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DOI: 10.1103/physrevlett.52.997

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2272	Some remarks on the time-dependent Thomas-Fermi theory. 1985 , 322, 655-664		2
2271	The Constrained Search Formulation of Density Functional Theory. 1985 , 11-30		40
2270	Current-density-functional theory for time-dependent systems. 1985 , 31, 2682-2684		40
2269	Kohn-Sham equation for time-dependent ensembles. 1985 , 31, 3970-3971		15
2268	Hohenberg-Kohn theorem for time-dependent ensembles. 1985 , 31, 1950-1951		33
2267	Local density-functional theory of frequency-dependent linear response. <i>Physical Review Letters</i> , 1985 , 55, 2850-2852	7.4	832
2266	Time-dependent density-functional theory: Conceptual and practical aspects. <i>Physical Review Letters</i> , 1986 , 56, 1993-1995	7.4	36
2265	The probability and energy density currents as density functionals. 1986 , 19, 2549-2557		3
2264	Time-dependent density-functional theory for multicomponent systems. 1986 , 34, 529-532		33
2263	Density-functional theory for time-dependent systems. 1987 , 35, 442-444		119
2262	Time-dependent density-functional theory in the linear-response regime. <i>Physical Review Letters</i> , 1987 , 59, 2627-2630	7.4	14
2261	Density functional theory and electronic structure. 1987 , 134-145		
2260	Density-functional theory in strong magnetic fields. <i>Physical Review Letters</i> , 1987 , 59, 2360-2363	7.4	399
2259	Quantum chemistry in phase space: Some current trends. 1987 , 99, 21-28		4
2258	How can density functional theory be excited from the ground state?. 1987 , 99, 67-72		4
2257	Density-functional theory of many-electron systems subjected to time-dependent electric and magnetic fields. 1988 , 38, 1149-1158		165
2256	First-principles calculations of the optical properties of metals. 1988 , 18, 833-849		68

2255	Zeros of the frequency-dependent linear density response. <i>Physical Review Letters</i> , 1988 , 61, 1518	7.4	8
2254	Density-functional theory for the dynamical response of molecules adsorbed on metal surfaces. 1988 , 37, 10020-10028		13
2253	Dynamic linear response of many-electron systems: An integral formulation of density-functional theory. 1988 , 38, 5512-5519		17
2252	Time-dependent screening in the electron gas. 1988 , 38, 1647-1653		22
2251	Density-functional and hydrodynamical approach to ion-atom collisions through a new generalized nonlinear Schrödinger equation. 1989 , 39, 1696-1713		84
2250	Transport properties and a current-functional theory in the linear-response regime. <i>Physical Review Letters</i> , 1989 , 62, 2417-2420	7.4	21
2249	Linear optical response in silicon and germanium including self-energy effects. <i>Physical Review Letters</i> , 1989 , 63, 1719-1722	7.4	327
2248	Quantum Molecular Dynamics: a New Algorithm for Linear and Nonlinear Electron Transport in Disordered Materials. 1990 , 4, 22-33		10
2247	Time-Dependent Density-Functional Theory. 1990 , 21, 255-291		893
2246	Magnetic Fields and Density Functional Theory. 1990 , 235-253		85
2245	Simplified Green-function approximations: Further assessment of a polarization model for second-order calculation of outer-valence ionization potentials in molecules. 1991 , 44, 5773-5783		15
2244	Simulation of many-electron correlations in a resonant-tunneling diode. 1991 , 43, 9066-9069		15
2243	Density functional theory generalized to degenerate excited states. 1991 , 106, 711-719		
2242	Phase associated with the single-particle density of many-electron systems. 1991 , 40, 501-510		4
2241	Time-dependent quantum-fluid density-functional study of high-energy proton-helium collisions. 1991 , 43, 1248-1257		51
2240	Electron transport in disordered systems: A nonequilibrium quantum-molecular-dynamics approach. 1991 , 43, 10928-10932		6
2239	Quasiparticle calculation of the dielectric response of silicon and germanium. 1991 , 43, 4187-4207		199
2238	Simulation of correlated electron tunneling and a Coulomb blockade in a quantum-dot diode. 1991 , 44, 8121-8128		14

2237	Bounds to Atomic and Molecular Energy Functionals. 1991 , 22, 211-300	18
2236	Silicon plasmon resonances in the local-density approximation. 1992 , 45, 8970-8976	21
2235	Quantum-molecular-dynamics simulations of liquid metals and highly degenerate plasmas. 1992 , 4, 2044-2051	19
2234	Electron Density Distribution in Crystals: Theory Problems and Experiment. 1992 , 3, 31-102	1
2233	Ab initio simulations of sodium using time-dependent density-functional theory. 1992 , 46, 12990-13003	105
2232	Current density functional theory in a continuum and lattice lagrangians: Application to spontaneously broken chiral ground states. 1992 , 44, 359-370	2
2231	Quantum fluid dynamics of a classically chaotic oscillator. 1993 , 181, 225-231	29
2230	Photoabsorption of Fe, Co, and Ni atoms embedded in Al-clusters near 3p-ionization threshold of impurity. 1993 , 28, 215-222	1
2229	Time-dependent functional theory of coupled electron and electromagnetic fields in condensed-matter systems. 1994 , 50, 3759-3765	36
2228	Time-dependent density-functional theory for superconductors. <i>Physical Review Letters</i> , 1994 , 73, 2915-2918	56
2227	Quantum dynamics of molecular motions based on a functional method. 1994 , 50, 721-726	3
2226	New approach in the microscopic Fermi systems theory. 1994 , 249, 1-134	132
2225	Electronegativity dynamics in a chemical reaction. 1994 , 49, 705-725	26
2224	Density functional theory for Jahn-Teller systems. 1994 , 52, 809-816	4
2223	Hardness dynamics in a chemical reaction. 1994 , 217, 342-348	24
2222	A nonequilibrium time-dependent functional theory based on Liouvillean quantum field dynamics in condensed matter systems. 1994 , 195, 312-318	8
2221	A time-dependent functional theory for the electron-ion system. 1994 , 193, 363-369	5
2220	The electronic structure of magnetic transition metallic materials. 1994 , 57, 1289-1344	58

2219	Harmonic-potential theorem: Implications for approximate many-body theories. <i>Physical Review Letters</i> , 1994 , 73, 2244-2247	7.4	268
2218	Vibrational and elastic properties of the hot solid related to the static and dynamic structure of the liquid within density functional theory. 1994 , 69, 833-848		9
2217	Plasmons and dynamic screening effects in crystals. 1994 , 69, 901-910		1
2216	Sum rule for the linear density response of a driven electronic system. 1995 , 209, 206-210		38
2215	A nonequilibrium time-dependent functional theory based on Liouvillean quantum field dynamics. 1995 , 56, 389-397		1
2214	Frequency-dependent linear response of superconducting systems. 1995 , 56, 521-533		5
2213	Time-dependent quantum fluid dynamics of the photoionization of the He atom under an intense laser field. 1995 , 56, 707-732		40
2212	Nonlinear effects in interactions of swift ions with solids. 1995 , 96, 610-618		6
2211	Electronegativity and hardness profiles of a chemical process: Comparison between quantum fluid density functional theory and ab initio SCF method. 1995 , 45, 65-73		6
2210	Time-dependent density-matrix functional in Liouville space and the optical response of many-electron systems. 1995 , 52, 3601-3621		12
2209	Center of mass and relative motion in time dependent density functional theory. <i>Physical Review Letters</i> , 1995 , 74, 3233-3236	7.4	110
2208	Nonequilibrium time-dependent functional theory for coupled interacting fields. 1995 , 51, 1883-1897		10
2207	Density functional-time-dependent local density approximation calculations of autoionization resonances in noble gases. 1995 , 28, 4973-4999		63
2206	Density Functional Theory of Time-Dependent Systems. 1995 , 149-171		55
2205	Time-dependent optimized effective potential. <i>Physical Review Letters</i> , 1995 , 74, 872-875	7.4	211
2204	A density functional theory study of frequency-dependent polarizabilities and Van der Waals dispersion coefficients for polyatomic molecules. 1995 , 103, 9347-9354		301
2203	Density-Functional Approach to Atoms in Strong Laser Pulses. 1995 , 99, 488-497		69
2202	Improved density functional theory results for frequency-dependent polarizabilities, by the use of an exchange-correlation potential with correct asymptotic behavior. 1996 , 105, 3142-3151		184

2201	Density functional theory of time-dependent phenomena. 1996 , 81-172		424
2200	Current-Dependent Exchange-Correlation Potential for Dynamical Linear Response Theory. <i>Physical Review Letters</i> , 1996 , 77, 2037-2040	7.4	391
2199	Excitation energies from time-dependent density-functional theory. <i>Physical Review Letters</i> , 1996 , 76, 1212-1215	7.4	1324
2198	Popular Electronic Structure Principles in a Dynamical Context. 1996 , 100, 16126-16130		297
2197	Spin dynamics in magnets: Equation of motion and finite temperature effects. 1996 , 54, 1019-1035		254
2196	Time-Dependent Density Functional Response Theory of Molecular Systems: Theory, Computational Methods, and Functionals. 1996 , 391-439		231
2195	Generalized functional theory of interacting coupled Liouvillean Quantum Fields of condensed matter. 1996 , 173-209		1
2194	Density-functional theory of the nonlinear optical susceptibility: Application to cubic semiconductors. 1996 , 53, 15638-15642		71
2193	Spin-multiplet energies from time-dependent density functional theory. 1996 , 60, 1393-1401		28
2192	Linear response and quasiparticle calculations as probes of the Kohn-Sham eigenvalues in metals. 1996 , 60, 1457-1468		2
2191	Treatment of electronic excitations within the adiabatic approximation of time dependent density functional theory. 1996 , 256, 454-464		4711
2190	Application of time-dependent density functional response theory to Raman scattering. 1996 , 259, 599-604		53
2189	Dielectric function and plasmon excitations in silver. 1996 , 193, 231-238		6
2188	Exchange-correlation potential for the local density-functional theory of frequency-dependent linear response. 1996 , 193, K11-K14		3
2187	Determination of frequency-dependent polarizabilities using current density-functional theory. 1996 , 53, 1316-1322		79
2186	The Sham-Schlüter equation in time-dependent density-functional theory. <i>Physical Review Letters</i> , 1996 , 76, 3610-3613	7.4	54
2185	Calculation of the dielectric matrix of Si. 1997 , 9, 1225-1240		9
2184	Time-Dependent Density Functional Theory beyond Linear Response: An Exchange-Correlation Potential with Memory. <i>Physical Review Letters</i> , 1997 , 79, 1905-1908	7.4	90

2183	Time-dependent Density Functional Results for the Dynamic Hyperpolarizability of C60. <i>Physical Review Letters</i> , 1997 , 78, 3097-3100	7.4	178
2182	Density-functional theory with optimized effective potential and self-interaction correction for ground states and autoionizing resonances. 1997 , 55, 3406-3416		147
2181	Two-dimensional, two-electron model atom in a laser pulse: Exact treatment, single-active-electron analysis, time-dependent density-functional theory, classical calculations, and nonsequential ionization. 1997 , 56, 3028-3039		130
2180	Time-dependent Kohn-Sham formalism. 1997 , 55, 2630-2639		37
2179	Density functional results for isotropic and anisotropic multipole polarizabilities and C6, C7, and C8 Van der Waals dispersion coefficients for molecules. 1997 , 106, 5091-5101		100
2178	Density Functional Theory of Metal Surfaces. 1997 , 5-48		1
2177	Time-Dependent Density Functional Theory Beyond the Adiabatic Local Density Approximation. <i>Physical Review Letters</i> , 1997 , 79, 4878-4881	7.4	199
2176	Dynamics of Chemical Reactivity Indices for a Many-Electron System in Its Ground and Excited States. 1997 , 101, 7893-7900		67
2175	CrCO Photodissociation in Cr(CO) ₆ : Reassessment of the Role of Ligand-Field Excited States in the Photochemical Dissociation of Metal-Ligand Bonds. 1997 , 119, 7324-7329		82
2174	The exchange - correlation potential for current-density functional theory of frequency-dependent linear response. 1997 , 9, L475-L482		25
2173	A Quantum Chemical View of Density Functional Theory. 1997 , 101, 5383-5403		511
2172	Adiabatic and non-adiabatic cluster collisions. 1997 , 110, 1201-1208		3
2171	Dynamic exchange-correlation potentials for the 2D electron gas. 1997 , 1, 188-190		3
2170	Variation-perturbation method in time-dependent density-functional theory. 1997 , 236, 525-532		8
2169	TDLDA calculations of photoionization cross-section and asymmetry parameter profiles of alkaline-earth atoms. 1997 , 222, 197-213		12
2168	Floquet formulation of time-dependent density functional theory. 1997 , 264, 466-476		26
2167	Alternating direction implicit technique and quantum evolution within the hydrodynamical formulation of Schrödinger's equation. 1998 , 297, 247-256		15
2166	Stripped ionHelium atom collision dynamics within a time-dependent quantum fluid density functional theory. 1998 , 67, 251-271		13

2165	Exact exchange kernel for time-dependent density-functional theory. 1998 , 69, 265-277	78
2164	Quantum fluid density functional theory of time-dependent processes. 1998 , 69, 279-291	23
2163	Time-dependent density-functional theory with optimized effective potential and self-interaction correction: Application to the study of coherent control of multiple high-order harmonic generation of He atoms in mixed laser fields. 1998 , 69, 293-303	12
2162	Generalized Floquet theoretical formulation of time-dependent density functional theory for many-electron systems in multicolor laser fields. 1998 , 69, 305-315	10
2161	Femtosecond quantum fluid dynamics of helium atom under an intense laser field. 1998 , 70, 441-474	24
2160	Excited-state potential energy curves from time-dependent density-functional theory: A cross section of formaldehyde's 1A1 manifold. 1998 , 70, 933-941	194
2159	Time-dependent linear response of an inhomogeneous Bose superfluid: microscopic theory and connection to current-density functional theory. 1998 , 254, 188-201	9
2158	Quantum-mechanical interpretation of time-dependent density-functional theory. 1998 , 247, 303-308	25
2157	Relations between action integral, response functions, and causality in density functional theory. 1998 , 250, 157-162	7
2156	Density functional. Theory and application to atoms and molecules. 1998 , 298, 1-79	130
2155	Causality and Symmetry in Time-Dependent Density-Functional Theory. <i>Physical Review Letters</i> , 1998 , 80, 1280-1283	7.4 207
2154	A guided tour of time-dependent density functional theory. 1998 , 116-146	22
2153	Time-dependent density-functional theory for strong-field multiphoton processes: Application to the study of the role of dynamical electron correlation in multiple high-order harmonic generation. 1998 , 57, 452-461	122
2152	Metal-to-Ligand Charge Transfer Photochemistry: Homolysis of the Mn-Cl Bond in the mer-Mn(Cl)(CO) ₃ (diimine) Complex and Its Absence in the fac-Isomer. 1998 , 37, 6244-6254	14
2151	Collective intersubband transitions in quantum wells: A comparative density-functional study. 1998 , 58, 15756-15765	22
2150	The GW method. 1998 , 61, 237-312	1333
2149	Molecular excitation energies to high-lying bound states from time-dependent density-functional response theory: Characterization and correction of the time-dependent local density approximation ionization threshold. 1998 , 108, 4439-4449	4178
2148	Generalized Floquet formulation of time-dependent current-density-functional theory. 1998 , 58, 4749-4756	15

2147	Linear Response Calculations of Spin Fluctuations. <i>Physical Review Letters</i> , 1998 , 81, 2570-2573	7.4	170
2146	Influence of electronic exchange on single and multiple processes in collisions between bare ions and noble-gas atoms. 1998 , 58, 2063-2076		54
2145	Density-functional-theory response-property calculations with accurate exchange-correlation potentials. 1998 , 57, 2556-2571		231
2144	Calculating frequency-dependent hyperpolarizabilities using time-dependent density functional theory. 1998 , 109, 10644-10656		181
2143	Surface energy of a bounded electron gas: Analysis of the accuracy of the local-density approximation via ab initio self-consistent-field calculations. 1998 , 57, 6329-6332		60
2142	Dynamic exchange-correlation potentials for the electron gas in dimensionality D=3 and D=2. 1998 , 58, 12758-12769		56
2141	Electronic stopping power of aluminum crystal. 1998 , 58, 10307-10314		42
2140	Kohn-Sham equations for multicomponent systems: The exchange and correlation energy functional. 1998 , 57, 2146-2152		28
2139	Differential virial theorem and quantum fluid dynamics. 1998 , 58, 1779-1782		12
2138	High-energy paramagnetic spin fluctuations in nickel. 1998 , 58, 12075-12080		
2137	Time-Dependent Thomas-Fermi Approach for Electron Dynamics in Metal Clusters. <i>Physical Review Letters</i> , 1998 , 80, 5520-5523	7.4	48
2136	Exact exchange-correlation kernel for dynamic response properties and excitation energies in density-functional theory. 1998 , 57, 3433-3436		70
2135	Excitation and Relaxation in Atom-Cluster Collisions. <i>Physical Review Letters</i> , 1998 , 80, 3213-3216	7.4	70
2134	Stepwise Explosion of Atomic Clusters Induced by a Strong Laser Field. <i>Physical Review Letters</i> , 1998 , 80, 1857-1860	7.4	36
2133	Electron correlation effects in the double ionization of He. 1998 , 31, L249-L256		123
2132	Nonlinear Response of Solids and Molecules in Intense Infrared Radiation. 1998 , 538, 453		
2131	Multiphoton absorption by multielectron atoms. 1999 , 357, 1331-1358		4
2130	Time-Dependent Variational Principle in Density Functional Theory. 1999 , 217-247		8

2129	Mapping from Densities to Potentials in Time-Dependent Density-Functional Theory. <i>Physical Review Letters</i> , 1999 , 82, 3863-3866	7.4	210
2128	Theoretical prediction of the peak structure in the EELS spectrum of palladium. 1999 , 59, 15085-15092		11
2127	Incommensurate and Commensurate Antiferromagnetic Spin Fluctuations in Cr and Cr Alloys from Ab Initio Dynamical Spin Susceptibility Calculations. <i>Physical Review Letters</i> , 1999 , 82, 3340-3343	7.4	80
2126	Exchange and Correlation Kernels at the Resonance Frequency: Implications for Excitation Energies in Density-Functional Theory. <i>Physical Review Letters</i> , 1999 , 82, 4416-4419	7.4	54
2125	Density-functional approach to electron dynamics: Stable simulation under a self-consistent field. 1999 , 59, 2579-2586		186
2124	Density-functional-theory calculations of the total energies, ionization potentials, and optical response properties with the van LeeuwenBaerends potential. 1999 , 60, 3599-3603		35
2123	Time-dependent density functional calculations on the electronic absorption spectrum of free base porphin. 1999 , 111, 2499-2506		158
2122	Frequency-dependent polarizabilities, hyperpolarizabilities, and excitation energies from time-dependent density-functional theory based on the quasienergy derivative method. 1999 , 111, 2878-2888 ²²		
2121	Exact Exchange Treatment for Molecules in Finite-Basis-Set Kohn-Sham Theory. <i>Physical Review Letters</i> , 1999 , 83, 5455-5458	7.4	252
2120	Density-functional theory beyond the Hohenberg-Kohn theorem. 1999 , 59, 3359-3374		169
2119	Analysis of causality in time-dependent density-functional theory. 1999 , 60, 5101-5104		11
2118	Exchange and correlation effects beyond the LDA on the dielectric function of silicon. 1999 , 60, 14224-14233		38
2117	ELECTRON DENSITY FUNCTIONAL THEORY. 1999 , 13, 511-523		3
2116	Stopping of heavy ions in plasmas at strong coupling. 1999 , 309, 117-208		150
2115	Developments in excited-state density functional theory. 1999 , 311, 47-94		87
2114	Time-dependent density functional theory for radicals. 1999 , 302, 375-382		398
2113	Connections between the correlation potential and the static correlation kernel for two-electron densities in high-density limit. 1999 , 308, 449-455		7
2112	Implementation of time-dependent density functional response equations. 1999 , 118, 119-138		561

2111	Selective bond-breaking in molecules by intense infrared radiation. 1999 , 303, 57-64	
2110	Accurate excitation energies from time-dependent density functional theory: assessing the PBE0 model for organic free radicals. 1999 , 314, 152-157	106
2109	Time-dependent density functional theory within the TammDancoff approximation. 1999 , 314, 291-299	1363
2108	Aspects of spin dynamics and magnetic interactions. 1999 , 200, 148-166	76
2107	Excitation Energies for Transition Metal Compounds from Time-Dependent Density Functional Theory. Applications to MnO ₄ ⁻ , Ni(CO) ₄ , and Mn ₂ (CO) ₁₀ . 1999 , 103, 6835-6844	180
2106	Density- and density-matrix-based coupled KohnBham methods for dynamic polarizabilities and excitation energies of molecules. 1999 , 110, 2785-2799	118
2105	Exact exchange-correlation potential for a time-dependent two-electron system. 1999 , 59, 7876-7887	32
2104	Perturbed ground state method for electron transfer. 1999 , 111, 7818-7827	33
2103	Analytical second derivatives for excited electronic states using the single excitation configuration interaction method: theory and application to benzo[a]pyrene and chalcone. 1999 , 96, 1533-1541	49
2102	Configuration interaction singles, time-dependent HartreeBock, and time-dependent density functional theory for the electronic excited states of extended systems. 1999 , 111, 10774-10786	169
2101	Time-dependent density functional study on the electronic excitation energies of polycyclic aromatic hydrocarbon radical cations of naphthalene, anthracene, pyrene, and perylene. 1999 , 111, 8904-8912	195
2100	Density-functional calculations of static and dynamic polarizabilities of thready cylindrical crystals. 1999 , 421, 237-245	2
2099	Ab Initio Methods. 1999 , 7-138	5
2098	Density-functional correction of random-phase-approximation correlation with results for jellium surface energies. 1999 , 59, 10461-10468	112
2097	Several Theorems in Time-Dependent Density Functional Theory. <i>Physical Review Letters</i> , 1999 , 82, 378-381	73
2096	Recent New Developments of Steady-State and Time-Dependent Density Functional Theories for the Treatment of Structure and Dynamics of Many-Electron Atomic, Molecular, and Quantum Dot Systems. 1999 , 46, 361-374	1
2095	Time-dependent differential virial theorems. 2000 , 78, 341-347	23
2094	Q-Chem 2.0: a high-performance ab initio electronic structure program package. 2000 , 21, 1532-1548	588

2093	Excitation energies from time-dependent density functional theory using exact and approximate potentials. 2000 , 80, 534-554	55
2092	Excited states of molecules from Green's function perturbation techniques. 2000 , 80, 807-815	28
2091	Novel substituted 1-amino-4,5,8-naphthalenetetracarboxylic acid-1,8-lactam-4,5-imides: experimental and theoretical study. 2000 , 13, 705-712	4
2090	Coherent-state representation of reduced density matrices of correlated electronic systems. 2000 , 327, 29-37	3
2089	A TDDFT study of the electronic spectrum of s-tetrazine in the gas-phase and in aqueous solution. 2000 , 330, 152-160	186
2088	Density functional theory study of the electronic absorption spectrum of Mg-porphyrin and Mg-etioporphyrin-I. 2000 , 317, 392-399	63
2087	Comparison of the electronic excitation spectra of chlorophyll a and pheophytin a calculated at density functional theory level. 2000 , 317, 545-552	80
2086	Dynamical response of correlated electrons in solids probed by inelastic scattering experiments: an ab initio theoretical perspective. 2000 , 61, 383-390	18
2085	Nonlinear electron dynamics in metal clusters. 2000 , 337, 493-578	401
2084	Calculations of single particle spectra in density functional theory. 2000 , 269, 337-342	9
2083	Electronic structure and polarizability of quantum metallic wires. 2000 , 42, 1898-1907	8
2082	Theory of inelastic lifetimes of low-energy electrons in metals. 2000 , 251, 1-35	291
2081	Time-dependent density-functional theory investigation of excitation spectra of open-shell molecules. 2000 , 527, 229-244	55
2080	Time-dependent Kohn-Sham approach to multiple ionization. 2000 , 501-502, 315-325	49
2079	Quantum fluid density functional theory of chemical reactivity in a two-state ensemble. 2000 , 501-502, 339-352	11
2078	Molecular excitation energies from time-dependent density functional theory. 2000 , 501-502, 353-367	69
2077	Mechanisms of giant resonance in 4d photoionization of Eu. 2000 , 33, 717-725	11
2076	Time-dependent mean-field description for multiple charge-transfer processes in Ar ⁸⁺ +Ar collisions. 2000 , 62,	27

2075	Exact relations of the quasienergy functional and the exchange-correlation potential from the Floquet formulation of time-dependent density functional theory. 2000 , 63,	11
2074	Hydrodynamic approach to time-dependent density functional theory; Response properties of metal clusters. 2000 , 113, 5614-5623	36
2073	First-principles calculations of hot-electron lifetimes in metals. 2000 , 61, 13484-13492	76
2072	Time-dependent density functional calculations of molecular photoionization cross sections: N ₂ and PH ₃ . 2000 , 112, 10871-10879	60
2071	Electron correlation energies from scaled exchange-correlation kernels: Importance of spatial versus temporal nonlocality. 2000 , 61, 13431-13437	105
2070	Local spin-density approximation for spin eigenspaces and its application to the excited states of atoms. 2000 , 61,	35
2069	Charge transfer and fragmentation in cluster-atom collisions. 2000 , 61,	24
2068	An efficient method for calculating molecular excitation energies by time-dependent density-functional theory. 2000 , 113, 2088-2099	112
2067	Spin fluctuations in nearly magnetic metals from ab initio dynamical spin susceptibility calculations: Application to Pd and Cr ₉₅ V ₅ . 2000 , 62, 1075-1082	21
2066	Density-matrix representation of nonadiabatic couplings in time-dependent density functional (TDDFT) theories. 2000 , 112, 3572-3579	161
2065	Cluster explosion in an intense laser pulse: Thomas-Fermi model. 2000 , 63,	41
2064	Application of time-dependent density-functional theory to the dielectric function of various nonmetallic crystals. 2000 , 62, 7071-7083	142
2063	Circular Dichroism of Helicenes Investigated by Time-Dependent Density Functional Theory. 2000 , 122, 1717-1724	431
2062	Linear response of embedded simple metal threads. 2000 , 469, 133-143	
2061	Electron-hole excitations and optical spectra from first principles. 2000 , 62, 4927-4944	1151
2060	Accurate and efficient evolution of nonlinear Schrödinger equations. 2000 , 62,	32
2059	Efficient real-space approach to time-dependent density functional theory for the dielectric response of nonmetallic crystals. 2000 , 112, 6517-6531	85
2058	Quasiparticle Calculations in Solids. 2000 , 54, 1-218	445

2057	On the density matrix based approach to time-dependent density functional response theory. 2001 , 114, 5982-5992	349
2056	Reactivity Dynamics in AtomField Interactions: A Quantum Fluid Density Functional Study. 2001 , 105, 169-183	70
2055	Current density functional theory for optical spectra: A polarization functional. 2001 , 115, 1995-1999	68
2054	Excitation energy transfer in condensed media. 2001 , 114, 3065-3072	166
2053	Jellium surface energy beyond the local-density approximation: Self-consistent-field calculations. 2001 , 63,	61
2052	Applications of B-splines in atomic and molecular physics. 2001 , 64, 1815-1943	534
2051	KEY CONCEPTS IN TIME-DEPENDENT DENSITY-FUNCTIONAL THEORY. 2001 , 15, 1969-2023	270
2050	C60 in intense femtosecond laser pulses: Nonlinear dipole response and ionization. 2001 , 64,	40
2049	Excitation and fragmentation mechanisms in ion-fullerene collisions. <i>Physical Review Letters</i> , 2001 , 86, 5258-61	7.4 88
2048	Excitation Energies from Time-Dependent Density Functional Theory for Linear Polyene Oligomers: Butadiene to Decapentaene. 2001 , 105, 451-458	247
2047	Spin currents and spin dynamics in time-dependent density-functional theory. <i>Physical Review Letters</i> , 2001 , 87, 206403	7.4 75
2046	Time-dependent density functional theory for molecules in liquid solutions. 2001 , 115, 4708-4717	1663
2045	Theoretical approach to the design of organic molecular and polymeric nonlinear optical materials. 2001 , 63-126	56
2044	The role of exchange and correlation in time-dependent density-functional theory for photoionization. 2001 , 114, 7816-7829	21
2043	Coupled-cluster theory, pseudo-Jahn-Teller effects and conical intersections. 2001 , 115, 10382	63
2042	Time-dependent density-functional theory for molecular processes in strong fields: Study of multiphoton processes and dynamical response of individual valence electrons of N2 in intense laser fields. 2001 , 64,	90
2041	Origin of substituent effects in the absorption spectra of peroxy radicals: time dependent density functional theory calculations. 2001 , 123, 11686-94	44
2040	The Role of the S1 State of Carotenoids in Photosynthetic Energy Transfer: The Light-Harvesting Complex II of Purple Bacteria. 2001 , 105, 11016-11025	81

2039	Many-body diagrammatic expansion in a kohn-sham basis: implications for time-dependent density functional theory of excited states. <i>Physical Review Letters</i> , 2001 , 86, 2078-81	7.4	56
2038	Molecular and electronic structure of [Mn(V)N(cyclam-acetato)]PF ₆ . A combined experimental and DFT study. 2001 , 40, 4191-8		19
2037	Time-Dependent Density Functional Theory Calculations of Photoabsorption Spectra in the Vacuum Ultraviolet Region. 2001 , 105, 4953-4962		209
2036	Photoabsorption spectra in the continuum of molecules and atomic clusters. 2001 , 114, 2550-2561		78
2035	Time-dependent tight binding. 2001 , 13, 10125-10148		56
2034	Reaction dynamics of metallic clusters colliding with atoms. 2001 , 40, 305-322		
2033	Electronic spectra and ionization potentials of a stable class of closed shell polycyclic aromatic hydrocarbon cations. 2001 , 57, 931-45		32
2032	Generalized spin density functional theory for noncollinear molecular magnetism II Influence of gradient correction and self-interaction correction. 2001 , 85, 421-431		19
2031	Relativistic effects in the optical response of HgSe by time-dependent density functionals theory. 2001 , 85, 449-454		10
2030	Time-Dependent Density Functional Theory Calculations of Photoabsorption of Fluorinated Cyclic Molecules in the Vacuum Ultraviolet Region. 2001 , 226, 69-77		9
2029	Recent progress in the computational many-body theory of metal surfaces. 2001 , 137, 123-142		23
2028	Efficient algorithm for TD-Schrödinger equation and TD-Kohn-Sham equation. 2001 , 142, 255-258		1
2027	Static polarizability of excited and charged alkali metal clusters. 2001 , 43, 792-798		3
2026	Charge transfer complexes between 4,4'-disubstituted diphenyldiacetylenes: experimental and theoretical study. 2001 , 540, 123-130		8
2025	Spin Dynamics and Noncollinearity in Molecules and Crystals. 2001 , 7-23		1
2024	HYBRID ENSEMBLE METHOD FOR THE UU COLLISION TERM. 2001 , 12, 1439-1451		4
2023	TIME-DEPENDENT DENSITY FUNCTIONAL THEORY BEYOND THE ADIABATIC APPROXIMATION. 2001 , 15, 1714-1723		10
2022	SELF-ENERGY REVISION OPERATOR THEORY FOR THE MANY-BODY PROBLEM: APPLICATION TO DYNAMICAL PROPERTIES OF THE ELECTRON GAS. 2001 , 15, 2595-2610		6

2021	A Rigorous Extension of the Kohn-Sham Equation for Strongly Correlated Electron Systems. 2001 , 70, 2038-2048	21
2020	Time-dependent density-functional theory for superfluids. 2001 , 53, 162-168	3
2019	Electronic excited-state wave functions for quantum Monte Carlo: Application to silane and methane. 2001 , 114, 7795-7804	25
2018	Exchange-correlation kernels for excited states in solids. 2001 , 63,	23
2017	Sum rules and properties in time-dependent density-functional theory. 2001 , 63,	21
2016	Self-interaction-free time-dependent density-functional theory for molecular processes in strong fields: High-order harmonic generation of H ₂ in intense laser fields. 2001 , 63,	105
2015	Finite-basis-set implementation of subspace density-functional theory for excited states. 2001 , 64,	17
2014	Response of C ₆₀ and C _n to ultrashort laser pulses. 2001 , 64,	72
2013	Demonstration of initial-state dependence in time-dependent density-functional theory. 2001 , 63,	62
2012	Sharp and window resonances in the 4d photoabsorption spectrum of Eu ⁺ ions. 2001 , 64,	5
2011	Reply to Comment on Analysis of causality in time-dependent density-functional theory 2001 , 63,	2
2010	A method for ab initio nonlinear electron-density evolution. 2001 , 114, 3385-3392	42
2009	Assessment of exchange-correlation functionals for the calculation of dynamical properties of small clusters in time-dependent density functional theory. 2001 , 115, 3006-3014	143
2008	Atomic clusters submitted to an intense short laser pulse: A density-functional approach. 2001 , 65,	68
2007	Differentiation of density functionals that conserves the normalization of the density. 2001 , 63,	21
2006	Linear-response calculation of dynamical spin susceptibility in doped CaCuO ₂ . 2001 , 63,	2
2005	Inclusion of vertex corrections in the self-consistent calculation of quasiparticles in metals. <i>Physical Review Letters</i> , 2001 , 87, 226402	7.4 44
2004	Excitonic optical spectrum of semiconductors obtained by time-dependent density-functional theory with the exact-exchange kernel. <i>Physical Review Letters</i> , 2002 , 89, 096402	7.4 100

2003	Local field effects in the electron energy loss spectra of rutile TiO ₂ . <i>Physical Review Letters</i> , 2002 , 88, 037601	7-4	77
2002	Many-body diagrammatic expansion for the exchange-correlation kernel in time-dependent density functional theory. 2002 , 65,		20
2001	Photoabsorption spectra of I and its ions in the 4d region. 2002 , 65,		8
2000	Exact Kohn-Sham exchange kernel for insulators and its long-wavelength behavior. 2002 , 66,		68
1999	Efficient method for simulating quantum electron dynamics under the time-dependent Kohn-Sham equation. 2002 , 65, 036705		23
1998	Density-functional approach to obtaining excited states: Study of some open-shell atomic systems. 2002 , 65,		12
1997	Spurious interactions, and their correction, in the ensemble-Kohn-Sham scheme for excited States. <i>Physical Review Letters</i> , 2002 , 88, 033003	7-4	85
1996	The energy as a functional of the charge density and the charge-density susceptibility: A simple, exact, nonlocal expression for the electronic energy of a molecule. 2002 , 116, 5440-5447		9
1995	Quantum-fluid-dynamics approach for strong-field processes: Application to the study of multiphoton ionization and high-order harmonic generation of He and Ne atoms in intense laser fields. 2002 , 65,		21
1994	Substrate effects in the magneto-optical second-harmonic generation from first principles: Fe/Cu(001). 2002 , 65,		10
1993	Efficient methods to calculate dynamic hyperpolarizability tensors by time-dependent density-functional theory. 2002 , 116, 9624-9640		24
1992	Maitra-Burke example of initial-state dependence in time-dependent density-functional theory. 2002 , 65,		2
1991	Multiphoton above-threshold detachment of Li ⁺ Exterior-complex-scaling generalized-pseudospectral method for calculations of complex-quasienergy resonances in Floquet formulation of time-dependent density-functional theory. 2002 , 66,		18
1990	A NONADIABATIC QUANTUM MECHANICAL MONTE CARLO METHOD. 2002 , 13, 909-915		
1989	A COMPUTATIONAL DESIGN OF SI-BASED DIRECT BAND-GAP MATERIALS. 2002 , 16, 4279-4284		1
1988	Excitation energies from time-dependent density-functional formalism for small systems. 2002 , 57, 683-689		27
1987	Quantum fluid dynamics approach for electronic structure calculation: application to the study of ground-state properties of rare gas atoms. 2002 , 35, 2075-2086		10
1986	Application of the subspace density functional theory to the excitation energies of molecules. 2002 , 35, 2313-2324		15

