## Giovanni F Caramori

## List of Publications by Citations

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107<br/>papers1,312<br/>citations21<br/>h-index30<br/>g-index116<br/>ext. papers1,527<br/>ext. citations4<br/>avg, IF4.61<br/>L-index

#	Paper	IF	Citations
107	No need for a re-examination of the electrostatic notation of the hydrogen bonding: a comment. <i>Angewandte Chemie - International Edition</i> , <b>2015</b> , 54, 2596-9	16.4	69
106	Nanoparticle translocation through a lipid bilayer tuned by surface chemistry. <i>Physical Chemistry Chemical Physics</i> , <b>2013</b> , 15, 2282-90	3.6	64
105	A computational study of [2.2]cyclophanes. <i>Journal of Organic Chemistry</i> , <b>2005</b> , 70, 3242-50	4.2	64
104	The Nature of the RuNO Bond in Ruthenium Tetraammine Nitrosyl Complexes. <i>Organometallics</i> , <b>2007</b> , 26, 5815-5825	3.8	53
103	Computational study about through-bond and through-space interactions in [2.2]cyclophanes. <i>Journal of Physical Chemistry A</i> , <b>2007</b> , 111, 1705-12	2.8	40
102	Nitro-substituted 4-[(phenylmethylene)imino]phenolates: solvatochromism and their use as solvatochromic switches and as probes for the investigation of preferential solvation in solvent mixtures. <i>Journal of Organic Chemistry</i> , <b>2012</b> , 77, 10668-79	4.2	39
101	Platinum-Triggered Bond-Cleavage of Pentynoyl Amide and -Propargyl Handles for Drug-Activation. <i>Journal of the American Chemical Society</i> , <b>2020</b> , 142, 10869-10880	16.4	38
100	Sesquiterpene lactones from Vernonia scorpioides and their in vitro cytotoxicity. <i>Phytochemistry</i> , <b>2010</b> , 71, 1539-44	4	37
99	Ureasil-poly(ethylene oxide) hybrid matrix for selective adsorption and separation of dyes from water. <i>Langmuir</i> , <b>2014</b> , 30, 3857-68	4	36
98	The nature of Ru-NO bonds in ruthenium tetraazamacrocycle nitrosyl complexesa computational study. <i>Dalton Transactions</i> , <b>2012</b> , 41, 7327-39	4.3	36
97	Quinolinyl and quinolinyl N-oxide chalcones: synthesis, antifungal and cytotoxic activities. <i>European Journal of Medicinal Chemistry</i> , <b>2011</b> , 46, 4448-56	6.8	32
96	The versatile ruthenium(II/III) tetraazamacrocycle complexes and their nitrosyl derivatives. <i>Coordination Chemistry Reviews</i> , <b>2016</b> , 306, 652-677	23.2	30
95	Synthesis and Solvatochromism of Substituted 4-(Nitrostyryl)phenolate Dyes. <i>Journal of Organic Chemistry</i> , <b>2015</b> , 80, 7971-83	4.2	29
94	The nature of the interactions between Pt4 cluster and the adsorbates *H, *OH, and H2O. <i>Journal of Physical Chemistry A</i> , <b>2008</b> , 112, 11731-43	2.8	29
93	Cyclic trinuclear copper(I), silver(I), and gold(I) complexes: a theoretical insight. <i>Dalton Transactions</i> , <b>2015</b> , 44, 377-85	4.3	26
92	Strontium mono-chloride IA new molecule for the determination of chlorine using high-resolution graphite furnace molecular absorption spectrometry and direct solid sample analysis. <i>Spectrochimica Acta, Part B: Atomic Spectroscopy</i> , <b>2014</b> , 102, 1-6	3.1	26
91	How the electron-deficient cavity of heterocalixarenes recognizes anions: insights from computation. <i>Physical Chemistry Chemical Physics</i> , <b>2017</b> , 19, 24696-24705	3.6	25

