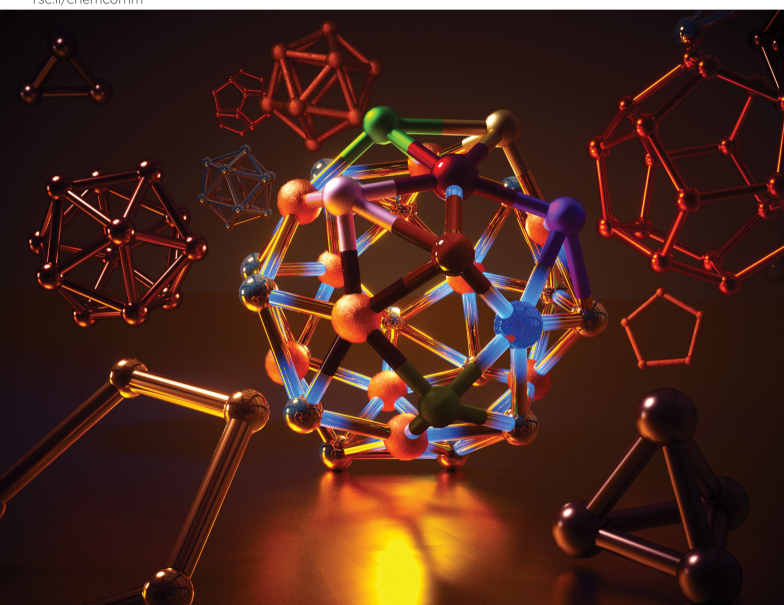
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Aromaticity of all-metal clusters

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The first all-metal aromatic cluster, Al_4^{2-} , face-capped by an M⁺ cation (M = Li, Na, Cu), was detected in 2001. Since then, several all-metal aromatic clusters have been extensively studied. Here, we review the most significant developments that have occurred mainly in our labs, including topics ranging from the difficulty of measuring aromaticity in metal clusters to the use of current techniques to investigate aromaticity in both small and large metal clusters.

1. Introduction

This year we celebrate the 200th anniversary of the discovery of benzene by Michael Faraday. This molecule has become a reference molecule that is used when discussing aromaticity with experiments and theoretical methods. Comparison with benzene is reasonable for classical aromatic organic molecules such as polycyclic aromatic hydrocarbons (PAHs),2 heteroaromatic organic compounds, 3,4 and nanographenes. 5 Such a

comparison can also be performed when a C-H group in these compounds is substituted by an isolobal metal fragment leading to metallobenzenes, or heteroatom-containing analogues such as metallapyridines or metallapentalenes. 6,7 However, it is not applicable to all-metal clusters, which are aggregates of only metal atoms that exhibit aromatic characteristics. The aromaticity of all-metal aromatic compounds containing σ -, π -, δ - and ϕ -electron delocalization is far more complex than that of typical aromatic organic molecules, which possess π electron delocalization only. Furthermore, these compounds have the ability to blend several forms of (anti)aromaticity, resulting in double or triple aromaticity—a phenomenon known as multifold (anti)aromaticity.8-10 They can also display aromaticity in one (or several) component(s) $(\sigma, \pi, \delta \text{ or } \phi)$ and antiaromaticity in other(s), giving rise to the conflicting

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aromaticity concept. 11,12 One can classify aromatic all-metal clusters based on the type of delocalized electrons (σ -, π -, δ -, ϕ -, and multifold) or take into account if they are relatively simple and planar or three-dimensional (3D). Here, we have preferred the latter classification.

In 2001, Li et al. 13 experimentally observed the $\mathrm{Al_4}^{2-}$ allmetal cluster in the form of the bimetallic clusters LiAl₄-, NaAl₄⁻, and CuAl₄⁻, which in their most stable isomeric form contains a square-planar ${\rm Al_4}^{2-}$ dianion. These clusters were obtained using a laser vaporization technique and characterized using their experimental and theoretical photoelectron spectra. To rationalize chemical bonding in this cluster the authors looked at the chemical bonding in Al₄²⁻, as they assumed that bonding between countercations (Li⁺, Na⁺, or Cu^+) and Al_4^{2-} is primarily ionic. For Al_4^{2-} , the total number of valence electrons is $3 \times 4 + 2 = 14$. As can be seen in Fig. 1, the four 3s atomic orbitals (AOs) of the Al atoms combine to give the σ_S system with four molecular orbitals (MOs): one bonding without nodes, two degenerate non-bonding with one node, and one antibonding with two nodes. These orbitals are completely filled with eight electrons. Same combinations of AOs take place with the set of four $3p_x$, $3p_y$, and $3p_z$ AOs generating the σ -radial (σ_r), σ -tangential (σ_t), and π systems, respectively, each of them occupied with two-electrons. 14 Therefore, with two electrons, the π -system follows the 4N + 2 Hückel rule^{15–18} and is considered aromatic. Although this is not the case for the σ -system with four electrons, the two pairs of delocalized σ electrons belong to MOs that follow orthogonal radial (σ_R) orbital $2a_{1g}$) and tangential (σ_T , orbital $1b_{2g}$) directions, which makes them independent, 19 thus separately following the 4N + 2 rule. Therefore, according to its electronic structure, Al₄²⁻ can be considered a three-fold aromatic system (σ_T , σ_R , and π). Thus, the Al₄²⁻ all-metal cluster is an example of multifold aromaticity, with the π -MOs resembling those of organic aromatic species, and the σ_r -MOs being similar to those found in σ -aromatic species such as $H_3^{+,20}$ Calculations of multicenter indices (MCI), nucleus-independent chemical shifts (NICS), resonance energies, and ring currents support the aromaticity of Al₄²⁻. ^{19,21-28} The discovery of all-metal aromaticity in Al₄²⁻ meant a complete revolution in the field, with many other allmetal aromatic clusters being discovered afterwards. 8,29-32

Al₄²⁻ is the paradigmatic example of all-metal aromatic clusters. From the beginning, aromaticity has been associated

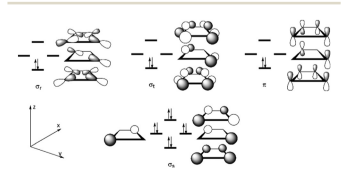


Fig. 1 Valence molecular orbitals of the Al_4^{2-} all-metal cluster. For the σ_{T^-} , σ_{R^-} , and π -MOs, only one of two degenerate MOs are displayed.

with kinetically and thermodynamically stable compounds. By connecting the concept of aromaticity with stability, Hoffmann concluded that all-metal clusters such as ${\rm Al_4}^{2-}$ cannot be classified as aromatic because "aromatic molecules are 'bottleable". 33 This is an interesting controversy. Only stable compounds can be aromatic? Several examples show that this is not the case. For instance, transition states of allowed pericyclic reactions are obviously nonbottleable species, but their aromaticity, which is widely accepted by the chemical community, is responsible, in part, for the relatively low barriers observed for this type of reaction.34-40 As another example, certain excited states are aromatic and nonbottleable compounds, like those following Baird's rule. 41-43 This rule states that the lowest-lying triplet states of $4N \pi$ -electrons, such as the T_1 state of cyclooctatetraene, are aromatic. We do not think that the 'bottleable' criterion should be the one that prevails when classifying compounds as aromatic. In our opinion, following the definition by Chen et al., 44 species in their ground or excited state showing cyclic π -electron delocalization that results in energy lowering and a tendency toward bond length equalization, particular magnetic behaviour, and characteristic spectroscopic features have to be classified as aromatic irrespective of whether they are 'bottleable'.

