Heterometallic clusters have attracted broad interests in the synthetic chemistry due to their various coordination modes and potential applications in heterogeneous catalysis. Here we report the synthesis, experimental, and theoretical characterizations of four ternary clusters ([M$_2$(CO)$_6$Sn$_2$Sb$_5$]$^{3-}$ (M = Cr, Mo), and [(MSn$_2$Sb$_5$)$_2$]$^{4-}$, (M = Cu, Ag)) in the process of capturing the hypho- [Sn$_2$Sb$_5$]$^{3-}$ in ethylenediamine (en) solution. We show that the coordination of the binary anion to transition-metal ions or fragments provides additional stabilization due to the formation of locally σ-aromatic units, producing a spherical aromatic shielding region in the cages. While in the case of [Mo$_2$(CO)$_6$Sn$_2$Sb$_5$]$^{3-}$ stabilization arises from locally σ-aromatic three-centre and five-centre two-electron bonds, aromatic islands in [(AgSn$_2$Sb$_5$)$_2$]$^{4-}$ and [(CuSn$_2$Sb$_5$)$_2$]$^{4-}$ render them globally antiaromatic. This work describes the coordination chemistry of the versatile building block [Sn$_2$Sb$_5$]$^{3-}$, thus providing conceptual advances in the field of metal-metal bonding in clusters.
Zintl-precursores play a key role in the construction of numerous metal clusters via complex coordination. The first structural determination of a Zintl anion was made by Kummer and Diehl in the 1970s, when they revealed the presence of $\text{Sn}_3^-$, a cluster which was subsequently used as a precursor for the synthesis of many polystannides. Afterward, many more homo- and heteroatomic polyanions were discovered including $[\text{E}_9]^{4-}$ (E = Si-Pb), $\text{[Pt}_4\text{I}_3]^-$ (Pt = P-BI), and $[\text{Tt}_4\text{Pt}_6]^{2-}$ (Tt = Ge-Pb), and a number of new clusters were produced using them as precursors. In the majority of these clusters, the relationship between structure, bonding, and valence electron count can be understood in terms of either the Zintl-Klemm concept addressing electron-precise structures composed of formal two-center two-electron (2c-2e) bonds, or the Wade-Mingos rules, which were originally applied to electron-deficient boranes with delocalized bonding, and later introduced into Zintl clusters. According to Wade-Mingos rules, $\text{[B}_{10}\text{H}_{10}]^{2-}$ is a classic closo-type polyhedron (Fig. 1a), isosstructural with $[\text{Ge}_{9}]^{6-}$ and $[\text{Pb}_{9}]^{3-}$. The most commonly used Zintl-precurors $[\text{E}_n]^{4-}$ (E = Si-Pb) of the nido- type can be formed by removing one 4-connected vertex from the closo-type (Fig. 1b). Lately, Dehnen and coworkers reported the first arachno-Zintl ion $[\text{Sn}_3\text{Bi}_3]^{3-}$, which fills the gap in the series of the known compounds (Fig. 1c). The only reported hypho-Zintl anion, $[\text{Sn}_3\text{Bi}_3]^{3-}$ with high negative charges, was observed only in liquid ammonia. However, the hypho-cage derived from the 10-vertex $[\text{B}_{10}\text{H}_{10}]^{2-}$ closo-type has not yet been observed.

Recently, we adopted self-assembly as a synthetic strategy to synthesize a series of Au/Pb clusters $[\text{Au}_x\text{Pb}_y]^{4-}$, $[\text{Au}_x\text{Pb}_y\text{I}_z]^{4-}$, constructed from a nido-type cluster $[\text{Au}_x\text{Pb}_y]^{4-}$, which was first reported by us. As part of this ongoing study, we aim to build metal clusters using reaction of $\text{K}_8\text{SnSb}_4$ with oxidizing organometallic compounds, which are stable towards electrons ejections and may serve as a potential building unit for further reactions. According to the Wade-Mingos rules, $[\text{E}_n]^{4-}$ (E = Si-Pb) of the nido-type can be formed by removing one 4-connected vertex from the closo-type (Fig. 1b). Lately, Dehnen and coworkers reported the first arachno-Zintl ion $[\text{Sn}_3\text{Bi}_3]^{3-}$, which fills the gap in the series of the known compounds (Fig. 1c). The only reported hypho-Zintl anion, $[\text{Sn}_3\text{Bi}_3]^{3-}$ with high negative charges, was observed only in liquid ammonia. However, the hypho-cage derived from the 10-vertex $[\text{B}_{10}\text{H}_{10}]^{2-}$ closo-type has not yet been observed.

