## Mercedes Alonso

## List of Publications by Year in descending order

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270111 299063 2,052 76 25 42 citations h-index g-index papers 82 82 82 2120 docs citations times ranked citing authors all docs

| #  | Article   | IF   | CITATIONS |
|----|---|------|-----------|
| 1  | Imine hydrogenation with simple alkaline earth metal catalysts. Nature Catalysis, 2018, 1, 40-47.   | 16.1 | 151       |
| 2  | A benchmark for the non-covalent interaction (NCI) index or $\hat{a} \in \ \mid \ $ is it really all in the geometry?. Theoretical Chemistry Accounts, 2016, 135, 1.  | 0.5  | 124       |
| 3  | Understanding the Fundamental Role of π <i>/</i> §, Ïf <i>/</i> §f, and Ïf <i>/</i> §F Dispersion Interactions in Shaping Carbonâ€Based Materials. Chemistry - A European Journal, 2014, 20, 4931-4941.   | 1.7  | 109       |
| 4  | Simple Alkalineâ€Earth Metal Catalysts for Effective Alkene Hydrogenation. Angewandte Chemie -<br>International Edition, 2018, 57, 15177-15182.   | 7.2  | 87        |
| 5  | New electron delocalization tools to describe the aromaticity in porphyrinoids. Physical Chemistry Chemical Physics, 2018, 20, 2787-2796.   | 1.3  | 86        |
| 6  | From Dibismuthenes to Three―and Twoâ€Coordinated Bismuthinidenes by Fine Ligand Tuning: Evidence for Aromatic BiC <sub>3</sub> N Rings through a Combined Experimental and Theoretical Study. Chemistry - A European Journal, 2015, 21, 16917-16928.  | 1.7  | 76        |
| 7  | Alkene Transfer Hydrogenation with Alkalineâ€Earth Metal Catalysts. Angewandte Chemie -<br>International Edition, 2019, 58, 4248-4253.  | 7.2  | 65        |
| 8  | Conductance Switching in Expanded Porphyrins through Aromaticity and Topology Changes. Journal of the American Chemical Society, 2018, 140, 1313-1326.  | 6.6  | 56        |
| 9  | Revisiting sulfur H-bonds in proteins: The example of peroxiredoxin AhpE. Scientific Reports, 2016, 6, 30369.   | 1.6  | 52        |
| 10 | Stibinidene and Bismuthinidene as Twoâ€Electron Donors for Transition Metals (Co and Mn). Chemistry - A European Journal, 2016, 22, 7376-7380.  | 1.7  | 51        |
| 11 | Topology Switching in [32]Heptaphyrins Controlled by Solvent, Protonation, and <i>meso</i> â€Substituents. Chemistry - A European Journal, 2013, 19, 1617-1628.   | 1.7  | 49        |
| 12 | Viability of Möbius Topologies in [26]―and [28]Hexaphyrins. Chemistry - A European Journal, 2012, 18, 10916-10928.  | 1.7  | 48        |
| 13 | A comparative study of the structure and bonding in heavier pnictinidene complexes $[(ArE)M(CO) < sub > n < /sub > ]$ (E = As, Sb and Bi; M = Cr, Mo, W and Fe). Dalton Transactions, 2017, 46, 3556-3568.  | 1.6  | 44        |
| 14 | Performance of Localized Coupled Cluster Methods in a Moderately Strong Correlation Regime: HÃ⅓ckel–Möbius Interconversions in Expanded Porphyrins. Journal of Chemical Theory and Computation, 2020, 16, 3641-3653.  | 2.3  | 44        |
| 15 | Exploring the structure–aromaticity relationship in Hückel and Möbius N-fused pentaphyrins using DFT. Physical Chemistry Chemical Physics, 2014, 16, 14396-14407.   | 1.3  | 41        |
| 16 | Global and local aromaticity of acenes from the information-theoretic approach in density functional reactivity theory. Physical Chemistry Chemical Physics, 2019, 21, 18195-18210.   | 1.3  | 41        |
| 17 | Different Products of the Reduction of (N),C,N $\hat{a}\in\mathbb{C}$ helated Antimony(III) Compounds: Competitive Formation of Monomeric Stibinidenes versus $1< i> H \hat{a}\in\mathbb{C}$ , $1\hat{a}\in\mathbb{B}$ enzazastiboles. Chemistry - A European Journal, 2017, 23, 2340-2349. | 1.7  | 39        |
| 18 | A universal scale of aromaticity for Ï€â€organic compounds. Journal of Computational Chemistry, 2010, 31, 917-928.  | 1.5  | 38        |