2. Quantifying aromaticity in all-metal clusters

From a theoretical perspective, assessing the aromaticity of allmetal aromatic compounds is necessary for a thorough investigation. The measurement of aromaticity in all-metal clusters is considerably more difficult than in classical organic compounds due to the presence of multifold and conflicting aromaticities and the lack of all-metal aromatic clusters that can act as inorganic reference systems (like benzene does in classical aromatic organic molecules). In fact, the majority of the methods for measuring aromaticity that are currently available were designed to assess the aromaticity of organic molecules using benzene or other aromatic organic compounds as a reference for their definitions. For example, the electronicbased descriptors like the bond order index of aromaticity (BOIA), 45 the aromatic fluctuation (FLU) index, 46,47 the aromaticity descriptor $\theta^{48,49}$ or the structure-based harmonic oscillator model of aromaticity (HOMA)50,51 are examples of this type of aromaticity indicator. Similarly, the absence of suitable reference systems21,52 makes it challenging to compute energy-based markers like resonance energy (RE) or aromatic stabilization energy (ASE)53 in all-metal clusters. In general, reference-system-based indices of aromaticity, without additional improvements, cannot be used directly to assess the aromaticity of all-metal clusters. Moreover, most of these reference-based methods do not allow for a study of the different types of aromaticity in species having multifold aromaticity, which represents another important disadvantage of these methods.

At this moment, there are three main groups of methods used to assess all-metal aromaticity. First, the magnetic-based indicators of aromaticity, which are based on the ring currents ChemComm Feature Article

that are generated when applying external magnetic fields to these compounds. Ring currents are not observable, but their effects are (in NMR spectra, for instance). The calculated ring currents are origin dependent. Several attempts have been performed to circumvent the gauge origin problem. The gaugeincluding magnetically induced currents (GIMIC)⁵⁴ and the continuous transformation of the origin of the current density (CTOCD)^{55,56} methods are among the most widely used. To get additional information, one can depict these ring currents on the surface of the so-called anisotropy of the magnetically induced current density tensor (ACID). 57,58 Among the magnetic descriptors, one of the most widely used is the nucleusindependent chemical shift (NICS)59 and its different variants such as NICS(0)zz and NICS(1)zz (the out-of-plane tensor component) or NICS(0)_{πzz} and NICS(1)_{πzz} (the π -orbital contribution),⁴⁴ as well as the NICS-scans analysis, 60-62 the NICS decomposition into canonical molecular orbital contributions (CMO-NICS),63 and the different representations of the NICS values. 64-66 Among the NICS's many benefits, one finds its ease of use and accessibility and the clear separation that provides among aromatic (negative NICS values), non-aromatic (values close to zero), and antiaromatic systems (positive values), and the possibility of separation into σ -, π -, and δ -contributions. 60 It is worth noting that NICS and ring currents can be affected by the relativistic corrections when molecules that contain heavy elements are studied.⁶⁷ Moreover, the core electrons in heavy atoms can also generate NICS and ring currents that result in spurious aromaticity assessments.68

The second group corresponds to the methods that measure the cyclic delocalization of mobile electrons in closed circuits of two or three dimensions.⁶⁹ Cyclic delocalization is one of the basic and crucial characteristics of aromatic compounds. Since this electronic delocalization is not observable, there is no experimental property that permits its direct measurement. Because of this, there is no single, widely accepted computational method to measure it. Among the methods that provide measures of electronic delocalization⁷⁰ without making use of reference values, we can mention the multicentre delocalization index in a ring (I_{ring}) , 71 which is a generalization of the delocalization index between two atoms, 72-74 the multicentre index (MCI), 45 an extension of the I_{ring} index, the electron localization function (ELF)⁷⁵ using the bifurcation values as a measure of aromaticity, 76,77 the localized orbital locator (LOL) 88-80 and its variant, LOL- π , which can be used to investigate π -electron delocalization,81 and the electron density of delocalized bonds (EDDB)82 that yields the number of electrons delocalized in a closed circuit as an indicator of aromaticity.83 The advantages and drawbacks of all these methods have been discussed recently.⁵ In particular, both the separation into α and β components and σ -, π -, and δ -contributions are possible.

The third group refers to the indicators of aromaticity that analyse the molecular structure. They are based on the fact that, in aromatic systems, there is usually an observed equalization of bond lengths. Then, one can use the degree of bond length alternation (BLA) as a quantitative measure of aromatic character in molecules.84 This descriptor can be used only in rings

made with a single metal, and, for this reason, is less used for the analysis of all-metal clusters than those of the two previous groups mentioned.

For the correct assessment of the aromaticity of all-metal clusters, we recommend the use of at least one indicator from each of the two former groups mentioned above. Moreover, together with the calculation of magnetic and electronic descriptors to verify electron delocalization, it is advisable to contextualize aromaticity in terms of the fulfilment of electron counting rules such as the 4N + 2 Hückel's rule, $^{15-18}$ the 4N Baird's rule for the lowest-lying triplet excited states, 85 the $2(N+1)^2$ Hirsch rule of spherical aromaticity, 86 etc. using either valence canonical MOs or, even better, MOs obtained from an adaptive natural density partitioning (AdNDP) analysis⁸⁷ or other localization schemes.

