

# CITATION REPORT

List of articles citing

## Self-Consistent Equations Including Exchange and Correlation Effects

DOI: 10.1103/physrev.140.a1133  
Physical Review, 1965, 140, A1133-A1138.

**Source:** <https://exaly.com/paper-pdf/8618323/citation-report.pdf>

**Version:** 2024-04-27

This report has been generated based on the citations recorded by exaly.com for the above article. For the latest version of this publication list, visit the link given above.

The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

#	Paper	IF	Citations
2200	One-Particle Properties of an Inhomogeneous Interacting Electron Gas. <i>Physical Review</i> , <b>1966</b> , 145, 561-567		799
2199	Crystal Potential and Correlation for Energy Bands in Valence Semiconductors. <i>Physical Review</i> , <b>1966</b> , 149, 597-613		97
2198	Statistical Approximation for Exchange in Self-Consistent-Field Calculations of the Ground State of Neutral Argon. <i>Physical Review</i> , <b>1966</b> , 144, 5-7		92
2197	Application of a Self-Consistent Scheme Including Exchange and Correlation Effects to Atoms. <i>Physical Review</i> , <b>1966</b> , 144, 1-4		232
2196	Modified Interionic Potential for the Alkali Metals. <i>Physical Review</i> , <b>1967</b> , 163, 667-675		49
2195	Electron Photodetachment from O and Elastic Scattering from Atomic Oxygen. <i>Physical Review</i> , <b>1967</b> , 153, 28-35		35
2194	Charge Density of Diamond. <i>Physical Review</i> , <b>1967</b> , 164, 1100-1105		17
2193	Self-Consistent-Field Model of Bimetallic Interfaces. I. Dipole Effects. <i>Physical Review</i> , <b>1967</b> , 160, 541-553		85
2192	Metallic Interfaces. II. Influence of the Exchange-Correlation and Lattice Potentials. <i>Physical Review</i> , <b>1967</b> , 162, 578-588		73
2191	"Muffin-Tin" Potential in Band Calculations. <i>Physical Review</i> , <b>1967</b> , 153, 704-705		22
2190	Approximate Wave Functions for Molecules and Crystals Using Perturbation Theory. <b>1967</b> , 39, 203-227		33
2189	Exchange Potential for Nearly Free Electrons. <i>Physical Review</i> , <b>1967</b> , 157, 515-517		18
2188	Relationship between the "Proton-Proton" and "Proton-Electron" Dielectric Constants. <i>Physical Review</i> , <b>1967</b> , 158, 670-671		14
2187	Miscibility of Molten Lithium and Sodium. <i>Physical Review</i> , <b>1967</b> , 159, 108-113		9
2186	Correlation Effects in Complex Spectra. II. Transition Probabilities for the Magnesium Isoelectronic Sequence. <b>1967</b> , 47, 3561-3572		92
2185	Relativistic Corrections to the Electron Density at the Nuclear Surface and to the Alkali Halide Overlap Integrals. <b>1967</b> , 46, 1929-1934		16
2184	Some non-structural aspects of the theory of metals. <b>1967</b> , 91, 701-723		100

2183	Low-Energy Interband Transitions and Band Structure in Nickel. <b>1967</b> , 19, 16-20	66
2182	Atomic Self-Consistent-Field Calculations Using Statistical Approximations for Exchange and Correlation. <i>Physical Review</i> , <b>1967</b> , 163, 54-61	246
2181	Energy Bands in Ferromagnetic Nickel. <i>Physical Review</i> , <b>1967</b> , 159, 415-426	301
2180	A density functional representation of quantum chemistry. I. Motivation and general formalism. <b>1967</b> , 1, 493-519	16
2179	Band structure, deformation potentials, and exciton states in solid xenon. <b>1967</b> , 28, 2067-2085	53
2178	High energy K conversion coefficients. <b>1968</b> , 28, 105-106	3
2177	Pseudopotential and the binding energy of aluminium. <b>1968</b> , 28, 61-62	3
2176	An application of the Kohn potential to electron states of crystalline Si. <b>1968</b> , 27, 467-468	
2175	Extension of the Korringa-Kohn-Rostoker method to the impurity problem. <b>1968</b> , 27, 138-139	9
2174	Internal conversion tables part I: K-, L-, M-shell conversion coefficients for Z=30 to Z=103. <b>1968</b> , 4, 1-11	1319
2173	Surface potential of a free electron metal in the RPA. <b>1968</b> , 11, 419-429	15
2172	The effects of screened exchange and correlation on the surface potential of an electron gas. <b>1968</b> , 11, 465-478	16
2171	Correlation Energy of an Electron Gas with a Slowly Varying High Density. <i>Physical Review</i> , <b>1968</b> , 165, 18-31	352
2170	A structural expansion of the cohesive energy of simple metals in an effective Hamiltonian approximation. <b>1968</b> , 1, 1620-1632	101
2169	Localization of electrons in condensed matter. <b>1968</b> , 31, 533-621	33
2168	Exchange Potential for Electrons in Atoms and Solids. <i>Physical Review</i> , <b>1968</b> , 171, 1-3	110
2167	Magnetic Ordering and the Electronic Properties of the Heavy Rare-Earth Metals. <i>Physical Review</i> , <b>1968</b> , 167, 497-503	62
2166	Application of the Orthogonalized-Plane-Wave Method to Lithium Chloride, Sodium Chloride, and Potassium Chloride. <i>Physical Review</i> , <b>1968</b> , 175, 1147-1155	90

2165	Perturbation-Theoretic Approach To Atoms and Molecules. <i>Physical Review</i> , <b>1968</b> , 173, 93-107	65
2164	l-Averaged Exchange Term in the Hartree-Fock Equations. <b>1968</b> , 48, 4140-4143	10
2163	Numerical Solution of Quantum-Mechanical Pair Equations. <b>1968</b> , 48, 5514-5523	46
2162	Exchange in Spin-Polarized Energy Bands. <i>Physical Review</i> , <b>1968</b> , 165, 658-669	51
2161	Relativistic Calculations of Electron Binding Energies by a Modified Hartree-Fock-Slater Method. <i>Physical Review</i> , <b>1968</b> , 176, 114-125	202
2160	Statistical model calculations of atoms embedded in an electron gas. <b>1968</b> , 1, 53-55	10
2159	Atomic susceptibilities from statistical electron models with kinetic, exchange and correlation corrections. <b>1969</b> , 2, 995-1000	1
2158	Electron-Ion Interaction and the Fermi Surfaces of the Alkali Metals. <i>Physical Review</i> , <b>1969</b> , 178, 953-965	102
2157	Crystal Potentials in Energy-Band Calculations of Noble Metals. <i>Physical Review</i> , <b>1969</b> , 184, 639-642	9
2156	Self-Consistent Augmented-Plane-Wave Method. <i>Physical Review</i> , <b>1969</b> , 181, 1024-1032	106
2155	Variation of Lattice Constant in Augmented-Plane-Wave Energy-Band Calculation for Lithium. <i>Physical Review</i> , <b>1969</b> , 181, 1033-1035	95
2154	Influence of the Lattice Potential on the Electronic Structure of Metallic Interfaces: Dipole Effects. <i>Physical Review</i> , <b>1969</b> , 188, 1060-1068	18
2153	Band Structure and Properties of Cesium Metal. <i>Physical Review</i> , <b>1969</b> , 183, 674-686	43
2152	Phonon Frequencies of Alkali Metals. <i>Physical Review</i> , <b>1969</b> , 187, 808-820	29
2151	Absolute Experimental X-Ray Form Factor of Aluminum. <i>Physical Review</i> , <b>1969</b> , 184, 607-613	31
2150	Symmetrized Relativistic Augmented-Plane-Wave Method: Gray Tin and the Warped Muffin-Tin Potential. <i>Physical Review</i> , <b>1969</b> , 188, 1049-1059	67
2149	Self-Consistent Orthogonalized-Plane-Wave and Empirically Refined Orthogonalized-Plane-Wave Energy-Band Models for Cubic ZnS, ZnSe, CdS, and CdSe. <i>Physical Review</i> , <b>1969</b> , 179, 740-751	144
2148	Energy-Band Structure of Aluminum Arsenide. <i>Physical Review</i> , <b>1969</b> , 188, 1193-1196	76

2147	Nonintegral Occupation Numbers in Transition Atoms in Crystals. <i>Physical Review</i> , <b>1969</b> , 184, 672-694	254
2146	Variational Principle and Slater's Generalized Hartree-Fock Theory for Nuclei. <i>Physical Review</i> , <b>1969</b> , 188, 1583-1589	5
2145	Electronic Structure of Point Defects and Impurities in Metals. <i>Physical Review</i> , <b>1969</b> , 186, 683-696	30
2144	Comparison of Dielectric Screening Methods Used in Phonon-Frequency Calculation of Normal Metals. <i>Physical Review</i> , <b>1969</b> , 185, 913-923	10
2143	Comparison of Several Exchange Potentials for Electrons in the Cu+Ion. <i>Physical Review</i> , <b>1969</b> , 179, 28-38	229
2142	Nonrelativistic Partial-Wave Scattering of Electrons by Ionized Atoms. <i>Physical Review</i> , <b>1969</b> , 177, 204-211	8
2141	Calculation of Constant-Energy Surfaces for Aluminum by the Korringa-Kohn-Rostoker Method. <i>Physical Review</i> , <b>1969</b> , 178, 914-917	26
2140	Local Exchange Approximation and the Virial Theorem. <i>Physical Review</i> , <b>1969</b> , 184, 10-17	54
2139	Pressure Calculations and the Virial Theorem for Modified Hartree-Fock Solids and Atoms. <i>Physical Review</i> , <b>1969</b> , 179, 612-615	45
2138	Interpolation between Slater's and Kohn-Sham's exchange approximations. <b>1969</b> , 19, 1199-1203	
2137	High-energy K-conversion coefficients for <sup>114</sup> Cd and <sup>150</sup> Sm. <b>1969</b> , 125, 637-640	4
2136	Theory of dielectric screening of an impurity at the surface of an electron gas. <b>1969</b> , 30, 2307-2319	38
2135	Some Numerical Results on Quasiparticle Properties in the Electron Gas. <b>1969</b> , 32, 273-280	156
2134	Relativistic and non-relativistic configuration interaction calculations for atoms having a closed core and two valence spin-orbitals. <b>1969</b> , 1, 113-134	7
2133	Band calculations for vanadium. <b>1969</b> , 7, 1439-1441	25
2132	Phonon dispersion-relations in beryllium calculated from a pseudopotential approach. <b>1969</b> , 7, 295-297	13
2131	Self-consistent properties of the electron distribution at a metal surface. <b>1969</b> , 7, 1047-1050	92
2130	A local approximation for exchange and correlation in band structure calculations. <b>1969</b> , 30, 1113-1118	10

2129	On the calculation of transition probabilities in the argon I spectrum. <b>1969</b> , 9, 1493-1506	11
2128	Reduction factor to the Slater exchange potential. <b>1969</b> , 29, 647-648	1
2127	The determination of the crystal potential for a calculation of phonon frequencies of normal metals. <b>1969</b> , 30, 123-124	7
2126	Lattice Dynamics of Metals. <b>1969</b> , 22, 159-312	43
2125	Optimized model potential: exchange and correlation corrections and calculation of magnesium phonon spectrum. <b>1969</b> , 2, 2071-2088	102
2124	Scattering Model of Molecular Electronic Structure. <b>1969</b> , 22, 1168-1171	65
2123	Analytic Independent-Particle Model for Atoms. <i>Physical Review</i> , <b>1969</b> , 184, 1-9	408
2122	Self-Consistent Many-Electron Theory of Electron Work Functions and Surface Potential Characteristics for Selected Metals. <i>Physical Review</i> , <b>1969</b> , 181, 522-529	418
2121	Improved Statistical Exchange Approximation for Inhomogeneous Many-Electron Systems. <b>1969</b> , 22, 807-811	329
2120	A first principle calculation of the total binding energy and c/a ratio of magnesium. <b>1970</b> , 32, 395-396	4
2119	The effect of different exchange approximations on a pseudopotential calculation of phonon dispersion relations in beryllium. <b>1970</b> , 31, 150-151	4
2118	Band structure and pressure-induced electronic transitions in calcium. <b>1970</b> , 8, 2109-2112	18
2117	Non-local pseudopotential calculation of the phonon dispersion relations in magnesium. <b>1970</b> , 8, 1199-1201	3
2116	Theory of Metal Surfaces: Charge Density and Surface Energy. <b>1970</b> , 1, 4555-4568	1700
2115	Relativistic calculation of atomic structures. <b>1970</b> , 19, 747-811	746
2114	Note on the minimization problem in the March-Young plane wave variational density matrix derivation of the Thomas-Fermi-Weizsäcker equation. <b>1970</b> , 32, 515-516	4
2113	Conduction-Electron Exchange and Correlation in Pseudopotential Calculations. <b>1970</b> , 41, K35-K38	4
2112	Relativistic electron band structure of gold. <b>1970</b> , 3, S109-S119	29

2111	Exchange effects in low-energy electron scattering. <b>1970</b> , 3, 536-538	11
2110	KKR-Z calculation of band structure in elementary semiconductors. <b>1970</b> , 3, 1884-1899	6
2109	Pseudopotentials and the Thomas-Fermi method for metals. <b>1970</b> , 3, S348-S354	7
2108	Inhomogeneity terms in exchange and correlation. <b>1970</b> , 3, L17-L20	1
2107	Calculation of Spin-Orbit Coupling Constants and Other Radial Parameters for the Actinide Ions Using Relativistic Wavefunctions. <b>1970</b> , 53, 809-820	69
2106	Comparison of Theoretical and Experimental Charge Densities for C, Si, Ge, and ZnSe. <b>1970</b> , 1, 756-763	63
2105	Screening of a Foreign Charge by an Electron Gas. <b>1970</b> , 1, 3645-3648	6
2104	Lattice Dynamics of Beryllium from a First-Principles Nonlocal Pseudopotential Approach. <b>1970</b> , 2, 1733-1742	42
2103	First-Principles Calculation of the Bulk Modulus of Diamond. <b>1970</b> , 1, 2574-2581	15
2102	Studies of the Statistical Exchange Approximation in First-Transition-Row Atoms and Ions: Mn <sup>+2</sup> Ion. <b>1970</b> , 2, 620-634	36
2101	Inhomogeneity Correction to Exchange Potential. <b>1970</b> , 2, 545-546	4
2100	Local Exchange Approximations and the Virial Theorem. <b>1970</b> , 1, 969-970	45
2099	Many-Body Contributions to Atomic Correlation Energies. <b>1970</b> , 1, 755-764	22
2098	Single-Parameter Free-Electron Exchange Approximation in Free Atoms. <b>1970</b> , 1, 37-38	69
2097	Spin-Polarized Energy Bands in Eu Chalcogenides by the Augmented-Plane-Wave Method. <b>1970</b> , 1, 4589-4603	140
2096	Cluster-Wave Approach to the Electronic Structures of Complex Molecules and Solids. <b>1970</b> , 24, 139-142	40
2095	Self-Consistent Energy Bands of Calcium by the Green's-Function Method. <b>1970</b> , 2, 4852-4856	17
2094	Orthogonalized-Plane-Wave Convergence of Some Tetrahedral Semiconductors. <b>1970</b> , 1, 4692-4700	34

2093	Energy-Band Structure of BeS, BeSe, and BeTe. <b>1970</b> , 2, 1852-1858	71
2092	Exchange Potentials in a Nonuniform Electron Gas. <b>1970</b> , 2, 874-876	47
2091	Effect of the Dielectric Function on the Phonon Spectrum of Magnesium. <b>1970</b> , 2, 3947-3952	14
2090	Exchange Study of Atomic Krypton and Tetrahedral Semiconductors. <b>1970</b> , 1, 779-790	25
2089	Proposed X <sup>α</sup> Method for Solids. <b>1970</b> , 2, 1506-1508	7
2088	Energy-Band Structure and Optical Spectrum of Grey Tin. <b>1970</b> , 2, 352-363	64
2087	Self-Consistent Energy Bands and Related Properties of Boron Phosphide. <b>1970</b> , 1, 4791-4797	22
2086	Tight-Binding Calculations for d Bands. <b>1970</b> , 1, 297-304	22
2085	Self-Consistent Orthogonalized-Plane-Wave Energy-Band Study of Silicon. <b>1970</b> , 1, 1635-1643	114
2084	Phonon Frequencies of the Hexagonal Metals Be, Mg, and Zn. <b>1970</b> , 1, 1468-1478	27
2083	New Statistical Theory of the Nuclear Surface. I. <b>1970</b> , 1, 98-111	20
2082	Band Structure, Spin Splitting, and Spin-Wave Effective Mass in Nickel. <b>1970</b> , 1, 305-314	40
2081	Electronic Structure and Optical Spectrum of Boron Arsenide. <b>1970</b> , 1, 3458-3463	36
2080	Band Structure and Fermi Surface of Ferromagnetic Nickel. <b>1970</b> , 1, 244-263	169
2079	Self-Consistent Orthogonalized-Plane-Wave Band Calculation on GaAs. <b>1970</b> , 1, 724-730	49
2078	SCF Relativistic Hartree-Fock Calculations on the Superheavy Elements 118-131. <b>1970</b> , 53, 2397-2406	90
2077	Beyond the Local-Density Approximation: Surface Properties of (110) W. <b>1970</b> , 25, 1023-1026	45
2076	Effects of Electron-Electron and Electron-Phonon Interactions on the One-Electron States of Solids. <b>1970</b> , 23, 1-181	568



2075	Pseudopotential Theory of Cohesion and Structure. <b>1970</b> , 24, 249-463	287
2074	New Potential Function for Atomic and Solid-State Calculations. <b>1970</b> , 2, 244-248	63
2073	Relativistic Calculation of Anomalous Scattering Factors for X Rays. <b>1970</b> , 53, 1891-1898	4330
2072	Gravitation-Induced Electric Field near a Metal. II. <b>1970</b> , 1, 4649-4654	27
2071	Exchange and correlation in the theory of simple metals. <b>1970</b> , 3, 1140-1158	169
2070	Explicit local exchange-correlation potentials. <b>1971</b> , 4, 2064-2083	3034
2069	Nonrelativistic Auger Rates, X-Ray Rates, and Fluorescence Yields for the 2p Shell. <b>1971</b> , 4, 2164-2170	98
2068	Need for a Nonlocal Correlation Potential in Silicon. <b>1971</b> , 4, 1910-1917	48
2067	The Calculation of Electronic Energy Bands by the Augmented Plane Wave Method. <b>1971</b> , 26, 103-274	72
2066	Ab Initio Calculation of the Electronic Structure and Optical Properties of Diamond Using the Discrete Variational Method. <b>1971</b> , 4, 3610-3622	198
2065	Theory of Work-Function Changes Induced by Alkali Adsorption. <b>1971</b> , 4, 4234-4244	383
2064	Simplified Theory of Electron Correlations in Metals. <b>1971</b> , 3, 1888-1898	303
2063	Incorporation of correction terms into the self-consistent calculation of screening in metals. <b>1971</b> , 23, 1329-1338	6
2062	A first principle calculation of binding energies, structures and lattice constants of simple metals. <b>1971</b> , 37, 335-336	5
2061	A first principles calculation of the Knight shift of sodium. <b>1971</b> , 36, 133-134	8
2060	A first principle pseudopotential calculation of the elastic shear constants of magnesium. <b>1971</b> , 32, 761-768	12
2059	Local exchange-correlation potentials. <b>1971</b> , 9, 537-541	93
2058	Electron correlations at metallic densities. <b>1971</b> , 52, 481-488	30

2057	Effects of exchange on the band structure of zinc and cadmium. <b>1971</b> , 3, 34-44	21
2056	A self-consistent muffin-tin potential. <b>1971</b> , 2, 608-610	1
2055	Theory of Metal Surfaces: Work Function. <b>1971</b> , 3, 1215-1223	1166
2054	Relativistic self-consistent field program for atoms and ions. <b>1971</b> , 2, 107-113	430
2053	Thermodynamic properties of compressed solid hydrogen. <b>1971</b> , 7, 621-636	68
2052	Nonrelativistic K-shell auger rates and matrix elements for 4 <b>1971</b> , 3, 301-315	62
2051	Bounds for the Hartree-Fock effective exchange interaction and the static dielectric constants. <b>1971</b> , 4, L165-L168	6
2050	Effect of exchange potential on the band structure of gold. <b>1971</b> , 1, 169-176	9
2049	Density functional theory and the von Weizsacker method. <b>1971</b> , 4, 1322-1330	95
2048	Inert Gases in Solids: Interatomic Potentials and Their Influence on Rare-Gas Mobility. <b>1971</b> , 3, 3984-3992	87
2047	Self-Consistent Electronic Structure of Titanium. II. <b>1971</b> , 4, 4261-4273	34
2046	Lattice Spacings and Compressibilities vs Pauling Radii and Valencies. <b>1971</b> , 4, 3287-3291	41
2045	Low-Energy Electron-Diffraction Intensity Calculations for Beryllium with a Realistic Crystal Potential. <b>1971</b> , 3, 3228-3243	68
2044	Lattice Dynamics of Magnesium from a First-Principles Nonlocal Pseudopotential Approach. <b>1971</b> , 3, 2485-2496	33
2043	Self-Consistent Energy Bands of Cu via the X $\alpha$ Method. <b>1971</b> , 3, 3093-3096	9
2042	Comparison of the Energy-Band Structure of Ge-Si with Those of Si and Ge. <b>1971</b> , 3, 3347-3356	27
2041	Self-Consistent Energy Bands and Cohesive Energy of Be Using Two Forms of the Warped-Muffin-Tin Potential. <b>1971</b> , 4, 3284-3287	14
2040	Virial Theorem in Self-Consistent-Field Calculations. <b>1971</b> , 3, 2081-2082	131

2039	Self-Consistent Warped-Muffin-Tin-Potential Energy Bands of Be with Various Exchange Approximations. <b>1971</b> , 3, 2418-2424	9
2038	Calculated Spin-Orbit Splittings of Some Group IV, III-V, and II-VI Semiconductors. <b>1971</b> , 4, 1296-1306	77
2037	Exact Screened Calculations of Atomic-Field Pair Production and Low-Energy Screening. <b>1971</b> , 4, 1835-1843	48
2036	Perturbation Treatment of the Hartree-Fock-Slater (X <sub>F</sub> ) Equations. <b>1971</b> , 4, 1774-1778	9
2035	Scattering of Slow Electrons by Neon and Argon. <b>1971</b> , 4, 1425-1431	38
2034	Quadrupole Antishielding Factors, and the Nuclear Quadrupole Moments of Several Alkali Isotopes. <b>1971</b> , 3, 837-848	131
2033	Relativistic One-Electron Calculations of Shielded Atomic Hyperfine Constants. <b>1971</b> , 3, 6-12	17
2032	Z Dependence of the KLL Auger Rates. <b>1971</b> , 3, 519-520	40
2031	Nonrelativistic Auger Rates, X-Ray Rates, and Fluorescence Yields for the K Shell. <b>1971</b> , 3, 1919-1927	162
2030	Validity of the Potentials of Hg. <b>1971</b> , 3, 1793-1795	3
2029	Evaluation of Approximations to Hartree-Fock Exchange. <b>1971</b> , 3, 1535-1543	9
2028	Effects of the Coulomb Correlation on the Calculated Results for Atoms with and without Spin Polarization. <b>1971</b> , 3, 1224-1230	43
2027	Exact Screened Calculations of Atomic-Field Bremsstrahlung. <b>1971</b> , 3, 100-115	243
2026	Lattice Dynamics of Transition Metals—Application to Paramagnetic Nickel. <b>1971</b> , 4, 1770-1778	17
2025	Electrical Resistivity of Potassium from 1 to 25 °K. <b>1971</b> , 4, 4215-4225	94
2024	Relativistic effects in low energy electron scattering from atoms. <b>1971</b> , 20, 257-323	213
2023	On the inclusion of Coulomb correlation effects in atomic and band structure calculations. <b>1972</b> , 5, L188-L190	2
2022	The stability of negative ions in liquid metals. <b>1972</b> , 2, 289-296	9

2021	Crystal potentials and the band electrons of iridium. <b>1972</b> , 2, 1033-1045	21
2020	A study of the electronic energy bands for Band Cerium. <b>1972</b> , 2, 450-458	7
2019	A k dependent potential for bands in metals I. Theory. <b>1972</b> , 2, 1055-1061	3
2018	A selfconsistent calculation of the rigid neutral atom density according to the auxiliary neutral atom model. <b>1972</b> , 5, 2333-2344	54
2017	Comparison of Screened Exchange with the Slater Approximation for Silicon. <b>1972</b> , 5, 1493-1499	38
2016	Renormalized Atoms and the Band Theory of Transition Metals. <b>1972</b> , 5, 3953-3971	247
2015	Calculation of the Cohesive Energies and Bulk Properties of the Alkali Metals. <b>1972</b> , 6, 3637-3642	64
2014	SCF Dirac-Slater Calculations for an Ion in a Dielectric Medium. <b>1972</b> , 57, 967-972	7
2013	Variational Calculation of the Image Potential near a Metal Surface. <b>1972</b> , 6, 1122-1130	154
2012	Augmented-Plane-Wave Calculations of the Electronic Structure of Intermetallic Compounds YCu and YZn. <b>1972</b> , 6, 939-945	61
2011	Nuclear Contact Densities for Electrons in Sodium. <b>1972</b> , 6, 1195-1198	3
2010	Spin-Orbit Effects in the Index-of-Refractive Formalism for Polarized Neutrons. <b>1972</b> , 28, 596-599	1
2009	Screening Effects in Atomic Pair Production below 5 MeV. <b>1972</b> , 6, 2049-2056	40
2008	Analysis of the Electronic Exchange in Atoms. <b>1972</b> , 5, 542-550	86
2007	Behavior of Electron Wave Functions near the Atomic Nucleus and Normalization Screening Theory in the Atomic Photoeffect. <b>1972</b> , 5, 1063-1072	72
2006	Local Exchange-Correlation Potentials and the Fermi Surface of Copper. <b>1972</b> , 6, 4367-4370	66
2005	Metallic-Field Effect and Its Consequences in Field Emission, Field Ionization, and the Capacitance of a Capacitor. <b>1972</b> , 6, 801-812	64
2004	Exchange Approximations Used in the Energy-Band Calculations of Metals. <b>1972</b> , 5, 1221-1225	10

2003	Self-Consistent Energy Bands in Vanadium at Normal and Reduced Lattice Spacings. <b>1972</b> , 5, 1214-1221	105
2002	Kohn-Sham Self-Consistent Calculation of the Structure of Metallic Sodium. <b>1972</b> , 6, 1189-1194	39
2001	Correlation Potentials in a Nonuniform Electron Gas. <b>1972</b> , 5, 2799-2810	29
2000	Semi-empirical band structure theory. <b>1972</b> , 35, 883-947	13
1999	Dynamic Contributions to the Surface Energy of Simple Metals. <b>1972</b> , 6, 1134-1142	67
1998	V Optical Absorption Strengths of Defects in Insulators. <b>1972</b> , 10, 165-228	13
1997	X-Ray Fluorescence Yields, Auger, and Coster-Kronig Transition Probabilities. <b>1972</b> , 44, 716-813	1724
1996	Self-Consistent Electronic Structure of Solid Surfaces. <b>1972</b> , 6, 2166-2177	227
1995	Relativistic Effects in the Hyperfine Structure of the Alkali Atoms. <b>1972</b> , 6, 109-121	69
1994	The Auger Effect. <b>1972</b> , 8, 163-284	49
1993	Hellmann-Feynman and Virial Theorems in the X $\alpha$ Method. <b>1972</b> , 57, 2389-2396	183
1992	Electron Distribution around Mobile and Fixed Point Charges in Metals. <b>1972</b> , 5, 2109-2118	120
1991	Self-Consistent-FieldX $\alpha$ Cluster Method for Polyatomic Molecules and Solids. <b>1972</b> , 5, 844-853	797
1990	Statistical Exchange-Correlation in the Self-Consistent Field. <b>1972</b> , 6, 1-92	909
1989	Energy Bands in Ferromagnetic Nickel. <b>1972</b> , 5, 124-134	77
1988	An analytic estimate of the field penetration into metal surfaces. <b>1972</b> , 2, 1124-1136	21
1987	Kanzaki Forces and Electron Theory of Displaced Charge in Relaxed Defect Lattices. <b>1972</b> , 111-138	1
1986	Self-consistent-field X $\alpha$ cluster calculations for the ground state Ne <sub>2</sub> molecule. <b>1972</b> , 16, 81-85	32

1985	Exchange and correlation potentials for K-shell electrons. <b>1972</b> , 14, 12-18	9
1984	A correction to the Liberman approximation for the exchange energy. <b>1972</b> , 40, 315-316	1
1983	Correlation and exchange contributions to the charge density variations of an electron gas in the metallic range. <b>1972</b> , 10, 279-285	27
1982	Method for calculating integrals over Fermi surface orbits. <b>1972</b> , 10, 833-836	12
1981	Theory of inhomogeneous magnetic electron gas. <b>1972</b> , 10, 1157-1160	111
1980	Screening in a spin-polarized electron liquid. <b>1972</b> , 11, 149-153	47
1979	Pseudopotential Calculation of Binding Energies, Structures, and Lattice Constants of Simple Metals. <b>1972</b> , 51, 107-114	16
1978	Dependence of the Atomic Magnetic Moment on the Lattice Parameter in Iron. <b>1972</b> , 53, K121-K124	1
1977	Resolution of apparent APW/APW energy band differences in NaCl. <b>1972</b> , 49, 173-177	9
1976	Berechnung der Bandstruktur vom Vanadiumcarbid VC. <b>1972</b> , 24, 170-190	21
1975	On the influence of different statistical exchange approximations on a pseudopotential calculation of the cohesive energies, structures and bulk properties of magnesium and aluminium. <b>1972</b> , 5, 503-507	3
1974	A local exchange-correlation potential for the spin polarized case. i. <b>1972</b> , 5, 1629-1642	4762
1973	Properties of Pseudoatoms Derived from the Shaw Model Potential. <b>1973</b> , 55, 537-546	3
1972	On the Influence of Electronic Exchange and Correlation on Pseudopotential Calculations I. Form Factors, Energy-Wave Number Characteristics, and Effective Interionic Pair Potentials. <b>1973</b> , 56, 579-590	20
1971	Magnetic Moment Calculation in Disordered Systems: $\beta$ Manganese. <b>1973</b> , 57, 571-585	4
1970	The Band Structure of VO by the APW-LCAO Method. <b>1973</b> , 60, 587-594	7
1969	Calculations of potential barriers using the SCF X $\alpha$ method NH <sub>3</sub> and H <sub>2</sub> O <sub>2</sub> . <b>1973</b> , 20, 246-250	19
1968	Nonlinear response corrections for screening in the RPA. <b>1973</b> , 46, 65-66	3

1967	Self-consistent molecular Hartree-Fock-Blater calculations II. The effect of exchange scaling in some small molecules. <b>1973</b> , 2, 52-59	405
1966	Compressibility of copper via the self-consistent APW method. <b>1973</b> , 13, 1775-1777	3
1965	Thermo-electrostatic effects in superconductors. <b>1973</b> , 65, 587-605	11
1964	Total electronic energy of a noble metal ordered alloy in the pseudopotential method. <b>1973</b> , 16, 1053-1059	
1963	Theory of Metal Surfaces: Induced Surface Charge and Image Potential. <b>1973</b> , 7, 3541-3550	796
1962	Influence of the surface charge on the metal surface energy. <b>1973</b> , 38, 499-502	4
1961	An analytical model for evaluating the metal-metal surface energy. <b>1973</b> , 38, 503-505	17
1960	A theory of adhesion at a bimetallic interface: Overlap effects. <b>1973</b> , 38, 77-92	85
1959	The electron distribution in silicon - II. Theoretical interpretation. <b>1973</b> , 332, 239-254	48
1958	Phonon Frequencies and Cohesive Energies of Copper, Silver, and Gold. <b>1973</b> , 8, 5532-5544	33
1957	Surface Forces and the Jellium Model. <b>1973</b> , 31, 1218-1221	119
1956	Exchange and Correlation in the Electron Gas. <b>1973</b> , 7, 4357-4363	61
1955	Self-Consistent Band Structure of Niobium at Normal and Reduced Lattice Spacings. <b>1973</b> , 7, 5115-5121	87
1954	Self-Consistent Calculation of Energy Bands in Ferromagnetic Nickel. <b>1973</b> , 7, 1096-1103	177
1953	Atomic Photoelectric Effect Above 10 keV. <b>1973</b> , 45, 273-325	305
1952	The effective exchange and correlation potential for metal surfaces. <b>1973</b> , 3, 2143-2156	53
1951	Some electronic properties of solid surfaces. <b>1973</b> , 4, 261-277	2
1950	Surface energy and curvature energy of electron-hole droplets. <b>1973</b> , 6, L433-L436	23

1949	An APW calculation of the energy band in diamond using different approximations for the exchange energy. <b>1973</b> , 6, 3430-3437	7
1948	Dielectric screening and phonon dispersion in hcp zinc: a pseudopotential approach. <b>1973</b> , 3, 709-720	20
1947	Pseudopotential calculation of charge densities in Na and Li: comparison with selfconsistent results. <b>1973</b> , 3, 1678-1682	7
1946	Screened ions in perfect solids-a model calculation. <b>1973</b> , 6, 3385-3402	8
1945	Lattice Dynamics of Metallic Lithium. <b>1973</b> , 34, 26-35	15
1944	Inhomogeneous Electron Gas. <b>1973</b> , 7, 1912-1919	616
1943	Hartree-Fock Lattice Constant and Bulk Modulus of Diamond. <b>1973</b> , 8, 4019-4025	45
1942	Energy Bands in Ferromagnetic Iron. <b>1973</b> , 7, 4242-4252	149
1941	General Crystalline Hartree-Fock Formalism: Diamond Results. <b>1973</b> , 7, 818-831	123
1940	Wannier Functions in a Simple Nonperiodic System. <b>1973</b> , 8, 2485-2495	106
1939	Electronic structure calculations for large planar molecules by SCF-scattered wave method. <b>1973</b> , 59, 3723-3731	29
1938	Theory of Physisorption: He on Metals. <b>1973</b> , 8, 5484-5495	102
1937	Analysis of Low-Energy-Electron Diffraction Intensity Profiles from the (100) and (111) Faces of Nickel. <b>1973</b> , 8, 515-527	56
1936	Relativistic Energy Band Structure and Properties of Uranium. <b>1973</b> , 7, 4454-4463	39
1935	One-Electron Interpretation of Optical Absorption and Soft-X-Ray Data in MgO. <b>1973</b> , 8, 5920-5933	47
1934	Charge Densities in the Presence of Strong Attractive Impurity Potentials. <b>1973</b> , 7, 4478-4484	6
1933	Soft-X-Ray Absorption Threshold in Metals, Semiconductors, and Alloys. <b>1973</b> , 7, 2215-2229	22
1932	Dispersion contributions to surface energy. <b>1973</b> , 59, 6157-6162	42



1931	Orbital-Correction Method. <b>1973</b> , 7, 1876-1888	9
1930	Augmented-Plane-Wave Hartree-Fock First-Principles Calculation of the Fermi Surfaces of Li, Na, and K. <b>1973</b> , 8, 1281-1292	53
1929	Electronic Structure of Calcium as a Function of the Lattice Constant. <b>1973</b> , 7, 674-684	49
1928	Total Energy as a Function of Lattice Parameter for Copper via the Self-Consistent Augmented-Plane-Wave Method. <b>1973</b> , 8, 5391-5397	74
1927	Low-Energy-Electron Scattering Amplitudes. II. Validity of Free-Electron-Gas Exchange Approximations Applied to Electrons Incident on Hg. <b>1973</b> , 7, 1251-1256	6
1926	Energy Bands in Paramagnetic Chromium. <b>1973</b> , 8, 5398-5403	85
1925	Study of the Electronic Structure and the Optical Properties of the Solid Rare Gases. <b>1973</b> , 8, 779-794	91
1924	Electronic-Structure Studies of Solids. II. "Exact" Hartree-Fock Calculations for Cubic Atomic-Hydrogen Crystals. <b>1973</b> , 7, 2850-2866	61
1923	Influence of Conduction-Band Core Electron Exchange and Correlation on Pseudopotential Calculations. <b>1973</b> , 8, 1303-1307	17
1922	Exchange Energies and Potentials at Finite Temperatures. <b>1973</b> , 7, 1429-1431	5
1921	Ionic soft sphere parameters from Hartree-Fock charge densities. <b>1973</b> , 58, 1905-1907	12
1920	Direct Comparison of Core-Electron Binding Energies of Surface and Bulk Atoms of Ti, Cr, and Ni. <b>1973</b> , 30, 846-849	55
1919	Electron Exchange Energy in Si Inversion Layers. <b>1973</b> , 30, 278-280	132
1918	Application de la méthode de l'atome neutre auxiliaire au calcul de l'énergie des métaux simples. <b>1973</b> , 34, 879-889	28
1917	Nonlocal Exchange and Correlation Effects on the Fermi Surface of Metals. <b>1974</b> , 32, 297-301	35
1916	Investigations of nonlocal exchange and correlation effects in metals via the density-functional formalism. <b>1974</b> , 10, 4195-4204	44
1915	Independent-particle-model study of the elastic scattering of low-energy electrons by positive ions. <b>1974</b> , 10, 1623-1632	16
1914	Polarization correlations in atomic-field pair production. <b>1974</b> , 9, 752-761	10

