

Silvana Botti

List of Publications by Year in descending order

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130
papers

6,960
citations

81900
39
h-index

62596
80
g-index

140
all docs

140
docs citations

140
times ranked

8011
citing authors

#	ARTICLE	IF	CITATIONS
1	Electronic Structure and Epitaxy of CdTe Shells on InSb Nanowires. <i>Advanced Science</i> , 2022, 9, e2105722.	11.2	7
2	Roadmap on Machine learning in electronic structure. <i>Electronic Structure</i> , 2022, 4, 023004.	2.8	69
3	A dataset of 175k stable and metastable materials calculated with the PBEsol and SCAN functionals. <i>Scientific Data</i> , 2022, 9, 64.	5.3	8
4	Machine-learning correction to density-functional crystal structure optimization. <i>MRS Bulletin</i> , 2022, 47, 765-771.	3.5	7
5	Superconductivity in S-rich phases of lanthanum sulfide under high pressure. <i>Physical Review Materials</i> , 2022, 6, .	2.4	3
6	Absorption mechanism of dopamine/DOPAC-modified TiO ₂ nanoparticles by time-dependent density functional theory calculations. <i>Materials Today Energy</i> , 2021, 19, 100571.	4.7	2
7	Bishop's hat silicene: a planar square silicon bilayer decorated with adatoms. <i>Physical Chemistry Chemical Physics</i> , 2021, 23, 16942-16947.	2.8	0
8	A high-throughput study of oxynitride, oxyfluoride and nitrofluoride perovskites. <i>Journal of Materials Chemistry A</i> , 2021, 9, 8501-8513.	10.3	18
9	Predicting stable crystalline compounds using chemical similarity. <i>Npj Computational Materials</i> , 2021, 7, .	8.7	41
10	Halogen molecular modifications at high pressure: the case of iodine. <i>Physical Chemistry Chemical Physics</i> , 2021, 23, 3321-3326.	2.8	5
11	Direct insight into the structure-property relation of interfaces from constrained crystal structure prediction. <i>Nature Communications</i> , 2021, 12, 811.	12.8	10
12	First-Principles Identification of Single Photon Emitters Based on Carbon Clusters in Hexagonal Boron Nitride. <i>Journal of Physical Chemistry A</i> , 2021, 125, 1325-1335.	2.5	51
13	Efficient strain-induced light emission in lonsdaleite germanium. <i>Physical Review Materials</i> , 2021, 5, .	2.4	16
14	A Global-Optimization Study of the Phase Diagram of Free-Standing Hydrogenated Two-Dimensional Silicon. <i>Journal of Physical Chemistry C</i> , 2021, 125, 6298-6305.	3.1	4
15	Unveiling Planar Defects in Hexagonal Group IV Materials. <i>Nano Letters</i> , 2021, 21, 3619-3625.	9.1	8
16	Point defects in hexagonal silicon. <i>Physical Review Materials</i> , 2021, 5, .	2.4	4
17	Electronic Structure of Molecules, Surfaces, and Molecules on Surfaces with the Local Modified Becke-Johnson Exchange-Correlation Potential. <i>Journal of Chemical Theory and Computation</i> , 2021, 17, 4746-4755.	5.3	10
18	Defect levels from SCAN and MBJ meta-GGA exchange-correlation potentials. <i>Physical Review B</i> , 2021, 104, .	3.2	5