Simple all-metal aromatic species

Since the landmark realization of all-metal aromaticity in 2001, this field triggered considerable research studies driven by the extension of the concept of aromaticity offering unprecedented cases and unexpected structures setting solid bases for the current understanding of this transversal concept as summarized in earlier reviews and books. 8,31,88 Interestingly, their appearance is given even in biological relevant macromolecules as reported for the human copper chaperone Atox1.89 When silver ions were mixed with copper chaperone Atox1, the resulting σ-aromatic [Ag₄]²⁺ core was found to attenuate cancer cell proliferation.⁹⁰

Regarding the smallest all-metal aromatic ring structure provided by M_3 , some earlier examples are given by the cyclo- $[Mg_3]^{2-}$ cluster that can be stabilized with alkaline and alkaline-earth cations. The cyclo-[Mg₃]²⁻ cluster has a triplet ground state with the singlet closed-shell state being almost degenerate. 91-93 The singlet closed-shell cyclo- $[Mg_3]^{2-}$ cluster is σ -aromatic and undergoes a dramatic aromaticity change to π-aromaticity when interacting with cations to form X_nMg_3 (n = 1, 2; $X = Li^+$, Na^+ , K^+ , Be^{2+} , Mg^{2+} , Ca^{2+}). Interestingly, the aromaticity of X_nMg_3 species can be tuned by modifying the X-Mg distance. 91 Another example of σ aromaticity confirmed by NICS and MCI (Table 1) calculations is given by the Cu₃⁺ species.^{27,94} Multifold aromaticity in M₃ species is found in the d-orbital $\sigma + \pi$ double aromaticity of valence isoelectronic Y₃⁻ and La₃⁻ clusters as confirmed by NICS, MCI, and resonance energies, 27,95 whereas Hf₃ shows three-fold σ -, π -, and δ-aromaticity according to both molecular orbital analysis and MCI results.27 Even more interesting is the multifold aromaticity in the open-shell molecule ⁵Ta₃⁻. The quintuplet ⁵Ta₃⁻ is the lowest-lying spin state of Ta_3^- , and it exhibits three-fold σ -, π -, and δ-aromaticity based on MCI findings and molecular orbital analysis. Its σ - and π -aromaticities are of the open-shell Baird-type, whereas its δ -aromaticity is of the Hückel-type (a pair of closedshell electrons).²⁷ On the other hand, the [Ga₃(Mes₂C₆H₃)₃]²⁻ complex⁹⁶ displays an homometallic equilateral [Ga₃]⁺ triangle with Ga-Ga distances of 2.441 Å. The metallic core exhibits a populated delocalized π -orbital fulfilling the Hückel rule, the overall structure being described as a metalloaromatic from related computational models.97 Similarly, the isoelectronic and

Table 1 MCI, MCI $_{\pi}$, MCI $_{\sigma}$, and MCI $_{\delta}$ indices for Cu $_{3}^{+}$, Y $_{3}^{-}$, La $_{3}^{-}$, Hf $_{3}$, and ${}^{5}\text{Ta}_{3}^{-}$ at the B3LYP/X/Stuttgart + 2f (X = Cu, Y, and La) level of theory. All MCI values in electrons and bond distances in Å. All molecules are in their

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ground states. Values taken from ref. 27

| | Cu ₃ ⁺ | Y ₃ - 3.116 | La ₃ | Hf ₃ | ⁵ Ta ₃ - |
|----------------|------------------------------|------------------------|-----------------|-------------------------|--------------------------------|
| MCI | 0.189 | 0.754 | 0.750 | 1.037 | 0.776 |
| MCI_{σ} | 0.188 | 0.458 | 0.454 | 0.445 | 0.362 |
| MCI_{π} | 0.001 | 0.296 | 0.296 | 0.296 | 0.178 |
| MCI_{δ} | 0.000 | 0.000 | 0.000 | 0.295 | 0.235 |
| Aromaticity | σ | $\sigma + \pi$ | $\sigma + \pi$ | $\sigma + \pi + \delta$ | $\sigma + \pi + \delta$ |

isostructural aluminium counterpart [Al₃(Mes₂C₆H₃)₃]²⁻ reported by Power et al. 98 retains the aromatic characteristics in the $[Al_3]^+$ core. In addition, the boron-dinitrogen and boron-carbonyl cations generated in the gas phase, exhibit cyclic structures of the form $[B_3(NN)_3]^+$ and $[B_3(CO)_3]^+$ featuring a related $[B_3]^+$ core which similarly leads to two π -electrons, as the smallest π -aromatic kernel.⁹⁹ Such examples expose that for these species, it is plausible to shift from boron to gallium retaining similar aromatic characteristics and keeping the same aromatic motif decorated with different supporting ligands. A recent review on triangular all-metal aromatic cores has been provided. 100

With respect to four membered metalloaromatic rings, Al₄²⁻ is the archetypal all-metal aromatic cluster. It has a total MCI value of 0.356 a.u. computed at the B3LYP/6-311+G(d) level of theory.²⁷ This value can be decomposed into a MCI_{\sigma} value of 0.169 a.u. and a MCI_{π} value of 0.187 a.u. indicating that the π -delocalization is slightly larger than the σ -delocalization, in agreement with NICS and ELF analyses.76,101

Among the different distortions that a benzene ring can suffer, the Kekulean or bond length alternation (BLA) distortion of the in-plane b_{2u} symmetry that changes the D_{6h} symmetry of benzene into a Kekulé-like D_{3h} symmetry structure is particularly interesting because it was found to be favoured by the π -electrons. ^{84,102–104} In the case of Al₄^{2–}, the π -electrons are also distortive, but their preference for a localized D_{2h} structure is weaker than in benzene. As for benzene, the σ -electrons enforce the regular D_{4h} equilibrium geometry with delocalized electron pairs. 105

Ge₄²⁺ is an all-metal cluster that is valence isoelectronic with ${\rm Al_4}^2-$. In fact, it has similar MCI (MCI = 0.386, MCI $_{\sigma}$ = 0.187, $MCI_{\pi} = 0.199 \text{ a.u.}$) and NICS values. ¹⁰¹ By performing successive substitution of Al atoms by more electronegative Ge atoms in Al_4^{2-} , we expect the following trend of aromaticity: Al_4^{-2-} > $Al_3Ge^- \ge Al_2Ge_2 \le AlGe_3^+ < Ge_4^{2+}$. The same trend is expected for all the series of four-membered ring valence isoelectronic clusters $[X_nY_{4-n}]^{q\pm}$ (X, Y = Al, Ga, Si, and Ge; n=0-4). This series was used to evaluate the reliability of NICS and MCI to provide the correct trends in all-metal and semimetal aromatic clusters. 101 It was found that MCI was superior to NICS in reproducing the expected trends in aromaticity. Among the different NICS-based indicators, the NICS(0)_{π} was the one that performed the best.