90	Cucurbituril-Mediated Catalytic Hydrolysis: A Kinetic and Computational Study with Neutral and Cationic Dioxolanes in CB7. ACS Catalysis, <b>2018</b> , 8, 12067-12079	3.1	25
89	Ruthenium(II)/4,6-dimethyl-2-mercaptopyrimidine complexes: Synthesis, characterization, X-ray structures and in vitro cytotoxicity activities on cancer cell lines. <i>Polyhedron</i> , <b>2014</b> , 68, 312-318	7	23
88	Role of the cation formal charge in cation[Interaction. A survey involving the [2.2.2]paracyclophane host from relativistic DFT calculations. <i>New Journal of Chemistry</i> , <b>2015</b> , 39, 9963-936	88	22
87	Aromaticity and Homoaromaticity in Methano[10]annulenes. <i>Journal of Organic Chemistry</i> , <b>2007</b> , 72, 76-85	2	22
86	[2.2.2]Paracyclophane, preference for 16 or 118 coordination mode including Ag(I) and Sn(II): a survey into the cation interaction nature through relativistic DFT calculations. RSC Advances, 2015, 5, 7803-7811	7	21
85	Polyacetylenes from the leaves of Vernonia scorpioides (Asteraceae) and their antiproliferative and antiherpetic activities. <i>Phytochemistry</i> , <b>2013</b> , 95, 375-83		21
84	Isatin-Schiff base copper(II) complexes DFT study of the metal-ligand bonding situation.  International Journal of Quantum Chemistry, 2012, 112, 625-646	1	21
83	Oxindole-Schiff base copper(II) complexes interactions with human serum albumin: spectroscopic, oxidative damage, and computational studies. <i>Journal of Inorganic Biochemistry</i> , <b>2009</b> , 103, 1331-41	2	21
82	Anion Recognition by Organometallic Calixarenes: Analysis from Relativistic DFT Calculations.  Organometallics, 2018, 37, 2167-2176	8	20
81	Tuning Heterocalixarenes to Improve Their Anion Recognition: A Computational Approach. <i>Journal of Physical Chemistry A</i> , <b>2018</b> , 122, 3328-3336	8	19
8o	Ru-NO and Ru-NO2 bonding linkage isomerism in cis-[Ru(NO)(NO)(bpy)2](2+/+) complexes - a theoretical insight. <i>Dalton Transactions</i> , <b>2014</b> , 43, 8792-804	3	18
79	A computational study of tetrafluoro-[2.2]cyclophanes. <i>Journal of Physical Chemistry A</i> , <b>2008</b> , 112, 1178428	800	17
78	Copaifera duckei Oleoresin and Its Main Nonvolatile Terpenes: In Vitro Schistosomicidal Properties.  Chemistry and Biodiversity, <b>2016</b> , 13, 1348-1356	5	16
77	Ruthenophanes: Evaluating Cation[Interactions in [Ru([6-C16H12R4)(NH3)3]2+/3+Complexes. A Computational Insight. <i>Organometallics</i> , <b>2014</b> , 33, 2301-2312	8	16
76	Peculiar reactivity of a di-imine copper(II) complex regarding its binding to albumin protein. <i>Dalton Transactions</i> , <b>2013</b> , 42, 6386-96	3	15
75	Polar Order and Symmetry Breaking at the Boundary between Bent-Core and Rodlike Molecular Forms: When 4-Cyanoresorcinol Meets the Carbosilane End Group. <i>Chemistry - A European Journal</i> , 4.2 <b>2016</b> , 22, 8181-97	8	15
74	A ruthenium polypyridyl complex with the antihypertensive drug valsartan: Synthesis, theoretical calculations and interaction studies with human serum albumin. <i>Polyhedron</i> , <b>2016</b> , 114, 232-241	7	13
73	The two faces of hydrogen-bond strength on triple AAA-DDD arrays. <i>ChemPhysChem</i> , <b>2013</b> , 14, 3994-400 <sub>3</sub> 1.	2	13

72	Interactions of di-imine copper(II) complexes with albumin: competitive equilibria, promoted oxidative damage and DFT studies. <i>Journal of the Brazilian Chemical Society</i> , <b>2010</b> , 21, 1303-1317	1.5	13