1985	What is Time-Dependent Density Functional Theory? Successes and Challenges. 2002 , 285-298		
1984	Generalized Floquet Formulation of Time-Dependent Density Functional Theory for Multiphoton Processes in Intense Laser Fields. 2002 , 49, 737-750		
1983	Time-Dependent Density Functional Theory Investigation of the Ground and Excited States of Coumarins 102, 152, 153, and 343. 2002 , 106, 12117-12123		148
1982	Memory in time-dependent density functional theory. <i>Physical Review Letters</i> , 2002 , 89, 023002	7.4	137
1981	Vibrational and Electronic Spectroscopy of the Fluorene Cation. 2002 , 106, 63-73		38
1980	Time-dependent current-density-functional theory for the linear response of weakly disordered systems. 2002 , 65,		64
1979	Calculation of the optical response of atomic clusters using time-dependent density functional theory and local orbitals. 2002 , 66,		161
1978	Electronic excitations: density-functional versus many-body Green's-function approaches. 2002 , 74, 601-659		2843
1977	FIRST PRINCIPLES MOLECULAR DYNAMICS INVOLVING EXCITED STATES AND NONADIABATIC TRANSITIONS. 2002 , 01, 319-349		125
1976	Ab initiomolecular dynamics: basic concepts, current trends and novel applications. 2002 , 14, R1297-R1355		187
1975	Combined Spectroelectrochemical and Theoretical Study of a Vinylene-Bridged Sexithiophene Cooligomer: Analysis of the π -Electron Delocalization and of the Electronic Defects Generated upon Doping. 2002 , 106, 3872-3881		60
1974	Exchange-correlation kernel in time-dependent density functional theory. 2002 , 66,		26
1973	Excitonic effects in solids described by time-dependent density-functional theory. <i>Physical Review Letters</i> , 2002 , 88, 066404	7.4	248
1972	Atomistic simulations of complex materials: ground-state and excited-state properties. 2002 , 14, 3015-3047		384
1971	First-principles density-functional calculations for optical spectra of clusters and nanocrystals. 2002 , 65,		147
1970	Optical absorption spectra of semiconductors and insulators including electron-hole correlations: An ab initio study within the LAPW method. 2002 , 66,		113
1969	Ab Initio calculations of the anisotropic dielectric tensor of GaAs/AlAs superlattices. <i>Physical Review Letters</i> , 2002 , 89, 216803	7.4	34
1968	Time-dependent independent-particle model calculation of multiple capture and ionization processes in pAr, pAr, and He2+Ar collisions. 2002 , 66,		53

1967	Correlation in time-dependent density-functional theory. 2002 , 117, 72-81		41
1966	Quinonoid oligothiophenes as electron-donor and electron-acceptor materials. A spectroelectrochemical and theoretical study. 2002 , 124, 12380-8		101
1965	Theoretical determination of chromophores in the chromogenic effects of aromatic neurotoxicants. 2002 , 124, 2744-52		35
1964	The virtual chemistry lab for reactions at surfaces: Is it possible? Will it be useful?. 2002 , 500, 347-367		70
1963	Density-functional theory of linear and nonlinear time-dependent molecular properties. 2002 , 117, 9630-9645	338	
1962	Theoretical Investigation of the Ground and Excited States of Coumarin 151 and Coumarin 120. 2002 , 106, 9294-9305		148
1961	Ab initio study of field emission from graphitic ribbons. <i>Physical Review Letters</i> , 2002 , 88, 127601	7.4	101
1960	First-principles computation of material properties: the ABINIT software project. 2002 , 25, 478-492		2556
1959	Ultranonlocality in time-dependent current-density-functional theory: application to conjugated polymers. <i>Physical Review Letters</i> , 2002 , 88, 186401	7.4	203
1958	Electronic Excitations in Pyrrole: A Test Case for Determination of Chromophores in the Chromogenic Effects of Neurotoxic Hydrocarbons by Time-Dependent Density Functional Theory and Single-Excitation Configuration Interaction Methods. 2002 , 216, 81-89		12
1957	Current correlation functions of ideal fermi gas at finite temperature. 2002 , 58, 703-711		3
1956	Tight-binding methods for transport and optical properties in realistic nanostructures. 2002 , 314, 211-219		20
1955	Theory of field evaporation and field emission from carbon nanotubes. 2002 , 323, 214-215		2
1954	Time-dependent density functional theory calculations of molecular static and dynamic polarizabilities, cauchy coefficients and their anisotropies with atomic numerical basis functions. 2002 , 591, 255-266		5
1953	Ab-initio study of the electromagnetic response and polarizability properties of carbon chains. 2002 , 357, 459-513		31
1952	Quantum electrodynamical density-matrix functional theory and group theoretical consideration of its solution. 2002 , 90, 273-281		4
1951	Time-dependent density functional theory simulation of UV/visible absorption spectra of zirconocene catalysts. 2002 , 354, 449-457		56
1950	Excitation and dissipation of interacting many-electron system. 2002 , 355, 400-404		5

1949	Photoionization of oriented molecules: a time dependent density functional approach. 2002 , 356, 153-160	15
1948	On the Floquet formulation of time-dependent density functional theory. 2002 , 359, 237-240	18
1947	Non-adiabatic couplings by time-dependent density functional theory. 2002 , 364, 75-79	88
1946	Electronic spectra of indolyl radicals: a time-dependent DFT study. 2002 , 365, 15-21	27
1945	Anisotropy and interplane interactions in the dielectric response of graphite. <i>Physical Review Letters</i> , 2002 , 89, 076402	7.4 106
1944	Self-energy-functional approach: Analytical results and the Mott-Hubbard transition. 2003 , 36, 335-348	148
1943	Non-adiabatic quantum molecular dynamics: General formalism and case study H ₂ + in strong laser fields. 2003 , 25, 15-24	48
1942	Time dependent density functional theory study of charge-transfer and intramolecular electronic excitations in acetone-water systems. 2003 , 119, 12417-12431	128
1941	Density functional theory and multiscale materials modeling. 2003 , 26, 3-12	4
1940	Chemical reactivity of the compressed noble gas atoms and their reactivity dynamics during collisions with protons. 2003 , 115, 195-218	11
1939	Influence of the dynamical correlations on the ionization of irradiated metal clusters. 2003 , 205, 350-354	
1938	QM/MM Car-Parrinello molecular dynamics study of the solvent effects on the ground state and on the first excited singlet state of acetone in water. 2003 , 4, 1177-82	105
1937	A dynamic mean field theory for dissipative interacting many-electron systems: Markovian formalism and its implementation. 2003 , 24, 2083-92	3
1936	Gas phase photoisomerization of urocanic acid: a theoretical study. 2003 , 370, 625-630	10
1935	Time dependent density functional theory of core electrons excitations. 2003 , 373, 115-123	202
1934	Calculations of two-photon absorption cross sections by means of density-functional theory. 2003 , 374, 446-452	121
1933	Another important coordinate in the photoisomerization of cis-stilbene. 2003 , 378, 323-329	31
1932	Time-dependent density functional theory calculations of the photoabsorption of fluorinated alkanes. 2003 , 122, 27-35	18

1931	Absorption spectra of germanium nanocrystals. 2003 , 127, 361-365	26
1930	Optical excitations in small hydrogenated silicon clusters: comparison of theory and experiment. 2003 , 239, 19-25	16
1929	Breathing modes of 2-D quantum dots with elliptical shape in magnetic fields. 2003 , 91, 483-489	5
1928	Dual transformation for non-BornOppenheimer time-dependent density functional theory. 2003 , 91, 105-112	6
1927	Ab initio electrical conductance of a molecular wire. 2003 , 91, 524-532	47
1926	Ground- and excited-states reactivity dynamics of hydrogen and helium atoms. 2003 , 91, 633-650	27
1925	Exact expressions for energy functional in the time-dependent density functional theory. 2003 , 92, 229-233	2
1924	Corollary to the HohenbergKohn theorem. 2003 , 95, 387-393	11
1923	Specific features of determination of the energy of excited electronic states in the formalism of density functional theory. 2003 , 95, 25-28	1
1922	Linear-scaling time-dependent density-functional theory. 2003 , 68,	81
1921	Photodegradation of Sulfonylurea Molecules: Analytical and Theoretical DFT Studies. 2003 , 107, 10032-10038	11
1920	Excitations in time-dependent density-functional theory. <i>Physical Review Letters</i> , 2003 , 90, 043005	7.4 148
1919	Electronic excitation energies of Zn(i)O(i) clusters. 2003 , 125, 9494-9	89
1918	Low-Lying Excited States of Light-Harvesting System II in Purple Bacteria. 2003 , 107, 9589-9600	13
1917	Time-Dependent Density Functional Study of the Electronic Excited States of Polycyclic Aromatic Hydrocarbon Radical Ions. 2003 , 107, 4940-4951	86
1916	Electronic Absorption Spectra of Neutral Perylene (C ₂₀ H ₁₂), Terrylene (C ₃₀ H ₁₆), and Quaterylene (C ₄₀ H ₂₀) and Their Positive and Negative Ions: Ne Matrix-Isolation Spectroscopy and Time-Dependent Density Functional Theory Calculations. 2003 , 107, 3660-3669	134
1915	Ab Initio and Density Functional Study of the Electronic Transitions of Indoline and Indoline-2-Carboxylic Acid. 2003 , 107, 5670-5680	14
1914	Current-density functional theory of the response of solids. 2003 , 68,	82

1913	Spectroscopic and Theoretical Study of PushPull Chromophores Containing Thiophene-Based Quinonoid Structures as Electron Spacers. 2003 , 107, 12175-12183	34
1912	Computational Insights into the Chemical Structures and Mechanisms of the Chromogenic and Neurotoxic Effects of Aromatic Ediketones. 2003 , 107, 2853-2861	21
1911	Excited state nuclear forces from the TammDancoff approximation to time-dependent density functional theory within the plane wave basis set framework. 2003 , 118, 3928-3934	154
1910	The spinflip approach within time-dependent density functional theory: Theory and applications to diradicals. 2003 , 118, 4807-4818	489
1909	Optical and loss spectra of carbon nanotubes: depolarization effects and intertube interactions. <i>Physical Review Letters</i> , 2003 , 91, 046402	7.4 166
1908	Nitro-functionalized oligothiophenes as a novel type of electroactive molecular material: spectroscopic, electrochemical, and computational study. 2003 , 125, 2524-34	101
1907	Time-dependent density-functional calculations for the optical spectra of molecules, clusters, and nanocrystals. 2003 , 15, R1517-R1547	56
1906	Self-consistent mean-field models for nuclear structure. 2003 , 75, 121-180	1666
1905	HSAB principle applied to the time evolution of chemical reactions. 2003 , 125, 2705-10	142
1904	Electronic structure of the trimethylenemethane diradical in its ground and electronically excited states: Bonding, equilibrium geometries, and vibrational frequencies. 2003 , 118, 6874-6883	83
1903	A New, Self-Contained Asymptotic Correction Scheme To Exchange-Correlation Potentials for Time-Dependent Density Functional Theory. 2003 , 107, 10154-10158	100
1902	The electronic spectroscopy of transition metal di-hydrides H ₂ M(CO) ₄ (M = Fe, Os): a theoretical study based on CASSCF/MS-CASPT2 and TD-DFT. 2003 , 5, 2948-2953	13
1901	Solution of Poisson's equation for finite systems using plane-wave methods. 2003 , 81, 1151-1164	34
1900	A discrete solvent reaction field model for calculating molecular linear response properties in solution. 2003 , 119, 3800-3809	66
1899	Localized-density-matrix implementation of time-dependent density-functional theory. 2003 , 119, 8794-8803	47
1898	Restricted density functional theory of linear time-dependent properties in open-shell molecules. 2003 , 119, 34-46	90
1897	Density Functional Theories and Self-energy Approaches. 2003 , 185-217	4
1896	Theoretical and experimental study of the dynamical electronic response of Ag. 2003 , 68,	6

1895	Ab initio calculation of the exchange-correlation kernel in extended systems. 2003 , 68,		71
1894	Density functional response approach for the linear and nonlinear electric properties of molecules. 2003 , 118, 10861-10866		18
1893	Resonant nonlinear polarizabilities in the time-dependent density functional theory. 2003 , 119, 8809-8823		137
1892	Spin virial theorem in the time-dependent density-functional theory. 2003 , 68,		4
1891	Macroscopic and microscopic components of exchange-correlation interactions. 2003 , 68,		31
1890	Time-Dependent Density Functional Theory. 2003 , 144-184		50
1889	Excited states of the 3d transition metal monoxides. 2003 , 118, 9608-9613		54
1888	Accurate Rydberg excitations from the local density approximation. <i>Physical Review Letters</i> , 2003 , 91, 263001	7-4	37
1887	Long-range behavior and frequency dependence of exchange-correlation kernels in solids. 2003 , 67,		45
1886	Dynamical ionization ignition of clusters in intense short laser pulses. 2003 , 68,		29
1885	Bound excitons in time-dependent density-functional theory: optical and energy-loss spectra. <i>Physical Review Letters</i> , 2003 , 91, 256402	7-4	136
1884	Parameter-free calculation of response functions in time-dependent density-functional theory. <i>Physical Review Letters</i> , 2003 , 91, 056402	7-4	141
1883	A discrete solvent reaction field model for calculating frequency-dependent hyperpolarizabilities of molecules in solution. 2003 , 119, 12998-13006		41
1882	Local exchange-correlation vector potential with memory in time-dependent density functional theory: The generalized hydrodynamics approach. 2003 , 67,		24
1881	Application of time-dependent current-density-functional theory to nonlocal exchange-correlation effects in polymers. 2003 , 118, 1044-1053		101
1880	Laser-driven electron transfer through metal-insulator-metal contacts: Time-dependent configuration interaction singles calculations for a jellium model. 2003 , 68,		79
1879	A theoretical investigation of valence and Rydberg electronic states of acrolein. 2003 , 119, 12323-12334		80
1878	Ionization and high-order harmonic generation in aligned benzene by a short intense circularly polarized laser pulse. 2003 , 68,		61

1877	Quasiparticle density-matrix representation of nonlinear time-dependent density-functional response functions. 2003 , 67,	21
1876	Exchange-correlation potentials in the adiabatic connection fluctuation-dissipation framework. 2003 , 68,	74
1875	Role of exchange interaction in Coulomb quantum kinetics. 2003 , 67,	12
1874	Theoretical Study of Photosynthetic Light-Harvesting Processes: Application of Time-Dependent Density Functional Theory. 2003 , 50, 745-756	4
1873	Many-Electron Problem in Terms of the Density: From Thomas-Fermi to Modern Density-Functional Theory. 2003 , 02, 301-322	9
1872	Density Functional Theory versus the Hartree-Fock Method: Comparative Assessment. 2003 , 68, C133-C140	4
1871	Time-Dependent Density Functional Theory in Atomic Collisions. 2003 , 205-220	6
1870	Time-dependent density functional theory study of the electronic states of BI. 2003 , 36, 2283-2290	3
1869	Description of Dynamic Properties of Finite Electron Systems in Density Functional Theory. 2003 , 68, C10-C17	1
1868	Exploring dynamical magnetism with time-dependent density-functional theory: From spin fluctuations to Gilbert damping. 2003 , 61, 354-360	16
1867	Dynamics of molecule-surface interactions from first principles. 2003 , 11, 1-26	5
1866	Time-dependent Density Functional Resonance Theory. 2003 , 511-517	5
1865	A density-functional approach to fermionization in the 1D Bose gas. 2004 , 37, S287-S300	23
1864	The Bethe-Salpeter equation: a first-principles approach for calculating surface optical spectra. 2004 , 16, S4313-S4322	23
1863	Time-dependent density-functional theory approach to nonlinear particle-solid interactions in comparison with scattering theory. 2004 , 16, 8621-8631	4
1862	Theory of time-resolved photoelectron imaging. Comparison of a density functional with a time-dependent density functional approach. 2004 , 120, 1172-80	5
1861	Calculations of static and dynamic polarizabilities of excited states by means of density functional theory. 2004 , 121, 7595-600	28
1860	Exchange and correlation effects in the relaxation of hot electrons in noble metals. 2004 , 69,	20

1859	Time-dependent exchange-correlation current density functionals with memory. 2004 , 121, 8731-41	43
1858	Double excitations within time-dependent density functional theory linear response. 2004 , 120, 5932-7	372
1857	Potential-energy surfaces for excited states in extended systems. 2004 , 120, 4593-602	70
1856	Ab initio study of the alternating current impedance of a molecular junction. 2004 , 120, 3387-96	112
1855	Conductance of molecular wires and transport calculations based on density-functional theory. 2004 , 69,	176
1854	Singlet-triplet transitions in three-atomic molecules studied by time-dependent MCSCF and density functional theory. 2004 , 102, 1391-1406	22
1853	Optical excitation and absorption spectra of C ₅₀ Cl ₁₀ . 2004 , 121, 2849-51	23
1852	Real-time study of the adiabatic energy loss in an atomic collision with a metal cluster. 2004 , 121, 6341-5	28
1851	Theoretical investigation of substituted anthraquinone dyes. 2004 , 121, 1736-43	102
1850	Photodesorption of oxygen from carbon nanotubes. 2004 , 70,	18
1849	High-order harmonic generation from silver clusters: Laser-frequency dependence and the screening effect of d electrons. 2004 , 70,	36
1848	Excitation energies for a benchmark set of molecules obtained within time-dependent current-density functional theory using the Vignale-Kohn functional. 2004 , 120, 8353-63	40
1847	Lack of Hohenberg-Kohn theorem for excited states. <i>Physical Review Letters</i> , 2004 , 93, 173001	7.4 69
1846	Investigation of photoinduced electron transfer in model system of vitamin E-duroquinone by time-dependent density functional theory. 2004 , 120, 10025-32	8
1845	Mapping from current densities to vector potentials in time-dependent current density functional theory. 2004 , 70,	89
1844	Analytic second variational derivative of the exchange-correlation functional. 2004 , 69,	7
1843	Long-wavelength behavior of the dynamical spin-resolved local-field factor in a two-dimensional electron liquid. 2004 , 70,	2
1842	Excitonic effects in time-dependent density-functional theory: An analytically solvable model. 2004 , 70,	35

1841	Density functional calculation of the electronic absorption spectrum of Cu ⁺ and Ag ⁺ aqua ions. 2004 , 121, 11885-99	40
1840	Adiabatic connection for near degenerate excited states. 2004 , 69,	22
1839	Time-dependent density-functional calculations with asymptotically correct exchange-correlation potentials. 2004 , 69,	15
1838	Density-functional theory in one dimension for contact-interacting fermions. 2004 , 70,	52
1837	Electronic excitations of stable fullerene-like GaP clusters. 2004 , 854, U3.4.1	
1836	Optical Responses of Conjugated Polymers by TDDFT in Real-Space and Real-Time Approach. 2004 , 846, DD2.3.1	4
1835	Femtosecond spectroscopy in semiconductors: a key to coherences, correlations and quantum kinetics. 2004 , 67, 433-512	157
1834	Potential Energy Curves for the Isomerization of Protonated Schiff Base of Retinal on the Triplet State Surface. 2004 , 30, 917-921	1
1833	Quantal Density Functional Theory. 2004 , 49-98	
1832	Excited states dynamics in time-dependent density functional theory. 2004 , 28, 211-218	114
1831	Ionization dynamics of simple metal clusters in intense fields by the Thomas-Fermi-Vlasov method. 2004 , 29, 367-378	64
1830	Magnetic susceptibility, exchange interactions and spin-wave spectra in the local spin density approximation. 2004 , 16, 7439-7446	45
1829	Time-dependent density functional theory. 2004 , 55, 427-55	932
1828	Atomistic theory of transport in organic and inorganic nanostructures. 2004 , 67, 1497-1561	246
1827	Ab initio study of the optical absorption and wave-vector-dependent dielectric response of graphite. 2004 , 69,	159
1826	Photon-assisted tunneling versus tunneling of excited electrons in metal-insulator-metal junctions. 2004 , 78, 189-199	26
1825	Optical absorption and electron energy loss spectra of carbon and boron nitride nanotubes: a first-principles approach. 2004 , 78, 1157-1167	99
1824	Dynamics of femtosecond laser interactions with dielectrics. 2004 , 79, 1695-1709	320

1823	The application of the single-channel random phase approximation to radiative properties of dense He and Li plasmas. 2004 , 83, 83-92	5
1822	Ionization process of the hydrogen atom in intense laser fields: NonBorn-Oppenheimer 1D model calculations. 2004 , 97, 891-895	2
1821	Ground-state electronic energies and densities of atomic systems in strong magnetic fields through a time-dependent hydrodynamical equation. 2004 , 97, 701-712	14
1820	On Ehrenfest's theorem. 2004 , 97, 953-960	1
1819	A theoretical study of solvent effects on tautomerism and electronic absorption spectra of 3-hydroxy-2-mercaptopyridine and 2,3-dihydroxypyridine. 2004 , 25, 1833-9	26
1818	Field emission and electronic structures of carbon allotropes. 2004 , 464-465, 354-359	4
1817	Theoretical study of the external heavy atom effect on phosphorescence of free-base porphin molecule. 2004 , 60, 3213-24	30
1816	Time-dependent density functional theory of excitation energies of closed-shell quantum dots. 2004 , 22, 486-489	2
1815	Beyond the Floquet theorem: generalized Floquet formalisms and quasienergy methods for atomic and molecular multiphoton processes in intense laser fields. 2004 , 390, 1-131	309
1814	Experimental and ab initio calculational studies on 2,3-diketo-benzopiperazine. 2004 , 705, 133-139	5
1813	A dressed TDDFT treatment of the 21Ag states of butadiene and hexatriene. 2004 , 389, 39-42	168
1812	Rotational aspects of short-pulse population transfer in diatomic molecules. 2004 , 392, 23-27	11
1811	Calculations of the third-order nonlinear optical responses in pushpull chromophores with a time-dependent density functional theory. 2004 , 392, 444-451	38
1810	HartreeBock exchange in time dependent density functional theory: application to charge transfer excitations in solvated molecular systems. 2004 , 394, 141-146	49
1809	Remarkable solvent effects on first hyperpolarizabilities of zwitterionic merocyanine dyes: ab initio TD-DFT/PCM approach. 2004 , 395, 269-273	38
1808	Calculation of the optical spectrum of the Ti8C12 and V8C12 Met-Cars. 2004 , 398, 292-296	9
1807	Time-dependent four-component relativistic density functional theory for excitation energies. 2004 , 121, 6658-66	85
1806	Excitation energies of some d1 systems calculated using time-dependent density functional theory: an implementation of open-shell TDDFT theory for doubletdoublet excitations. 2004 , 102, 2585-2595	54

1805	W(CO)5-pyridine π -acceptor complexes: theoretical calculations and a laser photolysis study. 2004 , 28, 1514-1519	9
1804	Using on-top pair density for construction of correlation functionals for multideterminant wave functions. 2004 , 102, 2207-2216	46
1803	Band-gap energy in the random-phase approximation to density-functional theory. 2004 , 70,	46
1802	Excitation energies of π -conjugated oligomers within time-dependent current-density-functional theory. 2004 , 121, 10707-14	20
1801	Quasiparticle effects and optical absorption in small fullerenelike GaP clusters. 2004 , 70,	21
1800	Testing the multi-configuration time-dependent Hartree-Fock method. 2004 , 37, 763-773	97
1799	Structures and Properties of Polyphosphinoborane: an Oligomeric Theoretical Study. 2004 , 37, 1009-1015	28
1798	Regioselectivity in the Chemical Reactions between Molecules and Protons: A Quantum Fluid Density Functional Study. 2004 , 108, 658-664	18
1797	First Principles Treatment of Configuration Optimizations, Excited-State Properties, and Dynamic Third-Order Polarizabilities of Chloro-Metal Phthalocyanines MPcCl (M = Al, Ga, In). 2004 , 108, 1837-1843	13
1796	Experimental and Theoretical Studies of the Electronic Spectra of Mixed Framework Phosphates of Zr and Co. 2004 , 108, 6969-6980	3
1795	The Photophysics of Free-Base Hemiporphyrizine: A Theoretical Study. 2004 , 108, 3926-3931	10
1794	Excitation energies from time-dependent density-functional theory beyond the adiabatic approximation. 2004 , 121, 28-35	60
1793	First Principle Treatments on Site- and Size-Dependent Supramolecular Interactions and Nonlinear Optical Properties of Polymer of 2-Methyl-4-Nitroaniline. 2004 , 108, 12658-12664	19
1792	VIS/NIR Absorption Spectra of Positively Charged Oligo(phenylenevinylene)s and Comparison with Time-Dependent Density Functional Theory Calculations. 2004 , 108, 19967-19975	13
1791	Effects of Chain Substitution on the Structures and Properties of Polyphosphinoborane. 2004 , 37, 5040-5046	3
1790	Electronic structure and solvation of copper and silver ions: a theoretical picture of a model aqueous redox reaction. 2004 , 126, 3928-38	183
1789	Chromogenic and Neurotoxic Effects of an Aliphatic β -Diketone: Computational Insights into the Molecular Structures and Mechanism. 2004 , 108, 6098-6104	18
1788	Enhanced Absorption Induced by a Metallic Nanoshell. 2004 , 4, 85-88	68

1787	Long-range contribution to the exchange-correlation kernel of time-dependent density functional theory. 2004 , 69,	155
1786	Excitations, optical absorption spectra, and optical excitonic gaps of heterofullerenes. I. C60, C59N+, and C48N12: theory and experiment. 2004 , 120, 5133-47	44
1785	Time-dependent partition-free approach in resonant tunneling systems. 2004 , 69,	249
1784	Band structure versus dynamical exchange-correlation effects in surface plasmon energy and damping: a first-principles calculation. <i>Physical Review Letters</i> , 2004 , 93, 176801	7.4 50
1783	Bonding Patterns in Benzene Triradicals from Structural, Spectroscopic, and Thermochemical Perspectives. 2004 , 108, 6581-6588	65
1782	Effects beyond the random-phase approximation in calculating the interaction between metal films. 2004 , 70,	64
1781	Tuning spectral properties of fullerenes by substitutional doping. 2004 , 69,	34
1780	Electronic modulation of dithienothiophene (DTT) as pi-center of D-pi-D chromophores on optical and redox properties: analysis by UV-Vis-NIR and Raman spectroscopies combined with electrochemistry and quantum chemical DFT calculations. 2004 , 126, 13363-76	48
1779	Ab Initio Investigation of the Structures and Properties of Polyaminoborane. 2004 , 108, 9616-9624	19
1778	Kinetic theory of quantum transport at the nanoscale. 2004 , 70,	41
1777	Time-dependent quantum transport: An exact formulation based on TDDFT. 2004 , 67, 14-20	113
1776	Asymmetry of above-threshold ionization of metal clusters in two-color laser fields: A time-dependent density-functional study. 2004 , 69,	28
1775	Electronic Spectroscopy and Photoreactivity of Transition Metal Complexes: Quantum Chemistry and Wave Packet Dynamics. 119-165	7
1774	Excitation Energies of Metal Complexes with Time-dependent Density Functional Theory. 2004 , 49-116	69
1773	Progress in classical and quantum variational principles. 2004 , 67, 159-208	23
1772	Electronic structure and electron energy-loss spectroscopy of ZrO2 zirconia. 2004 , 70,	121
1771	Transport in nanoscale systems: the microcanonical versus grand-canonical picture. 2004 , 16, 8025-8034	97
1770	Accurate and efficient non-adiabatic quantum molecular dynamics approach for laser-matter interactions. 2004 , 37, 2883-2901	16

1769	Effects of the crystal structure in the dynamical electron density-response of hcp transition metals. 2004 , 30, 104-109	5
1768	The Performance of Hybrid Density Functionals in Solid State Chemistry. 2004 , 171-232	164
1767	Combined Spectroscopic and Theoretical Study of Narrow Band Gap Heterocyclic Co-oligomers Containing Alternating Aromatic Donor and Quinoid Acceptor Units. 2004 , 108, 2516-2526	61
1766	Electronic, Structural, and Optical Properties of Conjugated Polymers Based on Carbazole, Fluorene, and Borafluorene. 2004 , 108, 3123-3129	67
1765	Field emission mechanisms of graphitic nanostructures. 2004 , 70,	51
1764	Exact-exchange density-functional calculations for noble-gas solids. 2004 , 69,	65
1763	Electronic absorption spectra of PAHs up to vacuum UV. 2004 , 426, 105-117	89
1762	Restricted Density Functional Response Theory for Open-Shell Systems. 2005 , 50, 271-288	1
1761	Theoretical electron affinities of PAHs and electronic absorption spectra of their mono-anions. 2005 , 432, 585-594	50
1760	Theoretical spectral properties of PAHs: towards a detailed model of their photophysics in the ISM. 2005 , 6, 178-184	3
1759	Theoretical Study on Acylacetanilide Azomethine Dyes: A Relationship between Electronic Absorption Properties and Molecular Structures. 2005 , 78, 1929-1938	4
1758	Electronic excitations in NiO: LSDA+U-based calculations vs. inelastic X-ray scattering and ellipsometry measurements. 2005 , 66, 2281-2289	7
1757	Spectroscopy and photophysics of flavin-related compounds: 3-ethyl-lumiflavin. 2005 , 170, 267-272	13
1756	Long and short time quantum dynamics: I. Between Green's functions and transport equations. 2005 , 29, 154-174	25
1755	Theoretical methods that help understanding the structure and reactivity of gas phase ions. 2005 , 240, 37-99	95
1754	Computational and spectroscopic studies on luminescence of [Ag(PPh ₃) ₂ (NMP)]NO ₃ . 2005 , 751, 133-138	13
1753	Time-dependent density functional theory calculations for collisions of bare ions with helium. 2005 , 233, 240-243	11
1752	TD-DFT studies on the electronic structure of imidazole bound vanadate in vanadium containing haloperoxidases (VHPO). 2005 , 725, 163-175	29

1751	Theoretical investigation of the absorption spectrum of thioindigo dyes. 2005 , 731, 67-72	41
1750	A TDDFT study of the low-lying excitation energies of polycyclic cinnolines and their carbocyclic analogues. 2005 , 732, 21-32	2
1749	Electronic and optical properties of functionalized carbon chains with the localized Hartree-Fock and conventional Kohn-Sham methods. 2005 , 309, 77-87	64
1748	Linear response at the 4-component relativistic density-functional level: application to the frequency-dependent dipole polarizability of Hg, AuH and PtH ₂ . 2005 , 311, 187-201	66
1747	Theoretical DFT study of phosphorescence from porphyrins. 2005 , 315, 215-239	84
1746	A joint theoretical and experimental study of phenylene-ethynylene molecular wires. 2005 , 401, 149-156	61
1745	Multiple ionization of a silver diatomic molecule in an intense laser field. 2005 , 404, 365-369	5
1744	Time-dependent density functional approach to chemical reactions induced by electronic double excitations. 2005 , 405, 234-239	9
1743	Substitution effects on the visible spectra of 1,4-diNHP-9,10-anthraquinones. 2005 , 405, 429-433	51
1742	A TD-DFT study of the absorption spectra of fast dye salts. 2005 , 410, 254-259	105
1741	Theoretical investigations of the UV spectra of coumarin derivatives. 2005 , 415, 20-24	48
1740	Importance of electronic self-consistency in the TDDFT based treatment of nonadiabatic molecular dynamics. 2005 , 35, 467-477	44
1739	Single-reference ab initio methods for the calculation of excited states of large molecules. 2005 , 105, 4009-37	2042
1738	Multidisciplinary physicochemical analysis of oligothiophenes end-capped by nitriles: electrochemistry, UV-vis-near-IR, IR, and Raman spectroscopies and quantum chemistry. 2005 , 109, 10115-25	40
1737	Spin dependence of density functionals that respect spin-rotational invariance. 2005 , 294, e17-e20	
1736	Computations on the A-X transition of isoprene-OH-O ₂ peroxy radicals. 2005 , 26, 836-45	3
1735	Experimental and Density Functional Theory Computational Studies on 2,3-Diaryl-tetrazole-5-thione. 2005 , 23, 548-556	2
1734	Dynamic evolution of Kohn-Sham electron density in the real-time domain with finite basis expansion. 2005 , 6, 655-62	3

1733	Regioselectivity in the [2 + 2] cyclo-addition reaction of triplet carbonyl compounds to substituted alkenes (Paterno-B�hi reaction): A spin-polarized conceptual DFT approach. 2005 , 117, 561-571	19
1732	Theoretical and Experimental Studies on a Metallorganic Complex of (Isopropylxanthato)(Phenyl)Mercury(II) [Hg(II)(C ₆ H ₅)(C ₄ H ₇ OS ₂)]. 2005 , 16, 469-474	3
1731	Experimental Studies on Isonicotinato Cadmium(II) Complex [Cd(C ₆ H ₄ NO ₂) ₂ (H ₂ O) ₄] and Density Functional Calculations. 2005 , 16, 529-533	7
1730	Experimental and Theoretical Studies on (p-methoxyphenyl)thiosemicarbazide. 2005 , 16, 361-367	7
1729	Quantum Chemical Calculation Studies on 4-Phenyl-1-(Propan-2-Ylidene)Thiosemicarbazide. 2005 , 16, 635-639	79
1728	Modeling the active centers of V ₂ O ₅ /SiO ₂ and V ₂ O ₅ /TiO ₂ supported catalysts. DFT theoretical analysis of optical properties. 2005 , 46, 577-590	8
1727	Synthesis, Crystal Structure and Luminescence of [Ag(PPh ₃) ₂ (H ₂ TMT)]. 2005 , 631, 1532-1535	20
1726	Electron transport with dissipation: A quantum kinetic approach. 2005 , 101, 564-571	6
1725	Soft cohesive forces. 2005 , 101, 579-598	78
1724	Avoiding asymptotic divergence of the potential from orbital- and energy-dependent exchange-correlation functionals. 2005 , 101, 635-644	19
1723	Optical properties of molecules in solution via hybrid TDDFT/MM simulations. 2005 , 101, 671-682	46
1722	Simulation of UV/visible absorption spectra of (Ethylenimine)nickel(II) catalysts by time-dependent density functional theory. 2005 , 101, 840-848	9
1721	Refractive index and third-order nonlinear susceptibility of C ₆₀ in the condensed phase calculated with the discrete solvent reaction field model. 2005 , 102, 612-619	21
1720	Memory formulas for perturbations in time-dependent density functional theory. 2005 , 102, 573-581	7
1719	TDDFT from molecules to solids: The role of long-range interactions. 2005 , 102, 684-701	58
1718	Spectroscopic and DFT studies of donor-acceptor molecules containing phenylquinoline and phenothiazine moieties in various redox states. 2005 , 104, 635-644	6
1717	DFT with effective potential expressed as a mapping of the external potential: Applications to closed-shell molecules. 2005 , 104, 538-550	14
1716	Electronic excitations in solids: Density functional and Green's function theory. 2005 , 242, 2737-2750	11

1715	Many-body perturbation theory combined with time dependent DFT: A new method for the calculation of the dielectric function of solids. 2005 , 242, 2729-2736	5
1714	First-principles optical spectra of low dimensional systems. 2005 , 242, 3032-3039	4
1713	A Local-Scaling Time-Dependent Current Density Functional Theory. 2005 , 74, 2864-2865	1
1712	Electronic structure and magnetic properties of solids. 2005 , 220,	11
1711	Ab initio density functional theory: the best of both worlds?. 2005 , 123, 62205	143
1710	Plane wave/pseudopotential implementation of excited state gradients in density functional linear response theory: a new route via implicit differentiation. 2005 , 122, 144101	28
1709	Analysis of the viscoelastic coefficients in the Vignale-Kohn functional: The cases of one- and three-dimensional polyacetylene. 2005 , 71,	18
1708	Molecular-dynamics simulations of non-adiabatic processes at surfaces. 2005 , 78, 773-786	15
1707	Time-dependent density-functional theory for the stopping power of an interacting electron gas for slow ions. 2005 , 71,	33
1706	Explicit inclusion of paramagnetic current density in the exchange-correlation functionals of current-density functional theory. 2005 , 71,	26
1705	Variational solution of the T-matrix integral equation. 2005 , 71,	18
1704	Dynamical exchange-correlation potentials in the spin channel for the two-dimensional electron liquid. 2005 , 72,	2
1703	Modeling of configurations and third-order nonlinear optical properties of methyl silsesquioxanes. 2005 , 122, 204709	8
1702	Cubic response functions in time-dependent density functional theory. 2005 , 122, 54107	67
1701	Generalized time-dependent density-functional-theory response functions for spontaneous density fluctuations and nonlinear response: Resolving the causality paradox in real time. 2005 , 71,	21
1700	Calculation of nonadiabatic couplings in density-functional theory. 2005 , 122, 34105	23
1699	Time-dependent current-density-functional theory for the metallic response of solids. 2005 , 71,	40
1698	Coordinate scaling in time-dependent current-density-functional theory. 2005 , 72,	6

1697	Time-dependent electron localization function. 2005 , 71,		101
1696	Derivative discontinuities in time-dependent density-functional theory. <i>Physical Review Letters</i> , 2005 , 95, 203004	7.4	61
1695	Recent development of self-interaction-free time-dependent density-functional theory for nonperturbative treatment of atomic and molecular multiphoton processes in intense laser fields. 2005 , 123, 62207		73
1694	Exciton sizes of conducting polymers predicted by time-dependent density functional theory. 2005 , 71,		179
1693	Non-V-representability of currents in time-dependent many-particle systems. 2005 , 71,		16
1692	Time-dependent configuration-interaction calculations of laser-pulse-driven many-electron dynamics: controlled dipole switching in lithium cyanide. 2005 , 123, 074105		138
1691	Efficient linear-response method circumventing the exchange-correlation kernel: theory for molecular conductance under finite bias. 2005 , 123, 204105		18
1690	Density functional theory of the electrical conductivity of molecular devices. <i>Physical Review Letters</i> , 2005 , 94, 146803	7.4	114
1689	Investigation of chromophore-chromophore interaction by electro-optic measurements, linear dichroism, x-ray scattering, and density-functional calculations. 2005 , 72, 036610		35
1688	Conserving approximations in time-dependent density functional theory. 2005 , 72,		72
1687	Time-dependent density-functional theory for molecular photoionization with noniterative algorithm and multicenter B-spline basis set: CS ₂ and C ₆ H ₆ case studies. 2005 , 122, 234301		82
1686	Exact-exchange time-dependent density-functional theory for static and dynamic polarizabilities. 2005 , 71,		23
1685	Calculation of vibrational frequencies within the real space pseudopotential approach. 2005 , 71,		3
1684	Molecular effects in the ionization of N ₂ , O ₂ , and F ₂ by intense laser fields. 2005 , 71,		29
1683	Continuum states from time-dependent density functional theory. 2005 , 122, 144103		15
1682	Spectroscopic properties of Na clusters embedded in a rare-gas matrix. 2005 , 71,		27
1681	Emergence of collective plasmon excitation in a confined one-dimensional electron gas. 2005 , 72,		29
1680	Undoing static correlation: long-range charge transfer in time-dependent density-functional theory. 2005 , 122, 234104		150

