  
citations

126858  
33  
h-index

88593  
70  
g-index

107  
all docs

107  
docs citations

107  
times ranked

5407  
citing authors

#	ARTICLE	IF	CITATIONS
1	Giant Anharmonicity and Nonlinear Electron-Phonon Coupling in MgB <sub>2</sub> : A Combined First-Principles Calculation and Neutron Scattering Study. <i>Physical Review Letters</i> , 2001, 87, 037001.	2.9	381
2	Systematic study of adsorption of single atoms on a carbon nanotube. <i>Physical Review B</i> , 2003, 67, .	1.1	305
3	Noncrystalline Structures of Ultrathin Unsupported Nanowires. <i>Physical Review Letters</i> , 1998, 80, 3775-3778.	2.9	269
4	Mo <sub>2</sub> C as a high capacity anode material: a first-principles study. <i>Journal of Materials Chemistry A</i> , 2016, 4, 6029-6035.	5.2	249
5	Elasticity of iron at the temperature of the Earth's inner core. <i>Nature</i> , 2001, 413, 57-60.	13.7	240
6	Systematic ab initio study of curvature effects in carbon nanotubes. <i>Physical Review B</i> , 2002, 65, .	1.1	235
7	Tunable Adsorption on Carbon Nanotubes. <i>Physical Review Letters</i> , 2001, 87, 116802.	2.9	184
8	MXenes/graphene heterostructures for Li battery applications: a first principles study. <i>Journal of Materials Chemistry A</i> , 2018, 6, 2337-2345.	5.2	173
9	Accuracy of equation-of-state formulations. <i>American Mineralogist</i> , 2000, 85, 338-344.	0.9	160
10	Premelting of thin wires. <i>Physical Review B</i> , 1995, 51, 7377-7380.	1.1	145
11	Added row model of TiO <sub>2</sub> (110)1-2. <i>Physical Review B</i> , 1998, 58, 1586-1589.	1.1	132
12	High-pressure thermoelasticity of body-centered-cubic tantalum. <i>Physical Review B</i> , 2002, 65, .	1.1	122
13	Reversible band-gap engineering in carbon nanotubes by radial deformation. <i>Physical Review B</i> , 2002, 65, .	1.1	121
14	Pressure-induced interlinking of carbon nanotubes. <i>Physical Review B</i> , 2000, 62, 12648-12651.	1.1	116
15	Energetics and Electronic Structures of Individual Atoms Adsorbed on Carbon Nanotubes. <i>Journal of Physical Chemistry B</i> , 2004, 108, 575-582.	1.2	116
16	Oxygenation of carbon nanotubes: Atomic structure, energetics, and electronic structure. <i>Physical Review B</i> , 2003, 67, .	1.1	109
17	Effects of hydrogen adsorption on single-wall carbon nanotubes: Metallic hydrogen decoration. <i>Physical Review B</i> , 2002, 66, .	1.1	104
18	Functionalized carbon nanotubes and device applications. <i>Journal of Physics Condensed Matter</i> , 2004, 16, R901-R960.	0.7	104