1913	Momentum density and Compton profile of the inhomogeneous interacting electronic system. I. Formalism. <b>1974</b> , 9, 5122-5127	161
1912	Lattice dynamics of aluminum: An investigation of exchange and correlation effects. <b>1974</b> , 9, 4138-4150	36
1911	Electronic structure and optical properties of layered dichalcogenides: TiS <sub>2</sub> and TiSe <sub>2</sub> . <b>1974</b> , 9, 481-486	120
1910	Charge on an electron-hole drop. <b>1974</b> , 9, 1540-1546	49
1909	Comment on the Hartree-Fock theory for hyperfine interactions. <b>1974</b> , 9, 2238-2241	10
1908	Calculation of excitation energies of atomic systems using the operator $O^A O^A$ . <b>1974</b> , 10, 1034-1040	22
1907	Optical properties of molybdenum. II. Theory. <b>1974</b> , 10, 1290-1297	32
1906	Simplification of total-energy and pressure calculations in solids. <b>1974</b> , 9, 3985-3988	192
1905	Quadrupole shielding and antishielding factors for the excited nd states of the alkali-metal atoms. <b>1974</b> , 9, 1783-1793	22
1904	Reduced fermion density matrices. II. Electron density of Kr. <b>1974</b> , 61, 3394-3400	8
1903	Exchange and correlation potentials in a Fermi gas. <b>1974</b> , 10, 4918-4924	8
1902	Nonlinear, Self-Consistent Theory of Proton Screening in Metals Applied to Hydrogen in Al and Mg. <b>1974</b> , 33, 1164-1167	120
1901	Analysis of low-energy-electron diffraction intensity profiles from the (100) and (111) faces of copper. <b>1974</b> , 9, 1204-1215	43
1900	Calculation of the total energy in the multiple scattering-X $\beta$ method. I. General theory. <b>1974</b> , 61, 3063-3070	86
1899	Zero-order pseudoatoms and the generalized pseudopotential theory. <b>1974</b> , 10, 3075-3091	59
1898	Effect of nonadditive forces on the binding of closed-shell adsorbates on closed-shell substrates. I. Ar/Ar(100). <b>1974</b> , 9, 741-747	17
1897	Gradient term in the Kohn-Sham exchange-correlation potential. <b>1974</b> , 10, 2221-2225	54
1896	Localized wave functions in the theory of cohesion. I. General formalism. <b>1974</b> , 9, 3169-3179	7

1895	Momentum density and Compton profile of the inhomogeneous interacting electron system. II. Application to atoms. <b>1974</b> , 9, 5128-5132	36
1894	Relativistic Effects in The Many-Electron Atom. <b>1974</b> , 10, 1-52	47
1893	Dependence of atomic Compton profiles upon the Xalphaexchange parameter. <b>1974</b> , 7, 3655-3660	16
1892	Calculation of the band structure, Fermi surface, and interband optical conductivity of lithium. <b>1974</b> , 4, 380-393	20
1891	Electronic Energy Band Structure of the Actinide Metals. <b>1974</b> , 51-108	25
1890	A first-principles generalized pseudopotential calculation for the phonon spectra of zinc. <b>1974</b> , 4, L106-L110	22
1889	Green function theory of a dense electron gas with surface. I. Work function. <b>1974</b> , 7, 2221-2233	9
1888	Band structure of solid o-H2: a simplified KKR approach. <b>1974</b> , 7, 2467-2475	8
1887	Chemical potentials and workfunctions of metal surfaces in the theory of an inhomogeneous electron gas. <b>1974</b> , 7, L370-L373	27
1886	Transition State Calculations of Oscillator Strengths in theXlocal Exchange Approximation. <b>1974</b> , 9, 104-108	35
1885	Electronic and optical properties of metallic tin. <b>1974</b> , 4, 1359-1370	23
1884	The crystal-field integrals of V3Ga. <b>1974</b> , 7, 1117-1135	7
1883	The calculation of heavy atom ionization potentials by the relativistic transition state approximation. <b>1974</b> , 28, 482-484	3
1882	Effects of the exchange parameters on the binding energy in the SCF Xscattered wave method. <b>1974</b> , 27, 415-418	9
1881	Nonlocal XExchange for s- and d-Electrons in Transition Metals. <b>1974</b> , 62, 193-200	11
1880	Density Functional Theory and Statistical Pseudopotentials. <b>1974</b> , 62, K79-K81	
1879	Effects of Pressure on the Electronic Structure of Transition d-Metals. <b>1974</b> , 63, 11-50	38
1878	Linearized Effective Exchange and Correlation Potentials for One-Particle Property Calculations in Simple Metals. <b>1974</b> , 65, 481-491	

1877	Critical Investigation of Potentials for Band Structure Calculations. <b>1974</b> , 65, 493-503	3
1876	Adsorption of xenon on group VIII and IB metals studied by photoelectric work function measurements. <b>1974</b> , 5, 418-428	51
1875	Contribution of collective excitations to the surface energy of metals. <b>1974</b> , 14, 979-981	8
1874	The band structure of VO by the APW-LCAO method. <b>1974</b> , 15, 991-995	5
1873	A self-consistent theory for exchange and correlation effects in electron gas. <b>1974</b> , 14, 313-315	9
1872	On the susceptibility sum rule in the electron liquid. <b>1974</b> , 11, 156-158	4
1871	Scattering form factors of atoms, molecules and solids. <b>1974</b> , 11, 159-160	
1870	Application of Harrison's first-principle pseudopotential method for semiconductors. <b>1974</b> , 36, 297-310	2
1869	A survey on exchange potentials in atoms and solids. <b>1974</b> , 37, 219-252	11
1868	Statistical exchange for electron in shell and the X $\alpha$ method. <b>1974</b> , 35, 213-218	65
1867	Effect of correlation on the band structure of zinc. <b>1974</b> , 20, 236-246	10
1866	Correlated crystal potentials in band structure calculations. <b>1974</b> , 23, 1-14	16
1865	Exchange and correlation contribution to one-body potential in metallic Be. <b>1974</b> , 23, 15-26	17
1864	Quasi-particle energies in aluminium. <b>1974</b> , 23, 37-50	11
1863	Optical constants of transition metals: Ti, V, Cr, Mn, Fe, Co, Ni, and Pd. <b>1974</b> , 9, 5056-5070	1191
1862	Contribution to the cohesive energy of simple metals: Spin-dependent effect. <b>1974</b> , 10, 1319-1327	292
1861	Wannier functions and self-consistent metal calculations. <b>1974</b> , 10, 382-383	27
1860	Band structure of nickel: Spin-orbit coupling, the Fermi surface, and the optical conductivity. <b>1974</b> , 9, 4897-4907	396

1859	Electronic structure and optical properties of metallic calcium. <b>1974</b> , 10, 1369-1383	38
1858	Electronic impurity levels in semiconductors. <b>1974</b> , 37, 1099-1210	232
1857	Electron Bremsstrahlung from Neutral Atoms. <b>1974</b> , 33, 516-518	18
1856	The Density-Functional Formalism and the Electronic Structure of Metal Surfaces. <b>1974</b> , 28, 225-300	311
1855	Theory of lattice contraction at aluminium surfaces. <b>1974</b> , 4, L37-L41	296
1854	Theory of electronic properties of surfaces. <b>1975</b> , 1-39	7
1853	Calculation of the knight shift in the system CaCd <sub>1-x</sub> Tl <sub>x</sub> by using relativistic APW wave functions. <b>1975</b> , 79, 1064-1071	5
1852	Effect of correlation on the band structure of cerium. <b>1975</b> , 4, 45-52	11
1851	An improved model potential for liquid lithium. <b>1975</b> , 30, 253-256	1
1850	Multiple scattering formalism for a molecular Dirac equation. <b>1975</b> , 35, 51-56	16
1849	Non-linear self-consistent calculation of the electron density distribution in metallic lithium. <b>1975</b> , 55, 57-58	3
1848	Calculation method for the inhomogeneous electron gas. <b>1975</b> , 17, 327-330	34
1847	Exchange-correlation in inhomogeneous systems. A comment on the surface energy problem. <b>1975</b> , 17, 581-584	41
1846	Self-consistent Hartree-Fock calculation of exchange effects in a weakly inhomogeneous electron gas. <b>1975</b> , 16, 243-246	42
1845	The exchange-correlation energy of a metallic surface. <b>1975</b> , 17, 1425-1429	723
1844	Self-consistent electron densities in metal vacancies. <b>1975</b> , 16, 831-834	22
1843	A study on the tight-binding method. <b>1975</b> , 36, 557-561	4
1842	Contact potential difference measurements on polycrystalline Ni during and after deposition with evaporated Ni in the temperature range 230° ? T ? 450°C. <b>1975</b> , 28, 93-106	17

1841	Exchange and correlation in nearly free-electron metals. <b>1975</b> , 68, 645-649	
1840	Electrons in a Well Potential: Density of States, Surface Energy, and Phase Rule for Wave Functions. <b>1975</b> , 70, 555-566	25
1839	Densité de valence et énergie de liaison d'un métal simple par la méthode de l'atome neutre : le potentiel ionique Hartree-Fock. <b>1975</b> , 36, 521-529	18
1838	Green's-function method for crystal films and surfaces. <b>1975</b> , 11, 3756-3760	37
1837	Inherently self-consistent procedure for obtaining electronic structure: Results for a lithium particle. <b>1975</b> , 12, 4238-4246	17
1836	Analytic calculation of metal surface dipole moments in the step-potential approximation. <b>1975</b> , 12, 3503-3508	23
1835	Approximations for the exchange potential in electron scattering. <b>1975</b> , 63, 2182-2191	427
1834	Extended Anderson-Grimley model for chemisorption: Solutions self-consistent within a surface complex. <b>1975</b> , 11, 3739-3755	18
1833	X-ray photoelectron spectrum and band structure of TiC. <b>1975</b> , 12, 5465-5471	70
1832	Orthogonalized-Plane-Wave Band Structure of Polymeric Sulfur Nitride, (SN) <sub>x</sub> . <b>1975</b> , 35, 1799-1803	75
1831	Gradient expansion in the density functional approach to an inhomogeneous electron system. <b>1975</b> , 12, 3129-3135	20
1830	Electrons and positrons in metal vacancies. <b>1975</b> , 12, 4012-4022	211
1829	Self-consistent pseudopotential calculations for Si (111) surfaces: Unreconstructed (1 $\times$ 1) and reconstructed (2 $\times$ 1) model structures. <b>1975</b> , 12, 4200-4214	423
1828	Tip region of the bremsstrahlung spectrum from incident electrons of kinetic energy 50 keV. <b>1975</b> , 11, 1797-1803	59
1827	Influence of the electron density profile on surface plasmons in a hydrodynamic model. <b>1975</b> , 11, 2118-2121	103
1826	Linear response theory and molecular vibrations: The Thomas-Fermi model. <b>1975</b> , 63, 438-442	18
1825	Ground-state thermomechanical properties of some cubic elements in the local-density formalism. <b>1975</b> , 12, 1257-1261	362
1824	Self-Consistent Theory of the Chemisorption of H, Li, and O on a Metal Surface. <b>1975</b> , 34, 531-534	209

1823	Self-consistent theory of the martensitic phase transformation in metallic lithium. <b>1975</b> , 12, 2194-2211	11
1822	Calculations of the energy loss of 4He ions in solid elements. <b>1975</b> , 46, 2817-2820	23
1821	de Haas-van Alphen effect and the band structure of URh3. <b>1975</b> , 12, 4102-4112	53
1820	Charge densities and interionic potentials in simple metals: Nonlinear effects. I. <b>1975</b> , 11, 2717-2725	167
1819	Spin-orbit coupling, Fermi surface, and optical conductivity of ferromagnetic iron. <b>1975</b> , 11, 287-294	128
1818	Exchange and correlation in the electron gas. <b>1975</b> , 11, 113-120	24
1817	Self-consistent field calculation of the electronic structure of the uranyl ion (UO <sub>2</sub> <sup>++</sup> ). <b>1975</b> , 63, 638-642	49
1816	Application of coherent potential approximation to Ni-rich Ni-Cu alloys. <b>1975</b> , 11, 2642-2650	10
1815	Electronic structure and optical properties of 3C-SiC. <b>1975</b> , 11, 1537-1546	48
1814	The electronic structure and Knight shift of EGa. <b>1975</b> , 5, 1667-1675	14
1813	RELATIVISTIC SELF-CONSISTENT-FIELD CALCULATIONS WITH APPLICATION TO ATOMIC HYPERFINE INTERACTION PART I: RELATIVISTIC SELF-CONSISTENT FIELDS PART II: RELATIVISTIC THEORY OF ATOMIC HYPERFINE INTERACTION. <b>1975</b> , 93-196	22
1812	Density Matrices. <b>1975</b> , 429-486	
1811	Self-consistent screening of impurity atoms in nearly-free-electron metals. <b>1975</b> , 31, 97-104	5
1810	Comparison of atomic Compton profiles obtained from four model local density functionals. <b>1975</b> , 8, 2593-2600	16
1809	New local self-consistency scheme for atoms. <b>1975</b> , 8, 2588-2592	2
1808	Hydrogen Chemisorption by the Spin-density Functional Formalism. I. <b>1975</b> , 11, 97-103	94
1807	Self-consistent electromagnetic modes at a metal slab. <b>1975</b> , 8, 1593-1613	
1806	Theoretical evaluation of the g shift in the alkali metals. <b>1975</b> , 5, 2300-2306	13

1805	Correlated potentials for simple metals: aluminium. <b>1975</b> , 5, 1155-1175	34
1804	Surface densities of states in the planar uniform background model. <b>1975</b> , 8, 3817-3824	9
1803	Energy band structure of beryllium and magnesium. <b>1975</b> , 5, 2089-2097	41
1802	Charge distribution in heterovalent alloys. <b>1975</b> , 5, 1502-1511	23
1801	On the role of the orthogonalization hole potential in Harrison's first principles pseudopotential theory. <b>1975</b> , 5, 1801-1816	23
1800	Influence of effective mass on pseudopotential theory. <b>1975</b> , 5, L187-L193	4
1799	Hartree-Fock Studies of Electronic Structures of Crystalline Solids. <b>1975</b> , 1, 147-218	36
1798	Calculations of low-energy electron diffraction intensities from the polar faces of ZnO. <b>1975</b> , 50, 605-614	77
1797	Determination of the structure of ordered adsorbed layers by analysis of LEED spectra. <b>1975</b> , 53, 501-522	255
1796	The structure of the p(2x2) tellurium overlayer on the (001) surface of copper investigated by leed. <b>1975</b> , 53, 523-537	29
1795	Theory of the electronic structure. <b>1975</b> , 28, 24-30	56
1794	The Van der Waals interaction between metals. <b>1975</b> , 5, 1475-1489	30
1793	Hohenberg-Kohn theorem for nonlocal external potentials. <b>1975</b> , 12, 2111-2120	589
1792	Transverse magnetic susceptibility in the local exchange approximation. <b>1975</b> , 5, 2119-2128	50
1791	Work functions of alkali metals. <b>1975</b> , 5, 1301-1306	10
1790	Ab Initio Calculation of the Spin Susceptibility for the Alkali Metals Using the Density-Functional Formalism. <b>1975</b> , 35, 1725-1728	59
1789	Self-consistent band theory of the Fermi-surface, optical, and photoemission properties of copper. <b>1975</b> , 11, 1522-1536	276
1788	The band structure of lithium with a screened non-local crystal potential. <b>1975</b> , 20, 261-274	19



1787	The use of discontinuous trial functions in the computation of the electronic structure of molecules: Calculation of the ESCA spectrum of tetrathiofulvalene (TTF). <b>1975</b> , 63, 697-701	9
1786	Removal of the discontinuities of the trial function in the computation of the electronic structure of molecules by the intersecting spheres model: Evaluation of the equilibrium geometry of N <sub>2</sub> and H <sub>2</sub> O. <b>1976</b> , 64, 1442-1445	8
1785	Quasispin model of itinerant magnetism. <b>1976</b> , 13, 3962-3971	25
1784	(110) surface states in III-V and II-VI zinc-blende semiconductors. <b>1976</b> , 13, 826-834	149
1783	Friedel oscillation in the spin density at the surface of an itinerant ferromagnet. <b>1976</b> , 1, 351-354	1
1782	Concentration-dependent absorption and spontaneous emission of heavily doped GaAs. <b>1976</b> , 47, 631-643	499
1781	A model for alkali chemisorption. <b>1976</b> , 59, 333-360	55
1780	Density-functional calculation of subband structure on semiconductor surfaces. <b>1976</b> , 58, 128-134	28
1779	A theory of thin metal films: electron density, potentials and work function. <b>1976</b> , 55, 427-444	331
1778	Local densities of states and bonding properties of 3d-transition metal clusters. <b>1976</b> , 61, 255-271	17
1777	Electronic structure of iron and models of the Earth's core. <b>1976</b> , 3, 45-48	24
1776	The effect of pressure on the physics and chemistry of potassium. <b>1976</b> , 3, 491-494	73
1775	The Electronic Structure of Alloys. <b>1976</b> , 31, 149-286	169
1774	Band model for magnetism of transition metals in the spin-density-functional formalism. <b>1976</b> , 6, 587-606	421
1773	The Fermi hole and the exchange parameter in X <sub>2</sub> theory. <b>1976</b> , 14, 1-10	96
1772	Chemisorption on Metals and Its Implications. <b>1976</b> , 32, 611-619	3
1771	Electron bremsstrahlung spectrum, 1-500 keV. <b>1976</b> , 13, 1714-1727	93
1770	Perturbation theory of structure in mixtures near phase separation. <b>1976</b> , 13, 859-871	47

1769	Chemisorption theory. <b>1976</b> , 6, 239-252	9
1768	Exchange and correlation energy of an inhomogeneous electron gas at metallic densities. <b>1976</b> , 13, 1477-1488	127
1767	Density-functional calculation of sub-band structure in accumulation and inversion layers. <b>1976</b> , 13, 3468-3477	230
1766	Study of sorption of oxygen on Al. <b>1976</b> , 14, 1446-1449	99
1765	Electronic structure of a metal-semiconductor interface. <b>1976</b> , 13, 2461-2469	387
1764	On the electronic structure of iron at core pressures. <b>1976</b> , 13, 57-66	22
1763	Hydrodynamic model for surface plasmons in metals and degenerate semiconductors. <b>1976</b> , 14, 1347-1361	97
1762	On the Shape of Electron-Hole Droplets in Highly Excited Anisotropic Semiconductors Application to Uniaxially Stressed Ge and Si. <b>1976</b> , 41, 91-99	1
1761	Exchange and correlation in atoms, molecules, and solids by the spin-density-functional formalism. <b>1976</b> , 13, 4274-4298	3137
1760	Theoretical studies of electron energy spectra in metals and metal alloys. <b>1976</b> , 19, 977-987	
1759	Neutral-atom electron binding energies from relaxed-orbital relativistic Hartree-Fock-Slater calculations. <b>1976</b> , 18, 243-291	440
1758	The interaction of rare gases with metals. <b>1976</b> , 18, 819-822	33
1757	Comments on chemisorption at a metal surface. <b>1976</b> , 20, 1073-1075	4
1756	Phenomenological representation of correlations in an electron gas. <b>1976</b> , 18, 265-267	1
1755	Non-local corrections to the electronic structure of metal surfaces. <b>1976</b> , 37, 99-104	39
1754	Variational-principle density-functional evaluation of the spin susceptibility for the alkali metals. <b>1976</b> , 25, 27-41	29
1753	The MTXB method: a semi-empirical version of the multiple scattering XBScheme with application to Li <sub>2</sub> . <b>1976</b> , 44, 321-324	13
1752	Gradient expansion of the atomic kinetic energy functional. <b>1976</b> , 43, 409-412	99

1751	The effect of the superconducting transition on the electronic work function of niobium. <b>1976</b> , 85, 277-282	2
1750	Analytically approximate screened calculations of atomic-field pair production cross sections. <b>1976</b> , 56, 27-28	5
1749	Self-consistent LCAO local density determination of anisotropic compton profile and X-ray structure factors in diamond. <b>1976</b> , 57, 457-459	9
1748	Exchange and correlation in atoms, molecules and solids. <b>1976</b> , 59, 177-179	70
1747	Exchange and correlation energy of an inhomogeneous electron gas. <b>1976</b> , 18, 549-552	19
1746	Ahoma <del>Б-tyhhehob-to-a</del> <del>Б-мак-сметлн.</del> "ohta" <del>Б-металл</del> <del>Б-обпн.</del> <b>1976</b> , 18, 1623-1630	5
1745	A limitation on the pseudopotential method. <b>1976</b> , 20, 151-153	8
1744	Metal interfaces: Adhesive energies and electronic barriers. <b>1976</b> , 20, 393-396	21
1743	Surface energy of electron-hole liquid in germanium at zero and finite<111> uniaxial stress. <b>1976</b> , 19, 935-938	13
1742	Calculation of ground state surface properties of simple metals including non-local effects of the exchange and correlation energy. <b>1976</b> , 19, 619-622	32
1741	First principles calculations of the volume dependence of the spin susceptibility for Li and Na. <b>1976</b> , 18, 85-91	17
1740	Role of the potential in LEED; application to the Na(110) surface. <b>1976</b> , 9, 3193-3204	21
1739	Pressure cell-boundary relation, Fermi levels and surface dipoles for transition metals. <b>1976</b> , 6, 573-585	106
1738	A spherical cell model of cerium. <b>1976</b> , 6, 797-805	5
1737	Helium heat of solution in Al and Mg using non-linear self-consistent screening of the nucleus. <b>1976</b> , 6, 1259-1268	25
1736	Green function theory of an electron gas with surface. II. The Hartree energy functional. <b>1976</b> , 9, 809-817	4
1735	Mode Gruneisen parameters in potassium. II. <b>1976</b> , 6, 1915-1922	19
1734	Electronic structure of YMg cubic compound by the self-consistent SAPW method. <b>1976</b> , 6, 789-796	18

1733	Phonon frequencies of lithium from a local effective potential. <b>1976</b> , 6, 1421-1431	14
1732	Self-consistency in calculations of the electronic band structure and the lattice dynamics of metals. <b>1976</b> , 6, 1503-1512	9
1731	Formation energy of heterovalent alloys. <b>1976</b> , 6, L173-L176	4
1730	Effect of correlation on the bandstructure of zirconium. <b>1976</b> , 6, 1639-1645	18
1729	The spherical-solid model: An application to x-ray edges in Li, Na, and Al. <b>1976</b> , 13, 3307-3319	174
1728	Effect of the atomic core on the fine-structure splitting for excited np states of the alkali-metal atoms. <b>1976</b> , 14, 1595-1602	25
1727	Study of the density gradient expansion in the surface energy calculation for metals. <b>1976</b> , 13, 5330-5337	41
1726	Condensed Phase of Lithium Impurities in Silicon. <b>1976</b> , 37, 354-357	14
1725	Chemical Trends in Atomic Adsorption on Simple Metals. <b>1976</b> , 37, 212-215	152
1724	Binding Energies for Different Adsorption Sites of Hydrogen on Simple Metals. <b>1976</b> , 37, 292-295	111
1723	Calculation of $g$ and $g^?$ for iron and nickel. <b>1976</b> , 14, 1214-1220	42
1722	Volume dependence of the Pauli susceptibility and the amplitude of the wave functions for Li and Na. <b>1976</b> , 13, 5136-5153	54
1721	Screening of a proton in an electron gas. <b>1976</b> , 14, 2250-2254	251
1720	Effect of ground-state and excitation potentials on energy levels of Ni metal. <b>1976</b> , 13, 1463-1467	34
1719	The electronic structure of solid surfaces. <b>1976</b> , 48, 479-496	215
1718	Surface of a spin-polarized electron gas. <b>1976</b> , 14, 2017-2031	33
1717	Cluster model for lattice distortion effects on electronic structure: VO and VO <sub>2</sub> . <b>1976</b> , 13, 3405-3418	16
1716	Prediction of the Fermi surface as a test of density-functional approximations to the self - energy. <b>1976</b> , 14, 4399-4406	16

1715	Theory of the heat of solution of hydrogen in Al and Mg using nonlinear screening. <b>1976</b> , 13, 590-602	133
1714	Equation for the direct determination of the density matrix. <b>1976</b> , 14, 41-50	216
1713	Relativistic electronic structure of the NaCl polymorph of CdS. <b>1976</b> , 13, 1675-1680	13
1712	Experimental and theoretical studies of extended x-ray absorption fine structure in bromine salts (NaBr and KBr). <b>1976</b> , 14, 741-746	12
1711	Local exchange approximations for ab initio construction of positive ion pseudopotentials: Alkali Rydberg states. <b>1976</b> , 65, 4321-4322	7
1710	An asymptotically exact form of the intersecting spheres model of molecules: Formulation. <b>1976</b> , 65, 253-256	11
1709	Self-consistent screening of a proton in jellium. <b>1977</b> , 7, 1763-1772	143
1708	Electron density of states of interstitial impurities in transition metals: H in Pd. <b>1977</b> , 7, 799-820	16
1707	The effect of microvoid size on positron annihilation characteristics and residual resistivity in metals. <b>1977</b> , 35, 973-981	166
1706	A KKRZ calculation on indium. I. An ab initio approach. <b>1977</b> , 7, 2067-2075	6
1705	Exchange and correlation potential in silicon. <b>1977</b> , 10, 987-999	25
1704	The energy of interstitial impurities in transition metals: H in Pd. <b>1977</b> , 7, 789-798	22
1703	On the energetics of the $f$ -phase transition in cerium metal: Determination of the atomic excitation $f_{d^2}$ to $d_{s^2}$ . <b>1977</b> , 7, 877-883	24
1702	Comparison of statistical and Lindgren conduction-core exchange approximations in a priori pseudopotential calculations for the lattice dynamics of metals. <b>1977</b> , 7, L169-L174	10
1701	The resonant model potential. II. Total energy: Theory and application to copper, silver, gold and calcium. <b>1977</b> , 7, 1167-1191	68
1700	An electron statistical vacancy model for the noble metals. <b>1977</b> , 7, 1383-1402	6
1699	Effect of correlation on the bandstructures of Th and Os. <b>1977</b> , 7, 2307-2322	19
1698	Hartree-Fock-Slater procedure for the calculation of optimised model potential parameters for aluminium. <b>1977</b> , 7, 1207-1217	3

1697	Self-consistent energy bands and Fermi surface of YZn. <b>1977</b> , 7, 1229-1244	19
1696	Charge and momentum density, and Compton profile of Se. <b>1977</b> , 10, 1875-1886	12
1695	The exchange and correlation energy of a spin-polarised electron gas. <b>1977</b> , 10, 3167-3178	6
1694	An investigation of the influence of the potential in LEED structural determination. <b>1977</b> , 10, 625-634	14
1693	Some muffin-tin calculations of electrical resistivity in molten palladium. <b>1977</b> , 35, 1609-1616	11
1692	Recursion method for electronic structure calculations in aperiodic materials. <b>1977</b> , 16, 5506-5511	10
1691	Energy bands and bonding in LaB <sub>6</sub> and YB <sub>6</sub> . <b>1977</b> , 15, 1859-1866	62
1690	Thomas-Fermi dielectric screening in semiconductors. <b>1977</b> , 16, 2717-2722	231
1689	Reply to "Further comments on atomic central-potential models". <b>1977</b> , 15, 1322-1325	6
1688	Muffin-tin orbitals and the total energy of atomic clusters. <b>1977</b> , 15, 3027-3038	131
1687	Hyperfine field of positive muons in Ni. <b>1977</b> , 15, 1560-1567	39
1686	Density-functional formulation of the generalized pseudopotential theory. <b>1977</b> , 16, 2537-2555	81
1685	Theoretical estimate of the exchange-correlation enhancement of the spin susceptibility of scandium. <b>1977</b> , 16, 777-784	17
1684	Energy bands in ferromagnetic iron. <b>1977</b> , 16, 2095-2105	368
1683	Convergence of the gradient expansion in metallic surface-energy calculations. <b>1977</b> , 15, 580-583	19
1682	Failure of the local exchange approximation in the evaluation of the H <sub>2</sub> ground state. <b>1977</b> , 15, 2858-2861	114
1681	Core holes in chemisorbed atoms. <b>1977</b> , 16, 2408-2419	248
1680	Ground-state electronic properties of diamond in the local-density formalism. <b>1977</b> , 15, 5049-5065	116

1679	Supposed Failure of the Boltzmann Equation in Nb. <b>1977</b> , 39, 225-228	22
1678	Renormalized atoms: Cohesion in transition metals. <b>1977</b> , 15, 1613-1628	178
1677	Density of states and spin susceptibility of a finite metal: Normal and giant surface effects. <b>1977</b> , 16, 1525-1535	11
1676	Orbital electron capture by the nucleus. <b>1977</b> , 49, 77-221	338
1675	Self-consistent relativistic APW calculation of the electronic structure of niobium with a non-muffin-tin potential. <b>1977</b> , 15, 3620-3632	82
1674	The virial theorem and the surface energy of an electron gas. <b>1977</b> , 15, 5302-5306	18
1673	Many-body effects in n-type Si inversion layers. II. Excitations to higher subbands. <b>1977</b> , 15, 3947-3958	98
1672	Electronic band structure of fluorite. <b>1977</b> , 16, 925-933	34
1671	Metal surface energies in the infinite and step-potential approximations. <b>1977</b> , 15, 1929-1935	26
1670	Effect of nonlocal contributions to the energy on properties of the electron-hole droplet in germanium and silicon. <b>1977</b> , 15, 4804-4816	3
1669	Effect of nonlocality on the Fermi surface of copper. <b>1977</b> , 15, 3714-3719	17
1668	Corrected spherical model for vacancies in simple metals. <b>1977</b> , 16, 4335-4342	19
1667	Self-consistent numerical-basis-set linear-combination-of-atomic-orbitals model for the study of solids in the local density formalism. <b>1977</b> , 15, 4716-4737	133
1666	Ground- and excited-state properties of LiF in the local-density formalism. <b>1977</b> , 16, 2901-2926	185
1665	Properties of the electron-hole liquid beyond the local density approximation. <b>1977</b> , 15, 979-988	10
1664	Study of the diamond (100) surface using the self-consistent-field extended tight-binding method. <b>1977</b> , 15, 3254-3259	24
1663	Nonresonant and resonant electron distributions from fast particle collisions: Application to sodium. <b>1977</b> , 16, 2232-2247	27
1662	Metal surface properties in the linear potential approximation. <b>1977</b> , 15, 1941-1949	57

1661	Low-energy-electron-diffraction analysis of the atomic geometry of ZnO (10 $\bar{1}$ 0). <b>1977</b> , 15, 4865-4873	65
1660	Investigation of the spin-density functional method for calculating spin-magnetic-moment densities. <b>1977</b> , 15, 1839-1846	21
1659	Electron gas exchange for atoms. <b>1977</b> , 16, 1358-1362	38
1658	Quasispin model of itinerant magnetism: High-temperature theory. <b>1977</b> , 15, 4281-4287	37
1657	Analytical structure of the wave-number-dependent susceptibility of many-fermion systems at low temperature and long wavelength. <b>1977</b> , 15, 1523-1532	12
1656	Self-consistent numerical-basis-set linear-combination-of-atomic-orbitals investigation of the electronic structure and properties of TiS <sub>2</sub> . <b>1977</b> , 16, 906-924	166
1655	Effect of self-consistency and exchange on the electronic structure of the transition metals, V, Nb, and Ta. <b>1977</b> , 15, 3685-3693	97
1654	Gradient expansion in the density-functional approach to an inhomogeneous electron system. II. <b>1977</b> , 16, 601-604	9
1653	Auger and radiative deexcitation of the 1s2l3l' configurations of lithium-like neon. <b>1977</b> , 15, 2318-2323	17
1652	Corrections to the Local Density Approximation: Gradient Expansion versus Wave-Vector Analysis for the Metallic Surface Problem. <b>1977</b> , 38, 1030-1033	80
1651	Calculation of the bandstructure of aluminium using the model potential method. <b>1977</b> , 7, 1467-1475	12
1650	XVI. Representations of the Electron Density and its Topographical Features. <b>1977</b> , 16, 187-197	51
1649	A Pseudopotential Approach to the Crystal Energy of Si. <b>1977</b> , 43, 1879-1883	9
1648	Band structure and x-ray photoelectron spectrum of ZrB <sub>2</sub> . <b>1977</b> , 16, 726-730	64
1647	Exchange-correlation energy of a metallic surface: Wave-vector analysis. <b>1977</b> , 15, 2884-2901	826
1646	Energy bands in ferromagnetic nickel. <b>1977</b> , 15, 298-306	395
1645	Theoretical Determination and Analysis of Electronic Charge Distributions. <b>1977</b> , 15, 147-162	95
1644	A theoretical equation of state for the inner core. <b>1977</b> , 14, 333-344	42



1643	Spin susceptibilities and the theory of itinerant-electron antiferromagnetism. <b>1977</b> , 26, 393-485	35
1642	Gradient corrections to the exchange-correlation energy of electrons at metal surfaces. <b>1977</b> , 15, 1801-1810	58
1641	Study of the density gradient expansion for the kinetic energy. <b>1977</b> , 16, 4249-4255	60
1640	Self-Consistent Calculation of Work Function, Charge Densities, and Local Densities of States for Cu(100). <b>1977</b> , 38, 561-564	104
1639	Atomic chemisorption on simple metals: Chemical trends and core-hole relaxation effects. <b>1977</b> , 68, 138-148	11
1638	Theoretical studies of atomic adsorption on nearly-free-electron-metal surfaces. <b>1977</b> , 68, 158-166	64
1637	Calculation of geometries and chemisorption energies of adatoms on simple metals. <b>1977</b> , 63, 348-357	51
1636	Electronic spectrum of ordered monolayers over metal surfaces. <b>1977</b> , 66, 238-252	1
1635	Low energy electron diffraction from Na(110) and Na <sub>2</sub> O(111) surfaces. <b>1977</b> , 65, 539-551	84
1634	I. Basic Concepts of Quantum Chemistry for Electron Density Studies. <b>1977</b> , 16, 87-102	29
1633	Statistical atomic models with piecewise exponentially decaying electron densities. <b>1977</b> , 16, 891-902	148
1632	Relation between the total energy and eigenvalue sum for neutral atoms and molecules. <b>1977</b> , 67, 4618-4619	32
1631	Energy bandstructure and Fermi surface of LaB <sub>6</sub> by a self-consistent APW method. <b>1977</b> , 7, 1245-1260	160
1630	A self-consistent theory for exchange and correlation effects in an electron gas. <b>1977</b> , 10, 3293-3303	10
1629	A physicist's view of the energetics of transition metals. <b>1977</b> , 1, 305-324	102
1628	Uniform susceptibilities of metallic elements. <b>1977</b> , 16, 255-262	704
1627	Local density theory of metallic cohesion. <b>1977</b> , 15, 2854-2857	216
1626	Quantum theory of adsorption of isolated adatoms. <b>1977</b> , 8, 103-122	12