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19	Bandgap of two-dimensional materials: Thorough assessment of modern exchange–correlation functionals. <i>Journal of Chemical Physics</i> , 2021, 155, 104103.	3.0	26
20	Layered CuI: a path to 2D p-type transparent conducting materials. <i>Journal of Materials Chemistry C</i> , 2021, 9, 11284-11291.	5.5	7
21	From pseudo-direct hexagonal germanium to direct silicon-germanium alloys. <i>Physical Review Materials</i> , 2021, 5, .	2.4	7
22	Crystal graph attention networks for the prediction of stable materials. <i>Science Advances</i> , 2021, 7, eabi7948.	10.3	37
23	Superconducting hydrogen tubes in hafnium hydrides at high pressure. <i>Physical Review B</i> , 2021, 104, .	3.2	11
24	Exchange-correlation functionals for band gaps of solids: benchmark, reparametrization and machine learning. <i>Npj Computational Materials</i> , 2020, 6, .	8.7	156
25	Validation of Pseudopotential Calculations for the Electronic Band Gap of Solids. <i>Journal of Chemical Theory and Computation</i> , 2020, 16, 3620-3627.	5.3	25
26	Electronic and Optical Properties of Small Metal Fluoride Clusters. <i>ACS Omega</i> , 2020, 5, 13268-13277.	3.5	8
27	Accurate electronic band gaps of two-dimensional materials from the local modified Becke-Johnson potential. <i>Physical Review B</i> , 2020, 101, .	3.2	21
28	Local Modified Becke-Johnson Exchange-Correlation Potential for Interfaces, Surfaces, and Two-Dimensional Materials. <i>Journal of Chemical Theory and Computation</i> , 2020, 16, 2654-2660.	5.3	42
29	Direct-bandgap emission from hexagonal Ge and SiGe alloys. <i>Nature</i> , 2020, 580, 205-209.	27.8	231
30	Novel two-dimensional silicon–carbon binaries by crystal structure prediction. <i>Physical Chemistry Chemical Physics</i> , 2020, 22, 8442-8449.	2.8	8
31	Atomic scale structure and its impact on the band gap energy for Cu ₂ Zn(Sn,Ge)Se ₄ kesterite alloys. <i>JPhys Energy</i> , 2020, 2, 035004.	5.3	3
32	Recent advances and applications of machine learning in solid-state materials science. <i>Npj Computational Materials</i> , 2019, 5, .	8.7	1,289
33	Large-Scale Benchmark of Exchange–Correlation Functionals for the Determination of Electronic Band Gaps of Solids. <i>Journal of Chemical Theory and Computation</i> , 2019, 15, 5069-5079.	5.3	151
34	Computational acceleration of prospective dopant discovery in cuprous iodide. <i>Physical Chemistry Chemical Physics</i> , 2019, 21, 18839-18849.	2.8	34
35	Mechanism of surface passivation of methylammonium lead tribromide single crystals by benzylamine. <i>Applied Physics Reviews</i> , 2019, 6, 031401.	11.3	34
36	On the calculation of the bandgap of periodic solids with MGGA functionals using the total energy. <i>Journal of Chemical Physics</i> , 2019, 151, 161102.	3.0	10