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regarded as derived from the 10-vertex closo deltahedra by removing one 4-connected and two 5-connected vertices. Clusters 1 and 2 are isostructural, and contain nine atoms in a distorted mono-capped square antiprismatic geometry, with Sb atoms occupying all five 4-connected vertices and two Sn and Mo atoms at the 5-connected vertices. The two-electron donor transition metal fragments Mo(CO)₃ coordinate to the 5-membered rings of the Sn₂Sb₂ cluster and form an expanded square plane Mo₂Sb₂ (3.1788(8)–3.2407(8) Å) compared with the bottom Sn₂ plane (2.803(9)–2.925(9) Å). All Mo–Sb bond lengths fall in a narrow range, from 2.925(8) Å to 2.953(8) Å, comparable to those reported in the [Sn₂Mo(CO)₃]₃ anion (2.942(8)–2.951(9) Å). 31

The anionic clusters [(AgSn₂Sb₃)₂]⁻ and [(CuSn₂Sb₃)₂]⁻ are also isostructural and can be described as a dimer of two arachno-cages (MSn₂Sb₅)²⁻ sharing an M₂Sb₂ diamond plane. From the structural perspective, the hypho-[Sn₂Sb₅]¹⁻ is very similar to that in 1 and 2 and the comparison of selected bond distances are listed in Table 1. By introducing a Cu⁺ or Ag⁺ ion to occupy one of the above-mentioned 5-connected vertices, 1 is transformed to an arachno-cage (MSn₂Sb₅)²⁻, which is isostructural with the recently reported Zintl ion [Sn₂Sb₅]¹⁻. 23. The two (MSn₂Sb₅)²⁻ cages are arranged upside down resulting in two parallel Sn₂ planes. In the Ag system (3), the Ag–Sn distances range from 2.970(3) Å to 3.025(3) Å, similar to those in [Ag(Sn₉-Sn₉)]⁵⁻ (2.880(2)–3.010(1) Å) 32, but longer than the typical Ag–Sb single bonds. There are limited opportunities to compare the Ag–Sb bonds with literature precedents because only a few organometallic compounds such as [Ag₂(Sb(SiMe₃))₆]³⁺(PiPr₃)₆ (2.722 (10)–2.746(12) Å) are available as ref.33. There are three types of Ag–Sb bond lengths in cluster 3. Ag₁–Sb₁–Ag₁’–Sb’₁ construct a diamond plane, in which Ag₁–Sb₁ and Ag₁’–Sb’₁ are both 2.758(14) Å, similar to most Ag–Sb single bonds. However, Ag₁–Sb₁ and Ag₁’–Sb’₁ are 2.917(15) Å, significantly longer than those of the reported complexes. The remaining Ag–Sb bonds lie in a narrow range of 2.874(2)–2.875(3) Å. The only Ag–Ag bond is 2.802(2) Å, shorter than the sum of covalent radii for a single Ag–Ag bond, i.e., 2.88 Å. However, the Cu,Sb₂ diamond plane is compressed compared to Ag₂Sb₂ in 4. The bond length of Cu–Sb₁ (2.669 Å) is akin to Cu–Sb’₁ (2.616 Å), close to those of [(CuSn₂Sb₅)²⁻] (2.619(2)–2.641(2) Å) 34, but considerably longer than typical Cu–Sb single bond, 2.554(9) Å in [CuCu(SbPh₃)₂]³⁵. The Cu–Cu distance is 2.545 Å, well comparable with those of [Cu₂@Sn₅]⁻ (2.529(12)–2.551(13) Å) and [(CuGe₉Mes₃]⁴⁻ (2.521(7) Å) 36,37.

Structure and chemical bonding. According to Wade-Mingos and mno rules 18-20,38, every naked Sn atom bearing a lone pair provides two electrons for cluster bonding (2 × 2), and each Sb atom owns three skeleton electrons (3 × 5), plus three negative charges, leading to a total of 22 skeleton electrons (2n + 8). The cluster contains one polyhedron (m = 1), seven atoms (n = 7) and three missing vertices (o = 3), totally generating 11 orbitals.