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19	Exohydrogenated single-wall carbon nanotubes. <i>Physical Review B</i> , 2001, 64, .	1.1	103
20	Alkali Metal Intercalation in MXene/Graphene Heterostructures: A New Platform for Ion Battery Applications. <i>Journal of Physical Chemistry Letters</i> , 2019, 10, 727-734.	2.1	88
21	Thermal equation of state of tantalum. <i>Physical Review B</i> , 2001, 63, .	1.1	87
22	Pentagonal nanowires: A first-principles study of the atomic and electronic structure. <i>Physical Review B</i> , 2002, 65, .	1.1	75
23	First principles study of structural phase stability of wide-gap semiconductors MgTe, MgS and MgSe. <i>Computational Materials Science</i> , 2009, 47, 593-598.	1.4	71
24	Electronic structure of the contact between carbon nanotube and metal electrodes. <i>Applied Physics Letters</i> , 2003, 83, 3180-3182.	1.5	61
25	Variable and reversible quantum structures on a single carbon nanotube. <i>Physical Review B</i> , 2000, 62, R16345-R16348.	1.1	50
26	Peculiar Piezoelectric Properties of Soft Two-Dimensional Materials. <i>Journal of Physical Chemistry C</i> , 2016, 120, 13948-13953.	1.5	50
27	Structural and electronic properties of MoS <sub>2</sub> , WS <sub>2</sub> , and WS <sub>2</sub> /MoS <sub>2</sub> heterostructures encapsulated with hexagonal boron nitride monolayers. <i>Journal of Applied Physics</i> , 2017, 122, .	1.1	49
28	Understanding the plasmonic properties of dewetting formed Ag nanoparticles for large area solar cell applications. <i>Optics Express</i> , 2013, 21, 18344.	1.7	47
29	Relaxations of fluorouracil tautomers by decorations of fullerene-like SiCs: DFT studies. <i>Physics Letters, Section A: General, Atomic and Solid State Physics</i> , 2016, 380, 2160-2166.	0.9	42
30	Adsorption of Pt and Bimetallic PtAu Clusters on the Partially Reduced Rutile (110) TiO <sub>2</sub> Surface: A First-Principles Study. <i>Journal of Physical Chemistry C</i> , 2012, 116, 5735-5746.	1.5	40
31	Rich complex behaviour of self-assembled nanoparticles far from equilibrium. <i>Nature Communications</i> , 2017, 8, 14942.	5.8	40
32	Validation of inter-atomic potential for WS <sub>2</sub> and WSe <sub>2</sub> crystals through assessment of thermal transport properties. <i>Computational Materials Science</i> , 2018, 144, 92-98.	1.4	36
33	DFT explorations of quadrupole coupling constants for planar 5-fluorouracil pairs. <i>Computational and Theoretical Chemistry</i> , 2016, 1090, 67-73.	1.1	35
34	A distinct correlation between the vibrational and thermal transport properties of group VA monolayer crystals. <i>Nanoscale</i> , 2018, 10, 7803-7812.	2.8	35
35	First principles force field for metallic tantalum. <i>Modelling and Simulation in Materials Science and Engineering</i> , 2004, 12, S445-S459.	0.8	34
36	A systematical ab-initio review of promising 2D MXene monolayers towards Li-ion battery applications. <i>JPhys Energy</i> , 2020, 2, 032006.	2.3	34