The four-membered cyclic systems $M_2A_2^{2-}$ (M and A = B, Al, and Ga) are also valence isoelectronic with $\mathrm{Al_4}^{2-}$. For the cases $M \neq A$, the clusters can adopt $cis(C_{2v})$ and $trans(D_{2h})$ configurations. With the induced magnetic field and MCI calculations, the double $\sigma + \pi$ aromatic character of these rings was confirmed. ¹⁰⁶ In general, between C_{2v} and D_{2h} structures, the most aromatic ring is also the most stable, except for $Al_2B_2^{\ 2-}$ and $Ga_2B_2^{\ 2-}$, for which the strong B-B bond present in their C_{2v} structures has an important stabilization role. The $M_2A_2^{2+}$ (M and A = C, Si, and Ge) species are also valence isoelectronic with the $M_2A_2^{2-}$ (M and A = B, Al, and Ga) clusters and they also show σ - and π -aromaticity. However, contrary to what was found for group 13 M₂A₂²⁻ clusters, the linear isomer of group 14 M2A22+ clusters is the most stable for two of the clusters (C₂Si₂²⁺ and C₂Ge₂²⁺) and it is isoenergetic with the cyclic D_{4h} isomer in the case of C_4^{2+} (see Fig. 2).107

The first all-metal with π -antiaromaticity was Li_3Al_4^- , which contains a planar and rectangular Al₄⁴⁻ unit. 11 Interestingly, the singlet (S_0) and the lowest-lying triplet (T_1) states of Al_4^{4-} are almost degenerate, with the singlet state being more stable by just 1.6 kcal mol⁻¹ at the (U)B3LYP/6-311+G(d) level of theory. ¹⁰⁸ In S_0 , the two electrons added to ${\rm Al_4}^{2-}$ to get ${\rm Al_4}^{4-}$ goes to one of the degenerate π -MOs of Al₄²⁻ (Fig. 1). With these two electrons, the π -system with four electrons becomes antiaromatic, whereas the σ -system keeps its aromaticity. Therefore, Al_4^{4-} in its closedshell singlet state has conflicting aromaticity since it has an aromatic σ -component and an antiaromatic π -component. In the

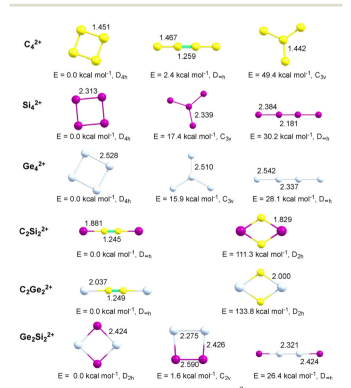


Fig. 2 Optimized geometries of most stable M₂A₂²⁺ clusters of group 14 computed using the PBE/TZ2P method. Bond lengths in Å and CCSD(T)/6-311G*//PBE/TZ2P relative energies in kcal mol⁻¹. Reprinted with permission from ref. 107. Copyright Royal Society of Chemistry, 2016.

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 T_1 state, the two unpaired electrons of Al_4^{4-} occupy σ_r degenerate MOs (Fig. 1) in a D_{4h} symmetric structure. In this state, the cluster is Baird σ_r -aromatic and Hückel σ_t - and π -aromatic. The extra

stabilization of the T₁ state comes from this Baird aromaticity. ¹⁰⁸ A similar situation is found for the T_1 states of Be_2B_6 and $Be_2B_7^+$, which are Baird σ -aromatic and Hückel π -aromatic. However, in these molecules, the lowest-lying triplet is clearly the ground state. 108 In contrast, Be₂(a)Be₆H₆, which has also a triplet ground state, has Hückel σ -aromaticity and Baird π -aromaticity. ¹⁰⁹

Finally, a σ-aromatic neutral rhombic Al₂Pd₂ cluster has been recently reported. Its aromaticity has been confirmed by AdNDP, NICS, ACID, ELF, EDDB, GIMIC and MO analysis. The system presents a 4c-2e σ -bond that provides the σ -aromaticity. ¹¹⁰

4. All-metal clusters with spherical aromaticity

Spherical all-metal clusters expose the delicate and challenging balance between both electronic shells and structural features, as a result of the quantum confinement of electrons, 111-114 which are highly desired as building blocks in molecularly conceived materials. Since the discovery of buckminsterfullerene (C₆₀) and related species, ^{115,116} an increasing interest has been focused on the use of stable three-dimensional cages with particular properties, leading to them being widely explored, where fullerenes have been proved to be highly versatile building blocks in nanotechnology. 117,118 The unique spherical structure of fullerenes, along with the surface full of π -electrons, possesses many fascinating properties and generates a wide range of applications in biology, medicine, and electronics. 119-121 Moreover, the internal cavity of fullerenes provides a suitable region for hosting atoms and molecules, giving rise to a whole new class of endohedral clusters, named endofullerenes. 122-124 Despite the structural stability, the 60 π -electrons in the curved C_{60} initially ascribed as a prototypical spherical aromatic molecule, 115,125-127 shifted to a more appropriate non-aromatic character¹²⁸ owing to the local aromatic and antiaromatic character displayed by hexagons and pentagons, respectively. Moreover, it is endowed with spherical aromatic character after reduction, as highlighted for the hexaanion C_{60}^{6-} formed in alkali-metal fullerene phases. 129-132

The rapid progress in fullerene-related clusters and the extensive applications for fullerene-based materials encourage the promising exploration of analogous hollow spheres composed of main-group or transition metal elements known as inorganic fullerenes. One of the earliest proposals was provided by Johansson and coworkers, who, following the Hirsch rule of spherical aromaticity $2(N+1)^2$ with N=3, were able to predict and locate the existence of an all-metal aromatic fullerene counterpart given by Au₃₂, ¹³³ featuring a sizable HOMO-LUMO gap in a symmetrical icosahedral structure. The reported central NICS value inside the cage is -100 ppm, 133 largely increased in comparison to that for C_{60} at the same level of theory (-2 ppm), supporting the spherical aromatic characteristic of this golden fullerene proposal, where the icosahedral structure is favoured as denoted in earlier works and

in the characterization from laser vaporization of a gold foil by Wang. 134,135 Using the same approach, Pyykkö and coworkers 136 were able to account for larger related species, highlighting the chiral structure of the Au₇₂ golden fullerene satisfying the Hirsch rule $2(N+1)^2$ with N=5, featuring 72-cluster electrons, giving rise to a spherical aromatic hollow cluster with a central NICS value of -111 ppm. For Au₅₀, satisfying the $2(N+1)^2$ rule with N=4, also depicts a central shielding NICS value of -88.5 ppm, where in contrast, the also spherical structure of Au42 shows antiaromatic character as seen from the NICS value of 125 ppm.