Table 1 Comparison of selected atomic distances for clusters 1, 2, 3, and 4.

| Cluster | Sb–Sb | Sn–Sb | M–Sn | M–Sb/Sb¹ |
|---------|-------|-------|------|----------|
| 1       | 2.785–2.948 | 2.784–2.934 | 3.125–3.203 |       |
| 2       | 2.803–2.926 | 2.813–2.967 | 3.179–3.241 |       |
| 3       | 2.790–2.930 | 2.790–2.945 | 2.970–3.025 | 2.917–2.758 |
| 4       | 2.816–2.942 | 2.816–2.913 | 2.815–2.860 | 2.669–2.616 |
available for cluster bonding. By introducing a $d^{10}$ metal atom (Ag or Cu) to occupy one of the missing 5-connected vertices, each half of dimeric 3 and 4 attains an arachno-type structure. In light of this, Ag$^+$ or Cu$^+$ provides one electron for cluster bonding, leading to 22 skeleton electrons (2$n + 6$) and fitting into $m + n + o$ rule as well. Two Mo$^0$ atoms each coordinated by three carbonyl groups occupy two 5-connected vertices, yielding a distorted monocapped square antiprism nido-type cluster 2. Thereby, the skeletal electron count is consistent with the above two cases.

In order to understand the chemical bonding of clusters 1, 2, 3, and 4, we started our analysis from the hypoh-[Sn$_2$Sb$_3$]$_{3}^-$ unit. The 36 valence electrons were distributed performing adaptive natural partitioning (AdNDP) analysis$^{39}$ as implemented in AdNDP 2.0 code$^{40}$. The AdNDP method can represent a chemical bonding pattern in terms of both Lewis bonding elements (lone pairs (1c–2e) and 2c–2e bonds) as well as delocalized bonding elements (such as nc–2e ($n > 2$) bonds), while the later are usually associated with the concepts of aromaticity and antiaromaticity. This technique has been widely used in describing chemical bonding pattern of various Zintl clusters$^{41-47}$. The chemical bonding of [Sn$_2$Sb$_3$]$_3^-$ is entirely described in terms of the classical 1c–2e and 2c–2e bonds (Supplementary Fig. 31). Specifically, seven s-type lone pairs with $\text{ON} = 1.98–1.89$ [e] were found on each Sn and Sb atoms, and one p-type lone pair with $\text{ON} = 1.71$ [e] was localized on the apex Sb-atom. The remaining 20 electrons form ten 2c–2e σ-bonds with $\text{ON} = 1.99–1.92$ [e], which are responsible for the shape of the entire frame. Since the initial precursor is rather unstable in solution, it can be expected that the transition-metal coordination increases the stability of these clusters. According to the comparative chemical bonding analyses of complexes 1–4, the stabilization effect can be explained by the formation of locally σ-aromatic fragments between transition metal atoms and [Sn$_2$Sb$_3$]$_3^-$ unit.

The optimized geometry of the [Mo$_2$(CO)$_6$Sn$_2$Sb$_5$]$_3^-$ has $C_{py}$ symmetry, and all bond lengths are within 0.07 Å of their crystallographic counterparts. Overall, 124 valence electrons can be found as 62 two-electron bonding elements. From the optimized geometry of the cluster, it is obvious that the geometry of [Sn$_2$Sb$_3$]$_3^-$ unit is preserved, and, hence, similar bonding fragments are anticipated in this structure. Indeed, seven 1c–2e s-type lone pairs and four 2c–2e Sb–Sb σ-bonds were also found in [Mo$_2$(CO)$_6$Sn$_2$Sb$_5$]$_3^-$, similar to the [Sn$_2$Sb$_3$]$_3^-$ cluster (Fig. 4a). However, the rest of the bonding elements were found as delocalized: four 2c–2e σ-bonds with $\text{ON} = 1.91$ [e] were found over the Mo–Sb–Sn units, and the remaining six electrons form three 5c–2e bonds that are responsible for the locally σ-aromatic behavior of the Mo$_3$Sn$_2$Sb cap fragment. Thus, it is the formation of the multicerter 3c–2e and 5c–2e bonding elements that provides extra stabilization of the complex. In addition to AdNDP, the topology analysis of the electron localization function$^{48}$ (ELF) was performed for this cluster, as additional valuable method for determination of chemical bonding of Zintl clusters$^{49}$. The ELF results are fully consistent with the AdNDP picture (Supplementary Fig. 39). The same bonding pattern was found for the cognate [Cr$_2$(CO)$_6$Sn$_2$Sb$_5$]$_3^-$ complex, although with slightly different ON values (Supplementary Figs. 33–34).

