Fanica Cimpoesu

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	AÂBinuclearÂFe(III)Dy(III)ÂSingleÂMoleculeÂMagnet.ÂQuantumÂEffectsÂandÂModels. Journal of the American Chemical Society, 2006, 128, 9008-9009.	6.6	252
2	Mechanism of Ferromagnetic Coupling in Copper(II)-Gadolinium(III) Complexes. Journal of the American Chemical Society, 2004, 126, 3321-3331.	6.6	141
3	Density Functional Theory (DFT) Study of Coumarin-based Dyes Adsorbed on TiO2 Nanoclusters—Applications to Dye-Sensitized Solar Cells. Materials, 2013, 6, 2372-2392.	1.3	74
4	Chiral Crystallization of a Heterodinuclear Ni-Ln Series: Comprehensive Analysis of the Magnetic Properties Inorganic Chemistry, 2012, 51, 11279-11293.	1.9	72
5	Syntheses, Structures, and Surface Aromaticity of the New Carbaalane [(AlH)6(AlNMe3)2(CCH2R)6] (R =) Tj ETQo of the American Chemical Society, 2002, 124, 5441-5448.	1 1 0.78 1 6.6	4314 rgBT 59
6	Lone-Pair-Electron-Driven Ionic Displacements in a Ferroelectric Metal–Organic Hybrid. Inorganic Chemistry, 2016, 55, 10337-10342.	1.9	51
7	Ligand field density functional theory calculation of the 4f2 → 4f15d1 transitions in the quantum cutter Cs2KYF6:Pr3+. Physical Chemistry Chemical Physics, 2013, 15, 13902.	1.3	50
8	Geometry, bonding and magnetism in planar triangulene graphene molecules with D3h symmetry: Zigzag (m=2,…,15). Chemical Physics, 2008, 354, 1-15.	0.9	47
9	Prospecting Lighting Applications with Ligand Field Tools and Density Functional Theory: A First-Principles Account of the 4f ⁷ –4f ⁶ 5d ¹ Luminescence of CsMgBr ₃ :Eu ²⁺ . Inorganic Chemistry, 2015, 54, 8319-8326.	1.9	39
10	Ferroelectric polarization of hydroxyapatite from density functional theory. RSC Advances, 2017, 7, 21375-21379.	1.7	37
11	Calculation of the 4f1→4f05d1 transitions in Ce3+-doped systems by Ligand Field Density Functional Theory. Chemical Physics Letters, 2013, 588, 260-266.	1.2	36
12	The theoretical account of the ligand field bonding regime and magnetic anisotropy in the DySc ₂ N@C ₈₀ single ion magnet endohedral fullerene. Physical Chemistry Chemical Physics, 2014, 16, 11337-11348.	1.3	36
13	New Synthetic and Structural Aspects in the Chemistry of Alkylaluminum Fluorides. The Mutual Influence of Hard and Soft Ligands and the Hybridization as Rigorous Structural Criterion#. Inorganic Chemistry, 2001, 40, 4947-4955.	1.9	28
14	The angular overlap model extended for two-open-shell f and d electrons. Physical Chemistry Chemical Physics, 2014, 16, 12282-12290.	1.3	28
15	Structure and Magnetism in Fe–Gd Based Dinuclear and Chain Systems. The Interplay of Weak Exchange Coupling and Zero Field Splitting Effects. Inorganic Chemistry, 2012, 51, 40-50.	1.9	27
16	Lightest Member of the Basic Carboxylate Structural Pattern:Â [Al3(μ3-O)(μ-O2CCF3)6(THF)3][(Me3Si)3CAl(O2CCF3)3]·C7H8. Inorganic Chemistry, 2002, 41, 1022-1025.	1.9	24
17	Influence of surface chemistry on the electronic properties of graphene nanoflakes. Chemical Physics Letters, 2011, 503, 91-96.	1.2	24
18	Ligand field density functional theory for the prediction of future domestic lighting. Physical Chemistry Chemical Physics, 2014, 16, 14625-14634.	1.3	24

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19	Synthesis, Structural Characterization, and Theoretical Treatment of an Unusual Organozirconium Hydroxide with the [Zr6(μ4-O)(μ-O)4(μ-OH)8] Coreậ€. Organometallics, 2003, 22, 3034-3038. Bonding and Magnetism in High Symmetry Nano-Sized Graphene Molecules: Linear Acenes Cost Bactor Bactor Structure (SUB-Cost and Structure) Structure (Sub-Structure) Structure (Sub-Structure) Structure (Sub-Structure) Structure) Structure (Sub-Structure) Structure (Sub-Structure) Structure (Sub-Structure) Structure) Structure (Sub-Structur	1.1	23
20	C _{6<1>m SUP>**2} H _{6<1>m } (<1>m =2,…10); Crenelated Hexangulenes C_{6(3<1>m SUP>**2−3<1>m }H_{6(2<1>m}	0.4	23
21	(<1>m=2,&nellip6); Zigzag Thangulenes C <sub< i="">m=2,&nellip6); Zigzag Thangulenes C<sub< i="">=2, Allow (I) > SUB> (I) = 2, 4(I) = 1, (I) = 1, (I) = 2, (I) =</sub<></sub<>	1.9	23