1625	Application of the intersecting spheres model to periodic structures: Computation of the energy bands of diamond. <b>1977</b> , 22, 783-787	6
1624	Positron annihilation in small metal voids. <b>1977</b> , 21, 293-296	30
1623	Electronic structure of H and He in metal vacancies. <b>1977</b> , 24, 691-693	54
1622	Local and nonlocal contributions to adhesive energies at metallic interfaces. <b>1977</b> , 23, 527-529	7
1621	Screening of light impurities in simple metals. <b>1977</b> , 23, 795-798	58
1620	An oxygen pseudopotential: Application to the electronic structure of ZnO. <b>1977</b> , 22, 351-354	97
1619	Inter-subband optical transitions in a surface space-charge layer. <b>1977</b> , 21, 133-136	94
1618	Charge transfer and stability of NFE random alloys. <b>1977</b> , 38, 1307-1313	
1617	Charge transfer in binary alloys and to impurities in iron. <b>1977</b> , 38, 869-876	22
1616	The Stoner model in the spin-density-functional formalism. <b>1977</b> , 91, 329-336	34
1615	The local exchange model of itinerant-electron ferromagnetism: Energy bands in nickel and iron. <b>1977</b> , 91, 337-343	12
1614	Investigation of the interaction between molecules at medium distances. <b>1977</b> , 19, 209-216	38
1613	Theoretical L-shell Coster-Kronig energies 11 $\geq$ 103. <b>1977</b> , 19, 97-151	124
1612	Inter-subband optical absorption in space-charge layers on semiconductor surfaces. <b>1977</b> , 26, 263-272	167
1611	The exchange energy of a nonuniform electron gas. <b>1977</b> , 20, 770-773	
1610	The use of pseudopotentials within local-density formalism calculations for atoms: Some results for the first row. <b>1977</b> , 49, 367-373	33
1609	Non-muffin-tin MS XE total energies for H <sub>2</sub> , C <sub>2</sub> , N <sub>2</sub> and CO. <b>1977</b> , 45, 150-154	31
1608	Density functional approach to physical adsorption of rare gases on metal surfaces. <b>1977</b> , 23, 623-627	38

1607	Sign reversals of charge on electron-hole drop. <b>1977</b> , 24, 171-174	9
1606	Exchange and correlation in inhomogeneous electron systems. <b>1977</b> , 24, 765-768	130
1605	Semi-statistical model for metals. <b>1977</b> , 38, 307-310	11
1604	Coulomb energies and nuclear radii in the energy density formalism. <b>1977</b> , 283, 67-77	10
1603	Energies and fermi level of electrons in thin size-quantized metal films. <b>1977</b> , 79, 149-153	28
1602	Contribution of the d-Electron Screening to the Properties of Impurities in Transition Metals. <b>1977</b> , 81, 681-690	7
1601	The effective screening function in insulating solids. <b>1977</b> , 83, 543-553	3
1600	Calculation of a resonant model potential for copper, silver, and gold. <b>1977</b> , 84, 311-324	29
1599	Resistivity and thermoelectric power of liquid lithium and sodium from Harrison's first principle. <b>1977</b> , 84, 797-801	5
1598	Electronic-structure studies of solides. V. Rigorous Hartree-Fock treatment of metallic hydrogen using a plane-wave basis. <b>1977</b> , 12, 145-160	9
1597	Experimental and theoretical studies of optical properties on alloys of the intermetallic systems $\text{Li}_2\text{Ag}_2\text{In}_x$ and $\text{Li}_2\text{Cd}_2\text{In}_x$ . <b>1978</b> , 16, 255-269	15
1596	Binding-energy calculations for multiple bound excitons in silicon. <b>1978</b> , 85, K53-K56	8
1595	Binding-energy calculations for multiple bound excitons. <b>1978</b> , 86, 505-515	16
1594	Effects of Core and Correlation Corrections on the Electrical Resistivity of Lithium. <b>1978</b> , 87, 145-150	1
1593	The energy of interstitial hydrogen in Zirconium. <b>1978</b> , 50, 203-211	18
1592	On the first principles Hartree-Fock and local density pseudopotentials. <b>1978</b> , 30, 423-443	16
1591	Electronic contribution to thermodynamic properties of metals. <b>1978</b> , 93, 149-164	2
1590	Chemisorption on metals. <b>1978</b> , 9, 1-43	360

1589	Electronic structure of vacancies and interstitials in metals. <b>1978</b> , 69-70, 157-175	17
1588	Energetics of vacancies in non-transition metals. <b>1978</b> , 69-70, 633-635	9
1587	Attachment of an electron-hole droplet to a donor. <b>1978</b> , 27, 331-333	8
1586	Contraction of diatomic molecules upon chemisorption. <b>1978</b> , 28, 899-902	12
1585	Self-consistent calculation of impurity scattering in inversion layers. <b>1978</b> , 28, 861-863	23
1584	The use of OPW's in the Lindgren approximation of conduction-core exchange and the lattice dynamical properties of rubidium. <b>1978</b> , 27, 835-837	9
1583	Impurity interactions and pseudo-molecule formation in metals. <b>1978</b> , 25, 995-998	19
1582	Non-spherical contributions to the vacancy formation energy. <b>1978</b> , 27, 33-35	7
1581	Variational calculation of the electronic density at surfaces. <b>1978</b> , 27, 1193-1197	6
1580	On the feasibility of using positive muons as probes of defect structure. <b>1978</b> , 27, 1249-1253	7
1579	A self-consistent energy band study of FeO, CoO and NiO. <b>1978</b> , 25, 9-12	35
1578	Hyperfine fields at positive muons in metals. <b>1978</b> , 4, 282-300	17
1577	Muonic and hydrogenic knight shifts in free electron paramagnets. <b>1978</b> , 4, 301-306	22
1576	The extended rigorous cellular method: Theory and applications to thin films. <b>1978</b> , 29, 353-360	6
1575	Surface plasmon dispersion relation for spherical metal particles. <b>1978</b> , 11, 135-144	9
1574	Some resonance model calculations of electrical resistivity in molten titanium. <b>1978</b> , 8, 1703-1712	4
1573	Band structure of US and UN by SAPW. <b>1978</b> , 28, 177-182	4
1572	Theoretical models of the electronic structure at surface irregularities. <b>1978</b> , 13, 541-556	1

1571	Self-consistent field electronic structure calculations for compressed magnesium oxide. <b>1978</b> , 39, 255-257	11
1570	On the effect of the local exchange in calculations of the hyperfine interaction parameters $p(0)$ and $(r\bar{B})$ . <b>1978</b> , 57, 458-462	4
1569	Itinerant ferromagnetism in fcc cobalt. <b>1978</b> , 25, 53-55	166
1568	Electron states near surfaces of solids. <b>1978</b> , 113, 184-218	7
1567	BNDPKG. A package of programs for the calculation of electronic energy bands by the LCGO method. <b>1978</b> , 14, 327-365	100
1566	On the calculation of correlation energies in the spin-density functional formalism. <b>1978</b> , 49, 143-149	229
1565	The electronic structure of impurities and other point defects in semiconductors. <b>1978</b> , 50, 797-858	587
1564	Density-functional theory of Wigner crystallization. <b>1978</b> , 18, 6506-6509	42
1563	Individual orbital contributions to the SCF virial in homonuclear diatomic molecules. <b>1978</b> , 69, 2930	39
1562	Local-density self-consistent energy-band structure of cubic CdS. <b>1978</b> , 17, 4850-4863	66
1561	Self-consistent augmented-plane-wave electronic-structure calculations for the A15 compounds V3X and Nb3X, X=Al,Ga,Si,Ge,andSn. <b>1978</b> , 18, 6411-6438	204
1560	Static semiclassical response of a bounded electron gas. II. The finite barrier model. <b>1978</b> , 68, 5010-5019	10
1559	Core-Level Binding-Energy Shifts in Metals. <b>1978</b> , 40, 954-957	294
1558	Proof that $\bar{E}_{ii} = \bar{E}_{in}$ in density-functional theory. <b>1978</b> , 18, 7165-7168	1542
1557	Inhomogeneous relativistic electron gas. <b>1978</b> , 11, L943-L948	242
1556	Hyperfine field systematics of nonmagnetic ions in ferromagnetic alloys. <b>1978</b> , 8, 99-111	27
1555	Calculation of low-energy-electron-diffraction intensities from ZnO(101 $\bar{1}$ 0). II. Influence of calculational procedure, model potential, and second-layer structural distortions. <b>1978</b> , 18, 4225-4240	133
1554	A review of the theoretical and experimental analyses of electron spin polarization in ferromagnetic transition metals. <b>1978</b> , 75, 401-489	27

1553	A review of the theoretical and experimental analyses of electron spin polarization in ferromagnetic transition metals. <b>1978</b> , 75, 490-528	11
1552	Crystallography by LEED. <b>1978</b> , 78, 339-370	31
1551	Theory of the Thermodynamics of Simple Liquid Metals. <b>1978</b> , 33, 1-81	199
1550	Electronic structure of vacancies and vacancy clusters in simple metals. <b>1978</b> , 8, 2243-2260	43
1549	A new approximation method for electron systems. <b>1978</b> , 11, 4945-4956	74
1548	Compton profiles of Ne, Ar, and Kr. <b>1978</b> , 18, 552-558	20
1547	Energy bands, Compton profile, and optical conductivity of vanadium. <b>1978</b> , 17, 455-461	80
1546	First-principles nonlocal-pseudopotential approach in the density-functional formalism: Development and application to atoms. <b>1978</b> , 18, 5449-5472	178
1545	A simple, useful analytical form of the static electron gas dielectric function. <b>1978</b> , 8, 1699-1702	233
1544	Electronic structure of the noble gas dimer ions. I. Potential energy curves and spectroscopic constants. <b>1978</b> , 69, 5151-5162	132
1543	Density-Functional Approach to the Metal-Insulator Transition in Doped Semiconductors. <b>1978</b> , 41, 1569-1572	43
1542	Theory of atomic chemisorption on simple metals. <b>1978</b> , 18, 616-636	529
1541	Surfaces of real metals by the variational self-consistent method. <b>1978</b> , 17, 2595-2611	196
1540	Statistical correlation energies in atomic and molecular systems. <b>1978</b> , 11, 2589-2600	6
1539	Linear response theory of surface electronic structure. <b>1978</b> , 11, 1865-1875	9
1538	The Compton profile as a criterion for the choice of the local exchange parameter $\Xi$ . <b>1978</b> , 11, 385-389	9
1537	First ionisation potentials of atoms obtained with local-density schemes. <b>1978</b> , 11, 1339-1351	50
1536	Spin density functional theory for the spin wave stiffness coefficient in ferromagnets. <b>1978</b> , 8, 1539-1556	16

1535	Bloch wave effects in the density response function. <b>1978</b> , 11, 1813-1824	5
1534	Large-q Form Factors for Light Atoms. <b>1978</b> , 17, 547-548	10
1533	Dynamic longitudinal spin susceptibility of Bloch electrons. <b>1978</b> , 8, 2569-2577	19
1532	The transverse dynamical susceptibility of ferromagnets and antiferromagnets within the local exchange approximation. <b>1978</b> , 8, 1501-1512	25
1531	Subband Structure and Inter-Subband Absorption in an Accumulation Layer in Strong Magnetic Fields. <b>1978</b> , 44, 475-481	42
1530	Electronic structure of ScRu, ScRh, ScPd, and ScAg. <b>1978</b> , 8, 2301-2311	37
1529	Relationship between the electronic potential energy of a crystal and X-ray scattering. <b>1978</b> , 38, 289-293	5
1528	Polarization Propagator Calculations. <b>1978</b> , 11, 275-352	328
1527	Theory of energy bands and related properties of 4d transition metals. III. s and d contributions to the equation of state. <b>1978</b> , 8, 219-230	151
1526	Structurally Induced Semimetal-to-Semiconductor Transition in 1T-TiSe <sub>2</sub> . <b>1978</b> , 40, 1155-1158	29
1525	Charge Density and Structural Properties of Covalent Semiconductors. <b>1978</b> , 40, 950-953	111
1524	Exchange Splitting in Nickel. <b>1978</b> , 40, 892-896	76
1523	Linear-response functions in spin-density-functional theory. <b>1978</b> , 17, 2980-2988	39
1522	Ab initio self-consistent study of the electronic structure and properties of cubic boron nitride. <b>1978</b> , 17, 2030-2042	83
1521	Electronic structure and optical transition energies of the U center in alkaline-earth fluorides. <b>1978</b> , 18, 1977-1985	2
1520	Self-consistent screening of a positive muon in a spin-polarized electron gas. <b>1978</b> , 17, 301-307	38
1519	Study of a neutrally charged electron gas with a surface. <b>1978</b> , 18, 3931-3945	39
1518	Self-consistent electronic structure of transition-metal surfaces: The Mo (001) surface. <b>1978</b> , 18, 5473-5483	54

1517	Variational approach to inhomogeneous electron liquids: Application to metallic hydrogen. <b>1978</b> , 18, 4048-4063	10
1516	Surface structure of electron-hole drops in germanium and silicon. <b>1978</b> , 17, 2655-2672	45
1515	Hydrogen Chemisorption by the Spin-Density Functional Formalism. II. Rle of thesp-conduction Electrons of Metal Surfaces. <b>1978</b> , 18, 481-493	83
1514	Electronic structure and proton spin-lattice relaxation in PdH. <b>1978</b> , 17, 3029-3039	107
1513	Nonlinear electron-density distribution around point defects in simple metals. II. Applications. <b>1978</b> , 18, 2723-2732	17
1512	Density functional theory of 3d-transition element atoms. <b>1978</b> , 68, 3316-3317	68
1511	Atoms in jellium. <b>1978</b> , 17, 1744-1757	84
1510	Exact solution of metal surface properties in square barrier and linear one-electron potential models. <b>1978</b> , 18, 2674-2682	9
1509	Electron diffraction from gaseous tellurium hexafluoride at 20, 90, and 150 °C. Molecular structure and three-atom scattering. <b>1978</b> , 68, 3548-3552	22
1508	Band structure of thin films by the linear augmented-plane-wave method. <b>1978</b> , 18, 605-615	106
1507	Electronic properties of a simple metal-metal interface. <b>1978</b> , 17, 3904-3918	49
1506	Exchange instabilities in an n-type silicon inversion layer. <b>1978</b> , 17, 1383-1387	4
1505	Anharmonic properties of Li. <b>1978</b> , 18, 2643-2655	22
1504	First-principles theoretical study on the electronic properties of the B32 intermetallic compound LiAl. <b>1978</b> , 17, 2582-2594	68
1503	Approach to the embedding problem in chemisorption in a self-consistent-field-molecular-orbital formalism. <b>1978</b> , 17, 3143-3153	73
1502	Nonlocal exchange and correlation in an inhomogeneous electron system. <b>1978</b> , 17, 2429-2435	22
1501	Nonlinear electron-density distribution around point defects in simple metals. I. Formulation. <b>1978</b> , 18, 2712-2722	20
1500	Density functional theory and molecular bonding. II. Alkali dimers. <b>1978</b> , 68, 1190-1193	68



1499	Electric-Field Gradient at Cu Nuclei Due to an Interstitial Positive Muon. <b>1978</b> , 40, 264-266	27
1498	Pseudopotentials in Density-Functional Theory. <b>1978</b> , 41, 191-194	48
1497	Calculated Bulk Properties of the Actinide Metals. <b>1978</b> , 41, 42-45	245
1496	Multiplet structure and charge distributions in silicon and germanium dimers. <b>1978</b> , 18, 2159-2166	40
1495	Self-consistent calculation of surface properties of electron-hole droplets. <b>1978</b> , 17, 1884-1892	20
1494	Heat of solution of hydrogen in Al and Mg. <b>1978</b> , 17, 1592-1595	18
1493	Electronic structure of hydrogen in simple metals. <b>1978</b> , 17, 3518-3524	99
1492	Magnetic susceptibility of an electron gas in the random-phase approximation. <b>1978</b> , 17, 385-395	22
1491	Density response theory of nonbonded interactions. <b>1978</b> , 68, 1242-1247	6
1490	Nonlocal approximation to the exchange potential and kinetic energy of an inhomogeneous electron gas. <b>1978</b> , 17, 3735-3743	275
1489	Chapter 3 Electronic structure of rare earth metals. <b>1978</b> , 1, 233-335	2
1488	Theory of interface energies. <b>1978</b> , 18, 2583-2590	20
1487	Change-in-self-consistent-field theory of the work function. <b>1978</b> , 18, 656-666	103
1486	Band magnetism in the spin-density-functional formalism. <b>1978</b> , 49, 1399-1404	28
1485	Electron liquid in collective description. IV. Proton in an electron gas. <b>1979</b> , 12, 3013-3023	21
1484	Linear augmented plane wave method for self-consistent calculations. <b>1979</b> , 9, 661-672	98
1483	Application of fast self-consistent cluster calculations for large systems to the electronic structure of solids. <b>1979</b> , 9, 1589-1611	46
1482	Spherical solid model for muon and hydrogen in metals. <b>1979</b> , 9, 1333-1348	90

1481	Lattice effects in the ferromagnetic response functions. <b>1979</b> , 12, 3025-3032	
1480	Single-particle Hamiltonian and self-consistency in random binary alloys. <b>1979</b> , 9, 253-260	3
1479	Core charge polarisation effects in atomic systems. <b>1979</b> , 12, 1091-1102	7
1478	The energy density functional formalism for excited states. <b>1979</b> , 12, 5419-5430	284
1477	On some 'local' force densities and stress tensors in molecular quantum mechanics. <b>1979</b> , 12, 3857-3871	33
1476	Density-functional approximation for the quasiparticle dynamics of Rb. <b>1979</b> , 9, L99-L106	7
1475	On the origin of the Hume-Rothery rules for phase stability in $\alpha$ and $\beta$ brasses. <b>1979</b> , 9, 1939-1960	36
1474	On H (and He) impurities in Al and Mg. <b>1979</b> , 9, 629-635	3
1473	The influence of different exchange-correlation potentials on the energy band structure of ScP. <b>1979</b> , 12, 5441-5451	11
1472	Self-Consistent Calculation of Molecular Chemisorption on Metals. <b>1979</b> , 20, 192-201	61
1471	Thomas-Fermi approach to diatomic systems. II. Correlation diagrams for N-N and Ne-Ne. <b>1979</b> , 20, 1808-1815	23
1470	Electromagnetic potential in Thomas-Fermi-Dirac atoms. <b>1979</b> , 20, 44-47	10
1469	Electron structure of single and interacting hydrogen impurities in free-electron-like metals. <b>1979</b> , 20, 446-454	118
1468	Simple model of hydrogen and lithium chemisorption on jellium substrates. <b>1979</b> , 19, 1270-1282	30
1467	Theory of intersubband cyclotron combined resonances in the silicon space-charge layer. <b>1979</b> , 19, 2106-2116	68
1466	Theory of metallic adhesion. <b>1979</b> , 19, 3911-3920	54
1465	Intra-atomic correlation energies in cubic metals with canonical bands. <b>1979</b> , 20, 4637-4644	21
1464	Local density functional for kinetic energy. <b>1979</b> , 20, 3165-3171	7

1463	Simplified local-density theory of the cohesive energy of metals. <b>1979</b> , 19, 609-619	26
1462	Self-consistent mixed-basis approach to the electronic structure of solids. <b>1979</b> , 19, 1774-1782	408
1461	Correlation effects on the energy band of Ni. <b>1979</b> , 19, 1295-1298	31
1460	Spin-polarized band-structure calculations for Ni. <b>1979</b> , 20, 3172-3185	70
1459	Chemical potential and electronegativity in terms of electron density. <b>1979</b> , 71, 1004-1006	16
1458	Effect of electron correlations on photoemission from narrow-band metals. <b>1979</b> , 50, 1944-1949	86
1457	Comparison of theoretical calculations of angular distributions of photoelectrons emitted from rare-gas atoms. <b>1979</b> , 19, 734-740	5
1456	Statistical calculation of jellium surface properties. <b>1979</b> , 19, 1290-1294	47
1455	Liquid structure of the simple alkali metals from a first-principles pseudopotential calculation. <b>1979</b> , 19, 328-333	27
1454	Descriptions of exchange and correlation effects in inhomogeneous electron systems. <b>1979</b> , 20, 3136-3164	514
1453	Interaction-Induced Transition at Low Densities in Silicon Inversion Layer. <b>1979</b> , 43, 1529-1532	56
1452	Study of the exchange energy of an inhomogeneous electron gas at a surface. <b>1979</b> , 20, 2291-2302	17
1451	Theory of the deviation from the Ruedenberg energy relation for molecules. <b>1979</b> , 71, 1495-1497	18
1450	Dynamical calculation of low-energy electron diffraction intensities from GaAs(110): Influence of boundary conditions, exchange potential, lattice vibrations, and multilayer reconstructions. <b>1979</b> , 19, 5194-5205	134
1449	Two-site interaction model for a ferromagnetic metal. <b>1979</b> , 19, 4661-4676	
1448	Self-consistent embedded-cluster model for magnetic impurities: Fe, Co, and Ni in $\bar{2}$ -NiAl. <b>1979</b> , 20, 1198-1207	108
1447	Self-consistent field model for condensed matter. <b>1979</b> , 20, 4981-4989	309
1446	Density functionals in unrestricted Hartree-Fock theory. <b>1979</b> , 71, 490-496	65

1445	Heavy alkali atoms in jellium. <b>1979</b> , 19, 2801-2812	23
1444	Self-consistent calculation of the structural properties of silicon. <b>1979</b> , 20, 4251-4255	21
1443	Electronic energy bands in Eberium. <b>1979</b> , 19, 6274-6278	5
1442	Electron bremsstrahlung angular distributions in the 1-500 keV energy range. <b>1979</b> , 19, 187-195	65
1441	Electronic structure of thin films by the self-consistent numerical-basis-set linear combination of atomic orbitals method: Ni(001). <b>1979</b> , 19, 793-805	99
1440	Thomas-Fermi approach to diatomic systems. I. Solution of the Thomas-Fermi and Thomas-Fermi-Dirac-Weizsäcker equations. <b>1979</b> , 20, 1798-1807	54
1439	Theory of Auger relaxation energies in metals. <b>1979</b> , 20, 1369-1376	120
1438	Impurity Scattering on the Fermi Surfaces: A (More) Realistic Calculation. <b>1979</b> , 42, 1174-1178	7
1437	Valence and Rydberg excited states of H <sub>2</sub> S: An SCF-XESW molecular orbital study. <b>1979</b> , 70, 1177-1186	47
1436	Positrons in metals: A real-space approach. <b>1979</b> , 20, 21-32	11
1435	Theory of structural properties of covalent semiconductors. <b>1979</b> , 19, 5251-5264	169
1434	Dielectric-constant enhancement as the insulator-metal transition is approached from the insulating side—reply. <b>1979</b> , 20, 3508-3510	
1433	Rayleigh-Ritz variational calculations of real-metal-surface properties. <b>1979</b> , 19, 1840-1854	43
1432	Hydrodynamic model of linear response for a jellium surface: Nonretarded limit. <b>1979</b> , 20, 4872-4882	42
1431	Subband energies in n-channel inversion layers on (111) Ge. <b>1979</b> , 20, 2395-2397	11
1430	Theory of the hydrogen interstitial impurity in germanium. <b>1979</b> , 20, 5050-5058	35
1429	Gradient correction to the statistical electronic free energy at nonzero temperatures: Application to equation-of-state calculations. <b>1979</b> , 20, 586-594	155
1428	Molecular bonding in Group IIA dimers Be <sub>2</sub> Ba <sub>2</sub> . <b>1979</b> , 71, 1300-1308	150

1427	Density functional theory and molecular bonding. III. Iron-series dimers. <b>1979</b> , 70, 830	233
1426	Theoretical and experimental studies of band formation in CO adlayers. <b>1979</b> , 20, 801-808	45
1425	Simple and accurate statistical calculation of jellium-metal surface properties. <b>1979</b> , 20, 3511-3514	10
1424	Calculation of equilibrium geometries and ionization energies of sodium clusters up to Na <sub>8</sub> . <b>1979</b> , 71, 3042	103
1423	Local exchange potential for atomic systems. <b>1979</b> , 62, 31-33	15
1422	Numerical tests of a local exchange potential for atomic systems. <b>1979</b> , 67, 408-411	1
1421	Total energies of molecules with the local density functional approximation and gaussian basis sets. <b>1979</b> , 65, 206-211	29
1420	Zero-temperature equation of state of metals in the statistical model with density gradient correction. <b>1979</b> , 98, 555-565	17
1419	Role of gradient corrections in interpolation formulae for surface energies. <b>1979</b> , 73, 401-403	6
1418	A lower bound for Coulomb energies. <b>1979</b> , 70, 444-446	99
1417	A modification of the Thomas-Fermi method. <b>1979</b> , 72, 312-314	
1416	Application of the ab initio Hartree-Fock-Slater method to the calculation of molecular conformations: Ozone and thiozone. <b>1979</b> , 36, 323-325	19
1415	First-principles pseudopotential in the local-density-functional formalism. <b>1979</b> , 39, 75-90	12
1414	Effect of zero point motion on the hyperfine field at an interstitial positive muon site in ferromagnetic Ni. <b>1979</b> , 31, 1003-1007	15
1413	On the electronic specific heat of transition metals. <b>1979</b> , 30, 785-789	12
1412	Electronic structure of K-(001) surface by the density matrix method. <b>1979</b> , 30, 301-304	23
1411	The gradient approximation to the exchange-correlation energy functional: A generalization that works. <b>1979</b> , 31, 567-571	58
1410	Self-consistent electronic structure and ground state properties of TlCl. <b>1979</b> , 32, 591-594	3

1409	The small-core approximation and structural phase stability in metals. <b>1979</b> , 31, 881-884	3
1408	Ground state energy of small electron-hole drops. <b>1979</b> , 29, 389-393	8
1407	Adsorbate band formation: The chemisorption of CO on Pd (100). <b>1979</b> , 31, 257-260	54
1406	A model for light impurities in metals. <b>1979</b> , 32, 1297-1301	35
1405	A solid state approach to the electronic structure of molecules: Self-consistent pseudopotential calculation of O <sub>2</sub> . <b>1979</b> , 32, 309-312	12
1404	Multiband and multivalley effective-mass theory for impurities in semiconductors. <b>1979</b> , 30, 65-70	11
1403	Size effects in small electron-hole drops. <b>1979</b> , 30, 697-701	6
1402	An efficient alternative method of LEED analysis for structural determinations of surfaces. <b>1979</b> , 29, 405-416	13
1401	A simulation of the exchange potential in unrestricted and restricted Hartree-Fock calculations studied on atoms. <b>1979</b> , 51, 339-348	1
1400	Isomer shifts at rare-earth impurities in s-p metals. <b>1979</b> , 37, 241-256	5
1399	A study of the electronic structure of molecules by the self-consistent discrete variation X $\beta$ method in a basis of numerical hartree-fock functions. <b>1979</b> , 19, 843-851	
1398	Bound excitons in local density Approximation. <b>1979</b> , 91, 331-337	18
1397	Harrison's Method Electrical Transport Properties of Alkali Metals. <b>1979</b> , 93, 391-395	6
1396	Density functional approach to point defect properties of coppergermanium alloys. <b>1979</b> , 96, 225-231	8
1395	Spin-density functional method and the ground state of solids. IV. <b>1979</b> , 22, 600-605	
1394	Electron densities at muons in simple metals. <b>1979</b> , 6, 29-32	1
1393	Cohesive and superconducting properties of La-In compounds from electronic-structure calculations. <b>1979</b> , 35, 27-33	11
1392	Binding energy calculations for metal aggregates. <b>1979</b> , 33, 13-20	10

1391	Effects of the local electron speed on the exchange-correlation phenomena. <b>1979</b> , 25, 485-488	1
1390	On the electronic properties and electrical conductivity of thin metal layers. <b>1979</b> , 34, 17-27	
1389	The nature of the liquid-vapour interface and other topics in the statistical mechanics of non-uniform, classical fluids. <b>1979</b> , 28, 143-200	2173
1388	Momentum-space formalism for the total energy of solids. <b>1979</b> , 12, 4409-4422	1305
1387	Band Structure Calculations for Metal Hydrogen Systems*. <b>1979</b> , 117, 89-112	333
1386	Remarks on the density functional approach to the inhomogeneous electron gas. <b>1979</b> , 95, 547-560	2
1385	The canonical exchange energy modulation parameters in molecular MTX calculations. Some first row homonuclear diatomic molecules. <b>1979</b> , 65, 494-499	3
1384	Orbital functional for exchange and correlation: self-interaction correction to the local density approximation. <b>1979</b> , 64, 127-130	148
1383	Models of electronic structure of hydrogen in metals: Pd-H. <b>1979</b> , 20, 3543-3551	52
1382	On first-row diatomic molecules and local density models. <b>1979</b> , 71, 4993	560
1381	Work function of metals. <b>1979</b> , 1-150	156
1380	Phonon dispersion in transition metals. <b>1979</b> , 19, 6142-6154	214
1379	A relativistic density functional formalism. <b>1979</b> , 12, 2977-2990	475
1378	Cohesive properties of metallic compounds: Augmented-spherical-wave calculations. <b>1979</b> , 19, 6094-6118	1126
1377	Linearized augmented plane-wave method for the electronic band structure of thin films. <b>1979</b> , 19, 1706-1719	311
1376	Spin-density gradient expansion for the kinetic energy. <b>1979</b> , 20, 397-403	166
1375	Electronic properties, chemical bonding, and lattice dynamics of semiconductors. <b>1979</b> , 21-42	4
1374	The generalised pseudoatom potential in solids: relation to screening and lattice dynamics. <b>1979</b> , 12, 1491-1503	17

1373	The influence of the hole and exchange terms in the Dirac-Slater potential on the photoionisation cross sections and internal conversion coefficients. <b>1979</b> , 12, 3187-3200	3
1372	On some approximations in applications of X $\alpha$ theory. <b>1979</b> , 71, 3396-3402	1192
1371	Calculation of the electronic properties of Mo in a first-principles nonlocal-pseudopotential approach. <b>1979</b> , 20, 581-593	33
1370	Self-consistent pseudopotential calculation of the bulk properties of Mo and W. <b>1979</b> , 19, 568-582	68
1369	First-principles nonlocal-pseudopotential approach in the density-functional formalism. II. Application to electronic and structural properties of solids. <b>1979</b> , 20, 4082-4108	180
1368	Electronic Properties of a Semiconductor Superlattice. I. Self-Consistent Calculation of Subband Structure and Optical Spectra. <b>1979</b> , 47, 1518-1527	129
1367	Local-density theory of multiplet structure. <b>1979</b> , 20, 1693-1703	336
1366	Calculated energies and geometries for hydrogen impurities in Al and Mg. <b>1979</b> , 9, 1975-1982	63
1365	Optical conductivities of iron and nickel. <b>1979</b> , 20, 1134-1138	52
1364	Effective-mass approximation in the presence of an interface. <b>1979</b> , 20, 734-747	117
1363	Spin density functional theory of the temperature-dependent spin susceptibility: Pd and Pt. <b>1979</b> , 12, 43-57	36
1362	The polarization of electrons by tunneling through a spin-dependent surface potential. <b>1979</b> , 90, 102-108	10
1361	Hydrogen chemisorption on Al, Mg and Na surfaces [Calculation of adsorption sites and binding energies. <b>1979</b> , 81, 539-561	97
1360	Variational calculation of the electron density and potential in a thin metal film. <b>1979</b> , 84, 121-128	3
1359	États de surface du sodium dans l'approximation des électrons quasi-libres. <b>1979</b> , 79, 489-497	7
1358	Theoretical description of molecule-metal interaction and surface reactions. <b>1979</b> , 89, 196-225	146
1357	Electronic structure of magnetic 3d metals: Ground state, Fermi surface and photoemission properties. <b>1979</b> , 50, 7423	65
1356	Self-Consistent Pseudopotential Calculation of the Band Structure and the Crystal Energy of As. <b>1979</b> , 47, 547-550	13



1355	The Energy and Electron Density of States of Hydrogen Impurities in Metals. <b>1979</b> , 115, 239-246	1
1354	Clustering in the approach to the metal-insulator transition. <b>1980</b> , 42, 859-872	53
1353	Self-consistency condition for calculations based on the Kohn-Sham formalism. <b>1980</b> , 79, 437-438	5
1352	Energy band theory and the lattice dynamics of rare gas crystals. <b>1980</b> , 38, 393-411	6
1351	Hohenberg-Kohn theorem. <b>1980</b> , 18, 1029-1035	21
1350	Multiple scattering mass operator method for molecular orbital calculations. <b>1980</b> , 18, 1165-1173	1
1349	Fermi Energy, Density of States, and Electronic Properties of Alkali Metals Exchange and Correlation Effects. <b>1980</b> , 100, 103-109	4
1348	A Density-Matrix Functional Approach to Bound Multiexciton Complexes. <b>1980</b> , 100, 191-200	2
1347	Electronic Energy Band Structure of Osmium Metal. <b>1980</b> , 100, K155-K158	5
1346	Dynamic Spin Susceptibility of an Interacting Electron Gas. <b>1980</b> , 101, 311-320	1
1345	Equation d'état et structure de bandes d'énergie de l'argent métallique. <b>1980</b> , 101, 741-747	5
1344	Charge and Spin Density Waves in Jellium. <b>1980</b> , 102, 283-293	22
1343	Electron Density Distribution at Point Defects. Application of the Quadratic Response Theory. <b>1980</b> , 102, 521-531	2
1342	Thomas-Fermi kinetic-energy density with gradient corrections. <b>1980</b> , 343, 91-108	10
1341	Critical temperature of superconducting metallic hydrogen. <b>1980</b> , 80, 193-194	3
1340	Some aspects of the electronic structure of uranium pnictides and chalcogenides. <b>1980</b> , 102, 51-58	31
1339	Cohesive properties of uranium dioxide. <b>1980</b> , 102, 81-83	8
1338	Band structure, Compton profile and the $f$ -transition in cerium. <b>1980</b> , 102, 348-352	7

1337	APW-XE Compton profiles in crystalline rare gases. <b>1980</b> , 41, 623-630	5
1336	An application of the exchange-correlation mass operator approximation to the study of s-doublet structure of some 3d-transition metal ions. <b>1980</b> , 41, 953-958	3
1335	Density-functional approach to charge-transfer insulators. <b>1980</b> , 33, 463-466	5
1334	A simple way of eliminating the exchange parameter in the multiple scattering method <b>1980</b> , 35, 329-331	1
1333	A simplified treatment of exchange and correlation in semiconducting surface inversion layers. <b>1980</b> , 34, 423-426	8
1332	A comparison of electronic properties of various modifications of graphite. <b>1980</b> , 33, 817-820	21
1331	A self-interaction corrected approach to many-electron systems: Beyond the local spin density approximation. <b>1980</b> , 34, 933-936	140
1330	Temperature dependence of impurity resistivity due to hydrogen in metals. <b>1980</b> , 34, 179-182	12
1329	Transition metal ions in crystals: A refined treatment and deduction of coulomb and exchange interaction constants. <b>1980</b> , 33, 381-384	6
1328	Self-consistent surface calculation of electron-hole drops in gallium phosphide. <b>1980</b> , 34, 121-124	1
1327	Allowance for kinematic selection rules in the electron-density functional method and ground-state quasiparticles <b>1980</b> , 23, 980-983	
1326	Lattice static properties of vacancy clusters and interstitials in hcp magnesium: Computer simulation studies. <b>1980</b> , 15, 189-205	6
1325	Dense phases of hydrogen. <b>1980</b> , 531-536	
1324	On the fermi surfaces of random substitutional alloys <b>1980</b> , 366-387	
1323	Light impurities in a uniform electron gas. <b>1980</b> , 13, 4137-4156	12
1322	Quantenmechanische Modelle zur Wechselwirkung einfacher Adsorbate mit Festkörpersoberflächen. <b>1980</b> , 36, 227-231	6
1321	Compton profiles for neon and argon from X-ray wavefunctions. <b>1980</b> , 13, 3075-3080	8
1320	High-field spin susceptibility for ion in the local spin-density functional formalism. <b>1980</b> , 10, 913-921	10

1319	The Cluster Approach in Theoretical Study of Chemisorption. <b>1980</b> , 12, 103-158	41
1318	Electronic magnetic and cohesive properties of some nickel-aluminium compounds. <b>1980</b> , 10, 427-440	143
1317	The virial and Hellmann-Feynman theorems of an inhomogeneous electron gas. <b>1980</b> , 13, 3625-3637	14
1316	Density Functional Calculations for Atoms, Molecules and Clusters. <b>1980</b> , 21, 394-401	216
1315	An Approximate Differential Equation for Calculating the Electron Density in Closed Shell Atoms and in Molecules. <b>1980</b> , 21, 402-408	17
1314	Local-Density Theory of Exchange and Correlation Energies for Valence Electrons. <b>1980</b> , 21, 585-588	10
1313	Self-Consistent Valence Charge Density of the 1T Layer Compounds TiS <sub>2</sub> , TiSe <sub>2</sub> , ZrS <sub>2</sub> and ZrSe <sub>2</sub> . <b>1980</b> , 22, 523-527	5
1312	Extended tight-binding calculations of bulk copper and (001) Cu film. <b>1980</b> , 13, 2807-2816	14
1311	Variation of Adatom Valence-Level Positions With the Distance to a Metal Surface. <b>1980</b> , 22, 165-170	39
1310	Self-consistent relativistic bandstructure for gold. <b>1980</b> , 10, 1135-1148	16
1309	Volume dependence of the magnetic properties of alkali metals. <b>1980</b> , 10, 1567-1573	13
1308	Density functional approximation for the quasiparticle properties of simple metals. II. Application to Li, Rb and Cs. <b>1980</b> , 10, 1737-1751	47
1307	Fermi surface of YAl <sub>2</sub> . <b>1980</b> , 10, 2207-2216	27
1306	Positron-electron correlations near a metal surface and the local density approximation. <b>1980</b> , 10, 253-263	11
1305	A constructive definition of doubly occupied Wannier functions for metals; application to BCC Na. <b>1980</b> , 10, 1167-1175	3
1304	Electron density in simple metals. Relation to bulk and surface properties. <b>1980</b> , 10, 1995-2008	16
1303	Stability of the collinear spin structure of Manganese. <b>1980</b> , 10, 2035-2040	39
1302	3d impurities in Al: density functional results. <b>1980</b> , 10, L123-L127	59