1679	N-representability and stationarity in time-dependent density-functional theory. 2005 , 71,	5
1678	Correlated many-electron dynamics: Application to inelastic electron scattering at a metal film. 2005 , 72,	31
1677	Self-consistent solution of the Dyson equation for atoms and molecules within a conserving approximation. 2005 , 122, 164102	93
1676	Generic Galilean-invariant exchange-correlation functionals with quantum memory. 2005 , 72,	24
1675	Dynamical corrections to the DFT-LDA electron conductance in nanoscale systems. <i>Physical Review Letters</i> , 2005 , 94, 186810	7-4 155
1674	Microscopic and macroscopic polarization within a combined quantum mechanics and molecular mechanics model. 2005 , 122, 34103	36
1673	Rydberg transition frequencies from the local density approximation. <i>Physical Review Letters</i> , 2005 , 95, 163006	7-4 28
1672	HARMONIC GENERATIONS OF CLUSTER Na ₂ IN ULTRASHORT INTENSE LASER PULSES. 2005 , 19, 2687-2692	1
1671	COUPLED-CLUSTER EQUATION OF MOTION STUDY FOR THE ELECTRONIC AND OPTICAL PROPERTIES OF CONJUGATED SYSTEMS. 2005 , 04, 603-622	4
1670	ELECTRONIC PATHWAYS IN PHOTOACTIVATED REPAIR OF UV MUTATED DNA. 2005 , 19, 473-487	10
1669	IONIZATION OF Na ₂ BY HIGHLY CHARGED PARTICLES. 2005 , 19, 2886-2891	4
1668	Chapter 2 Time-Dependent Density Functional Theory in Quantum Chemistry. 2005 , 19-30	44
1667	Ab initio absorption spectra of Ge nanocrystals. 2005 , 71,	37
1666	The first hyperpolarizability of p-nitroaniline in 1,4-dioxane: a quantum mechanical/molecular mechanics study. 2005 , 123, 074307	62
1665	The Ab Initio Calculation of Optical Rotation and Electronic Circular Dichroism. 2005 , 50, 185-212	101
1664	Fermi-Amaldi model for exchange-correlation: atomic excitation energies from orbital energy differences. 2005 , 103, 2061-2072	51
1663	Hole localization in [AlO ₄] ⁰ defects in silica materials. 2005 , 122, 144704	68
1662	Propagator corrections to adiabatic time-dependent density-functional theory linear response theory. 2005 , 122, 54111	102

1661	Many-body perturbation theory using the density-functional concept: beyond the GW approximation. <i>Physical Review Letters</i> , 2005 , 94, 186402	7.4	115
1660	Five Possible Isocyanoazulenes and Electron-Rich Complexes Thereof: A Quantitative Organometallic Approach for Probing Electronic Inhomogeneity of the Azulenic Framework. 2005 , 24, 2386-2397		20
1659	Ground- and excited-state electronic structure of an iron-containing molecular spin photoswitch. 2005 , 123, 94709		9
1658	Excitation energies from time-dependent density functional theory with accurate exchange-correlation potentials. 2005 , 103, 711-717		19
1657	Parallel implementation of density functional theory within the real space pseudopotential approach.		
1656	Linear Response Properties Required to Simulate Vibrational Spectra of Biomolecules in Various Media: (R)-Phenyloxirane (A Comparative Theoretical and Spectroscopic Vibrational Study). 2005 , 91-124		11
1655	Energy dependence of the exchange-correlation kernel of time-dependent density functional theory: A simple model for solids. 2005 , 72,		42
1654	Nonlinear optical properties of zwitterionic merocyanine aggregates: role of intermolecular interaction and solvent polarity. 2005 , 109, 9095-103		50
1653	Synthesis and characterization of a novel terthiophene-based quinodimethane bearing a 3,4-ethylenedioxythiophene central unit. 2005 , 109, 22308-18		17
1652	Time-dependent density functional theory description of on-site electron repulsion and ligand field effects in the optical spectrum of hexaaquoruthenium(II) in solution. 2005 , 109, 12222-6		5
1651	Quantum chemical study on excited states and electronic coupling matrix element in a catechol-bridge-dicyanoethylene system. 2005 , 109, 4154-61		15
1650	Calculation of excitation energies of open-shell molecules with spatially degenerate ground states. I. Transformed reference via an intermediate configuration Kohn-Sham density-functional theory and applications to d1 and d2 systems with octahedral and tetrahedral symmetries. 2005 , 123, 144105		29
1649	On the origin of optical activity in tris-diamine complexes of Co(III) and Rh(III): a simple model based on time-dependent density function theory. 2005 , 127, 975-85		71
1648	Local-density approximation for the exchange energy functional in excited-state density functional theory. 2005 , 38, 3765-3777		14
1647	Characterizing the dimerizations of phenalenyl radicals by ab initio calculations and spectroscopy: sigma-bond formation versus resonance pi-stabilization. 2005 , 109, 11261-7		82
1646	Time-dependent quasirelativistic density-functional theory based on the zeroth-order regular approximation. 2005 , 123, 144101		58
1645	Optical properties and delocalization of excess negative charges on oligo(phenylenevinylene)s: a quantum chemical study. 2005 , 109, 5644-52		21
1644	Linear response theory in the continuum for deformed nuclei: Green's function vs time-dependent Hartree-Fock with the absorbing boundary condition. 2005 , 71,		140

1643	Synthesis and characterization of three novel perfluoro-oligothiophenes ranging in length from the trimer to the pentamer. 2005 , 109, 20737-45		16
1642	Time-dependent Kohn-Sham theory with memory. <i>Physical Review Letters</i> , 2005 , 95, 086401	7.4	47
1641	DFT/TDDFT studies of the geometry, electronic structure and spectra of (12S)-1,4,7,10-tetraazadicyclo[10,3,0]-pentadecane-3,11-dione and its derivatives. 2005 , 109, 2878-86		13
1640	Spin-polarized conceptual density functional theory study of the regioselectivity in the [2+2] photocycloaddition of enones to substituted alkenes. 2005 , 109, 6335-43		41
1639	Alternated quinoid/aromatic units in terthiophenes building blocks for electroactive narrow band gap polymers. Extended spectroscopic, solid state, electrochemical, and theoretical study. 2005 , 109, 16616-27		41
1638	A TDDFT study of the optical response of DNA bases, base pairs, and their tautomers in the gas phase. 2005 , 109, 2373-80		89
1637	Time-dependent four-component relativistic density-functional theory for excitation energies. II. The exchange-correlation kernel. 2005 , 123, 054102		80
1636	Time-dependent density functional theory of classical fluids. <i>Physical Review Letters</i> , 2005 , 94, 183001	7.4	54
1635	Single-molecule manipulation and chemistry with the STM. 2005 , 17, S1049-S1074		59
1634	Molecular dynamics in electronically excited states using time-dependent density functional theory. 2005 , 103, 963-981		120
1633	Quantum many-body dynamics in a Lagrangian frame: II. Geometric formulation of time-dependent density functional theory. 2005 , 71,		53
1632	Quantum many-body dynamics in a Lagrangian frame: I. Equations of motion and conservation laws. 2005 , 71,		65
1631	Ab initio theory of superconductivity. II. Application to elemental metals. 2005 , 72,		199
1630	Finite lifetime effects on the polarizability within time-dependent density-functional theory. 2005 , 122, 224115		146
1629	Time-dependent density functional theory: past, present, and future. 2005 , 123, 62206		665
1628	Time-dependent density functional theory for nonadiabatic processes. 2005 , 45, 161-170		26
1627	Laser-driven electron dynamics at interfaces. 2005 , 45, 205-215		12
1626	Exact time-dependent exchange-correlation potentials for strong-field electron dynamics. <i>Physical Review Letters</i> , 2005 , 94, 143003	7.4	130

1625	Direct optimization method to study constrained systems within density-functional theory. 2005 , 72,	400
1624	Carbon-based nanotechnology on a supercomputer. 2005 , 17, R413-R459	16
1623	The multiconfiguration time-dependent Hartree-Fock method for quantum chemical calculations. 2005 , 122, 124102	178
1622	Time-dependent quantum transport: A practical scheme using density functional theory. 2005 , 72,	275
1621	Molecular electronic excitations calculated from a solid-state approach: Methodology and numerics. 2005 , 72,	61
1620	Trajectory surface hopping in the time-dependent Kohn-Sham approach for electron-nuclear dynamics. <i>Physical Review Letters</i> , 2005 , 95, 163001	7.4 505
1619	Raman and theoretical study of the solvent effects on the sizable intramolecular charge transfer in the push-pull 5-(dimethylamino)-5'-nitro-2,2'-bithiophene. 2005 , 109, 8724-31	26
1618	Incisive structure-spectroscopic correlation in oligothiophenes functionalized with (+/-) inductive/mesomeric fluorine groups: joint Raman and DFT study. 2005 , 127, 13364-72	28
1617	Excited electronic states of small water clusters. 2005 , 122, 44111	64
1616	Density Functional Methods for Excited States: Equilibrium Structure and Electronic Spectra. 2005 , 16, 93-128	169
1615	Role of donor-acceptor strengths and separation on the two-photon absorption response of cytotoxic dyes: a TD-DFT study. 2005 , 109, 7276-84	52
1614	Calculation of optical rotation with time-periodic magnetic-field-dependent basis functions in approximate time-dependent density-functional theory. 2005 , 123, 114103	83
1613	Electron-hole and plasmon excitations in 3d transition metals: Ab initio calculations and inelastic x-ray scattering measurements. 2005 , 72,	38
1612	Fluctuation-dissipation theorem density-functional theory. 2005 , 122, 164106	180
1611	Fast electron correlation methods for molecular clusters in the ground and excited states. 2005 , 103, 2255-2265	125
1610	Approach to steady-state transport in nanoscale conductors. 2005 , 5, 2569-72	89
1609	Mesitylthio-oligothiophenes in various redox states. Molecular and electronic views as offered by spectroscopy and theory. 2005 , 109, 11275-84	19
1608	Synthesis and X-ray structure of the MnIIICl ₂ and MnIIIF ₂ complexes of N,N'-dimethyl-2,11-diaza[3,3](2,6)pyridinophane. High-field electron paramagnetic resonance and density functional theory studies of the MnIII complex. Evidence for a low-lying spin triplet state. 2005 , 44, 6059-66	29

1607	Theoretical study of the effects of solvent environment on photophysical properties and electronic structure of paracyclophane chromophores. 2005 , 122, 224505		58
1606	First-principles theoretical description of electronic transport including electron-electron correlation. 2005 , 72,		33
1605	Ab initio Ehrenfest dynamics. 2005 , 123, 084106		269
1604	Simulating molecular conductance using real-time density functional theory. 2006 , 74,		135
1603	TD-DFT investigation of the UV spectra of pyranone derivatives. 2006 , 110, 8144-50		69
1602	Microsolvation effects on the excited-state dynamics of protonated tryptophan. 2006 , 128, 16938-43		130
1601	Real-time ab initio simulations of excited carrier dynamics in carbon nanotubes. <i>Physical Review Letters</i> , 2006 , 97, 126104	7-4	43
1600	Conical intersections and double excitations in time-dependent density functional theory. 2006 , 104, 1039-1051		479
1599	Ion formation and kinetic electron emission during the impact of slow atomic metal particles on metal surfaces. 2006 , 73,		34
1598	Calculating state-to-state transition probabilities within time-dependent density-functional theory. 2006 , 74,		23
1597	Beyond time-dependent exact exchange: the need for long-range correlation. 2006 , 124, 144113		31
1596	Drug-Target Binding Investigated by Quantum Mechanical/Molecular Mechanical (QM/MM) Methods. 2006 , 449-479		7
1595	A TDDFT study of the excited states of DNA bases and their assemblies. 2006 , 110, 7129-38		104
1594	Orbital magnetization in crystalline solids: Multi-band insulators, Chern insulators, and metals. 2006 , 74,		168
1593	Ab initio calculation of optical spectra of liquids: many-body effects in the electronic excitations of water. <i>Physical Review Letters</i> , 2006 , 97, 137402	7-4	71
1592	Electrostatic modification of novel materials. 2006 , 78, 1185-1212		421
1591	Theoretical study of phosphorescence in dye doped light emitting diodes. 2006 , 125, 234704		31
1590	First hyperpolarizability of a sesquifulvalene transition metal complex by time-dependent density-functional theory. 2006 , 110, 1014-21		26

1589	Comparison of frozen-density embedding and discrete reaction field solvent models for molecular properties. 2006 , 8, 2349-59	80
1588	Protein-bound chromophores astaxanthin and phytochromobilin: excited state quantum chemical studies. 2006 , 8, 4053-71	29
1587	Efficient approach to time-dependent density-functional perturbation theory for optical spectroscopy. <i>Physical Review Letters</i> , 2006 , 96, 113001	7.4 166
1586	Nonequilibrium Green function theory for excitation and transport in atoms and molecules. 2006 , 35, 324-339	21
1585	Substitution and chemical environment effects on the absorption spectrum of indigo. 2006 , 124, 74104	100
1584	Time-dependent density functional theory investigation of the absorption, fluorescence, and phosphorescence spectra of solvated coumarins. 2006 , 125, 164324	104
1583	2-Phenylpyridine: To Twist or Not To Twist?. 2006 , 2, 1530-7	9
1582	Unified semiclassical theory for the two-state system: an analytical solution for general nonadiabatic tunneling. 2006 , 125, 44104	9
1581	Kohn-Sham Time-Dependent Density Functional Theory with Applications to Linear and Nonlinear Properties. 2006 , 151-209	1
1580	Bond length alternation and energy band gap of polyyne. 2006 , 110, 9771-4	108
1579	Spin and molecular electronics in atomically generated orbital landscapes. 2006 , 73,	551
1578	Density functional theory/time-dependent DFT studies on the structures, trend in DNA-binding affinities, and spectral properties of complexes [Ru(bpy) ₂ (p-R-pip)] ²⁺ (R = -OH, -CH ₃ , -H, -NO ₂). 2006 , 110, 8174-80	49
1577	Electronic excitations in metals and at metal surfaces. 2006 , 106, 4160-206	206
1576	Optical absorption and luminescence energies of F centers in CaO from ab initio embedded cluster calculations. 2006 , 125, 074710	27
1575	Structure-property relationships in push-pull amino/cyanovinyl end-capped oligothiophenes: quantum chemical and experimental studies. 2006 , 71, 7509-20	80
1574	Electronic structure and excitations in oligoacenes from ab initio calculations. 2006 , 124, 134901	95
1573	Effects of conjugation in length and dimension on spectroscopic properties of fluorene-based chromophores from experiment and theory. 2006 , 110, 13172-82	40
1572	Octopolar chromophores based on donor- and acceptor-substituted 1,3,5-tris(phenylethynyl)benzenes: impact of meta-conjugation on the molecular and electronic structure by means of spectroscopy and theory. 2006 , 110, 19198-206	30

1571	Spectroscopy of the cyano radical in an aqueous environment. 2006 , 110, 4854-65	18
1570	Combined quantum chemical density functional theory and spectroscopic Raman and UV-vis-NIR study of oligothienoacenes with five and seven rings. 2006 , 110, 5058-65	37
1569	Exploration of ground and excited electronic states of aromatic and quinoid S,S-dioxide terthiophenes. Complementary systems for enhanced electronic organic materials. 2006 , 128, 10134-44	53
1568	Synthesis and characterization of amphiphilic phenylene ethynylene oligomers and their Langmuir-Blodgett films. 2006 , 22, 8813-20	16
1567	Toward a Theoretical Quantitative Estimation of the Φ_{max} of Anthraquinones-Based Dyes. 2006 , 2, 434-40	86
1566	Electronic and geometric properties of ETS-10: QM/MM studies of cluster models. 2006 , 110, 8959-64	23
1565	Investigation of the UV/visible absorption spectra of merocyanine dyes using time-dependent density functional theory. 2006 , 110, 13007-13	62
1564	Excited-state properties and emission spectra of nonplanar heterocyclic helicenes. 2006 , 110, 11018-24	15
1563	A self-consistent field quantum hydrodynamic approach for molecular clusters. 2006 , 110, 5333-41	10
1562	Time-dependent density-functional theory beyond the adiabatic approximation: insights from a two-electron model system. 2006 , 125, 234108	97
1561	Truncation of periodic image interactions for confined systems. 2006 , 73,	207
1560	Ab initio investigation of the $n \rightarrow \pi^*$ transitions in thiocarbonyl dyes. 2006 , 110, 9145-52	59
1559	Time-dependent density-functional studies of the D2 coulomb explosion. 2006 , 110, 8443-50	28
1558	Synthesis, properties, and reactions of a series of stable dialkyl-substituted silicon-chalcogen doubly bonded compounds. 2006 , 128, 16914-20	114
1557	An ab initio study of the absorption spectra of indirubin, isoindigo, and related derivatives. 2006 , 110, 5629-35	72
1556	Electronic structure and optical properties of charged oligofluorenes studied by VIS/NIR spectroscopy and time-dependent density functional theory. 2006 , 110, 5984-93	39
1555	Origin of the photoinduced optical second harmonic generation arising in N-phenyl microcrystalline films. 2006 , 110, 6492-8	16
1554	Effect of conjugation path length on quadratic nonlinear optical properties of monomer and aggregates of zwitterionic merocyanine dyes. 2006 , 110, 8963-9	15

1553	Quantum dynamical approach to ultrafast molecular desorption from surfaces. 2006 , 106, 4116-59	128
1552	Two-photon excitation of substituted enedynes. 2006 , 110, 241-51	43
1551	Nonadiabatic electron dynamics in time-dependent density-functional theory. 2006 , 73,	43
1550	Theoretical study of structure, vibrational frequencies, and electronic spectra of polychlorinated dibenzo-p-dioxins. 2006 , 110, 4524-34	16
1549	Effect of double bonds on the conducting properties of ciguatoxin 3C and tetrahydropyrane-based polymers: a theoretical study. 2006 , 110, 1172-8	2
1548	Hybrid organic semiconductors including chalcogen atoms in pi-conjugated skeletons. Tuning of optical, redox, and vibrational properties by heavy atom conjugation. 2006 , 110, 7422-30	23
1547	Solution of the time-dependent, multi-particle Schrödinger equation using Monte Carlo and numerical integration. 2006 , 38, 231-239	2
1546	Molecular conduction: do time-dependent simulations tell you more than the Landauer approach?. 2006 , 124, 214708	67
1545	Density functional for spectroscopy: no long-range self-interaction error, good performance for Rydberg and charge-transfer states, and better performance on average than B3LYP for ground states. 2006 , 110, 13126-30	1001
1544	Density Functional Theory Based Ab Initio Molecular Dynamics Using the Car-Parrinello Approach. 2006 , 223-285	9
1543	Cation- π Interactions in Serotonin: Conformational, Electronic Distribution, and Energy Decomposition Analysis. 2006 , 2, 746-60	23
1542	Extracting electron transfer coupling elements from constrained density functional theory. 2006 , 125, 164105	260
1541	Quantum Mechanics and Molecular Mechanics Studies of Host-Guest Stabilization and Reactivity in Cyclodextrin Nanocavities. 2006 , 155-179	
1540	Excitation spectra of dibenzoborole containing pi-electron systems: controlling the electronic spectra by changing the p(pi)-pi* conjugation. 2006 , 110, 2434-9	26
1539	Propagating the Kadanoff-Baym equations for atoms and molecules. 2006 , 35, 340-348	15
1538	An exact ab initio theory of quantum transport using TDDFT and nonequilibrium Green's functions. 2006 , 35, 17-24	8
1537	Simple Iterative Procedure for Optimized Effective Potential Method in Density Functional Theory and in Current Density Functional Theory. 2006 , 75, 014302	2
1536	Symmetry preserving optimised effective potential theory (application to atoms). 2006 , 30, 249-261	6

1535	Time-dependent density-functional approaches to ultrafast intersubband dynamics in quantum wells. 2006 , 3, 2498-2501	1
1534	First principles calculation of optical and electronic properties with inclusion of exciton effects. 2006 , 3, 3365-3372	4
1533	Double-pole approximation in time-dependent density functional theory. 2006 , 106, 2840-2847	6
1532	Assessment of PBE0 for evaluating the absorption spectra of carbonyl molecules. 2006 , 106, 1853-1859	18
1531	Theoretical analysis of low-lying charge transfer states in [Ru(X) (Me)(CO) ₂ (Me-DAB)] (X=Cl, I; DAB=1,4-diaza-1,3-butadiene) complexes by TDDFT and CASSCF/CASPT2 methods. 2006 , 106, 2458-2469	22
1530	Circular dichroism spectrum of [Co(en) ₃] ³⁺ in water: A discrete solvent reaction field study. 2006 , 106, 2479-2488	30
1529	Three real-space discretization techniques in electronic structure calculations. 2006 , 243, 1016-1053	82
1528	PARSEC II: the pseudopotential algorithm for real-space electronic structure calculations: recent advances and novel applications to nano-structures. 2006 , 243, 1063-1079	242
1527	Real-time, real-space implementation of the linear response time-dependent density-functional theory. 2006 , 243, 1121-1138	165
1526	octopus: a tool for the application of time-dependent density functional theory. 2006 , 243, 2465-2488	659
1525	How much double excitation character do the lowest excited states of linear polyenes have?. 2006 , 329, 39-49	170
1524	Qprop: A Schrödinger-solver for intense laser-atom interaction. 2006 , 174, 396-421	155
1523	Failure of time-dependent density functional methods for excitations in spatially separated systems. 2006 , 419, 557-562	97
1522	Coherent states/density functional theory approach to molecular dynamics. 2006 , 420, 54-59	5
1521	Absorption and emission spectra in gas-phase and solution using TD-DFT: Formaldehyde and benzene as case studies. 2006 , 421, 272-276	66
1520	TDDFT investigation of the optical properties of cyanine dyes. 2006 , 425, 105-109	83
1519	A new challenge for time-dependent density functional theory. 2006 , 431, 410-414	8
1518	Analysis of Floquet formulation of time-dependent density-functional theory. 2006 , 433, 204-210	7

1517	Spectroscopic properties of dipicolinic acid and its dianion. 2006 , 322, 254-268	30
1516	On the role of second number-conserving functional derivatives. 2006 , 355, 148-151	5
1515	Quantum transport from the perspective of quantum open systems. 2006 , 357, 449-453	30
1514	On the linear dynamic response of average atom in plasma. 2006 , 99, 84-101	7
1513	Density functional theory study of vibronic structure of the first absorption Qx band in free-base porphin. 2006 , 65, 308-23	64
1512	Modelling of UV-molecular spectra of several bis-pyrazolopyridines derivatives. 2006 , 65, 511-6	1
1511	Kinetic excitation of solids: The concept of electronic friction. 2006 , 246, 333-339	42
1510	Molecular dynamics of nonadiabatic processes at surfaces: Chemisorption of H/Al(111). 2006 , 600, 3624-3628	32
1509	Time-dependent density-functional molecular-dynamics study of the isotope effect in chemicurrents. 2006 , 600, 5068-5073	19
1508	Energy and lifetime of surface plasmon from first-principles calculations. 2006 , 81, 186-191	13
1507	Investigation of the structures and absorption spectra for some hemicyanine dyes with pyridine nucleus by TD-DFT/PCM approach. 2006 , 778, 15-20	14
1506	Electrophilicity index. 2006 , 106, 2065-91	1155
1505	Heterobimetallic Zn(II)-Ln(III) phenylene-bridged schiff base complexes, computational studies, and evidence for singlet energy transfer as the main pathway in the sensitization of near-infrared Nd ³⁺ luminescence. 2006 , 45, 9315-25	149
1504	Constrained Density Functional Theory and Its Application in Long-Range Electron Transfer. 2006 , 2, 765-74	225
1503	Thioindigo dyes: highly accurate visible spectra with TD-DFT. 2006 , 128, 2072-83	220
1502	AB-INITIO STUDY ON THE EXCITON BINDING ENERGIES IN ORGANIC SEMICONDUCTORS. 2006 , 20, 261-280	28
1501	Synthesis, crystal structure and density functional theoretical studies on phenyl-thiocarbamic acid-O-pyridin-4-ylmethyl ester. 2006 , 17, 539-545	8
1500	Quantum-chemical analysis of the CuCl ₂ molecule. 2006 , 47, 404-412	6

1499	Quantum mechanics at the core of multi-scale simulations. 2006 , 13, 89-109	5
1498	A critical evaluation of DFT, including time-dependent DFT, applied to bioinorganic chemistry. 2006 , 11, 702-11	302
1497	Calculation of electronic circular dichroism spectra by rotating wave approximation. 2006 , 324, 622-630	1
1496	Time-dependent density functional theory determination of the absorption spectra of naphthoquinones. 2006 , 328, 324-332	29
1495	A TD-DFT study on the electronic spectrum of Ru(II)L ₂ [L=bis(5'-methyl-2,2'-bipyridine-6-carboxylato)] in the gas phase and DMF solution. 2006 , 330, 204-211	42
1494	Electronic interactions in a new pi-extended tetrathiafulvalene dimer. 2006 , 12, 2709-21	29
1493	Tuning of electronic properties in thienyl-phosphole pi-conjugated systems through P-functionalization monitored by Raman spectroscopy. 2006 , 12, 3759-67	24
1492	The interplay of inverted redox potentials and aromaticity in the oxidized states of new pi-electron donors: 9-(1,3-dithiol-2-ylidene)fluorene and 9-(1,3-dithiol-2-ylidene)thioxanthene derivatives. 2006 , 12, 3389-400	32
1491	Optical, redox, and NLO properties of tricyanovinyl oligothiophenes: comparisons between symmetric and asymmetric substitution patterns. 2006 , 12, 5458-70	37
1490	Prediction of spectroscopic constants for diatomic molecules in the ground and excited states using time-dependent density functional theory. 2006 , 27, 163-73	13
1489	Quantum chemical methods for the investigation of photoinitiated processes in biological systems: theory and applications. 2006 , 7, 2259-74	66
1488	Quantum Chemical Investigations of Reaction Paths of Metalloenzymes and Biomimetic Models □ The Hydrogenase Example. 2006 , 1-46	7
1487	Electronic Structure Calculations for Nanomolecular Systems. 2006 , 77-116	2
1486	Chapter 3 Ab initio Molecular Dynamics: Dynamics and Thermodynamic Properties. 2006 , 2, 55-95	1
1485	On correlated electron-nuclear dynamics using time-dependent density functional theory. 2006 , 125, 014110	24
1484	Non-adiabatic quantum molecular dynamics: ionization of many-electron systems. 2006 , 39, 2989-3008	20
1483	Quantum dynamics of laser-induced desorption from metal and semiconductor surfaces, and related phenomena. 2006 , 18, S1425-S1459	10
1482	Quantum chemical study of electronic and structural properties of retinal and some aromatic analogs. 2006 , 125, 144901	11

1481	Photoabsorption spectra of Ti ₈ C ₁₂ metallocarbohedrynes: Theoretical spectroscopy within time-dependent density functional theory. 2006 , 125, 074311		14
1480	Calculation of two-photon absorption spectra of donor- π -acceptor compounds in solution using quadratic response time-dependent density functional theory. 2006 , 125, 094103		55
1479	Calculation of nonadiabatic couplings with restricted open-shell Kohn-Sham density-functional theory. 2006 , 125, 224103		18
1478	Long-range excitations in time-dependent density functional theory. 2006 , 125, 184111		52
1477	The quantum defect: the true measure of time-dependent density-functional results for atoms. 2006 , 124, 94102		13
1476	Structural implications of ring shape, dimension, and metal atom insertion in nanosized cyclic oligothiophenes: joint Raman and density functional theory study. 2006 , 125, 44518		9
1475	Density functional theory of complex transition densities. 2006 , 125, 124104		26
1474	Exchange interactions, spin waves, and transition temperatures in itinerant magnets. 2006 , 86, 1713-1752		101
1473	Electronic excitation energies of Zn(i)S(i) nanoparticles. 2006 , 17, 4100-5		16
1472	Adiabatic approximation of the correlation function in the density-functional treatment of ionization processes. <i>Physical Review Letters</i> , 2006 , 97, 203001	7.4	41
1471	First-principles description of correlation effects in layered materials. <i>Physical Review Letters</i> , 2006 , 96, 136404	7.4	178
1470	Signatures of short-range many-body effects in the dielectric function of silicon for finite momentum transfer. <i>Physical Review Letters</i> , 2006 , 97, 237602	7.4	38
1469	Optimal control of ultrafast laser driven many-electron dynamics in a polyatomic molecule: N-methyl-6-quinolone. 2006 , 124, 144310		48
1468	Zero-bias molecular electronics: Exchange-correlation corrections to Landauer's formula. 2006 , 73,		160
1467	Quantum memory effects in the dynamics of electrons in gold clusters. 2006 , 73,		36
1466	Excitation spectra of nitro-diphenylaniline: accurate time-dependent density functional theory predictions for charge-transfer dyes. 2006 , 124, 204321		47
1465	Real-time propagation time-dependent density functional theory study on the ring-opening transformation of the photoexcited crystalline benzene. 2006 , 124, 124507		20
1464	Exact-exchange Kohn-Sham formalism applied to one-dimensional periodic electronic systems. 2006 , 74,		16

1463	Transient absorption spectroscopy and quantum-chemical studies of matrix-isolated perylene derivatives. 2006 , 73,		33
1462	Electronic density response of liquid water using time-dependent density functional theory. 2006 , 73,		60
1461	Exact-exchange time-dependent density-functional theory with the frequency-dependent kernel. 2006 , 73,		29
1460	Construction of model dielectric functions for two- and three-dimensional electron liquids from density functionals. 2006 , 73,		5
1459	Performance of the Vignale-Kohn functional in the linear response of metals. 2006 , 74,		19
1458	Electron correlation of one-dimensional H ₂ in intense laser fields: Time-dependent extended Hartree-Fock and time-dependent density-functional-theory approaches. 2006 , 73,		30
1457	Time-dependent density functional theory with ultrasoft pseudopotentials: Real-time electron propagation across a molecular junction. 2006 , 73,		66
1456	Time-dependent density-functional theory beyond the local-density approximation. <i>Physical Review Letters</i> , 2006 , 97, 036403	7.4	19
1455	Ab initio simulation of the spin transition during chemisorption: H/Al(111). <i>Physical Review Letters</i> , 2006 , 97, 216101	7.4	53
1454	Photoabsorption spectra of cationic mercury clusters. 2006 , 74,		9
1453	Structural, electronic, and optical properties of the diindenoperylene molecule from first-principles density-functional theory. 2006 , 74,		20
1452	Modified linear response for time-dependent density-functional theory: Application to Rydberg and charge-transfer excitations. 2006 , 74,		36
1451	Optimized effective potential in real time: Problems and prospects in time-dependent density-functional theory. 2006 , 74,		29
1450	Classical nuclear motion in quantum transport. <i>Physical Review Letters</i> , 2006 , 97, 046603	7.4	47
1449	Current-induced magnetization dynamics in disordered itinerant ferromagnets. 2006 , 74,		123
1448	Ab-initio studies of functionalized semiconductor quantum dots. 2006 ,		
1447	Magnetoelasticity theory of incompressible quantum Hall liquids. 2006 , 73,		19
1446	Bounds on the overlap of the Hartree-Fock, optimized effective potential, and density functional approximations with the exact energy eigenstates. 2006 , 124, 204109		14

1445	TIME-DEPENDENT CURRENT-DENSITY-FUNCTIONAL THEORY APPLIED TO ATOMS AND MOLECULES. 2006 , 20, 3419-3463	20
1444	Role of the exchange-correlation potential in ab initio electron transport calculations. 2007 , 126, 201102	99
1443	DETERMINATION OF THE ELECTRONIC GROUND STATE OF MOLECULAR SYSTEMS WITH THE MULTI-CONFIGURATION TIME-DEPENDENT HARTREE-FOCK METHOD. 2007 , 06, 563-574	14
1442	Time-dependent density functional theory scheme for efficient calculations of dynamic (hyper)polarizabilities. 2007 , 126, 184106	94
1441	Plasmon excitation in beryllium: inelastic x-ray scattering experiments and first-principles calculations. 2007 , 19, 046207	11
1440	Effects of solvation on one- and two-photon spectra of coumarin derivatives: a time-dependent density functional theory study. 2007 , 126, 094303	45
1439	Time-Dependent Density-Functional Calculations for Optical Spectra of Na ₂ and Na ₄ Clusters. 2007 , 47, 901-904	2
1438	Charged polycyclic aromatic hydrocarbon clusters and the galactic extended red emission. 2007 , 104, 5274-8	97
1437	Photoinduced electric currents in ring-shaped molecules by circularly polarized laser pulses. 2007 , 75,	64
1436	Path integral formulation for quantum nonadiabatic dynamics and the mixed quantum classical limit. 2007 , 126, 134107	17
1435	Optimized effective potential method for individual low-lying excited states. 2007 , 126, 174106	32
1434	Localized exchange-correlation potential from second-order self-energy for accurate Kohn-Sham energy gap. 2007 , 126, 214102	30
1433	Nonadiabatic couplings from time-dependent density functional theory: formulation in the Casida formalism and practical scheme within modified linear response. 2007 , 127, 064103	80
1432	Theoretical understanding of the increment of beta upon protonation of pyridine peripheral octupolar molecules: toward nonlinear optical sensors. 2007 , 127, 164704	10
1431	On the Electronic States of S ₄ ⁺ and S ₄ ⁻ Isomers. 2007 , 72, 83-99	6
1430	Excitonic effects in a time-dependent density functional theory. 2007 , 127, 114902	55
1429	Improved exchange-correlation potential for polarizability and dissociation in density functional theory. 2007 , 126, 191106	19
1428	Real-time propagation of the reduced one-electron density matrix in atom-centered Gaussian orbitals: application to absorption spectra of silicon clusters. 2007 , 127, 234107	68

1427	Dynamics simulation of a π -conjugated light-harvesting dendrimer. 2007 , 19, 365242	7
1426	Multiscale modelling of a nanoelectromechanical shuttle. 2007 , 9, 51-51	4
1425	Efficient ab initio calculations of bound and continuum excitons in the absorption spectra of semiconductors and insulators. 2007 , 76,	39
1424	Magnetization damping in a local-density approximation. 2007 , 75,	33
1423	Strong variation of dielectric response and optical properties of lithium under pressure. 2007 , 75,	15
1422	Time-dependent density-matrix-functional theory. 2007 , 75,	76
1421	Multicomponent density-functional theory for time-dependent systems. 2007 , 76,	27
1420	Analysis of the Vignale-Kohn current functional in the calculation of the optical spectra of semiconductors. 2007 , 75,	25
1419	Electronic viscosity in a quantum well: A test for the local-density approximation. 2007 , 76,	10
1418	Influence of hydrogen absorption on low-energy electronic collective excitations in palladium. 2007 , 76,	22
1417	Quantum Monte Carlo calculations of the optical gaps of Ge nanoclusters using core-polarization potentials. 2007 , 75,	7
1416	Atomic-orbital-based approximate self-interaction correction scheme for molecules and solids. 2007 , 75,	139
1415	Alternative perspective on density-functional perturbation theory. 2007 , 76,	5
1414	Time-dependent density functional calculation of e-H scattering. <i>Physical Review Letters</i> , 2007 , 99, 043005	20
1413	Excited-state forces within time-dependent density-functional theory: A frequency-domain approach. 2007 , 76,	17
1412	Optical properties of MoS ₃ nanowires. 2007 , 76,	13
1411	Time-dependent density functional theory: Derivation of gradient-corrected dynamical exchange-correlational potentials. 2007 , 76,	16
1410	Violation of the zero-force theorem in the time-dependent Krieger-Li-Iafrate approximation. 2007 , 75,	48

1409	Lorentz shear modulus of a two-dimensional electron gas at high magnetic field. 2007 , 76,	52
1408	Macroscopic limit of time-dependent density-functional theory for adiabatic local approximations of the exchange-correlation kernel. 2007 , 76,	7
1407	Time-dependent deformation functional theory. 2007 , 75,	37
1406	Quantum master equation scheme of time-dependent density functional theory to time-dependent transport in nanoelectronic devices. 2007 , 75,	36
1405	Simple dynamic exchange-correlation kernel of a uniform electron gas. 2007 , 75,	43
1404	Density-functional calculation for the tunnel ionization rate of hydrocarbon molecules. 2007 , 75,	20
1403	Momentum distributions in time-dependent density-functional theory: Product-phase approximation for nonsequential double ionization in strong laser fields. 2007 , 76,	29
1402	Stochastic time-dependent current-density-functional theory. <i>Physical Review Letters</i> , 2007 , 98, 226403 7.4	66
1401	Organic Materials for Multiphoton Absorption: Time-Dependent Density Functional Theory Calculations. 2007 ,	
1400	Including nonlocality in the exchange-correlation kernel from time-dependent current density functional theory: Application to the stopping power of electron liquids. 2007 , 76,	46
1399	Microscopic current dynamics in nanoscale junctions. 2007 , 75,	57
1398	Bound states in ab initio approaches to quantum transport: A time-dependent formulation. 2007 , 75,	73
1397	Self-energy and lifetime of Shockley and image states on Cu(100) and Cu(111): Beyond the GW approximation of many-body theory. 2007 , 76,	8
1396	Resonance Raman study of polyynes encapsulated in single-wall carbon nanotubes. 2007 , 76,	43
1395	Photoelectron spectra of anionic sodium clusters from time-dependent density-functional theory in real time. 2007 , 76,	49
1394	Photoselected electron transfer pathways in DNA photolyase. 2007 , 104, 802-7	68
1393	Time-dependent quantum Monte Carlo: preparation of the ground state. 2007 , 9, 70-70	39
1392	Calculation of static and dynamic linear magnetic response in approximate time-dependent density functional theory. 2007 , 126, 024101	48