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|----|---|-----|-----------|
| 19 | Aromaticity as a Guiding Concept for Spectroscopic Features and Nonlinear Optical Properties of Porphyrinoids. Molecules, 2018, 23, 1333.   | 1.7 | 38        |
| 20 | Simple Alkalineâ€Earth Metal Catalysts for Effective Alkene Hydrogenation. Angewandte Chemie, 2018, 130, 15397-15402.   | 1.6 | 33        |
| 21 | Fingerprint of Aromaticity and Molecular Topology on the Photophysical Properties of Octaphyrins.<br>Journal of Physical Chemistry C, 2019, 123, 7318-7335.   | 1.5 | 32        |
| 22 | Neural Networks as a Tool To Classify Compounds According to Aromaticity Criteria. Chemistry - A European Journal, 2007, 13, 3913-3923.   | 1.7 | 29        |
| 23 | Substituent effects on the aromaticity of carbocyclic five-membered rings. Physical Chemistry Chemical Physics, 2010, 12, 1305-1317.  | 1.3 | 29        |
| 24 | Understanding the molecular switching properties of octaphyrins. Physical Chemistry Chemical Physics, 2016, 18, 11885-11900.  | 1.3 | 29        |
| 25 | Molecular dynamics based descriptors for predicting supramolecular gelation. Chemical Science, 2020, 11, 4226-4238.   | 3.7 | 29        |
| 26 | Conformational Control in [22]- and [24]Pentaphyrins(1.1.1.1.1) by Meso Substituents and their N-Fusion Reaction. Journal of Organic Chemistry, 2013, 78, 4419-4431.  | 1.7 | 25        |
| 27 | Implementing the mechanical force into the conceptual DFT framework: understanding and predicting molecular mechanochemical properties. Physical Chemistry Chemical Physics, 2019, 21, 7378-7388.                               | 1.3 | 25        |
| 28 | Switching between $H\tilde{A}\frac{1}{4}$ ckel and $M\tilde{A}\P$ bius aromaticity: a density functional theory and information-theoretic approach study. Physical Chemistry Chemical Physics, 2020, 22, 4715-4730.             | 1.3 | 25        |
| 29 | Alkene Transfer Hydrogenation with Alkalineâ€Earth Metal Catalysts. Angewandte Chemie, 2019, 131, 4292-4297.  | 1.6 | 24        |
| 30 | Chemical applications of neural networks: aromaticity of pyrimidine derivatives. Physical Chemistry Chemical Physics, 2011, 13, 20564.  | 1.3 | 23        |
| 31 | Hydrosilylation Induced by N→Si Intramolecular Coordination: Spontaneous Transformation of Organosilanes into 1â€Azaâ€Siloleâ€Type Molecules in the Absence of a Catalyst. Chemistry - A European Journal, 2014, 20, 2542-2550. | 1.7 | 23        |
| 32 | Ambident Nucleophilic Substitution: Understanding Nonâ€HSAB Behavior through Activation Strain and Conceptual DFT Analyses. Chemistry - A European Journal, 2020, 26, 3884-3893.  | 1.7 | 23        |
| 33 | Synthesis and Structural Characterization of Heteroboroxines with MB <sub>2</sub> O <sub>3</sub> Core (M = Sb, Bi, Sn). Inorganic Chemistry, 2013, 52, 1424-1431.   | 1.9 | 22        |
| 34 | A computational study on the role of noncovalent interactions in the stability of polymer/graphene nanocomposites. Journal of Molecular Modeling, 2017, 23, 43.   | 0.8 | 22        |
| 35 | Performance of Electronic Structure Methods for the Description of HÃ⅓ckel–Möbius<br>Interconversions in Extended π-Systems. Journal of Physical Chemistry A, 2020, 124, 2380-2397.   | 1.1 | 22        |
| 36 | Studies on aromatic compounds: inhibition of calpain I by biphenyl derivatives and peptide-biphenyl hybrids. Bioorganic and Medicinal Chemistry Letters, 2004, 14, 2753-2757.   | 1.0 | 21        |