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37	Stable Ordered Phases of Cuprous Iodide with Complexes of Copper Vacancies. <i>Chemistry of Materials</i> , 2019, 31, 7877-7882.	6.7	17
38	Double perovskites as p-type conducting transparent semiconductors: a high-throughput search. <i>Journal of Materials Chemistry A</i> , 2019, 7, 14705-14711.	10.3	32
39	Structural prediction of stabilized atomically thin tin layers. <i>Npj 2D Materials and Applications</i> , 2019, 3, .	7.9	14
40	Out-of-plane excitons in two-dimensional crystals. <i>Physical Review B</i> , 2019, 99, .	3.2	30
41	Rare-earth magnetic nitride perovskites. <i>JPhys Materials</i> , 2019, 2, 025003.	4.2	25
42	Accurate electronic and optical properties of hexagonal germanium for optoelectronic applications. <i>Physical Review Materials</i> , 2019, 3, .	2.4	41
43	Stable hybrid organic-inorganic halide perovskites for photovoltaics from <i>ab initio</i> high-throughput calculations. <i>Journal of Materials Chemistry A</i> , 2018, 6, 6463-6475.	10.3	40
44	High-pressure phases of VO ₂ from the combination of Raman scattering and ab initio structural search. <i>Physical Review B</i> , 2018, 97, .	3.2	9
45	Local Hybrid Density Functional for Interfaces. <i>Journal of Chemical Theory and Computation</i> , 2018, 14, 939-947.	5.3	21
46	The ground state of two-dimensional silicon. <i>2D Materials</i> , 2018, 5, 035010.	4.4	25
47	Size-dependent optical absorption of Cu ₂ ZnSn(Se,S) ₄ quantum dot sensitizers from ab initio many-body methods. <i>European Physical Journal B</i> , 2018, 91, 1.	1.5	3
48	Predicting the stability of ternary intermetallics with density functional theory and machine learning. <i>Journal of Chemical Physics</i> , 2018, 148, 241728.	3.0	30
49	Nitrogen-hydrogen-oxygen ternary phase diagram: New phases at high pressure from structural prediction. <i>Physical Review Materials</i> , 2018, 2, .	2.4	17
50	Anisotropic layered Bi ₂ Te ₃ -In ₂ Te ₃ composites: control of interface density for tuning of thermoelectric properties. <i>Scientific Reports</i> , 2017, 7, 43611.	3.3	18
51	Structure and Optical Properties of Small (TiO ₂) _n Nanoparticles, <i>n</i> = 21-24. <i>Journal of Physical Chemistry C</i> , 2017, 121, 9528-9536.	3.1	7
52	Predicting the Thermodynamic Stability of Solids Combining Density Functional Theory and Machine Learning. <i>Chemistry of Materials</i> , 2017, 29, 5090-5103.	6.7	217
53	High-throughput search of ternary chalcogenides for p-type transparent electrodes. <i>Scientific Reports</i> , 2017, 7, 43179.	3.3	46
54	Pressure-induced radial collapse in few-wall carbon nanotubes: A combined theoretical and experimental study. <i>Carbon</i> , 2017, 125, 429-436.	10.3	27

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55	From mesoscale to nanoscale mechanics in single-wall carbon nanotubes. <i>Carbon</i> , 2017, 123, 145-150.		10.3	41
56	Structural prediction of two-dimensional materials under strain. <i>2D Materials</i> , 2017, 4, 045009.		4.4	19
57	Chemically Tunable Properties of Graphene Covered Simultaneously with Hydroxyl and Epoxy Groups. <i>Journal of Physical Chemistry C</i> , 2017, 121, 27603-27611.		3.1	6
58	Thermodynamic, electronic, and optical properties of graphene oxide: A statistical <i>ab initio</i> approach. <i>Physical Review B</i> , 2017, 95, .		3.2	16
59	Nanostructured water and carbon dioxide inside collapsing carbon nanotubes at high pressure. <i>Physical Chemistry Chemical Physics</i> , 2016, 18, 19926-19932.		2.8	16
60	Rigamonti <i>et Al.</i> Reply. <i>Physical Review Letters</i> , 2016, 117, 159702.		7.8	6
61	Research Update: Stable single-phase Zn-rich Cu ₂ ZnSnSe ₄ through In doping. <i>APL Materials</i> , 2016, 4, 070701.		5.1	11
62	Prediction and Synthesis of a Non-Zintl Silicon Clathrate. <i>Chemistry of Materials</i> , 2016, 28, 3711-3717.		6.7	15
63	Radial collapse of carbon nanotubes for conductivity optimized polymer composites. <i>Carbon</i> , 2016, 106, 64-73.		10.3	28
64	Publisher's Note: Optical properties of Cu-chalcogenide photovoltaic absorbers from self-consistent $\langle mml:math \text{ xmlns:mml="http://www.w3.org/1998/Math/MathML" } \rangle \langle mml:mrow \rangle \langle mml:mi \rangle G \langle /mml:mi \rangle \langle mml:mi \rangle W \langle /mml:mi \rangle \langle mml:mrow \rangle \langle mml:mi \rangle \text{the Bethe-Salpeter equation [Phys. Rev. B} \langle b \rangle 91 \langle /b \rangle, 075134 (2015)] \langle /mml:mrow \rangle \langle /mml:math \rangle$. <i>Physical Review B</i> , 2016, 93, .			
65	Topological Crystalline Insulator in a New Bi Semiconducting Phase. <i>Scientific Reports</i> , 2016, 6, 21790.		3.3	12
66	Investigation of new phases in the Ba–Si phase diagram under high pressure using ab initio structural search. <i>Physical Chemistry Chemical Physics</i> , 2016, 18, 8108-8114.		2.8	15
67	Stability and electronic properties of new inorganic perovskites from high-throughput ab initio calculations. <i>Journal of Materials Chemistry C</i> , 2016, 4, 3157-3167.		5.5	235
68	Novel crystal structures for lithium–silicon alloy predicted by minima hopping method. <i>Journal of Alloys and Compounds</i> , 2016, 655, 147-154.		5.5	21
69	Low-density silicon allotropes for photovoltaic applications. <i>Physical Review B</i> , 2015, 92, .		3.2	70
70	Detection of Cu ₂ Zn ₅ SnSe ₈ and Cu ₂ Zn ₆ SnSe ₉ phases in co-evaporated Cu ₂ ZnSnSe ₄ thin-films. <i>Applied Physics Letters</i> , 2015, 107, .		3.3	6
71	Optical properties of Cu-chalcogenide photovoltaic absorbers from self-consistent $\langle mml:math \text{ xmlns:mml="http://www.w3.org/1998/Math/MathML" } \rangle \langle mml:mrow \rangle \langle mml:mi \rangle G \langle /mml:mi \rangle \langle mml:mi \rangle W \langle /mml:mi \rangle \langle mml:mrow \rangle \langle mml:mi \rangle \text{the Bethe-Salpeter equation. Physical Review B}, 2015, 91, .$			
72	Identification of Novel Cu, Ag, and Au Ternary Oxides from Global Structural Prediction. <i>Chemistry of Materials</i> , 2015, 27, 4562-4573.		6.7	56