1391	Time-dependent quantum Monte Carlo and the stochastic quantization. 2007 , 127, 134110	11
1390	Time-dependent approach to electronically excited states of molecules with the multiconfiguration time-dependent Hartree-Fock method. 2007 , 126, 214106	62
1389	Time-Dependent Schrödinger Equation Approach and Bethe-Salpeter Equation Approach. 2007 , 48, 649-652	5
1388	New approach for a complete experiment: C1s photoionization in CO ₂ molecules. 2007 , 40, F241-F250	15
1387	Theoretische Chemie 2006. 2007 , 55, 313-315	
1386	Chapter 10 Time-dependent transport phenomena. 2007 , 17, 247-284	20
1385	Role of charge transfer interaction and conjugation length on electrical polarizability of doped trans-polyacetylene oligomers. 2007 , 111, 11867-72	1
1384	Tetrathiafulvalene-Based Materials for Organic Field Effect Transistors. Inspection of Their Semiconductor Properties by Means of Molecular Spectroscopy and Quantum Chemistry. 2007 , 111, 10110-10118	17
1383	Investigation of crown-containing styrylthiophene derivatives which are optically and electrochemically sensitive to the presence of metal cations. 2007 , 157, 885-893	10
1382	Trajectory surface hopping within linear response time-dependent density-functional theory. <i>Physical Review Letters</i> , 2007 , 98, 023001	7.4 303
1381	Electronic excitations: Ab initio calculations of electronic spectra and application to zirconia ZrO ₂ , titania TiO ₂ and cuprous oxide Cu ₂ O. 2007 , 38, 482-493	16
1380	Photochromic properties of dithienylazoles and other conjugated diarylethenes. 2007 , 192, 211-219	51
1379	A coupled cluster study of the electronic spectroscopy and photochemistry of Cr(CO) ₆ . 2007 , 9, 6115-22	40
1378	Theoretical study of structure, vibrational frequencies, and electronic spectra of dibenzofuran and its polychlorinated derivatives. 2007 , 111, 1339-50	9
1377	Linear and Nonlinear Optical Properties of Pyridine-Based Octopolar Chromophores Designed for Chemical Sensing. Joint Spectroscopic and Theoretical Study. 2007 , 111, 18778-18784	23
1376	Average excitation energies from time-dependent density functional response theory. 2007 , 126, 074112	20
1375	Thiophene- and selenophene-based heteroacenes: combined quantum chemical DFT and spectroscopic Raman and UV-Vis-NIR study. 2007 , 111, 7488-96	31
1374	Synthesis and radical coupling of pyridine-bridged pi-extended tetrathiafulvalene (TTF)-type donors and push-pull analogues. 2007 , 5, 1201-9	14

1373	An Introduction to Density Functional Theory. 2007 , 187-216		23
1372	Transforming nonlocality into a frequency dependence: a shortcut to spectroscopy. <i>Physical Review Letters</i> , 2007 , 99, 057401	7.4	32
1371	Role of electronic excitations in ion collisions with carbon nanostructures. <i>Physical Review Letters</i> , 2007 , 99, 016104	7.4	122
1370	A well-tempered density functional theory of electrons in molecules. 2007 , 9, 2932-41		321
1369	Scaled second-order perturbation corrections to configuration interaction singles: efficient and reliable excitation energy methods. 2007 , 111, 5314-26		167
1368	Optical properties of real surfaces: Local-field effects at oxidized Si(100)(2 \times 8) computed with an efficient numerical scheme. 2007 , 75,		10
1367	Formulation of magnetically perturbed time-dependent density functional theory. 2007 , 127, 134108		24
1366	Dependence of Spurious Charge-Transfer Excited States on Orbital Exchange in TDDFT: Large Molecules and Clusters. 2007 , 3, 976-87		259
1365	Time-dependent density-functional theory for extended systems. 2007 , 70, 357-407		168
1364	Critique of the foundations of time-dependent density-functional theory. 2007 , 75,		64
1363	Ultrasoft pseudopotentials in time-dependent density-functional theory. 2007 , 127, 164106		56
1362	Water-molecule fragmentation induced by charge exchange in slow collisions with He ⁺ and He ²⁺ ions in the keV-energy region. 2007 , 75,		29
1361	Update 1 of: Electrophilicity Index. 2007 , 107, PR46-PR74		256
1360	Theoretical studies on structures and spectroscopic properties of a series of novel mixed-ligand Ir(III) complexes [Ir(Mebib)(ppy)X]. 2007 , 1922-8		19
1359	Frequency-dependent response properties and excitation energies from one-electron density matrix functionals. 2007 , 9, 5956-65		14
1358	Intersystem crossing processes in nonplanar aromatic heterocyclic molecules. 2007 , 111, 10490-9		187
1357	Quantum chemical insights in energy dissipation and carotenoid radical cation formation in light harvesting complexes. 2007 , 9, 2917-31		22
1356	Inner-shell spectroscopy by the Gaussian and augmented plane wave method. 2007 , 9, 1599-610		69

1355	A visible light photocatalyst: effects of vanadium substitution on ETS-10. 2007 , 9, 5096-104	20
1354	Advanced correlation functionals: application to bulk materials and localized systems. 2007 , 111, 12458-65	34
1353	Push-pull bithienyl chromophore with an unusual transverse path of conjugation. 2007 , 111, 841-51	5
1352	Electronic and molecular structures of trigonal truxene-core systems conjugated to peripheral fluorene branches. Spectroscopic and theoretical study. 2007 , 111, 4026-35	33
1351	Signature of the conformational preferences of small peptides: a theoretical investigation. 2007 , 111, 8650-8	20
1350	Accurate evaluation of valence and low-lying Rydberg states with standard time-dependent density functional theory. 2007 , 111, 5549-56	104
1349	Generator coordinate method in time-dependent density-functional theory: memory made simple. 2007 , 127, 124101	17
1348	Reductive cleavage mechanism of methylcobalamin: elementary steps of Co-C bond breaking. 2007 , 111, 7638-45	44
1347	Optical spectra of Cu(II)-azurin by hybrid TDDFT-molecular dynamics simulations. 2007 , 111, 10248-52	36
1346	Electronic stopping power in LiF from first principles. <i>Physical Review Letters</i> , 2007 , 99, 235501	7.4 124
1345	New Hybrid Materials Made from Dipolar Fe(II) Arylacetylide Complexes Exhibiting Temperature-Dependent Second Harmonic Generation Properties. 2007 , 111, 12094-12099	12
1344	Ab initio optical absorption spectra of size-expanded xDNA base assemblies. 2007 , 111, 14012-21	46
1343	Electron turbulence at nanoscale junctions. 2007 , 7, 1789-92	13
1342	Optical spectra of Cu, Ag, and Au monomers and dimers at regular sites and oxygen vacancies of the MgO(001) surface. A systematic time-dependent density functional study using embedded cluster models. 2007 , 111, 6870-80	9
1341	First-principles determinations and investigations of the electronic absorption and third-order polarizability spectra of electron donor-acceptor chromophores tetraalkylammonium halide/carbon tetrabromide. 2007 , 111, 9249-54	2
1340	Carotenoid radical cations as a probe for the molecular mechanism of nonphotochemical quenching in oxygenic photosynthesis. 2007 , 111, 3481-7	67
1339	On the Convergence of the Physicochemical Properties of [n]Helicenes. 2007 , 111, 14948-14955	65
1338	Assessment of time-dependent density functional schemes for computing the oscillator strengths of benzene, phenol, aniline, and fluorobenzene. 2007 , 127, 084103	76

1337	Origin of the anomalous two-photon absorption in fluorescent protein DsRed. 2007 , 111, 505-7	21
1336	Electronic Properties of DNA Base Molecules Adsorbed on a Metallic Surface. 2007 , 111, 14541-14551	53
1335	Predictions of novel two-photon absorption bands in fluorescent proteins. 2007 , 111, 14043-50	40
1334	Solving the Kadanoff-Baym equations for inhomogeneous systems: application to atoms and molecules. <i>Physical Review Letters</i> , 2007 , 98, 153004	7.4 111
1333	Theoretical studies on the excited states, DNA photocleavage, and spectral properties of complex [Ru(phen)(2)(6-OH-dppz)](2+). 2007 , 111, 273-80	32
1332	Opto-Electronic Properties of Fluorene-Based Derivatives as Precursors for Light-Emitting Diodes. 2007 , 111, 5812-5820	20
1331	Theoretical study of the photoabsorption spectrum of small chromium clusters. 2007 , 76,	12
1330	Time-dependent density-functional theory for open systems. 2007 , 75,	150
1329	Theoretical Studies on Structures and Spectroscopic Properties of Bis-Cyclometalated Iridium Complexes. 2007 , 26, 143-149	72
1328	Synthesis, characterization, and electronic structures of a series of two-dimensional trimetallic cluster complexes, Ru ₃ (CO) ₉ (μ-SnPh ₂) ₃ [Pt(PBu(t) ₃)] _x , x = 0-3. 2007 , 129, 12328-40	37
1327	Theoretical studies on structures and spectroscopic properties of a series of novel cationic [trans-(C/N) ₂ Ir(PH ₃) ₂] ⁺ (C/N = ppy, bzq, ppz, dfppy). 2007 , 111, 8724-30	74
1326	TD-DFT investigation of diarylethene dyes with cyclopentene, dihydrothiophene, and dihydropyrrole bridges. 2007 , 111, 5528-35	32
1325	Density Functional Theory Molecular Cluster Study of Copper Interaction with Nitric Oxide Dimer in Cu/ZSM-5 Catalysts. 2007 , 111, 3080-3089	22
1324	Chapter 13 Chemical reactivity dynamics in ground and excited electronic states. 2007 , 19, 269-286	8
1323	Efficient calculation of van der Waals dispersion coefficients with time-dependent density functional theory in real time: application to polycyclic aromatic hydrocarbons. 2007 , 127, 014107	28
1322	Gas phase spectra of all-benzenoid polycyclic aromatic hydrocarbons: triphenylene. 2007 , 126, 084304	14
1321	Real-time time-dependent density functional theory approach for frequency-dependent nonlinear optical response in photonic molecules. 2007 , 127, 154114	132
1320	Photolysis of methylcobalamin: identification of the relevant excited states involved in Co-C bond scission. 2007 , 111, 2419-22	43

1319	Theoretical investigation of the excited states of coumarin dyes for dye-sensitized solar cells. 2007 , 111, 5544-8	152
1318	Properties of phase-coherent energy shuttling on the nanoscale. 2007 , 126, 014705	7
1317	Computational study of the absorption spectra of green fluorescent protein mutants. 2007 , 85, 253-63	21
1316	Photoswitching of the fluorescent protein asFP595: mechanism, proton pathways, and absorption spectra. 2007 , 46, 530-6	89
1315	Photoswitching of the Fluorescent Protein asFP595: Mechanism, Proton Pathways, and Absorption Spectra. 2007 , 119, 536-542	25
1314	Analytical excited state forces for the time-dependent density-functional tight-binding method. 2007 , 28, 2589-601	36
1313	Vertical excitation energies for ribose and deoxyribose nucleosides. 2007 , 28, 1776-82	23
1312	Description of core excitations by time-dependent density functional theory with local density approximation, generalized gradient approximation, meta-generalized gradient approximation, and hybrid functionals. 2007 , 28, 2067-74	35
1311	Matrix model to predict specific optical rotations of acyclic chiral molecules. 2007 , 63, 2292-2314	30
1310	Ground and excited state properties of naphthazarin: Absorption spectroscopy and theoretical modeling study. 2007 , 803, 79-87	6
1309	On the basis set convergence of TD-DFT oscillator strengths: Dinitrophenylhydrazones as a case study. 2007 , 804, 31-34	17
1308	Conformational analysis and electronic transition of carbazole-based oligomers as explained by density functional theory. 2007 , 807, 109-119	21
1307	Performance of Time Dependent Density Functional Theory on excitations of medium sized molecules Test on ionic forms of anthraquinone dihydroxy derivatives. 2007 , 823, 78-86	27
1306	Study of the non-covalent interactions in Langmuir-Blodgett films: An interplay between π - π and dipole-dipole interactions. 2007 , 516, 58-66	24
1305	Modelling of electronic absorption spectrum of Pb(II)-caffeate complex by time-dependent density functional theory. 2007 , 438, 41-46	14
1304	Ab initio time-resolved density functional theory for lifetimes of excited adsorbate states at metal surfaces. 2007 , 439, 199-203	33
1303	Comment on Analysis of Floquet formulation of time-dependent density-functional theory [Chem. Phys. Lett. 433 (2006) 204]. 2007 , 441, 167-169	8
1302	Density functional theory study of . 2007 , 441, 309-313	3

1301	Time-dependent density functional theory with the multilayer fragment molecular orbital method. 2007 , 444, 346-350	59
1300	The triplet state of indigo: Electronic structure calculations. 2007 , 449, 11-17	8
1299	Experimental and theoretical study on the structure and electronic spectra of imiquimod and its synthetic intermediates. 2007 , 17, 4942-6	10
1298	Time-dependent density functional study of the electronic spectra of oligoacenes in the charge states $\bar{1}$, 0, +1, and +2. 2007 , 340, 43-58	92
1297	Vibrational spectra of nonlinear optical chromophores based on octopolar C3-symmetric 1,3,5 trisalkynylbenzenes. 2007 , 834-836, 369-373	2
1296	Electronic spectroscopy study and molecular docking simulation of the interaction of terthiophene with DNA. 2007 , 834-836, 176-181	4
1295	Synthesis, characterization, crystal structure and antiproliferative activity of platinum(II) complexes with 2-acetylpyridine-4-cyclohexyl-thiosemicarbazone. 2007 , 26, 2871-2879	29
1294	Syntheses, crystallography and spectroelectrochemical studies of ruthenium azomethine complexes. 2007 , 26, 3675-3685	27
1293	Ab initio tools for the accurate prediction of the visible spectra of anthraquinones. 2007 , 67, 334-41	42
1292	Numerical aspects of real-space approaches to strong-field electron dynamics. 2007 , 226, 89-103	13
1291	Analysis of self-interaction correction for describing core excited states. 2007 , 107, 23-29	36
1290	Density functional theory study of tautomerization of 2-aminothiazole in the gas phase and in solution. 2007 , 107, 247-258	13
1289	Associated and correlated quantum fluid dynamics in laser-induced ionization of helium. 2007 , 107, 647-656	4
1288	Constrained optimized potential method and second-order correlation energy for excited states. 2007 , 107, 2604-2615	12
1287	Femtosecond electron-ion dynamics in excited nano-materials: Real-time propagation based on the time-dependent density functional theory. 2007 , 204, 1925-1930	3
1286	Identification of gold oxide cluster structures in Au/Al ₂ O ₃ catalysts for low-temperature CO oxidation. 2007 , 413, 75-80	4
1285	Post-Hartree-Fock methods and dynamic correlation in atoms and molecules. 2007 , 103, 717-722	1
1284	Inclusion of the exchange-correlation effects in Ab initio methods for calculating the plasmon dispersion and line width in metals. 2007 , 49, 1820-1828	4

1283	Time-dependent density functional theory based upon the fragment molecular orbital method. 2007 , 127, 104108	85
1282	Electronic stopping in insulators: a simple model. 2007 , 19, 275211	20
1281	A CAS-DFT study of fundamental degenerate and nearly degenerate systems. 2007 , 105, 2667-2679	28
1280	Identification of fullerene-like CdSe nanoparticles from optical spectroscopy calculations. 2007 , 75,	45
1279	Theoretical investigation of excited states of molecules. An application on the nitrogen molecule. 2007 , 118, 637-642	7
1278	Quantum chemical calculation studies on 4-phenyl-1-(1-phenylethylidene)thiosemicarbazide. 2007 , 18, 325-330	1
1277	Synthesis, characterization, and quantum chemical calculational studies on 3-p-methylphenyl-4-amino- 1, 2, 4-triazole-5-thione. 2007 , 18, 993-1000	15
1276	Reaction of functionalized maleimides with versatile nucleophiles. Synthesis, electronic spectra and molecular orbital study. 2007 , 72, 271-284	7
1275	Theoretical studies of the electronic structures and optical properties of stable blue-emitting polymer based on 4H-cyclopenta-[def]-phenanthrene. 2008 , 49, 2077-2084	4
1274	TD-DFT investigation on the electronic spectra of novel N-methylmaleimides linked with indolizine ring system. 2008 , 855, 92-101	12
1273	Study of intramolecular charge transfer of Michler's ketone using time dependent density functional theory. 2008 , 860, 8-12	5
1272	The M06 suite of density functionals for main group thermochemistry, thermochemical kinetics, noncovalent interactions, excited states, and transition elements: two new functionals and systematic testing of four M06-class functionals and 12 other functionals. 2008 , 120, 215-241	19079
1271	On the TD-DFT UV/vis spectra accuracy: the azoalkanes. 2008 , 120, 405-410	50
1270	Theoretical studies of the spectroscopic properties of blue emitting iridium complexes. 2008 , 121, 155-164	21
1269	The role of bound states in time-dependent quantum transport. 2008 , 93, 355-364	24
1268	Theoretical investigation on dual fluorescence and intramolecular charge transfer of 5-phenyl-5H-phenanthridin-6-one. 2008 , 51, 632-639	1
1267	Theoretical studies on Ru(fppz) ₂ (CO)L (L = N-heterocyclic ligand): Electronic structure, absorption, phosphorescence, and solvatochromism. 2008 , 51, 1211-1220	1
1266	Ab-initio optical spectra of complex systems. 2008 , 5, 2543-2550	2

1265	Analysis of chemical bonding in electronic excited states using parity function. 2008 , 108, 1-14	5
1264	Time-dependent DFT-PCM investigation of the photophysics of ESIPT-exhibiting benzazole dyes. 2008 , 108, 2334-2339	3
1263	New perspectives on the fundamental theorem of density functional theory. 2008 , 108, 2756-2762	14
1262	Characterization of optical spectra of interacting systems: Application to oxide-supported metal clusters. 2008 , 108, 2978-2990	3
1261	Vibrational analysis of trans-stilbene in the excited singlet state by time-dependent density functional theory: calculations of the Raman, infrared, and fluorescence excitation spectra. 2008 , 39, 1584-1591	21
1260	Electronic and vibronic contributions to two-photon absorption in donor-acceptor-donor squaraine chromophores. 2008 , 14, 11082-91	41
1259	Time-dependent density functional theory molecular dynamics simulations of liquid water radiolysis. 2008 , 9, 2099-103	40
1258	The origin of the improved efficiency and stability of triphenylamine-substituted anthracene derivatives for OLEDs: a theoretical investigation. 2008 , 9, 2601-9	84
1257	Polarizable continuum model with the fragment molecular orbital-based time-dependent density functional theory. 2008 , 29, 2667-76	48
1256	Theoretical Studies on [Ru(bpy) ₂ (NN)] ²⁺ [NN = Hydrazone and Azine]: Ground- and Excited-State Geometries, Electronic Structures, Absorptions, and Phosphorescence Mechanisms. 2008 , 2008, 1268-1276	9
1255	Aminopyrimidine-Based Donor-Acceptor Chromophores: Push-Pull versus Aromatic Behaviour. 2008 , 2008, 99-108	20
1254	Thermo-Solvatochromism of Merocyanine Polarity Probes [What Are the Consequences of Increasing Probe Lipophilicity through Annulation?]. 2008 , 2008, 1165-1180	37
1253	Enhanced Two-Photon Absorption of Organic Chromophores: Theoretical and Experimental Assessments. 2008 , 20, 4641-4678	441
1252	Self-interaction correction in a simple model. 2008 , 372, 5598-5602	7
1251	Computational studies of CO and CO ⁺ : Density functional theory and time-dependent density functional theory. 2008 , 109, 2546-2560	7
1250	Synthesis, characterization, crystal structure and DFT studies on 1-acetyl-3-(2,4-dichloro-5-fluoro-phenyl)-5-phenyl-pyrazoline. 2008 , 69, 647-53	22
1249	Theory of current-driven magnetization dynamics in inhomogeneous ferromagnets. 2008 , 320, 1282-1292	119
1248	Theoretical studies on structures and spectroscopic properties of bis-cyclometalated iridium complexes [Ir(ppy) ₂ X ₂]. 2008 , 693, 947-956	10

1247	Two photon absorption of extended substituted phenylenevinylene oligomers: A TDDFT study. 2008 , 450, 322-328	11
1246	Mid-gap excitations in Anderson polyoxometalates. 2008 , 450, 391-395	6
1245	The luminescent [Mo ₆ X ₈ (NCS) ₆] ₂ [X=Cl, Br, I] clusters?: A computational study based on time-dependent density functional theory including spin-orbit and solvent-polarity effects. 2008 , 455, 38-41	24
1244	Theoretical study of the lowest electronic transitions of sulfur-bearing mesoionic compounds in gas-phase and in dimethyl sulfoxide. 2008 , 457, 119-123	18
1243	Electronic structure and molecular properties of the [Mo ₆ X ₈ L ₆] ₂ [X = Cl, Br, I; L = F, Cl, Br, I] clusters. 2008 , 460, 438-441	52
1242	Origin of electronic structure and time-dependent state averaging in the multi-configuration time-dependent Hartree-Fock approach to electron dynamics. 2008 , 463, 263-266	19
1241	Pigment Yellow 101: A showcase for photo-initiated processes in medium-sized molecules. 2008 , 347, 472-482	35
1240	Theoretical study of 2,5-diphenyl-1,4-distyrylbenzene (A model compound of PPV): A comparison of the electronic structure and photophysical properties of cis- and trans-isomers. 2008 , 345, 23-31	3
1239	Theoretical study on the structures and electronic spectra of novel quinoid 1,3,4-oxadiazole derivatives. 2008 , 848, 1-8	4
1238	DFT/TDDFT studies on electronic absorption and emission spectra of [Ru(bpy) ₂ (L)] ²⁺ (L = pip, o-mopip and p-mopip) in aqueous solution. 2008 , 855, 77-81	16
1237	Hydrogen atom transfer in the reaction of hydroxycinnamic acids with OH and HO ₂ radicals: DFT study. 2008 , 862, 53-59	17
1236	Theoretical studies on the optical properties and substituent effects of osmium (II) complexes Os(N [^] N)(CN) ₂ (PH ₃) ₂ . 2008 , 869, 11-18	
1235	Application of Lanczos-based time-dependent density-functional theory approach to semiconductor nanoparticle quantum dots. 2008 , 66, 7-15	13
1234	Carbonyl sulphide under strong laser field: Time-dependent density functional theory. 2008 , 49, 43-49	8
1233	Orbital-dependent density functionals: Theory and applications. 2008 , 80, 3-60	941
1232	First-principles electron dynamics simulation for optical breakdown of dielectrics under an intense laser field. 2008 , 77,	127
1231	Solvent effects on optical properties of molecules: a combined time-dependent density functional theory/effective fragment potential approach. 2008 , 129, 144112	89
1230	Frequency and Solvent Dependence of Nonlinear Optical Properties of Molecules. 2008 , 112, 8016-8021	35

1229	Investigation on the Second-Order Nonlinear Optical Responses in the Keto-Enol Equilibrium of Anil Derivatives. 2008 , 112, 5638-5645	74
1228	Configurational thermodynamics of alloys from first principles: effective cluster interactions. 2008 , 71, 046501	234
1227	Excitation energies in density functional theory: an evaluation and a diagnostic test. 2008 , 128, 044118	1066
1226	Theoretical methods of investigation of excited states of organic molecules. 2008 , 78, 774-783	9
1225	Evolution of the electron structure and excitation spectrum in palladium as a result of hydrogen absorption. 2008 , 53, 318-322	8
1224	Identification of excited singlet states of chlorophenol isomers. 2008 , 105, 327-332	6
1223	Absorption spectrum of OH radical in water. 2008 , 112, 13372-81	35
1222	Mechanism of Ir(ppy) ₂ (N--N) ⁺ (N--N = 2-phenyl-1H-imidazo[4,5-f][1,10]phenanthroline) sensor for F ⁻ , CF ₃ COOH, and CH ₃ COO ⁻ : density functional theory and time-dependent density functional theory studies. 2008 , 112, 8254-62	21
1221	A beta-naphthaleneimide-modified terthiophene exhibiting charge transfer and polarization through the short molecular axis. Joint spectroscopic and theoretical study. 2008 , 112, 6732-40	25
1220	Hartree-Fock and Kohn-Sham time-dependent response theory in a second-quantization atomic-orbital formalism suitable for linear scaling. 2008 , 129, 054106	41
1219	On the origin of alternating bond distortions and the emergence of chirality in polyoxometalate anions. 2008 , 130, 8223-33	98
1218	Time-dependent DFT studies of metal core-electron excitations in Mn complexes. 2008 , 112, 11223-34	17
1217	Mechanisms for ultrafast nonradiative relaxation in electronically excited eumelanin constituents. 2008 , 95, 4396-402	37
1216	Adiabatic approximation in nonperturbative time-dependent density-functional theory. <i>Physical Review Letters</i> , 2008 , 100, 153004	7.4 107
1215	Computational studies of semiconductor quantum dots. 2008 , 10, 4535-50	33
1214	TDDFT-Generalized kick perturbations and monitoring observables for calculation of excitation energies. 2008 , 88, 2817-2827	
1213	Melanin absorption spectroscopy: new method for noninvasive skin investigation and melanoma detection. 2008 , 13, 014017	153
1212	Natural dyes adsorbed on TiO ₂ nanowire for photovoltaic applications: enhanced light absorption and ultrafast electron injection. 2008 , 8, 3266-72	181

1211	DFT/TDDFT studies on the electronic structures and spectral properties of rhenium(I) pyridinybenzoimidazole complexes. 2008 , 112, 11190-7	69
1210	Fast and reliable theoretical determination of pKa* for photoacids. 2008 , 112, 794-6	32
1209	Self-consistent field method to obtain potential energy surfaces of excited molecules on surfaces. 2008 , 78,	153
1208	Electronic structure calculations for inhomogeneous systems: Interfaces, surfaces, and nanocontacts. 2008 , 17, 525	
1207	Structure, spectroscopy, and reactivity properties of porphyrin pincers: a conceptual density functional theory and time-dependent density functional theory study. 2008 , 112, 305-11	24
1206	Synthesis, characterization and density functional calculations on dichloro- bis (4-dimethylaminopyridine) cobalt(II) complex [Co(II)(C7H10N2)2Cl2]. 2008 , 61, 705-714	5
1205	Comment on "Critique of the foundations of time-dependent density-functional theory" 2008 , 78,	29
1204	Reply to "Comment on "Critique of the foundations of time-dependent density-functional theory" 2008 , 78,	29
1203	Theoretical analysis of the intermolecular interaction effects on the excitation energy of organic pigments: solid state quinacridone. 2008 , 112, 10887-94	39
1202	Investigating interaction-induced chaos using time-dependent density-functional theory. 2008 , 77,	4
1201	Self-consistent field calculations of excited states using the maximum overlap method (MOM). 2008 , 112, 13164-71	335
1200	Optimization and Parallelization of DFT and TDDFT in GAMESS on DoD HPC Machines. 2008 ,	1
1199	TD-DFT investigation of the electronic spectra of (DOEO)4HgBr4 ·TCE salt. 2008 , 106, 33-42	4
1198	Time-dependent approach to electron pumping in open quantum systems. 2008 , 77,	94
1197	Linear optical response of current-carrying molecular junction: a nonequilibrium Green's function-time-dependent density functional theory approach. 2008 , 128, 124705	44
1196	Rydberg energies using excited state density functional theory. 2008 , 129, 124112	40
1195	Double excitations and state-to-state transition dipoles in π excited singlet states of linear polyenes: Time-dependent density-functional theory versus multiconfigurational methods. 2008 , 77,	54
1194	Direct observation of photoinduced bent nitrosyl excited-state complexes. 2008 , 112, 8505-14	15

1193	One- and two-photon absorptions in asymmetrically substituted free-base porphyrins: a density functional theory study. 2008 , 128, 074302	5
1192	Performance of a nonempirical meta-generalized gradient approximation density functional for excitation energies. 2008 , 128, 084110	30
1191	Exciton scattering approach for branched conjugated molecules and complexes. I. Formalism. 2008 , 129, 174111	17
1190	Real-time, local basis-set implementation of time-dependent density functional theory for excited state dynamics simulations. 2008 , 129, 054110	151
1189	Electrochemical, spectral and theoretical studies of two new methyl-thieno[3,2-b]benzothiophenes and their polymers electrosynthesized in organic and micellar media. 2008 , 158, 6-24	14
1188	Cohesive energy curves for noble gas solids calculated by adiabatic connection fluctuation-dissipation theory. 2008 , 77,	292
1187	A density matrix-based quasienergy formulation of the Kohn-Sham density functional response theory using perturbation- and time-dependent basis sets. 2008 , 129, 214108	93
1186	Testing the numerical stability of time-dependent density functional simulations using the Suzuki-Trotter formula. 2008 , 77,	32
1185	Optimizing organic photovoltaics using tailored heterojunctions: A photoinduced absorption study of oligothiophenes with low band gaps. 2008 , 77,	95
1184	Mixed time-dependent density-functional theory/classical trajectory surface hopping study of oxirane photochemistry. 2008 , 129, 124108	165
1183	Correlated dynamics of electrons with reduced two-electron density matrices. 2008 , 78,	12
1182	Density functional calculations of nanoscale conductance. 2008 , 20, 083203	104
1181	Ab initio methods for the optical properties of CdSe clusters. 2008 , 77,	40
1180	Time-dependent density-functional theory in the projector augmented-wave method. 2008 , 128, 244101	158
1179	Theoretical studies of the tautomers of pyridinethiones. 2008 , 112, 3231-8	8
1178	Efficient formalism for large-scale ab initio molecular dynamics based on time-dependent density functional theory. <i>Physical Review Letters</i> , 2008 , 101, 096403	7.4 85
1177	Anion-arene adducts: C-H hydrogen bonding, anion-pi interaction, and carbon bonding motifs. 2008 , 2417-28	351
1176	Approximate spin symmetries in single determinantal approximations of many electron systems. 2008 , 104, 012020	1

1175	Optical properties in conjugated polymers. 2008 , 20, 064231		15
1174	A combined computational and experimental study on DNA-photocleavage of Ru(II) polypyridyl complexes [Ru(bpy) ₂ (L)] ²⁺ (L = pip, o-mopip and p-mopip). 2008 , 291-301		31
1173	Ab initio study of the dielectric response of crystalline ropes of metallic single-walled carbon nanotubes: Tube-diameter and helicity effects. 2008 , 78,		16
1172	Extended Hückel tight-binding approach to electronic excitations. 2008 , 129, 044107		9
1171	Nonradiative quenching of fluorescence in a semiconducting carbon nanotube: a time-domain ab initio study. <i>Physical Review Letters</i> , 2008 , 100, 197402	7-4	115
1170	Spin-flip scattering in time-dependent transport through a quantum dot: Enhanced spin-current and inverse tunneling magnetoresistance. 2008 , 78,		35
1169	A combined theoretical-experimental study on the acidity of WO(x)-ZrO(2) systems. 2008 , 10, 4181-8		15
1168	Real-time electron dynamics with exact-exchange time-dependent density-functional theory. <i>Physical Review Letters</i> , 2008 , 100, 056404	7-4	52
1167	Time-dependent V-representability on lattice systems. 2008 , 129, 044105		41
1166	Molecular engineering of torsional potentials in fluorogenic dyes via electronic substituent effects. 2008 , 112, 9692-701		12
1165	Theoretical Investigation of the Geometries and UV-vis Spectra of Poly(l-glutamic acid) Featuring a Photochromic Azobenzene Side Chain. 2008 , 4, 637-45		22
1164	Time-dependent density-functional theory for ultrafast interband excitations. 2008 , 77,		20
1163	Benchmarks for electronically excited states: time-dependent density functional theory and density functional theory based multireference configuration interaction. 2008 , 129, 104103		433
1162	Absorption spectra of blue-light-emitting oligoquinolines from time-dependent density functional theory. 2008 , 112, 13701-10		10
1161	Electronic, Optical, and Vibrational Properties of Bridged Dithienylethylene-Based NLO Chromophores. 2008 , 112, 3109-3120		44
1160	Ultrafast intersystem crossing in 1-nitronaphthalene. An experimental and computational study. 2008 , 112, 358-65		91
1159	Electrochemical, Magnetic, and Electrical Properties of H-Capped Sexithiophene Films. Part 3. Conduction in Poly(bis-terthienyl-B)s (B = Ethane, Disulfide, Diacetylene, Acetylene, Ethylene). 2008 , 20, 6847-6856		12
1158	Circular dichroism of trigonal dihedral chromium(III) complexes: a theoretical study based on open-shell time-dependent density functional theory. 2008 , 47, 11656-68		25

1157	Electronic and optical properties of ordered BexZn1-xSe alloys by the FPLAPW method. 2008 , 20, 075205	19
1156	Exact dynamics of dissipative electronic systems and quantum transport: Hierarchical equations of motion approach. 2008 , 128, 234703	271
1155	Single-Reference Methods for Excited States in Molecules and Polymers. 2008 , 15-64	7
1154	Assessment of the Accuracy of TD-DFT Absorption Spectra: Substituted Benzenes. 2008 , 73, 898-908	4
1153	Modeling the doubly excited state with time-dependent Hartree-Fock and density functional theories. 2008 , 129, 204107	49
1152	ON TIME DEPENDENT DFT WITH SIC. 2008 , 22, 4666-4673	1
1151	Combining DFT and many-body methods to understand correlated materials. 2008 , 20, 293201	85
1150	Reliability of range-separated hybrid functionals for describing magnetic coupling in molecular systems. 2008 , 129, 184110	73
1149	Dipole switching in large molecules described by explicitly time-dependent configuration interaction. 2008 , 128, 234307	21
1148	Dynamic correlations with time-dependent quantum Monte Carlo. 2008 , 128, 244106	19
1147	Current through single conjugated molecules: calculations versus measurements. 2008 , 129, 024901	8
1146	Molecular-orbital-free algorithm for excited states in time-dependent perturbation theory. 2008 , 129, 064114	15
1145	A revised electronic Hessian for approximate time-dependent density functional theory. 2008 , 129, 184114	62
1144	Local electron and ionic heating effects on the conductance of nanostructures. 2008 , 20, 374102	15
1143	Quasidegenerate scaled opposite spin second order perturbation corrections to single excitation configuration interaction. 2008 , 128, 164106	41
1142	Molecular dynamics with time dependent quantum Monte Carlo. 2008 , 129, 214107	11
1141	Electron dynamics in molecules: a new combination of nuclear quantum dynamics and electronic structure theory. 2008 , 41, 074006	57
1140	Theory of inelastic lifetimes of surface-state electrons and holes at metal surfaces. 2008 , 20, 304207	2

1139	Energy loss spectra of lithium under pressure. 2008 , 10, 053035	16
1138	On the mapping of time-dependent densities onto potentials in quantum mechanics. 2008 , 128, 044103	38
1137	Atomistic structure dependence of the collective excitation in metal nanoparticles. 2008 , 129, 034710	18
1136	Time-dependent configuration-interaction calculations of laser-driven dynamics in presence of dissipation. 2008 , 129, 084302	43
1135	Quantum Drude friction for time-dependent density functional theory. 2008 , 129, 134106	14
1134	Are Octahedral Ruthenium(II/III) and Osmium(II/III) Complexes Always Low-Spin?. 2008 , 73, 1231-1244	
1133	Pseudospectral time-dependent density functional theory. 2008 , 128, 104103	50
1132	Strong-field ionization in time-dependent density functional theory. 2008 , 84, 43001	14
1131	Dynamical generalization of a solvable family of two-electron model atoms with general interparticle repulsion. 2008 , 41, 085304	4
1130	The decay of excited He from stochastic density-functional theory: a quantum measurement theory interpretation. 2008 , 20, 395214	4
1129	A many-body approach to quantum transport dynamics: Initial correlations and memory effects. 2008 , 84, 67001	120
1128	Anomalous angular dependence of the dynamic structure factor near Bragg reflections: graphite. <i>Physical Review Letters</i> , 2008 , 101, 266406	7.4 21
1127	Density-functional theory of nonequilibrium tunneling. 2008 , 78,	8
1126	Dielectric properties and excitons for extended systems from hybrid functionals. 2008 , 78,	267
1125	Pairing vibrations study with the time-dependent Hartree-Fock-Bogoliubov theory. 2008 , 78,	87
1124	Adapting approximate-memory potentials for time-dependent density functional theory. 2008 , 77,	13
1123	Real-time resolution of the causality paradox of time-dependent density-functional theory. 2008 , 77,	66
1122	Turbo charging time-dependent density-functional theory with Lanczos chains. 2008 , 128, 154105	207