A different origin of stabilization was found in the [(CuSn$_2$Sb$_3$)$_3$]$_4^-$ and [(CuSn$_2$Sb$_3$)$_3$]$_4^-$ clusters. As one would expect, the same bonding patterns were revealed for these species due to the valence isoelectronic Ag and Cu, in agreement with their similar geometrical structures. For the sake of simplicity, we will describe only the Cu-containing cluster below, noting that the same conclusions hold for the Ag-containing counterpart. Some of the bonds relevant to the discussion of the bonding pattern in the [(CuSn$_2$Sb$_3$)$_3$]$_4^-$ cluster are shown in Fig. 4b. As in the case of the Mo-based complex, the optimized geometry of [CuSn$_2$Sb$_3$]$_4^-$ contains two Sn$_2$Sb$_5$ fragments that are almost identical to the isolated [Sn$_2$Sb$_3$]$_3^-$ unit geometry. However, in this case, the similarity is even more traced. Thus, fourteen s-type lone pairs with $\text{ON} = 1.97–1.89$ [e] and twenty 2c–2e bonds with $\text{ON} = 1.98–1.80$ [e] were localized, which almost completely coincides with the pattern found in [Sn$_2$Sb$_3$]$_3^-$ unit. Further localization leads to five doubly occupied d-type lone-pairs on each Cu-atom with $\text{ON} = 1.99–1.97$ [e]. The remaining 4 electrons form two 4c–2e Sb–Cu–Cu–Sb σ-bonds (Supplementary Fig. 36), whose combination can give two 3c–2e Cu–Cu–Sb σ-bonds (Fig. 4b).

The complete bonding patterns for the Ag-containing and Cu-containing complexes are shown in Supplementary Figs. 37, 38, respectively. The shape of the Cu$_2$Sb$_2$ fragment, the chemical bonding picture, and the number of electrons (4e) hint at the antiaromatic character of this diamond-shaped unit. Positive NICS$_{zz}$ values at the center of Cu$_2$Sb$_2$ confirm the antiaromatic behavior of this unit (Supplementary Table 8). This is also in agreement with our previous studies on the [CuGe$_3$Mes$_3$]$_3^-$ cluster featuring a similar Cu$_2$Ge$_2$ fragment that was found to exhibit the antiaromatic character$^{37}$. Similarly, it is noted that the antiaromaticity of fragments with four atoms leads to the formation of two locally σ-aromatic 3c–2e rings providing stabilization (Supplementary Fig. 36). This is denoted by their negative NICS$_{zz}$ values at the center of the M$_2$Sb$_2$ triangle (Supplementary Table 8), as manifestation of the characteristics central shielding for aromatic rings, with a complementary deshielding region (positive NICS$_{zz}$ values) which overlaps at the center of the M$_2$Sb$_2$ diamond.

To further explore the potential global aromatic/antiaromatic characteristics of 2–4, we report the induced magnetic field (B$^{\text{ind}}$) as a magnetic criterion of aromaticity (Fig. 5)$^{50-52}$. Spherical aromatic clusters are able to sustain a long-ranged shielding cone as a distinctive property involving the overall structure. It is useful to compare directly the induced field for all four clusters despite their different shape and composition. The isotropic term (B$_{\text{iso}}^{\text{ind}}$), accounting for the average of different orientations of the applied field resembling the constant molecular tumbling in solution-state, exhibits a shielding surface for [Mo$_2$(CO)$_6$Sn$_2$Sb$_5$]$_3^-$ owing to the presence of the multiple delocalized bonding elements (Fig. 4). It is depicted by 3c–2e and 5c–2e bonds within the cluster cage, which share common atom centers (certain atoms contribute to both multi-center bonds), resulting in a spherical-like shielding surface as observed for other Zintl-clusters and intermetalloids$^{41,53}$. This situation is similar to the [Ge$_4$]$_4^-$ case$^{42}$, where the presence of multiple σ-aromatic bonding elements, sharing common atom centers, generates a spherical-like aromatic behavior given by the characteristic shielding surface (Supplementary Fig. 42). To achieve the inherent characteristics of spherical-like aromatic clusters related to the observation of a featured shielding cone, the analysis of the different terms of B$^{\text{ind}}$ accounting for specific orientations of the applied field is given. This behavior builds a direct relation between planar and spherical aromatic compounds, which is now extended to spherical-like aromatic species bearing multiple aromatic circuits with common atom centers in the same cage. In the case of planar aromatic species, a shielding cone is established solely when the field is oriented perpendicularly to the ring, in contrast to its appearance under different orientations of the field in three-dimensional aromatic species$^{54,55}$.