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73	Superconductivity in an expanded phase of ZnO: an <i>ab initio</i> study. <i>New Journal of Physics</i> , 2015, 17, 043034.		2.9	10
74	Optimized Exchange and Correlation Semilocal Functional for the Calculation of Energies of Formation. <i>Journal of Chemical Theory and Computation</i> , 2015, 11, 3844-3850.		5.3	21
75	Materials Design On-the-Fly. <i>Journal of Chemical Theory and Computation</i> , 2015, 11, 3955-3960.		5.3	25
76	Estimating Excitonic Effects in the Absorption Spectra of Solids: Problems and Insight from a Guided Iteration Scheme. <i>Physical Review Letters</i> , 2015, 114, 146402.		7.8	56
77	Novel phases of lithium-aluminum binaries from first-principles structural search. <i>Journal of Chemical Physics</i> , 2015, 142, 024710.		3.0	14
78	Prediction of Stable Nitride Perovskites. <i>Chemistry of Materials</i> , 2015, 27, 5957-5963.		6.7	102
79	Comment on "Towards Direct-Gap Silicon Phases by the Inverse Band Structure Design Approach". <i>Physical Review Letters</i> , 2014, 112, 199801.		7.8	2
80	Local versus global electronic properties of chalcopyrite alloys: X-ray absorption spectroscopy and <i>ab initio</i> calculations. <i>Journal of Applied Physics</i> , 2014, 116, 093703.		2.5	12
81	First-principles predicted low-energy structures of NaSc(BH ₄) ₄ . <i>Journal of Chemical Physics</i> , 2014, 140, 124708.		3.0	25
82	Benchmark Many-Body <i>GW</i> and Bethe-Salpeter Calculations for Small Transition Metal Molecules. <i>Journal of Chemical Theory and Computation</i> , 2014, 10, 3934-3943.		5.3	98
83	Density-functional tight-binding study of the collapse of carbon nanotubes under hydrostatic pressure. <i>Carbon</i> , 2014, 69, 355-360.		10.3	40
84	Carbon structures and defect planes in diamond at high pressure. <i>Physical Review B</i> , 2013, 88, .		3.2	32
85	Prediction of a novel monoclinic carbon allotrope. <i>European Physical Journal B</i> , 2013, 86, 1.		1.5	13
86	Conducting Boron Sheets Formed by the Reconstruction of the B_{12} Cluster. <i>Physical Review Letters</i> , 2013, 111, 136101.		7.8	40
87	Comment on "Topological Insulators in Ternary Compounds with a Honeycomb Lattice". <i>Physical Review Letters</i> , 2013, 110, 129701.		7.8	4
88	Ab Initio Electronic Gaps of Ge Nanodots: The Role of Self-Energy Effects. <i>Journal of Physical Chemistry C</i> , 2013, 117, 14229-14234.		3.1	12
89	Atomic and electronic properties of quasi-one-dimensional MoS ₂ nanowires. <i>Journal of Materials Research</i> , 2013, 28, 240-249.		2.6	20
90	Sodium-gold binaries: novel structures for ionic compounds from an <i>ab initio</i> structural search. <i>New Journal of Physics</i> , 2013, 15, 115007.		2.9	58