1121	Wave function and external potential from constrained search in density-functional theory. 2008 , 78,		
1120	Theory of electronic transport in random alloys with short-range order: Korringa-Kohn-Rostoker nonlocal coherent potential approximation. 2008 , 77,	27	
1119	Stochastic time-dependent current-density-functional theory: A functional theory of open quantum systems. 2008 , 78,	34	
1118	Calculating interaction between a highly charged high-speed ion and a solid surface. 2008 , 77,	9	
1117	Electronic excitation in an Ar ⁷⁺ ion traversing a graphene sheet: Molecular dynamics simulations. 2008 , 77,	24	
1116	Ground-state reference systems for expanding correlated fermions in one dimension. 2008 , 78,	51	
1115	Dynamical effects in the interaction of ion beams with solids. <i>Physical Review Letters</i> , 2008 , 100, 103201	7.4	21
1114	Time-resolved Dyson equations in the context of time-dependent density-functional theory: Extension to solid systems. 2008 , 78,	3	
1113	Successes and failures of Bethe ansatz density functional theory. 2008 , 78,	23	
1112	Linear density response function within the time-dependent exact-exchange approximation. 2008 , 78,	59	
1111	Ultrafast manipulation of electron spins in a double quantum dot device: A real-time numerical and analytical study. 2008 , 78,	17	
1110	Time-dependent density-functional theory with a self-interaction correction. <i>Physical Review Letters</i> , 2008 , 101, 096404	7.4	39
1109	High-level correlated approach to the jellium surface energy, without uniform-gas input. <i>Physical Review Letters</i> , 2008 , 100, 036401	7.4	63
1108	First-principles calculations of dielectric and optical properties of MgB ₂ . 2008 , 78,	21	
1107	Collective excitations in one-dimensional ultracold Fermi gases: Comparative study. 2008 , 78,	24	
1106	Accurate density functional calculations of near-edge x-ray and optical absorption spectra of liquid water using nonperiodic boundary conditions: the role of self-interaction and long-range effects. <i>Physical Review Letters</i> , 2008 , 100, 107401	7.4	42
1105	Spin-charge separation in molecular wire conductance simulations. 2008 , 78,	23	
1104	Time-dependent density-functional theory and strongly correlated systems: insight from numerical studies. <i>Physical Review Letters</i> , 2008 , 101, 166401	7.4	105

1103	Optical excitations of defects in realistic nanoscale silica clusters: comparing the performance of density functional theory using hybrid functionals with correlated wavefunction methods. 2008 , 129, 014706	8
1102	Time-dependent density functional theory study of cobalt corrinoids: Electronically excited states of methylcobalamin. 2008 , 129, 085101	28
1101	Synthesis, Crystal Structure and Density Functional Studies on 1N-Phenyl-3-(2,4-Dichlorophenyl)-5-(4-Chlorophenyl)-2-Pyrazoline. 2008 , 55, 183-188	4
1100	Bibliography. 2008 , 297-314	
1099	Synthesis, crystal structure and quantum chemical study on 3-phenylamino-4-phenyl-1,2,4-triazole-5-thione. 2009 , 14, 608-20	16
1098	Alloying effects on the optical properties of Ge _{1-x} Si _x nanocrystals from time-dependent density functional theory and comparison with effective-medium theory. 2009 , 79,	5
1097	Time-dependent density-functional study of field emission from tipped carbon nanotubes. 2009 , 80,	9
1096	Density functional theory and quantum computation. 2009 , 79,	8
1095	Electron-hole spectra created by adsorption on metals from density functional theory. 2009 , 79,	31
1094	Exact-exchange kernel of time-dependent density functional theory: frequency dependence and photoabsorption spectra of atoms. 2009 , 131, 044110	35
1093	Effects of quantum tunneling in metal nanogap on surface-enhanced Raman scattering. 2009 , 94, 243102	62
1092	Kohn-Sham scheme for frequency-dependent linear response. 2009 , 79,	
1091	Local Hartree-exchange and correlation potential defined by local force equations. 2009 , 80,	16
1090	Density-functional-theory investigation of antiproton-helium collisions. 2009 , 80,	12
1089	Time-dependent internal density functional theory formalism and Kohn-Sham scheme for self-bound systems. 2009 , 80,	5
1088	Absorption and emission properties of phenylene ethynylene oligomers: effect of substitution and E-conjugation length. 2009 , 107, 1629-1639	16
1087	Nonuniform light-matter interaction theory for near-field-induced electron dynamics. 2009 , 80,	25
1086	Ground and excited-state fermions in a one-dimensional double-well: Exact and density-functional solutions. 2009 , 79,	13

- 1085 Linear continuum mechanics for quantum many-body systems. *Physical Review Letters*, **2009**, 103, 086401. 7.4 20
- 1084 Rabi oscillations and few-level approximations in time-dependent density functional theory. *Physical Review Letters*, **2009**, 102, 233001 7.4 61
- 1083 Blindness of the exact density response function to certain types of electronic excitations: implications for time-dependent density-functional theory. *Physical Review Letters*, **2009**, 102, 233003 7.4 12
- 1082 Collective modes of trapped superfluid Fermi gases in the BCS-BEC crossover beyond the mean-field approximation. **2009**, 79, 13
- 1081 Photoabsorption spectra of small cationic xenon clusters from time-dependent density functional theory. **2009**, 131, 214302 4
- 1080 Large amplitude dynamics of the pairing correlations in a unitary fermi gas. *Physical Review Letters*, **2009**, 102, 085302 7.4 36
- 1079 Low-energy collective electronic excitations in Pd metal. **2009**, 80, 22
- 1078 Trajectory surface-hopping study of methane photodissociation dynamics. **2009**, 131, 224320 26
- 1077 Real space calculation of optical constants from optical to x-ray frequencies. **2009**, 80, 26
- 1076 Many-particle Hamiltonian for open systems with full Coulomb interaction: Application to classical and quantum time-dependent simulations of nanoscale electron devices. **2009**, 79, 42
- 1075 Exchange and correlation in molecular wire conductance: nonlocality is the key. **2009**, 131, 034106 15
- 1074 Modulation of alternating electric field inside photoexcited carbon nanotubes. **2009**, 95, 053109 19
- 1073 Exact dynamical exchange-correlation kernel of a weakly inhomogeneous electron gas. *Physical Review Letters*, **2009**, 102, 113001 7.4 16
- 1072 SYNTHESIS AND NLO PROPERTIES OF NEW CHROMOPHORES BASED ON IMIDAZO[1,2-A]PYRIDINE. **2009**, 196, 1466-1474 4
- 1071 High-order harmonics from C60 fullerenes -toward attosecond dynamics of complex systems. **2009**, 19
- 1070 Simplification of the time-dependent generalized self-interaction correction method using two sets of orbitals: Application of the optimized effective potential formalism. **2009**, 80, 9
- 1069 Assessment of advanced energy functionals in an exactly solvable model system. **2009**, 79, 5
- 1068 Hydrodynamic perspective on memory in time-dependent density-functional theory. **2009**, 79, 19

1067	On the existence of effective potentials in time-dependent density functional theory. 2009 , 42, 425207	13
1066	REAL-TIME APPROACH TO TIME-DEPENDENT DENSITY-FUNCTIONAL THEORY IN THE LINEAR AND NONLINEAR REGIME. 2009 , 08, 561-574	15
1065	TDDFT APPROACH TO PHOTOABSORPTION IN EVEN-EVEN NUCLEI. 2009 , 24, 2159-2167	
1064	TS-1 from first principles. 2009 , 113, 15006-15	47
1063	Time-dependent current density functional theory via time-dependent deformation functional theory: a constrained search formulation in the time domain. 2009 , 11, 4621-30	14
1062	Calculation of atomic excitation energies by time-dependent density functional theory within a modified linear response. 2009 , 21, 064229	3
1061	Double excitations in finite systems. 2009 , 130, 044108	85
1060	Representation independent algorithms for molecular response calculations in time-dependent self-consistent field theories. 2009 , 130, 054111	81
1059	Coupled-cluster and density functional theory studies of the electronic excitation spectra of trans-1,3-butadiene and trans-2-propeniminium. 2009 , 131, 024301	43
1058	Dynamics simulation of a π -conjugated light-harvesting dendrimer II: phenylene-based dendrimer (phDG2). 2009 , 21, 064217	2
1057	Theoretical study of the time-dependent phenomenon of photon-assisted tunneling through a charged quantum dot. 2009 , 21, 064230	2
1056	First-principles calculation of the electron dynamics in crystalline SiO(2). 2009 , 21, 064224	37
1055	Double spin-flip approach within equation-of-motion coupled cluster and configuration interaction formalisms: Theory, implementation, and examples. 2009 , 130, 044103	82
1054	An exploration of electronic structure and nuclear dynamics in tropolone: II. The A (1)B2 (π^* π) excited state. 2009 , 130, 144304	21
1053	Relationships between the third-order reactivity indicators in chemical density-functional theory. 2009 , 130, 244105	31
1052	All-electron calculation of nonadiabatic couplings from time-dependent density functional theory: Probing with the Hartree-Fock exact exchange. 2009 , 131, 114101	28
1051	Atoms in boxes: From confined atoms to electron-atom scattering. 2009 , 131, 104108	22
1050	Must Kohn-Sham oscillator strengths be accurate at threshold?. 2009 , 131, 114308	4

1049	Excited state properties of liquid water. 2009 , 21, 033101	21
1048	Electronic transport calculations for rough interfaces in Al, Cu, Ag, and Au. 2009 , 21, 315001	5
1047	Ferrocenyl-ended thieno-vinylene oligomers: donor-acceptor polarization and mixed-valence properties with emphasis on the raman mapping of localized-to-delocalized transitions. 2009 , 15, 2548-59	19
1046	trans-cis photoisomerization of azobenzene-conjugated dithiolato-bipyridine platinum(II) complexes: extension of photoresponse to longer wavelengths and photocontrollable tristability. 2009 , 15, 1429-39	48
1045	Thiophene-diazine molecular semiconductors: synthesis, structural, electrochemical, optical, and electronic structural properties; implementation in organic field-effect transistors. 2009 , 15, 5023-39	76
1044	Synthesis and properties of salen-aluminum complexes as a novel class of color-tunable luminophores. 2009 , 15, 6478-87	46
1043	Oxidation of end-capped pentathienoacenes and characterization of their radical cations. 2009 , 15, 12346-61	16
1042	Synthesis, spectroscopy, nonlinear optics, and theoretical investigations of thienylethynyl octopoles with a tunable core. 2009 , 15, 8223-34	13
1041	Conformational Analysis and Optical Rotation of Carene β -Amino Alcohols: A DFT Study. 2009 , 2009, 4600-4605	8
1040	Theoretical study of electronic properties of organic photovoltaic materials. 2009 , 30, 1027-37	17
1039	Ab initio theory for treating local electron excitations in molecules and its performance for computing optical properties. 2009 , 30, 2213-30	13
1038	Time-dependent density functional calculations on the electronic spectra of the neutral nickel complex [Ni(LISQ) ₂] (LISQ = 3,5-di-tert-butyl-o-diiminobenzosemiquinonate(1-)) and its monoanion and dication. 2009 , 30, 2087-98	5
1037	Excited states of OsO ₄ : a comprehensive time-dependent relativistic density functional theory study. 2010 , 31, 532-51	18
1036	DFT/TD-DFT investigation on Ir(III) complexes with N-heterocyclic carbene ligands: geometries, electronic structures, absorption, and phosphorescence properties. 2010 , 31, 628-38	25
1035	Evaluation of exchange-correlation functionals for time-dependent density functional theory calculations on metal complexes. 2010 , 31, 1008-14	24
1034	Intramolecular electronic communication in a dimethylaminoazobenzene-fullerene C ₆₀ dyad: an experimental and TD-DFT study. 2010 , 31, 1182-94	5
1033	Azure A chloride: computational and spectroscopic study. 2009 , 40, 176-182	16
1032	Theoretical modeling and experimental studies on N-n-decyl-2-oxo-5-nitro-1-benzylidene-methylamine. 2009 , 15, 1281-90	39

1031	Phase-space explorations in time-dependent density functional theory. 2009 , 914, 30-37	10
1030	Time-dependent closed and open-shell density functional theory from the perspective of partitioning techniques and projections. 2009 , 914, 50-59	1
1029	Time-dependent density-functional theory for molecules and molecular solids. 2009 , 914, 1-2	27
1028	Theoretical study of orientation-dependent multiphoton ionization of polyatomic molecules in intense ultrashort laser fields: A new time-dependent Voronoi-cell finite difference method. 2009 , 366, 91-102	33
1027	Theoretical study on the electronic structures and optical properties of blue phosphorescent iridium(III) complexes. 2009 , 919, 204-209	3
1026	Heating electrons with ion irradiation: A first-principles approach. 2009 , 267, 590-593	4
1025	ABINIT: First-principles approach to material and nanosystem properties. 2009 , 180, 2582-2615	2006
1024	The challenge of predicting optical properties of biomolecules: What can we learn from time-dependent density-functional theory?. 2009 , 10, 469-490	19
1023	Foreword. 2009 , 10, 465-468	
1022	Almost completely dedoped electrochemically deposited luminescent films exhibiting excellent LED performance. 2009 , 54, 7006-7011	27
1021	Electron impact ionisation and UV absorption study of α - and β -pinene. 2009 , 280, 169-173	15
1020	Dissipative Time Dependent Density Functional Theory. 2009 , 48, 2660-2664	5
1019	Introduction to theory/modeling methods in photosynthesis. 2009 , 102, 437-41	8
1018	Comparative study on two 2-pyrazoline derivatives with experimental and theoretical methods. 2009 , 20, 443-451	6
1017	Synthesis, characterization and spectra studies on Zn(II) and Cu(II) complexes with thiocarbamide ligand containing Schiff base group. 2009 , 20, 995-1003	22
1016	Theoretical study of the electronic structures and spectroscopic properties of $[(4\text{-X-C}_6\text{H}_4\text{C}_6\text{H}_4\text{N}_3)\text{PtCl}]^+$ (trpy = 2,2':6',2''-terpyridine; X = H, Me and Ph). 2009 , 52, 2226-2236	1
1015	Time-dependent relativistic density functional study of Yb and YbO. 2009 , 52, 1945-1953	23
1014	Theoretical studies on electronic structures and spectroscopic properties of a series of novel β -diketonate Os(II) complexes. 2009 , 122, 31-42	4

1013	Current density functional theory for one-dimensional fermions. 2009 , 246, 941-947	11
1012	Electronic properties of extended graphene nanomaterials from GW calculations. 2009 , 246, 2572-2576	23
1011	A DFT/TD-DFT study for the ground and excited states of peramine and some pyrrolopyrazinone compounds. 2009 , 22, 1058-1064	2
1010	Prediction of excited state energies for molecular nitrogen using quantum Monte Carlo methods. 2009 , 109, 43-49	2
1009	Computational studies on the spectroscopic properties of the 2-pyridylpyrazolate-based platinum(II) complexes with modified pyrazolate fragment. 2009 , 109, 308-319	6
1008	Theoretical study on the spectroscopic properties and electronic structures of heteroleptic phosphorescent Ir(III) complexes. 2009 , 109, 1167-1176	18
1007	Intramolecular charge-transfer excitation energies from range-separated hybrid functionals using the Yukawa potential. 2009 , 109, 1905-1914	28
1006	Low-lying excited states of 7-aminocoumarin derivatives: A theoretical study. 2009 , 109, 1940-1949	12
1005	Electron invariants and excited state structural analysis for electronic transitions within CIS, RPA, and TDDFT models. 2009 , 110, NA-NA	35
1004	Relativistic adiabatic time-dependent density functional theory using hybrid functionals and noncollinear spin magnetization. 2009 , 109, 2091-2112	76
1003	Identification of singlet excited electronic states of hydroxybenzoic acid isomers. 2009 , 106, 334-342	9
1002	Identification of singlet excited electronic states of toluidine isomers. 2009 , 107, 244-251	3
1001	Photostability and spectral properties of fluorinated fluoresceins and their biarsenical derivatives: a combined experimental and theoretical study. 2009 , 85, 1082-8	13
1000	Microscopic studies of atom-water collisions. 2009 , 285, 143-148	27
999	Attosecond control of the dissociative ionization via electron localization: A comparison between D2 and CO. 2009 , 366, 139-147	43
998	Synthesis, crystal structures and density functional theory study of dimeric copper(II) complexes of a tridentate Schiff-base ligand. 2009 , 362, 4081-4086	9
997	Linear fully conjugated meso-aryl pentapyrrins. 2009 , 50, 6909-6912	6
996	Role played by the organometallic fragment on the first hyperpolarizability of iron-acetylide complexes: A TD-DFT study. 2009 , 900, 110-117	17

995	Illustration of a TDDFT spatial overlap diagnostic by basis function exponent scaling. 2009 , 914, 110-114	35
994	From linear quaterthiophene to sulflower: A comparative theoretical study. 2009 , 912, 27-31	9
993	The absorption, emission spectra as well as ground and excited states calculations of some dimethine cyanine dyes. 2009 , 906, 50-55	19
992	Prevalence of the adiabatic exchange-correlation potential approximation in time-dependent density functional theory. 2009 , 914, 19-21	18
991	Application of time-dependent density-functional theory to molecules and nanostructures. 2009 , 914, 115-129	12
990	Is charge transfer transitions really too difficult for standard density functionals or are they just a problem for time-dependent density functional theory based on a linear response approach. 2009 , 914, 106-109	53
989	Approximate time-dependent density functional theory. 2009 , 914, 38-49	66
988	Environmental effects on electronic absorption spectra using DFT: An organic and positively charged fused polycyclic chromophore as a case study. 2009 , 914, 94-99	22
987	Time-dependent density-functional theory for molecules and molecular solids. 2009 , 914, 3-18	476
986	A theoretical study on the ground and excited state behaviors of TTBC related carbocyanine dyes. 2009 , 915, 149-159	11
985	Inelastic X-ray scattering and first-principles study of electron excitations in MgB ₂ . 2009 , 149, 1706-1711	1
984	Synthesis, DNA-binding and spectral properties of novel complexes [RuL ₂ (idpq)] ²⁺ (L = bpy, phen) with embedded CO. 2009 , 920, 163-171	16
983	A discrete action principle for electrodynamics and the construction of explicit symplectic integrators for linear, non-dispersive media. 2009 , 228, 3421-3432	6
982	A novel series of iridium complexes with alkenylquinoline ligands: Theoretical study on electronic structure and spectroscopic property. 2009 , 694, 150-156	4
981	Electronic structures and optical properties of neutral substituted fluorene-based cyclometalated platinum(II)acetylide complexes: A DFT exploration. 2009 , 694, 1848-1860	24
980	Four-coordinate boron compounds derived from 2-(2-pyridyl)phenol ligand as novel hole-blocking materials for phosphorescent OLEDs. 2009 , 694, 1922-1928	31
979	Experimental and theoretical characterization of Ru(II) complexes with polypyridine and phosphine ligands. 2009 , 694, 3781-3792	9
978	Synthesis, photophysical properties, and theoretical studies on pyrrole-containing bromo Re(I) complex. 2009 , 694, 3742-3748	63

977	Solvent effects on the absorption and emission of [Re(R2bpy)(CO)3X] complexes and their sensitivity to CO2 in solution. 2009 , 204, 174-182	29
976	Photoinduced electron dynamics at the chromophore-semiconductor interface: A time-domain ab initio perspective. 2009 , 84, 30-68	155
975	Experimental and theoretical comparative studies on two 2-pyrazoline derivatives. 2009 , 74, 87-93	11
974	Ortho effects on the change in electronic absorption spectrum of pyridinium salts of saturated bromohydrocarbon. 2009 , 74, 1084-9	2
973	Molecular structure and vibrational assignments of hippuric acid: a detailed density functional theoretical study. 2009 , 74, 1197-203	46
972	Quantum chemical and spectroscopic investigations of 5-aminoquinoline. 2009 , 74, 1215-23	19
971	Excited state geometry optimizations by time-dependent density functional theory based on the fragment molecular orbital method. 2009 , 474, 227-232	30
970	Pyridine as axial ligand on the [Mo6Cl8]4+ core switches off luminescence. 2009 , 475, 232-234	20
969	Ab initio absorption spectra of 3-tert-butylcyclohexene. 2009 , 10, 491-503	1
968	Photoelectron and UV absorption spectroscopy for determination of electronic configurations of negative molecular ions: Chlorophenols. 2009 , 171, 37-46	10
967	On the exact treatment of time-dependent self-interaction correction. 2009 , 324, 955-976	24
966	QUANTUM ESPRESSO: a modular and open-source software project for quantum simulations of materials. 2009 , 21, 395502	13251
965	Extensive TD-DFT Benchmark: Singlet-Excited States of Organic Molecules. 2009 , 5, 2420-35	799
964	Mixed Si-Ge nanoparticle quantum dots: a density functional theory study. 2009 , 72, 193-201	5
963	Optical absorption spectra of Ag11 isomers. 2009 , 52, 199-202	6
962	The role of homophase and heterophase interfaces on transport properties in structured materials. 2009 , 177, 83-101	7
961	Electrophilicity index within a conceptual DFT framework. 2009 , 105, 13	143
960	Accurate simulation of optical properties in dyes. 2009 , 42, 326-34	404

959	A TD-DFT investigation of ground and excited state properties in indoline dyes used for dye-sensitized solar cells. 2009 , 11, 11276-84	144
958	Different electronic structures and spectroscopic properties of cationic $[M(ppy)_2(N-N)]^+$ ($M = Rh, Ir$; $N-N = Hcmbpy, H2dcbpy$), a DFT study. 2009 , 11, 6051-9	21
957	Strong field ionization of many electron systems: A quantum chemical challenge. 2009 , 19, 1521-1534	9
956	Excited states of biological chromophores studied using many-body perturbation theory: Effects of resonant-antiresonant coupling and dynamical screening. 2009 , 80,	80
955	Photodissociation of Co-C bond in methyl- and ethylcobalamin: an insight from TD-DFT calculations. 2009 , 113, 6898-909	44
954	Revisiting Molecular Dissociation in Density Functional Theory: A Simple Model. 2009 , 5, 770-80	71
953	Time-dependent density-functional theory simulation for electron-ion dynamics in molecules under intense laser pulses. 2009 , 21, 064222	21
952	Polarization and spectral shift of benzophenone in supercritical water. 2009 , 113, 5112-8	24
951	ONIOM calculation on azurin: effect of metal ion substitutions. 2009 , 113, 8615-25	7
950	Excitation Energies in Time-Dependent (Current-) Density-Functional Theory: A Simple Perspective. 2009 , 5, 859-65	12
949	Luminescence vapochromism of a platinum(II) complex for detection of low molecular weight halohydrocarbon. 2009 , 48, 10202-10	73
948	Excited states of DNA base pairs using long-range corrected time-dependent density functional theory. 2009 , 113, 9761-5	63
947	Effect of periodic replacement of the heteroatom on the spectroscopic properties of indole and benzofuran derivatives. 2009 , 113, 464-73	28
946	Strong deviations from jellium behavior in the valence electron dynamics of potassium. 2009 , 80,	12
945	Dynamic current suppression and gate voltage response in metal-molecule-metal junctions. 2009 , 9, 2671-5	28
944	Temperature-dependent resistivity of ferromagnetic $Ga_{1-x}Mn_xAs$: Interplay between impurity scattering and many-body effects. 2009 , 80,	4
943	Photoexcitation of a light-harvesting supramolecular triad: a time-dependent DFT study. 2009 , 113, 5345-9	39
942	Unrestricted algebraic diagrammatic construction scheme of second order for the calculation of excited states of medium-sized and large molecules. 2009 , 130, 024104	60

941	Two-Photon Absorption Properties of New Fluorene-Based Singlet Oxygen Photosensitizers. 2009 , 113, 4706-4711	45
940	Asymmetry between absorption and photoluminescence line shapes of TPD: spectroscopic fingerprint of the twisted biphenyl core. 2009 , 113, 315-24	32
939	Many-Body Perturbation Theory Extended to the Quantum Mechanics/Molecular Mechanics Approach: Application to Indole in Water Solution. 2009 , 5, 1822-8	13
938	Solvent fluctuations drive the hole transfer in DNA: a mixed quantum-classical study. 2009 , 113, 13107-17	68
937	Circular dichroism of chiral 1,8,15,22-tetra(alkoxyl)phthalocyaninato lead and yttrium complexes: time-dependent density functional theory calculations. 2009 , 113, 12179-86	8
936	Modified Ehrenfest Formalism for Efficient Large-Scale ab initio Molecular Dynamics. 2009 , 5, 728-42	95
935	Photodissociation of [Fe(x)(C ₂₄ H ₁₂) _y] ⁺ complexes in the PIRENEA setup: iron-polycyclic aromatic hydrocarbon clusters as candidates for very small interstellar grains. 2009 , 113, 4878-88	26
934	Qualitatively Correct Charge-Transfer Excitation Energies in HeH(+) by Time-Dependent Density-Functional Theory Due to Exact Exchange Kohn-Sham Eigenvalue Differences. 2009 , 5, 781-5	20
933	Dynamics simulations of atmospherically relevant molecular reactions. 2009 , 28, 207-222	33
932	DFT-based methods in the design of two-photon operated molecular switches. 2009 , 113, 7080-9	48
931	Ab initio excited state properties and dynamics of a prototype sigma-bridged-donor-acceptor molecule. 2009 , 113, 9595-602	19
930	High-order harmonic generation from C ₆₀ -rich plasma. 2009 , 80,	46
929	Transient electronic dynamics of noninteracting open systems beyond linear response. 2009 , 21, 355301	13
928	Theoretical studies on electronic structures, spectra and charge transporting properties of a series of Pt(CN) ₂ complexes. 2009 , 159, 1090-1098	6
927	Time-independent excited-state density functional theory: study of 1s22p3(4S) and 1s22p3(2D) states of the boron isoelectronic series up to Ne ⁵⁺ . 2009 , 42, 015003	13
926	The influence of initial conditions on charge transfer dynamics. 2009 , 11, 10293-8	20
925	An Extremely High Molar Extinction Coefficient Ruthenium Sensitizer in Dye-Sensitized Solar Cells: The Effects of Conjugation Extension. 2009 , 113, 14559-14566	111
924	Spin-Orbit Coupling: Effects in Heavy Element Chemistry. 2009 ,	

923	Density functional theory for transition metals and transition metal chemistry. 2009 , 11, 10757-816	1248
922	Singlet-Triplet Transitions in Real-Time Time-Dependent Hartree-Fock/Density Functional Theory. 2009 , 5, 2415-9	43
921	Long-range intermolecular charge transfer induced by laser pulses: an explicitly time-dependent configuration interaction approach. 2009 , 11, 3875-84	24
920	The effect of local environment on photoluminescence: a time-dependent density functional theory study of silanone groups on the surface of silica nanostructures. 2009 , 131, 034705	28
919	Time-dependent density functional theory versus Bethe-Salpeter equation: an all-electron study. 2009 , 11, 4451-7	55
918	Time-dependent density functional theory for resonant properties: resonance enhanced Raman scattering from the complex electric-dipole polarizability. 2009 , 11, 4539-48	38
917	Autoionizing resonances in time-dependent density functional theory. 2009 , 11, 4655-63	17
916	Time-dependent density functional theory of high excitations: to infinity, and beyond. 2009 , 11, 4437-50	27
915	Excitation energies of zinc porphyrin in aqueous solution using long-range corrected time-dependent density functional theory. 2009 , 113, 6041-3	58
914	Application of quantum chemistry to nanotechnology: electron and spin transport in molecular devices. 2009 , 38, 2319-33	109
913	Optoelectronic properties of natural cyanin dyes. 2009 , 113, 8801-10	22
912	Charge-transfer excitation energies with a time-dependent density-functional method suitable for orbital-dependent exchange-correlation kernels. 2009 , 80,	46
911	Quantum Mechanical Studies of the Photophysics of DNA and RNA Bases. 2009 , 285-339	7
910	Prediction of spin-spin coupling constants in solution based on combined density functional theory/molecular mechanics. 2009 , 130, 134508	46
909	Enhancement of high-order harmonic generation using a two-color pump in plasma plumes. 2009 , 80,	75
908	Excitation energies with time-dependent density matrix functional theory: Singlet two-electron systems. 2009 , 130, 114104	50
907	Excited States from Time-Dependent Density Functional Theory. 2009 , 91-165	115
906	Time-independent (static) density-functional theories for pure excited states: Extensions and unification. 2009 , 80,	40

905	All-electron density functional theory and time-dependent density functional theory with high-order finite elements. 2009 , 131, 054103	43
904	Two-component relativistic density functional method for computing nonsingular complex linear response of molecules based on the zeroth order regular approximation. 2009 , 130, 194102	38
903	Absorption and fluorescence properties of oligothiophene biomarkers from long-range-corrected time-dependent density functional theory. 2009 , 11, 4498-508	131
902	Solvent effects on isolated formamide and its monohydrated complex: observations from PCM study. 2009 , 33, 1709	12
901	Kadanoff-Baym approach to quantum transport through interacting nanoscale systems: From the transient to the steady-state regime. 2009 , 80,	170
900	Examination of DFT and TDDFT methods II. 2009 , 113, 10873-9	18
899	Boron carbides from first principles. 2009 , 176, 012002	68
898	Tuning electronic properties of functionalized polyhedral oligomeric silsesquioxanes: a DFT and TDDFT study. 2009 , 113, 9707-14	42
897	Novel Re(II) dendrimers: synthesis, characterization and theoretical studies. 2009 , 10592-600	22
896	Photodegradation mechanism of nonsteroidal anti-inflammatory drugs containing thiophene moieties: suprofen and tiaprofenic acid. 2009 , 113, 11306-13	20
895	Selectivity in binary fluid mixtures: static and dynamical properties. 2009 , 80, 021409	25
894	Spectroscopic Properties Obtained from Time-Dependent Density Functional Theory (TD-DFT). 2009 ,	1
893	Electronic Structure and Charge-Transport Parameters of Functionalized Tetracene Crystals: Impact of Partial Fluorination and Alkyl or Alkoxy Derivatization. 2009 , 21, 3593-3601	41
892	Luminescent boron-contained ladder-type pi-conjugated compounds. 2009 , 48, 7230-6	82
891	Molecular design of photoactive acenes for organic photovoltaics. 2009 , 130, 194701	18
890	Quartic-Scaling Analytical Gradient of Quasidegenerate Scaled Opposite Spin Second-Order Perturbation Corrections to Single Excitation Configuration Interaction. 2009 , 5, 1224-36	23
889	Performance of quasi-degenerate scaled opposite spin perturbation corrections to single excitation configuration interaction for excited state structures and excitation energies with application to the Stokes shift of 9-methyl-9,10-dihydro-9-silaphenanthrene. 2009 , 113, 10564-76	21
888	On the relation between time-dependent and variational density functional theory approaches for the determination of excitation energies and transition moments. 2009 , 130, 154102	90

887	One- and two-photon spectra of platinum acetylide chromophores: a TDDFT study. 2009 , 113, 13943-52	26
886	Fragment molecular orbital calculations on red fluorescent proteins (DsRed and mFruits). 2009 , 113, 1153-61	36
885	Photoabsorption spectra from adiabatically exact time-dependent density-functional theory in real time. 2009 , 11, 4631-9	27
884	Ultrafast electronic excitations of small sodium clusters and the onset of electron thermalization. 2009 , 11, 349-57	25
883	Nonlinear time-dependent density functional theory studies of the ionization of CO ₂ by ultrashort intense laser pulses. 2009 , 87, 1081-1089	2
882	Porphyrin dimers: A theoretical understanding of the impact of electronic coupling strength on the two-photon absorption properties. 2009 , 19, 7545	21
881	Physical signatures of discontinuities of the time-dependent exchange-correlation potential. 2009 , 11, 4647-54	19
880	Chemical and protein shifts in the spectrum of the photoactive yellow protein: a time-dependent density functional theory/molecular mechanics study. 2009 , 11, 4556-63	27
879	Density functional study of collective electron localization: detection by persistent current. 2009 , 21, 155602	2
878	A time-dependent density-functional approach to nonadiabatic electron-nucleus dynamics: formulation and photochemical application. 2009 , 11, 4570-8	27
877	Excitation energies from ground-state density-functionals by means of generator coordinates. 2009 , 11, 4564-9	3
876	Effect of ring fusion on the amplified spontaneous emission properties of oligothiophenes. 2009 , 19, 6556	14
875	Time-dependent current-density functional theory for generalized open quantum systems. 2009 , 11, 4509-22	36
874	Spectral properties of self-assembled squaraine-tetralactam: a theoretical assessment. 2009 , 11, 1258-62	57
873	Confinement effects on excitation energies and regioselectivity as probed by the Fukui function and the molecular electrostatic potential. 2009 , 11, 2862-8	28
872	Correlation-induced corrections to the band structure of boron nitride: a wave-function-based approach. 2009 , 131, 044119	12
871	Excited state surfaces in density functional theory: a new twist on an old problem. 2009 , 131, 091101	150
870	Nonadiabatic coupling vectors within linear response time-dependent density functional theory. 2009 , 130, 124107	102

869	Optimizing conical intersections by spin-flip density functional theory: application to ethylene. 2009 , 113, 12749-53	131
868	Exciton-plasmon States in nanoscale materials: breakdown of the Tamm-Dancoff approximation. 2009 , 9, 2820-4	107
867	Time-dependent density-functional theory for nonadiabatic electronic dynamics. <i>Physical Review Letters</i> , 2009 , 102, 053002	7.4 7
866	Improving approximate-optimized effective potentials by imposing exact conditions: Theory and applications to electronic statics and dynamics. 2009 , 80,	8
865	Injection, transport, absorption and phosphorescence properties of a series of blue-emitting Ir(III) emitters in OLEDs: a DFT and time-dependent DFT study. 2009 , 48, 7740-9	103
864	TDDFT diagnostic testing and functional assessment for triazene chromophores. 2009 , 11, 4465-70	140
863	Density-functional studies of plasmons in small metal clusters. 2009 , 130, 174701	33
862	Reconstructing the adiabatic exchange-correlation kernel of time-dependent density-functional theory. 2009 , 80,	8
861	Prediction of charge-transfer excitations in coumarin-based dyes using a range-separated functional tuned from first principles. 2009 , 131, 244119	270
860	Density functional theory calculation of ionization in antiproton-helium collisions. 2009 , 194, 072016	
859	Laser-Driven Molecular Dissociation: Time-Dependent Density Functional Theory and Molecular Dynamics Simulations. 2009 , 78, 094707	12
858	Extension of Frozen-orbital Analysis to the TammDancoff Approximation to Time-dependent Density Functional Theory. 2009 , 38, 528-529	9
857	Photoinduced conductivity of a porphyrin-gold composite nanowire. 2009 , 113, 4549-56	33
856	Artificial damping in the Kadanoff-Baym dynamics of small Hubbard chains. 2010 , 220, 012016	1
855	Visualizing ElectronHole Separation in Photo-excited Polar Crystals. 2010 , 3, 047202	2
854	Application of Real-time Time-dependent Density Functional Theory with the CVB3LYP Functional to Core Excitations. 2010 , 39, 407-409	15
853	DFT calculations of molecular excited states using an orbital-dependent nonadiabatic exchange kernel. 2010 , 110, 199-210	2
852	Quantum-chemical and correlation study of deprotonation and complexation of 1-amino-4-hydroxyanthraquinone. 2010 , 80, 1986-1995	7

851	Modelling non-adiabatic processes using correlated electron-ion dynamics. 2010 , 77, 305-329	24
850	Comparison of the directional characteristics of swift ion excitation for two small biomolecules: glycine and alanine. 2010 , 60, 71-76	11
849	Exploration of dynamical regimes of irradiated small protonated water clusters. 2010 , 58, 131-136	15
848	Optical spectrum of proflavine and its ions. 2010 , 493, 33-36	5
847	High-efficiency dye-sensitized solar cells: the influence of lithium ions on exciton dissociation, charge recombination, and surface states. 2010 , 4, 6032-8	492
846	Acceleration of the Rate of the Heck Reaction through UV- and Visible-Light-Induced Palladium(II) Reduction. 2010 , 2, 1467-1476	29
845	Dynamic structure factor and dielectric function of silicon for finite momentum transfer: Inelastic x-ray scattering experiments and ab initio calculations. 2010 , 81,	39
844	Theoretical characterization of photoinduced electron transfer in rigidly linked donor-acceptor molecules: the fragment charge difference and the generalized Mulliken-Hush schemes. 2010 , 108, 2775-2789	15
843	Dependence of Excited State Potential Energy Surfaces on the Spatial Overlap of the Kohn-Sham Orbitals and the Amount of Nonlocal Hartree-Fock Exchange in Time-Dependent Density Functional Theory. 2010 , 6, 2315-24	94
842	Near-field-induced optical force on a metal particle and C60: Real-time and real-space electron dynamics simulation. 2010 , 82,	3
841	Modifying the variational principle in the action-integral-functional derivation of time-dependent density-functional theory. 2010 , 82,	5
840	General Runge-Gross-type theorem for dipole laser-matter interaction. 2010 , 81,	6
839	Brief review related to the foundations of time-dependent density functional theory. 2010 , 125, 427-432	10
838	The mechanisms of excited states in enzymes. 2010 , 125, 345-352	
837	Role of quantum chemical calculations in molecular biophysics with a historical perspective. 2010 , 125, 121-144	5
836	The K-band ϵ_{max} values of the ultraviolet-visible spectra of some hydrazones in ethanol by a TD-DFT/PCM approach. 2010 , 494, 198-201	3
835	Protolytic fluorescein species evaluated using chemometry and DFT studies. 2010 , 86, 15-24	56
834	Electron spectroscopies and inelastic processes in nanoclusters and solids: Theory and experiment. 2010 , 493, 237-319	43