One of the guiding principles for identifying systems with spherical aromaticity is the $2(N+1)^2$ Hirsch rule. An alternative to this rule is to look for whether the system under consideration (usually a molecular cluster) has, for the valence electrons, an electronic structure with a closed-shell configuration on the basis of the jellium model. 137-139 The energy levels of the valence electrons for such a model are $1S^21P^61D^{10}2S^21F^{14}2P^61G^{18}2D^{10}3S^2...$, where S, P, D, F, and G letters denote the angular momentum and numbers 1, 2, and 3 indicate the radial nodes. The abundance of alkali metals, alkaline earth metals, and gold clusters bearing 2, 8, 18, 20, 34, 40, 58, 68, 70, 92,... electrons found in experimental mass spectra are justified by taking into account the fact that these numbers of electrons reach closed-shell electronic structures in the jellium model. 140,141 An extension of the Baird's rule for the jellium model was also proposed in 2019.142 Clusters whose last energy level of valence electrons is half-filled with same-spin electrons in the jellium electronic structure are aromatic and present extra stability compared to those that do not have this electronic structure. This situation is reached for the magic numbers of valence electrons of 1 (S = 1/2), 5 (S = 3/2), 13 (S = 5/2), 19 (S = 1/2), 27 (S = 7/2), 37 (S = 3/2), 49 (S = 9/2),... Na₁₉ with S =1/2 or Be₁₃ clusters with S = 7/2 are examples of this open-shell jellium stability. Interestingly, some metal clusters follow more than a single rule. For instance, Li_6^+ (S = 3/2) follows both the $2N^2 +$ 2N + 1 with S = N + 1/2 (N = 1) for open-shell spherical aromaticity and the open-shell jellium rules¹⁴³ (vide infra).

The quest for all-metal fullerene counterparts is promising, encouraging their synthesis from wet methods to further explore their novel chemistries. In this respect, it is worth mentioning the crystallization of the [K@Au₁₂Sb₂₀]⁵⁻ anion obtained after reaction of the Zintl phase K₈SnSb₄ with Au(PPh₃)Me in the ethylenediamine solution by Sun and coworkers. 144 The resulting structure reveals a highly symmetrical cluster featuring a dodecahedral Sb₂₀ cage with each five-membered face decorated with an Au atom, sustaining a backbone composed of Au-Sb bonds, where a K⁺ cation is allocated inside the cluster. Interestingly, the AdNDP analysis reveals twenty 1c-2e lone pairs at each Sb atom, thirty 4c-2e σ -bonds lying at the cluster surface, in addition to the sixty Au-5d lone pairs. The remaining 18 electrons inside the cage (cluster electrons) give rise to nine occupied orbitals featuring a superatomic shell structure given by the 1S21P61D10 configuration, which fulfils both the jellium closed-shell structure and the $2(N+1)^2$ rule with N=2, satisfying the spherical aromatic requirements contributing to the overall stabilization of the cluster, as pointed out from the electronic criteria of aromaticity. To further evaluate the spherical aromatic in the [K@Au₁₂Sb₂₀]⁵⁻

a) NICS B_z^{ind} B_x^{ind} B_y^{ind} b) NICS B_z^{ind} 15.0Å 7.5 0.0

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Fig. 3 Magnetic response properties of $[K@Au_{12}Sb_{20}]^{5-}$ from NICS isosurfaces and under specific orientations of the external field accounting for $B_z^{\rm ind}$, $g_x^{\rm ind}$, and $g_y^{\rm ind}$ isosurfaces, denoting both isosurface (a) and contour plot (b) representations. Isosurfaces are set at ± 3.0 ppm, blue – shielding; red – deshielding.

0.0

-10.0 -5.0

5.0

anion, the NICS isosurface was calculated (Fig. 3), leading to a spherical-like shielding surface ascribed to the cluster backbone, which is common in spherical aromatic fullerenes, ¹⁴⁵ resulting in a common characteristic in both organic and all-metal spherical aromatic fullerenes. In addition, under specific orientations of the external field, long-range shielding cone characteristics are enabled as obtained from representative orientations aligned with the z-, x-, and y-axis, which shows values of -20.0 ppm at 7.5 Å from the centre of the structure, and of -2.6 ppm at 15.0 Å. Such shielding cone characteristics enabled from any orientation of the external field are inherent to spherical aromatic species, ¹⁴⁶ as an exclusive aspect of spherical aromatic species which is not obtained in planar aromatic species such as benzene. ^{145,147,148}

Thus, finding persistent structures leading to suitable building blocks may be guided by both electronic and magnetic criteria of aromaticity. The resulting chemical and physical characteristics of the designed clusters can be explored further. For instance, Mondal and Chattaraj denote aromatic clusters as interesting species that can be used in hydrogen storage applications. ¹⁴⁹

5. Octahedral aromaticity in all-metal clusters

Systems described as spherical aromatics have either a closed-shell structure (*e.g.* Au₃₂) or a half-filled shell with same spin structure (*e.g.* Be₁₃⁻ with S = 7/2). This recipe was applied in the quest of all-metal clusters with octahedral ($O_{\rm h}$) aromaticity. To this end, we analysed all octahedral clusters of the type X_6^q with X = Li-C and Be–Si and q = -2 to +4 in $^{2S+1}\text{A}_{1g}$ electronic states (S = 0 to 3). ¹⁵⁰ A typical octahedral cluster with a closed-shell electronic structure has the MO energy levels shown in Fig. 4. This is the situation of, for instance, Si₆²⁻¹⁵¹ With this MO

structure, the series 2, 8, 12, 14, 20, 26, 32, and so on could be suggested as the magic numbers that result in closed-shell octahedral aromatic species based on these MOs. The magic numbers for open-shell clusters could be 1, 5, 10, 13, 17, 23, and 29. Unfortunately, because the energy order of the MOs in Fig. 4 varies according to the X atoms, as well as the multiplicity and charge of the $O_h^{2S+1}A_{1g}X_6^q$ clusters, these series of magic numbers cannot be generalized. For the clusters of the second period (X = Li, Be, and B), the $2a_{1g}$ and $1t_{2g}$ MOs invariably become more stable than the $1e_{\rm g}$ ones. Furthermore, the $2a_{\rm 1g}$ are more stable than the $1t_{2g}$ in some clusters, and vice versa in others. In general, the ordering of the various orbitals varies from cluster to cluster, and the energy difference between 1eg, the radial $2a_{1g}$, and the tangential $1t_{2g}$ is minor. This leads to the conclusion that it is not possible to derive a general rule for octahedral aromaticity similar to those of spherical aromaticity.