1301	Density functional approximation for the quasiparticle properties of simple metals. I. Theory and electron gas calculations. <b>1980</b> , 10, 1719-1736	38
1300	Pressure dependence of the energy spectrum of trigonal Se. <b>1980</b> , 13, 4953-4963	10
1299	Single particle lifetime in copper. <b>1980</b> , 10, L231-L234	10
1298	Pseudopotential spin-density functional calculation of the ground state properties of Na <sub>2</sub> and Na <sub>2</sub> <sup>+</sup> . <b>1980</b> , 73, 4511-4516	19
1297	Extension of liquid-metal theory to dense partially ionized plasmas. <b>1980</b> , 21, 3140-3151	23
1296	Dynamical analysis of low-energy-electron diffraction intensities from InP (110). <b>1980</b> , 22, 6171-6183	61
1295	Hartree-Fock theory of the inhomogeneous electron gas at a jellium metal surface: Rigorous upper bounds to the surface energy and accurate work functions. <b>1980</b> , 22, 5987-5996	25
1294	Prediction of Fermi-surface pressure dependence in Rb and Cs. <b>1980</b> , 21, 5584-5593	22
1293	Electronic structure and Mössbauer hyperfine interactions of Au(I) compounds. <b>1980</b> , 22, 4203-4214	36
1292	Resonant Photoemission in Barium and Cerium. <b>1980</b> , 45, 204-207	223
1291	Knight Shifts in Simple Metals: A Single-Ion Approach. <b>1980</b> , 44, 175-178	33
1290	Charge-density waves in two- and three-dimensional jellium. <b>1980</b> , 21, 2739-2744	34
1289	5f-electron Delocalization in Americium. <b>1980</b> , 44, 1230-1233	119
1288	Self-consistent study of surfaces of simple metals by the density-matrix method: (100) and (110) surfaces of Na, K, Rb, and Cs. <b>1980</b> , 22, 1806-1817	31
1287	Scattering-theoretic method for defects in semiconductors. II. Self-consistent formulation and application to the vacancy in silicon. <b>1980</b> , 21, 3545-3562	219
1286	One-parameter electronic densities in atoms. <b>1980</b> , 22, 343-347	20
1285	Ab initio self-consistent calculation of ground-state properties, Wannier functions and electronic structure of TiCl. <b>1980</b> , 21, 3571-3580	13
1284	Annihilation of a positron in a vacancy in aluminum. <b>1980</b> , 22, 4572-4589	40

1283	Rayleigh scattering by neutral atoms, 100 eV to 10 MeV. <b>1980</b> , 22, 1970-2004	185
1282	Lattice dynamics and superconducting Tc for dilute H and D interstitials in Al. <b>1980</b> , 21, 5096-5110	14
1281	Self-consistent local-orbital method for calculating surface electronic structure: Application to Cu (100). <b>1980</b> , 21, 2201-2221	186
1280	Pressure-induced structural transitions in partially ionic semiconductors: Self-consistent pseudopotential approach to ZnSe. <b>1980</b> , 22, 4816-4824	36
1279	Spin-dependent correlated atomic pseudopotentials. <b>1980</b> , 22, 649-662	34
1278	Elastic-electron-scattering cross sections for N2 from 0 to 1000 eV. Energy-dependent exchange potentials. <b>1980</b> , 21, 85-94	39
1277	Theory of vacancies near a bimetallic interface. <b>1980</b> , 22, 4776-4783	10
1276	Localized-muffin-tin-orbital basis for atomic-cluster calculations within the local-density formalism. <b>1980</b> , 22, 2614-2625	28
1275	Band calculation of the effect of magnetic impurity atoms on the properties of superconductors. <b>1980</b> , 22, 3165-3172	25
1274	Gradient expansion of the exchange energy density functional: A complementary expansion of the atomic energy functional. <b>1980</b> , 73, 1340-1343	33
1273	Comparative study of the gradient expansion of the atomic kinetic energy functional-neutral atoms. <b>1980</b> , 72, 429-433	88
1272	Electronic states in thorium under pressure. <b>1980</b> , 21, 1489-1496	61
1271	Dependence of the one-electron eigenvalues, $\epsilon_i$ and the total energy, $E$ , on the parameter in the Hartree-Fock-Slater scheme for atoms. <b>1980</b> , 22, 1375-1382	8
1270	Approximate exchange energy as a functional of the electron density. Light atoms. <b>1980</b> , 72, 6299-6302	10
1269	Inhomogeneous electron gas at nonzero temperatures: Exchange effects. <b>1980</b> , 21, 2064-2068	42
1268	Local approximations for the exchange interaction between valence and core electrons. <b>1980</b> , 22, 22-27	17
1267	Self-consistent Al (111) film calculations. <b>1980</b> , 22, 5768-5773	69
1266	Pseudopotential study of alkali metals: Unified approach. <b>1980</b> , 22, 1856-1865	24

1265	Self-consistent calculation of hyperfine fields at impurity sites in ferromagnetic host. <b>1980</b> , 22, 2411-2419	5
1264	"Thomas-Fermi-pseudopotential" approach for calculating the static properties of simple metals. <b>1980</b> , 21, 3074-3086	34
1263	Influence of an improved local-spin-density correlation-energy functional on the cohesive energy of alkali metals. <b>1980</b> , 22, 3812-3815	97
1262	Electronic structure of Sr monochalcogenides. <b>1980</b> , 13, 1995-1999	60
1261	Application of variational principles for single-particle expectation values to the metal surface problem. <b>1980</b> , 22, 1843-1855	1
1260	Screening effects in pair production near threshold. <b>1980</b> , 21, 454-457	24
1259	Self-consistent electronic spectrum of trigonal Te. Pressure dependence of energy spectrum. <b>1980</b> , 22, 2945-2954	15
1258	Diamagnetic shielding of impurities in simple metals. <b>1980</b> , 22, 5490-5494	12
1257	Heat of solution of hydrogen in aluminum. <b>1980</b> , 21, 5594-5600	37
1256	$r \rightarrow$ -space method for the total energy applied to silicon. <b>1980</b> , 21, 4600-4607	10
1255	Hypernetted-chain Euler-Lagrange equations and the electron fluid. <b>1980</b> , 22, 2353-2372	110
1254	Electronic structure of zinc-blende-wurtzite interfaces: ZnS-ZnS (111-0001) and ZnSe-ZnSe (111-0001). <b>1980</b> , 22, 2060-2065	25
1253	Initial-state screening effects in metal Auger spectra: Be. <b>1980</b> , 21, 430-435	75
1252	Application of virial theorem for the total binding energy of neutral atomic systems. <b>1980</b> , 72, 1404-1406	6
1251	Dependence of the He-Scattering Potential at Surfaces on the Surface-Electron-Density Profile. <b>1980</b> , 45, 807-810	381
1250	Dynamical Correlation Effects on the Quasiparticle Bloch States of a Covalent Crystal. <b>1980</b> , 45, 290-294	136
1249	Spin-density functional calculations for chromium. <b>1980</b> , 20, 277-284	90
1248	LEED analysis of the surface structure of Mo(001). <b>1980</b> , 91, 131-152	56

1247	Adatom mobility on the surface planes of a simple metal. <b>1980</b> , 97, 53-72	6
1246	Electrostatic energy and screened charge interaction near the surface of metals with different Fermi surface shape. <b>1980</b> , 94, 179-203	45
1245	A quantitative approach to ionic adsorption. <b>1980</b> , 92, 325-335	9
1244	Interaction-induced transition at low densities in silicon inversion layer. <b>1980</b> , 98, 250-255	8
1243	Nonlocal electrostatic approach to the double layer and adsorption at the electrode-electrolyte interface. <b>1980</b> , 101, 23-48	51
1242	Energy-dependent exchange potentials in low-energy electron diffraction from GaAs(110). <b>1980</b> , 97, 512-528	33
1241	High pressure metallization of CaO: Tentative evidence. <b>1980</b> , 7, 227-230	9
1240	Pressure induced metallization of SrO and BaO: Theoretical estimate of transition pressures. <b>1980</b> , 7, 689-692	25
1239	The Renaissance and Quantitative Development of the Tight-Binding Method. <b>1980</b> , 35, 129-214	89
1238	Scattering of electrons by atoms. <b>1980</b> , 29, 771-867	39
1237	Effective-medium theory of chemical binding: Application to chemisorption. <b>1980</b> , 21, 2131-2136	414
1236	Chemisorption theory for metallic surfaces: Electron localization and the description of surface interactions. <b>1980</b> , 21, 4357-4367	154
1235	Theory of nonuniform electronic systems. I. Analysis of the gradient approximation and a generalization that works. <b>1980</b> , 21, 5469-5493	466
1234	Linear-response theory within the density-functional formalism: Application to atomic polarizabilities. <b>1980</b> , 21, 12-23	284
1233	Theory of the silicon vacancy: An Anderson negative-U system. <b>1980</b> , 21, 5662-5686	380
1232	Calculated specific volumes and magnetic moments of the 3d transition metal monoxides. <b>1980</b> , 15-18, 861-862	2
1231	Electronic structure of conducting $\pi$ -electron systems. <b>1980</b> , 1, 193-212	47
1230	Density-functional approach to local-field effects in finite systems: Photoabsorption in the rare gases. <b>1980</b> , 21, 1561-1572	892

1229	Many-particle effects in the optical spectrum of a semiconductor. <b>1980</b> , 21, 4656-4673	384
1228	Density-functional theory of the metal-insulator transition. <b>1980</b> , 21, 3037-3042	45
1227	Effect of pressure on bonding in MgO. <b>1980</b> , 85, 285-292	41
1226	Microscopic Theory of the Phase Transformation and Lattice Dynamics of Si. <b>1980</b> , 45, 1004-1007	465
1225	Quasiatoms: An approach to atoms in nonuniform electronic systems. <b>1980</b> , 22, 1564-1583	392
1224	Validity of the frozen-core approximation and pseudopotential theory for cohesive energy calculations. <b>1980</b> , 21, 2222-2228	89
1223	A theoretical study of the electronic properties of intercalated graphite. <b>1980</b> , 1, 233-247	5
1222	Relativistic energy bands of NaCl structure uranium compounds. <b>1980</b> , 15-18, 873-874	31
1221	On the Quantum-Mechanical Kinetic Energy as a Measure of the Information in a Distribution. <b>1980</b> , 19, 165-173	192
1220	Metal-insulator transition and local moment formation: A spin-density functional approach. <b>1980</b> , 15-18, 955-956	
1219	Electronic structure calculations and magnetic properties. <b>1980</b> , 15-18, 847-852	63
1218	Extensions of the LSD approximation in density functional calculations. <b>1980</b> , 72, 5357-5362	71
1217	Self-consistent electronic structure of surfaces: Surface states and surface resonances on W(001). <b>1980</b> , 21, 5601-5612	254
1216	Nonlocal pseudopotential calculation of the electronic properties of relaxed GaAs (110) surface. <b>1980</b> , 22, 959-969	66
1215	The theory of electron-molecule collisions. <b>1980</b> , 52, 29-119	668
1214	Many-body effects in semiconductors. <b>1980</b> , 13, 5515-5527	12
1213	Calculation of structurally related properties of bulk and surface Si. <b>1980</b> , 21, 1527-1536	68
1212	Atoms in jellium: background corrections. <b>1980</b> , 10, 321-333	11



1211	Systematization of the stable crystal structure of all AB-type binary compounds: A pseudopotential orbital-radii approach. <b>1980</b> , 22, 5839-5872	235
1210	Direct and reverse transformations between electron density and electron momentum density. <b>1981</b> , 24, 2906-2912	50
1209	Magnetism of nickel. <b>1981</b> , 23, 5974-5977	147
1208	Self-consistent calculation of Compton profiles, X-ray structure factors, and band structure for silicon. <b>1981</b> , 14, 347-351	11
1207	The role of single-particle density in chemistry. <b>1981</b> , 53, 95-126	134
1206	Self-interaction correction to density-functional approximations for many-electron systems. <b>1981</b> , 23, 5048-5079	16702
1205	Method for calculating wave functions in a nonspherical potential. <b>1981</b> , 23, 6301-6306	33
1204	Interaction between Closed-Shell Systems and Metal Surfaces. <b>1981</b> , 46, 842-845	265
1203	Iron clusters: Electronic structure and magnetism. <b>1981</b> , 24, 5673-5692	163
1202	Chemisorption of molecular hydrogen on simple metal surfaces. <b>1981</b> , 104, 510-526	105
1201	Variational spherical model of small metallic particles. <b>1981</b> , 106, 265-271	183
1200	Ground-state properties of diamond. <b>1981</b> , 24, 6121-6124	129
1199	Atoms embedded in an electron gas: Immersion energies. <b>1981</b> , 24, 3037-3047	329
1198	Orthonormal orbitals for the representation of an arbitrary density. <b>1981</b> , 24, 680-682	288
1197	Structural-energy calculations based on norm-conserving pseudopotentials and localized Gaussian orbitals. <b>1981</b> , 24, 4745-4752	150
1196	Itinerant electron magnetism. <b>1981</b> , 44, 329-409	258
1195	Ab initio calculation of the static structural properties of Al. <b>1981</b> , 24, 4224-4229	93
1194	The gaps of the ideal TiS <sub>2</sub> and TiSe <sub>2</sub> . <b>1981</b> , 14, L75-L80	20

1193	Pseudopotential spin-density-functional calculation of the electronic properties of small lithium and sodium clusters. <b>1981</b> , 106, 280-286	32
1192	The electronic structure of antiferromagnetic chromium. <b>1981</b> , 11, 97-111	113
1191	Calculations for 4f atoms using new exchange-correlation schemes. <b>1981</b> , 52, 2149-2151	6
1190	Positron bremsstrahlung. <b>1981</b> , 24, 1358-1363	12
1189	Slater transition state calculations of valence electron spin-orbit splitting in atoms. <b>1981</b> , 75, 5971-5971	2
1188	Full-potential self-consistent linearized-augmented-plane-wave method for calculating the electronic structure of molecules and surfaces: O <sub>2</sub> molecule. <b>1981</b> , 24, 864-875	1836
1187	Universal Binding Energy Relations in Metallic Adhesion. <b>1981</b> , 7, 19-30	10
1186	First-principles electronic structure of Si, Ge, GaP, GaAs, ZnS, and ZnSe. II. Optical properties. <b>1981</b> , 24, 3417-3429	199
1185	Density-functional theory of the correlation energy in atoms and ions: A simple analytic model and a challenge. <b>1981</b> , 23, 2785-2789	104
1184	On the Disappearance of Ferromagnetism in Disordered Fe-Al Alloys. <b>1981</b> , 50, 70-76	15
1183	Optical Properties of 3d Transition Metals: V, Cr, Fe and Ni. <b>1981</b> , 50, 835-842	10
1182	Method of approximate natural orbitals. <b>1981</b> , 22, 429-431	
1181	Comparative studies of static dipole polarizabilities of ionic crystals on the basis of different self-consistent potentials. <b>1981</b> , 59, 299-307	8
1180	Self-Consistent APW $\bar{\mu}$ p method. II. Application to NaCl. <b>1981</b> , 20, 933-949	3
1179	Simple Non-Local Calculation of Jellium Metal Surface Properties. <b>1981</b> , 105, 147-153	7
1178	Variational Hartree-Fock Calculations of the Ground State Energy of Metallic Hydrogen. <b>1981</b> , 105, 769-775	3
1177	A Muffin-Tin Calculation of the Electronic d-Bandwidth of F. C. C. Palladium. <b>1981</b> , 106, K33-K38	
1176	Electrical Resistivity of Liquid Ti at Its Melting Point. <b>1981</b> , 108, K65-K68	

1175	Work function of metals: Relation between theory and experiment. <b>1981</b> , 11, 293-338	54
1174	Density-functional method for the inhomogeneous electron-hole plasma in polar substances. <b>1981</b> , 109, 161-177	2
1173	One-electron formulation of the electron-phonon coupling problem by the density functional approach. <b>1981</b> , 107, 669-670	1
1172	Correlated wave function theory of metal surfaces. <b>1981</b> , 108, 871-872	
1171	A note on finite temperature Thomas-Fermi, Hartree-Fock-Slater calculations. <b>1981</b> , 81, 169-171	5
1170	Relativistic gradient expansion of the kinetic energy density. <b>1981</b> , 81, 447-450	11
1169	Exchange splitting of ferromagnetic nickel within the local potential approximation. <b>1981</b> , 86, 45-47	21
1168	A mixed bonding band structure calculation for GaAs and AlAs using the APW-k $\cdot$ p method. <b>1981</b> , 42, 291-296	5
1167	Theory of donor-bound multi-exciton complexes in Germanium. <b>1981</b> , 40, 165-168	3
1166	Density matrix approach to the study of a monovacancy in sodium. <b>1981</b> , 40, 37-39	10
1165	Face dependent surface energies of simple metals. <b>1981</b> , 37, 91-96	44
1164	Choice of $\epsilon_n$ scattering from $X_\mu$ potentials. <b>1981</b> , 38, 961-963	4
1163	Comment on the self-interaction correction to the local spin density exchange splitting. <b>1981</b> , 38, 989-990	1
1162	Microscopic theory of the static structural properties and phase transformation of Ge. <b>1981</b> , 38, 625-627	49
1161	Self-interaction corrections in the density functional formalism. <b>1981</b> , 37, 249-252	50
1160	Density functional calculation of stopping power of an electron gas for slow ions. <b>1981</b> , 37, 779-781	367
1159	Self-consistent energy band calculations. <b>1981</b> , 44, 139-212	105
1158	Computation of the total energy of transition metals by a model electron density function. <b>1981</b> , 24, 569-573	

1157	Hydrogen in metals: Electronic properties. <b>1981</b> , 8, 437-444	6
1156	RPA response to a static point charge near a metal surface: Finite-barrier treatment. <b>1981</b> , 31, 465-468	2
1155	Accuracy of transition states of many-electron systems on the basis of a quantum-statisticalab initio theory. <b>1981</b> , 50, 125-129	
1154	Identification and Properties of Defects in GaP. <b>1981</b> , 47, 413-416	51
1153	Theory of Electric Field Gradient Due to a Point Defect in a Cubic Metal. <b>1981</b> , 46, 610-614	25
1152	Foundations of Walsh's rules for molecular shape. <b>1981</b> , 74, 2973-2974	15
1151	Spin-Polarized Band-Structure Determination of the Si <sub>2</sub> Molecular Ground State by the Method of Full-Potential Linearized Augmented Plane Waves. <b>1981</b> , 47, 705-708	11
1150	Easily Implementable Nonlocal Exchange-Correlation Energy Functional. <b>1981</b> , 47, 446-450	281
1149	Effect of the local exchange-correlation correction to electron density on the properties of inorganic solids. <b>1981</b> , 74, 6319-6341	10
1148	Self-consistent calculations of oxygen monolayers on Al(111) films. <b>1981</b> , 23, 4960-4964	16
1147	Localized orbitals for band-structure calculations in complex semiconductors. <b>1981</b> , 24, 5949-5959	14
1146	Nonlocal density approximation to exchange in relativistic-augmented-plane-wave band-structure calculations and consistent interpretation of photoemission data on Rh(110). <b>1981</b> , 23, 3113-3120	42
1145	Exchange-correlation effects in silicon (111) inversion layers: Strain-enhanced valley-occupancy phase transition. <b>1981</b> , 23, 1839-1842	11
1144	Use of a local-density approximation for exchange-correlation potentials in multichannel atomic quantum-defect calculations. <b>1981</b> , 23, 2761-2775	20
1143	Band structure, Fermi surface, Compton profile, and optical conductivity of paramagnetic chromium. <b>1981</b> , 23, 4977-4987	84
1142	Virial theorem in the density-functional formalism: Forces in H <sub>2</sub> . <b>1981</b> , 24, 6795-6800	56
1141	Use of local exchange potentials in the calculation of photoionization and electron-ion scattering. <b>1981</b> , 23, 2905-2913	7
1140	Sixth-order term of the gradient expansion of the kinetic-energy density functional. <b>1981</b> , 24, 1682-1688	128

1139	Initial oxidation of the Al(001) surface: Self-consistent electronic structure of clean Al(001) and Al(001)H <sub>2</sub> O. <b>1981</b> , 23, 3859-3876	57
1138	Positron annihilation in metal-vacancy-hydrogen complexes. <b>1981</b> , 24, 2884-2887	21
1137	The surface photoeffect. <b>1981</b> , 24, 628-638	76
1136	Dielectric formulation of strongly coupled electron liquids at metallic densities. V. Possibility of a charge-density-wave instability. <b>1981</b> , 24, 3226-3230	21
1135	A semiempirical XPS calculation of the KVV Auger line shape of O <sub>2</sub> . <b>1981</b> , 75, 300-305	26
1134	Potentials, band structures, and Fermi surfaces in the noble metals. <b>1981</b> , 23, 2684-2696	126
1133	Self-consistent non-muffin-tin augmented-plane-wave calculation of the band structure of silicon. <b>1981</b> , 23, 1652-1663	20
1132	First-order phase transitions in intermediate-valence solids— theory based on metallic hydrogen. <b>1981</b> , 24, 5664-5672	0
1131	Self-consistent band-structure theory of the metal-insulator transition. <b>1981</b> , 24, 4879-4882	28
1130	An atoms in molecules approach to density functional theory. <b>1981</b> , 75, 828-833	22
1129	Free-atom-metal shifts in the M <sub>4,5</sub> N <sub>4,5</sub> N <sub>4,5</sub> Auger spectra of Ag, Cd, In, Sn, Sb, and Te. <b>1981</b> , 23, 4362-4368	19
1128	Single-particle energy levels in doped semiconductors at densities below the metal-nonmetal transition. <b>1981</b> , 23, 1920-1935	69
1127	Nonadjustable local model potentials for the exchange interaction between valence and core electrons. <b>1981</b> , 24, 649-655	12
1126	Influence of relativistic contributions to the effective potential on the electronic structure of Pd and Pt. <b>1981</b> , 23, 6377-6398	96
1125	Metal-insulator transition in dilute alkali-metal systems. <b>1981</b> , 23, 552-560	15
1124	Comparison of methods for the calculation of phase stability in silicon. <b>1981</b> , 24, 7210-7216	25
1123	Ab initio calculation of interatomic potentials and electronic properties of a simple metal—Al. <b>1981</b> , 24, 7057-7070	62
1122	Effects of electron-positron correlation on positron annihilation: Self-consistent band-structure calculations in Al. <b>1981</b> , 24, 7423-7426	45

1121	Cohesive Energies of Simple Metals as Determined from Atomic Kinetic Energies. <b>1981</b> , 47, 387-390	19
1120	Band structure measurements and multi-electron effects (satellites) for nearly-filled d-band metals: Fe, Co, Ni, Cu, Ru, and Pd (invited). <b>1981</b> , 52, 1658-1663	28
1119	Analytic velocity-dependent potential for bound and scattering states of electrons and atoms. <b>1981</b> , 24, 3010-3018	50
1118	Momentum eigenfunctions in the complex momentum plane. IV. The construction of local potential functions. Perturbation series. <b>1981</b> , 74, 1225-1238	6
1117	Electronic structure of the rotation twin stacking fault in $\alpha$ -Sn. <b>1981</b> , 23, 2563-2566	13
1116	Density-functional theory of the electron-hole liquid in doped Ge. <b>1981</b> , 23, 4029-4034	2
1115	Atomic structure and properties of polar Ge-GaAs(100) interfaces. <b>1981</b> , 24, 3445-3455	141
1114	Improved correlation corrections to the local-spin-density approximation. <b>1981</b> , 24, 4264-4270	93
1113	Theory of dense hydrogen. <b>1981</b> , 24, 1624-1635	64
1112	Force calculations in the density functional formalism. <b>1981</b> , 75, 3904-3908	57
1111	Elastic moduli of rhodium: Correct prediction by a new theoretical method. <b>1981</b> , 24, 2254-2256	43
1110	Phonon optics in semiconductors: Phonon generation and electron-phonon scattering in n-GaAs epilayers: I. Theory. <b>1981</b> , 24, 4692-4713	24
1109	Theoretical study of the electronic structure of GaP(110). <b>1981</b> , 24, 6029-6042	86
1108	Self-consistent second-order perturbation treatment of multiplet structures using local-density theory. <b>1981</b> , 24, 943-954	49
1107	Bound excited states in density-functional theory. <b>1981</b> , 23, 2127-2133	43
1106	Variational calculations of low-index crystal face-dependent surface energies and work functions of simple metals. <b>1981</b> , 23, 6512-6523	50
1105	Resonant two-electron excitation in copper. <b>1981</b> , 24, 4121-4127	52
1104	Density-functional calculation of static and dynamic properties of GaAs. <b>1981</b> , 24, 2311-2314	156

1103	Direct method of calculation of dynamic effective charges: Application to GaAs. <b>1981</b> , 24, 2081-2088	50
1102	Electron-electron correlation in beryllium and related properties. <b>1981</b> , 11, L1-L6	4
1101	Density matrix calculations for molecules and clusters I. Theoretical foundations. <b>1981</b> , 14, 4441-4450	5
1100	Spin polarised ultra-thin metallic films. <b>1981</b> , 14, 89-95	40
1099	Exchange and correlation potentials for finite temperature quantum calculations at intermediate degeneracies. <b>1981</b> , 14, 629-646	84
1098	The dynamic response of a non-uniform distribution of electrons. <b>1981</b> , 14, L167-L169	2
1097	The cellular method for graphite. <b>1981</b> , 14, L213-L220	22
1096	A self-consistent theory of inhomogeneous liquid metals: Calculations of the electron and ion density profiles and the liquid-vapour surface tension of the alkali metals. <b>1981</b> , 14, 5225-5246	55
1095	Self-consistent screening of hydrogen in zirconium. <b>1981</b> , 11, 1023-1033	13
1094	Electronic structure of $(\text{SN})_x$ and its intercalate $(\text{SNBr}_y)_x$ . <b>1981</b> , 14, 5091-5107	15
1093	The surface structure of Mo(110) determined by LEED. <b>1981</b> , 14, 5391-5410	39
1092	Density functional-pseudopotential approach to the heat of formation in alloys of alkali metals. <b>1981</b> , 11, 2045-2053	15
1091	Theory of spin polarised inhomogeneous relativistic electron gas. <b>1981</b> , 14, 4291-4302	35
1090	The electronic structure of calcium. <b>1981</b> , 11, 805-820	43
1089	Density matrix study of $\text{Cs}_2$ , $\text{Au}_2$ and $\text{CsAu}$ . <b>1982</b> , 15, 2569-2578	6
1088	Ab initio calculation of the phonon dispersion relation: Application to Si. <b>1982</b> , 25, 4317-4320	116
1087	Density-functional theory of hydrogen plasmas. <b>1982</b> , 26, 2096-2104	183
1086	Electron density in sodium fluoride by $\bar{\Gamma}$ -ray diffractometry. <b>1982</b> , 25, 2545-2549	9

1085	Optical response of uniaxial semiconductors. II. Optical and electron-energy-loss spectra of ZrS <sub>2</sub> and ZrSe <sub>2</sub> . <b>1982</b> , 26, 5807-5814	13
1084	Inelastic scattering of rare-gas atoms from metal surfaces. Excitation of electron-hole pairs. <b>1982</b> , 25, 2514-2521	54
1083	Asymptotic freedom in solids: A theorem. <b>1982</b> , 25, 3474-3481	40
1082	Multiple-scattering theory of itinerant electron magnetism in random muffin-tin alloys. <b>1982</b> , 26, 367-378	5
1081	Bonding in the first-row diatomic molecules within the local spin-density approximation. <b>1982</b> , 26, 1781-1790	116
1080	Electronic structure of semiconductor surface inversion layers at finite temperature. The Si(100)-SiO <sub>2</sub> system. <b>1982</b> , 26, 960-974	31
1079	Temperature-dependent nonlinear screening of a proton in an electron gas. <b>1982</b> , 25, 489-495	35
1078	Influence of charged impurities on Si inversion-layer electrons. <b>1982</b> , 26, 6808-6825	54
1077	X-ray form factors and the electronic structure of graphite. <b>1982</b> , 26, 5382-5390	200
1076	Remarks on local and nonlocal exchange and correlation-energy calculations of surface energies and work functions. <b>1982</b> , 25, 6275-6280	10
1075	Dynamical aspects of correlation corrections in a covalent crystal. <b>1982</b> , 25, 2867-2888	229
1074	Universal binding-energy relation in chemisorption. <b>1982</b> , 25, 1419-1422	88
1073	Quasiband crystal-field method for calculating the electronic structure of localized defects in solids. <b>1982</b> , 26, 846-895	64
1072	A radially restricted Thomas-Fermi theory for atoms. <b>1982</b> , 76, 5043-5050	9
1071	Phonon spectra and crystal structure of Zn and Cd using the resonant model-potential approach. <b>1982</b> , 26, 743-752	3
1070	Electronic structure of chemisorbed chalcogen atoms on Ni (hkl) surfaces. <b>1982</b> , 25, 2124-2137	35
1069	Spin response of a charge-density wave. <b>1982</b> , 26, 1671-1681	5
1068	Theory of spin-orbit and many-body effects on the Knight shift. <b>1982</b> , 25, 3091-3116	40



1067	Improvement on the hypernetted-chain equations for dense plasmas. <b>1982</b> , 25, 2434-2436	55
1066	Theory of donor-bound multiexciton complexes in germanium and silicon. <b>1982</b> , 25, 3897-3914	12
1065	Many-body theory of magnetic susceptibility of electrons in solids. <b>1982</b> , 26, 1903-1928	28
1064	A new gradient expansion of the exchange energy to be used in density functional calculations on atoms. <b>1982</b> , 76, 6057-6059	15
1063	Multiple-scattering approach to band theory. II. Fast band theory. <b>1982</b> , 26, 1597-1607	25
1062	Embedded-atom calculations of Auger and x-ray photoemission shifts for metallic elements. <b>1982</b> , 25, 67-77	30
1061	Self-consistent electronic structure of the contracted tungsten (001) surface. <b>1982</b> , 25, 755-761	38
1060	On the atomic kinetic energy functionals with full Weizsacker correction. <b>1982</b> , 76, 1467-1472	74
1059	Theory of ab initio pseudopotential calculations. <b>1982</b> , 25, 7403-7412	138
1058	Ferrell's criterion on the correlation energy of the electron gas: Generalization and contributions of the exchange and correlation terms. <b>1982</b> , 25, 1374-1376	5
1057	On the nature of the correction to the Weizsacker term. <b>1982</b> , 76, 3157-3160	42
1056	On the functional derivative of the kinetic energy density functional. <b>1982</b> , 77, 4576-4585	44
1055	Magnetism of surfaces and interfaces. <b>1982</b> , 53, 1997-2001	29
1054	Theory of amorphous SiO <sub>2</sub> and SiO <sub>x</sub> . II. Electron states in an intrinsic glass. <b>1982</b> , 26, 6622-6632	64
1053	Numerical Hartree-Fock calculations on diatomic molecules. <b>1982</b> , 76, 6037-6045	118
1052	Positron annihilation as a probe for impurities trapped by vacancy clusters. <b>1982</b> , 26, 5264-5267	7
1051	Density-functional formulation of the generalized pseudopotential theory. II. <b>1982</b> , 26, 1754-1780	111
1050	Electronic structure and chemical bonding in metallic clusters of binary and ternary transition metal chalcogenides. I. SCF MS X-ray study including relativistic effect of PbMo <sub>6</sub> S <sub>8</sub> . <b>1982</b> , 76, 6060-6066	18

1049	Spherical solid model in nonsimple metals: Application to hydrogen impurity in copper. <b>1982</b> , 25, 7331-7340	9
1048	Off-diagonal occupation numbers in local-density theory. <b>1982</b> , 26, 3203-3210	4
1047	Electrodynamic response of a bounded electron gas in hydrodynamic formalism: Theory and applications. <b>1982</b> , 26, 6559-6570	9
1046	Nrepresentability of electron density and first-order density matrix. <b>1982</b> , 26, 1845-1847	21
1045	New approach to the calculation of ground-state properties in solids. <b>1982</b> , 26, 1512-1526	8
1044	Statistical atomic models with minimal-basis-set-type electron densities. <b>1982</b> , 25, 107-112	12
1043	Compton Profile of Beryllium. <b>1982</b> , 49, 1452-1455	36
1042	Depolarisation studies of positive muons in copper, vanadium, niobium and tantalum single crystals. <b>1982</b> , 12, 875-893	21
1041	Surface electronic structure. <b>1982</b> , 45, 223-284	133
1040	Generalized Ewald Potential in the Non Muffin-Tin APW Method. <b>1982</b> , 51, 116-123	4
1039	Orbital self-interaction in Hartree-Fock and density functional theories. <b>1982</b> , 15, L1183-L1186	9
1038	Self-consistent pseudopotential calculations of the equilibrium bulk properties of diamond-type semiconductors. <b>1982</b> , 15, L739-L742	2
1037	The electronic structure of actinide monocarbides. <b>1982</b> , 15, 6361-6378	17
1036	Density functional formalism in the problem of mixed-valence compounds. <b>1982</b> , 15, L1125-L1130	1
1035	Electronic Properties of Point Defects in Metals. <b>1982</b> , 25, 703-707	1
1034	Density-Functional Calculations of Auger and X-Ray Photoemission Shifts for Metallic Elements. <b>1982</b> , 25, 708-712	3
1033	Hypernetted Chain Theory of Charged Impurity. <b>1982</b> , 25, 943-951	48
1032	Transition rate calculations of some forbidden lines in X-ray emission spectra. <b>1982</b> , 15, 2339-2348	1

1031	Interaction of helium with a metal surface. <b>1982</b> , 15, 2275-2291	114
1030	Pseudopotential perturbation theory for the surface tension of liquid metals. <b>1982</b> , 15, 353-375	30
1029	Phonon calculations for KCl ab initio. <b>1982</b> , 15, 2093-2103	5
1028	A note on face-dependent surface properties of simple metals. <b>1982</b> , 15, 4717-4725	11
1027	Aspects of self-consistent procedures in surface pseudopotential calculations. <b>1982</b> , 15, 3627-3637	3
1026	Estimates of non-local corrections to total, ionisation, and single-particle energies. <b>1982</b> , 15, 2139-2150	65
1025	Self-consistent pseudopotential calculation of the electronic properties of the InP (110) surface. <b>1982</b> , 15, 1099-1109	31
1024	Phonon spectra in the Frolich approach-application to the transition metals. <b>1982</b> , 12, 1577-1602	3
1023	Theory of the heat of formation in homovalent disordered solid alloys of non-transition metals. <b>1982</b> , 12, 1907-1921	15
1022	Soft X-ray emission and the d band width in nickel. <b>1982</b> , 12, 1539-1546	7
1021	Cohesion, Compound Formation and Phase Diagrams from First Principles. <b>1982</b> , 19, 17	1
1020	Density Functional Theory of Interplane Cohesion in Graphite and Graphite Intercalation Compounds. <b>1982</b> , 20, 123	
1019	Gaussian Quadrature XMethod for Self-Consistent Hartree-Fock-Slater Equation. I. Application to the Total Energy Calculation of Diatomic Molecule. <b>1982</b> , 51, 4028-4035	11
1018	Recent Development in the Statistical Theory of High-Density Plasmas. <b>1982</b> , T2A, 198-205	1
1017	Theory of static structural properties, crystal stability, and phase transformations: Application to Si and Ge. <b>1982</b> , 26, 5668-5687	794
1016	Electronic structure of the actinide nitride series. <b>1982</b> , 29, 257-261	27
1015	Schrödinger fluid dynamics of many-electron systems in a time-dependent density-functional framework. <b>1982</b> , 77, 342-348	139
1014	Electronic structure of $\text{BaAl}_2\text{O}_3$ . <b>1982</b> , 15, 5399-5410	92