22	DFT Study of Binding and Electron Transfer from a Metal-Free Dye with Carboxyl, Hydroxyl, and Sulfonic Anchors to a Titanium Dioxide Nanocluster. International Journal of Photoenergy, 2013, 2013, 1-15.	1.4	23
23	Development and applications of the LFDFT: the non-empirical account of ligand field and the simulation of the fâ \in d transitions by density functional theory. Physical Chemistry Chemical Physics, 2015, 17, 18547-18557.	1.3	23
24	Vibrational properties of noble gas endohedral fullerenes. Physical Chemistry Chemical Physics, 2011, 13, 9609.	1.3	22
25	On The Density Functional Theory Treatment of Lanthanide Coordination Compounds: A Comparative Study in a Series of Cu–Ln (Ln = Gd, Tb, Lu) Binuclear Complexes. Inorganic Chemistry, 2017, 56, 9474-9485.	1.9	22
26	Tailoring the optical properties of lanthanide phosphors: prediction and characterization of the luminescence of Pr ³⁺ -doped LiYF ₄ . Physical Chemistry Chemical Physics, 2015, 17, 9116-9125.	1.3	18
27	Valence Bond Account of Triangular Polyaromatic Hydrocarbons with Spin: Combining Ab Initio and Phenomenological Approaches. Journal of Physical Chemistry C, 2019, 123, 6869-6880.	1.5	18
28	On exchange coupling and bonding in the Gd ₂ @C ₈₀ and Gd ₂ @C ₇₉ N endohedral dimetallo-fullerenes. Molecular Physics, 2015, 113, 1712-1727.	0.8	17
29	The angular overlap model of ligand field theory for f elements: An intuitive approach building bridges between theory and experiment. Coordination Chemistry Reviews, 2021, 441, 213981.	9.5	17
30	A ligand field theory-based methodology for the characterization of the Eu 2+ [Xe]4f 6 5d 1 excited states in solid state compounds. Chemical Physics Letters, 2015, 622, 120-123.	1.2	14
31	Metal-Organic Frameworks with d–f Cyanide Bridges: Structural Diversity, Bonding Regime, and Magnetism. Structure and Bonding, 2014, , 185-229.	1.0	11
32	Disorder, exchange and magnetic anisotropy in the room-temperature molecular magnet V[TCNE]x – A theoretical study. Computational Materials Science, 2014, 91, 320-328.	1.4	11
33	Noble gas endohedral fullerenes, Ng@C60 (Ng=Ar, Kr): a particular benchmark for assessing the account of non-covalent interactions by density functional theory calculations. Theoretical Chemistry Accounts, 2016, 135, 1.	0.5	11
34	Properties Design: Prediction and Experimental Validation of the Luminescence Properties of a New Eu ^{II} â€Based Phosphor. Chemistry - A European Journal, 2018, 24, 16276-16281.	1.7	11
35	The magnetic anisotropy and assembling of the lanthanide coordination units in [Fe(bpca)2][Er(NO3)3(H2O)4]NO3. Polyhedron, 2007, 26, 2069-2073.	1.0	10
36	The DFT rationalization of exchange and anisotropy in one-dimensional d-p magnets: The [MnIII(porphyrin)][TCNE] case study. Polyhedron, 2009, 28, 2039-2043.	1.0	10

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#	Article	IF	CITATIONS
37	Magnetic anisotropy and molecular assembling in d complex cation–f complex anion type coordination compounds. Solid State Sciences, 2009, 11, 760-765.	1.5	9
38	Ab initio study of exchange coupling for the consistent understanding of the magnetic ordering at room temperature in V[TCNE] x. Theoretical Chemistry Accounts, 2014, 133, 1.	0.5	9
39	Theoretical study of DHEA: comparative HF and DFT calculations of the electronic properties of a complex between DHEA and serotonin. Journal of Molecular Modeling, 2006, 12, 146-151.	0.8	8
40	DFT study of structure–properties correlations in [MnTPP][TCNE] quasi-one-dimensional molecular magnets. Theoretical Chemistry Accounts, 2011, 129, 847-857.	0.5	8
41	Broken symmetry DFT calculations of exchange coupling constants for manganese–porphyrin quasi-one-dimensional molecular magnets. Theoretical Chemistry Accounts, 2012, 131, 1.	0.5	8
42	Noncovalent effects in the coordination and assembling of the[Fe(bpca)2][Er(NO3)3(H2O)4]NO3 system. Open Chemistry, 2010, 8, 519-529.	1.0	7
43	Intra- and Inter-Molecular Spin Coupling in Phenalenyl Dimeric Systems. Journal of Physical Chemistry A, 2021, 125, 6893-6901.	1.1	7