Calculations showed that, in general, $O_{\rm h}$ systems with closed-shells or open-shells half-filled with same spin electron systems have large multicentre indices and negative NICS values, as expected for aromatic compounds. These results confirm the existence of octahedral aromaticity in all-metal clusters like those found previously in Be $_{\rm 6}$ in its quintet state ¹⁵² and the singlet ${\rm Au_6}^{2-}$ and ${\rm Al_6}^{2-}$ clusters. ^{114,153}

Interestingly, despite B and Al belonging to the same group 13 elements, ${\rm Al_6}^{2-}$ favours the $O_{\rm h}$ structure, whereas the ${\rm B_6}^{2-}$ cluster prefers the planar $D_{\rm 2h}$ geometry. Energy decomposition analysis based on the turn-upside-down approach concluded that is the orbital interaction term, which combined with the electrostatic component do $({\rm Al_6}^{2-})$ or do not $({\rm B_6}^{2-})$ compensate the higher Pauli repulsion of the $O_{\rm h}$ form. These findings are consistent with the tendency for more localized bonding in

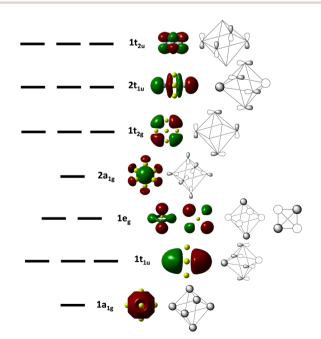


Fig. 4 Schematic representation of the molecular orbitals in an octahedral cluster. Reprinted with permission from ref. 150. Copyright Royal Society of Chemistry, 2016.

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B metal clusters and a dominant delocalization force in Al clusters. Similar to B_6^{2-} , mixed clusters $Al_xB_y^{2-}$ (x + y = 6) with y > 2 favour the planar structure.

A special example of octahedral aromaticity is given by ⁴A_{1g} Li₆⁺ and ⁵A_{1g} Be₆ all-metal clusters. ¹⁴³ These two species have a large number of non-nuclear attractors (NNAs), 156 with all or almost all valence electrons located in these NNAs. The chemical bonding arrangement of these systems is reminiscent of solid metals, where metal cations are encircled by a "sea" of delocalized valence electrons. This is because these NNAs exhibit extremely delocalized electron densities. These new types of compounds were named metal cluster electrides, 143 to differentiate them from the molecular electrides, 157 which have an electron (or a high portion of an electron) that cannot be assigned to any nucleus of the molecule and is located in a NNA.

6. Zintl-ion clusters

Zintl-ion chemistry 158-161 extends the rich structural and composition diversity of all-metal clusters, constantly challenging our notion of chemical bonding¹⁵⁹⁻¹⁶⁷ and related properties, featuring electron-deficient cages and multicentre bonding elements. 167-169 The relation between electron counts and structural characteristics has been classically understood in terms of the Wade-Mingos rules, 170-172 which correlate the number of vertices (n) and cluster electrons (ce) in such species. Deltahedral clusters displaying a -2 charge form electronprecise species fulfilling the Wade's 2n + 2 skeleton electron count for closo-species, and 2n lone pairs of electrons, according to the vertex number, n, as denoted for representative Zintlion bare clusters. 168,173-177

In 2000, using the proposal of the $2(N+1)^2$ Hirsch rule, 86,178 the spherical aromaticity of the representative E_4^{4-} and $E_9^{2-/4-}$ (E = Si, Ge, Sn, Pb) clusters, ¹⁷⁹ was rationalized in terms of separating between $2(N+1)^2$ σ - and $2(N+1)^2$ π -electron kernels, involving both tangential and radial orbitals, respectively. In this sense, the notion of spherical aromaticity in Zintl ions favours the recognition of such a property in a variety of threedimensional structures of different shapes, composition, and bonding characteristics, 167 favouring the extension of this concept to all-metal three-dimensional clusters.

For highly symmetric clusters given by the icosahedral stannaspherene and plumbaspherene ($[Sn_{12}]^{2-}$ and $[Pb_{12}]^{2-}$) cages, $^{180-182}$ their electron count meets both Wade's 2n + 2 and the Hirsch $2(N + 1)^2$ (N = 4) electron counts, resulting in representative spherical aromatic clusters as probed by the magnetic criteria of aromaticity, where the role of relativistic corrections¹⁸³ and core electrons has been discussed.⁶⁸ The aromatic characteristics of the hollow icosahedral cages are retained for metal encapsulated counterparts given by $[M@Pb_{12}]^q$ (M = Co, Rh, Ir, Au; q = -3, and, M = Ni, Pd, Pt; q = -2), ^{175,184} also noted as intermetalloid clusters, ¹⁸⁵ highlighting the formation of a long-ranged shielding cone under the application of an orientation fixed external magnetic field. In addition, a good agreement is achieved in the calculation of

²⁰⁷Pb-NMR parameters, which further supports the quantities obtained for the magnetic criteria of aromaticity. 175 Moreover, for the proposed [Ca²⁺@Pb₁₂²⁻] cluster, ¹⁸⁶ a long-range shielding region is enabled for particular orientations of the external field, indicating that this behavior is inherent to the [Pb₁₂]²⁻ cage. Similar features have been described for $[E_9]^{4-}$ species, as given for $[Ge_9]^{4-.187}$

The chemical bonding analysis provided by the adaptive natural density partitioning (AdNDP) algorithm, 87 offers a realization of the bonding elements acting in the overall cluster, which has been employed to evaluate the $[Pb_4]^{4-}$, $[Pb_5]^{2-}$, $[Pb_9]^{4-}$, $[Pb_{10}]^{2-}$, and $[Pb_{12}]^{2-}$ clusters. For $[Pb_4]^{4-}$, the 20 ce are distributed in four 1c-2e lone-pairs and six 2c-2e σ -bonds; similarly, the 40 ce in [Pb₉]⁴⁻ shows nine 1c-2e lone-pairs and eleven multicentre σ -bonds, showing 3×4 c-2e, 3×5 c-2e, and 5×8 c-2e bonding elements in its C_{4v} geometry and $2 \times 3c$ -2e and $9 \times 5c$ -2e in the D_{3h} structure. In addition, for $[Pb_{12}]^{2-}$ an alternative bonding pattern is obtained using the AdNDP method, in comparison to canonical molecular orbitals, 181 revealing a set of twelve 1c-2e lone-pairs, six 5c-2e σ-bonds, and seven 10c-2e σbonds. 188 In addition, a more delocalized view of AdNDP for the [Pb₁₂]²⁻ cluster¹⁸⁶ has been provided for the related [Ca²⁺@ Pb₁₂²⁻] cluster, given by twelve 1c-2e lone-pairs, nine 13c-2e and four 12c-2e delocalized bonds. From both magnetic and electronic descriptors, the spherical aromaticity in such species is supported, providing a qualitative proof of these characteristics.