1013	Charge interaction in layered systems with spatial dispersion. <b>1982</b> , 121, 375-395	18
1012	Adatom binding at the surface ledges of a jellium metal. <b>1982</b> , 116, 522-538	59
1011	On the feasibility of studying surface magnetism by spin-polarized low-energy electron diffraction. <b>1982</b> , 117, 285-293	35
1010	Theory of space-charge layers in narrow-gap semiconductors. <b>1982</b> , 113, 131-136	84
1009	Green's-function methods for electronic-structure calculations. <b>1982</b> , 26, 5433-5444	213
1008	New approach for solving the density-functional self-consistent-field problem. <b>1982</b> , 26, 3114-3137	151
1007	Electron densities in search of Hamiltonians. <b>1982</b> , 26, 1200-1208	502
1006	Dielectric matrix scheme for fast convergence in self-consistent electronic-structure calculations. <b>1982</b> , 25, 4260-4262	91
1005	Calculated electron affinities of the elements. <b>1982</b> , 25, 1265-1271	144
1004	Universal behavior of exchange-correlation energy in electron-hole liquid. <b>1982</b> , 25, 6492-6495	245
1003	The Density Functional Description OP Atomic Clusters. <b>1982</b> , 33-41	
1002	Study of the density-gradient expansion for the exchange energy. <b>1982</b> , 26, 4371-4377	210
1001	Pseudopotentials and Total Energy Calculations. <b>1982</b> , T1, 5-10	332
1000	Band structure calculation and photoemission analysis of iridium. <b>1982</b> , 12, 921-933	44
999	Nonlinear ionic pseudopotentials in spin-density-functional calculations. <b>1982</b> , 26, 1738-1742	2046
998	Electronic structure of simple deep-level defects in semiconductors. <b>1982</b> , 115-148	6
997	Pseudopotentials that work: From H to Pu. <b>1982</b> , 26, 4199-4228	2984
996	Covalent effects in the effective-medium theory of chemical binding: Hydrogen heats of solution in the 3d metals. <b>1982</b> , 26, 2875-2885	390

- 995 Strongly coupled plasmas: high-density classical plasmas and degenerate electron liquids. **1982**, 54, 1017-1059 1164
- 994 Ab initio calculation of the tetragonal shear moduli of the cubic transition metals. **1982**, 26, 1527-1537 37
- 993 Comparison of the orbitals of Neon, Argon and Krypton calculated by the hartree-fock and the X $\beta$  methods with several values of  $\beta$  **1982**, 53, 247-254 10
- 992 Pressure effects on bonding in CaO: Comparison with MgO. **1982**, 87, 303-310 40
- 991 New model dielectric function and exchange-correlation potential for semiconductors and insulators. **1982**, 25, 6310-6316 211
- 990 Insulating Nickel at a Pressure of 34 TPa. **1982**, 49, 1198-1201 67
- 989 Non-muffin-tin and relativistic interaction effects on the electronic structure of noble metals. **1982**, 25, 713-725 91
- 988 Theory of lattice-dynamical properties of solids: Application to Si and Ge. **1982**, 26, 3259-3272 271
- 987 Magnetism in iron and nickel. **1982**, 25, 5766-5777 93
- 986 Total-energy all-electron density functional method for bulk solids and surfaces. **1982**, 26, 4571-4578 861
- 985 Density-Functional Theory for Fractional Particle Number: Derivative Discontinuities of the Energy. **1982**, 49, 1691-1694 2236
- 984 Time-dependent Kohn-Sham density-functional theory. **1982**, 26, 2243-2244 87
- 983 Generalisation of the virial theorem and related theorems for jellia with arbitrary background densities and application to special geometries and profiles. I. General theory and application to the jellium sphere and the semi-infinite electron gas. **1982**, 15, 4807-4820 11
- 982 Electron-electron interaction and single-particle properties in copper. **1982**, 12, 281-292 21
- 981 Density functional theory. **1982**, 35, 36-43 97
- 980 Ab initio calculation of phonon frequencies of Al. **1982**, 25, 6139-6145 79
- 979 Surface properties of simple metals via inhomogeneous linear electronic response. I. Theory. **1982**, 15, 7429-7456 27
- 978 Bond analysis of heats of formation: application to some group VIII and IB hydrides. **1982**, 12, 141-161 100

977	Fixed-shell statistical atomic models with piecewise exponentially decaying electron densities. <b>1982</b> , 25, 2901-2912	18
976	SCF-X $\alpha$ -SW electron densities with the overlapping sphere approximation. <b>1982</b> , 46, 449-463	4
975	Electron removal energies in Kohn-Sham density-functional theory. <b>1982</b> , 26, 5445-5450	100
974	High-Pressure Structural Phase Transitions in Na, Mg, and Al. <b>1982</b> , 48, 809-812	124
973	Electronic properties of two-dimensional systems. <b>1982</b> , 54, 437-672	6270
972	Efficient and accurate expansion methods for molecules in local density models. <b>1982</b> , 76, 1949-1960	399
971	An alternative one-particle approach to exchange and correlation in N-electron systems. <b>1982</b> , 48, 209-218	28
970	Self-consistent-field-Korringa-Kohn-Rostoker-coherent-potential approximation for random alloys. <b>1982</b> , 46, 95-98	70
969	Energy of intermetallide formation for 3d transition metals. <b>1982</b> , 25, 420-424	
968	Calculation of the formation energy of alloys using the method of the model functional of electron density. <b>1982</b> , 25, 30-33	
967	Electronic structure of transition metals under pressure. <b>1982</b> , 25, 1112-1122	1
966	Electron configuration and x-ray emission spectra of intermetallic compounds with CsCl structure. <b>1982</b> , 25, 1135-1148	1
965	n-Particle correlation functions of an interacting electron gas. <b>1982</b> , 50, 288-296	3
964	Isotope effect in chemical shifts of $\mu$ and H $^+$ : MuBr versus HBr and MuHO versus H $_2$ O. <b>1982</b> , 12, 261-278	5
963	Statistical theory of diffusion. Quantum states, configurational shape of a potential, and two types of noncoherent transitions of a hydrogen atom in metals. <b>1982</b> , 50, 81-93	2
962	Density functional formalism at finite temperatures with some applications. <b>1982</b> , 87, 259-311	133
961	Effective core potential calculations on the NiH $_4$ ion as a test case for studying rotational barriers. <b>1982</b> , 66, 453-458	20
960	Comments on the electronic structure of dimeric copper, as calculated with the hartree-fock-later method. <b>1982</b> , 86, 316-319	7

959	X-ray excited Auger studies of metals and alloys. <b>1982</b> , 11-12, 730-760	55
958	Systematics of final state screening in 5d metals. <b>1982</b> , 43, 257-260	20
957	Ab initio calculation of the static structural properties of Be. <b>1982</b> , 42, 861-863	34
956	Self-consistent band structures of higher stage graphite intercalation compounds. <b>1982</b> , 44, 761-765	11
955	Cellular theory of ordered AB alloys. <b>1982</b> , 44, 931-935	3
954	Generalization of the Kohn-Sham $\epsilon$ result for excited states. <b>1982</b> , 42, 147-148	11
953	Inner electron binding energies of chemisorbed atoms. <b>1982</b> , 44, 99-104	9
952	Energy bands and Fermi surfaces of quasi-one-dimensional transition metal chalcogenides Nb <sub>3</sub> X <sub>4</sub> . <b>1982</b> , 43, 607-612	40
951	Calculation of the lattice dynamical properties of Ge. <b>1982</b> , 43, 391-393	5
950	Self-consistent calculations for the electronic structure of a vacancy in copper. A solution of the embedding problem. <b>1982</b> , 42, 701-704	30
949	Ab-initio calculation of the heat of formation in the metallic transition metal monoxides. <b>1982</b> , 44, 339-343	3
948	Self-interaction correction for energy band calculations: Application to LiCl. <b>1982</b> , 41, 827-829	63
947	Inferno: A better model of atoms in dense plasmas. <b>1982</b> , 27, 335-339	81
946	Densities, density-functionals and electron fluids. <b>1982</b> , 92, 1-44	169
945	Pseudopotential theory of spin susceptibility of metals. <b>1982</b> , 92, 300-304	6
944	The surface exchange energy of a metal for the full range of wave vector fluctuations. <b>1982</b> , 89, 27-30	3
943	Effective core potential calculations on small molecules containing transition metal atoms. <b>1982</b> , 66, 459-464	36
942	A comparative analysis of the applicability of X $\alpha$ methods for the calculation of molecules and clusters. <b>1982</b> , 23, 274-289	1

941	Local Density Approximation Applied to Bound Excitons in Polar Semiconductors. <b>1982</b> , 110, 157-167	4
940	Electronic Structure and Soft X-Ray Emission Spectra of 5d Transition Metals. <b>1982</b> , 111, 11-52	38
939	Effective Interactions in Jellium Including Exchange. <b>1982</b> , 111, 365-374	9
938	Electronic Relaxation Effects in Core Level Spectra of Solids. <b>1982</b> , 112, 9-49	54
937	A Modified Local Density Approximation. Electron Density in Inversion Layers. <b>1982</b> , 113, 165-178	134
936	The Electronic Structure of a Hydrogen Impurity in Aluminium I. The Jellium Model. <b>1982</b> , 113, 747-756	7
935	The electronic structure of a hydrogen impurity in aluminium II. The bandstructure model. <b>1982</b> , 114, 337-348	5
934	Some properties of reduced density matrices, correlated and uncorrelated, for pure and mixed states. <b>1982</b> , 21, 269-273	10
933	In defense of the Hohenberg-Kohn theorem and density functional theory. <b>1982</b> , 21, 511-513	16
932	Virial fragments and the Hohenberg-Kohn functional. <b>1982</b> , 21, 927-935	2
931	Quantum fluid dynamics of many-electron systems in three-dimensional space. <b>1982</b> , 22, 871-888	21
930	Electronic structure of hydrogen and muonium in simple metals. <b>1983</b> , 2, 883-897	4
929	Density-functional theory of the Compton profile anisotropy of copper metal. <b>1983</b> , 54, 17-24	32
928	Calculation of high-pressure phases of A1. <b>1983</b> , 27, 5986-5991	78
927	Electronic structure of semiconductors with doping superlattices. <b>1983</b> , 27, 3538-3546	130
926	Formation and coupling of magnetic moments in Heusler alloys. <b>1983</b> , 28, 1745-1755	756
925	Density-functional theory applied to phase transformations in transition-metal alloys. <b>1983</b> , 27, 5169-5172	844
924	A simple analytic representation for first-principles pseudopotentials. <b>1983</b> , 16, L29-L34	3



923	Collective modes in semiconductor superlattices. <b>1983</b> , 44, 807-812	25
922	Photoelectron spectroscopy of chemisorbed atoms. <b>1983</b> , 23, 561-566	9
921	Energy-band structure and charge distribution for BaC6. <b>1983</b> , 23, 1223-1230	22
920	Recent investigations on the electronic structure of the fourth and fifth group transition metal monocarbides, mononitrides, and monoxides. <b>1983</b> , 23, 1317-1353	178
919	Gradient expansion correction to the Dirac exchange term in statistical models for the Na atom with shell structure. <b>1983</b> , 23, 1973-1978	2
918	The modified Hartree-Hock self-consistent field equation. <b>1983</b> , 23, 1979-1987	
917	Density functionals for coulomb systems. <b>1983</b> , 24, 243-277	866
916	The electronic structure of a hydrogen impurity in aluminium III. The energy and the metal-impurity interaction. <b>1983</b> , 115, 53-61	11
915	Generalized pseudoatom model of inhomogeneous systems and its application to a simple-metal surface. <b>1983</b> , 116, 179-190	2
914	Density Functional Approach for Metals. The Matheiss Prescription. <b>1983</b> , 118, 373-379	5
913	K and L absorption spectra of aluminium from APW calculation. <b>1983</b> , 119, 269-277	9
912	Electronic structure of impurities in dilute alloys of Al. <b>1983</b> , 119, 381-389	9
911	The Relativistic Linear Muffin-Tin Orbital Method Application to Au. <b>1983</b> , 120, 283-296	124
910	Ab initio calculation of electron transfer rates in a model of the metal-electrolyte interface. <b>1983</b> , 150, 355-363	3
909	Predictions of the bond length and vibrational frequency of Ge2. <b>1983</b> , 102, 440-441	27
908	MS X <sub>23</sub> and LCAO X <sub>23</sub> calculations of ionization potentials. A comparison with Koopmans' theorem and accurate calculations. <b>1983</b> , 76, 185-193	8
907	Local force variations caused by isoelectric impurities: Method of determination from first principles. <b>1983</b> , 116, 52-57	2
906	Ab initio determination of static, dynamic and dielectric properties of semiconductors. <b>1983</b> , 117-118, 511-516	10

905	Microscopic calculation of the phonon frequencies of silicon, including exchange and correlation. <b>1983</b> , 117-118, 537-539	
904	Analytic calculation of the structural properties of semiconductors. <b>1983</b> , 117-118, 579-581	3
903	Semiconductor surface structures. <b>1983</b> , 3, 193-300	243
902	Polarizabilities of heavy closed-shell atoms from the relativistic local density approximation. <b>1983</b> , 99, 172-174	9
901	Theoretical aspects of adsorption and heterogeneous catalysis. <b>1983</b> , 33, 639-649	22
900	The origin of Schottky barriers on the cleavage plane of III $\bar{V}$ semiconductors: Review of some recent theoretical work. <b>1983</b> , 104, 301-316	24
899	Bond contraction in monolayers: 2D-hexagonal Cs. <b>1983</b> , 134, L487-L493	
898	Structural properties of Si and GaP. <b>1983</b> , 46, 11-15	4
897	Variational calculation of the work function for small metal spheres. <b>1983</b> , 47, 845-849	49
896	Normal spin polarized photoemission from ferromagnetic Fe(100), Fe(110) and Fe(111). <b>1983</b> , 47, 29-31	
895	First-principles calculation of the insulator-to-metal transition pressure in CsI. <b>1983</b> , 47, 855-857	21
894	Calculation of the nonlinear susceptibilities of interacting homogeneous electron gas by the density functional method. <b>1983</b> , 48, 85-89	2
893	On the cohesive energy and charge density of uranium dioxide. <b>1983</b> , 45, 689-692	33
892	Nonlinear matrix solution of the Kohn-Hohenberg-Sham equation for electrons in a jellium slab. <b>1983</b> , 45, 125-128	3
891	Force sum-rule and static electronic polarizability of atoms and microstructures. <b>1983</b> , 48, 989-993	33
890	Density-functional theory of the compton profiles of beryllium metal. <b>1983</b> , 47, 673-676	19
889	Exchange and correlation potentials in the one-electron equation. <b>1983</b> , 47, 641-644	2
888	Lcao-G $\bar{p}$ $\bar{p}$ -Kohn-Sham energy bands of trans-polyacetylene-chain. <b>1983</b> , 48, 1085-1087	7

887	Self-interaction correction for density-functional theory of electronic energy bands of solids. <b>1983</b> , 28, 5992-6007	175
886	The electronic band structures for zincblende and wurtzite BeO. <b>1983</b> , 16, 3475-3480	63
885	Electronic structure of the ternary chalcopyrite semiconductors CuAlS <sub>2</sub> , CuGaS <sub>2</sub> , CuInS <sub>2</sub> , CuAlSe <sub>2</sub> , CuGaSe <sub>2</sub> , and CuInSe <sub>2</sub> . <b>1983</b> , 28, 5822-5847	451
884	From band tailing to impurity-band formation and discussion of localization in doped semiconductors: A multiple-scattering approach. <b>1983</b> , 28, 4704-4715	93
883	Metal-Metal Bonding in Cr-Cr and Mo-Mo Dimers: Another Success of Local Spin-Density Theory. <b>1983</b> , 50, 488-491	137
882	Local Spin-Density Description of Multiple Metal-Metal Bonding: Mo <sub>2</sub> and Cr <sub>2</sub> . <b>1983</b> , 50, 1451-1454	58
881	A new ab initio SCF pseudopotential energy band structure for KC <sub>8</sub> (abstract). <b>1983</b> , 8, 215	1
880	All-electron self-consistent determination of spin-polarization and knight shift of A Pt(001)film. <b>1983</b> , 31-34, 891-892	8
879	Magnetism of linear chains. <b>1983</b> , 38, 23-33	63
878	Magnetism at surfaces and interfaces. <b>1983</b> , 38, 269-272	18
877	Surface properties of alkali-metal alloys. <b>1983</b> , 16, 6883-6896	15
876	Experimental band structure of cadmium sulfide. <b>1983</b> , 28, 3306-3319	61
875	Electronic Structure and Cohesion in the Rare Earth Metals. <b>1983</b> , 213-254	8
874	Lattice relaxation around interstitial hydrogen in aluminium. <b>1983</b> , 13, 1379-1392	21
873	Bond contraction in monolayers: 2d-hexagonal Cs. <b>1983</b> , 134, L487-L493	44
872	Atomistic studies of helium trapping in metals. <b>1983</b> , 78, 25-36	14
871	Magnetism of amorphous metal-metal alloys. <b>1983</b> , 35, 192-198	98
870	First-principles calculation of the equilibrium ground-state properties of transition metals: Applications to Nb and Mo. <b>1983</b> , 28, 5480-5486	484

869	Will Diamond Transform under Megabar Pressures?. <b>1983</b> , 50, 2006-2009	215
868	All-electron local density functional study of metallic monolayers. I. Alkali metals. <b>1983</b> , 13, 2313-2321	95
867	Density-functional theory of correlations in dense plasmas: Improvement on the hypernetted-chain scheme. <b>1983</b> , 27, 3241-3250	65
866	Physical Content of the Exact Kohn-Sham Orbital Energies: Band Gaps and Derivative Discontinuities. <b>1983</b> , 51, 1884-1887	1795
865	Density-Functional Theory of the Energy Gap. <b>1983</b> , 51, 1888-1891	1434
864	Energy of hydrogen in b.c.c. transition metals. <b>1983</b> , 91, 209-215	9
863	Theory of the two-center bond. <b>1983</b> , 27, 3592-3604	220
862	Exchange integral matrices and cohesive energies of transition metal atoms. <b>1983</b> , 13, L197-L202	65
861	Comparison of local density functional Hartree-Fock and configuration interaction in the STO-3G orbital basis for some first-row diatomics. <b>1983</b> , 16, 1125-1131	3
860	Rigid muffin-tin model for interatomic forces in metals. <b>1983</b> , 13, 805-816	3
859	Pressure dependence of the spin susceptibility in vanadium. <b>1983</b> , 13, L123-L125	9
858	Self-consistent non-local pseudopotential calculations for the ground-state properties of $\text{Eu}$ . <b>1983</b> , 16, 1649-1657	4
857	Elastic scattering of slow electrons from Xe atoms. <b>1983</b> , 16, 603-612	28
856	Relativistic corrections to the atomic volumes of the actinide metals. <b>1983</b> , 13, 103-108	80
855	A density-functional approach to the electronic structure of dissolved impurities in simple liquid metals. <b>1983</b> , 13, 1645-1655	4
854	On the calculation of cohesive energy using Compton profile. <b>1983</b> , 16, L705-L709	1
853	Effects of screening on crystal-field magnetoelastic coupling in rare-earth intermetallic compounds. <b>1983</b> , 16, 4637-4653	11
852	Application of the time-dependent local density approximation to the photoionisation of mercury. <b>1983</b> , 16, L375-L379	18

851	Accuracy of the Hartree-Fock and Local Density Approximations for Electron Densities: A study for Light Atoms. <b>1983</b> , 28, 389-393	12
850	Elastic scattering of slow electrons from Ar and Kr atoms. <b>1983</b> , 16, 1819-1825	17
849	Self-interaction correction in the local density approximation. <b>1983</b> , 16, 349-357	9
848	Non-local approximations to the kinetic energy functional. <b>1983</b> , 16, 677-689	18
847	Static and Dynamical Effects of Core Holes in KLV Auger, SXE, and SXA Spectra of Simple Metals. <b>1983</b> , 28, 107-124	51
846	Influence of shear strains on the electronic structure of vanadium. <b>1983</b> , 13, 761-777	29
845	Electron-electron interaction in 3d metals which b.c.c. structure. <b>1983</b> , 13, 1801-1806	5
844	Calculation of Knight shifts for liquid binary alloys of simple metals: charge-transfer effects. <b>1983</b> , 13, 503-511	5
843	Evidence for magnetically induced electronic and phonon anomalies in cubic Pd <sub>2</sub> MnSn (Mossbauer study). <b>1983</b> , 13, 1547-1558	4
842	Self-energy contributions to the band structure of alkali metals: application to potassium. <b>1983</b> , 13, 1895-1914	10
841	Convergence of momentum space, pseudopotential calculations for Si. <b>1983</b> , 28, 7346-7348	23
840	Electric field gradient and electronic structure of linear-bonded halide compounds. <b>1983</b> , 28, 3697-3705	33
839	Electronic structure of Ni and Pd alloys. I. X-ray photoelectron spectroscopy of the valence bands. <b>1983</b> , 27, 2145-2178	295
838	Self-consistency iterations in electronic-structure calculations. <b>1983</b> , 28, 5462-5472	110
837	Calculation of the Compton profile of beryllium. <b>1983</b> , 28, 1696-1700	16
836	Microscopic analysis of interatomic forces in transition metals with lattice distortions. <b>1983</b> , 28, 6687-6694	113
835	Pseudopotentials for the 3d transition-metal elements. <b>1983</b> , 28, 535-543	33
834	Symmetric lattice distortions around deep-level impurities in semiconductors: Vacancy and substitutional Cu in silicon. <b>1983</b> , 28, 4510-4518	36

833	Electronic properties of alkali trimers. <b>1983</b> , 78, 5646-5655	172
832	Nonadjustable exchange-correlation model for electron scattering from closed-shell atoms and molecules. <b>1983</b> , 27, 1893-1903	187
831	Orbital-moments analysis of ionic radii in silver and copper halides. <b>1983</b> , 27, 5919-5923	20
830	Electronic excitations in semiconductors with doping superlattices. <b>1983</b> , 27, 3547-3553	37
829	Ab initio study of structural and electronic properties of beryllium. <b>1983</b> , 28, 4179-4185	117
828	Magnetism of the Ni(110) and Ni(100) surfaces: Local-spin-density-functional calculations using the thin-slab linearized augmented-plane-wave method. <b>1983</b> , 28, 610-623	115
827	Interaction of helium with a metal surface. <b>1983</b> , 27, 4612-4616	127
826	Convergence study of the density-gradient expansion for the screened-Coulomb exchange energy. <b>1983</b> , 28, 3014-3021	4
825	Stability in ordered and amorphous transition-metal compounds. <b>1983</b> , 27, 7194-7198	78
824	Electronic band structures for zinc-blende and wurtzite CdS. <b>1983</b> , 28, 4736-4743	79
823	Quantum Motion of Chemisorbed Hydrogen on Ni Surfaces. <b>1983</b> , 51, 1081-1084	145
822	Simultaneous Relaxation of Nuclear Geometries and Electric Charge Densities in Electronic Structure Theories. <b>1983</b> , 50, 1684-1688	87
821	v-Representability and Density Functional Theory. <b>1983</b> , 51, 1596-1598	66
820	A pseudoatom theory for the liquid-vapor interface of simple metals: Computer simulation studies of sodium and cesium. <b>1983</b> , 78, 5225-5249	93
819	All-electron local-density theory of alkali-metal bonding on transition-metal surfaces: Cs on W(001). <b>1983</b> , 28, 3074-3091	261
818	Structural phase stability in third-period simple metals. <b>1983</b> , 27, 3235-3251	188
817	Pair potentials on metal surfaces. <b>1983</b> , 28, 6749-6757	7
816	Total energies in Se. I. The trigonal crystal. <b>1983</b> , 27, 6296-6301	32

815	X <sub>2</sub> Cr <sub>2</sub> , and the symmetry dilemma. <b>1983</b> , 27, 2217-2219	33
814	Calculation of self-consistent potentials for substitutionally disordered systems with application to the Ag <sub>x</sub> Bd <sub>1-x</sub> alloy series. <b>1983</b> , 27, 882-904	141
813	Ground- and excited-state properties of solid argon under pressure. <b>1983</b> , 28, 6812-6820	26
812	Phonon effects in deep-level spectra of metals: Beyond the linear-screening and linear-coupling models. <b>1983</b> , 28, 7362-7365	1
811	Pseudopotential local-spin-density calculations for Si <sub>2</sub> . <b>1983</b> , 28, 1945-1950	24
810	Self-consistent semirelativistic pseudopotential calculation of the energy bands, cohesive energy, and bulk modulus of W. <b>1983</b> , 27, 3152-3159	43
809	Static electronic susceptibility, $\chi(q  ,z,z?)$ , of the Lang-Kohn jellium surface. <b>1983</b> , 27, 6542-6544	22
808	Simplified self-interaction correction applied to the energy bands of neon and sodium chloride. <b>1983</b> , 28, 2135-2139	41
807	Generalized non-muffin-tin band theory. <b>1983</b> , 27, 4564-4579	75
806	First-principles phonon spectra in Ca and Sr. <b>1983</b> , 28, 4818-4821	24
805	Structural determination of Cl chemisorption on Si{111} and Ge{111} by total-energy minimization. <b>1983</b> , 28, 2302-2304	55
804	Basis set effects on spectroscopic constants for C <sub>2</sub> and Si <sub>2</sub> and the symmetry dilemma in the X <sub>2</sub> model. <b>1983</b> , 78, 4997-5003	64
803	Gradient-free representation of the Weizsäcker term for atoms. <b>1983</b> , 28, 1808-1809	9
802	Total energy of isolated point defects in solids. <b>1983</b> , 28, 2296-2298	28
801	Comparative studies of electronic structures of sodium metasilicate and $\beta$ - and $\gamma$ -phases of sodium disilicate. <b>1983</b> , 28, 4724-4735	39
800	Phase transitions in a mercury monolayer. <b>1983</b> , 28, 593-597	40
799	Large Orbital-Moment Contribution to 5f Band Magnetism. <b>1983</b> , 51, 1708-1711	201
798	Dynamical Density Response Function of a Metal Film in the Random-Phase Approximation. <b>1983</b> , 51, 1907-1910	117

797	Localized states and the electronic properties of a hydrogenated defect in amorphous silicon. <b>1983</b> , 28, 3246-3257	28
796	Energy bands and magnetization of a Ni(001) monolayer. <b>1983</b> , 27, 2092-2100	25
795	Atomic geometry and surface-state spectrum for Ge(111)-(2 $\times$ 1). <b>1983</b> , 27, 6553-6556	119
794	Effective proton-proton potential in hydrogen plasmas. <b>1983</b> , 28, 344-349	29
793	Static structure factor and electron-electron correlations in Be. A comparison of experiment with electron-gas theory. <b>1983</b> , 28, 1772-1780	17
792	Geometry of density matrices. IV. The relationship between density matrices and densities. <b>1983</b> , 27, 632-645	60
791	Density-functional study of interplanar binding in graphite. <b>1983</b> , 27, 2458-2469	44
790	Geometry of the Ag{001}-c(2 $\times$ 2)Cl structure as determined by He diffraction. <b>1983</b> , 28, 494-503	45
789	Electronic structure of transition-atom impurities in semiconductors: Substitutional3dimpurities in silicon. <b>1983</b> , 27, 1191-1227	84
788	Electric field gradient of single impurities in aluminum. <b>1983</b> , 27, 5110-5112	5
787	General operator ground-state expectation values in the Hohenberg-Kohn-Sham density-functional formalism. <b>1983</b> , 27, 5912-5918	69
786	Energy profiles for light impurities in simple metals. <b>1983</b> , 27, 642-658	27
785	Augmented Gaussian-orbital basis for atomic-cluster calculations within the density-functional formalism: Application to Cu <sub>2</sub> . <b>1983</b> , 28, 5536-5548	43
784	Total energy of Al within the neutral-pseudoatom model. <b>1983</b> , 28, 3582-3584	4
783	Relationship between gradient expansion terms in the kinetic and exchange energy-density functionals in statistical models of an atom with shell structure. <b>1983</b> , 27, 1184-1186	7
782	Spin-density calculations for core-electron photoemission and Auger electron line shapes, x-ray-edge exponents, and solid-state shifts. <b>1983</b> , 28, 3182-3192	28
781	Relativistic spin polarization and magnetization: Knight shift of Pt(001). <b>1983</b> , 28, 6262-6269	59
780	Density-functional calculation of the static electronic polarizability of a small metal sphere. <b>1983</b> , 28, 5702-5710	93



779	Pressure and virial theorem in pseudopotential formalism. <b>1983</b> , 27, 7769-7771	14
778	Energy-band-structure study of the (100), (110), and (111) surfaces of TiC. <b>1983</b> , 27, 5934-5942	34
777	Density functional calculations for H <sub>2</sub> O, NH <sub>3</sub> , and CO <sub>2</sub> using localized muffin-tin orbitals. <b>1983</b> , 79, 1874-1884	50
776	Cu and Ag as one-valence-electron atoms: Pseudopotential results for Cu <sub>2</sub> , Ag <sub>2</sub> , CuH, AgH, and the corresponding cations. <b>1983</b> , 79, 5532-5542	121
775	Density functional calculations for low-lying states of CO <sub>2</sub> . <b>1983</b> , 79, 1885-1890	13
774	Band Structure of NbSe <sub>3</sub> . <b>1983</b> , 52, 578-586	34
773	A Single-Site Picture of Valence Fluctuation in Periodic Anderson Hamiltonian with Large Coulomb Correlation. <b>1983</b> , 52, 3886-3896	24
772	Band structure and optical properties of NiO. I. Band structure calculations. <b>1983</b> , 16, 6713-6721	19
771	A comparison of self-interaction-corrected local correlation energy functionals. <b>1983</b> , 16, 3687-3702	61
770	Electron gas models and density functional theory. <b>1983</b> , 229-258	1
769	Molecular field theory of nematics: density functional approach. I. Bulk effects. <b>1983</b> , 16, 1539-1553	71
768	Comparison of Local-Density and Quantal Hypernetted-Chain Approximations in the Calculation of Electron Density Distribution. <b>1983</b> , 70, 331-342	6
767	General Density Functional Theory. <b>1983</b> , 79-147	109
766	The density of conduction electrons at a cylindrical dilatation in a metal. <b>1983</b> , 48, 449-456	
765	Atomic and Molecular Orbital Electronegativity Models Based on the Transition State and Transition Operator Approaches. <b>1983</b> , 87, 925-932	8
764	Beyond the local-density approximation in calculations of ground-state electronic properties. <b>1983</b> , 28, 1809-1834	918
763	Cohesive Properties of Alkali Halides and Simple Oxides in the Local-Density Formalism. <b>1983</b> , 52, 3506-3513	54
762	Electronic Structure of Quasi-One-Dimensional Transition Metal Chalcogenide Nb <sub>3</sub> X <sub>4</sub> . <b>1983</b> , 52, 587-596	39

761	Band Structures and Charge Distributions along the c-Axis of Higher Stage Graphite Intercalation Compounds. <b>1983</b> , 52, 223-232	44
760	Band Theory of the Intermetallic Compound UGe <sub>3</sub> . <b>1984</b> , 53, 3929-3937	19
759	AN OVERVIEW OF DENSITY-FUNCTIONAL THEORY. <b>1984</b> , 3-49	10
758	Density Functional Theory for Solids. <b>1984</b> , 67-140	17
757	Self-interaction-correction theory for density functional calculations of electronic energy bands for the lithium chloride crystal. <b>1984</b> , 17, 1853-1866	46
756	Atomic-number-dependent pseudopotentials: general theory. <b>1984</b> , 17, 2705-2711	2
755	Electronic states of transition metal impurities in II-VI and III-V semiconductors. <b>1984</b> , 17, 2333-2356	55
754	The relativistic time-dependent local-density approximation. <b>1984</b> , 17, 531-540	24
753	The influence of electron-electron interaction on inner-shell excitation processes in heavy-ion collisions. <b>1984</b> , 17, 615-640	32
752	Charge densities, interionic potentials and phonon frequencies in Al. <b>1984</b> , 14, 2017-2025	7
751	A new local exchange potential for low-energy electron scattering by atoms based on first principles. <b>1984</b> , 17, 1637-1657	24
750	A crystalline cluster model of the electronic structure of copper chloride. <b>1984</b> , 17, 1695-1702	5
749	All-electron local density functional study of metallic monolayers. III. Transition metals Sc to Cu. <b>1984</b> , 14, 2613-2624	19
748	Non-linear screening of neon and argon impurities in dense, hot hydrogen plasmas. <b>1984</b> , 17, 3617-3626	5
747	Local field corrections for the dielectric function of an electron liquid. <b>1984</b> , 14, 1673-1683	8
746	Electronic structure of NaCl-type compounds of the light actinides. I. UN, UC, and UO. <b>1984</b> , 14, 639-652	85
745	Calculated cohesive energies of the light actinide metals. <b>1984</b> , 14, 1157-1171	36
744	Dipole polarisabilities of atoms: an X $\alpha$ density matrix study. <b>1984</b> , 17, 1463-1475	19

743	Non-local approximation for the exchange part of the density functional. <b>1984</b> , 17, 2107-2120	10
742	A study of the self-energy of quasiparticles in an electron liquid. <b>1984</b> , 17, 6685-6699	6
741	Exchange-correlation potential in semiconductors and insulators. <b>1984</b> , 17, 1497-1510	48
740	Electronic Damping of Atomic and Molecular Vibrations at Metal Surfaces. <b>1984</b> , 29, 360-371	152
739	The direct correlation function of inhomogeneous quantum liquids. <b>1984</b> , 17, 1633-1642	34
738	Forces in pseudopotential molecular calculations. <b>1984</b> , 80, 1525-1528	8
737	Angle-resolved photoemission study of the electronic structure of beryllium: Bulk band dispersions and many-electron effects. <b>1984</b> , 30, 5500-5507	47
736	Polarizability of 5s25p(P122) atomic indium. <b>1984</b> , 29, 2977-2980	16
735	Interstitial-sphere linear muffin-tin orbital structural calculations for C and Si. <b>1984</b> , 30, 5835-5841	101
734	Effect of exchange on shallow-impurity calculations. <b>1984</b> , 30, 4583-4585	1
733	Si-III (BC-8) crystal phase of Si and C: Structural properties, phase stabilities, and phase transitions. <b>1984</b> , 30, 1773-1776	190
732	Structural properties of NaCl. <b>1984</b> , 29, 3770-3772	95
731	Linear coupled-cluster method. II. Analysis of local exchange-correlation potentials in beryllium and its isoelectronic series. <b>1984</b> , 29, 58-63	4
730	Rigorous formulation of the Kohn and Sham theory. <b>1984</b> , 30, 2183-2186	48
729	Local-density approximation for dynamical correlation corrections to single-particle excitations in insulators. <b>1984</b> , 30, 4719-4733	100
728	Electron correlation in the Coulomb hole model. Comparison of methods. <b>1984</b> , 81, 4008-4013	15
727	Ab initio calculation of the electronic structure of thallos halides. II. One-particle states: Wannier functions, band structures, and densities of states. <b>1984</b> , 29, 2246-2254	11
726	Ab initio calculation of the electronic structure of thallos halides. I. Ground state: Static structural properties. <b>1984</b> , 29, 2238-2245	5