44	Spin Coupling in the Supramolecular Structure of a New Tetra(Quinolineâ^'TEMPO)Yttrium(III) Complex. Inorganic Chemistry, 2007, 46, 660-669.	1.9	6
45	Monte Carlo simulation of magnetic ordering in the Gd3Fe5O12 Ising ferrite with garnet structure. Journal of Magnetism and Magnetic Materials, 2008, 320, 2149-2154.	1.0	6
46	Comparative computational IR, Raman and phosphorescence study of Ru- and Rh-based complexes. Molecular Physics, 2013, 111, 1526-1538.	0.8	6
47	Molecular and Supramolecular Interactions in Systems with Nitroxide-Based Radicals. International Journal of Molecular Sciences, 2019, 20, 4733.	1.8	6
48	New insights in the bonding regime and ligand field in Wernerian complexes. A density functional study. Polyhedron, 2013, 52, 183-195.	1.0	5
49	The Spin Coupling in the Polyaromatic Hydrocarbons and Carbon-Based Materials. , 2017, , 327-371.		5
50	Atoms in Generalized Orbital Configurations: Towards Atom-Dedicated Density Functionals. International Journal of Molecular Sciences, 2019, 20, 5943.	1.8	4
51	New Keys for Old Keywords. Case Studies within the Updated Paradigms of the Hybridization and Aromaticity. Monatshefte Für Chemie, 2005, 136, 1071-1085.	0.9	3
52	A MONTE CARLO SIMULATION OF MAGNETIC ORDERING IN ISING FERRITES OF FORMULA 5Fe2O3.3Y2O3 WITH GARNET STRUCTURE. Journal of Theoretical and Computational Chemistry, 2006, 05, 151-161.	1.8	3
53	New Syntheses, Analytic Spin Hamiltonians, Structural and Computational Characterization for a Series of Tri-, Hexa- and Hepta-Nuclear Copper (II) Complexes with Prototypic Patterns. Chemistry, 2021, 3, 411-439.	0.9	3
54	On the thermal stability and non-isothermal decomposition kinetics of some coordination compounds of cobalt, nickel and copper with a l²-dicarbonylic compound and 4-benzoylpyridine or 4,4'-dipyridyl as ligands. Thermochimica Acta, 1993, 221, 237-253.	1.2	2

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55	The association of dehydro-epiandrosterone and adenosine triphosphate acid: A DFT study of interactions between prototypic biologically active molecules. Computational and Theoretical Chemistry, 2009, 912, 32-37.	1.5	2
56	Monte Carlo Simulations of the Magnetic Behavior, Ordering Temperature and Magnetocaloric Effects in 1D, 2D and 3D Ferrimagnetic Systems. Journal of Nanoscience and Nanotechnology, 2015, 15, 263-268.	0.9	2
57	Spin Coupling and Magnetic Anisotropy in 1D Complexes with Manganese(III) Units and Carboxylate Bridges – Synthesis, Analysis, Calculations, and Models. European Journal of Inorganic Chemistry, 2018, 2018, 1409-1418.	1.0	1
58	Coordination Bonding: Electronic Structure and Properties. , 2018, , 503-612.		1
59	Electronic Structure of Linear Polyacenes. Current Organic Chemistry, 2018, 21, .	0.9	1
60	Characterizing the E⊗e Jahn–Teller Potential Energy Surfaces by Differential Geometry Tools. Symmetry, 2022, 14, 436.	1.1	1
61	The Density Functional Theory Account of Interplaying Long-Range Exchange and Dispersion Effects in Supramolecular Assemblies of Aromatic Hydrocarbons with Spin. Molecules, 2022, 27, 45.	1.7	1
62	Computational study of the molecular complexes between cholesterol and two isomers of the pralidoxime (PAM). Computational and Theoretical Chemistry, 2007, 813, 3-8.	1.5	0
63	Theoretical Calculations of Structure and Exchange Coupling of a Room-Temperature Molecular Magnet. , 2010, , .		Ο
64	New Keys for Old Keywords: Hybridization and Aromaticity, Graphs and Topology. , 2018, , 389-501.		0
65	The Modeling in Molecular Magnetism. , 2018, , 613-679.		0
66	Bonding in Rings and Clusters. , 2018, , 681-723.		0
67	Wave Function Theories and Electronic Structure Methods: Quantum Chemistry, from Atoms to Molecules. , 2018, , 107-220.		Ο
68	Electron Transfer and Dye Regeneration in Dye-Sensitized Solar Cells. , 2018, , .		0
69	Case Studies in the Challenge of Properties Design at Nanoscale. Advances in Chemical and Materials Engineering Book Series, 2017, , 148-184.	0.2	О