833	Time-dependent density functional theory as a thermodynamics. 2010 , 943, 48-52	7
832	A coarse-grained density functional theory, chemical potential equalization and electric response in molecular systems. 2010 , 943, 178-182	4
831	Synthesis, spectroscopic properties, crystal structure and density functional studies of Cu(II) complex with 2-((dehydroabietylamine)methyl)-6-methoxyphenol. 2010 , 975, 194-199	9
830	Molecular orbital evaluation of charge flow dynamics in natural pigments based photosensitizers. 2010 , 16, 523-33	21
829	Experimental and quantum chemical calculational studies on 2-[(4-Fluorophenylimino)methyl]-3,5-dimethoxyphenol. 2010 , 16, 577-87	68
828	Effect of the Enhanced Cyan Fluorescent Protein framework on the UV/visible absorption spectra of some chromophores. 2010 , 2, 38-47	25
827	Theoretical studies on electronic spectroscopy and dynamics with the real-time time-dependent density functional theory. 2010 , 5, 11-28	7
826	Synthesis, Crystal Structure, Photophysical Properties, and DFT Calculations of a Bis(tetrathia-calix[4]arene) Tetracadmium Complex. 2010 , 21, 867-878	14
825	Correlated and idempotent Dirac first-order density matrices with identical diagonal Fermion density: a route to extract a one-body potential energy in TDDFT. 2010 , 47, 505-519	2
824	Synthesis, structures, and property studies on Zn(II), Ni(II), and Cu(II) complexes with a Schiff base ligand containing thiocarbamide group. 2010 , 21, 977-987	28
823	Experimental and computational studies on zwitterionic (E)-2-(1-(2-(4-methylphenylsulfonamido)ethyliminio)ethyl) phenolate. 2010 , 21, 1027-1036	7
822	The nature of transitions in electronic absorption spectra of radical cations of dipyrroles with phenylene bridging groups. 2010 , 59, 779-783	
821	. 2010 , 12, 18-27	7
820	Application of DFT Methods in the Study of VIVO2+Peptide Interactions. 2010 , 2010, 4697-4710	40
819	SoluteSolvent Hydrogen Bond Formation in the Excited State. Experimental and Theoretical Evidence. 2010 , 79-123	0
818	Relaxation Dynamics in Image Potential States at Solid Interfaces. 2010 , 75-97	
817	Low-Energy Collective Electronic Excitations at Metal Surfaces. 2010 , 167-188	
816	Quinoidal oligothiophenes: towards biradical ground-state species. 2010 , 16, 470-84	63

815	A theoretical and spectroscopic study of conformational structures of piroxicam. 2010 , 75, 901-7	12
814	Synthesis, characterization, crystal structure and DFT studies on 3-(1H-benzo[d][1,2,3]triazol-1-yl)-1-(4-ethylphenyl)-1-oxopropan-2-yl-4-ethylbenzoate. 2010 , 75, 1051-6	1
813	DFT computations and spectroscopic analysis of a pesticide: chlorothalonil. 2010 , 77, 36-44	54
812	Fluorescence emission and enhanced photochemical stability of Zn(II)-5-triethyl ammonium methyl salicylidene ortho-phenylendiiminate interacting with native DNA. 2010 , 104, 765-73	12
811	Scandium 2-mercaptobenzothiazolate: Synthesis, structure and electroluminescent properties. 2010 , 29, 400-404	10
810	X-ray structure, electronic properties and density functional calculations: trans-Dihalo (1-(4-phenylimino)-1-(phenylhydrazono)-propan-2-one) (4,4'-di-tert-butyl-2,2'-bipyridine) ruthenium(II) complexes. 2010 , 29, 3214-3219	9
809	Electronic characterization of all-thiophene conducting dendrimers: Molecules and assemblies. 2010 , 51, 308-315	9
808	Microscopic details of the sensing ability of 15-crown-5-ether functionalized poly(bithiophene). 2010 , 51, 4267-4272	11
807	Excited-state properties from ground-state DFT descriptors: A QSPR approach for dyes. 2010 , 28, 465-71	22
806	Product of alaptide synthesis: determination of the absolute configuration. 2010 , 53, 958-61	16
805	Synthesis, structure, spectroscopic properties, and theoretical studies of alkaline earth metal complexes of 1,3-bis(carboxymethyl)benzimidazolium. 2010 , 984, 39-50	11
804	Visible photodissociation spectroscopy of PAH cations and derivatives in the PIRENEA experiment. 2010 , 371, 16-23	34
803	Theoretical approximations to X-ray absorption spectroscopy of liquid water and ice. 2010 , 177, 135-157	118
802	Variations in electron density and bonding in the lowest $1\bar{g}$ state of H ₂ molecule under strong magnetic fields by using a time-dependent density functional theory. 2010 , 943, 65-70	14
801	Electronic structure and vibrational frequencies in dehydroacetic acid (DHA) transition-metal complexes: A DFT study. 2010 , 945, 78-84	14
800	DFT studies on thiophene acetylide Ru(II) complexes for nonlinear optics: Structure-function relationships and solvent effects. 2010 , 946, 33-42	22
799	DFT study on the electronic structure, energetics and spectral properties of several bis(organohydrazido(2-)) molybdenum complexes containing substituted phosphines and chloro atoms as ancillary ligands. 2010 , 957, 126-132	8
798	Theoretical study of 2-phenylbenzoxazole derivatives and derived phenolic Schiff compounds in gas and solution phases: Electronic structures and optical properties. 2010 , 960, 106-114	4

797	Phase field modeling of defects and deformation. 2010 , 58, 1212-1235	322
796	Absorption and emission spectra of 1,8-naphthalimide fluorophores: A PCM-TD-DFT investigation. 2010 , 372, 61-66	53
795	A fresh look at excitonically coupled chromophores from a Jahn-Teller perspective. 2010 , 377, 78-85	7
794	Real-time electron dynamics simulation of the adsorption of an oxygen molecule on Pt and Au clusters. 2010 , 486, 48-52	5
793	Photochromic molecular wires: Insights from theory. 2010 , 488, 193-197	31
792	Electronic absorption spectra of photoreaction intermediates of 7-hydroxyquinoline monomer in a low-temperature argon matrix and time-dependent density-functional-theory calculations. 2010 , 490, 46-49	7
791	Regioselectivity on the cooxidation of 5,6-dihydroxyindole and its 2-carboxy derivative from the quantum chemical calculations. 2010 , 490, 226-229	6
790	Effect of molecular adsorbates on the plasmon resonance of metallic nanoparticles. 2010 , 494, 255-259	12
789	Time-dependent density-functional theory method in the electron nuclear dynamics framework. 2010 , 496, 188-195	13
788	Density functional study on the effect of substituent group for the monomer of donor-acceptor copolymer. 2010 , 48, 2099-2107	14
787	Coulomb interactions and spin transport in semiconductors: The spin Coulomb drag effect. 2010 , 247, 235-247	7
786	Modification of response properties of the Be(0001) surface upon adsorption of a potassium monolayer: An ab initio calculation. 2010 , 247, 1849-1857	10
785	Many-body meets QM/MM: Application to indole in water solution. 2010 , 247, 1920-1924	3
784	Ab initio calculation of second harmonic generation in solids. 2010 , 247, 1984-1991	5
783	Is Cu involved in prion oligopeptide stability? Experiments and numerical simulations. 2010 , 110, 656-680	3
782	Theoretical study on the influence of ancillary ligand on the spectroscopic properties and electronic structures of phosphorescent Pt(II) complexes. 2010 , 110, 1142-1151	1
781	Absorption spectra of recently synthesised organic dyes: A TD-DFT study. 2010 , 110, 2121-2129	23
780	Molecular excitation spectra by TDDFT with the nonadiabatic exact exchange kernel. 2010 , 110, 2202-2220	23

- 779 Continuum, discrete, and explicit solvation models for describing the low-lying absorption spectrum of the pterin acid in aqueous environment. **2010**, 110, 2371-2377 3
- 778 Analytical energy gradients of Coulomb-attenuated time-dependent density functional methods for excited states. **2010**, 110, 2247-2255 19
- 777 Solvatochromic shift of the $\pi \rightarrow \pi^*$ transition in all-trans, cis-13, cis-11, cis-9, and cis-7 retinal isomers induced by water and methanol. **2010**, 110, 2076-2087 3
- 776 Fourteen easy lessons in density functional theory. **2010**, 110, 2801-2807 38
- 775 The adiabatic approximation in time-dependent density matrix functional theory: response properties from dynamics of phase-including natural orbitals. **2010**, 133, 174119 29
- 774 Collective Electron Dynamics in Metallic and Semiconductor Nanostructures. **2010**, 1-44 13
- 773 Harnessing molecular excited states with Lanczos chains. **2010**, 22, 074204 14
- 772 Synthesis, Crystal Structure and Density Functional Calculations on 1-Phenyl-3-p-fluorophenyl-5-p-chlorophenyl-2-pyrazoline. **2007**, 28, 1539-1544 9
- 771 Comparative Studies on Two Fluoro-Substituted 2-Pyrazoline Derivatives with Experimental and Theoretical Methods. **2009**, 30, 1061-1066 1
- 770 Experimental and Theoretical Studies on a Thiocarbamide Derivative Containing Schiff Base Groups. **2009**, 30, 1667-1670 5
- 769 Synthesis, Crystal Structure, Spectra Characterization and DFT Studies on a Di-Cycle Pyrazoline Derivative. **2010**, 31, 1875-1880 4
- 768 Experimental and Computational Approaches to the Molecular Structure of 3-(2-Mercaptopyridine)phthalonitrile. **2011**, 32, 673-680 52
- 767 The dynamic process of two-electron atom irradiated by intense laser pulse using time dependent quantum Monte Carlo method. **2012**, 61, 163203 2
- 766 NIR absorbing aromatic (antiaromatic) vinylogous carbasapphyrins (3.3.1.0.1) with built-in fused dipolar aromatic hydrocarbon: synthesis and characterization. 0
- 765 Color tuning of di-boron derived TADF emitters: molecular design and property prediction. **2021**, 9, 15309-15330 0
- 764 Hydrogen bonding interactions can decrease clar sextet character in acridone pigments. **2021**, 19, 9619-9623 0
- 763 Ultrafast relaxation investigated by photoelectron circular dichroism: an isomeric comparison of camphor and fenchone. **2021**, 23, 25612-25628 1
- 762 Introduction. **2021**, 1-8

761	Synthesis, Spectral Characterization, Electronic Structure and Biological Activity Screening of the Schiff Base 4-((4-Hydroxy-3-Methoxy-5-Nitrobenzylidene)Amino)-N-(Pyrimidin-2-yl)Benzene Sulfonamide from 5-Nitrovaniline and Sulphadiazene. 1-18	7
760	Direct orange 26 dye environmental degradation: experimental studies (UV, mass, and thermal) in comparison with computational exploration hydrogen bonding analysis of TD-DFT calculations. 2021 , 27, 325	
759	Thermal Fluctuations in Conjugation and their Effect on Calculated Excitation Energies: A Case Study on the Astaxanthin Carotenoid.	
758	Density functional theory calculations of large systems: Interplay between fragments, observables, and computational complexity. e1574	5
757	Algorithm advances and applications of time-dependent first-principles simulations for ultrafast dynamics. e1577	0
756	Multi-directions pure even harmonic simultaneous generation from planar molecules in linearly polarized laser fields. 2021 , 119, 151105	0
755	Exploring Dyson's Orbitals and Their Electron Binding Energies for Conceptualizing Excited States from Response Methodology. 2021 , 12, 9963-9972	2
754	Computational Protocol to Calculate the Phosphorescence Energy of Pt(II) Complexes: Is the Lowest Triplet Excited State Always Involved in Emission? A Comprehensive Benchmark Study. 2021 , 60, 17230-17240	6
753	Colloquium: Nonthermal pathways to ultrafast control in quantum materials. 2021 , 93,	20
752	Theoretical Insights into Energy Absorption and Charge Draining during Field Evaporation Assisted by Femtosecond Laser Pulses. 2021 , 125, 9338-9345	
751	Three-Dimensional Tomographic Imaging of CO Molecular Orbitals Revealing Multi-Electron Effects.	0
750	Recent Progress in Advanced Organic Photovoltaics: Emerging Techniques and Materials.	1
749	Local approaches for electric dipole moments in periodic systems and their application to real-time time-dependent density functional theory. 2021 , 155, 134116	2
748	Ketone Incorporation Extends the Emission Properties of the Xanthene Scaffold Beyond 1000 nm. 2021 ,	2
747	Excited-state theorem in both case of a nuclear potential and a more general external potential in correspondence to the Hohenberg-Kohn theorem. 2022 , 161, 110406	0
746	Quantal Density Functional Theory. 2002 , 355-377	
745	Numerical Simulation of Current Dependence on Well Widths in AlGaAs/GaAs/AlGaAs Double Barrier Diode Structures. 2003 , 103, 47-56	
744	Schrödinger Theory from the Perspective of Classical Fields Derived from Quantal Sources. 2004 , 7-47	

- 743 The Hohenberg-Kohn Theorems and Kohn-Sham Density Functional Theory. **2004**, 99-123 o
- 742 Defect and Carrier Dynamics in Nanotubes under Electronic Excitations: Time-Dependent Density Functional Approaches. **2004**, 141-148
- 741 References. **2005**, 405-440
- 740 Comparative Study of Time-Dependent and Scattering-State Ab Initio Calculations for Field Emission. **2005**, 3, 457-460 1
- 739 Ultra-Fast Dynamics in Nanocarbon Explored by TDDFT-MD Simulations. **2007**, 125-130
- 738 Ab Initio GW Calculations Using an All-Electron Approach. **2008**, 171-188
- 737 Einführung in die computergestützte Quantenchemie. **2008**, 391-413
- 736 Electronic Transport in Nanowires at Different Length Scales. **2008**, 404-420
- 735 Density Functional Theory and Car-Parrinello Molecular Dynamics Methods. **2008**, 487-499
- 734 Time-Dependent Density Functional Theory of Many-Electron Systems. **2009**,
- 733 High-Order Harmonic Generation from Nanostructured Material Using the Laser-Plasma Method. **2010**, 145-160
- 732 An Additive Long-range Potential to Correct for the Charge-transfer Failure of Time-dependent Density Functional Theory. **2010**, 21-34
- 731 Density-Functional Theory with Orbital-Dependent Functionals: Exact-exchange Kohn-Sham and Density-Functional Response Methods. **2010**, 35-52
- 730 A theoretical study on UV-spectroscopy, electronic structure and reactivity properties of sesquiterpenes.
- 729 Theoretical and Computational Methods. **2011**, 21-55
- 728 First-Principles Calculation for Photoionization Processes in Time-Dependent Density-Functional Theory. **2011**, 54, 522-528 1
- 727 A Novel Rhenium(I) Tricarbonyl Complex with 4,5-Diazafluoren-Driven Ligand: Synthesis, Spectroscopic and Theoretical Studies. **2011**, 83-88
- 726 First-principles Calculations of Interaction between Intense Optical Field and Materials. **2011**, 54, 518-521

725 Theoretical Background. **2011**, 67-100

724 Some Practical Considerations for Density Functional Theory Studies of Chemistry at Metal Surfaces. **2011**,

723 DFT and Time-dependant DFT Investigation of eElectronic Structure, Phosphorescence and Electroluminescence Properties of Iridium (III) Quinoxaline Complexes. **2011**, 55, 354-363

722 An Introduction to Linear-Scaling Ab Initio Calculations. **2012**, 1-35

721 Tensorial Aspects of Calculating Hubbard U Interaction Parameters. **2012**, 169-195

720 Geometric Aspects of Representation Optimisation. **2012**, 125-149

719 Simulation of X-RayMatter Interaction. 215-237

718 Electron correlation methods based on the random phase approximation. **2012**, 103-120

717 First-Principles Calculations of Physical Properties of Planetary Ices. **2013**, 149-169

716 Effects of mutations on the absorption spectra of copper proteins: a QM/MM study. **2014**, 39-47

715 The Role of Trajectories in Quantum Chemistry and Chemical Physics. **2012**, 235-301

714 Massive Computation for Femtosecond Dynamics in Condensed Matters. **2013**, 155-168

713 Time-dependent density functional theory study of charge transfer in collisions. **2014**, 241-250

712 Multi-configuration time dependent Hartree Fock method in three different representations. **2013**, 62, 123202

711 Theoretical Spectroscopy. **2013**, 45-58

1

710 Theoretical Methods. **2013**, 45-65

709 A Safari Through Density Functional Theory. **2013**, 465-478

1

708 High order harmonics generation in laser surface ablation: current trends. **2013**, 183, 815-847

3

- 707 Theoretical study of the ground and excited state properties of newly designed size-expanded adenine analogue x-adenine. **2013**, 62, 107102 1
- 706 Ablation of Clusters from Surfaces for Harmonic Generation of Laser Radiation. **2014**, 181-221
- 705 Quantum Chemistry. **2014**, 1-33 1
- 704 Real Time Method. **2014**, 35-39
- 703 Effective Usage of the Maximum Entropy Method in Optical Spectrum Analysis of Real-Time TDDFT. **2014**, 13, 314-316 1
- 702 Many-Body Systems and Quantum Hydrodynamics. **2014**, 271-303
- 701 Improved Technique of Optical Spectrum Analysis of Real-Time TDDFT Using Maximum Entropy Method. **2014**, 13, 190-192 2
- 700 Stopping of Slow Ions. **2014**, 343-415 1
- 699 Kohn-Sham Method. **2014**, 79-99
- 698 Unidirectional Photo-induced Charge Separation and Thermal Charge Recombination of Cofacially Aligned Donor-Acceptor System Probed by Ultrafast Visible-Pump/Mid-IR-Probe Spectroscopy. **2014**, 35, 587-596 1
- 697 Patterns of Field Electron Emission from Carbon Nanotubes: Ab Initio Simulations by Time-Dependent Density Functional Theory. **2014**, 0
- 696 Manifestations of the Quantum Confinement Effect in the Phototransport Properties of Ensembles of Semiconductor Quantum Dots. **2014**, 335-408 1
- 695 Time-dependent density functional theory. **1986**, 51-57
- 694 Current Problems in Density Functional Theory. **1987**, 339-357
- 693 Generalized Nonlinear Schrödinger Equations in Quantum Fluid Dynamics. **1988**, 359-365
- 692 Aspects of Density Functional Approach to Many-Electron Systems. **1989**, 541-557
- 691 Formalism. **1990**, 9-35
- 690 Quantum Fluid Dynamical Approach to Chemical Dynamics. **1991**, 185-200

689 Contemporary Topics in Band Theory. **1992**, 1-45

688 Charge and Current Conserving Mesoscopic Transport. **1995**, 657-672

2

687 Ab initio Studies of Electronic Excitations in Real Solids. **1999**, 329-359

686 Towards a Time-Dependent Density-Functional Description of Multiple Multiphoton-Ionization of Helium in Strong Laser Fields. **1999**, 142-157

685 Quantum-Mechanical Treatment of Responses to Electric Fields in Molecular Systems. **2015**, 9-28

684 Quasiparticle Electronic Structures. **2015**, 351-393

1

683 Encyclopedia of Nanotechnology. **2015**, 1-10

682 Time-Domain Ab Initio Modeling of Charge and Exciton Dynamics in Nanomaterials. **2015**, 353-392

681 Modeling of Quasi-One-Dimensional Carbon Nanostructures with Density Functional Theory. **2015**, 1-41

680 Optical properties of Cu doped TiO₂ using First Principle calculations. **2015**,

679 Progress of surface plasmon research based on time-dependent density functional theory. **2015**, 64, 077303

678 Optical Spectral Analysis of Real-Time TDDFT Using Maximum Entropy Method. **2015**, 14, 71-73

1

677 Time-Dependent Density Functional Theory: A Tool to Explore Excited States. **2015**, 1-35

676 The investigation of the photophysical properties of 6-chlorocurcumin and 6-methylcurcumin. **2015**, 6, 279-286

0

675 Combined approach to uv-vis study of 2-allyl and 2-ethylthioquinolines in various solutions. **2016**, 8, 34-40

674 Encyclopedia of Nanotechnology. **2016**, 3798-3806

673 Quantal Density Functional Theory. **2016**, 67-133

672 Quantum Mechanics for Quantum Chemistry. **2016**, 357-549

- 671 - Electronic Excitation Energies of Molecular Systems from the Bethe-Salpeter Equation: Example of the H₂ Molecule. **2016**, 404-427
- 670 Density Functional Theory (DFT): Periodic Advancement and New Challenges. **2016**, 219-230
- 669 Quantum Chemical Computational Studies on a vic-Dioxime Ligand and Its Nickel Complex. 1-1
- 668 Theoretical Study of N-Methyl-3-Phenyl-3-(4-(Trifluoromethyl) Phenoxy) Propan as a Drug and Its Five Derivatives. **2018**, 06, 80-98 1
- 667 Theoretical Methods of Surface Dynamics. **2018**, 117-142
- 666 Real-time time dependent density functional theory with numerical atomic orbital basis set: methodology and applications. **2018**, 67, 120201 2
- 665 Quantum chemical study of electron structure and charge transport properties of symmetric acenequinones. **2018**, 11, 83-93 0
- 664 Selective colorimetric molecular probe for cyanide ion detection in aqueous solution. **2018**, 9, 338-346 1
- 663 On local aromaticity of selected model aza-[n]circulenes (n = 6, 7, 8 and 9): Density functional theoretical study. **2019**, 12, 70-81 0
- 662 Structural, Spectroscopic (FT-IR, Raman, NMR and UV-Vis.) and Computational Studies on N-phenylpropanamide. 823-834
- 661 Molecular Structural and Vibrational Spectroscopic Assignments of n5-(4-Methoxyphenyl)-3-(1-methylindol-3-yl)-isoxazole using DFT Theory Calculations. **2019**, 4, 147-151
- 660 Parallel transport time-dependent density functional theory calculations with hybrid functional on summit. **2019**, 3
- 659 Many-Body Spin Excitations in Ferromagnets from First Principles. **2020**, 1-39 0
- 658 Effect of zirconium doping on the properties of defect titanium dioxide films: quantum chemical calculations. **2019**, 11(26), 372-381
- 657 Solar elements based on organic and organo-inorganic materials. **2019**, 11(26), 270-343
- 656 Accurate determination of quasi-particle electronic and optical spectra of anatase titanium dioxide. **2020**,
- 655 Predicting band gaps of semiconductors with quantum chemistry. **2020**, 246, 00006
- 654 Mechanisms of Tryptophan Transformations Involving the Photochemical Formation of Silver Nanoparticles. **2020**, 237-244 1

- 653 Exact nonadiabatic part of the Kohn-Sham potential and its fluidic approximation. **2020**, 4, 1
- 652 Electronic Excitation Energies of Molecular Systems from the Bethe-Salpeter Equation: Example of the H₂ Molecule. **2020**, 402-425
- 651 Indium complex with task-specific ionic liquid ligands: Ligand to ligand charge transfer in the excited state investigation and reliable DFT predictions. **2020**, 225, 117391 2
- 650 Influence of the electron spill-out and nonlocality on gap plasmons in the limit of vanishing gaps. **2021**, 104, 0
- 649 Improved singlet oxygen generation in Rhenium(I) complexes functionalized with a pyridinyl selenoether ligand. **2021**, 211, 115548
- 648 Non-covalent interactions in hexanuclear polyoxidometalates [VIV₆B₂₀O₅₀H₈]⁸⁻: An experimental and theoretical approach. **2021**, 211, 115553 0
- 647 Scrutinizing GW-Based Methods Using the Hubbard Dimer. **2021**, 9, 751054 6
- 646 Directional dependency of electronic stopping in nickel, projectile-excited charge state and momentum transfer. **2021**, 75, 1 3
- 645 Laser-Induced Electronic and Vibronic Dynamics in the Pyrene Molecule and Its Cation. **2021**, 125, 9619-9631 2
- 644 Breakdown of the ionization potential theorem of density functional theory in mesoscopic systems. **2021**, 155, 194105 1
- 643 Prototyping Ultrafast Charge Separation by Means of Time-Dependent Density Functional Methods. **2020**, 325-343
- 642 Electrically Controlling the Kondo Effect. 13,
- 641 Multiscale simulation of terahertz radiation process in benzimidazole crystal by impulsive stimulated Raman scattering. **2020**, 153, 244506 1
- 640 Assessing the Performance of DFT Functionals for Excited-State Properties of Pyridine-Thiophene Oligomers. **2021**, 125, 115-125 5
- 639 Assessment of long-range corrected density functional theory on the absorption and vibrationally resolved fluorescence spectrum of carbon nanobelts. **2021**, 42, 505-515 0
- 638 n-Tolilbenin (n= o, m, p) Yapısal, Spektroskopik ve Elektronik Özellikleri Üzerine DFT Analizleri. 479-492
- 637 An Efficient Multi-GPU Implementation for Linear-Response Time-Dependent Density Functional Theory. **2020**, 0
- 636 Properties of Conjugated Materials from Quantum Chemistry Coupled to Molecular Dynamics Generated Ensembles. **2020**, 124, 10667-10677 0

635	Molecular Structure-Optical Property Relationship of Salicylidene Derivatives: A Study on the First-Order Hyperpolarizability. 2021 , 125, 99-105	1
634	Highly red-emissive salenIridium complexes: impact of 4-amino-substitution on the photophysical properties.	0
633	A functional model for quercetin 2,4-dioxygenase: Geometric and electronic structures and reactivity of a nickel(II) flavonolate complex. 2022 , 226, 111632	2
632	Synthesis and characterization of metal complex amino acid using spectroscopic methods and theoretical calculation. 2022 , 1250, 131670	3
631	Schiff base-type copper(I) complexes exhibiting high molar extinction coefficients: Synthesis, characterization and DFT studies. 2022 , 1249, 131638	2
630	Recent Developments in Density Functional Approximations. 2020 , 213-226	2
629	Challenges and Opportunities in Modeling Oxides for Energy and Information Devices. 2020 , 1001-1012	2
628	Ground State Structural, Elastic, Electronic Properties and Pressure-Induced Structural Phase Transition of XCoSb (X = Sc, Ti, V, Cr and Mn). 2020 , 33, 1821-1829	0
627	Fundamentals of the Analysis Tools. 2020 , 101-153	
626	Understanding Novel Superconductors with Ab Initio Calculations. 2020 , 73-112	
625	Optimal Control Theory for Electronic Structure Methods. 2020 , 469-489	
624	Optical Properties of Metallic Nanoalloys: From Clusters to Nanoparticles. 2020 , 151-192	1
623	Theoretical Methods. 2020 , 31-39	
622	Singlet fission relevant energetics from optimally tuned range-separated hybrids. 2020 , 22, 27060-27076	2
621	Structural, Spectroscopic (FT-IR, Raman, NMR and UV-Vis.) and Theoretical Investigations of Cyclopentolate.	
620	All-electron real-time and imaginary-time time-dependent density functional theory within a numeric atom-centered basis function framework. 2021 , 155, 154801	3
619	Recent advances in ensemble density functional theory and linear response theory for strong correlation.	2
618	Classical electronic and molecular dynamics simulation for optical response of metal system. 2021 , 155, 174118	1

- 617 Second-order perturbative correlation energy functional in the ensemble density-functional theory. **2021**, 104, 1
- 616 Inq, a Modern GPU-Accelerated Computational Framework for (Time-Dependent) Density Functional Theory. **2021**, 1
- 615 Density Functional Theory Half-Electron Self-Energy Correction for Fast and Accurate Nonadiabatic Molecular Dynamics. **2021**, 12, 10886-10892 5
- 614 Time-dependent approaches for high-harmonic generation spectroscopy. **2021**, 34, 2
- 613 Accelerating molecular property calculations with semiempirical preconditioning. **2021**, 155, 204111 2
- 612 Deformation potential extraction and computationally efficient mobility calculations in silicon from first principles. **2021**, 104, 2
- 611 Quantum Nano-Plasmonics. **2020**, 1
- 610 Theoretical Models. **2004**, 29-70
- 609 First-Principles Simulation on Femtosecond Dynamics in Condensed Matters Within TDDFT-MD Approach. **2006**, 63-75
- 608 Charge localization induced by reorientation of FA cations greatly suppresses nonradiative electron-hole recombination in FAPbI₃ perovskites: A time-domain Ab Initio study. **2020**, 33, 642-648
- 607 INVESTIGATING THE EFFECTS OF THE EXTERNAL ELECTRIC FIELD ON OSMABENZENE IN THE GROUND (S₀) AND FIRST EXCITED SINGLET (S₁) STATES: INSIGHT INTO STRUCTURES, ENERGY, AND PROPERTIES. **2020**, 61, 1691-1699 0
- 606 Vibrational (FT-IR, FT Raman), electronic and docking studies and wave function analysis with quantum chemical computation on 3-Bromophenyl acetic acid: A potential amidase inhibitor. **2022**, 50, 2853-2864
- 605 Aggregation-induced emission spectra of triphenylamine salicylaldehyde derivatives excited-state intramolecular proton transfer revealed by molecular spectral and dynamics simulations.. **2021**, 11, 37171-37180¹
- 604 Three decades of unveiling the complex chemistry of C-nitroso species with computational chemistry. 1
- 603 Gate induced modulation of electronic states in monolayer organic field-effect transistor. **2021**, 119, 223301
- 602 Ensemble Density Functional Theory of Neutral and Charged Excitations : Exact Formulations, Standard Approximations, and Open Questions. **2021**, 380, 4 1
- 601 Customized Luminescent Multiplexed Quick-Response Codes as Reliable Temperature Mobile Optical Sensors for eHealth and Internet of Things. 2100206 7
- 600 Symmetry Dilemma of Doubly Hybrid Density Functionals for Equilibrium Molecular Property Calculations. **2021**, 0


- 599 Parallel transport dynamics for mixed quantum states with applications to time-dependent density functional theory. **2021**, 451, 110850 1
- 598 Synthesis, spectroscopic (FT-IR, FT-Raman, NMR & UV-Vis), reactive (ELF, LOL, Fukui), drug likeness and molecular docking insights on novel 4-[3-(3-methoxy-phenyl)-3-oxo-propenyl]-benzonitrile by experimental and computational methods. **2021**, 7, e08429 2
- 597 Robust broadband directional plasmons in a MoOCl₂ monolayer. **2021**, 104, 1
- 596 Numerical integration of quantum time evolution in a curved manifold. **2021**, 3, 0
- 595 Luminescent Complexes of Europium (III) with 2-(Phenylethynyl)-1,10-phenanthroline: The Role of the Counterions. **2021**, 26, 0
- 594 Zwitterionic Aromaticity on Azulene Extrapolated to carbo-Azulene. **2021**, 2021, 6450 1
- 593 Application of TD-DFT Theory to Studying Porphyrinoid-Based Photosensitizers for Photodynamic Therapy: A Review. **2021**, 26, 1
- 592 Tunable Cr Molecular Color Centers. **2021**, 10
- 591 Modeling dark- and light-induced crystal structures and single-crystal optical absorption spectra of ruthenium-based complexes that undergo SO-linkage photoisomerization.. **2021**, 155, 234111
- 590 YO:Eu and the Mössbauer isomer shift coefficient of Eu compounds from simulations. **2021**, 34, 1
- 589 Corroboration and proposal of isomers for M@C₂(9)-C₈₂(Adamantylidene) (M = La, Y, Sc) endohedral metallofullerenes. **2021**, 1207, 113530
- 588 Simulation of Low-Lying Singlet and Triplet Excited States of Multiple-Resonance-Type Thermally Activated Delayed Fluorescence Emitters by Delta Self-Consistent Field (SCF) Method. **2021**, 125, 10373-10378¹
- 587 Two-step Brillouin zone sampling for efficient computation of electron dynamics in solids. **2021**, 0
- 586 A dynamical study of the principle of maximum hardness. **1994**, 106, 229-249 3
- 585 Mechanistic photophysics and photochemistry of unnatural bases and sunscreen molecules: insights from electronic structure calculations. **2021**, 1
- 584 Computational Methods in Organometallic Chemistry. **2021**, 1
- 583 Probing the nature of donor-acceptor effects in conjugated materials: a joint experimental and computational study of model conjugated oligomers. **2021**, 23, 26534-26546 0
- 582 Ab Initio Computational Approach for Nanophotonics Based on Time-Dependent Density Functional Theory. **2021**, 103-133

- 581 Lactic acid photochemistry following excitation of S0 to S1 at 220 to 250 nm. e4316 0
- 580 Collision site effect on the radiation dynamics of cytosine induced by proton. 1
- 579 Large-scale ab initio simulation of light-matter interaction at the atomic scale in Fugaku. 109434202110657 0
- 578 Introduction to theory of high-harmonic generation in solids: tutorial. 3
- 577 A tale of two vectors: A Lanczos algorithm for calculating RPA mean excitation energies.. **2022**, 156, 014102
- 576 Effect of protein matrix on CP29 spectra and energy transfer pathways.. **2021**, 1863, 148521 1
- 575 First-principle investigation of boron nitride nanobelt. **2022**, 1208, 113571
- 574 The role of π -donors/acceptors in molecular rotors towards development of ambient blue light sensors - A density functional theory study. **2022**, 277, 125563
- 573 Orange-red emissive Cu(I) complexes bearing Schiff base ligands: Synthesis, structures, and photophysical properties. **2022**, 1252, 132180 0
- 572 DFT analysis of different substitutions on optoelectronic properties of carbazole-based small acceptor materials for Organic Photovoltaics. **2022**, 140, 106381 1
- 571 A study of DNA/BSA interaction and catalytic potential of oxidovanadium(V, IV) complexes incorporating dibenzofuran based $O^{\wedge}N^{\wedge}O$ ligand. **2022**, 961, 122244 0
- 570 Nuclear Norm Based Spectrum Estimation for Molecular Dynamic Simulations. **2020**,
- 569 Molecular-Level Understanding of Dual-RTP via Host-Sensitized Multiple Triplet-to-Triplet Energy Transfers and Data Security Application.. **2022**, 7, 3722-3730 0
- 568 Search for long-lasting electronic coherence using on-the-fly ab initio semiclassical dynamics.. **2022**, 156, 034104 2
- 567 Ab-Initio Prediction of a Negative Barkas Coefficient for Slow Protons and Antiprotons in LiF.. *Physical Review Letters*, **2022**, 128, 043401 7.4 1
- 566 Heisenberg Spin Hamiltonian Derived from a Multiple Grand Canonical Spin Density Functional Theory with a Principal Nonlocal Exchange-Correlation Energy Functional. **2022**, 91,
- 565 The surprising effects of sulfur: achieving long excited-state lifetimes in heteroleptic copper(i) emitters.. **2022**, 10, 3089-3102 2
- 564 Exploring Intra- and Intermolecular Interactions in Selected -Oxides-The Role of Hydrogen Bonds.. **2022**, 27, 1

563	Multireference Density Functional Theory for Describing Ground and Excited States with Renormalized Singles.. 2022 , 894-903		1
562	Correlating Interfacial Charge Transfer Rates with Interfacial Molecular Structure in the Tetraphenyldibenzoperiflanthene/C Organic Photovoltaic System.. 2022 , 763-769		0
561	First-principles wave-vector- and frequency-dependent exchange-correlation kernel for jellium at all densities. 2022 , 105,		1
560	Optical Control of Multistage Phase Transition via Phonon Coupling in MoTe ₂ .. <i>Physical Review Letters</i> , 2022 , 128, 015702	7.4	2
559	DFT and TD-DFT study of adsorption behavior of Zejula drug on surface of the B12N12 nanocluster. 2022 , 1-16		0
558	Theoretical Study on the Light-Emitting Mechanism of Multifunctional Thermally Activated Delayed Fluorescence Molecules. 2022 , 126, 2437-2446		1
557	Thermal excitation signals in the inhomogeneous warm dense electron gas.. 2022 , 12, 1093		4
556	Ultrafast optical control over spin and momentum in solids. 2022 , 120, 032403		1
555	Manipulation of the high-order harmonic generation in monolayer hexagonal boron nitride by two-color laser field.. 2022 , 156, 074701		0
554	Ultrafast nonadiabatic dynamics of DNA upon low energy proton irradiation. 2022 , 120, 043702		0
553	Dissymmetric Chiral Poly(diphenylacetylene)s: Secondary Structure Elucidation and Dynamic Luminescence.. 2022 ,		2
552	Photoelectron elliptical dichroism spectroscopy of resonance-enhanced multiphoton ionization the 3s, 3p and 3d Rydberg series in fenchone.. 2022 ,		1
551	Enhancing the photocatalytic hydrogen generation performance and strain regulation of the vertical GeI ₂ /C ₂ N van der Waals heterostructure: insights from first-principles study.		1
550	Excited States Computation of Models of Phenylalanine Protein Chains: TD-DFT and Composite CC2/TD-DFT Protocols.. 2022 , 23,		
549	Real-Time, Time-Dependent Density Functional Theory Study on Photoinduced Isomerizations of Azobenzene Under a Light Field.. 2022 , 427-432		2
548	Manifold curvature and Ehrenfest forces with a moving basis. 2022 , 12,		
547	Accurate Spectral Properties within Double-Hybrid Density Functional Theory: A Spin-Scaled Range-Separated Second-Order Algebraic-Diagrammatic Construction-Based Approach.. 2022 ,		2
546	Design of dyes for energy transformation: From the interaction with biological systems to application in solar cells. 2022 , 79-114		