725	Total-energy full-potential linearized augmented-plane-wave method for bulk solids: Electronic and structural properties of tungsten. <b>1984</b> , 30, 561-569	647
724	Repulsive interaction of the helium atom with a metal surface. <b>1984</b> , 29, 2314-2316	113
723	Comment on "Electron removal energies in Kohn-Sham density-functional theory". <b>1984</b> , 30, 3523-3524	10
722	Static response of a jellium surface: The image potential and indirect interaction between two charges. <b>1984</b> , 30, 5449-5459	64
721	Efficient solution of Poisson's equation in linear combination of atomic orbitals (LCAO) electronic structure calculations. <b>1984</b> , 81, 5864-5872	21
720	Fock potentials. I. Separation of short- and long-range electronic interactions. <b>1984</b> , 29, 6-10	9
719	Parameter-free model of the correlation-polarization potential for electron-molecule collisions. <b>1984</b> , 29, 1742-1748	330
718	Self-consistent calculation of the polarizability of small jellium spheres. <b>1984</b> , 30, 6935-6942	138
717	Total energy and pressure in the Gaussian-orbitals technique. I. Methodology with application to the high-pressure equation of state of neon. <b>1984</b> , 29, 6425-6433	28
716	Total energy of the adatom and pyramidal-cluster models for Si(111). <b>1984</b> , 29, 1966-1969	41
715	Electronic structure and identification of deep defects in GaP. <b>1984</b> , 29, 3269-3282	40
714	Electron-positron enhancement factors for an electron gas of high density within the Kahana formalism. <b>1984</b> , 30, 2490-2497	23
713	Anisotropic electron-impurity scattering in dilute AlLi: (Small) corrections to the local-density approximation. <b>1984</b> , 30, 4796-4799	2
712	Generalized Wannier functions as a way to study the electron-phonon interaction in silicon. <b>1984</b> , 30, 2104-2111	1
711	Calculated electrical conductivity and thermopower of silver-palladium alloys. <b>1984</b> , 29, 4217-4223	79
710	Energetics of Surface Multilayer Relaxation on W (001): Evidence for Short-Range Screening. <b>1984</b> , 53, 675-678	88
709	Internal density functional theory of molecular systems. <b>1984</b> , 81, 2088-2102	16
708	Density-Functional Calculations for Ozone: Striking Results for an Important Molecule. <b>1984</b> , 52, 2002-2005	16

707	Cluster Study of the Interaction of a Water Molecule with an Aluminum Surface. <b>1984</b> , 53, 2493-2496	86
706	High-pressure behavior of MgO: Structural and electronic properties. <b>1984</b> , 30, 4774-4781	193
705	Magnetism at the Ni(001) surface: A high-precision, all-electron local-spin-density-functional study. <b>1984</b> , 30, 3113-3123	162
704	Local-density Hartree-Fock theory of electronic states of molecules with self-interaction correction. <b>1984</b> , 80, 1972-1975	183
703	Bonding of surface states on W(001): All-electron local-density-functional studies. <b>1984</b> , 29, 5267-5278	72
702	Interface magnetism in metals: Ag/Fe(001). <b>1984</b> , 30, 36-43	173
701	Fast method for calculating the self-consistent electronic structure of random alloys. <b>1984</b> , 30, 5508-5515	43
700	Graphitic spheres by pyrolyzing tolueneferrocenehiophene in a chemical vapor deposition experiment. <b>2023</b> , 58, 2170-2187	0
699	Adsorption properties of a paracyclophane molecule on NaCl/Au surfaces: a first-principles study.	0
698	Theoretical investigation on optical properties of Möbius carbon nanobelts in one- and two-photon absorption. <b>2023</b> , 18,	0
697	Magnetic Transition State Searching: Beyond the Static Ion Approximation. <b>2023</b> , 9, 42	1
696	Oxidation of metals and formation of defects by theoretical modeling. <b>2023</b> , 129-160	0
695	Triggering the mechanism of the initial reaction of energetic materials under pressure based on Raman intensity analysis.	0
694	Optoelectronic properties and lattice thermal conductivity of Cs <sub>2</sub> CuBiX <sub>6</sub> (X = F, Cl, Br, I) double perovskites: Thermodynamic and ab initio approaches. <b>2023</b> , e00791	0
693	Polarons and electrical leakage in BaZrO <sub>3</sub> and BaCeO <sub>3</sub> . <b>2023</b> , 7,	0
692	Improved calculations of mean ionization states with an average-atom model. <b>2023</b> , 5,	0
691	The aid of calorimetry for the thermochemical and kinetic study of the π-hole bonding leading to I <sub>2</sub> and 4-(dimethylamino) pyridine complexes in solution.	0
690	Comment on "Theoretical study of the NO <sub>3</sub> radical reaction with CH <sub>2</sub> ClBr, CH <sub>2</sub> ICl, CH <sub>2</sub> IBr, CHCl <sub>2</sub> Br, and CHClBr <sub>2</sub> " by I. Alkorta, J. M. C. Plane, J. Elguero, J. Z. Dávalos, A. U. Acuña and A. Saiz-Lopez, Phys. Chem. Chem. Phys. 2022, 24, 14365. <b>2023</b> , 25, 4355-4356	0

- 689 Structural, electronic and thermoelectric properties of LiAlX<sub>2</sub> (X=S and Se) chalcopyrites: promising for thermoelectric power generators. **2023**, 20, 73-83 ○
- 688 Enabling Room-Temperature Triferroic Coupling in Dual Transition-Metal Dichalcogenide Monolayers Via Electronic Asymmetry. **2023**, 145, 2485-2491 1
- 687 Ab initio vacancy formation energies and kinetics at metal surfaces under high electric field. **2023**, 107, ○
- 686 Mixed-domain Charge Transport in the S-Se System from First Principles. **2023**, ○
- 685 Passivation of Hematite by a Semiconducting Overlayer Reduces Charge Recombination: An Insight from Nonadiabatic Molecular Dynamics. **2023**, 14, 879-887 1
- 684 Tuning the Optical Properties of the Metal-organic Framework UiO-66 via Ligand Functionalisation. ○
- 683 Understanding the mechanism and regio- and stereo selectivity of [3 + 2] cycloaddition reactions between substituted azomethine ylide and 3,3,3- trifluoro -1-nitroprop-1-ene, within the molecular electron density theory. ○
- 682 A wide-range semiclassical self-consistent average atom model. **2023**, 30, 012711 ○
- 681 DFT study of cobalt based quaternary full-Heusler compound for spintronics and thermoelectric technologies. ○
- 680 Hardness and Mechanical Properties of Wurtzite B<sub>2</sub>X<sub>2</sub> Compounds. **2023**, 127, 2581-2588 ○
- 679 Interpolative Separable Density Fitting for Accelerating Two-Electron Integrals: A Theoretical Perspective. ○
- 678 Reparameterization of the Chemical-Potential Equalization Model with DFTB3: A Practical Balance Between Accuracy and Transferability. ○
- 677 The core ionization energies calculated by delta SCF and Slater's transition state theory. ○
- 676 Janus Ga<sub>2</sub>SeTe and In<sub>2</sub>SeTe nanosheets: Excellent photocatalysts for hydrogen production under neutral pH. **2023**, ○
- 675 Molecular structure and spectroscopic properties of two radicals of C<sub>4</sub>H<sub>2</sub>N: a DFT study. **2023**, 29, ○
- 674 Properties of carbon up to 10 million kelvin from Kohn-Sham density functional theory molecular dynamics. **2023**, 107, 1
- 673 Single-Atom Catalysis Enabled by High-Energy Metastable Structures. ○
- 672 The Origin of Amphipathic Nature of Short and Thin Pristine Carbon NanotubesFully Recyclable 1D Water-in-Oil Emulsion Stabilizers. 2202407 1

671	Tracking the Role of Defect Types in Co <sub>3</sub> O <sub>4</sub> Structural Evolution and Active Motifs during Oxygen Evolution Reaction. <b>2023</b> , 145, 2271-2281	3
670	Electron-Phonon effects and temperature-dependence of the electronic structure of monoclinic EGa <sub>2</sub> O <sub>3</sub> . <b>2023</b> , 11, 011106	0
669	Experimental and theoretical analysis for the structural, FT-IR, NLO, NBO and RDG properties of lindane using DFT technique. <b>2023</b> ,	0
668	Effect of Bi <sup>3+</sup> Doping on the Electronic Structure and Thermoelectric Properties of (Sr <sub>0.889-x</sub> La <sub>0.111</sub> Bi <sub>x</sub> )TiO <sub>2.963</sub> : First-Principles Calculations. <b>2023</b> , 13, 178	0
667	Third-Order Elastic Constants of Germanium and Silicon using Adiabatic-Connection Fluctuation-Dissipation Theorem in Random Phase Approximation.	1
666	First-principles calculations of plasmon excitations in graphene, silicene, and germanene. <b>2023</b> , 107,	0
665	An introduction to quantum chemistry. <b>2023</b> , 21-36	0
664	First-principles thermal equation of state of fcc iridium. <b>2023</b> , 107,	0
663	Engineering Chemo-Mechanical Properties of Zn Surfaces via Alucone Coating. <b>2023</b> , 127, 2481-2492	0
662	Machine Learning Identification of Active Sites in Graphite-Conjugated Catalysts. <b>2023</b> , 127, 2303-2313	0
661	Investigation of structural, elastic, and piezoelectric properties of NbSbO <sub>4</sub> crystal.	0
660	Phonon Dominated Thermal Transport in Metallic Niobium Diselenide from First Principles Calculations. <b>2023</b> , 13, 315	0
659	Effect of Defects in Graphene/Cu Composites on the Density of States. <b>2023</b> , 16, 962	0
658	First-principles calculations of Mg <sub>2</sub> FeH <sub>6</sub> under high pressures and hydrogen storage properties. <b>2023</b> , 29,	0
657	Topological Impact of Delocalization on the Stability and Band Gap of Partially Oxidized Graphene. <b>2023</b> , 8, 5124-5135	0
656	Metallic water: transient state under ultrafast electronic excitation.	0
655	The Role of M <sup>3+</sup> Substitutional Doping (M = In, Sb, Bi) in the Passivation of the $\sqrt{3}\times\sqrt{3}$ PbI <sub>3</sub> (100) Surface. <b>2023</b> , 127, 1713-1721	0
654	Phonon transport in Janus monolayer siblings: a comparison of 1T and 2H-ISbTe. <b>2023</b> , 13, 4202-4210	0

- 653 Novel Ultrahard Extended Hexagonal C10, C14 and C18 Allotropes with Mixed sp<sup>2</sup>/sp<sup>3</sup> Hybridizations: Crystal Chemistry and Ab Initio Investigations. **2023**, 9, 11 ○
- 652 Modeling phonons in nanomaterials. **2023**, 125-149 ○
- 651 First principles study on thermal conductivity of nitrogen substituted diamane. **2023**, 2431, 012057 ○
- 650 BN-Doped Carbon Nanotubes and Nanoribbons as Nonlinear-Optical Functional Materials for Application in Second-Order Nonlinear Optics. **2023**, 6, 1549-1561 ○
- 649 ??????????. **2023**, ○
- 648 Molecular Structure. **2023**, 487-506 ○
- 647 Synthesis, crystal structure, DFT, and photovoltaic studies of BaCeCuS<sub>3</sub>. ○
- 646 Revealing the electronic, optical and photocatalytic properties of PN-M<sub>2</sub>CO<sub>2</sub> (P = Al, Ga; M = Ti, Zr, Hf) heterostructures. ○
- 645 Physics-informed machine learning combining experiment and simulation for the design of neodymium-iron-boron permanent magnets with reduced critical-elements content. 9, ○
- 644 Thomas-Fermi and Other Density-Functional Theories. **2023**, 297-308 ○
- 643 Solids that are also liquids: elastic tensors of superionic materials. **2023**, 9, ○
- 642 Noncollinear density functional theory. **2023**, 5, ○
- 641 Ordered vacancy compounds: the case of the Mangli phases of TiO<sub>2</sub>. **2023**, 533-565 ○
- 640 Modeling metamaterials: Planar heterostructures based on graphene, silicene, and germanene. **2023**, 27-50 ○
- 639 Electronic structure and density functional theory. **2023**, 3-35 ○
- 638 Electronic and magnetic properties of TATA-DNA sequence driven by chemical functionalization. ○
- 637 Catching the Killer: Dynamic Disorder Design Rules for Small-Molecule Organic Semiconductors. 2213370 ○
- 636 Computational Design of an Ultra-Strong High-Entropy Alloy. **2023**, 43-50 ○



- 635 Structure and Characterization of  $K_2Na_3B_2P_3O_{13}$ , a New Nonlinear Optical Borophosphate with One-Dimensional Chain Structure and Short Ultraviolet Cutoff Edge. **2023**, 62, 2480-2488 ○
- 634 Strong anharmonicity and high thermoelectric performance of cubic thallium-based fluoride perovskites  $TlXF_3$  (X = Hg, Sn, Pb). ○
- 633 Graph-based Quantum Response Theory and Shadow Born-Oppenheimer Molecular Dynamics. 1
- 632 Effect of vacancy defects on the electronic and mechanical properties of two-dimensional  $MoSi_2N_4$ . **2023**, 13, 5307-5316 ○
- 631 Understanding the Effects of Amine and Morpholine Adsorption on Unglazed Earthenware Using Density Functional Theory. ○
- 630 Computational Molecular Design to Assist Modification of Single Walled Carbon Nanotubes with B, N, Al, Si, P, and S Dopant Atoms for  $Cl_2$  Gas Sensor Application. **2023**, 122264 ○
- 629 Bismuth and oxygen vacancies induce  $(2 \times 1)$  reconstructions in bismuth oxyhalide ( $BiOX$ , X = Cl, Br, I)  $(0\ 0\ 1)$  surfaces. **2023**, 618, 156583 ○
- 628 Elucidating l-tyrosine crystal phase transitions by Raman spectroscopy and ab initio calculations. **2023**, 176, 111234 ○
- 627 First principles calculations to investigate structural, electronic, mechanical, thermoelectric and optical properties of Bi- and Se-doped SnTe. **2023**, 176, 111232 ○
- 626 Solvatochromism and solution  $\pi$ -stacking of N-(4-pyridyl)-1,8-naphthalimide and its corresponding triruthenium coordination complex. **2023**, 292, 122420 ○
- 625 A novel series of thiosemicarbazone hybrid scaffolds: Design, synthesis, DFT studies, metabolic enzyme inhibition properties, and molecular docking calculations. **2023**, 1280, 135077 ○
- 624 A DFT study of hydrogen adsorption on Pt modified carbon nanocone structures: Effects of modification and inclination of angles. **2023**, ○
- 623 Feature-rich electronic and magnetic properties in silicene monolayer induced by nitrogenation: A first-principles study. **2023**, 568, 111844 ○
- 622 Ab-initio simulation of the structural, electronic and optical properties for the vacancy-ordered double perovskites  $ATl$  (A = Cs or NH); a time-dependent density functional theory study. **2023**, 176, 111262 ○
- 621 First-principles approach to the structural, physical, electronic, magnetic and optical properties of honeycomb ordered antimonates  $Na_3Fe_2SbO_6$ . **2023**, 176, 111258 ○
- 620 Stability, electronic, optical and thermoelectric properties of site-substituted  $LaV O_3$ . **2023**, 35, 105510 ○
- 619 Electronic Effect on Phenoxide Migration at a Nickel(II) Center Supported by a Tridentate Bis(phosphinophenyl)phosphido Ligand. ○
- 618 Phenolic compounds extraction from propolis using imidazole based ionic liquids: a theoretical and experimental study. ○

- 617 Efficient CO<sub>2</sub> utilization and sustainable energy conversion via aqueous Zn-CO<sub>2</sub> batteries. **2023**, 109, 108242 ○
- 616 First-principles study of different oxidation process on Al(111) and Cu(111): Metal pulled-off effect. **2023**, 731, 122260 ○
- 615 Ab Initio Simulation of Dielectric and Optical Properties of Ices Ih and III and Lattice Frameworks of Hydrates sl and sH. **2022**, 64, 576-586 ○
- 614 2.5 Million-Atom Ab Initio Electronic-Structure Simulation of Complex Metallic Heterostructures with DGDFE. **2022**, ○
- 613 Exchange-Correlation Excitons in Plasma Media: A Monte Carlo Method and the Sign Problem. **2022**, 60, S315-S324 ○
- 612 Searching for d0 spintronic materials: bismuthene monolayer doped with IVA-group atoms. **2023**, 13, 5885-5892 ○
- 611 Modulation of chiral spectral deflection by van der Waals force-induced molecular electropolarization in catenane oligomers. **2023**, 13, 11055-11061 ○
- 610 Efficient photo-Fenton catalysis using magnetic iron nanoparticles decorated boron nitride quantum dots: theoretical and experimental investigations. **2023**, 13, 6779-6792 ○
- 609 Effect of hydrostatic pressure on structural, mechanical, and electronic properties of energetic molecular perovskite (C<sub>6</sub>H<sub>14</sub>N<sub>2</sub>)(NH<sub>2</sub>NH<sub>3</sub>)(ClO<sub>4</sub>)<sub>3</sub>: A DFT-D insight. ○
- 608 Fluorescent sensing of non-steroidal anti-inflammatory drugs naproxen and ketoprofen by dansylated squaramide-based receptors. **2023**, 21, 2968-2975 ○
- 607 Combined experimental and DFT approach to BiNbO<sub>4</sub> polymorphs. **2023**, 13, 5576-5589 ○
- 606 AIO<sub>3</sub>·H<sub>6</sub>TeO<sub>6</sub> (A = NH<sub>4</sub><sup>+</sup>, Rb<sup>+</sup>): Two telluric acid and iodate Co-crystalline compounds with second harmonic generation response. **2023**, 42, 100025 ○
- 605 Metal-to-Insulating Transition in the Perovskite System YSr<sub>2</sub>Cu<sub>2</sub>FeO<sub>8</sub>(0 &lt; &lt; 1) Modeled by DFT Methods. **2023**, 62, 3445-3456 ○
- 604 Advanced Design of Metal Nanoclusters and Single Atoms Embedded in C 1 N 1 -Derived Carbon Materials for ORR, HER, and OER. 2300405 1
- 603 Comparison of three cyclodextrins to optimize bisphenol A extraction from source water: Computational, spectroscopic, and analytical studies. 2300012 ○
- 602 Investigation of Structural, Mechanical, Optoelectronic, and Thermoelectric Properties of BaXF<sub>3</sub> (X = Co, Ir) Fluoro-Perovskites: Promising Materials for Optoelectronic and Thermoelectric Applications. **2023**, 8, 5274-5284 ○
- 601 Optimizing Vanadium Redox Reaction in Na<sub>3</sub>V<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub> Cathodes for Sodium-Ion Batteries by the Synergistic Effect of Additional Electrons from Heteroatoms. ○
- 600 Electronic properties and hydrogen evolution reaction performance of silicene with precious metal-doped nonmetal ring: first-principles calculations. **2023**, 58, 4487-4498 ○

- 599 Lattice Distorted Rhodium Nanocrystals in Porous Nanofiber toward Aqueous Zinc-CO<sub>2</sub> System. **2023**, 5, 1271-1280 ○
- 598 Graphynes and Graphdiynes for Energy Storage and Catalytic Utilization: Theoretical Insights into Recent Advances. ○
- 597 Oxygenated Triazine-Heptazine Heterostructure Creates an Enormous Ascension to the Visible Light Photocatalytic Hydrogen Evolution Performance of Porous C<sub>3</sub>N<sub>4</sub> Nanosheets. 2301579 ○
- 596 Atomically thin metallic Si and Ge allotropes with high Fermi velocities. **2023**, 107, ○
- 595 Time-dependent exchange-correlation hole and potential of the electron gas. **2023**, 107, ○
- 594 Inter- vs. Intra-Molecular Hydrogen Bond in Complexes of Nitrophthalic Acids with Pyridine. **2023**, 24, 5248 ○
- 593 Analysis of Bonding by Quantum Chemistry-Resolving Delocalization Stabilization in a Mechanistic Basis and New Hückel Model. ○
- 592 Hydrogen adsorption on Au (111), U (110), and nAu/U (110) alloy surfaces: A first-principles study. **2023**, 577, 154331 ○
- 591 Modeling the Photo-Absorption Properties of Noble Metal Nanoclusters: A Challenge for Density-Functional Theory. ○
- 590 Vibrational Spectra of HNIW and its Isotopologues: A Combined Experimental and Computational Study. **2023**, 48, ○
- 589 First-principles prediction of superconductivity in MgB<sub>3</sub>C<sub>3</sub>. **2023**, 107, ○
- 588 Heteroleptic and Homoleptic Iron(III) Complexes with a Tris(N-Heterocyclic Carbene) Borate Ligand: Synthesis, Characterization, and Catalytic Application. ○
- 587 Step Structures and Adatom Diffusion on SiC Surfaces. **2023**, 66, 239-244 ○
- 586 Electronic Structure and Photocatalytic Water Splitting Application of a New Type II g-ZnO/Ga<sub>2</sub>SSe van der waals Heterostructure. ○
- 585 An ab initio study of novel quaternary Heusler alloys for spin polarized and waste heat recycling systems. **2023**, 571, 170541 ○
- 584 First-Principles Studies on the Physical Properties of the Half Heusler RbNbCd and RbNbZn Compounds: A Promising Material for Thermoelectric Applications. **2023**, 13, 618 ○
- 583 Spectroscopic study on size-dependent optoelectronics of N-type ultra-high conductive polymer PBFDO. **2023**, 122744 ○
- 582 Application of molecular dynamics simulation for exploring the roles of plant biomolecules in promoting environmental health. **2023**, 869, 161871 ○

- 581 Vacancy Assisted Bilayer Graphene Contact for Monolayer Graphene Channel Devices. **2023**, 44, 666-669 ○
- 580 Metal-Organic Framework with a Redox-Active Bridge Enables Electrochemically Highly Selective Removal of Arsenic from Water. ○
- 579 How Regiochemistry Influences Aggregation Behavior and Charge Transport in Conjugated Organosulfur Polymer Cathodes for Lithium-Sulfur Batteries. ○
- 578 Understanding formation of the InPd<sub>3</sub> polymorphs: a DFT study. **2023**, ○
- 577 New Strategy to Improve Photocatalytic Activity and Mechanistic Aspect for Water Splitting. ○
- 576 Investigation of the electronic, optical, elastic, mechanical and vibrational properties of CuI using HSE03. ○
- 575 Mechanism of ring-opening polymerization of L-lactide by lanthanide aryloxide: A theoretical study on the effect of the aryloxide ligands on the process. ○
- 574 Effect of Surface Termination on Carrier Dynamics of Metal Halide Perovskites: Ab Initio Quantum Dynamics Study. ○
- 573 Thematic Exordium for Special Issue Density Functional Theory Application on Chemical Calculation **2023**, 16, 2904 ○
- 572 Towards a better understanding of commonly used medicinal plants from Turkiye: Detailed phytochemical screening and biological activity studies of two Teucrium L. species with in vitro and in silico approach. **2023**, 116482 ○
- 571 Role of spin-orbit coupling on the physical properties of APb<sub>3</sub> (A = Na, Ca, Y, and Th) superconductors. **2023**, 608, 1354250 ○
- 570 Gas sensing selectivity of SnO<sub>2</sub>-xNiO sensors for homogeneous gases and its selectivity mechanism: Experimental and theoretical studies. **2023**, 354, 114273 ○
- 569 Performance of an anticonvulsant drug, expired gabapentin, on zinc corrosion in an acidic environment. ○
- 568 Chemo- and regioselectivities of the TBAF-catalyzed C-F bond allylation of trifluoromethylalkenes: A theoretical view. **2023**, 542, 113111 ○
- 567 Theoretical study of the mechanism of the hydrogen evolution reaction on the V<sub>2</sub>C MXene: Thermodynamic and kinetic aspects. **2023**, 421, 252-263 ○
- 566 Al<sup>3+</sup> ions co-doped Ba<sub>2</sub>YSbO<sub>6</sub>: Mn<sup>4+</sup> phosphors with high thermal stability and strong far-red emission for plant growth LEDs. **2023**, 320, 123854 ○
- 565 Structural, spectroscopic, electronic, Hirshfeld, QTAIM and biological predications of a hybrid 2,6-dichloropurine compound: A detailed density functional theoretical study. **2023**, 100, 100984 ○
- 564 Evaluating residual stresses in compositionally graded TiN films via ab initio and Rietveld simulation. **2023**, 28, 101715 ○

- 563 A 4-aminophthalimide derive smart molecule for sequential detection of aluminum ions and picric acid. **2023**, 439, 114593 1
- 562 Mechanism of sulfur poisoning to Ru-based catalysts in supercritical water gasification of glycerol: From experiment to combined DFT and kinetics studies. **2023**, 464, 142622 0
- 561 Determination of the relative orientation between  $^{15}\text{N}$ - $^1\text{H}$  dipolar coupling and  $^1\text{H}$  chemical shift anisotropy tensors under fast MAS solid-state NMR. **2023**, 350, 107428 0
- 560 First-principles investigation of the Metal-insulator transitions in Ti-substituted tetragonal rutile  $\text{VO}_2$ . **2023**, 211, 111922 0
- 559 Oxidation susceptibility of UN  $\bar{B}$ (210) grain boundary with Al/Cr/Ni dopant: A first-principles study. **2023**, 578, 154344 0
- 558 Negative electron affinity driven topological spin-polarized electrons in cesium adsorbed H-GaBi surface. **2023**, 151, 110577 0
- 557 Metal-oxoacid-mediated oxyhydroxide with proton acceptor to break adsorption energy scaling relation for efficient oxygen evolution. **2023**, 80, 594-602 0
- 556 Assessing the high concentration of vacancies in refractory high entropy alloys. **2023**, 28, 101764 0
- 555 Fluorine substitution and pre-sodiation strategies to boost energy density of V-based NASICON-structured SIBs: Combined theoretical and experimental study. **2023**, 463, 142464 0
- 554 Atomic-scale friction of black phosphorus/degraded Cu substrate: A route to robust superlubricity obtained by the critical load. **2023**, 619, 156749 0
- 553 Dependence of predicted bulk properties of hexagonal hydroxyapatite on exchange-correlation functional. **2023**, 224, 112153 0
- 552 Theoretical investigation into activation of hydroperoxides by excited quinones under ultraviolet irradiation. **2023**, 463, 142423 0
- 551 A Kohn-Sham scheme based neural network for nuclear systems. **2023**, 840, 137870 0
- 550 Compatibility of DFT+U with non-collinear magnetism and spin-orbit coupling within a framework of numerical atomic orbitals. **2023**, 286, 108684 0
- 549 Structural phase stability and thermodynamical properties of transition metal complex hydrides  $\text{Na}_2\text{MgTMH}_7$  (TM=Sc/Ti) for hydrogen storage applications. **2023**, 321, 123867 0
- 548 An exceptionally strong, ductile and impurity-tolerant austenitic stainless steel prepared by laser additive manufacturing. **2023**, 250, 118868 0
- 547 Visible light responsive  $\text{MoS}_2/\text{Ag}@\text{WO}_3/\text{EG}$  photoanode with highly stable Z-scheme induced circular electron motion pioneered by Exfoliated graphite for bisphenol a photoelectrodegradation. **2023**, 464, 142462 0
- 546 Insights into the adsorption and interfacial products improving the wetting of the Ag-Ti/graphite and Cu-Ti/graphite systems: A first-principles calculation. **2023**, 38, 102840 0

- 545 On the enhanced performance of Pt-based high-entropy alloys catalyst during water-gas shift reaction: A density functional theory study. **2023**, 623, 157023 ○
- 544 Investigation of electronic and optical properties of the ternary chalcogenides for optoelectronic applications: A TB-mBJ DFT study. **2023**, 49, 151-157 ○
- 543 Electronic band structure and optical properties of Li<sub>2</sub>In<sub>2</sub>GeSe<sub>6</sub> crystal. **2023**, 35, 105798 ○
- 542 Spectroscopic characterization of a mononuclear oxovanadium (IV) Schiff base complex. Oxidation catalysis applications and antibacterial activities. **2023**, 1281, 135131 ○
- 541 Predicting formation of chemically graded metal/ceramic interfaces. **2023**, 224, 112155 ○
- 540 Machine learning insight into h-BN growth on Pt(111) from atomic states. **2023**, 621, 156893 ○
- 539 Tensile strain as an efficient way to tune transport properties of Graphdiyne/Borophene hetero-bilayers; a first principle investigation. **2023**, 224, 112161 ○
- 538 Analysis of the electronic structure of SiO<sub>2</sub> intrinsic defects based on Density Functional Theory. **2023**, 35, 105554 ○
- 537 Dehydroxylation of smectites from Nudged Elastic Band and Born-Oppenheimer Molecular Dynamics simulations. **2023**, 238, 106880 ○
- 536 Effect of four-phonon scattering on anisotropic thermal transport in bulk hexagonal boron nitride by machine learning interatomic potential. **2023**, 207, 124011 ○
- 535 A comparative study of cubic methylammonium lead iodide (CH<sub>3</sub>NH<sub>3</sub>PbI<sub>3</sub>) perovskite by using density functional theory. **2023**, 35, 105814 ○
- 534 Alkali cation controlled tuning of the metal-insulator transition in vanadium hollandites. **2023**, 292, 116408 ○
- 533 Strain effects on the electronic, magnetic and optical properties of CrI<sub>3</sub>: An ab-initio investigation. **2023**, 292, 116409 ○
- 532 Ab initio molecular dynamics benchmarking study of machine-learned potential energy surfaces for the HBr + HCl reaction. **2023**, 11, 100257 ○
- 531 First-principles calculations to investigate electronic structure and optical spectra of CdxZn1-xS ternary semiconductor alloys. **2023**, 6, 100177 ○
- 530 First-principles prediction of two-dimensional Janus XMnZ<sub>2</sub> (X = Cl, Br, I; M =. **2023**, 623, 157020 ○
- 529 First-principles calculations integrated with experimental optical and electronic properties for MoS<sub>2</sub>-graphene heterostructures and MoS<sub>2</sub>-graphene-Au heterointerfaces. **2023**, 623, 156948 ○
- 528 Na-modified carbon nitride as a leach-resistant and cost-effective solid base catalyst for biodiesel production. **2023**, 341, 127548 ○

- 527 A novel smart framework for sustainable nanocomposite electrolytes based on ionic liquids of dye-sensitized solar cells by a covalently multifunctional graphene oxide-vinyl imidazole/4-tert-butylpyridine cobalt complex. **2023**, 945, 169241 ○
- 526 Assimilation of electronic, elastic, mechanical, optical, and thermal profiles in metal halide perovskite CsPbCl<sub>3</sub>, for optoelectronic applications. **2023**, 35, e00804 ○
- 525 DFT and DFPT calculations of the structural, electronic, optical, vibrational and thermodynamic properties of silicon tetraborate. **2023**, 35, e00796 ○
- 524 First principles insight into physical properties of CaX<sub>2</sub>O<sub>4</sub> (X = In, Gd) spinels for optical and spintronic applications. **2023**, 322, 123999 ○
- 523 Structure, stability, and properties of cyclo[18]carbon-Zinc super sandwich complexes (C<sub>18</sub>-Zn-C<sub>18</sub>). **2023**, 991, 122668 ○
- 522 Nature and coordination geometry of geologically relevant aqueous Uranium(VI) complexes up to 400 °C: A review and new data. **2023**, 452, 131309 ○
- 521 Exact solutions of nonlinear dynamical equations for large-amplitude atomic vibrations in arbitrary monoatomic chains with fixed ends. **2023**, 120, 107176 ○
- 520 Process and mechanism of enhanced HCl leaching of platinum group metals from waste three-way catalysts by Li<sub>2</sub>CO<sub>3</sub> calcination pretreatment. **2023**, 452, 131348 ○
- 519 First-principles calculations: Structural stability, electronic structure, optical properties and thermodynamic properties of AlBN<sub>2</sub>, Al<sub>3</sub>BN<sub>4</sub> and Al<sub>3</sub>N<sub>4</sub> nitrides. **2023**, 160, 107400 ○
- 518 New insights into the reactivity of the triscyclopentadienyl monothiolate uranium(IV) complexes: CS<sub>2</sub> and CO<sub>2</sub> insertion and redox properties. A DFT theoretical approach. **2023**, 992, 122692 ○
- 517 Short bandgap of porphyrin molecules (Py) filled in a semiconducting single-walled carbon nanotube (Py@NT<sub>17</sub>) for highly efficient organic photovoltaic cells. **2023**, 293, 116456 ○
- 516 Metalloids (B, Si) and non-metal (N, P, S) doped graphene nanosheet as a supercapacitor electrode: A density functional theory study. **2023**, 35, 105905 ○
- 515 Prediction of novel final phases in aged uranium-niobium alloys. **2023**, 579, 154394 ○
- 514 First-principles study of Xe behavior in U<sub>2</sub>Zr<sub>2</sub>. **2023**, 579, 154387 ○
- 513 Efficient biobased carboxylic acids synthesis by synergistic electrocatalysis of multi-active sites on bimetallic Cu-Co oxide/oxyhydroxide. **2023**, 331, 122709 ○
- 512 A revisit to the role of Mo in an MP35N superalloy: An experimental and theoretical study. **2023**, 157, 60-70 ○
- 511 Tuned electronic band structure and intensified phonon scattering of Ge<sub>2</sub>Sb<sub>2</sub>Te<sub>5</sub> by strain engineering for thermoelectric performance. **2023**, 35, 105839 ○
- 510 Effect of Mn:Mg ratio on sintering behavior and microwave dielectric properties of (Mn, Mg)V<sub>2</sub>O<sub>6</sub> ceramics at ultra-low sintering temperature. **2023**, 43, 4060-4065 ○

- 509 Unusual mechanical properties of CO<sub>2</sub>V: Auxetic potential in a high-pressure polymorph of carbon dioxide. **2023**, 178, 111349 ○
- 508 Vanadium tuning amorphous iron phosphate encapsulated iron phosphide on phosphorous-doped graphene promoted oxygen reactions for flexible zinc air batteries. **2023**, 331, 122674 ○
- 507 Exploring particular electronic and optical properties of Tl<sub>2</sub>HgSnSe<sub>4</sub>, promising chalcogenide for solar photovoltaics and optoelectronics: A complex experimental and theoretical study. **2023**, 952, 170093 ○
- 506 Bandgap tuning and analysis of the electronic structure of the Cu<sub>2</sub>NiXS<sub>4</sub> (X=Sn, Ge, Si) system: mBJ accuracy with DFT expense. **2023**, 1, 100001 ○
- 505 Heterocyclic boronates as colorimetric and chemical sensors for dialkyltin derivatives. **2023**, 1285, 135515 ○
- 504 First-principles prediction of electronic, optical, and thermodynamic properties of c-TiAl<sub>3</sub> with M doping (M = V, Nb, Ta). ○
- 503 Marangoni Flow Driven via Hole Structure of Soluble Acene Polymer Blends for Selective Nitrogen Dioxide Sensing. ○
- 502 3D microstructure evolution in Na<sub>x</sub>FePO<sub>4</sub> storage particles for sodium-ion batteries. **2023**, 565, 232902 ○
- 501 Switching Rashba spin-splitting by reversing electric-field direction. **2023**, 7, ○
- 500 Density Functional Theory Studies of the Direct Conversion of Methane to Methanol Using O<sub>2</sub> on Graphitic MN<sub>4</sub>G-BN (M = Fe, Co, Cu) and CuN<sub>4</sub>G-PN Single-Atom Catalysts. ○
- 499 Molecular modeling and solubility of olopatadine hydrochloride polymorphs. **2023**, 1224, 114110 ○
- 498 Highly furosemide uptake employing magnetic graphene oxide: DFT modeling combined to experimental approach. **2023**, 379, 121652 ○
- 497 Conformational control of morphology for perylene diimide dimer as electron transporting material at perovskite surface. **2023**, 441, 114705 ○
- 496 Improved activity and significant SO<sub>2</sub> tolerance of SbPdV oxides on N-doped TiO<sub>2</sub> for CB/NO synergistic degradation. **2023**, 329, 138613 ○
- 495 Ba<sub>3</sub>GeTeS<sub>4</sub>: A new quaternary heteroanionic chalcogenide semiconductor. **2023**, 323, 124028 ○
- 494 First-principles calculations investigation on different coverage of H<sub>2</sub>O adsorption on the Mg-montmorillonite (0 1 0) edge surface. **2023**, 626, 157232 ○
- 493 CeO<sub>2</sub> promotes electrocatalytic formic acid oxidation of Pd-based alloys. **2023**, 948, 169665 ○
- 492 Insights into structural, elastic, mechanical, opto-electronic, and thermoelectric properties of rubidium-based fluoroperovskites RbXF<sub>3</sub> (X = Zn, Cd, Hg). **2023**, 178, 111357 ○