Besides spherical aromatics, Zintl ions or intermetalloids also exhibit planar aromatic characteristics, 189-191 among other types of aromaticities, 192 unravelling examples of the increased versatility in the chemistry of these clusters. The characterization of $[Sn_5]^{6-}$ and $[Pb_5]^{6-}$ as five-membered planar rings has been provided by Sevov, 193,194 showing that such species resemble the π -orbitals from the 6π aromatic cyclopentadienyl anion, C₅H₅⁻, ascribing such species to heavy-metal aromatic rings. The overall 26 ce are distributed in five 1c-2e lone-pairs, five 2c-2e σ -bonds, and in a set of three 5c-2e delocalized π -bonds, which meets the 4N + 2 Hückel rule. It is noteworthy that aromatic 2π electron Ga_5 rings in $[Ga_5(CH(SiMe_3)_2)_5]^{2-}$ have been characterized, 195 which exhibit a single 5c-2e delocalized π-bonding element, enabling electronic delocalization as probed by magnetic descriptors from magnetically induced ring currents and NICS values. Thus, two isostructural rings are able to sustain aromatic characteristics despite the different π -electron count fulfilling the Hückel rule.

Recently, in two separate reports, the characterization of five-membered rings Sb₅⁻ and Bi₅⁻ is provided, ^{189,190} highlighting the presence of induced ring currents from magnetic descriptors, with diatropic currents with a strength of 14.4 nA T^{-1} , which is comparable to the calculated for the cyclopentadienyl anion (12.8 nA T⁻¹). In addition, the AdNDP analysis exhibits five 1c-2e lone-pairs, five 2c-2e σ -bonds, and a set of six π -electrons, thus, supporting the planar aromatic characteristics of both magnetic and electronic descriptors. In addition, the role of relativistic effects, particularly the spin-orbit coupling in the [M₅] series (M = N, P, As, Sb, Bi, Mc) has been discussed in the literature, ¹⁹⁶ denoting a very relevant role in the heavier members, exposing

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the requirement of taking into account such effects to achieve a proper magnetic evaluation of these systems. Thus, extending the aromaticity concept to cases where relativistic effects are crucial favours an equal footing treatment and evaluation of aromatic species across the periodic table. Moreover, it is shown that [Sb₅] and [Bi₅] rings enable the formation of M-M bonds from coordinating metal atoms located on the faces of such rings, 189 in analogy to the recent characterization of the B₀ cluster. ¹⁹⁷ M₅ aromatic rings are recursive motifs, as has been found in several species mimicking the coordinating chemistry of cyclopentadienyl anions, which has been collected and discussed in a previous review from Boldyrev and Sun.167

In addition to 2π and 6π aromatic rings, solid phases of the stoichiometry of Ba₄Li₂Si₆ and Ba₄Li₂Ge₆ reveal six-membered cyclic Zintl-ions motifs of the [Si₆]¹⁰⁻ and [Ge₆]¹⁰⁻ forms, respectively, highlighting the formation of aromatic 10π Hückel aromatic rings. 198

Smaller aromatic ring members have also been isolated, as for example, the σ -aromatic $[Bi_4]^{4+}$ ring sustaining an induced ring current strength of 9.1 nA T⁻¹, lower in comparison to the seminal [Al₄]⁴⁻ cluster (14.8 nA T⁻¹).¹⁹⁹ And aromatic threemembered rings have been characterized in [As₃Nb(As₃Sn₃)]³⁻ and $[Sb_3Au_3Sb_3]^{3-}$, among others. These examples expose both structural and chemical bonding versatility of Zintl-ions, enabling them to achieve an aromatic behaviour favouring a stable situation (vide infra). Given this rich diversity, the choice of an appropriate aromaticity descriptor is challenging, where the recommendation to involve at least both magnetic and electronic descriptors is suggested to be particularly useful in providing reliable results from different points of view, owing to the multidimensional character of this concept.²⁰⁰

Other outstanding species have been characterized involving several structural layers, which serves as an extension of the spherical aromaticity in endohedral M@E₁₂ species. The intermetalloid cluster coined as "a bronze matryoska" characterized by Fässler, involves three concentric structural layers, ²⁰¹ featuring an inner Sn atom, encapsulated into a Cu12 icosahedron coated by a Sn_{20} dodecahedron with the formula $[Sn@Cu_{12}@Sn_{20}]^{12-}$. This prototypical spherical structure is based on the icosahedron-dodecahedron duality according to the number of vertices and faces, where the icosahedron has 20 faces and 12 vertices, and the dodecahedron has 12 faces and 20 vertices. Besides its aesthetically pleasing structure, this outstanding architecture serves as a prototypical case study to unravel the aromatic characteristics in multilayer clusters. The AdNDP analysis of $[Sn@Cu_{12}@Sn_{20}]^{12-}$ reveals thirty 4c-2e σ -bonds as bonding elements connecting the two atoms from the outer Sn20 cage (Fig. 5) to two adjunct Cu atoms from the internal Cu₁₂ icosahedral cage.²⁰² Interestingly, the remaining 8 electrons are delocalized over the overall Sn@Cu₁₂ cage, resulting in a set of four 13c-2e bonds which satisfy the Hirsch rule of spherical aromaticity, ¹⁷⁹ suggesting that both inner structural layers feature electronic delocalization giving rise to spherical aromatic properties.

Furthermore, the plausible formation of a spherical aromatic cluster for [Sn@Cu₁₂@Sn₂₀]¹²⁻ has been evaluated via the magnetic criteria of aromaticity, which shows a shielding

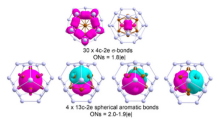


Fig. 5 Bonding pattern of [Sn@Cu₁₂@Sn₂₀]¹²⁻. Copper atoms are brown. Reproduced with permission from ref. 202. Copyright Wiley, 2020.

region ascribed to the structural backbone, as noted from the isotropic term (B_{iso}^{ind}) , also noted as the NICS isosurface (Fig. 6). From specific orientations of the external field, a long-ranged shielding cone characteristic is enabled, which extends to 11.5 Å from the centre of the structure with shielding values of -3.0 ppm and to 16.0 Å with values of -1.0 ppm. Such observations further support the spherical aromaticity of this multi-layered cluster, serving as a prototypical case for further identification of similar aromatic clusters. Interestingly, such observation is similar to that given for fullerenes and endohedral metallofullerenes, 203-206 suggesting the equal footing treatment of aromaticity in these structures based on the magnetic criteria of aromaticity.