71	In vitro cytotoxicity and structure-activity relationship approaches of ent-kaurenoic acid derivatives against human breast carcinoma cell line. <i>Phytochemistry</i> , <b>2018</b> , 156, 214-223	4	13
70	Shedding Light on the Nature of Host@uest Interactions in PAHs-ExBox4+ Complexes. <i>Journal of Physical Chemistry C</i> , <b>2016</b> , 120, 15480-15487	3.8	12
69	Analysis of the metalligand bonds in $[Mo(X)(NH2)3]$ (X = P, N, PO, and NO), $[Mo(CO)5(NO)]+$ , and $[Mo(CO)5(PO)]+$ . Theoretical Chemistry Accounts, <b>2008</b> , 120, 351-361	1.9	12
68	Synthesis, characterization and biological evaluation of new manganese metal carbonyl compounds that contain sulfur and selenium ligands as a promising new class of CORMs. <i>Dalton Transactions</i> , <b>2019</b> , 48, 5574-5584	4.3	11
67	Synthetic pathway for a new series of liquid crystal 2,6-disubstituted imidazo[2,1-b][1,3,4]thiadiazole. <i>Liquid Crystals</i> , <b>2013</b> , 40, 570-580	2.3	11
66	In vivo and in silico anti-inflammatory mechanism of action of the semisynthetic (-)-cubebin derivatives (-)-hinokinin and (-)-O-benzylcubebin. <i>Bioorganic and Medicinal Chemistry Letters</i> , <b>2017</b> , 27, 176-179	2.9	10
65	The ability of Ex2Box4+ to interact with guests containing Electron-rich and Electron-poor moieties. <i>International Journal of Quantum Chemistry</i> , <b>2018</b> , 118, e25607	2.1	9
64	Ruthenium(II) complexes of N-heterocyclic carbenes derived from imidazolium-linked cyclophanes. <i>Dalton Transactions</i> , <b>2014</b> , 43, 14710-9	4.3	9
63	Mechanism of Palladium(II)-Mediated Uncaging Reactions of Propargylic Substrates. <i>ACS Catalysis</i> , <b>2019</b> , 9, 3792-3799	13.1	8
62	Proton-induced generation of remote N-heterocyclic carbene-Ru complexes. <i>Chemistry - A European Journal</i> , <b>2015</b> , 21, 106-10	4.8	8
61	Bonding Analysis in Homo- and Hetero-Trihalide Species: A Charge Displacement Study. <i>European Journal of Inorganic Chemistry</i> , <b>2016</b> , 2016, 3804-3812	2.3	8
60	Metalligand Bonding Situation in Ruthenophanes Containing Multibridged Cyclophanes. <i>Organometallics</i> , <b>2017</b> , 36, 3465-3470	3.8	8
59	Quest for Insight into Ultrashort C-HIPProximities in Molecular "Iron Maidens". <i>Journal of Organic Chemistry</i> , <b>2018</b> , 83, 5114-5122	4.2	7
58	ESI-QTof-MS characterization of hirsutinolide and glaucolide sesquiterpene lactones: Fragmentation mechanisms and differentiation based on Na /H adducts interactions in complex mixture. <i>Journal of Mass Spectrometry</i> , <b>2019</b> , 54, 915-932	2.2	7
57	Electronic properties and metal-ligand bonding situation in Eu(III) complexes containing tris(pyrazolyl)borate and phenantroline ligands. <i>Journal of Luminescence</i> , <b>2017</b> , 182, 137-145	3.8	7
56	Reverse Solvatochromism of Imine Dyes Comprised of 5-Nitrofuran-2-yl or 5-Nitrothiophen-2-yl as Electron Acceptor and Phenolate as Electron Donor. <i>Chemistry - A European Journal</i> , <b>2018</b> , 24, 9364-937	76 <sup>4.8</sup>	6
55	Bond Analysis in DihalogenHalide and DihalogenDimethylchalcogenide Systems. <i>European Journal of Inorganic Chemistry</i> , <b>2018</b> , 2018, 1007-1015	2.3	6