- 491 Plasma-assisted in-situ engineering carambola-like indium-doped Ni-Co selenides for robust hydrogen evolution. **2023**, 625, 157198 ○
- 490 Improving strong scalability of electronic structure simulations with reduced overhead of communications. **2023**, 288, 108747 ○
- 489 Growth of BiSb on GaAs (001) and (111)A surfaces: A joint experimental and theoretical study. **2023**, 622, 156688 ○
- 488 Investigation on non-radioactive behavior of an acylhydrazone-based fluorescent probe: Coexistence of PET and TICT mechanisms. **2023**, 295, 122603 ○
- 487 A new carbon allotrope: Biphenylene as promising anode materials for Li-ion and Li O<sub>2</sub> batteries. **2023**, 395, 116214 ○
- 486 Similarities and trends in adsorbate induced reconstruction structure and stability of FCC iron and cobalt surface carbides. **2023**, 626, 157245 ○
- 485 Decreasing the W-Cr solid solution decomposition rate: Theory, modelling and experimental verification. **2023**, 576, 154288 ○
- 484 Kinetically versus thermodynamically controlled factors governing elementary pathways of GaP(111) surface oxidation. **2023**, 560, 232663 ○
- 483 Electronic structure of covalent networks of triangular graphene flakes embedded in hBN. **2023**, 62, 025001 ○
- 482 Dynamic investigations on hydrogenHelium interaction around the vacancy in BCC iron from ab-initio calculations. **2023**, 63, 046005 ○
- 481 Computational studies for boosting nitrate electroreduction activity of Fe-N<sub>4</sub>-C Single-Atom catalyst via axial fifth ligand. **2023**, 616, 156440 ○
- 480 Thickness-dependent oxygen chemisorption behaviors on (1 1 1) surfaces of two-dimensional FCC metals Al and Cu: First-principles study. **2023**, 219, 112022 ○
- 479 The effect of the spin-orbit coupling on the physical properties of ScGa<sub>3</sub> and LuGa<sub>3</sub> superconductors. **2023**, 210, 111864 ○
- 478 Density functional theory study of bulk properties of transition metal nitrides. **2023**, 25, 5156-5163 ○
- 477 Theoretical analysis and experimental preparation of nanodiamond yttrium vacancy color center. **2023**, 98, 035801 ○
- 476 First-principles calculations of hematite (Fe<sub>2</sub>O<sub>3</sub>) by self-consistent DFT+U+V. **2023**, 26, 106033 ○
- 475 Anisotropic elastic properties of triclinic 2D materials using density functional theory with application to rhenium disulfide. **2023**, 34, e00790 ○
- 474 Strong anharmonic phonon scattering and superior thermoelectric properties of Li<sub>2</sub>NaBi. **2023**, 31, 100990 ○

- 473 Coexistence of Bulk-Nodal and Surface-Nodeless Cooper Pairings in a Superconducting Dirac Semimetal. **2023**, 130, ○
- 472 Pressure-dependent semiconductor-metal transition and elastic, electronic, optical, and thermophysical properties of orthorhombic SnS binary chalcogenide. **2023**, 45, 106236 ○
- 471 Stochastic algorithms for self-consistent calculations of electronic structures. **2023**, 92, 1693-1728 ○
- 470 The effect of Cr-substitution on the structural, electronic and magnetic properties of rutile VO<sub>2</sub>. **2023**, 569, 170449 ○
- 469 Fragmentation route of doubly ionized benzene, aniline, and nitroanilines monomers using a novel protocol from density functional theory and QTAIM. **2023**, 29, ○
- 468 In-situ formation of onion-like carbon film by tribo-induced catalytic degradation of hydrocarbon: Effect of lubrication condition and load. **2023**, 459, 141566 ○
- 467 A coupled quantum-molecular mechanics approach for performance analysis of defective Silicon based photovoltaic solar cells. **2023**, 98, 035007 ○
- 466 First principles study on structural, vibrational, electronic and elastic properties of 2D alkaline-earth carbides as a metallic material. **2023**, 293, 117281 ○
- 465 GW+EDMFT investigation of Pr<sub>1-x</sub>Sr<sub>x</sub>NiO. **2023**, 107, ○
- 464 Band offsets at the interfaces between TiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub>. **2023**, 7, ○
- 463 Recent advances in the ab initio theory of solid-state defect qubits. **2023**, 12, 359-397 1
- 462 Structural, elastic, and electronic properties of BC12 carbon under pressure. **2023**, 568, 111838 ○
- 461 Nickel ion removal system using activated carbon: A theoretical investigation. **2023**, 1221, 114055 ○
- 460 Recent progress of theoretical research on inorganic solid state electrolytes for Li metal batteries. **2023**, 561, 232720 1
- 459 A feedforward unitary equivariant neural network. **2023**, 161, 154-164 ○
- 458 Novel ultrahard sp<sup>2</sup>/sp<sup>3</sup> hybrid carbon allotrope from crystal chemistry and first principles: Body-centered tetragonal C<sub>6</sub> (Beoglitte) **2023**, 133, 109747 ○
- 457 Ab-initio investigation of Structural, Opto-Electronic and Thermodynamic properties of ZnAl<sub>2</sub>Se<sub>4</sub> for photovoltaic applications. ○
- 456 Optimization study of pharmaceuticals pollutants adsorption onto large surface area walnut shells activated carbon: Experimental design, Mechanism and DFT calculations. ○

- 455 Effects of chemical complexity on the initial oxidation resistance of HfC<sub>1-x</sub> ceramics. **2023**, 220, 112037 ○
- 454 Hydroxide Diffusion in Functionalized Cylindrical Nanopores as Idealized Models of Anion Exchange Membrane Environments: An Ab Initio Molecular Dynamics Study. **2023**, 127, 2792-2804 ○
- 453 Chiral Magnetic Interactions in Small Fe Clusters Triggered by Symmetry-Breaking Adatoms. **2023**, 15, 397 ○
- 452 (ZnO)<sub>42</sub> nanocluster: a novel visibly active magic quantum dot under first principle investigation. **2023**, 142, ○
- 451 Emerging d<sub>xy</sub> orbital coupling between non-d-block main-group elements Mg and I at high pressure. **2023**, 26, 106113 ○
- 450 2D janus niobium oxydihalide NbOXY: Multifunctional piezoelectric semiconductor for electronics, photonics, sensing and sustainable energy applications. **2023**, 31, 101001 ○
- 449 Quantum Mechanics of Electrons in Crystals. **2023**, 219-255 ○
- 448 Use of an Ammonium Quaternary Salt Obtained from Avocado as Corrosion Inhibitor of a Duplex Stainless Steel in CO<sub>2</sub> Saturated NaCl Solution. ○
- 447 Non-empirical Mixing Coefficient for Hybrid XC Functionals from Analysis of the XC Kernel. **2023**, 14, 1326-1333 1
- 446 Designing All-Solid-State Batteries by Theoretical Computation: A Review. **2023**, 6, ○
- 445 Effects of dispersion corrections on the theoretical description of bulk metals. **2023**, 107, ○
- 444 First-principles prediction of configurational order-disorder phase transition in Mo<sub>2</sub>(Ti<sub>1-x</sub>V<sub>x</sub>)C<sub>2</sub> MXene alloys. **2023**, 2431, 012041 ○
- 443 Pyridinecarboxaldehydes: Structures, Vibrational Assignments and Molecular Characteristics Using Experimental and Theoretical Methods. **2023**, 53, ○
- 442 A conceptual DFT and information-theoretic approach towards QSPR modeling in polychlorobiphenyls. **2023**, 61, 1143-1164 ○
- 441 Optical and dielectric response of two-dimensional WX<sub>2</sub> (X = Cl, O, S, Se, Te) monolayers: A comprehensive study based on density functional theory. ○
- 440 Orthorhombic Na<sub>2/3</sub>Cu<sub>0.1</sub>Mn<sub>0.9</sub>O<sub>2</sub> cathode: Enhanced Na storage performances with the suppressed Mn-O bond anisotropy. **2023**, 460, 141744 ○
- 439 C and N effect on magnetic coupling and vibrational properties of bcc and fcc-FeMnCr alloys: DFT study. **2023**, 570, 170501 ○
- 438 Effects on the glassy phase and electronic structure of Ca<sub>3</sub>Co<sub>2</sub>O<sub>6</sub>. **2023**, 654, 414707 ○

- 437 DFT insights on the opto-electronic and thermoelectric properties of double perovskites  $K_2AgSbX_6$  ( $X = Cl, Br$ ) via halides substitutions for solar cell applications. **2023**, 290, 116338 ○
- 436 Synthesis of technetium hydride  $TcH_{1.3}$  at 27 GPa. **2023**, 107, ○
- 435 Development and Validation of a ReaxFF Reactive Force Field for Modeling Silicon/Carbon Composite Anode Materials in Lithium-Ion Batteries. **2023**, 127, 2818-2834 ○
- 434 Reconciling experimental and theoretical stacking fault energies in face-centered cubic materials with the experimental twinning stress. **2023**, 27, 101708 ○
- 433 Making and Breaking Insight into the Symmetry of Salen Analogues. **2023**, 15, 424 ○
- 432 Thermoelectric transport properties of metal phosphide  $XLiP$  ( $X = Sr, Ba$ ). **2023**, 35, 155501 ○
- 431 Theoretical study of thermal and magneto-electronic properties of  $YbX_2$  ( $X = Co$  and  $Fe$ ) intermetallic compounds. ○
- 430 Molecular dynamics simulation studies of 1,3-dimethyl imidazolium nitrate ionic liquid with water. **2023**, 158, 084505 ○
- 429 Vibrational, thermodynamic and acoustic properties of  $AgAlS_2$  crystal. **2023**, 654, 414731 ○
- 428 Supercapacitor Performance of Magnetite Nanoparticles Enhanced by a Catecholate Dispersant: Experiment and Theory. **2023**, 28, 1562 ○
- 427 Electronic, mechanical, optical and thermodynamic properties of the quaternary semiconductors  $Sr_3GeMgN_4$  and  $Ba_3GeMgN_4$ . **2023**, 290, 116292 ○
- 426 First-principles theory of electrochemical capacitance. **2023**, 444, 142016 ○
- 425 Average-atom model with Siegert states. **2023**, 107, ○
- 424 Interfacial interaction of monolayer  $MX_2$  ( $M = Mo, W$ ;  $X = S, Se, Te$ )/ $SiO_2$  interfaces for composite optical fibers. **2023**, 37, 102739 ○
- 423 Hydrogen molecule adsorption and sensing on lanthanide (La) doped/decorated carbon nanotube and graphene structures. **2023**, 41, 023202 ○
- 422 Experimental and Theoretical Exploration of ES IPT in a Systematically Constructed Series of Benzimidazole Based Schiff Base Probes: Application as Chemosensors. **2023**, 29, ○
- 421 Accelerating the density-functional tight-binding method using graphical processing units. **2023**, 158, 084802 ○
- 420 A variational formulation of the Harris functional as a correction to approximate Kohn-Sham density functional theory. **2023**, 158, 054111 ○

- 419 Change in the Nature of ZSM-5 Zeolite Depending on the Type of Metal Adsorbent The Analysis of DOS and Orbitals for Iron Species. **2023**, 24, 3374 ○
- 418 Toward routine Kohn-Sham inversion using the Lieb-response approach. **2023**, 158, 064102 ○
- 417 Machine learning based modeling of disordered elemental semiconductors: understanding the atomic structure of a-Si and a-C. **2023**, 38, 043001 ○
- 416 Atomic and electronic structures of interfaces between amorphous  $(Al_2O_3)_1x(SiO_2)_x$  and GaN polar surfaces revealed by first-principles simulated annealing technique. **2023**, 133, 065301 ○
- 415 All-Electron Plane-Wave Electronic Structure Calculations. **2023**, 19, 1300-1309 ○
- 414 On the self-consistency of DFT-1/2. **2023**, 158, 094103 ○
- 413 Effect of Hubbard U on the electronic and magnetic properties of  $Ca_2VMoO_6$  double perovskite. **2023**, 570, 170478 ○
- 412  $NO_2$  Physical-to-Chemical Adsorption Transition on Janus WSe Monolayers Realized by Defect Introduction. **2023**, 28, 1644 1
- 411 Monolayer group IV monochalcogenides T-MX (M = Sn, Ge; X = S, Se) with fine piezoelectric performance and stability. **2023**, 122, 062903 ○
- 410 Coherent description of the magnetic properties of  $SeCuO_3$  versus temperature and magnetic field. **2023**, 107, ○
- 409 Promoting Reversibility of Co-Free Layered Cathodes by Al and Cation Vacancy. **2023**, 13, 2204241 ○
- 408 One-body reduced density-matrix functional theory for the canonical ensemble. **2023**, 107, ○
- 407 Doping of Graphene Nanostructure with Iron, Nickel and Zinc as Selective Detector for the Toxic Gas Removal: A Density Functional Theory Study. **2023**, 9, 20 ○
- 406 Intrinsic layer-polarized anomalous Hall effect in bilayer  $MnBi_2Te_4$ . **2023**, 107, ○
- 405 DNA bases detection via  $MoS_2$  field effect transistor with a nanopore: first-principles modeling. **2023**, 114, 253-264 ○
- 404 Tuning the Surface Mn/Al Ratio and Crystal Crystallinity of  $MnAl$  Oxides by Calcination Temperature for Excellent Acetone Low-Temperature Mineralization. **2023**, 3, 487-499 ○
- 403 Strong anisotropic optical response in two-dimensional Mo-VIA and Mo-VIIA monolayer binary materials. **2023**, 53, 101114 ○
- 402 Exploration and investigation of stable novel  $Al_2O_3$  by high-throughput screening and density functional theory. **2023**, 23, 4244-4257 ○

- 401 Band Engineering of the Second Phase to Reach High Thermoelectric Performance in Cu<sub>2</sub>Se-Based Composite Material. 2210345 ○
- 400 Enhanced Magnetization in CoFe<sub>2</sub>O<sub>4</sub> Through Hydrogen Doping. 2212298 ○
- 399 Tuning Catalytic Performance of C<sub>2</sub>N/GaN Heterostructure for Hydrogen Evolution Reaction by Doping. **2023**, 6, ○
- 398 First-principles study on the thermoelectric properties of Sr<sub>2</sub>Si and Sr<sub>2</sub>Ge. **2023**, 32, 101015 ○
- 397 Surface Tamm States of 28 nm Nanodiamond via Raman Spectroscopy. **2023**, 13, 696 ○
- 396 Revealing Structural and Physical Properties of Polylactide: What Simulation Can Do beyond the Experimental Methods. 1-39 ○
- 395 Structural, mechanical, electronic, and thermoelectric properties of TiZrCo<sub>2</sub>Bi<sub>2</sub>, TiHfCo<sub>2</sub>Bi<sub>2</sub>, and ZrHfCo<sub>2</sub>Bi<sub>2</sub> double half Heusler semiconductors. ○
- 394 Tuning of magnetic, electronic and electrolytic water properties of silicene supported precious-metal by non-metal doping and vacancy defect. **2023**, 38, 100486 ○
- 393 Two-Photon Interface of Nuclear Spins Based on the Optonuclear Quadrupolar Effect. **2023**, 13, ○
- 392 Study on low hydrostatic pressure-dependent optoelectronic, mechanical, and anisotropic properties of heavy thallium perovskites TlPbX<sub>3</sub> (X = Cl, Br). **2023**, 38, 2007-2017 ○
- 391 Structure and Dynamics in Liquid Iron at High Pressure and Temperature. A First Principles Study. **2023**, 128, ○
- 390 Octahedral Distortion and Excitonic Behavior of Cs<sub>3</sub>Bi<sub>2</sub>Br<sub>9</sub> Halide Perovskite at Low Temperature. **2023**, 127, 3523-3531 ○
- 389 Polaron-assisted electronic transport in ZnP<sub>2</sub> nanowires. **2023**, 11, 4243-4253 ○
- 388 Chiral Dirac fermion in a collinear antiferromagnet. ○
- 387 How a Ferroelectric Layer Can Tune a Two-Dimensional Electron Gas at the Interface of LaInO<sub>3</sub> and BaSnO<sub>3</sub>: A First-Principles Study. **2023**, 15, 11314-11323 ○
- 386 Toward Pair Atomic Density Fitting for Correlation Energies with Benchmark Accuracy. **2023**, 19, 1499-1516 ○
- 385 Microstructure and wear properties of powder-pack borided Ti<sub>5</sub>Al<sub>0.5</sub>Sn alloy. **2023**, 23, 4032-4043 ○
- 384 A simple uric acid assay by using 3-hydroxytyramine as a chromogenic colorimetric sensor in human serum samples: Density functional theory supported mechanistic approach. **2023**, 70, 894-908 ○

- 383 Orbital Polarization-Dependent Fragment Twist-Induced Intramolecular Electric-Field-Driven Charge Transfer. **2023**, 28, 1801 ○
- 382 Origin of performance degradation in high-delithiation  $\text{Li}_x\text{CoO}_2$ : insights from direct atomic simulations using global neural network potentials. **2023**, 11, 5370-5379 1
- 381 Numerical stability and efficiency of response property calculations in density functional theory. **2023**, 113, ○
- 380 Continuous-flow electrosynthesis of ammonia by nitrogen reduction and hydrogen oxidation. **2023**, 379, 707-712 1
- 379 Structure elucidation of an aspidofractinine-type monoterpene indole alkaloid from *Melodinus reticulatus*. **2022**, ○
- 378 DFT insights of mechanical, optoelectronic and thermoelectric properties for  $\text{Cs}_2\text{ScTlX}_6$  (X = Cl, Br, I) double perovskites. **2023**, 55, ○
- 377 Layered Semiconductor  $\text{Cr}_{0.32}\text{Ga}_{0.68}\text{Te}_{2.33}$  with Concurrent Broken Inversion Symmetry and Ferromagnetism: A Bulk Ferrovalley Material Candidate. **2023**, 145, 4683-4690 ○
- 376 Designing for dopability in semiconducting  $\text{AgInTe}_2$ . **2023**, 11, 3832-3840 ○
- 375 Fast carrier diffusion via synergistic effects between lithium-ions and polarons in rutile  $\text{TiO}_2$ . **2023**, 25, 7519-7526 ○
- 374 Bottom-up building of two-dimensional magnetic materials with self-assembly of superatom  $\text{TM}@\text{Sn}_{12}$  (TM = Sc, Ti, V, Cr, Mn, Fe) clusters. **2023**, 56, 144001 ○
- 373 First-principles studies of electronic properties in a  $\text{Pt}_2\text{CdSe}_3/\text{Pt}_2\text{HgSe}_3$  Kane-Mele heterobilayer. **2023**, 107, ○
- 372 Impact of spin-orbit coupling on physical properties and superconductivity in noncentrosymmetric superconductors  $\text{Ru}_7\text{B}_3$  and Re. **2023**, 655, 414743 ○
- 371 Novel 2D sulfur-doped  $\text{V}_2\text{O}_5$  flakes and their applications in photoelectrochemical water oxidation and high-performance energy storage supercapacitors. **2023**, 461, 141935 ○
- 370 Density functional theory-based quantum-computational analysis on the strain-assisted electronic and photocatalytic properties of  $\text{BX-MSSe}$  (X = P, As and M = Mo, W) heterostructures. **2023**, 129, ○
- 369 On the Nature of Hydrophobic Organic Compound Adsorption to Smectite Minerals Using the Example of Hexachlorobenzene-Montmorillonite Interactions. **2023**, 13, 280 ○
- 368 Tensor Hypercontraction Form of the Perturbative Triples Energy in Coupled-Cluster Theory. **2023**, 19, 1476-1486 ○
- 367 Properties of the double half-Heusler alloy  $\text{ScNbNi}_2\text{Sn}_2$  with respect to structural, electronic, optical, and thermoelectric aspects. **2023**, 363, 115103 ○
- 366 Improving the performance of fermionic neural networks with the Slater exponential Ansatz. ○

- 365 First principles and mean field study on the magnetocaloric effect of YFe<sub>3</sub> and HoFe<sub>3</sub> compounds. **2023**, 13, ○
- 364 Generating Exchange-Correlation Functionals with a Simplified, Self-Consistent Correlation Factor Model. **2023**, 127, 2026-2033 ○
- 363 Tuned optoelectronic and thermoelectric properties of TIMX<sub>2</sub> through M=Ga,In X=S,Se,Te intercalation. **2023**, 20, 131-144 ○
- 362 First-principle insights of initial hydration behavior affected by copper impurity in alite phase based on static and molecular dynamics calculations. **2023**, 398, 136478 ○
- 361 Topography of high-speed steel substrates sputter cleaned by an Ar/Ti cathodic arc plasma. **2023**, 458, 129344 ○
- 360 Cesium-metalloid halide perovskites MBX<sub>3</sub> (M<sub>1</sub><sup>+</sup>= Cs; B<sub>2</sub><sup>+</sup>= Si, Ge, Sn, Pb; X<sub>B</sub><sup>-</sup> Cl, Br, I) as semiconductor photovoltaic materials for sustainable renewable-energy applications. **2023**, 19, 113-140 ○
- 359 Adsorption of juglone on pure and boron-doped C<sub>24</sub> fullerene-like nano-cage: A density functional theory investigation. **2023**, 1222, 114077 ○
- 358 Structural and Spectroscopic Effects of Li<sup>+</sup> Substitution for Na<sup>+</sup> in Li<sub>x</sub>Na<sub>1-x</sub>CaLa<sub>0.5</sub>Er<sub>0.05</sub>Yb<sub>0.45</sub>(MoO<sub>4</sub>)<sub>3</sub> Upconversion Scheelite-Type Phosphors. **2023**, 13, 362 1
- 357 Magnetic phase transition in a machine trained spin model: A study of hexagonal CrN monolayer. **2023**, 615, 128589 ○
- 356 DFT-Based Study for the Enhancement of CO<sub>2</sub> Adsorption on Metal-Doped Nitrogen-Enriched Polytriazines. **2023**, 8, 8876-8884 ○
- 355 The electron-phonon renormalization in the electronic structure calculation: Fundamentals, current status, and challenges. **2023**, 158, 130901 ○
- 354 Unraveling the effects of inter-site Hubbard interactions in spinel Li-ion cathode materials. **2023**, 25, 9061-9072 ○
- 353 Study of Structural, optoelectronic and elastic properties of MAX phase of Ti<sub>2</sub>BrX (X<sup>-</sup>=B, C and N) by density functional theory. **2023**, 150, 110515 ○
- 352 Multiple Quantum States Induced in 1T-TaSe<sub>2</sub> by Controlling the Stacking Order of Charge Density Waves. 2214583 ○
- 351 Insights on Potential Photoprotective Activity of Two Butylchalcone Derivatives: Synthesis, Spectroscopic Characterization and Molecular Modeling. **2023**, 10, 228 ○
- 350 How cation nature controls the bandgap and bulk Rashba splitting of halide perovskites. ○
- 349 High-temperature ferromagnetism in Cr<sub>1+x</sub>Pt<sub>5-x</sub>P. **2023**, 7, ○
- 348 Introduction. **2023**, 1-23 ○



- 347 Self-Supported MoO<sub>2</sub>/MoSi<sub>2</sub> Ceramic Electrode for High Current Density Hydrogen Evolution Reaction. **2023**, 11, 3769-3779 ○
- 346 Initial decomposition mechanisms of 2,4,6-triamino-1,3,5-trinitrobenzene (TATB) and their kinetic isotope effect. **2023**, 133, 075902 ○
- 345 Electronic structure and physical properties of the candidate topological material GdAgGe. **2023**, 107, ○
- 344 Toward self-organizing low-dimensional organic/inorganic hybrid perovskites: Machine learning-driven co-navigation of chemical and compositional spaces. **2023**, 48, 164-172 ○
- 343 Multiscale Modeling of Metal-Oxide-Metal Conductive Bridging Random-Access Memory Cells: From Ab Initio to Finite-Element Calculations. **2023**, 19, ○
- 342 Phase stability and physical properties of lanthanum dicarbide under pressure. 1-18 ○
- 341 Cooperative Pseudo Jahn Teller distortion derives phase transitions in bismuth oxide. **2023**, 299, 127534 ○
- 340 Y<sub>3</sub>Fe<sub>0.5</sub>SiSe<sub>7</sub>: A new cation-deficient quaternary mixed transition metal chalcogenide with extremely low thermal conductivity. **2023**, 138, 107133 ○
- 339 Self-Consistent-Charge Density-Functional Tight-Binding Parameters for Modeling an All-Solid-State Lithium Battery. **2023**, 19, 1381-1387 ○
- 338 Introduction. **2023**, 1-7 ○
- 337 Dynamic Precision Approach for Accelerating Large-Scale Eigenvalue Solvers in Electronic Structure Calculations on Graphics Processing Units. **2023**, 19, 1457-1465 ○
- 336 Effects of an external electric field on the electronic properties and optical excitations of germanane and silicane monolayers. **2023**, 35, 175502 ○
- 335 Boron vacancy-driven thermodynamic stabilization and improved mechanical properties of AlB<sub>2</sub>-type tantalum diborides as revealed by first-principles calculations. **2023**, 6, 025002 ○
- 334 Accurate non-covalent interaction energies on noisy intermediate-scale quantum computers via second-order symmetry-adapted perturbation theory. **2023**, 14, 3587-3599 ○
- 333 Ab initio study of the adsorption of SO<sub>2</sub> on single-atom Cu-decorated ZnO(0001) surface. **2023**, 29, ○
- 332 Vapor-Phase Indium Intercalation in van der Waals Nanofibers of Atomically Thin W<sub>6</sub>Te<sub>6</sub> Wires. **2023**, 17, 5561-5569 ○
- 331 Tunable magnetic interactions in mono-transition metal spinels MgCr<sub>2</sub>X<sub>4</sub> (X=O,S,Se). **2023**, 7, ○
- 330 CIMG-BERT: Pretraining Bidirectional Transformers with Chemistry Knowledge for Molecular Property Prediction. **2022**, ○

- 329 Theoretical Study on the Role of Solvents in Lithium Polysulfide Anchoring on Vanadium Disulfide Facets for Lithium Sulfur Batteries. **2023**, 127, 4416-4424 ○
- 328 Effect of Series Resistance and Temperature on PCE of Double Perovskite La<sub>2</sub>NiMnO<sub>6</sub>-Based PSC. **2023**, 407, ○
- 327 Unveiling the Multistep Electrochemical Desorption Mechanism of Cubic NiO Films for Transmissive-to-Black Electrochromic Energy Storage Devices. **2023**, 14, 2284-2291 ○
- 326 Multiscale simulations of nanofluidics: Recent progress and perspective. ○
- 325 Mixed-Valence Conductors from Ni Bis(diselenolene) Complexes with a Thiazoline Backbone. **2023**, 62, 4197-4209 ○
- 324 Gaussian basis functions for an orbital-free-related density functional theory of atoms. ○
- 323 Optical Absorption and Second-Harmonic Generation in Violet Phosphorene: Experimental and Theoretical Aspects. 2202770 ○
- 322 Phase formation of powders sputtered from X<sub>2</sub>BC targets and XC+XB powder mixtures {X = Nb, Ta, W}. **2023**, 458, 129379 ○
- 321 High-Energy Aqueous Magnesium Ion Batteries with Capacity-Compensation Evolved from Dynamic Copper Ion Redox. 2300148 ○
- 320 A density functional theory for ecology across scales. **2023**, 14, ○
- 319 DFT-aided machine learning-based discovery of magnetism in Fe-based bimetallic chalcogenides. **2023**, 13, ○
- 318 Analysis of XGaO<sub>3</sub> (X = Ba and Cs) cubic based perovskite materials for photocatalytic water splitting applications: a DFT study. **2023**, 9, e14112 ○
- 317 Mechanism of tungsten strengthening hydrogen transportation in Nb<sub>48</sub>Ti<sub>27</sub>Co<sub>25</sub> hydrogen permeable alloy membrane. **2023**, 23, 5413-5422 ○
- 316 Analytical Forces for the Optimized Effective Potential Calculations. **2023**, 19, 1744-1752 ○
- 315 Multilayer In-Plane Heterostructures Based on Transition Metal Dichalcogenides for Advanced Electronics. **2023**, 17, 6545-6554 ○
- 314 Electrical and thermal conductivity of fcc and hcp iron under conditions of the Earth's core from ab initio simulations. **2023**, 107, 1
- 313 Red-Fluorescing Paramagnetic Conjugated Polymer Nanoparticles-Triphenyl Methyl Radicals as Monomers in Cu Cross-Coupling Dispersion Polymerization. **2023**, 56, 2104-2112 ○
- 312 Strain effect on the high T<sub>c</sub> superconductor YBa<sub>2</sub>Cu<sub>3</sub>O<sub>7</sub>: an ab initio study comparing bulk and monolayer models. **2023**, 5, 015002 ○

- 311 Linear response based theories for Dzyaloshinskii-Moriya interactions. **2023**, 107, ○
- 310 Iridium Incorporation into MnO<sub>2</sub> for an Enhanced Electrocatalytic Oxygen Evolution Reaction. ○
- 309 Basic Concepts of Quantum Mechanics. **2023**, 35-86 ○
- 308 The first-principle study on certain structural, band-structural, elastic, optical and piezoelectric properties of the Ca, Zr and Ca/Zr-doped BaTiO<sub>3</sub>. **2023**, 37, ○
- 307 Electronic, spintronic, and piezoelectric properties of new Janus ZnAXY (A=Si,Ge,Sn, and . **2023**, 107, ○
- 306 Multimodal Luminescent Low-Dimension Cs<sub>2</sub>ZrCl<sub>6</sub>:xSb<sup>3+</sup> Crystals for White Light-Emitting Diodes and Information Encryption. **2023**, 39, 3792-3799 ○
- 305 Restricted multicanonical sampling for machine learning potential construction. **2023**, 107, ○
- 304 Remote Site-Selective C(sp<sup>3</sup>)H Monodeuteration of Unactivated Alkenes via Chain-Walking Strategy. **2023**, 13, 3644-3654 ○
- 303 Molecular Orbitals. **2023**, ○
- 302 Comparative Theoretical Study of Quinazolinone Derivatives in Different Solvents and Gas Phase. A Density Functional Theory (DFT) Calculations. ○
- 301 A heterogeneous processing-in-memory approach to accelerate quantum chemistry simulation. **2023**, 116, 103017 ○
- 300 Effect of Substitutional Oxygen on Properties of Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub>MXene Produced Using Recycled TiO<sub>2</sub> Source. 2201715 ○
- 299 First Principles Computation of New Topological B<sub>2</sub>X<sub>2</sub>Zn (X = Ir, Rh, Co) Compounds. **2023**, 6, 152-163 ○
- 298 Raman and Far-Infrared Synchrotron Nanospectroscopy of Layered Crystalline Talc: Vibrational Properties, Interlayer Coupling, and Symmetry Crossover. **2023**, 127, 5876-5885 1
- 297 High temperature spectroscopy of ensembles of nitrogen-vacancy centers in diamond. **2023**, 133, 094401 ○
- 296 Structural-Stability Study of Antiperovskite Na<sub>3</sub>OCl for . **2023**, 19, ○
- 295 Influence of Nitrogen Substitution on the Electronic Structure of Ti<sub>2</sub>O<sub>3</sub>: Insights into the Doping-Induced Insulator-to-Metal Transition. 2200495 ○
- 294 Self-screening corrections beyond the random-phase approximation: Applications to band gaps of semiconductors. **2023**, 107, ○

- 293 ELECTRONIC STRUCTURE AND MAGNETIC PROPERTIES OF THE ISOMERS OF BIS-PHENALENYL STILBENE DERIVATIVES: A QUANTUM CHEMICAL STUDY. **2023**, 64, 58-68 ○
- 292 Orbital magnetoelectric effect in nanoribbons of transition metal dichalcogenides. **2023**, 107, ○
- 291 Estimation of electron temperature dynamics in warm dense gold using measured and DFT-calculated plasma frequency. **2023**, 98, 045813 ○
- 290 Effect of Magnetic Coupling on the Optical Properties of Oxide Co Nanowires on Vicinal Pt Surfaces. **2023**, 9, 72 ○
- 289 Magnetism and  $T_c$  phase diagrams of Na- and Ag-substituted  $\text{EuCd}_2\text{As}_2$ . **2023**, 7, ○
- 288 Prediction of topological phases in metastable ferromagnetic  $\text{MPX}_3$  monolayers. **2023**, 107, ○
- 287 First-principles-based simulation of the electrocaloric effect. **2023**, 63-91 ○
- 286 Spatial Decay and Limits of Quantum Solute-Solvent Interactions. **2023**, 14, 2473-2480 ○
- 285 Triterpene Derivatives as Potential Inhibitors of the RBD Spike Protein from SARS-CoV-2: An In Silico Approach. **2023**, 28, 2333 ○
- 284 Exploring and machine learning structural instabilities in 2D materials. **2023**, 9, ○
- 283 Atomic-Scale Revealing the Structure Distribution between  $\text{LiMO}_2$  and  $\text{Li}_2\text{MnO}_3$  in Li-Rich and Mn-Based Oxide Cathode Materials. **2023**, 13, ○
- 282 The electronic structure of  $\text{TeO}_2$  as wide bandgap p-type oxide semiconductor. **2023**, 122, 101901 ○
- 281 Innate dynamics and identity crisis of a metal surface unveiled by machine learning of atomic environments. **2023**, 158, 124701 ○
- 280 Strongly constrained and appropriately normed density functional theory exchange-correlation functional applied to rare earth oxides. **2023**, 107, ○
- 279 Role of spin-orbit coupling effects in rare-earth metallic tetra-borides: a first principle study. **2023**, 96, ○
- 278 Enhancing spin splitting by symmetry and molecular orbital hybridization in  $\text{VO}_2$ . **2023**, 222, 112100 ○
- 277 Uncertainty-driven dynamics for active learning of interatomic potentials. **2023**, 3, 230-239 ○
- 276 The  $\text{P}(4S) + \text{NH}(3\Sigma^-)$  and  $\text{N}(4S) + \text{PH}(3\Sigma^-)$  reactions as sources of interstellar phosphorus nitride. **2023**, 40, ○

- 275 Low resistance electrical contacts to few-layered MoS<sub>2</sub> by local pressurization. **2023**, 10, 021003 ○
- 274 First-Principles Modeling of the Adsorption Mechanism of Carboxylic and Phosphonic Acids onto Pristine and Defective Delafossite CuAlO<sub>2</sub> Surfaces. ○
- 273 High reactivity of mesoporous CeO<sub>2</sub> to dissociate chemical warfare agent sarin. ○
- 272 Walter Kohn Centennial Nobel laureate for density functional theory. ○
- 271 Electronic and optical properties of boron nitride nanoribbons exploiting DFT. **2023**, 55, ○
- 270 High-pressure studies of atomically thin van der Waals materials. **2023**, 10, 011313 ○
- 269 Introducing the Random Phase Approximation Theory. **2023**, 9, 141 ○
- 268 Superconductivity in Te-Deficient ZrTe<sub>2</sub>. **2023**, 127, 5162-5168 ○
- 267 DELTA50: A Highly Accurate Database of Experimental <sup>1</sup>H and <sup>13</sup>C NMR Chemical Shifts Applied to DFT Benchmarking. **2023**, 28, 2449 ○
- 266 Magnetoelastic Coupling and Cryogenic Magnetocaloric Effect in Two-Site Disordered GdSrCoFeO<sub>6</sub> Double Perovskite. **2023**, 35, 2439-2455 ○
- 265 Floquet Engineering of Nonequilibrium Valley-Polarized Quantum Anomalous Hall Effect with Tunable Chern Number. **2023**, 23, 2166-2172 ○
- 264 Electronic Excited States in Extreme Limits via Ensemble Density Functionals. **2023**, 130, ○
- 263 An automated reaction route mapping for the reaction of NO and active species on Ag<sub>4</sub> clusters in zeolites. **2023**, 25, 8524-8531 ○
- 262 Coupling interface constructions of FeOOH/NiCo<sub>2</sub>S<sub>4</sub> by microwave-assisted method for efficient oxygen evolution reaction. ○
- 261 Hybrid Functional-Based Scissors Operator for Perovskite Oxide Nanostructures: A NaTaO<sub>3</sub> Case Study. **2023**, 127, 5604-5612 ○
- 260 Role of Chalcogen Defect Introducing Metal-Induced Gap States and Its Implications for Metal/MDs Interface Chemistry. **2023**, 8, 10176-10184 ○
- 259 Using Collocation to Solve the Schrödinger Equation. **2023**, 19, 1641-1656 ○
- 258 Valence Band Structure Degeneracy Enhanced Thermoelectric Performance in ECu<sub>2</sub>Se. **2023**, 127, 5576-5583 ○