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37	Theoretical analysis of STM experiments at rutile TiO <sub>2</sub> surfaces. Surface Science, 1997, 377-379, 150-154.		0.8	33
38	Metal nanoring and tube formation on carbon nanotubes. Physical Review B, 2002, 66, .		1.1	33
39	A comparative study of O <sub>2</sub> adsorbed carbon nanotubes. Chemical Physics Letters, 2003, 380, 1-5.		1.2	30
40	Effect of impurities on the mechanical and electronic properties of Au, Ag, and Cu monatomic chain nanowires. Physical Review B, 2011, 84, .		1.1	30
41	Formation of quantum structures on a single nanotube by modulating hydrogen adsorption. Physical Review B, 2003, 68, .		1.1	29
42	Pt-incorporated anatase $\langle\text{mml:math}\text{ xmlns:mml="http://www.w3.org/1998/Math/MathML"}\text{ display="inline"}\rangle\langle\text{mml:mrow}\rangle\langle\text{mml:msub}\rangle\langle\text{mml:mrow}\rangle\langle\text{mml:mtext} \text{TiO}\langle/\text{mml:mtext}\rangle\langle/\text{mml:mrow}\rangle\langle\text{mml:mn} \text{2}\langle/\text{mml:mn}\rangle\langle/\text{mml:msub}\rangle\langle/\text{mml:mrow}\rangle\langle/\text{mml:math}\rangle$ for solar cell applications: First-principles density functional theory calculations. Physical Review B, 2009, 79, .		1.1	28
43	<i>Ab initio</i> study of neutral (TiO <sub>2</sub> ) <sub>n</sub> clusters and their interactions with water and transition metal atoms. Journal of Physics Condensed Matter, 2012, 24, 305301.		0.7	27
44	An experimental and first-principles study of the effect of B/N doping in TiO <sub>2</sub> thin films for visible light photo-catalysis. Journal of Photochemistry and Photobiology A: Chemistry, 2013, 254, 25-34.		2.0	27
45	Piezoelectric Pb(Zr0.5Ti0.5)O <sub>3</sub> : Interplay of atomic ordering, ferroelectric soft modes, and pressure. Physical Review B, 2002, 66, .		1.1	26
46	First principles study of electronic and mechanical properties of molybdenum selenide type nanowires. Physical Review B, 2006, 74, .		1.1	26
47	Dye adsorbates BrPDI, BrGly, and BrAsp on anatase TiO <sub>2</sub> (001) for dye-sensitized solar cell applications. Physical Review B, 2009, 80, .		1.1	25
48	Analysis of Charge Transfer for in Situ Li Intercalated Carbon Nanotubes. Journal of Physical Chemistry C, 2012, 116, 11364-11369.		1.5	25
49	Vacancy formation enthalpy at high pressures in tantalum. Journal of Physics Condensed Matter, 2003, 15, 855-861.		0.7	24
50	Investigation of new two-dimensional materials derived from stanene. Computational Materials Science, 2017, 137, 208-214.		1.4	23
51	Effect of Molecular and Electronic Structure on the Light-Harvesting Properties of Dye Sensitizers. Journal of Physical Chemistry C, 2007, 111, 7539-7547.		1.5	22
52	First-principles investigation of pentagonal and hexagonal core-shell silicon nanowires with various core compositions. Physical Review B, 2009, 80, .		1.1	22
53	First-principles study of thin $\langle\text{mml:math}\text{ xmlns:mml="http://www.w3.org/1998/Math/MathML"}\text{ display="inline"}\rangle\langle\text{mml:mrow}\rangle\langle\text{mml:msub}\rangle\langle\text{mml:mrow}\rangle\langle\text{mml:mtext} \text{TiO}\langle/\text{mml:mtext}\rangle\langle/\text{mml:mrow}\rangle\langle\text{mml:mi}\times\langle/\text{mml:mi}\rangle\times\langle\text{mml:mi}\rangle\times\langle/\text{mml:mi}\rangle\langle/\text{mml:msub}\rangle\langle/\text{mml:mrow}\rangle\langle/\text{mml:math}\rangle$ bulklike rutile nanowires. Physical Review B, 2009, 80, .		1.1	21
54	Non-covalent functionalization of single wall carbon nanotubes and graphene by a conjugated polymer. Applied Physics Letters, 2014, 105, .		1.5	21