545	Modelling the physical properties of environmentally friendly optical magnetic switches: DFT and TD-DFT. 2022 , 355-384	2
544	Floquet Time-Dependent Configuration Interaction for Modeling Ultrafast Electron Dynamics.. 2022 ,	1
543	Nonlinear dynamics of electromagnetic field and valley polarization in WSe2 monolayer. 2022 , 120, 051108	0
542	Effects of molecular geometry on the efficiency of intramolecular charge transfer-based luminescence in o-carboranyl-substituted 1H-phenanthro[9,10-d]imidazoles. 2022 , 9, 501-513	0
541	Unravelling the origin of dual photoluminescence in Au2Cu6 clusters by triplet sensitization and photon upconversion.	2
540	Highly-anisotropic plasmons in two-dimensional hyperbolic copper borides.. 2022 , 30, 5596-5607	0
539	Computational chemistry and the study and design of catalysts. 2022 , 299-332	
538	Ab initio study of ultrafast spin dynamics in Gdx(FeCo) 1x alloys. 2022 , 120, 042401	0
537	Theoretical design of SnS-graphene heterojunctions with vacancy and impurity defects for multi-purpose photoelectric devices.. 2021 ,	0
536	Ultrafast transient absorption spectroelectrochemistry: femtosecond to nanosecond excited-state relaxation dynamics of the individual components of an anthraquinone redox couple.. 2022 , 13, 486-496	0
535	DFT calculation in design of near-infrared absorbing nitrogen-doped graphene quantum dots.. 2021 ,	1
534	Dissymmetric Chiral Poly(diphenylacetylene)s: Secondary Structure Elucidation and Dynamic Luminescence.	
533	Absorption Spectra and the Electronic Structure of Gallic Acid in Water at Different pH: Experimental Data and Theoretical Cluster Models.. 2022 , 126, 190-197	1
532	Excited State Dynamics in Dual-Defects Modified Graphitic Carbon Nitride.. 2022 , 1033-1041	5
531	First-principles modelling for time-resolved ARPES under different pump-probe conditions. 2022 , 254, 147152	1
530	Generalized Oscillator Strengths for the Valence Shell Excitations in Carbon Tetrachloride Studied by Fast Electron Impact.. 2022 ,	1
529	Time-dependent exchange-correlation potential in lieu of self-energy. 2022 , 105,	0
528	Effect of W on the thermal stability, mechanical properties and corrosion resistance of Fe-based bulk metallic glass. 2022 , 143, 107485	4

527	Synthesis, spectroscopic investigation and quantum chemical computation of 2-(2-arylamino-4-aminothiazol-5-yl) naphthalene derivatives. 2022 , 583, 126553	
526	A bowl-shaped phenoxido-bridged binuclear zinc complex: Experimental and theoretical studies. 2022 , 534, 120807	0
525	Stabilization and fluorescence light-up of G-quadruplex nucleic acids using indolyl-quinolinium based probes.. 2022 ,	2
524	Functionalized Crystalline -Trimethyltriindoles: Counterintuitive Influence of Peripheral Substituents on Their Semiconducting Properties.. 2022 , 27,	
523	First-principles calculations for transient absorption of laser-excited magnetic materials.	
522	Structural and theoretical studies on copper(II) and zinc(II) complexes with a 9-anthraldehyde-derived thiosemicarbazone. 2022 , 115724	1
521	Synthesis, Spectroscopic Characterization, Electronic and Docking Studies on Novel Chalcone Derivatives (3DPP and 5PPD) by Experimental and DFT Methods. 2022 , 132553	0
520	Computational analysis of transient XMCD sum rules for laser pumped systems: When do they fail?. 2022 , 120, 062409	1
519	Herzberg-Teller Effect in Single-Crystalline Hexacene at Finite Temperatures.	0
518	Do any types of double-hybrid models render the correct order of excited state energies in inverted singlet-triplet emitters?. 2022 , 156, 064302	1
517	New Cd(II) complex derived from (1-methylimidazol-2-yl)methanol: synthesis, crystal structure, spectroscopic study, DFT and TD-DFT calculations, antimicrobial activity and free-radical scavenging capacity. 2022 , 1257, 132583	0
516	A Concise Review on Recent Developments of Machine Learning for the Prediction of Vibrational Spectra.. 2022 ,	1
515	Absorption and Photoluminescence of Silicon Nanocrystals Investigated by Excited State DFT: Role of Embedding Dielectric and Defects.	1
514	Origin of the Failure of Density Functional Theories in Predicting Inverted Singlet-Triplet Gaps.. 2022 ,	5
513	Imprints of multielectron polarization effects in odd-even harmonic generation from CO molecules. 2022 , 105,	0
512	Spirovetivane- and Eudesmane-Type Sesquiterpenoids Isolated from the Culture Media of Two Cyanobacterial Strains.. 2022 ,	
511	Tetraphenyl Tetrel Molecules and Molecular Crystals: From Structural Properties to Nonlinear Optics.	0
510	Stationary and Time-Dependent Carbon Monoxide Stretching Mode Features in Carboxy Myoglobin: A Theoretical-Computational Reappraisal.. 2021 , 125, 13624-13634	2

- 509 Probing the Energy Conversion Pathways between Light, Carriers, and Lattice in Real Time with Attosecond Core-Level Spectroscopy. **2021**, 11, 1
- 508 Theoretical study on an intriguing excited-state proton transfer process induced by weakened intramolecular hydrogen bonds.. **2022**, 0
- 507 Excited-state properties of organic semiconductor dyes as electrically pumped lasing candidates from new optimally tuned range-separated models.. **2022**, 1
- 506 Luminescence of Fullerene-like Clusters by a first-principles study. **2022**,
- 505 Cyclo[18]carbon including zero-point motion: ground state, first singlet and triplet excitations, and hole transfer.. **2022**, 1
- 504 Plasmon-Induced Hot Electrons in Metallic Nanoparticles. **2022**, 155-175
- 503 Aerobic photolysis of methylcobalamin: unraveling the photoreaction mechanism.. **2022**, 0
- 502 Quantum simulations of thermally activated delayed fluorescence in an all-organic emitter.. **2022**, 0
- 501 Correlated organic-inorganic motion enhances stability and charge carrier lifetime in mixed halide perovskites.. **2022**, 6
- 500 Charge-Transfer Excitations within Density Functional Theory: How Accurate Are the Most Recommended Approaches?. **2022**, 1
- 499 Charge-Transfer Excitation within a Hybrid-(G)KS Framework through Cartesian Grid DFT.. **2022**, 126, 1448-1457 1
- 498 Energy Landscape of State-Specific Electronic Structure Theory.. **2022**, 3
- 497 Molecular Structural, Hydrogen Bonding Interactions, and Chemical Reactivity Studies of Ezetimibe-L-Proline Cocrystal Using Spectroscopic and Quantum Chemical Approach.. **2022**, 10, 848014 0
- 496  **2022**, 56, 179
- 495 Computational characterization of nanosystems. **2022**, 35, 1-15
- 494 Towards a quantum fluid theory of correlated many-fermion systems from first principles. **2022**, 12, 5
- 493 Signatures of atomic structure in subfemtosecond laser-driven electron dynamics in nanogaps. **2022**, 105, 1
- 492 Molecular Structure, Experimental and Theoretical Vibrational Spectroscopy, (HOMO-LUMO, NBO) Investigation, (RDG, AIM) Analysis, (MEP, NLO) Study and Molecular Docking of Ethyl-2-{{[4-Ethyl-5-(Quinolin-8-yloxyMethyl)-4H-1,2,4-Triazol-3-yl] Sulfanyl} Acetate. 1-25 3

- 491 Optical Properties of OLED Materials by TDDFT. **2022**, 2207, 012039
- 490 Power functional theory for many-body dynamics. **2022**, 94, 5
- 489 Comparison of semi-empirical and density functional approaches for the colour and constitution of anthraquinone dyes using X-ray structure.
- 488 Excited State Dynamics of Methylated Guanosine Derivatives Revealed by Femtosecond Time-resolved Spectroscopy.. **2022**,
- 487 Unified Time Evolution Approach for the Electronic Structure Calculation. **2022**, 2207, 012037
- 486 Application of chromium-silicon cluster for selective removal of arsenic and sulfide from wastewater.
- 485 Element-selective ultrafast magnetization dynamics of hybrid Stoner-Heisenberg magnets. **2022**, 105, 0
- 484 Thermally Activated Delayed Fluorescence and Room-Temperature Phosphorescence in Asymmetric Phenoxazine-Quinoline (D2A) Conjugates and Dual Electroluminescence. **2022**, 126, 5649-5657 2
- 483 Laser-Controlled Charge Transfer in a Two-Dimensional Organic/Inorganic Optical Coherent Nanojunction. 1
- 482 Single Excitation Energies Obtained from the Ensemble "HOMO-LUMO Gap": Exact Results and Approximations.. **2022**, 2452-2458 0
- 481 Working equation of linear response time-dependent density functional theory: First-order polarization propagator approximation. 0
- 480 Theoretical Modeling of Absorption and Fluorescent Characteristics of Cyanine Dyes. **2022**, 2, 202-216 0
- 479 Conversion of twisted light to twisted excitons using carbon nanotubes. **2022**, 8,
- 478 Optical spectrum of OLED materials by time-dependent density functional theory. **2022**, 7, 310
- 477 Influence of Linker Orientation and Regulative Factor(s) in Liposomal Gene Delivery: A Molecular Level Investigation.. **2022**, 126, 1816-1822
- 476 Estimating Phosphorescent Emission Energies in Ir Complexes Using Large-Scale Quantum Computing Simulations.. **2022**, e202116175 1
- 475 Relativistic Effects in Modeling the Ligand K-Edge X-ray Absorption Near-Edge Structure of Uranium Complexes.. **2022**, 0
- 474 Continuous Kubo-Greenwood formula: Theory and numerical implementation.. **2022**, 105, 035307

- 473 Conservation of angular momentum in ultrafast spin dynamics. **2022**, 105,
- 472 Mono Versus Dinuclear Vanadium(V) Complexes: Solvent Dependent Structural Versatility and Electro Syntheses of Mixed-Valence Oxovanadium(IV/V) Entities in Solution.. **2022**, 7, 11710-11721
- 471 Charge carrier nonadiabatic dynamics in non-metal doped graphitic carbon nitride.. **2022**, 156, 094702 4
- 470 Generalized Ab Initio Nonadiabatic Dynamics Simulation Methods from Molecular to Extended Systems.. **2022**, 1
- 469 Estimating Phosphorescent Emission Energies in Ir III Complexes Using Large-Scale Quantum Computing Simulations**. 0
- 468 Combining localized orbital scaling correction and Bethe-Salpeter equation for accurate excitation energies.. **2022**, 156, 154101 1
- 467 Prospects of quantum computing for molecular sciences. **2022**, 6, 4
- 466 Excited Electron and Spin Dynamics in Topological Insulator: A Perspective from Ab Initio Non-adiabatic Molecular Dynamics. **2022**,
- 465 Theoretical insights into the effects of RE doping on the structural, electronic, and optical properties of magnesium clusters. **2022**, 12, 035348 0
- 464 Experimental and Quantum Chemical Studies of Nicotinamide-Oxalic Acid Salt: Hydrogen Bonding, AIM and NBO Analysis.. **2022**, 10, 855132 1
- 463 Complete solution to the inverse Kohn-Sham problem: From the density to the energy. **2022**, 105,
- 462 Ultrastrong Coupling of a Single Molecule to a Plasmonic Nanocavity: A First-Principles Study.. **2022**, 9, 1065-1077 1
- 461 Orbital-resolved visualization of single-molecule photocurrent channels.. **2022**, 603, 829-834 6
- 460 Numerical scheme for a nonlinear optical response of a metallic nanostructure: quantum hydrodynamic theory solved by adopting an effective Schrödinger equation.. **2022**, 30, 11572-11587 0
- 459 Improving Lattice Rigidity and Charge Carrier Lifetime by Engineering Spacer Cation of Ruddlesden-Popper Perovskites: A Time-Domain Study.. **2022**, 2718-2724 6
- 458 Spectroscopic and topological analysis and in vitro antimicrobial activity of phenothiazine. **2022**, 55, 212-228 0
- 457 Interplay of Steric Effects and Aromaticity Reversals to Expand the Structural/Electronic Responses of Dihydrophenazines.. **2022**, 0
- 456 Synthesis, photochemistry and computational study of novel 1,2,3-triazole heterostilbenes: Expressed biological activity of their electrocyclization photoproducts.. **2022**, 121, 105701 2

- 455 Sensitivity of time-dependent density functional theory to initial conditions. **2022**, 105,
- 454 Polarization-resolved Analysis to Solid High-order Harmonic Generation. 0
- 453 Deep dive into machine learning density functional theory for materials science and chemistry. **2022**, 6, 5
- 452 n2v : A density-to-potential inversion suite. A sandbox for creating, testing, and benchmarking density functional theory inversion methods. 2
- 451 Deciphering the Spectral Tuning Mechanism in Proteorhodopsin: The Dominant Role of Electrostatics Instead of Chromophore Geometry.. **2022**, 0
- 450 First principles approach to solar energy conversion efficiency of semiconductor heterojunctions. **2022**, 236, 445-454 0
- 449 Finite-size and quantum effects in plasmonics: manifestations and theoretical modelling [Invited]. **2022**, 12, 1869 2
- 448 Photochromism of a Spiropyran in Low-Temperature Matrices: Unprecedented Bidirectional Switching between a Merocyanine and an Allene Intermediate.. **2022**, 1
- 447 Transient uniform electron gases.
- 446 Heteroleptic Copper(I) complexes of bipyridine glycoluril and phosphine ligands: photophysical and computational studies. **2022**, 120934 0
- 445 Restricting Bond Rotations by Ring Fusion: A Novel Molecular Design Strategy to Improve Photodynamic Antibacterial Efficacy of AIE Photosensitizers.. **2022**, 2
- 444 Improving Results by Improving Densities: Density-Corrected Density Functional Theory.. **2022**, 7
- 443 Structural, electronic properties (different solvents), chemical reactivity, ELF, LOL, spectroscopic insights, molecular docking and in vitro anticancer activity studies on methyl (4-nitro-1-imidazolyl)acetate. **2022**, 99, 100438 0
- 442 Insight into the spatial interaction of D-BA bridge derived cyanines and nitroreductase for fluorescent cancer hypoxia detection.. **2022**, 273, 121031
- 441 Triple proton transfer after water rearrangement in (2,6-aza)Ind[7](H₂O)₂. **2022**, 353, 118847 0
- 440 Computational hints for the simultaneous spectroscopic detection of common contaminants in water. **2022**, 355, 118908 1
- 439 A real-time TDDFT scheme for strong-field interaction in Cartesian coordinate grid. **2022**, 796, 139562 0
- 438 Synthesis, structures, and photophysical properties of three-coordinate copper(I) complexes bearing bidentate bis[(2-diphenylphosphino)phenyl]ether (POP) ligand and monodentate substituted-quinoline ligand. **2022**, 1257, 132642 0

437	Synthesis of novel antipyrine-azole-S-alkyl derivatives antimicrobial activity, molecular docking, and computational studies. 2022 , 1260, 132810	2
436	New silver(I) complex as antibiotic candidate: Synthesis, spectral characterization, DFT, QTAIM and antibacterial investigations and docking properties. 2022 , 1261, 132902	1
435	Crystal structure, spectroscopic studies, DFT calculations, and biological activity of 5-bromosalicylaldehydeBased Schiff bases. 2022 , 1262, 132976	0
434	Variational functionals for excited state saddle points versus traditional Hylleraas and McDonald higher roots, and a way to instantly improve a lowest state crude approximant. 2022 , 122,	
433	Quantum-chemical approaches in the study of fullerene and its derivatives by the example of the most typical cycloaddition reactions: A review. 2022 , 122,	1
432	Charge transfer excitations and constrained density functional theory. 2022 , 141, 1	0
431	The Impact of Retinal Configuration on the Protein-Chromophore Interactions in Bistable Jumping Spider Rhodopsin-1.. 2021 , 27,	1
430	Plasmon excitation in $\text{MoS}_2/\text{graphene}$ van der waals heterostructures. 2022 , 96, 1	
429	Electronic Structure of (Organic-)Inorganic Metal Halide Perovskites: The Dilemma of Choosing the Right Functional. 2022 , 5, 2100496	2
428	Double and Charge-Transfer Excitations in Time-Dependent Density Functional Theory.. 2021 ,	7
427	CO Adsorbate Promotes Polaron Photoactivity on the Reduced Rutile TiO(110) Surface.. 2022 , 2, 234-245	6
426	Chiral 1D MetalOrganic Materials Based on Cu(II) and Amino Acid Schiff Bases. 2022 , 22, 237-250	0
425	Deep Well Trapping of Hot Carriers in a Hexagonal Boron Nitride Coating of Polymer Dielectrics. 2021 ,	0
424	First-principles study of excitons in optical spectra of silver chloride. 2021 , 104,	0
423	Data-driven modeling of S -> S excitation energy in the BODIPY chemical space: High-throughput computation, quantum machine learning, and inverse design.. 2021 , 155, 244102	2
422	Accurate X-ray Absorption Spectra near L- and M-Edges from Relativistic Four-Component Damped Response Time-Dependent Density Functional Theory.. 2021 ,	4
421	Calculating Photoabsorption Cross-Sections for Atmospheric Volatile Organic Compounds.. 2022 , 6, 207-217	3
420	Charge-Compensated Doping Extends Carrier Lifetimes in SrTiO by Passivating Oxygen Vacancy Defects.. 2021 , 12, 12040-12047	3

419	Benchmarking Density Functional Approximations for Excited-State Properties of Fluorescent Dyes.. 2021 , 26,	2
418	An energetics assessment of benzo[a]tetracene and benzo[a]pyrene as triplet-triplet annihilation emitters.	0
417	Thermodynamic equilibrium between locally excited and charge-transfer states through thermally activated charge transfer in 1-(pyren-2?-yl)-o-carborane.	4
416	Electron dynamics in B-quartz induced by two-color 10-femtosecond laser pulses. 2022 , 105,	0
415	Relativistic Orbital-Optimized Density Functional Theory for Accurate Core-Level Spectroscopy.. 2022 , 3438-3449	5
414	Ab initio study of ultrafast demagnetization of elementary ferromagnets by terahertz versus optical pulses. 2022 , 105,	0
413	Shortcut to Self-Consistent Light-Matter Interaction and Realistic Spectra from First Principles.. <i>Physical Review Letters</i> , 2022 , 128, 156402	7.4 3
412	Probing the role of grain boundaries in single Cu nanoparticle oxidation by in situ plasmonic scattering. 2022 , 6,	0
411	Kinetic studies of exchange-correlation effect on the collective excitations of warm dense plasmas.. 2022 , 105, 045206	0
410	Superalkali X3O (X = Li, Na, K) doped B12N12 nano-cages as a new drug delivery platform for chlormethine: A DFT approach. 2022 , 113722	0
409	Review on the QM/MM Methodologies and Their Application to Metalloproteins.. 2022 , 27,	1
408	Effects of Donor Position and Multiple Charge Transfer Pathways in Asymmetric Pyridyl-Sulfonyl TADF Emitters. 2022 , 103550	1
407	The Linear Response Function. 2022 , 301-323	
406	A Combined Experimental and Theoretical Studies of two new Decavanadate: (C6N2H9)4[H2V10O28]·4H2O and (C7H9NF)4[H2V10O28]·4H2O. 2022 , 133085	
405	Synthesis, structures, and photophysical properties of orange-red emissive Cu(I) complexes of 9,9-dimethyl-4,5-bis(diphenylphosphino)-9H-xanthene (Xantphos) with the substituted N-heterocycle carbene (NHC) ligand. 2022 , 122356	
404	Effects of electronic correlation on the high harmonic generation in helium: A time-dependent configuration interaction singles vs time-dependent full configuration interaction study.. 2022 , 156, 174106	1
403	Linear and Nonlinear Optical Properties from TDOMP2 Theory.. 2022 ,	1
402	Table_1.DOCX. 2020 ,	

401 Data_Sheet_1.PDF. **2020**,

400 Automated Generation of Optimized Auxiliary Basis Sets for Long-Range-Corrected TDDFT Using the Cholesky Decomposition.. **2022**,

1

399 Conformational Analysis of Psilocybin: A Computational Approach.

398 Understanding the Photolysis of CH₃ONO₂ with the On-the-fly Nonadiabatic Dynamics Simulation at the ADC(2) Level.

397 From ground to excited electronic state dynamics of electron and ion irradiated graphene nanomaterials. **2022**, 87-107

396 Dissipation and spontaneous emission in quantum electrodynamical density functional theory based on optimized effective potential: A proof of concept study. **2022**, 105,

395 Theoretical Insights into Ultrafast Dynamics in Quantum Materials. **2022**, 2022, 1-16

3

394 Assessment of the Heat Capacity by Thermodynamic Approach Based on Density Functional Theory Calculations.

393 Transition-Based Constrained DFT for the Robust and Reliable Treatment of Excitations in Supramolecular Systems.. **2022**,

392 Accelerating materials discovery using artificial intelligence, high performance computing and robotics. **2022**, 8,

5

391 Electronic stopping power and electronic energy-loss mechanism for a low-energy ion in TiN under channeling conditions. **2022**, 105,

390 Multiple collisions in crystal high-order harmonic generation.

389 Time-dependent extension of the self-consistent band theory for neutron star matter: Anti-entrainment effects in the slab phase. **2022**, 105,

0

388 Variational Density Functional Calculations of Excited States: Conical Intersection and Avoided Crossing in Ethylene Bond Twisting.. **2022**, 3990-3999

2

387 Ultrafast dynamics of adenine following XUV ionization.

0

386 PhenoxazineQuinoline Conjugates: Impact of Halogenation on Charge Transfer Triplet Energy Harvesting via Aggregate Induced Phosphorescence.

0

385 Theoretical study on aggregation-induced emission of new multi-layer 3D chiral molecules. 1-10

384 Revisiting the Performance of Time-Dependent Density Functional Theory for Electronic Excitations: Assessment of 43 Popular and Recently Developed Functionals from Rungs One to Four.. **2022**,

4

383	Quantum materials out of equilibrium. 2022 , 75, 42-47	
382	Computing X-Ray Absorption Spectra from Linear-Response Particles atop Optimized Holes.	2
381	Influence of static correlation on the magnon dynamics of an itinerant ferromagnet with competing exchange interactions: First-principles study of MnBi. 2022 , 6,	0
380	Geometric structural insights for enhanced radiative efficiency: Spiro[fluorene]barbazole]-based ortho -carboranyl luminophores.	
379	Determination of H-Atom Positions in Organic Crystal Structures by NEXAFS Combined with Density Functional Theory: a Study of Two-Component Systems Containing Isonicotinamide.. 2022 ,	2
378	AIE mechanism of 2-(2-hydroxyphenyl) benzothiazole derivatives: CASPT2 and spin-flip study. 2022 , 110396	
377	An asymptotic Green's function method for time-dependent Schrödinger equations with application to Kohn-Sham equations. 2022 , 463, 111272	
376	Novel indolo [3,2-c]isoquinoline-5-one-6-yl [1,2,4]triazolo [3,4-b] [1,3,4]thiadiazole analogues: Design, synthesis, anticancer activity, docking with SARS-CoV-2 Omicron protease and MESP/TD-DFT approaches. 2022 , 1264, 133153	1
375	A new class of deep blue emitting four-coordinate NHC-Cu(I) complexes bearing nonconjugated triazolyl-imidazolylidene-type ligand: Synthesis, photophysical properties and theoretical investigations. 2022 , 431, 114008	0
374	Polarizable Embedding Complex Polarization Propagator in Four- and Two-Component Frameworks.. 2022 ,	2
373	Photoproduct formation in coenzyme B12-dependent CarH via a singlet pathway. 2022 , 112471	0
372	Quantum Chemistry Methods. 2022 , 3-44	
371	Assessing Recent Time-Dependent Double-Hybrid Density Functionals on Doublet-Doublet Excitations.	1
370	Evaluation of Molecular Photophysical and Photochemical Properties Using Linear Response Time-Dependent Density Functional Theory with Classical Embedding: Successes and Challenges.	1
369	Part 2. Vibronic effects accelerate the intersystem crossing processes of the through-space charge transfer states in the triptycene bridged acridine-triazine donor-acceptor molecule TpAT-tFFO.	4
368	Inelastic scattering of electrons in water from first principles: cross sections and inelastic mean free path for use in Monte Carlo track-structure simulations of biological damage. 2022 , 9,	2
367	Nonadiabatic Exchange-Correlation Potential for Strongly Correlated Materials in the Weak and Strong Interaction Limits. 2022 , 10, 77	
366	A many-body approach to transport in quantum systems: From the transient regime to the stationary state.	0

- 365 Mechanism of Ir(ppy)₃ Guest Exciton Formation with the Exciplex-Forming TCTA:TPBI Cohost within a Phosphorescent Organic Light-Emitting Diode Environment. **2022**, 23, 5940 1
- 364 Tribology at the Atomic Scale with Density Functional Theory. 0
- 363 Ultrafast photo-induced processes in complex environments: The role of accuracy in excited-state energy potentials and initial conditions. **2022**, 3, 021307 1
- 362 Atomic and Molecular Physics Methods for Nanosystems. **2022**, 15-93
- 361 Optical properties improvement for the [14]annulene using different donor-acceptor substituents. 1-8
- 360 Few-Femtosecond Dynamics of Free-Free Opacity in Optically Heated Metals. **2022**, 12,
- 359 Modeling Radiation Damage in Materials Relevant for Exploration and Settlement on the Moon.
- 358 Photoionization and core resonances from range-separated density-functional theory: General formalism and example of the beryllium atom. 0
- 357 Coumarinolignans with Reactive Oxygen Species (ROS) and NF- κ B Inhibitory Activities from the Roots of *Waltheria indica*. **2022**, 27, 3270
- 356 Understanding Competitive Photo-Induced Molecular Oxygen Dissociation and Desorption Dynamics atop a Reduced Rutile TiO₂(110) Surface: A Time-Domain Ab Initio Study. **2022**, 12, 6702-6711 3
- 355 Calibrating Out-of-Equilibrium Electron-Phonon Couplings in Photoexcited MoS₂. 1
- 354 Search for a density-based alternative quantum mechanics of many-electron systems. **1984**, 93, 965-976 1
- 353 [Pt₁Ag₃₇(SAdm)₂₁(Dppp)₃Cl₆]₂⁺: intercluster transformation and photochemical properties. 0
- 352 Photo-induced ligand substitution of Cr(CO)₆ in 1-pentanol probed by time resolved X-ray absorption spectroscopy. 0
- 351 Extended quasiparticle approach to non-resonant and resonant X-ray emission spectroscopy.
- 350 UPS, XPS, NEXAFS and Computational Investigation of Acrylamide Monomer. **2022**, 2, 463-478 1
- 349 Local Phonon Environment as a Design Element for Long-Lived Excitonic Coherence: Dithia-anthracenophane Revisited. **2022**, 126, 3765-3773
- 348 Nonequilibrium Charge-Density-Wave Melting in 1T-TaS₂ Triggered by Electronic Excitation: A Real-Time Time-Dependent Density Functional Theory Study. 5711-5718 0

- 347 Real-time first-principles calculations of ultrafast carrier dynamics of SnSe/TiO₂ heterojunction under Li⁺ implantation.
- 346 Electronic transport coefficients from density functional theory across the plasma plane. **2022**, 105, 0
- 345 Construction of anisotropic nanostructures by self-assembly of aggregation-induced emission driven from tris-branched [2]rotaxane based molecular zipper. **2022**, 24, 100997 0
- 344 Rational construction of deep-red fluorescent probe for rapid detection of HClO and its application in bioimaging and paper-based sensing. 0
- 343 Porphyrins: Electronic Structure and Ultraviolet/Visible Absorption Spectroscopy. **2022**, 505-586
- 342 First-principles method for nonlinear light propagation at oblique incidence. **2022**, 30, 23664
- 341 Dissociation mechanism of a C₆₀-Li⁺ complex by microscopic hydration: density functional theory study.
- 340 Depleted Oxygen Defect State Enhancing Tungsten Trioxide Photocatalysis: A Quantum Dynamics Perspective. **2022**, 13, 5571-5580 4
- 339 Light induced ultrafast magnetization dynamics in metallic compounds. **2022**, 169596 1
- 338 Describing Chemical Reactivity with Frontier Molecular Orbitals. 1
- 337 Applications of the dynamical generator coordinate method to quadrupole excitations. **2022**, 105,
- 336 Effects of Damping Constant of Electron and Size on Quantum-Based Frequency-dependent Dielectric Function of Small Metallic Plasmonic Devices. **2022**, 16, e01242 0
- 335 Tautomeric, spectroscopic, electronic and NLO analyses of purpald (4-amino-3-hydrazino-5-mercapto-1,2,4-triazole). **2022**, 32, 103862 2
- 334 Intriguing photophysical and mechanochromic characteristics of carboline-based benzothiadiazole donor-acceptor triad compounds. **2022**, 205, 110515
- 333 Quantal Density Functional Theory: A Local Effective Potential Theory Complement to Schrödinger Theory. **2022**, 161-213
- 332 Introduction. **2022**, 1-16
- 331 Modern Density Functional Theory. **2022**, 215-279
- 330 Unravelling a New Conformer of Psilocin Through Computational Methods.

- 329 Exploration of the photocatalytic cycle for sacrificial hydrogen evolution by conjugated polymers containing heteroatoms. 2
- 328 Local orbital formulation of the Floquet theory of projectile electronic stopping. **2022**, 105, 0
- 327 Following the density evolution using real time density functional theory and density based indexes: Application to model pushpull molecules. **2022**, 43, 1464-1473 0
- 326 Identifying Vibrations that Control Non-adiabatic Relaxation of Polaritons in Strongly Coupled MoleculeCavity Systems. **2022**, 13, 6259-6267 0
- 325 Crucial Factors Regulating Intramolecular Charge-Transfer-Based Radiative Efficiency in ortho-Carboranyl Luminophores: Planarity between Substituted Biphenyl Rings. **2022**, 7, 24027-24039 1
- 324 Theory of ultrafast magnetization of nonmagnetic semiconductors with localized conduction bands. **2022**, 105, 0
- 323 Comparison of (B?B)?M bond and luminescence behavior of group 11/12 metal Ediborene complexes: Theoretical investigation. 0
- 322 Benchmarking time-dependent density functional theory predictions of emission spectra and CIE color: A rainbow of error. 0
- 321 On the Origin of Photoluminescence Enhancement in Biicosahedral Ag x Au 25k Nanoclusters (x 3-003) and Their Application to TripletTriplet Annihilation Photon Upconversion. 2200864 3
- 320 Synergistic effect of atomic layer deposition-assisted cocatalyst and crystal facet engineering in SnS2 nanosheet for solar water oxidation. **2022**, 3
- 319 Reference Energies for Cyclobutadiene: Automerization and Excited States. 2
- 318 Spectroscopic and quantum mechanical study of a scavenger molecule: N,N-diethylhydroxylamine. **2022**, 121555 0
- 317 Efficient and interpretable graph network representation for angle-dependent properties applied to optical spectroscopy. **2022**, 8, 0
- 316 Transition of the generation mechanism of high-order harmonics in an extended neon system. **2022**, 7, 044403 0
- 315 Contribution of F-electron excitation to electronic stopping power of platinum for protons. **2022**, 106, 0
- 314 Synthesis, quantum mechanical calculations, molecular docking, Hirshfeld surface analysis and ADMET estimation studies of (E)-3-(anthracene-10-yl)-1-(naphthalen-1-yl)prop-2-en-1-one. **2022**, 133748 0
- 313 Prediction of Effective Photoelectron and Hole Separation in Type-I MoS2/PtSe2 van der Waals Junction. 6407-6411 0
- 312 Ultra-fast spectroscopy for high-throughput and interactive quantum chemistry. 0

311	Photoinduced small electron polarons generation and recombination in hematite. 2022 , 8,	4
310	Tuning the electron injection mechanism by changing the adsorption mode: the case study of Alizarin on TiO ₂ . 2022 , 101085	
309	Conformational and functional changes of the native neuropeptide somatostatin occur in the presence of copper and amyloid- β	1
308	Exchange Spin Coupling in Optically Excited States.	0
307	Schiff base-type Cu(I) complexes containing naphthylpyridyl-methanimine ligands featuring higher light-absorption capability: Synthesis, structures, and photophysical properties. 2022 , 224, 116002	1
306	Electronic structure calculations for inhomogeneous systems: Interfaces, surfaces, and nanocontacts. 2008 , 520, 525-560	
305	Exploring Structure-Property Relations of B,S-Doped Polycyclic Aromatic Hydrocarbons through the Trinity of Synthesis, Spectroscopy, and Theory.	1
304	Experimental and DFT Study of Transition Metal Doping in a Zn-BDC MOF to Improve Electrical and Visible Light Absorption Properties.	2
303	Going beyond the Electric-Dipole Approximation in the Calculation of Absorption and (Magnetic) Circular Dichroism Spectra including Scalar Relativistic and Spin-Orbit Coupling Effects.	0
302	An interpretation of quantum foundations based on density functional theory and polymer self-consistent field theory.	0
301	Spectroscopic Investigation and Density Functional Theory prediction of First and Second order Hyperpolarizabilities of 1-(4-Bromophenyl)-3-(2,4-dichlorophenyl)-prop-2-en-1-one. 2022 , 133807	2
300	Computational Analyses of Plasmonics of a Silver Nanoparticle in a Vacuum and in a Water Solution by Classical Electronic and Molecular Dynamics Simulations. 2022 , 126, 4762-4771	
299	Designing of PC 31 BM based acceptors for dye-sensitized solar cell.	0
298	Optical response of two-dimensional systems: Insights from classical electromagnetism to ab initio calculations. 2022 , 106,	0
297	Photochemical and thermochemical pathways to S ₂ and polysulfur formation in the atmosphere of Venus. 2022 , 13,	0
296	Time-dependent coherent light pulse absorption from macroscopic Maxwell equations. 2022 , 106,	0
295	Large Exciton-Driven Linear and Nonlinear Optical Processes in Monolayers of Nitrogen Arsenide and Nitrogen Antimonide.	0
294	turboMagnon Γ A code for the simulation of spin-wave spectra using the Liouville-Lanczos approach to time-dependent density-functional perturbation theory. 2022 , 108500	

293	High-order harmonic generation of 1-nonene under linearly polarized laser pulses. 2022 , 106,	3
292	Bandgap engineering and modulation of thermodynamic, and optical properties of III-N monolayers XN (X = In, Ga & Al) by mutual alloying. 2022 , 97, 095806	
291	Orbital-Free Methods for Plasmonics: Linear Response.	3
290	Insights in the Scavenging Mechanism of Trichloromethyldioxy Radical with Lycopene: a DFT/TD-DFT Study.	
289	A novel aureothin diepoxide derivative from Streptomyces sp. NIIST-D31 strain. 2022 , 75, 491-497	0
288	Photoinduced Small Hole Polarons Formation and Recombination in All-Inorganic Perovskite from Quantum Dynamics Simulation. 2022 , 13, 7532-7540	4
287	Extensibility of Hohenberg-Kohn Theorem to General Quantum Systems. 2200041	0
286	Electronic origin of x-ray absorption peak shifts. 2022 , 106,	
285	Benchmarking time-dependent density functional theory for singlet excited states of thermally activated delayed fluorescence chromophores. 2022 , 4,	1
284	Effect of bridging units on the photophysical properties of 4-NEt ₂ -appended salenRuthenium complexes.	0
283	Design of new hole transport materials based on triphenylamine derivatives using different linkers for the application in perovskite solar cells. A theoretical study. 10,	0
282	Koopmans Spectral Functionals in Periodic Boundary Conditions.	1
281	Solvent Dependent Nuclear Magnetic Resonance Molecular Parameters Based on a Polarization Consistent Screened Range Separated Hybrid Density Functional Theory Framework.	
280	Multiscale Strategy for Predicting Radiation Chemistry in Polymers.	1
279	Towards the description of charge transfer states in solubilised LHCII using subsystem DFT.	0
278	Multielectron effect in XUV light-driven strong-field ionization beyond the dipole approximation. 2022 , 39, 2486	0
277	Tetrathiafulvalene Derivatives as Hole-Transporting Materials in Perovskite Solar Cell. 2022 , 126, 5079-5088	1
276	Experimental and Computational Study of Novel Pyrazole Azo Dyes as Colored Materials for Light Color Paints. 2022 , 15, 5507	0