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|----|--|-----|-----------|
| 37 | The hunt for reactive alkynes in bio-orthogonal click reactions: insights from mechanochemical and conceptual DFT calculations. Chemical Science, 2020, 11, 1431-1439.   | 3.7 | 21        |
| 38 | Decabromobiphenyl (PBB-209) Activates the Aryl Hydrocarbon Receptor While Decachlorobiphenyl (PCB-209) Is Inactive: Experimental Evidence and Computational Rationalization of the Different Behavior of Some Halogenated Biphenyls. Chemical Research in Toxicology, 2008, 21, 643-658. | 1.7 | 19        |
| 39 | Aromaticity changes along the reaction coordinate connecting the cyclobutadiene dimer to cubane and the benzene dimer to hexaprismane. Structural Chemistry, 2007, 18, 773-783.  | 1.0 | 18        |
| 40 | Synthesis and reactivity of a germylene stabilized by a boraguanidinate ligand. RSC Advances, 2016, 6, 19377-19388.  | 1.7 | 18        |
| 41 | Qualitative Insights into the Transport Properties of Hýckel/Möbius (Anti)Aromatic Compounds: Application to Expanded Porphyrins. Journal of Physical Chemistry C, 2018, 122, 19842-19856.   | 1.5 | 17        |
| 42 | ACTIVATION OF THE ARYL HYDROCARBON RECEPTOR BY CARBARYL: COMPUTATIONAL EVIDENCE OF THE ABILITY OF CARBARYL TO ASSUME A PLANAR CONFORMATION. Environmental Toxicology and Chemistry, 2006, 25, 3141.  | 2.2 | 16        |
| 43 | Spontaneous Double Hydrometallation Induced by N→M Coordination in Organometallic Hydrides of Group 14 Elements. Chemistry - A European Journal, 2016, 22, 5620-5628.  | 1.7 | 16        |
| 44 | The active site architecture in peroxiredoxins: a case study on Mycobacterium tuberculosis AhpE. Chemical Communications, 2016, 52, 10293-10296.   | 2.2 | 16        |
| 45 | Biological and chemical studies on aryl hydrocarbon receptor induction by the p53 inhibitor pifithrin- $\hat{l}^{\pm}$ and its condensation product pifithrin- $\hat{l}^{2}$ . Life Sciences, 2011, 88, 774-783.   | 2.0 | 14        |
| 46 | Metalated Hexaphyrins: From Understanding to Rational Design. Chemistry - A European Journal, 2015, 21, 17631-17638.   | 1.7 | 14        |
| 47 | Mechanochemically Triggered Topology Changes in Expanded Porphyrins. Chemistry - A European<br>Journal, 2021, 27, 3397-3406.   | 1.7 | 14        |
| 48 | Oxidative Additions of Homoleptic Tin(II) Amidinate. Organometallics, 2015, 34, 606-615.   | 1.1 | 13        |
| 49 | Computational Tools to Rationalize and Predict the Self-Assembly Behavior of Supramolecular Gels. Gels, 2021, 7, 87.   | 2.1 | 13        |
| 50 | Comparison of reactivity of <i>C</i> , <i>N</i> -chelated and Lappert's stannylenes with trimethylsilylazide. Canadian Journal of Chemistry, 2014, 92, 434-440.  | 0.6 | 12        |
| 51 | 8-Aryl substituted boron-dipyrromethene dyes: crystal structures and computational studies. Journal of Molecular Structure, 2004, 697, 29-40.  | 1.8 | 11        |
| 52 | Reactivity of Single Transition Metal Atoms on a Hydroxylated Amorphous Silica Surface: A Periodic Conceptual DFT Investigation. Chemistry - A European Journal, 2021, 27, 6050-6063.  | 1.7 | 11        |
| 53 | Studies on calpain inhibitors. Synthesis of partially reduced isoquinoline-1-thione derivatives and conversion to functionalized 1-chloroisoquinolines. Tetrahedron Letters, 2008, 49, 2275-2279.  | 0.7 | 10        |
| 54 | Understanding conductivity in molecular switches: a real space approach in octaphyrins. Physical Chemistry Chemical Physics, 2016, 18, 11829-11838.  | 1.3 | 10        |