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55	Temperature-dependent phonon spectrum of transition metal dichalcogenides calculated from the spectral energy density: Lattice thermal conductivity as an application. <i>Physical Review B</i> , 2019, 100, .	1.1	21
56	Bias in bonding behavior among boron, carbon, and nitrogen atoms in ion implanted $\alpha$ -BN, $\alpha$ -BC, and diamond like carbon films. <i>Journal of Applied Physics</i> , 2011, 110, .	1.1	20
57	An experimental and theoretical examination of the effect of sulfur on the pyrolytically grown carbon nanotubes from sucrose-based solid state precursors. <i>Carbon</i> , 2011, 49, 508-517.	5.4	20
58	First-Principles Investigation of $\text{NO}_{\text{x}}$ and $\text{SO}_{\text{x}}$ Adsorption on Anatase-Supported BaO and Pt Overlays. <i>Journal of Physical Chemistry C</i> , 2012, 116, 6191-6199.	1.5	20
59	Electronic structure of half-metallic ferromagnet $\text{Co}_2\text{MnSi}$ at high-pressure. <i>European Physical Journal B</i> , 2010, 76, 321-326.	0.6	19
60	First-principles investigation of armchair stanene nanoribbons. <i>Physics Letters, Section A: General, Atomic and Solid State Physics</i> , 2018, 382, 180-185.	0.9	19
61	Theoretical and spectroscopic investigations on the structure and bonding in $\text{B}_x\text{N}$ thin films. <i>Thin Solid Films</i> , 2009, 518, 1459-1464.	0.8	18
62	Gate induced monolayer behavior in twisted bilayer black phosphorus. <i>2D Materials</i> , 2017, 4, 035025.	2.0	18
63	Modification of electronic structure by Au impurity investigated with density functional theory. <i>Physical Review B</i> , 2009, 80, .	1.1	16
64	A simple theory of 40K superconductivity in $\text{MgB}_2$ : first-principles calculations of $T_c$ , its dependence on boron mass and pressure. <i>Journal of Physics and Chemistry of Solids</i> , 2002, 63, 2201-2206.	1.9	14
65	Electronic structures and optical spectra of thin anatase $\text{TiO}_2$ through hybrid density functional and quasiparticle calculations. <i>Physical Review B</i> , 2014, 89, .	1.1	14
66	DFT study of noble metal impurities on $\text{TiO}_2(110)$ . <i>European Physical Journal B</i> , 2012, 85, 1.	0.6	12
67	Chemically uracil-functionalized carbon and silicon carbide nanotubes: Computational studies. <i>Materials Chemistry and Physics</i> , 2018, 205, 164-170.	2.0	12
68	Distributed contact flip chip InGaN/GaN blue LED; comparison with conventional LEDs. <i>Superlattices and Microstructures</i> , 2019, 128, 9-13.	1.4	12
69	DFT Studies of Graphene-Functionalised Derivatives of Capecitabine. <i>Zeitschrift Fur Naturforschung - Section A Journal of Physical Sciences</i> , 2017, 72, 1131-1138.	0.7	11
70	VIBRATIONAL MODES IN SMALL Agn, Aun CLUSTERS: A FIRST PRINCIPLE CALCULATION. <i>International Journal of Modern Physics B</i> , 2009, 23, 5819-5834.	1.0	10
71	Characterization of platinum nitride from first-principles calculations. <i>Journal of Physics Condensed Matter</i> , 2009, 21, 485403.	0.7	10
72	Nanotribological Properties of the h-BN/Au(111) Interface: A DFT Study. <i>Journal of Physical Chemistry C</i> , 2019, 123, 28411-28418.	1.5	10

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73	Electronic structure of strained Si <sub>n</sub> /Ge(001) superlattices. Solid State Communications, 1988, 65, 1285-1290.	0.9	9
74	Localization of acoustical modes due to the electron-phonon interaction within a two-dimensional electron gas. Journal of Physics Condensed Matter, 1993, 5, 589-598.	0.7	9
75	Hofstadter butterfly of graphene with point defects. Physical Review B, 2012, 85, .	1.1	9
76	DFT studies of CNTâ€“functionalized uracil-acetate hybrids. Physica E: Low-Dimensional Systems and Nanostructures, 2015, 73, 105-109.	1.3	8
77	On the structural and electronic properties of Ir-silicide nanowires on Si(001) surface. Journal of Applied Physics, 2016, 120, .	1.1	8
78	Double Perovskite Structure Induced by Co Addition to PbTiO <sub>3</sub> : Insights from DFT and Experimental Solid-State NMR Spectroscopy. Journal of Physical Chemistry C, 2019, 123, 27132-27139.	1.5	8
79	A study on the reconstruction of Ga terminated GaAs [1 1 1] surface. Solid State Communications, 1985, 56, 501-504.	0.9	7
80	Interaction of BrPDI, BrGly, and BrAsp with the Rutile TiO <sub>2</sub> (110) Surface for Photovoltaic and Photocatalytic Applications: A First-Principles Study. Journal of Physical Chemistry C, 2011, 115, 9220-9226.	1.5	7
81	Functionalization of (n, 0) CNTs (n = 3â€“16) by uracil: DFT studies. European Physical Journal B, 2018, 91, 1.	0.6	7
82	Electric-field effects on finite-length superlattices. Physical Review B, 1992, 46, 7621-7626.	1.1	6
83	In pursuit of barrierless transition metal dichalcogenides lateral heterojunctions. Nanotechnology, 2018, 29, 295202.	1.3	6
84	Local entanglement and string order parameter in dimerized models. Journal of Physics Condensed Matter, 2019, 31, 505602.	0.7	6
85	Anatase TiO <sub>2</sub> nanowires functionalized by organic sensitizers for solar cells: A screened Coulomb hybrid density functional study. Journal of Applied Physics, 2015, 118, 194301.	1.1	5
86	Elucidating the Barriers on Direct Water Splitting: Key Role of Oxygen Vacancy Density and Coordination over PbTiO <sub>3</sub> and TiO <sub>2</sub> . Journal of Physical Chemistry C, 2021, 125, 1874-1880.	1.5	5
87	Analysis of defects on BN nano-structures using high-resolution electron microscopy and density-functional calculations. Ultramicroscopy, 2008, 108, 1484-1489.	0.8	4
88	Thermoelectric efficiency of nanowires with long-range surface disorder. Physical Review B, 2012, 85, .	1.1	4
89	The integer quantum Hall effect of a square lattice with an array of point defects. Journal of Physics Condensed Matter, 2012, 24, 345501.	0.7	3
90	Synthesis of Phosphorus Included Multiwalled Carbon Nanotubes by Pyrolysis of Sucrose. Journal of Physical Chemistry C, 2013, 117, 24554-24560.	1.5	3