## (2020-2016)

54	Hydrazine decomposition on a small platinum cluster: the role of N2H5 intermediate. <i>Theoretical Chemistry Accounts</i> , <b>2016</b> , 135, 1	1.9	6	
53	Spectroscopy and theoretical studies of natural melanin (eumelanin) and its complexation by iron(III). <i>Journal of Coordination Chemistry</i> , <b>2014</b> , 67, 986-1001	1.6	6	
52	The influence of L ligands on the {RuNO}6/7 bonding situation in cis-[Ru(NO)(NO2)L1월]q complexes: a theoretical insight. <i>RSC Advances</i> , <b>2015</b> , 5, 69057-69066	3.7	5	
51	Ruthenium nitrosyl complexes containing pyridine-functionalized carbenes IA theoretical insight. <i>Journal of Organometallic Chemistry</i> , <b>2015</b> , 799-800, 54-60	2.3	5	
50	Coordination among Bond Formation/Cleavage in a Bifunctional-Catalyzed Fast Amide Hydrolysis: Evidence for an Optimized Intramolecular -Protonation Event. <i>Journal of Organic Chemistry</i> , <b>2020</b> , 85, 4663-4671	4.2	5	
49	Polylactic acid, maleic anhydride and dicumyl peroxide: NMR study of the free-radical melt reaction product. <i>Polymer Degradation and Stability</i> , <b>2018</b> , 155, 1-8	4.7	5	
48	Helicenes as Molecular Tweezers in the Formation of Cation-Complexes. Bonding and Circular Dichroism Properties from Relativistic DFT Calculations. <i>ChemPhysChem</i> , <b>2018</b> , 19, 2321-2330	3.2	5	
47	The usefulness of energy decomposition schemes to rationalize host-guest interactions. <i>Dalton Transactions</i> , <b>2020</b> , 49, 17457-17471	4.3	5	
46	On the cationLapabilities of small all sp2-carbon host structures. Evaluation of [6.8]3cyclacene from relativistic DFT calculations. <i>International Journal of Quantum Chemistry</i> , <b>2019</b> , 119, e25811	2.1	5	
45	How Do Secondary Phosphine Oxides Interact with Silver Nanoclusters? Insights from Computation. Journal of Physical Chemistry C, 2018, 122, 21449-21461	3.8	5	
44	Binding affinity studies of 1,2,3-triazole copper(II) complexes to human serum albumin. <i>Journal of Coordination Chemistry</i> , <b>2018</b> , 71, 1894-1909	1.6	5	
43	Evaluation of Electron Donation as a Mechanism for the Stabilisation of Chalcogenate-Protected Gold Nanoclusters. <i>ChemPhysChem</i> , <b>2016</b> , 17, 3102-3111	3.2	4	
42	Transport properties of ruthenophanes 🖪 theoretical insight. Chemical Physics, 2016, 478, 23-33	2.3	4	
41	Removal of the emerging contaminant bisphenol A by an ureasil-PEO hybrid membrane: experimental study and molecular dynamic simulation. <i>Environmental Science and Pollution Research</i> , <b>2017</b> , 24, 18421-18433	5.1	4	
40	Hydrogen bond and the resonance effect on the formamide water complexes. <i>International Journal of Quantum Chemistry</i> , <b>2012</b> , 112, 1401-1420	2.1	4	
39	How does the acidic milieu interfere in the capability of ruthenium nitrosyl complexes to release nitric oxide?. <i>New Journal of Chemistry</i> , <b>2020</b> , 44, 773-779	3.6	4	
38	Evaluation of Lignans from Piper cubeba against Schistosoma mansoni Adult Worms: A Combined Experimental and Theoretical Study. <i>Chemistry and Biodiversity</i> , <b>2019</b> , 16, e1800305	2.5	4	
37	In Silico Design of Cylindrophanes: The Role of Functional Groups in a Fluoride Selective Host. <i>ChemPhysChem</i> , <b>2020</b> , 21, 1989-2005	3.2	3	