Now, we turn our attention to structures seen as the aggregation of individual cluster building blocks. In this issue, the $[Pd_2@E_{18}]^{4-}$ clusters (E = Ge, Sn) has been rationalized as the fusion of parent spherical aromatic $[Pd@E_{12}]^{2-}$ building units, 207,208 resulting in an interesting intercluster bonding pattern, leading to an overall bond order of 2.70 and 2.31 for [Pd₂@Ge₁₈]⁴⁻ and [Pd2@Sn18]4-, respectively.209 Moreover, to evaluate the plausible aromatic behaviour in this structure involving two fused clusters, the characteristics of the induced magnetic field have been obtained, showing a related shielding surface contained within the oblate cage from the NICS isosurface (Fig. 7). When compared to the [Pd@E₁₂]²⁻ parent cluster, the NICS isosurface in $[Pd_2@E_{18}]^{4-}$ is centred on both sides, denoting aggregation of two building units. Under specific orientations of the external field the shielding cone characteristics in the parent cluster are enabled, which are enhanced after formation of the $[Pd_2@E_{18}]^{4-}$ clusters, denoting the spherical aromatic characteristics of the overall structure as the fusion of two spherical aromatic cluster units.

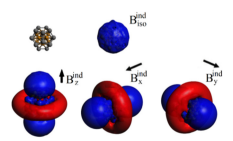


Fig. 6 Magnetic response properties of $[Sn@Cu_{12}@Sn_{20}]^{12-}$, denoting the isotropic (averaged) term ($B_{\rm iso}^{\rm ind}$), also noted as NICS isosurfaces, and under specific orientations of the external field (B_z^{ind} , B_x^{ind} , and B_y^{ind}). Isosurfaces set to ± 3 ppm, blue - shielding; red - deshielding. Reproduced with permission from ref. 202. Copyright Wiley, 2020.

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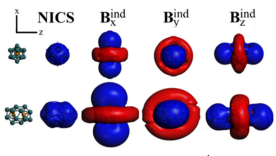
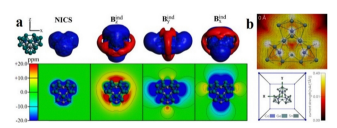


Fig. 7 Magnetic response properties of [Pd₂@Sn₁₈]⁴⁻, denoting the NICS isosurfaces, and from specific orientations of the external field (B_z^{ind}, B_x^{ind}) and B_{ν}^{ind}). Isosurfaces set to ± 3 ppm, blue – shielding; red – deshielding. Reprinted with permission from ref. 209. Copyright Royal Society of Chemistry, 2024.

Lastly, the aggregation of three cluster units in a cyclic array is accounted for through the recent [Co₃@Ge₆Sn₁₈]⁵⁻ structure, ²¹⁰ which can be viewed as the aggregation of three [Co@Ge₃Sn₆]⁴⁻ building units. This cluster features 32 electrons, retaining the bonding per each fused cluster unit, giving rise to three superatomic units with a filled 1S²1P⁶1D¹⁰1F¹⁴ electronic closed-shell. In addition, two electrons are delocalized in the overall cluster structure in a 27c-2e σ-bonding element, which fills a bonding combination of three 2S shells centred at each building unit, sustaining a bonding 2S + 2S + 2S combination leading to cyclic cluster-of-cluster bonding characteristics. Interestingly, this 27c-2e bonding element is analogous to the 3c-2e bonding in H₃⁺ and Li₃⁺, suggesting the extension of aromatic properties from these small triatomic rings to a large cluster-based aggregate. Interestingly, from the global electron density of delocalized bonds, 211-213 it is pointed out that the electronic delocalization is of global character, involves the entire cluster, which is also supported by calculating the magnetically induced currents from the GIMIC suite^{54,214} and analysis of the induced magnetic field (Fig. 8). From the NICS isosurface, the continuous shielding region is centred at each cluster building unit, resulting in three adjacent shielding regions. Under specific orientations of the external field, the overall [Co₃@Ge₆Sn₁₈]⁵⁻ cluster structure enables shielding cone characteristics when the external field is oriented through the



(a) Isosurface and contour plot representation of the magnetic response properties for $[Co_3@Ge_6Sn_{18}]^{5-}$, denoting the NICS (isotropic/ averaged) term, and from different orientations of the external field (B_x^{ind}) B_y^{ind} , and B_z^{ind}). Isosurfaces set to ± 8 ppm, blue: shielding; red: deshielding. (b) Current density of the $[Co_3@Ge_6Sn_{18}]^{5-}$ cluster at the central plane located at 0 Å, containing the $[Co_3Ge_3]$ ring. The calculations were performed at the PBEO/def2-TZVP level of theory, without including solvent effects. Reprinted with permission from ref. 210. Copyright 2025 American Chemical Society.

z-axis (B_z^{ind}) , in line with the appearance of the delocalized 27c-2e bonding element fulfilling the Hückel rule for aromaticity. Thus, this structure is ascribed as the first σ-bonded cluster trimer unravelling a σ-aromatic character, reported to date. Moreover, under parallel orientations (B_x^{ind} and B_y^{ind}), a similar long-range shielding cone is obtained originating at each building unit, denoting that besides the electronic delocalization in the overall [Co₃@Ge₆Sn₁₈]⁵⁻ cluster structure, the inherent spherical aromatic characteristics of the constituent building units are retained.

Note that from the results obtained for the [Pd₂@Sn₁₈]⁴⁻ dimer and [Co₃@Ge₆Sn₁₈]⁵⁻ trimer, under different orientations of the external field, long-range shielding regions are enabled which are complemented with a perpendicular deshielding region, which nicely resemble the classical shielding cone properties from planar aromatics. 147,215,216 but enabled from any orientation in spherical aromatic species owing to its three-dimensional character.

7. Conclusions

The exploration of aromaticity in all-metal clusters has deeply reshaped our understanding of this fundamental chemical concept. Unlike the straightforward rules governing classical organic molecules, all-metal clusters introduce remarkable complexity, often displaying multifold (anti)aromaticity in the σ -, π -, δ - or ϕ electron delocalization paths or conflicting aromaticity among several of these components. To navigate this intricate landscape, researchers usually combine magnetic-based indicators with electron delocalization-based indices to quantify the aromatic character of all-metal clusters. While, despite their relevance, resonance energies are rarely computed because of inherent difficulties in their definition. Supplementing these measures with an analysis of existing electron counting rules provides invaluable qualitative insights, even if these rules do not offer a quantitative metric.

The journey since the discovery of the first all-metal aromatic cluster, Al₄²⁻, clearly demonstrates how this concept has expanded across the periodic table. This significant progress is not just an academic curiosity; it directly challenges and broadens our foundational understanding of many aspects of the aromaticity concept. As the relevance of aromaticity continues to grow in various fields, it is highly probable that we will see further advancements in connecting the existing rules and formulating new ones in the years ahead. This ongoing evolution will move us closer to a unified theory of aromaticity, which hopefully will provide connections among apparently unrelated aromatic systems.

Author contributions

M. S.: conceptualization, writing - original draft, writing review & editing; A. M.-C.: conceptualization, writing - original draft, writing - review & editing.

Conflicts of interest

There are no conflicts to declare.

Data availability

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No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

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