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91	Hybrid functional calculated optical and electronic structures of thin anatase TiO <sub>2</sub> nanowires with organic dye adsorbates. <i>Applied Surface Science</i> , 2015, 354, 437-442.	3.1	3
92	Mixed ab initio and semiempirical study of hydrogen-terminated finite germanium nanowires. <i>European Physical Journal Plus</i> , 2018, 133, 1.	1.2	3
93	First-principles zone-center theory of superconductivity in MgB <sub>2</sub> . <i>Applied Physics A: Materials Science and Processing</i> , 2002, 74, s945-s947.	1.1	2
94	Ab-initio Atomic Scale Study of Nearly Frictionless Surfaces. , 2007, , 57-77.		2
95	Hall conductance in graphene with point defects. <i>Journal of Physics Condensed Matter</i> , 2013, 25, 055302.	0.7	2
96	Range-Separated Hybrid Density Functional Study of Organic Dye Sensitizers on Anatase TiO <sub>2</sub> Nanowires. <i>Journal of Physical Chemistry C</i> , 2014, 118, 24776-24783.	1.5	2
97	Deterministic phase transitions and self-organization in logistic cellular automata. <i>Physical Review E</i> , 2019, 100, 042216.	0.8	2
98	Tailoring Vibrational Signature and Functionality of 2D-Ordered Linear-Chain Carbon-Based Nanocarriers for Predictive Performance Enhancement of High-End Energetic Materials. <i>Nanomaterials</i> , 2022, 12, 1041.	1.9	2
99	Electronic structure of a Si delta-layer embedded in Ge(001). <i>Semiconductor Science and Technology</i> , 1991, 6, 1002-1005.	1.0	1
100	Electronic structure of Ge-Si superlattices grown on Ge (001). <i>Semiconductor Science and Technology</i> , 1991, 6, 638-641.	1.0	1
101	Pattern information extraction from crystal structures. <i>Computer Physics Communications</i> , 2007, 176, 486-506.	3.0	1
102	New Trends in Nanotribology. <i>Tribology Letters</i> , 2010, 39, 227-227.	1.2	1
103	DIPEPTIDE ADSORPTION ON <i>Si</i> (100)-2 Å—1 ASYMMETRIC SURFACE BY FIRST PRINCIPLES. <i>International Journal of Modern Physics C</i> , 2010, 21, 97-106.	0.8	1
104	Systematic spatial and stoichiometric screening towards understanding the surface of ultrasmall oxygenated silicon nanocrystal. <i>Applied Surface Science</i> , 2016, 387, 771-778.	3.1	1
105	Infrared spectrum and STM images of cyclohexene-2-ethanamine: First principle investigation. <i>Journal of Molecular Structure</i> , 2008, 886, 144-147.	1.8	0
106	Scattering analysis of silver nanoparticles for solar cell applications using integral equations. , 2018, , .		0
107	Theoretical study of germanium nanoclusters: significance of surface passivation. <i>European Physical Journal Plus</i> , 2022, 137, 1.	1.2	0