36	Understanding the interplay between Hand cation-Interactions in [janusene-Ag] host-guest systems: a computational approach. <i>Dalton Transactions</i> , <b>2019</b> , 48, 13281-13292	4.3	3
35	Investigating the Ritter Type Reaction of EMethylene-Ehydroxy Esters in Acidic Medium: Evidence for the Intermediacy of an Allylic Cation. <i>European Journal of Organic Chemistry</i> , <b>2013</b> , 2013, 5180-5187	3.2	3
34	A New Exploration of the Torsional Energy Surface of n-Pentane Using Molecular Models and Molecular Modeling Software. <i>Journal of Chemical Education</i> , <b>2005</b> , 82, 1800	2.4	3
33	Solvatochromism of new substituted 4-[(E)-(4-nitrophenyl)diazenyl]phenolate dyes. <i>Journal of Molecular Liquids</i> , <b>2020</b> , 301, 112330	6	3
32	On the recognition of chloride, bromide and nitrate anions by anthracene quaramide conjugated compounds: a computational perspective. <i>New Journal of Chemistry</i> , <b>2020</b> , 44, 17831-17839	3.6	3
31	Kinetics and adsorption calculations: insights into the MgO-catalyzed detoxification of simulants of organophosphorus biocides. <i>Journal of Materials Chemistry A</i> , <b>2020</b> , 8, 19011-19021	13	3
30	Shedding Light on the Hydrolysis Mechanism of cis, trans-[Ru(dmso)Cl] Complexes and Their Interactions with DNA-A Computational Perspective. <i>Journal of Physical Chemistry B</i> , <b>2019</b> , 123, 457-467	<b>7</b> 3.4	3
29	Shedding light on the electronic structure of [Ru(IJCH)(NH)] complex: a computational insight. <i>Journal of Molecular Modeling</i> , <b>2019</b> , 25, 11	2	3
28	Advances in bonding and properties of inorganic systems from relativistic calculations in Latin America. <i>International Journal of Quantum Chemistry</i> , <b>2019</b> , 119, e25777	2.1	3
27	Metal-ligand bonding situation in ruthenophanes containing i,j-xylylene-linked bis(NHC)cyclophane ligands. <i>Journal of Organometallic Chemistry</i> , <b>2017</b> , 830, 100-108	2.3	2
26	Synthesis, characterization and photoinduced CO-release by manganese(I) complexes. <i>New Journal of Chemistry</i> , <b>2020</b> , 44, 10892-10901	3.6	2
25	Tracking the role of trans-ligands in ruthenium NO bond lability: computational insight. <i>New Journal of Chemistry</i> , <b>2020</b> , 44, 11448-11456	3.6	2
24	Aromaticity and Homoaromaticity in Methano[10]annulenes <i>Journal of Organic Chemistry</i> , <b>2007</b> , 72, 2698-2698	4.2	2
23	Efeitos de substituintes na liga <b>ő</b> de hidrog <b>ő</b> io do 3-hidroxipropenal. <i>Quimica Nova</i> , <b>2006</b> , 29, 1187-1192	1.6	2
22	What is the driving force behind molecular triangles and their guests? A quantum chemical perspective about host-guest interactions. <i>Physical Chemistry Chemical Physics</i> , <b>2020</b> , 22, 19213-19222	3.6	2
21	Supersaturating drug delivery systems containing fixed-dose combination of two antihypertensive drugs: Formulation, in vitro evaluation and molecular metadynamics simulations. <i>European Journal of Pharmaceutical Sciences</i> , <b>2021</b> , 163, 105860	5.1	2
20	Solvation Enhances the Distinction between Carboxylated Armchair and Zigzag Single-Wall Carbon Nanotubes (SWNT-COOH). <i>Journal of Physical Chemistry C</i> , <b>2017</b> , 121, 9516-9527	3.8	1
19	The anionic recognition mechanism based on polyol and boronic acid receptors. <i>New Journal of Chemistry</i> , <b>2020</b> , 44, 5564-5571	3.6	1