- 257 Hydroxyapatite materials-synthesis routes, mechanical behavior, theoretical insights, and artificial intelligence models: a review. ○
- 256 Fluids and Electrolytes under Confinement in Single-Digit Nanopores. **2023**, 123, 2737-2831 ○
- 255 AisNet: A Universal Interatomic Potential Neural Network with Encoded Local Environment Features. **2023**, 63, 1756-1765 ○
- 254 Hydrothiolation of alkynes with thiolatechol derivatives catalysed by CuNPs/TiO<sub>2</sub>: exploring the reaction mechanism by DFT calculations. **2023**, 13, 8025-8033 ○
- 253 Dual-Channel Dielectric Tunability in Highly Textured BaTi<sub>0.99</sub>Fe<sub>0.01</sub>O<sub>3</sub> Ceramics With Micro-Twin Boundary. 2201243 ○
- 252 Interface synergism and engineering of Pd/Co@N-C for direct ethanol fuel cells. **2023**, 14, ○
- 251 Investigation of electronic and thermal properties of CoCrFe and CoCrFeNi high entropy alloys via extended tight-binding DFT computational method. ○
- 250 Symbolic Regression in Materials Science: Discovering Interatomic Potentials from Data. **2023**, 1-30 ○
- 249 Decreased spin-resolved anti-bonding states filling to accelerate CHO conversion into CH<sub>2</sub>O in transitional metal-doped Mo<sub>2</sub>C monolayers during CO<sub>2</sub> reduction. ○
- 248 Assessments of DFT-based energy decomposition analysis methods for intermolecular interactions. **2023**, 158, 124116 ○
- 247 Structure Defects and Photovoltaic Properties of TiO<sub>2</sub>:ZnO/CuO Solar Cells Prepared by Reactive DC Magnetron Sputtering. **2023**, 13, 3613 ○
- 246 Symmetry of Identical Particles, Modern Achievements in the Pauli Exclusion Principle, in Superconductivity and in Some Other Phenomena. **2023**, 15, 701 ○
- 245 Classical Density Functional Theory: Representability and Universal Bounds. **2023**, 190, ○
- 244 Unconventional gapless semiconductor in an extended martini lattice in covalent honeycomb materials. **2023**, 107, ○
- 243 Hydration Mechanisms of Tungsten Trioxide Revealed by Water Adsorption Isotherms and First-Principles Molecular Dynamics Simulations. **2023**, 127, 5584-5596 ○
- 242 PEA<sub>2</sub>PbI<sub>4</sub>: fast two-dimensional lead iodide perovskite scintillator with green and red emission. **2023**, 29, 101455 ○
- 241 The Effect of Cr Additive on the Mechanical Properties of Ti-Al Intermetallics by First-Principles Calculations. **2023**, 13, 488 ○
- 240 Generalized exciton with a noninteger particle and hole charge as an excitation order. **2023**, ○

- 239 Fluence dependent dynamics of excitons in monolayer MoSi<sub>2</sub>Z<sub>4</sub> (Z = pnictogen). **2023**, 35, 235701 ○
- 238 The Temperature Dependence of the Hexagonal Boron Nitride Oxidation Resistance, Insights from First-Principle Computations. **2023**, 13, 1041 ○
- 237 Impact of active sites on encapsulation of curcumin in Metal Organic Frameworks. **2023**, 10, 035102 ○
- 236 Current Status and Future Scope of Phase Diagram Studies. **2023**, 63, 407-418 ○
- 235 Simulating Highly Activated Sticking of H<sub>2</sub> on Al(110): Quantum versus Quasi-Classical Dynamics. **2023**, 127, 5395-5407 ○
- 234 A DFT study on how vanadium affects hydrogen storage kinetics in magnesium nickel hydride. **2023**, ○
- 233 Electrical conductivity of iron in Earth's core from microscopic Ohm's law. **2023**, 107, 1
- 232 Theoretical Research Methods Involved in This Book. **2023**, 19-43 ○
- 231 Structural and thermodynamic properties of quasi-2D Mo(1-x)W<sub>x</sub>(S, Se, Te)<sub>2</sub> monolayer alloys: a statistical first principle study. **2023**, 34, 275704 ○
- 230 Study of electronic, magneto-optical and transport properties of double perovskite Ca<sub>2</sub>XMnO<sub>6</sub> (X = Ti, Cr) under uniaxial compressive strain by using a DFT method. **2023**, 37, ○
- 229 An effective strategy for CO<sub>2</sub> reduction to C<sub>1</sub> products using Cu-embedded MoS<sub>2</sub> electrocatalyst: DFT study. **2023**, 47, 6932-6942 ○
- 228 Systematically improvable mean-field variational ansatz for strongly correlated systems: Application to the Hubbard model. **2023**, 107, ○
- 227 Grand Canonical Quantum Mechanics with Applications to Mechanisms and Rates for Electrocatalysis. ○
- 226 Electronic density response of warm dense matter. **2023**, 30, 032705 2
- 225 Low-rank approximations to accelerate hybrid functional enabled real-time time-dependent density functional theory within plane waves. **2023**, 5, 014008 ○
- 224 Engineering an iron atom-cluster nanostructure towards efficient and durable electrocatalysis. **2023**, 11, 8202-8212 ○
- 223 Electron-phonon interaction and phonon frequencies in two-dimensional doped semiconductors. **2023**, 107, ○
- 222 Study of pnictides for photovoltaic applications. **2023**, 25, 9626-9635 ○

- 221 Numerical Representations of Chemical Data for Structure-Based Machine Learning. **2023**, 173-200 ○
- 220 Ab initio investigation of Bi/BaTiO<sub>3</sub> and Bi/PbTiO<sub>3</sub> heterostructures for spintronic applications. **2023**, 605, 27-35 ○
- 219 Correlation energy of the paramagnetic electron gas at the thermodynamic limit. **2023**, 107, ○
- 218 Non-Uniformly Strained CoreShell InAs/InP Nanowires for Mid-Infrared Photonic Applications. **2023**, 6, 5460-5468 ○
- 217 Accelerating self-consistent field iterations in Kohn-Sham density functional theory using a low-rank approximation of the dielectric matrix. **2023**, 107, 1
- 216 Nanostructured system based on hydroxyapatite and curcumin: A promising candidate for osteosarcoma therapy. **2023**, ○
- 215 Observation of electron orbital signatures of single atoms within metal-phthalocyanines using atomic force microscopy. **2023**, 14, ○
- 214 Low-rank approximations for accelerating plane-wave hybrid functional calculations in unrestricted and noncollinear spin density functional theory. **2023**, 158, 134106 ○
- 213 Decomposing Chemical Space: Applications to the Machine Learning of Atomic Energies. **2023**, 19, 2029-2038 ○
- 212 Facile Tailoring of Surface Terminations of MXenes by Doping Nb Element: Toward Extraordinary Pseudocapacitance Performance. **2023**, 15, 15367-15376 ○
- 211 Open source variational quantum eigensolver extension of the quantum learning machine for quantum chemistry. 1
- 210 Solving Schrodinger equations using a physically constrained neural network\*. **2023**, 47, 054104 ○
- 209 Critical interphase overpotential as a lithium dendrite-suppression criterion for all-solid-state lithium battery design. ○
- 208 Absorption Modulation and Anomalous Thermal Transport in Two-dimensional X-AlN(X=C,Si,TC) Semiconductor. **2023**, 0 ○
- 207 Electrophilicity and nucleophilicity scales at different DFT computational levels. ○
- 206 A study of the structural, thermodynamic, magnetic, and optoelectronic properties of the Dy<sub>2</sub>Be<sub>2</sub>GeO<sub>7</sub> complex oxide via ab initio methods. **2023**, 138, ○
- 205 Temperature phase transitions in silver niobate and lithium tantalate-modified silver niobate ceramics. ○
- 204 Excimer Energies. **2023**, 14, 2917-2926 ○



- 203 Revealing the impact of organic spacers and cavity cations on quasi-2D perovskites via computational simulations. **2023**, 13,
- 202 Giant hyperferroelectricity in LiZnSb and its origin. **2023**, 107,
- 201 A systematic computational investigation of lithiation-induced structural phase transitions of O-functionalized MXenes. **2023**, 25, 9428-9436
- 200 Connecting Higher-Order Topology with the Orbital Hall Effect in Monolayers of Transition Metal Dichalcogenides. **2023**, 130,
- 199 Activating bulk nickel foam for the electrochemical oxidization of ethanol by anchoring MnO<sub>2</sub>@Au nanorods. **2023**, 11, 8101-8109
- 198 Possible electronic state quasi-half-valley metal in a VGe<sub>2</sub>P<sub>4</sub> monolayer. **2023**, 107,
- 197 Mathematical Modeling of the Perovskite and Double Perovskite Crystal Structure. **2022**, 51, 659-661
- 196 Magnetic properties of rare-earth-lean ThMn<sub>12</sub>-type (Nd,X)Fe<sub>11</sub>Ti (X: Y and Ce) compounds: A DFT study. **2023**, 572, 170645
- 195 Higher-order Rayleigh-quotient gradient effect on electron correlations. **2023**, 158, 134102
- 194 Effects of pressure on structural, mechanical, and electronic properties of chalcopyrite compound CuAlS<sub>2</sub>. **2023**, 20, 215-225
- 193 Investigation of the N<sup>+</sup>C Ligand Effects on Emission Characteristics in a Series of Bis-Metalated [Ir(N<sup>+</sup>C)<sub>2</sub>(N<sup>+</sup>N)]<sup>+</sup> Complexes. **2023**, 28, 2740
- 192 Indolyl imine substituted BODIPY systems; synthesis, photophysical, and biological properties. **2023**, 137, 133367
- 191 Engineering Catalytically Active Sites by Sculpting Artificial Edges on MoS<sub>2</sub> Basal Plane for Dinitrogen Reduction at a Low Overpotential. 2206357
- 190 Density-potential inversion from Moreau-Vosida regularization. **2023**, 5, 014009
- 189 Anion influence on electronic and optical properties of several iodobismuthates. **2023**, 223, 112138
- 188 Electronic signatures of successive itinerant, antiferromagnetic transitions in hexagonal La<sub>2</sub>Ni<sub>7</sub>. **2023**, 35, 245501
- 187 Speciation of borate in aqueous solutions studied experimentally by potentiometry and Raman spectroscopy and computationally by DFT calculations.
- 186 On the energetic and magnetic stability of neutral and charged lithium clusters doped with one and two yttrium atoms. **2023**, 25, 9656-9668


- 185 Ab Initio Study of the Electronic Structure and Lattice Dynamics of Scheelite Type  $\text{AgTcO}_4$ . ○
- 184 DFT Treatment of Some Cantharidine Isomers and Some Radicals from Them. 77-91 ○
- 183 Simulation of Hafnium-Based FinFET and Ferroelectricity of Related 2-Dimensional Hafnium-Based Materials. 29, 69-76 ○
- 182 Structural Bistability in RbI Monolayers on  $\text{Ag}(111)$ . **2023**, 14, 3023-3030 ○
- 181 Density functional theory. **2022**, ○
- 180 Gate-Induced Trans-Dimensionality of Carrier Distribution in Bilayer Lateral Heterosheet of  $\text{MoS}_2$  and  $\text{WS}_2$  for Semiconductor Devices with Tunable Functionality. **2023**, 6, 5434-5439 ○
- 179 Understanding the fundamentals of  $\text{TiO}_2$  surfaces Part II. Reactivity and surface chemistry of  $\text{TiO}_2$  single crystals. **2022**, 38, 846-906 ○
- 178 Electronic, Optical, Thermoelectric and Elastic Properties of  $\text{RbxCs}_{1-x}\text{PbBr}_3$  Perovskite. **2023**, 28, 2880 ○
- 177 Embedding vertex corrections in GW self-energy: Theory, implementation, and outlook. **2023**, 158, 144105 ○
- 176 Effects of High Pressure on the Bandgap and the  $d\bar{d}$  Crystal Field Transitions in Wolframite  $\text{NiWO}_4$ . **2023**, 127, 6543-6551 ○
- 175 Testing Koopmans spectral functionals on the analytically solvable Hooke  $\delta$  atom. **2023**, 158, 144113 ○
- 174 Intrinsic Nonlinear Planar Hall Effect. **2023**, 130, ○
- 173 Comparison of LANL2DZ and SBKJC Basis Sets, over DFT Calculations of a Typical Half-Sandwich Ruthenium Complex. **2022**, 96, 3161-3169 ○
- 172 Quantum computation for periodic solids in second quantization. **2023**, 5, ○
- 171 Automatic differentiation for orbital-free density functional theory. **2023**, 158, 124801 ○
- 170 Role of Non-Covalent Interactions in Carbonic Anhydrase III Opiramate Complex Based on QM/MM Approach. **2023**, 16, 479 ○
- 169 Size Limiting Elemental Ferroelectricity in Bi Nanoribbons: Observation, Mechanism, and Opportunity. **2023**, 14, 3160-3167 ○
- 168 Dual-Atom Doping Carbon Materials as Highly Efficient Electrocatalysts for Lithium Sulfur Batteries: Bimetallic Cooperation Mechanism. **2023**, 127, 6271-6279 ○

- 167 Superionic effect and anisotropic texture in Earth's inner core driven by geomagnetic field. **2023**, 14,
- 166 Substrate-Induced Changes on the Optical Properties of Single-Layer WS<sub>2</sub>. **2023**, 16, 2591
- 165 From ultra-low friction to superlubricity state of black phosphorus: Enabled by the critical oxidation and load.
- 164 ZrSe<sub>2</sub>-HfSe<sub>2</sub> lateral heterostructures: stability, fundamental properties, and interline defects. **2023**, 129,
- 163 DFT-Based Investigation of the Structural Stability, Elastic, Electronic, and Magnetic in Pd<sub>2</sub>CrGe Heusler Alloy.
- 162 Novel Bis-1,3,4-Thiadiazoles Derivatives: Synthesis, Spectroscopic Characterization, DFT Calculations and Evaluation of their Antimicrobial and Antioxidant Activities. **2023**, 44, 81-89
- 161 Complexation behaviour of piceatannol ligand with Ti(IV) and Zr(IV) metal ions: a combined DFT and deep learning investigation.
- 160 Electronic band structure and density of state modulation of amphetamine and ABW type zeolite adsorption system: DFT-CASTEP analysis. **2023**, 29,
- 159 Two-dimensional dichalcogenides of type XY<sub>2</sub> (X=Mo,W; Y=S,Se): A DFT study of the structural, optoelectronic, thermodynamic properties, infrared, and Raman spectra.
- 158 Single Selenium Atomic Vacancy Enabled Efficient Visible-Light-Response Photocatalytic NO Reduction to NH<sub>3</sub> on Janus WSSe Monolayer. **2023**, 28, 2959
- 157 Interface-vacancy synergy of Co(OH)<sub>2</sub>/CoN to boost alkaline water splitting.
- 156 Unveiling the local structure of the amorphous metal Fe<sub>1-x</sub>Zr<sub>x</sub> combining first-principles-based simulations and modelling of EXAFS spectra. **2023**, 13,
- 155 Resolving the polar interface of infinite-layer nickelate thin films. **2023**, 22, 466-473
- 154 Noncollinear and Spin-Flip TDDFT in Multicollinear Approach.
- 153 Synthesis and optical properties of phosphorus doped ZnO: X-ray absorption, X-ray emission, and X-ray excited optical luminescence studies. **2023**,
- 152 First-Principles Study of the Structural, Phase-Stability, Electronic, Magnetic, and Elastic Properties of Heusler Alloys VXRh<sub>2</sub> (X=Si, Ge, and Sn).
- 151 Predicting the Na<sup>+</sup> ion transport properties of NaSICON materials using density functional theory and Kinetic Monte Carlo.
- 150 Catalytic Activity Maps for Alloy Nanoparticles. **2023**, 145, 7352-7360

- 149 Intersite Coulomb Interactions in Charge-Ordered Systems. **2023**, 130, ○
- 148 Silicate Dissolution Mechanism from Metakaolinite Using Density Functional Theory. **2023**, 13, 1196 ○
- 147 Machine-learning accelerated annealing with fitting-search style for multicomponent alloy structure predictions. **2023**, 7, ○
- 146 A first principles study of structural and optoelectronic properties and photocatalytic performance of  $\text{GeCMX}_2$  ( $M = \text{Mo}$  and  $\text{W}$ ;  $X = \text{S}$  and  $\text{Se}$ ) van der Waals heterostructures. ○
- 145 Dopability and Magnetic Properties of 3d Transition Metals in an Atomic-Thick SnTe (001) Monolayer. **2023**, 127, 6389-6395 ○
- 144 Fractional Charge Density Functional Theory and Its Application to the Electro-inductive Effect. **2023**, 14, 3329-3334 ○
- 143 Experimental and Theoretical Investigations of Out-of-Plane Ordered Nanolaminate Transition Metal Borides:  $\text{M}_4\text{CrSiB}_2$  ( $M = \text{Mo}, \text{W}, \text{Nb}$ ). **2023**, 62, 5341-5347 ○
- 142 Scanning Intrinsic Superconductivity and the Spin Hall Effect in Niobium Borides. ○
- 141 Nanoscale fluctuation of stacking fault energy strengthens multi-principal element alloys. **2023**, 158, 218-225 ○
- 140 Ab initio calculation of carrier mobility in semiconductors including ionized-impurity scattering. **2023**, 107, ○
- 139 Magnetic iron-cobalt silicides discovered using machine-learning. **2023**, 7, ○
- 138 Variational atomic model of plasma accounting for ion radial correlations and electronic structure of ions. **2023**, 107, ○
- 137 First-principles investigation on the structural, vibrational, mechanical, electronic, and optical properties of  $\text{MSi}_2\text{Z}_4$  ( $M : \text{Pd}$  and  $\text{Pt}$ , . **2023**, 7, ○
- 136 Intercalation of  $\text{CO}_2$  Selected by Type of Interlayer Cation in Dried Synthetic Hectorite. **2023**, 39, 4895-4903 ○
- 135 meso-Carbazole decorated BODIPYs in an electron donor-acceptor system with excellent fluorosolvato/vapochromic behavior, aggregation-induced emission, and antileishmanial activity. ○
- 134 SbSeI'nin Anizotropik Elastik Ėelliklerine Basıncı Etkileri: DFT Hesaplaması ○
- 133 Fast general two- and three-body interatomic potential. **2023**, 107, ○
- 132 Activating the paddle-wheel effect towards lower temperature in a new sodium-ion solid electrolyte,  $\text{Na}_{3.5}\text{Si}_0.5\text{P}_0.5\text{Se}_4$ . ○

- 131 The Structure of Density-Potential Mapping. Part I: Standard Density-Functional Theory. ○
- 130 Polaron-induced metal-to-insulator transition in vanadium oxides from density functional theory calculations. **2023**, 107, ○
- 129 On the Donor: Acceptor Features for Poly(3-hexylthiophene): TiO<sub>2</sub> Quantum Dots Hybrid Materials Obtained via Water Vapor Flow Assisted Sol-Gel Growth. **2023**, 15, 1706 ○
- 128 Toward density functional theory on quantum computers?. **2023**, 14, ○
- 127 Pressure-Induced Semiconductor-to-Metallic Transition of Monoclinic KCa<sub>2</sub>Nb<sub>3</sub>O<sub>10</sub> Layered Perovskite: A Theoretical DFT Insight. ○
- 126 HfXO (X = S and Se) Janus monolayers as promising two-dimensional platforms for optoelectronic and spintronic applications. ○
- 125 Bibliography. **2023**, 431-464 ○
- 124 Influencing Bonding Interactions of the Neptunyl (V, VI) Cations with Electron-Donating and -Withdrawing Groups. **2023**, 62, 6055-6064 ○
- 123 Corrosion Inhibiting by Some Organic Heterocyclic Inhibitors Through Langmuir Adsorption Mechanism on the Al-X (X = Mg/Ga/Si) Alloy Surface: A Study of Quantum Three-Layer Method of CAM-DFT/ONIOM. **2023**, 9, ○
- 122 Current-induced spin polarization in Janus WSSe monolayer. **2023**, 107, ○
- 121 Investigating the Thermodynamics and Kinetics of Catechin Pyrolysis for Environmentally Friendly Binders. **2023**, 8, 12693-12701 ○
- 120 Lu doping nickel oxide thin films using sol-gel spin coated and density functional theory: optoelectronic and magnetic properties. **2023**, 9, e14874 ○
- 119 Quasi-2D FCC lithium crystals inside defective bi-layer graphene: insights from first-principles calculations. **2023**, 34, 101293 ○
- 118 Solubility and diffusivity of hydrogen and its isotopes in the BeO system. ○
- 117 Structure and mechanism of oxalate transporter OxIT in an oxalate-degrading bacterium in the gut microbiota. **2023**, 14, ○
- 116 Isolated Cu-Sn diatomic sites for enhanced electroreduction of CO<sub>2</sub> to CO. ○
- 115 Theory of x-ray absorption spectroscopy: A microscopic Bloch equation approach for two-dimensional solid states. **2023**, 5, ○
- 114 Experimental Investigation and Thermodynamic Assessment of the Ternary AlNiEr System. **2023**, 11, 1061 ○

- 113 Improved Hydrogen Storage Properties and Air Stability of Metal Hydrides by Constructing Heterophase Composites. ○
- 112 Absorption versus adsorption: high-throughput computation of impurities in 2D materials. **2023**, 7, ○
- 111 Quantum Embedding Method for the Simulation of Strongly Correlated Systems on Quantum Computers. **2023**, 14, 3491-3497 ○
- 110 Shift Current in Molecular Crystals Possessing Charge-Transfer Characteristics. **2023**, 19, ○
- 109 Tuning Multiple Landau Quantization in Transition-Metal Dichalcogenide with Strain. ○
- 108 Influence of defects on the valley polarization properties of monolayer MoS<sub>2</sub> grown by chemical vapor deposition. **2023**, 107, ○
- 107 Bulk and surface electronic structure of NiBi<sub>3</sub>. **2023**, 107, ○
- 106 First-Principles Insights into the Relative Stability, Physical Properties, and Chemical Properties of MoSe<sub>2</sub>. **2023**, 8, 13799-13812 ○
- 105 Growth, spectral and quantum chemical investigations on N-butyl-4-nitroaniline single crystal for nonlinear optical and optoelectronic device applications. **2023**, 34, ○
- 104 Imaginary-time correlation function thermometry: A new, high-accuracy and model-free temperature analysis technique for x-ray Thomson scattering data. **2023**, 30, 042707 ○
- 103 State-Specific Configuration Interaction for Excited States. ○
- 102 Density functional theory (DFT) simulation and approach to property-driven investigations in ceramic and composites materials. **2023**, 461-490 ○
- 101 Benchmark Study on Phosphorescence Energies of Anthraquinone Compounds: Comparison between TDDFT and UDFT. **2023**, 28, 3257 ○
- 100 Polyethylene Glycol 20k. Does It Fluoresce?. **2023**, 8, 14208-14218 ○
- 99 Sensory Sensitivity to the Form of Ga<sub>2</sub>O<sub>3</sub> Nanoparticles. **2022**, 20, ○
- 98 Phase stability of TiAl-based BCC high entropy alloys. **2023**, 158, 107893 ○
- 97 Colloquium : Room temperature superconductivity: The roles of theory and materials design. **2023**, 95, ○
- 96 Structural, vibrational, electronic, and elastic properties of 2D alkali carbide as a metallic material. **2023**, e00805 ○

- 95 Some problems in density functional theory. **2023**, 113,
- 94 Metastable Polymorphic Phases in Monolayer TaTe<sub>2</sub>.
- 93 Seven useful questions in density functional theory. **2023**, 113,
- 92 (MgO)<sub>60</sub>: A magic cluster active in UV range under DFT study. **2023**,
- 91 Electronic structure and microscopic model of Cu<sub>2</sub>(SeO<sub>3</sub>)F<sub>2</sub>: a 2-D AFM ladder compound. **2023**, 96,
- 90 Elucidating the Role of Noncovalent Interactions in Favipiravir, a Drug Active against Various Human RNA Viruses; a 1H-14N NQDR/Periodic DFT/QTAIM/RDS/3D Hirshfeld Surfaces Combined Study. **2023**, 28, 3308
- 89 Ab initio calculations of the 2p<sub>3/2</sub>->2s transition in He-, Li-, and Be-like uranium. **2023**, 107,
- 88 Adaptive Exploration and Optimization of Materials Crystal Structures.
- 87 Quantum version of the integral equation theory-based dielectric scheme for strongly coupled electron liquids. **2023**, 158, 141102
- 86 Modelling of the structural, electronic, magnetic properties and magnetocaloric effect of Cr<sub>2</sub>Ge<sub>2</sub>Te<sub>6</sub> and Cr<sub>2</sub>Si<sub>2</sub>Te<sub>6</sub> compounds: DFT combined with the Monte Carlo method.
- 85 Automated Active Space Selection with Dipole Moments.
- 84 Exploring structural, mechanical, and thermoelectric properties of half-Heusler compounds RhBiX (X = Ti, Zr, Hf): A first-principles investigation. **2023**, 13, 11513-11524
- 83 Interactions between curcumin and human salt-induced kinase 3 elucidated from computational tools and experimental methods. 14,
- 82 First-principles calculations of structural, electronic, elastic, and thermal properties of phase M<sub>2</sub>CdC (M = Sc, V, and Nb).
- 81 First-principles calculations to investigate optical properties of topological semimetal MX compounds (M = Ti, Zr, Hf and X = S, Se, Te). **2023**, 106001
- 80 Experimental and simulation study of microstructure and magnetic properties of (AlFeMnNi)<sub>1-x</sub>Ndx. 1-14
- 79 Spectral Downshifting and Passivation Effects Using 2D Perovskite (OAm)<sub>2</sub>SnBr<sub>4</sub> Films to Enhance the Properties of Si Nanowire Solar Cells.
- 78  **2016**, 121-126

- 77 Ab initio calculation of the reflectivity of molecular fluids under shock compression. **2023**, 107, ○
- 76 Non-Adiabatic Dynamics in Condensed Matter and Nanoscale Systems. **2023**, ○
- 75 Fifth Paradigm in Science: A Case Study of an Intelligence-Driven Material Design. **2023**, ○
- 74 Accurate relativistic density functional for exchange energy of atomic nuclei. **2023**, 841, 137913 ○
- 73 Controlling the carrier and phonon transport behavior of SnSe via stoichiometric adjustment. **2023**, 51, 13-21 ○
- 72 Phototunable Cell Killing by Photochromic Diarylethene of Thiazoyl and Thienyl Derivatives. ○
- 71 Giant intrinsic magnetoresistance in spin-filtered tunnel junctions with ferrimagnetic electrode. **2023**, 107, ○
- 70 On the coexistence of ferroelectric and antiferroelectric polymorphs in NaNbO<sub>3</sub> fibers at room temperature. ○
- 69 Ferroelectric order in hybrid organic-inorganic perovskite NH<sub>4</sub>PbI<sub>3</sub> with non-polar molecules and small tolerance factor. **2023**, 9, ○
- 68 Electronic Structure, Magnetism and Magnetocrystalline Anisotropy of Antiferromagnetic Semiconducting Chalcopyrite. 423, 23-31 ○
- 67 Structural, electronic properties and optical absorption of oxygen vacancy cluster defects in KDP crystals: hybrid density functional theory investigation. ○
- 66 Cation Distribution Effect on Structural and Electronic Properties of Oxygen Vacancy and Niobium Substituted Spinel Lithium Titanium Oxide Anode Material in Lithium-ion Batteries. 423, 57-66 ○
- 65 Shadow energy functionals and potentials in Born-Oppenheimer molecular dynamics. **2023**, 158, 154105 ○
- 64 Simulation and Measurement of Dielectric Constants of Crystal Alumina and Powder Alumina in Terahertz Waveband. **2022**, ○
- 63 Unraveling the structural and mechanical stability, electronic, and optical properties of (La<sub>x</sub>Sr<sub>1-x</sub>VO<sub>3</sub>)<sub>n</sub> (n = 1, 2; x = 0, 0.5, 1). **2023**, 115173 ○
- 62 Some Isomers of Nevirapine - A DFT Study. 93-109 ○
- 61 Temperature-dependent and magnetism-controlled Fermi surface changes in magnetic Weyl semimetals. **2023**, 5, ○
- 60 Electrochemical detection of arsenic (III) hazardous chemicals using cubic CsPbBr<sub>3</sub> single crystals: Structural insights from DFT study. **2023**, 115940 ○



- 59 Adsorption of sulfur on Au(111) surface: An extremely stable configuration. **2023**, 108494 ○
- 58 Acetylcholine Conformational Flexibility and Its Neutral Hydrolysis in Aqueous Solution. **2023**, 8, ○
- 57 A systematic first-principles exploration of the impact of metal doping on the electronic properties of MOF MIP-177(Ti). **2023**, 112607 ○
- 56 Magnetic Ordering in  $TlGa_{1-x}Fe_xSe_2$  Dilute Magnetic Semiconductors with Various Fe Dilution Ratios. ○
- 55 Atomic structure and peculiarities of the electronic properties of Br layers on Ag(111) at different coverages. **2023**, 122304 ○
- 54 Weak Electron-Phonon Renormalization Effect Caused by the Counteraction of the Different Phonon Vibration Modes in  $FeS_2$ . ○
- 53 The structural, electronic, and mechanical properties of FeH under extreme pressures: First-principles calculation. **2023**, 368, 115180 ○
- 52 Effect of aliovalent bismuth substitution on structure and optical properties of  $CsSnBr_3$ . **2023**, 6, ○
- 51 The influence of strong anharmonicity on high thermoelectric properties for the ternary compound  $NaMgX$  ( $X = As, Sb$ ). **2023**, 823, 140521 ○
- 50 Effect of crystal field engineering and Fermi level optimization on thermoelectric properties of  $Ge_{1.01}Te$ : Experimental investigation and theoretical insight. **2023**, 7, ○
- 49 Unconventional surface state pairs in a high-symmetry lattice with anti-ferromagnetic band-folding. **2023**, 6, ○
- 48 Environmental theoretical calculation for non-periodic systems. **2023**, ○
- 47 Sign-tunable exchange bias effect in proton-intercalated  $Fe_3GaTe_2$  nanoflakes. **2023**, 107, ○
- 46 Spectroscopic (FT-IR, FT-Raman, UV-vis and NMR) Investigation, Molecular Structure, Docking and Chemical Reactivity Elucidation of Antifungal Drug Tioconazole. 1-25 ○
- 45 Theoretical prediction by DFT on properties of  $\delta$ - $SrTa_2O_6$  crystal. **2023**, 17, 1-8 ○
- 44 Organic Stable Radical Oligomers as Spin Filters. ○
- 43 Density functional theory investigation on the structural, mechanical, lattice dynamical and thermal properties of nodal-line semimetals  $CaAgX$  ( $X: P, As$ ). **2023**, 46, ○
- 42 Orange/Red Benzo[1,2-b:4,5-b']dithiophene 1,1,5,5-Tetraoxide-Based Emitters for Luminescent Solar Concentrators: Effect of Structures on Fluorescence Properties and Device Performances. ○

- 41 Topological Insulator Bi<sub>2</sub>Se<sub>3</sub>-Assisted Heterostructure for Ultrafast Charging Sodium-Ion Batteries. ○
- 40 Amorphous Zirconia-doped Tantalum modeling and simulations using explicit multi-element spectral neighbor analysis machine learning potentials (EME-SNAP). **2023**, 7, ○
- 39 Parameter-free prediction of phase transition in PbTiO<sub>3</sub> through combination of quantum mechanics and statistical mechanics. **2023**, 232, 115480 ○
- 38 Pressure-induced transition from a Mott insulator to a ferromagnetic Weyl metal in La<sub>2</sub>O<sub>3</sub>Fe<sub>2</sub>Se<sub>2</sub>. **2023**, 14, ○
- 37 Electronic structure and optical properties of tin (IV) doped transparent perovskite crystal BaTiO<sub>3</sub> for efficient visible optoelectronic devices and solar cells. **2023**, 35, 106035 ○
- 36 Observation of gapless nodal-line states in NdSbTe. **2023**, 7, ○
- 35 Effect of an electric field on ferroelectric and piezoelectric properties of brownmillerite Ca<sub>2</sub>Al<sub>2</sub>O<sub>5</sub>. **2023**, 107, ○
- 34 Investigation of the solvent effect, regioselectivity, and the mechanism of the cycloaddition reaction between 2-chlorobenzimidazole and benzonitrile oxide. ○
- 33 DFT based comparative analysis of the physical properties of some binary transition metal carbides XC (X = Nb, Ta, Ti). **2023**, ○
- 32 Experimental investigation and thermodynamic modeling of Cu<sub>3</sub>Nb<sub>3</sub>Si system. **2023**, 33, 824-838 ○
- 31 Protection of Titanium Alloys Against High Temperature Oxidation During Closed-die Forging: Structural Analysis of the Boro-silicate Glass Coating/Ti-6Al-4V Alloy Interfacial Region by Correlative Imaging. **2023**, 111198 ○
- 30 Large intrinsic anomalous Hall effect in both Nb<sub>2</sub>FeB<sub>2</sub> and Ta<sub>2</sub>FeB<sub>2</sub> with collinear antiferromagnetism. **2023**, 107, ○
- 29 Mn(Pt<sub>1-x</sub>Pdx)<sub>5</sub>P : Isovalent tuning of Mn-sublattice magnetic order. **2023**, 107, ○
- 28 Realizing the potentials of density functional theory (DFT) and of the materials genome initiative (MGI). ○
- 27 Reconfigurable Elastic Metamaterials: Engineering Dispersion with Beyond Nearest Neighbors. **2023**, 19, ○
- 26 Hydrogen-Bonding Receptor substituted BODIPYs as Selective ON-OFF Fluorimetric Sensors for Fluoride Ions in Polar Aprotic Organic Solvents - A Molecular-level Understanding based on Experimental and Theoretical Studies. **2023**, 114780 ○
- 25 Construction of the Fast Potassiation Path in Sb<sub>x</sub>Bi<sub>1-x</sub>@NC Anode with Ultrahigh Cycling Stability for Potassium-Ion Batteries. ○
- 24 The first chrysin-based receptor for anions recognition: Experimental investigations, DFT studies and its in vitro antitumor activity. **2023**, 135637 ○

- 23 Thio/carbohydrazone derivatives from iso(thio)/cyanates: preparation, structure elucidation, DFT studies, antimicrobial activity and DNA interactions. ○
- 22 Site selective behaviour of B, C and N doping in MgO monolayers towards spintronic and optoelectronic applications. **2023**, 162, 107514 ○
- 21 Consistent wide-range equation of state of silicon by a unified first-principles method. **2023**, 107, ○
- 20  $A_2B_nPb_{n+1}$  (A = BA, PEA; B = MA; n = 1, 2): Engineering Quantum-Well Crystals for High Mass Density and Fast Scintillators. ○
- 19 Novel germanene-arsenene and germanene-antimonene lateral heterostructures: interline-dependent electronic and magnetic properties. ○
- 18 Understanding the effects of amine and morpholine adsorption on unglazed earthenware using density functional theory. **2023**, 61, 168-176 ○
- 17 Physical Mechanism of Nonlinear Spectra in Triangene. **2023**, 28, 3744 ○
- 16 A semilocal machine-learning correction to density functional approximations. **2023**, 158, ○
- 15 A Review of Trends in Corrosion-Resistant Structural Steels Research From Theoretical Simulation to Data-Driven Directions. **2023**, 16, 3396 ○
- 14 Communication: Non-adiabatic derivative coupling elements for the coupled cluster singles and doubles model. **2023**, 158, ○
- 13 Ab initio study of phononic thermal conduction in ScAgC half-Heusler. **2023**, 96, ○
- 12 Ab Initio Molecular Dynamics: A Guide to Applications. **2023**, ○
- 11 Predicting HP-HT Earth and Planetary Materials. **2023**, 131-151 ○
- 10 Inverse-perovskites  $Sc_3GaX$  (X = B, C, N): A comprehensive theoretical investigation at ambient and elevated pressures. **2023**, 35, e00808 ○
- 9 Relativistic Pseudopotentials. **2023**, ○
- 8 Novel interfacial lateral electron migration pathway formed by constructing metallized CoP<sub>2</sub>/CdS interface for excellent photocatalytic hydrogen production. **2023**, 334, 122860 ○
- 7 Unraveling complexation and separation of novel asymmetric uranyl-5-methoxy-2-(4-methoxy-6-quinazolin-2-yl)-[2,2'-bipyridin]-6-yl quinazoline to chiral fungicides R/S-metalaxyls and R/S-benalaxyls. 1-18 ○
- 6 Nonlinear Optical Materials: Predicting the First-Order Molecular Hyperpolarizability of Organic Molecular Structures. **2023**, 10, 545 ○

- 5 Theoretical insight of stabilities and optoelectronic properties of double perovskite Cs<sub>2</sub>CuIrF<sub>6</sub>: Ab-initio calculations. **2023**, 29,
- 4 ~~XXXXXXXXXX~~ Ga<sub>2</sub>O<sub>3</sub>.
- 3 Chemical Trend of Nonradiative Recombination in Cu(In,Ga)Se<sub>2</sub> Alloys. **2023**, 19,
- 2 First-principles study of SiC and GeC monolayers with adsorbed non-metal atoms. **2023**, 13, 14879-14886
- 1 Toward using collective x-ray Thomson scattering to study C<sub>H</sub> demixing and hydrogen metallization in warm dense matter conditions. **2023**, 30,