275	Screening the Band Shape of Molecules by Optimal Tuning of Range-Separated Hybrid Functional with TD-DFT: A Molecular Designing Approach. 2022 , 126, 5252-5264	0
274	Carbon Nanodots from an In Silico Perspective. 2022 , 122, 13709-13799	2
273	AI-Egens-NLOphores coumarin-triphenylamine chalcone derivatives: Synthesis, photophysical properties and DFT computational study. 2022 , 134009	0
272	Highly Efficient Near-Infrared Thermally Activated Delayed Fluorescent Emitters in Non-Doped Electroluminescent Devices.	0
271	Highly Efficient Near-Infrared Thermally Activated Delayed Fluorescent Emitters in Non-Doped Electroluminescent Devices.	
270	Effective lifetime of non-equilibrium carriers in semiconductors from non-adiabatic molecular dynamics simulations. 2022 , 2, 486-493	0
269	An optimally tuned range-separated hybrid starting point for ab initio GW plus Bethe-Salpeter equation calculations of molecules. 2022 , 157, 074103	6
268	Multistate Density Functional Theory of Excited States. 2022 , 13, 7762-7769	3
267	Field induced electron emission from graphene nanostructures. 2022 , 3, 034001	
266	Magnons in antiferromagnetic bcc Cr and Cr ₂ O ₃ from time-dependent density functional theory. 2022 , 106,	
265	UV-Visible Absorption Spectra of Solvated Molecules by Quantum Chemical Machine Learning. 2022 , 18, 4891-4902	1
264	Perspective: New directions in dynamical density functional theory.	0
263	Opto-electronic properties of isomers of azobispyridine. 2022 , 805, 139956	
262	Luminescence color tuning of the four-coordinate N-heterocyclic carbene (NHC) copper(I) complexes with imidazolylidene ligand functionalized by thiazole/benzoxazole moiety. 2022 , 977, 122469	0
261	In-depth analysis of the photophysics of BOPAHY dyes in solution, glass and film. 2022 , 206, 110662	0
260	Electronic, optical and thermoelectric properties of WSe ₂ /hN 2D interface: A DFT study. 2022 , 354, 114889	
259	Revealing the nature of localized and delocalized excitations of DNA. 2022 , 170, 110960	
258	The synthesis and spectral study of thiazolo[5,4-d]thiazole based small molecules using 1,3,4-oxadiazole as a linker for organic electronics. 2023 , 434, 114217	1

- 257 A new framework for frequency-dependent polarizable force fields. **2022**, 157, 124106 1
- 256 Ultrafast Charge-Transfer Dynamics in a Visible-Light-Excited Iridium(III) Terpyridine 2-Phenylpyridine Complex Studied by Femtosecond X-ray Absorption Spectroscopy. **2022**, 114267 0
- 255 Methoxylated flavones with potential therapeutic and photo-protective attributes: Theoretical investigation of substitution effect. **2022**, 203, 113387 1
- 254 Effect of oligothiophene spacer length on photogenerated charge transfer from perylene diimide to boron-doped diamond electrodes. **2022**, 248, 111984 0
- 253 The synthesis and photoelectrical performances of perylenediimide-based devices as an interface layer in metal-organic-semiconductors. **2022**, 286, 116036 0
- 252 Density-functional theory. **2023**, 27-65 0
- 251 The resolution-vs.-accuracy dilemma in machine learning modeling of electronic excitation spectra. 0
- 250 Relativistic Density-Functional Theories. **2022**, 1-32 0
- 249 Theoretical studies on the effects of bridge engineering on the photoelectric performance of Y6.. 0
- 248 Realization of switching between TADF and HLCT emissions through modulation of the intramolecular charge transfer character. **2022**, 10, 13124-13136 0
- 247 Donor-Acceptor naphthalimides and peryleneimides for all-solution-processed thin film lasers. 0
- 246 Toward highly efficient hyperfluorescence-based emitters through excited-states alignment using novel optimally tuned range-separated models. **2022**, 24, 23718-23736 0
- 245 Ni(II)/PA stabilization by hydrogen bond formation on the second coordination sphere: a DFT characterization. **2022**, 51, 12585-12595 0
- 244 First-principles simulations for spatial symmetry sensitive transient attosecond absorption of a polar semiconductor driven by an intense laser pulse. **2022**, 0
- 243 Theoretical study on the generation of single ultrashort attosecond pulse from the high-order harmonic generation of atom driven by optimized combined laser field. **2022**, 0 0
- 242 Rational designs of structurally similar TADF and HLCT emitters with benzo- or naphtho-carbazole units as electron donors. 0
- 241 Time-dependent density functional theory calculations of electronic friction in non-homogeneous media. **2022**, 24, 20239-20248 0
- 240 Advanced spectral analysis of complex molecular system. **2022**, 25-54 0

239	DFT Exchange: Sharing Perspectives on the Workhorse of Quantum Chemistry and Materials Science.	9
238	Ultrafast excited state dynamics of pyridine N-oxide derivative in solution; femtosecond fluorescence up-conversion and theoretical calculations. 2023 , 285, 121896	0
237	Time-Dependent Second-Order Green's Function Theory for Neutral Excitations. 2022 , 18, 5221-5232	0
236	Exchange-correlation effect in the charge response of a warm dense electron gas. 2022 , 106,	2
235	Carrier Multiplication in Transition Metal Dichalcogenides Beyond Threshold Limit. 2203400	1
234	Accurate Vertical Excitation Energies of BODIPY/Aza-BODIPY Derivatives from Excited-State Mean-Field Calculations.	0
233	Investigation of Ionization Potential in Quantum Dots Using the Stratified Stochastic Enumeration of Molecular Orbitals Method.	0
232	Tuning the Optical Properties and Conductivity of Bundles in Networks of Single-Walled Carbon Nanotubes. 2022 , 13, 8775-8782	2
231	Vibrationally resolved absorption and fluorescence spectra of flavins: A theoretical simulation in the gas phase.	0
230	First-principles simulation of light-ion microscopy of graphene. 2022 , 9, 045023	0
229	Predicting adsorption behavior of 6-mercaptopurine anticancer drug upon polyaxazoline nanocarrier: A theoretical study.	0
228	Density functionals for core excitations. 2022 , 157, 094107	1
227	Smart Covalent Organic Framework with Proton-Initiated Switchable Photocatalytic Aerobic Oxidation. 2022 , 12, 12398-12408	0
226	Charge-Transfer Transitions Govern the Reactivity and Photophysics of Vicinally Diphosphanyl-Substituted Diborapentacenes.	0
225	Cluster formation between an oxadiazole derivative with metal nanoclusters (Ag/Au/Cu), graphene quantum dot sheets, SERS studies, and solvent effects.	1
224	Making a case for femto-phono-magnetism with FePt. 2022 , 8,	1
223	Reliable prediction of the singlet-triplet gap in TADF molecules with GW/BSE approach. 2022 ,	0
222	Charge transfer in ultrafast isomerization of acetylene ions. 2022 , 106,	0

221	Electron dynamics in extended systems within real-time time-dependent density-functional theory.	2
220	The exact exchange–correlation potential in time-dependent density functional theory: Choreographing electrons with steps and peaks. 2022 , 3, 031307	0
219	Dynamics of the Energy Transfer Process in Eu(III) Complexes Containing Polydentate Ligands Based on Pyridine, Quinoline, and Isoquinoline as Chromophoric Antennae.	3
218	Ab initio simulation of laser-induced electronic and vibrational coherence. 2022 , 106,	0
217	Molecular Dynamics and TD-DFT Study of the Ternary Complexes of Cucurbit[8]uril with Aromatic Amino Acids and Auxiliary Ligands. 2022 , 7,	0
216	Conformational Landscape and Properties of Psilocybin: A Computational Approach. 2022 , 7,	1
215	Photo-induced phase-transitions in complex solids.	0
214	Combining machine learning and quantum chemical calculations for high-throughput virtual screening of thermally activated delayed fluorescence molecular materials: the impact of selection strategy and structural mutations. 2022 , 12, 30962-30975	1
213	A theoretical investigation of uranyl covalency via symmetry-preserving excited state structures.	0
212	General time-dependent configuration-interaction singles. I. Molecular case. 2022 , 106,	0
211	Theoretical study of infrared and ultraviolet spectra of fourteen isomers of C ₂₄ and comparison with astronomical observations.	0
210	Dramatic Plasmon Response to the Charge-Density-Wave Gap Development in 1T ₁ Se ₂ . 2022 , 129,	0
209	Aza-BODIPY chromophore as a unit of oligomers with outstanding optical properties. 2022 , 45,	0
208	Push-Pull Effect of Terpyridine Substituted by Triphenylamine Motive—Impact of Viscosity, Polarity and Protonation on Molecular Optical Properties. 2022 , 27, 7071	0
207	Constitutional isomerism of the linkages in donor–acceptor covalent organic frameworks and its impact on photocatalysis. 2022 , 13,	4
206	Double Excitation Energies from Quantum Monte Carlo Using State-Specific Energy Optimization.	0
205	Modelling of excited state potential energy surfaces with the Bethe–Salpeter equation formalism: The 4-(dimethylamino)benzonitrile twist.	3
204	Investigating a TADF Emitter by Time Resolved Near Infrared Spectroscopy.	0

203	Molecular Tuning in Diaryl-Capped Pyrrolo[2,3-d:5,4-d']bisthiazoles: Effects of Terminal Aryl Unit and Comparison to Dithieno[3,2-b:2',3'-d']pyrrole Analogues. 2022 , 27, 6638	0
202	Strain effects on high-harmonic generation in monolayer hexagonal boron nitride. 10,	0
201	The Role of Excited States of LNiII/III(Aryl)(Halide) Complexes in Ni-Halide Bond Homolysis in the Arylation of Csp ³ -H Bonds. 13215-13224	2
200	Minimal Active Space: NOSCF and NOSI in Multistate Density Functional Theory.	3
199	A Theoretical Investigation into the Homo- and Hetero-leptic Cu(I) Phosphorescent Complexes Bearing 2,9-dimethyl-1,10-phenanthroline and bis [2-(diphenylphosphino)phenyl]ether Ligand. 2022 , 15, 7253	0
198	Optical Properties and Light-Induced Charge Transfer in Selected Aromatic C60 Fullerene Derivatives and in Their Bulk Heterojunctions with Poly(3-Hexylthiophene). 2022 , 15, 6908	0
197	Relationship between the Molecular Geometry and the Radiative Efficiency in Naphthyl-Based Bis-Ortho-Carboranyl Luminophores. 2022 , 27, 6565	0
196	Combining Renormalized Singles GW Methods with the Bethe-Salpeter Equation for Accurate Neutral Excitation Energies.	2
195	DFT Studies of p-N,N-(Dimethylamino) Benzoic Acid with Para or Meta-Electron Withdrawing or Donating Moieties for Dye-Sensitized Solar Cells (DSSCs). 2022 , 2022, 1-12	0
194	Electrochemical Strategy for Proton Relay Installation Enhances the Activity of a Hydrogen Evolution Electrocatalyst.	0
193	Molecular Basis for the Difference in Singlet Oxygen Quantum Yield Between the First Genetically Encoded Photosensitizer, KillerRed, and its Monomeric Counterpart, SuperNova. 2022 , 7,	0
192	Transient Spin Injection Efficiencies at Ferromagnet/Metal Interfaces. 2201233	0
191	Dynamic interplay between thionine and DNA under carbon ion irradiation:a real-time first-principles study.	0
190	Nonadiabatic energy transfer in single wall carbon nanotube upon H ⁺ irradiation from time-dependent density-functional theory. 2022 , 531, 93-99	0
189	Molecular dynamics-based characterisation of early oxide in Fe/Cr alloys. 2022 , 9, 100087	0
188	Ab initio electronic stationary states for nuclear projectiles in solids. 2022 , 4,	0
187	Cyano-rich donor-acceptor-donor-type NLOphores containing dialkylated triazene and aniline groups. 2022 , 110894	0
186	PQKLP: Projected Quantum Kernel based Link Prediction in Dynamic Networks. 2022 , 196, 249-267	0



185	Computational approaches for nanocluster science. 2023 , 313-343	0
184	Quantum computational, spectroscopic, topological investigations and molecular docking studies on piperazine derivatives: A comparative study on Ethyl, Benzene and Furan sulfonyl Piperazine. 2023 , 1274, 134324	0
183	Static/dynamic first and second order hyperpolarizabilities, optimized structures, IR, UV-Vis, ¹ H and ¹³ C NMR spectra for effective charge transfer compounds: A DFT study. 2023 , 286, 122005	0
182	Quantum Transport in the Phase Space, the Wigner Equation. 2023 , 1559-1582	0
181	A theoretical study on the influence of N-containing heterocyclic ligands on the luminescence mechanisms (phosphorescence or TADF) of Au (III) complexes. 2023 , 113, 106676	0
180	Revisiting the Fluorescence of Benzothiadiazole Derivatives: Anti-Kasha Emission or Not?.	0
179	A Pragmatic Protocol for Determining Charge Transfer States of Molecules at Metal Surfaces by Constrained Density Functional Theory.	1
178	Real-space, real-time approach to quantum-electrodynamical time-dependent density functional theory. 2022 , 157, 194106	1
177	Extracting Inelastic Scattering Cross Sections for Finite and Aperiodic Materials from Electronic Dynamics Simulations.	0
176	Dynamical Born effective charges. 2022 , 106,	0
175	Franck-Condon factors within damped displacement harmonic oscillators: Solvent-enhanced absorption and fluorescence spectra.	0
174	Featuring a new computational protocol for the estimation of intensity and overall quantum yield in lanthanide chelates with applications to Eu(III) mercapto-triazole Schiff base ligands. 2022 , 100216	0
173	DFT, Molecular docking, Photocatalytic and Antimicrobial activity of coumarin enriched Cinnamon barkextract mediated silver nanoparticles. 2022 , 110176	0
172	Light-induced Weyl semiconductor-to-metal transition mediated by Peierls instability. 2022 , 106,	1
171	Cu-Catalyzed Atom Transfer Radical Polymerization: The Effect of Cocatalysts. 2200347	1
170	Synthesis and Spectroscopic Characterization of Selected Phenothiazines and Phenazines Rationalized Based on DFT Calculation. 2022 , 27, 7519	0
169	Influence of coherent vibrational excitation on the high-order harmonic generation of diatomic molecules. 2022 , 106,	0
168	High-order harmonic generation of the cyclo[18]carbon molecule irradiated by circularly polarized laser pulse.	3

167	Generalized Energy-Based Fragmentation Approach for the Electronic Emission Spectra of Large Systems.	1
166	Predicting adsorption behavior of Triacanthine anticancer drug with pure B12N12 nano-cage: A theoretical study. 2023 , 100, 100812	0
165	A bimetallic Ag ₁₅ Cu ₁₂ (S-c-C ₆ H ₁₁) ₁₈ (CH ₃ COO) ₃ nanocluster featuring an irregular Ag ₁₂ kernel.	0
164	Structural determination and anticholinesterase assay of C-glycosidic ellagitannins from Lawsonia inermis leaves: A study supported by DFT calculations and molecular docking. 2023 , 164, 105360	0
163	New coumarin derivative with potential antioxidant activity: Synthesis, DNA binding and in silico studies (Docking, MD, ADMET). 2023 , 16, 104440	1
162	Visualizing screening in noble-metal clusters: static vs. dynamic.	0
161	Theoretical study on the working mechanism and computational evidence of robust imine-based light-driven molecular motor. 2023 , 811, 140245	0
160	Modelling single atom catalysts for water splitting and fuel cells: A tutorial review. 2023 , 556, 232492	2
159	A full-potential and multiscale computational scheme for interactions between ultrafast intense laser pulses and condensed medium. 2023 , 284, 108633	0
158	Spectroscopic and DFT investigations: Mesogenic alpha-Naphthyl derivatives with central cinnamate and ester linkages. 2023 , 1275, 134629	0
157	First-Principles Calculation of Photoexcited Electron Dynamics of Nanostructures. 2022 , 1-35	0
156	Revealing incorporation of NH ₂ group into the edge of carbon dots for H ₂ O ₂ sensing via C=NH hydrogen bond interaction.	0
155	?????????????. 2022 ,	0
154	Frontier Molecular Orbitals Regulation Enables Efficient and Ultraviolet to Deep-Blue Narrowband Emission. 2202176	0
153	Either Accurate Singlet-Triplet Gaps or Excited-State Structures: Testing and Understanding the Performance of TD-DFT for TADF Emitters. 2022 , 18, 7702-7713	0
152	Enhanced energy deposition and carrier generation in silicon induced by two-color intense femtosecond laser pulses. 2022 , 106,	0
151	Study of optical absorption cross-section spectra and high-order harmonic generation of thymine, thymine glycol, and thymine dimer molecules. 2022 , 28,	0
150	Structure-property relationships of photofunctional diiridium(II) complexes with tetracationic charge and an unsupported Ir-Cl bond. 2022 , 5,	0

- 149 Control of Charge Carrier Relaxation at the Au/WSe₂ Interface by Ti and TiO₂ Adhesion Layers: Ab Initio Quantum Dynamics. 0
- 148 Theoretical Approach for Electron Dynamics and Ultrafast Spectroscopy (EDUS). 0
- 147 Dynamic Response of an Electron Gas: Towards the Exact Exchange-Correlation Kernel. **2022**, 129, 2
- 146 Implementation of real-time TDDFT for periodic systems in the open-source PySCF software package. 0
- 145 Modeling the electronic structure of organic materials: a solid-state physicist's perspective. **2023**, 6, 012001 0
- 144 Electron spill-out effect on third-order optical nonlinearity of metallic nanostructure. **2022**, 106, 0
- 143 Building on the Strengths of a Double-Hybrid Density Functional for Excitation Energies and Inverted Singlet-Triplet Energy Gaps. 0
- 142 Assessment of State-Averaged Driven Similarity Renormalization Group on Vertical Excitation Energies: Optimal Flow Parameters and Applications to Nucleobases. 1
- 141 Photolysis versus Photothermolysis of N₂O on a Semiconductor Surface Revealed by Nonadiabatic Molecular Dynamics. 3
- 140 Evidence of Graphene-like ZnO Nanostructures via Zinc Dimethoxide Hydrolysis/Condensation Under Ambient Conditions on a Au(111) Surface Using SERS: Simulation and Experiment. 0
- 139 QM/MM methods in studies of coinage metals: copper, silver, and gold interacting with biological and organic molecules. **2023**, 8, 0
- 138 Connections between many-body perturbation and coupled-cluster theories. **2022**, 157, 231102 3
- 137 Spectral features of the ferrous- H_2O complex in cytochrome P450: a revisit using TDDFT calculations. 0
- 136 Plane wave implementation of the magnetic force theorem for magnetic exchange constants: Application to bulk Fe, Co and Ni. 0
- 135 Impact of Hydrophobic/Hydrophilic Balance on Aggregation Pathways, Morphologies, and Excited-State Dynamics of Amphiphilic Diketopyrrolopyrrole Dyes in Aqueous Media. **2022**, 144, 22479-22492 0
- 134 Electron-electron interactions and high-order harmonics in solids. **2022**, 106, 0
- 133 Structure and Electronic Properties of Metalloboranes with General Formula $\text{Cp}^*_3(\text{E})\text{M}_3\text{B}_8\text{H}_8$ (M=Cr, Mo and W): The Effect of the Size of the Metal. **2022**, 7, 0
- 132 Highly Luminescent Blue Emitter with Balanced Hybridized-Locally and Charge-Transfer Excited-States Emission. 0

- 131 The subsystem quantum chemistry program Serenity. o
- 130 Photoionization and core resonances from range-separated time-dependent density-functional theory for open-shell states: Example of the lithium atom. o
- 129 Nonequilibrium two-particle self-consistent approach. **2022**, 106, o
- 128 Design and computational study of the novel nano-buds of C20@C60 with high NLO properties. **2023**, 134961 o
- 127 Fission dynamics, dissipation, and clustering at finite temperature. **2023**, 107, o
- 126 Spectroscopic and computational characterizations, Hirshfeld surface investigations, anticancer studies and molecular docking analysis of novel NLO 3-hydroxy-3',4',5,7-tetramethoxyflavone. **2023**, 46, o
- 125 Efficient fully-coherent quantum signal processing algorithms for real-time dynamics simulation. **2023**, 158, 024106 o
- 124 Dynamical analysis of attosecond molecular modes. **2023**, 107, o
- 123 4-Cyanotryptophan as a Sensitive Fluorescence Probe of Local Electric Field of Proteins. o
- 122 Practical Computational Chemistry Course for a Comprehensive Understanding of Organic, Inorganic, and Physical Chemistry: From Molecular Interactions to Chemical Reactions. o
- 121 State averaged CASSCF in AMOEBA polarizable water model for simulating nonadiabatic molecular dynamics with nonequilibrium solvation effects. **2023**, 158, 014101 o
- 120 Ultrafast hot electron dynamics in plasmonic nanostructures: experiments, modelling, design. **2023**, o
- 119 Electronic excited states in deep variational Monte Carlo. **2023**, 14, o
- 118 Ab Initio Calculations of XUV Ground and Excited States for First-Row Transition Metal Oxides. o
- 117 Exciton dispersion and exciton-phonon interaction in solids by time-dependent density functional theory. o
- 116 Emergence of Wigner oscillations in a model of real time cooling process: a time-dependent density-functional theory approach. **2023**, 35, 115602 o
- 115 Size- and Shape-Dependent Photoexcitation Electron Transfer in Metal Nanoclusters. **2023**, 127, 816-823 o
- 114 Conformational Landscape and Hydrogen Bonding Pattern of Psilocin: Computational Insights. **2023**, 8, o

- 113 Anisotropically Fused Clusters Form a 2D Superatomic Sheet Exhibiting Polarized Light Emission. o
- 112 Accelerating Parallel First-Principles Excited-State Calculation by Low-Rank Approximation with K-Means Clustering. **2022**, o
- 111 Synthesis, Photochemistry, Computational Study and Potential Application of New Styryl-Thiophene and Naphtho-Thiophene Benzylamines. **2023**, 24, 610 o
- 110 Electronic, Elastic, and Thermoelectric Properties of Half-Heusler Topological Semi-Metal HfIrAs from First-Principles Calculations. **2023**, 13, 37 o
- 109 Click-Type Synthesis of Homoconjugated Push-Pull Chromophores: Computational Investigation of Optical and Nonlinear Optical (NLO) Properties. o
- 108 Acoustic Plasmons in Nickel and Its Modification upon Hydrogen Uptake. **2023**, 13, 141 o
- 107 Investigation of parameters affecting the high harmonic generation from monolayer WS₂. o
- 106 Synthesis, Quantum Computational Analysis and Molecular Docking of 3-(2-Hydroxyphenyl)-1-Phenyl Propanone: A Combined Experimental and Theoretical Analysis. 1-30 o
- 105 Diquat based Dyes: A new Class of Photoredox Catalysts and Their Use in Aerobic Thiocyanation. o
- 104 Efficient All-Electron Time-Dependent Density Functional Theory Calculations Using an Enriched Finite Element Basis. o
- 103 A Deep Understanding on the Effective Generation of Twisted Intramolecular Charge Transfer by Protonation in Thiazolo[5,4-d]thiazole Derivatives. **2023**, 127, 902-912 1
- 102 Observable Error Bounds of the Time-Splitting Scheme for Quantum-Classical Molecular Dynamics. **2023**, 61, 26-44 o
- 101 Light-Enhanced Cytotoxicity of Doxorubicin by Photoactivation. **2023**, 12, 392 2
- 100 Mechanisms and Energetics for Hydrogen Abstraction of Thymine Photosensitized by Benzophenone from Theoretical Principle. o
- 99 Structure and emission properties of dinuclear copper(i) complexes with pyridyltriazole. **2023**, 13, 3899-3909 o
- 98 Quantum Electrodynamics of Intense Laser-Matter Interactions: A Tool for Quantum State Engineering. **2023**, 4, o
- 97 Rational Design of Non-Fullerene Acceptors via Side-Chain and Terminal Group Engineering: A Computational Study. o
- 96 Thomas-Fermi and Other Density-Functional Theories. **2023**, 297-308 o

- 95 Noncollinear density functional theory. **2023**, 5, [DOI](#) 
- 94 Insights in the Scavenging Mechanism of Trichloromethyldioxy Radical with Lycopene: a DFT/TD-DFT Study. [DOI](#) 
- 93 Impact of structural sampling, coupling scheme and state of interest on the energy transfer in CP29. [DOI](#) 
- 92 Double-Hybrid Density Functional Theory for Core Excitations: Theory and Benchmark Calculations. [DOI](#) 
- 91 Uncovering extreme nonlinear dynamics in solids through time-domain field analysis. **2023**, 107, [DOI](#) 
- 90 Borophene molecular plasmons. **2023**, 176, 111267 [DOI](#) 
- 89 Ab-initio simulation of the structural, electronic and optical properties for the vacancy-ordered double perovskites ATiI (A = Cs or NH); a time-dependent density functional theory study. **2023**, 176, 111262 [DOI](#) 
- 88 ML-based Performance Portability for Time-Dependent Density Functional Theory in HPC Environments. **2022**, [DOI](#) 
- 87 Computational Investigation of Stability and Molecular Properties of C18BN Corannulene Molecules. **2022**, 67, S158-S168 [DOI](#) 
- 86 Role of Pi-Electron Density at the Interface of Small Molecule-Sensitized Solar Cells. **2023**, 127, 3928-3939 [DOI](#) 
- 85 Efficient calculation of k-integrated electron energy loss spectra: Application to monolayers of MoS₂, hBN, and graphene. **2023**, 107, [DOI](#) 
- 84 Strategy for Modulating Dihedral Angle Distribution for Enhanced Anti-Kasha TADF Behavior. **2023**, 127, 5567-5575 [DOI](#) 
- 83 Time-dependent exchange-correlation hole and potential of the electron gas. **2023**, 107, [DOI](#) 
- 82 Systematic Theoretical Study on the pH-Dependent Absorption and Fluorescence Spectra of Flavins. **2023**, 28, 3315 [DOI](#) 
- 81 P1 Push-Pull Dye as a Case Study in QM/MM Theoretical Characterization for Dye-sensitized Solar Cell Organic Chromophores**. **2023**, 8, [DOI](#) 
- 80 Real-Space and Real-Time Propagation for Correlated Electron Nuclear Dynamics Based on Exact Factorization. [DOI](#) 
- 79 Spectroscopic investigations, hirshfeld surface analysis, anticancer and molecular docking studies of new novel NLO 3-hydroxy-3',4',5,7-tetramethoxyflavone. **2023**, 101000 [DOI](#) 
- 78 oeINDO: Efficient determination of excitation energies and UV-Vis absorption spectra of nano-sized Zn, Cd, S and their complexes. **2023**, 1223, 114096 [DOI](#) 

- 77 DFT design of novel nano-bud from B₁₂N₁₂ and C₆₀ fullerene. **2023**, 136, 109909 ○
- 76 Know your building blocks: Time-resolved EPR spectroscopy reveals NDI-T2 and not T-NDI-T to resemble the electronic structure of PNDIT2. **2023**, 117, 106790 ○
- 75 New semiconductor halocadmte [CdnXm](2nBh) crystal structure, molecular conformation and theoretical investigations. **2023**, 322, 123954 ○
- 74 Synthesis and luminescence properties of the four-coordinate N-heterocyclic carbene (NHC) copper(I) complexes with different bisphosphine ligands. **2023**, 1285, 135504 ○
- 73 Synthesis, spectroscopic, and molecular interaction study of lead(II) complex of DL-alanine using experimental techniques and quantum chemical calculations. **2023**, 1283, 135208 ○
- 72 Computational underpinnings for the dimerization of para-aminothiophenol to dimercaptoazobenzene on copper surface. **2023**, 571, 111910 ○
- 71 Revealing the supramolecular interactions of the bis(azopyrenyl) dibenzo-18-crown-6-ether system. **2023**, 374, 121298 ○
- 70 Recent advances in the ab initio theory of solid-state defect qubits. **2023**, 12, 359-397 1
- 69 Robustness of electronic screening effects in electron spectroscopies: Example of V2O5. **2023**, 107, ○
- 68 A Holographic Principle for Non-Relativistic Quantum Mechanics. **2023**, 62, ○
- 67 A Theoretical Study on Rate Constants of Excited State Proton Transfer Reaction in Anthracene-Urea Derivatives. **2023**, 96, 215-222 ○
- 66 Anti-inflammatory Quinoline Alkaloids from the Roots of *Waltheria indica*. **2023**, 86, 276-289 ○
- 65 Molecular L-amino butyric acid and its crystals: Structural, electronic and optical properties. **2023**, 321, 123900 ○
- 64 Synthesis and structural analysis of titanium-dinitrogen complex supported by di-anionic guanidinate ligands. ○
- 63 A pH-Activatable Prodrug and Metal Prodrug Conjugate of Gossypol: Synthesis, Emergent Photophysical, Nanoscopic, Computational, and in-Vitro Cellular Studies. ○
- 62 Eliminating Artificial Boundary Conditions in Time-Dependent Density Functional Theory Using Fourier Contour Deformation. **2023**, 19, 1409-1420 ○
- 61 Unexpected dipole instabilities in small molecules after ultrafast XUV irradiation. **2023**, 107, ○
- 60 Highly Efficient Purely Organic Phosphorescence Light-Emitting Diodes Employing a Donor-Acceptor Skeleton with a Phenoxaselenine Donor. 2207003 ○

- 59 Strong field driven extreme nonlinear photoemission from individual single-walled carbon nanotubes. **2023**, 107, ○
- 58 WS22 database, Wigner Sampling and geometry interpolation for configurationally diverse molecular datasets . **2023**, 10, ○
- 57 Molecular and electronic structures, bonding analysis, and UV-Vis spectra predictions of quinolino[3,2-b]benzodiazepine and quinolino[3,2-b]benzoxazepine metal transition M(L)2Cl2 and M(L)Cl2 complexes. ○
- 56 Rapid Interlayer Charge Separation and Extended Carrier Lifetimes due to Spontaneous Symmetry Breaking in Organic and Mixed Organic-Inorganic Dion-Jacobson Perovskites. **2023**, 145, 5297-5309 ○
- 55 Gaussian basis functions for an orbital-free-related density functional theory of atoms. ○
- 54 Ionization energies and ionization-induced structural changes in 2-phenylethylamine and its monohydrate. **2023**, 158, 114305 ○
- 53 NIR Absorbing Aromatic E-Ethylene Bridged Hexaphyrins (2.1.1.2.1.1): Synthesis, Characterization, and Protonation Studies. **2023**, 25, 1491-1496 ○
- 52 Molecular Orbitals. **2023**, ○
- 51 The Effect of Medium Acidity and Basicity on the Stability of Tryptophan and Formylkynurenine Species with Silver. **2023**, 3-15 ○
- 50 Experimental and quantum-chemical studies of electronic and spectral properties of titanium dioxide, modified with tin and lanthanum. ○
- 49 Exploring the attosecond laser-driven electron dynamics in the hydrogen molecule with different real-time time-dependent configuration interaction approaches. **2023**, ○
- 48 Synthesis, crystal structures, and DFT calculations: novel Mn(II), Co(II) and Ni(II) complexes of N-(pyridin-2-ylmethylene)methanamine with isothiocyanate as promising optical materials. **2023**, 55, ○
- 47 New Hybrid Compound Candidate as Photothermal Agent Based on DPP Derivatives and Toluidine Blue: A Theoretical Perspective. ○
- 46 Spin-Orbit Coupling Notably Retards Non-radiative Electron-Hole Recombination in Methylammonium Lead Triiodide Perovskites. **2023**, 14, 2715-2721 ○
- 45 Fluids and Electrolytes under Confinement in Single-Digit Nanopores. **2023**, 123, 2737-2831 ○
- 44 Assessing the Role of the Kohn-Sham Density in the Calculation of the Low-Lying Bethe-Salpeter Excitation Energies. **2023**, 127, 2618-2627 ○
- 43 Convergence of Møller-Plesset perturbation theory for excited reference states. **2023**, ○
- 42 Design of BPEA-based derivatives with high singlet fission performance: a theoretical perspective. **2023**, 25, 10071-10081 ○

- 41 Electronic density response of warm dense matter. **2023**, 30, 032705 2
- 40 Low-rank approximations to accelerate hybrid functional enabled real-time time-dependent density functional theory within plane waves. **2023**, 5, 014008 o
- 39 THz induced giant spin and valley currents. **2023**, 9, o
- 38 Hot carrier relaxation dynamics in non-stoichiometric CdSe quantum dots: computational insights. **2023**, 11, 8256-8264 o
- 37 Nanostructured system based on hydroxyapatite and curcumin: A promising candidate for osteosarcoma therapy. **2023**, o
- 36 Molecular Structure, Spectral Analysis, Molecular Docking and Physicochemical Studies of 3-Bromo-2-hydroxypyridine Monomer and Dimer as Bromodomain Inhibitors. **2023**, 28, 2669 o
- 35 Wavelength engineerable porous organic polymer photosensitizers with protonation triggered ROS generation. **2023**, 14, o
- 34 Synthesis, DFT calculations, α -glucosidase inhibitor activity, and docking studies on Schiff base metal complexes containing isothiocyanate. **2023**, 37, o
- 33 A DFT study of structural-stability, Mulliken charges, MEP, FMO, and NLO properties of trans alkenyl substituted chalcones conformers: theoretical study. o
- 32 Density functional theory. **2022**, o
- 31 Molecular recognition of tripeptides containing tryptophan by cucurbit[8]uril: A computational study. **2023**, 16, 104819 o
- 30 Multiple resonance induced thermally activated delayed fluorescence: effect of chemical modification. **2023**, 5, 014010 o
- 29 Laser-induced topological spin switching in a 2D van der Waals magnet. **2023**, 14, o
- 28 Design of new reversible and selective inhibitors of monoamine oxidase A and a comparison with drugs already approved. **2023**, 47, o
- 27 Noncollinear and Spin-Flip TDDFT in Multicollinear Approach. o
- 26 Curing the Divergence in Time-Dependent Density Functional Quadratic Response Theory. **2023**, 14, 3186-3192 o
- 25 Regulating the Electronic Structure of Metal Nanoclusters by Longitudinal Single-Dithiolate Substitution. **2023**, 14, 3216-3221 o
- 24 How Does Pseudo-Jahn-Teller Effect Induce the Photoprotective Potential of Curcumin?. **2023**, 28, 2946 o

- 23 Ab Initio Molecular Cavity Quantum Electrodynamics Simulations Using Machine Learning Models. ○
- 22 Insights into Laser-Matter Interaction from Inside: Wealth of Processes, Multiplicity of Mechanisms and Possible Roadmaps for Energy Localization. **2023**, 3-64 ○
- 21 State-Specific Configuration Interaction for Excited States. ○
- 20 Ultrafast Quantum Processes at the Nanoscale: Insights from Modeling. **2023**, 139-171 ○
- 19 Benchmark Study on Phosphorescence Energies of Anthraquinone Compounds: Comparison between TDDFT and UDFT. **2023**, 28, 3257 ○
- 18 Polyethylene Glycol 20k. Does It Fluoresce?. **2023**, 8, 14208-14218 ○
- 17 Some problems in density functional theory. **2023**, 113, ○
- 16 A combined spectroscopic and quantum chemical approach to study the molecular interaction between anti-inflammatory drug Hydrocortisone and amino acid L-Phenylalanine. **2023**, 135546 ○
- 15 Exploring Bethe-Salpeter Excited-State Dipoles: The Challenging Case of Increasingly Long Push-Pull Oligomers. **2023**, 14, 3727-3734 ○
- 14 Reimagining the Wave Function in Density Functional Theory: Exploring Strongly Correlated States in Pancake-Bonded Radical Dimers. ○
- 13 Vibrational and computational study on the characterization of Ag-Threonine complex. **2023**, ○
- 12 Non-Adiabatic Dynamics in Condensed Matter and Nanoscale Systems. **2023**, ○
- 11 Real-Time Time-Dependent Density Functional Theories With Large Time Step and Short Simulation Time. **2023**, ○
- 10 Localized Surface Plasmon Resonance in Metamaterials Composed of As_{1-x}Sb_x Semimetal Nanoparticles in Al_xGa_{1-x}As_{1-y}Sb_y Semiconductor Matrix. **2023**, 13, 1355 ○
- 9 Photoinduced Dynamics of 13,13'-Diphenylpropyl-β-carotene. **2023**, 28, 3505 ○
- 8 Enantiodivergent Photochemical Rearrangements Due to Different Coordination Modes at an Oxazaborolidine Lewis Acid Catalyst. 5896-5905 ○
- 7 Optical properties of quasi-two-dimensional objects from time-dependent density functional theory: Longitudinal versus transverse dielectric functions. **2023**, 107, ○
- 6 Synthesis, crystallographic and spectroscopic investigation, chemical reactivity, hyperpolarizabilities and in silico molecular docking study of (Z)-2N-(tert-butylimino)-3N-(4-methoxyphenyl) thiazolidin-4-one. **2023**, 135620 ○

- 5 Electronic stopping power in titanium for proton and helium ions from first-principle calculations. **2023**, 107, ○
- 4 Physical mechanisms of contact-electrification induced photon emission spectroscopy from interfaces. ○
- 3 Molecular Structure, Hydrogen Bonding Interactions and Docking Simulations of Nicotinamide (Monomeric and Trimeric Models) by Using Spectroscopy and Theoretical Approach. 1-19 ○
- 2 Synthesis, structure determination, NBO analysis and vibrational/electronic spectroscopic study of Iron(II) Bis(diethyldithiocarbamate) [Fe(DDTC)₂]. **2023**, 135618 ○
- 1 Biological Advantage of the Arrangements of C-Phycocyanin Chromophores in Phycobilisome from the Electronic Energy Transfer Viewpoint. **2023**, 96, 381-393 ○