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91	Strong Renormalization of the Electronic Band Gap due to Lattice Polarization in the CuInSe_3 Formalism. <i>Physical Review Letters</i> , 2013, 110, 226404.	7.8	56	
92	Low-Energy Polymeric Phases of Alanates. <i>Physical Review Letters</i> , 2013, 110, 135502.	7.8	38	
93	The crystal structure of p-type transparent conductive oxide CuBO ₂ . <i>MRS Communications</i> , 2013, 3, 157-160.	1.8	12	
94	Low-energy silicon allotropes with strong absorption in the visible for photovoltaic applications. <i>Physical Review B</i> , 2012, 86, .	3.2	138	
95	Feedback mechanism for the stability of the band gap of CuInSe ₃ . <i>Physical Review B</i> , 2012, 86, .	3.2	29	
96	<math>\langle \text{mml:math} \text{ xmlns:mml="http://www.w3.org/1998/Math/MathML"} \text{ display="block">\rangle \langle \text{mml:mi} \text{ p} \langle / \text{mml:mi} \rangle \langle / \text{mml:math} \rangle \text{ Doping in Expanded Phases of ZnO: An Ab initio Study. Physical Review Letters, 2012, 108, 115903.}	7.8	18	
97	Speeding up the solution of the Bethe-Salpeter equation by a double-grid method and Wannier interpolation. <i>Physical Review B</i> , 2012, 86, .	3.2	42	
98	High-Pressure Structures of Disilane and Their Superconducting Properties. <i>Physical Review Letters</i> , 2012, 108, 117004.	7.8	86	
99	Raman activity of Cu _n Si ₂ allotropes under pressure: A density functional theory study. <i>Physical Review B</i> , 2012, 85, .			
100	Novel Structural Motifs in Low Energy Phases of LiAlH ₄ . <i>Physical Review Letters</i> , 2012, 108, 205505.	7.8	43	
101	Crystal Structure of Cold Compressed Graphite. <i>Physical Review Letters</i> , 2012, 108, 065501.	7.8	292	
102	Modeling van der Waals interactions between proteins and inorganic surfaces from time-dependent density functional theory calculations. <i>Physical Chemistry Chemical Physics</i> , 2011, 13, 15055.	2.8	5	
103	Band structures of Cu ₂ ZnSnS ₄ and Cu ₂ ZnSnSe ₄ from many-body methods. <i>Applied Physics Letters</i> , 2011, 98, 241915.	3.3	112	
104	Density-based mixing parameter for hybrid functionals. <i>Physical Review B</i> , 2011, 83, .	3.2	338	
105	First-principles study of the band structure and optical absorption of CuGaS ₂ . <i>Physical Review B</i> , 2011, 84, .	3.2	74	
106	Enhancing the Superconducting Transition Temperature of BaSi ₂ by Structural Tuning. <i>Physical Review Letters</i> , 2011, 106, 087002.	7.8	17	
107	Superconductivity in layered binary silicides: A density functional theory study. <i>Physical Review B</i> , 2011, 84, .	3.2	21	
108	Band structures of delafossite transparent conductive oxides from a self-consistent approach. <i>Physical Review B</i> , 2010, 82, .	3.2	63	