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18	Shedding light on the bonding situation of triangular and square heterometallic clusters: computational insight. <i>New Journal of Chemistry</i> , <b>2020</b> , 44, 5079-5087	3.6	1
17	Tracking the absence of anionInteractions in modified [23](1,3,5)cyclophanes: insights from computation. <i>New Journal of Chemistry</i> , <b>2019</b> , 43, 13271-13281	3.6	1
16	The bonding situation in heteromultimetallic carbonyl complexes. <i>Dalton Transactions</i> , <b>2020</b> , 49, 16762	2-1463/7	1 1
15	A theoretical investigation on the aminolysis of pyromellitic and 1,4,5,8-naphthalenetetracarboxylic dianhydrides. <i>Computational and Theoretical Chemistry</i> , <b>2019</b> , 1147, 13-19	2	1
14	Can the relative positions (cistrans) of ligands really modulate the coordination of NO in ruthenium nitrosyl complexes?. <i>New Journal of Chemistry</i> , <b>2021</b> , 45, 1658-1666	3.6	1
13	The design of anion-Interactions and hydrogen bonds for the recognition of chloride, bromide and nitrate anions. <i>Physical Chemistry Chemical Physics</i> , <b>2021</b> , 23, 11455-11465	3.6	1
12	Box-Shaped Hosts: Evaluation of the Interaction Nature and Host Characteristics of ExBox Derivatives in Host-Guest Complexes from Computational Methods <b>2021</b> , 395-416		1
11	The simultaneous recognition mechanism of cations and anions using macrocyclic-iodine structures: insights from dispersion-corrected DFT calculations. <i>Physical Chemistry Chemical Physics</i> , <b>2020</b> , 22, 237	9 <i>5</i> -238	:03 <sup>3</sup>
10	Development of new dinuclear Fe(III) coordination compounds with nanomolar antitrypanosomal activity. <i>Dalton Transactions</i> , <b>2021</b> , 50, 12242-12264	4.3	O
9	Reverse solvatochromism in a family of probes having 2,6difertButylphenolate as electrondonor and 4ditrophenyl as electrondocceptor groups. <i>Dyes and Pigments</i> , <b>2022</b> , 203, 110376	4.6	O
8	Development, structural, spectroscopic and investigation of new complexes relevant as anti-toxoplasma metallopharmacs. <i>Journal of Molecular Structure</i> , <b>2022</b> , 133380	3.4	O
7	A ruthenium nitrosyl cyclam complex with appended anthracenyl fluorophore. <i>Polyhedron</i> , <b>2019</b> , 173, 114117	2.7	
6	Are DFT Methods Able to Predict Reduction Potentials of Ruthenium Nitrosyl Complexes Accurately?. <i>Journal of Physical Chemistry A</i> , <b>2020</b> , 124, 6186-6192	2.8	
5	The Nature of the RuNO Bond in Ruthenium Tetraammine Nitrosyl Complexes <i>Organometallics</i> , <b>2008</b> , 27, 1660-1660	3.8	
4	Nature of hydride and halide encapsulation in Ag cages: insights from the structure and interaction energy of [Ag(X){SP(OPr)}] (X = H, F, Cl, Br, I) from relativistic DFT calculations <i>Physical Chemistry Chemical Physics</i> , <b>2021</b> , 24, 452-458	3.6	
3	Theoretical study of chloride complexes with hybrid macrocycles. <i>New Journal of Chemistry</i> , <b>2021</b> , 45, 463-470	3.6	
2	Designing boron and metal complexes for fluoride recognition: a computational perspective. <i>Physical Chemistry Chemical Physics</i> , <b>2021</b> , 23, 22768-22778	3.6	
1	The Edonor/acceptor trans effect on NO release in ruthenium nitrosyl complexes: a computational insight. <i>New Journal of Chemistry</i> , <b>2021</b> , 45, 8949-8957	3.6	