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|----|---|-----|-----------|
| 55 | Toward the Design of Bithermoelectric Switches. Journal of Physical Chemistry C, 2018, 122, 24436-24444.  | 1.5 | 10        |
| 56 | Scrutinizing ion- $i \in A$ and ion- $i \in A$ interactions using the noncovalent index and energy decomposition analysis. Computational and Theoretical Chemistry, 2015, 1053, 150-164.  | 1.1 | 9         |
| 57 | Rationalising Supramolecular Hydrogelation of Bisâ€Ureaâ€Based Gelators through a Multiscale<br>Approach. ChemPlusChem, 2020, 85, 267-276.  | 1.3 | 9         |
| 58 | Single-molecule conductance in a unique cross-conjugated tetra(aminoaryl)ethene. Chemical Communications, 2021, 57, 591-594.  | 2.2 | 9         |
| 59 | Reactivity of Bis(organoamino)phosphanes with Aluminum(III) Compounds: Straightforward Access to Diiminophosphinates by Means of Hydrogen-Atom Migration - An Experimental and Theoretical Study. European Journal of Inorganic Chemistry, 2014, 2014, 5193-5203.       | 1.0 | 8         |
| 60 | Reactivity of Tin(II) Guanidinate with 1,2- and 1,3-Diones: Oxidative Cycloaddition or Ligand Substitution ?. Organometallics, 2015, 34, 2202-2211.   | 1.1 | 8         |
| 61 | Derivatives of 3-sec-Butyl-1-oxo-2,3-dihydroisoquinoline as Inhibitors of $\hat{l}$ 4-Calpain. ChemMedChem, 2006, 1, 710-714.   | 1.6 | 7         |
| 62 | A Benchmark of Density Functional Approximations For Thermochemistry and Kinetics of Hydride Reductions of Cyclohexanones. ChemistryOpen, 2019, 8, 788-806.   | 0.9 | 7         |
| 63 | SuFEx-enabled, chemoselective synthesis of triflates, triflamides and triflimidates. Chemical Science, 2022, 13, 2270-2279.   | 3.7 | 7         |
| 64 | X-ray Diffraction, Solution Structure, and Computational Studies on Derivatives of (3- <i>&gt;sec</i> -Butyl-2,3-dihydro-1 <i>H</i> -isoquinolin-4-ylidene)acetic Acid: Compounds with Activity as Calpain Inhibitors. Journal of Organic Chemistry, 2010, 75, 342-352. | 1.7 | 5         |
| 65 | Reactivity of bis(organoamino)phosphanes with magnesium( <scp>ii</scp> ) compounds. Dalton Transactions, 2015, 44, 4533-4545.   | 1.6 | 5         |
| 66 | Designing Force Probes Based on Reversible 6 ∈-Electrocyclizations in Polyenes Using Quantum Chemical Calculations. Journal of Organic Chemistry, 2021, 86, 7477-7489.  | 1.7 | 5         |
| 67 | Stereoselective Reductions of 3-Substituted Cyclobutanones: A Comparison between Experiment and Theory. Journal of Organic Chemistry, 2020, 85, 7803-7816.  | 1.7 | 5         |
| 68 | Fine-Tuning of Nonlinear Optical Contrasts of Hexaphyrin-Based Molecular Switches Using Inverse Design. Frontiers in Chemistry, 2021, 9, 786036.  | 1.8 | 5         |
| 69 | Understanding the Fundamental Role of $   \in    \in    \in    f    = 0$ Dispersion Interactions in Shaping Carbon-Based Materials. Chemistry - A European Journal, 2014, 20, 4845-4845.  | 1.7 | 3         |
| 70 | Towards the understanding of halogenation in peptide hydrogels: a quantum chemical approach. Materials Advances, 2021, 2, 4792-4803.  | 2.6 | 3         |
| 71 | Uncovering the Activity of Alkaline Earth Metal Hydrogenation Catalysis Through Molecular Volcano Plots. Topics in Catalysis, 2022, 65, 289-295.  | 1.3 | 3         |
| 72 | Towards the Design of Optically Active Thiophene Sâ€Oxides using Quantum Chemistry. Chemistry - A European Journal, 2019, 25, 2840-2851.  | 1.7 | 2         |

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| 73 | Rationalising Supramolecular Hydrogelation of Bisâ€Ureaâ€Based Gelators through a Multiscale Approach. ChemPlusChem, 2020, 85, 266-266.  | 1.3 | 1         |
| 74 | Synthesis and Reactivity of Spirocarbocycles as Scaffolds for Nucleoside Analogues. Journal of Organic Chemistry, 2020, 85, 14989-15005. | 1.7 | 1         |
| 75 | Quantifying aromaticity according to the energetic criterion. , 2021, , 195-235.   |     | 1         |
| 76 | Aryl hydrocarbon receptor induction by alpha- and ss-pifithrin. Toxicology Letters, 2010, 196, S258.                                     | 0.4 | 0         |