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109	Effects of Electronic and Lattice Polarization on the Band Structure of Delafossite Transparent Conductive Oxides. <i>Physical Review Letters</i> , 2010, 104, 136401.	7.8	88
110	Excitonic effects in the optical properties of CdSe nanowires. <i>Applied Physics Letters</i> , 2010, 96, .	3.3	24
111	Strong Interplay between Structure and Electronic Properties in $\text{CuIn}_x\text{Ga}_{1-x}\text{S}_3$. <i>Journal of the American Chemical Society</i> , 2009, 131, 10784-10791.	7.8	133
112	Alloying effects on the optical properties of $\text{Ge}_x\text{Si}_{1-x}$. <i>Physical Review B</i> , 2009, 79, .	8.0	111
113	Local atomic order and optical properties in amorphous and laser-crystallized GeTe. <i>Comptes Rendus Physique</i> , 2009, 10, 514-527.	0.9	17
114	Optical and magnetic properties of boron fullerenes. <i>Physical Chemistry Chemical Physics</i> , 2009, 11, 4523.	2.8	50
115	Cluster-surface and cluster-cluster interactions: <i>i</i> Ab initio <i>ii</i> calculations and modeling of asymptotic van der Waals forces. <i>Physical Review B</i> , 2008, 78, .	3.2	26
116	Time-dependent density functional theory scheme for efficient calculations of dynamic (hyper)polarizabilities. <i>Journal of Chemical Physics</i> , 2007, 126, 184106.	3.0	106
117	Time-dependent density-functional theory for extended systems. <i>Reports on Progress in Physics</i> , 2007, 70, 357-407.	20.1	201
118	Origin of the Optical Contrast in Phase-Change Materials. <i>Physical Review Letters</i> , 2007, 98, 236403.	7.8	162
119	Efficient calculation of van der Waals dispersion coefficients with time-dependent density functional theory in real time: Application to polycyclic aromatic hydrocarbons. <i>Journal of Chemical Physics</i> , 2007, 127, 014107.	3.0	31
120	Identification of fullerene-like CdSe nanoparticles from optical spectroscopy calculations. <i>Physical Review B</i> , 2007, 75, .	3.2	47
121	Growth dynamics of hydrogenated silicon nanoparticles under realistic conditions of a plasma reactor. <i>Computational Materials Science</i> , 2006, 35, 216-222.	3.0	21
122	TDDFT from molecules to solids: The role of long-range interactions. <i>International Journal of Quantum Chemistry</i> , 2005, 102, 684-701.	2.0	65
123	The planar-to-tubular structural transition in boron clusters from optical absorption. <i>Journal of Chemical Physics</i> , 2005, 123, 014310.	3.0	46
124	Comment on "Quantum Confinement and Electronic Properties of Silicon Nanowires". <i>Physical Review Letters</i> , 2005, 94, 219701; author reply 219702.	7.8	43
125	Energy dependence of the exchange-correlation kernel of time-dependent density functional theory: A simple model for solids. <i>Physical Review B</i> , 2005, 72, .	3.2	54
126	Ab initio and semiempirical dielectric response of superlattices. <i>Physical Review B</i> , 2004, 70, .	3.2	24

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127	Applications of Time-Dependent Density Functional Theory. <i>Physica Scripta</i> , 2004, T109, 54.	2.5	6
128	Long-range contribution to the exchange-correlation kernel of time-dependent density functional theory. <i>Physical Review B</i> , 2004, 69, .	3.2	184
129	Ab Initio Calculations of the Anisotropic Dielectric Tensor of GaAs/AlAs Superlattices. <i>Physical Review Letters</i> , 2002, 89, 216803.	7.8	35
130	Giant Optical Oscillator Strengths in Perturbed Hexagonal Germanium. <i>Physica Status Solidi - Rapid Research Letters</i> , 0, , 2100555.	2.4	4