For [Mo$_2$(CO)$_6$Sn$_2$Sb$_5$]$_3^-$, a long-ranged shielding response with a complementary perpendicular deshielding region is established under a field along the z-axis (B$^{\text{ext}}$). Interestingly, under different orientations given by B$_{\text{x}}^{\text{ext}}$ and B$_{\text{y}}^{\text{ext}}$, the same features are retained, indicating spherical aromatic behavior in [Mo$_2$(CO)$_6$Sn$_2$Sb$_5$]$_3^-$, which is not found in the hypho-[Sn$_2$Sb$_3$]$_3^-$ unit. This observation suggests that for the reactive precursor, the inclusion of Mo(CO)$_3$ fragments achieves a more favorable situation resulting in the above depicted multiple-local aromatic circuits, combining in a spherical-like aromatic cluster, able to capture the hypho-[Sn$_2$Sb$_3$]$_3^-$ motif. It
has been recently discussed by Solà and Teixidor on the basis of closo-C_{2}B_{10}H_{10}, that the spherical aromaticity in deltahedral clusters for a closo-cluster is able to be retained after the removal of a single vertex leading to an aromatic nido-counterpart\(^{56}\). Our results suggest that for a hypho-motif, the spherical-like aromatic behavior ascribed to the nido-[Mo_{2}(CO)_{6}Sn_{2}Sb_{5}]^{3–} is not able to be retained, as a direct consequence of the consecutive removal of cluster-vertices. Similar results are found for the Cr counterpart (Supplementary Fig. 42).

After aggregation of two hypho-[Sn_{2}Sb_{5}]^{3–} motifs linked by two Ag(I) or Cu(I) atoms, the resulting cluster shows two separated spherical aromatic regimes as suggested by the appearance of two spherical-like shielding regions ascribed to each MSn_{2}Sb_{5} side (Fig. 5b, c), from the \(B_{\text{ind}}\) term. Moreover, specific orientations of the field cause the shielding cone behavior to be exposed, which results in a dual spherical-spherical aromatic cluster with two separated spherical aromatic regimes bridged by an antiaromatic M_{2}Sb_{2} diamond, similar to our findings in \([\text{CuGe}_{9}\text{Mes}]^{2–}\) with an antiaromatic Cu_{2}Ge_{2} bridging unit\(^{37}\). This behavior is related to the one found for \([\text{CB}_{11}H_{11}]^{2–}\) with two separated spherical closo-carboranes\(^{57}\). The characterization of \([\text{AgSn}_{2}Sb_{5}]^{14–}\) and \([\text{CuSn}_{2}Sb_{5}]^{14–}\), retaining two spherical aromatic motifs resulting in a stable aggregate, suggests that the hypho-[Sn_{2}Sb_{5}]^{3–} is a useful structural unit to achieve stable building blocks prone to condensate in a controlled manner.

**Discussion**

In summary, we have synthesized four ternary clusters \([\text{M}_{2}(\text{CO})_{6}Sn_{2}Sb_{5}]^{3–}\) (M = Cr, Mo), and \([\text{MSn}_{2}Sb_{5}]^{14–}\) (M = Cu, Ag) by the coordination of the hypho-[Sn_{2}Sb_{5}]^{3–} to various transition metals. Through detailed theoretical analysis, we have shown that the origin of stabilization of the clusters lies in the formation of locally \(\sigma\)-aromatic regions that result in a spherical-like shielding surface. In this work, we have presented an effective way to lead the construction of intermediates that can be used in further studies. We believe that our findings will bring new ideas to search for unstable main-group clusters and will also contribute to our understanding of chemical bonding in heterometallic clusters.

**Methods**

All manipulations and reactions were performed under a nitrogen atmosphere using standard Schlenk or glovebox techniques. Et (Aldrich, 99%) and DMF (Aldrich, 99.8%) were freshly distilled by CaH\(_{2}\) prior to use, and stored in N\(_{2}\) prior to use. Tol (Aldrich, 99.8%) was distilled from sodium/benzophenone under nitrogen and stored under nitrogen. \([2.2.2]\)-crypt (4,7,13,16,21,24-Hexaoxa-1,10-diazabicyclo[8.8.8]hexacosane, purchased from Sigma-Aldrich, 98%) was dried in a vacuum for one day prior to use. K\(_{8}\)SnSb\(_{4}\) was prepared by heating a stoichiometric mixture of the elements (K: 625.6 mg, Sn: 237.4 mg, Sb: 974.4 mg; K: +99%, Sn: 99.99%, Sb: 99.9%, all from Strem) at a rate of 70 °C per hour to 700 °C and keeping it at 36 h in sealed niobium containers closed in evacuated quartz ampules according to the previous procedures\(^{26}\). The K\(_{8}\)SnSb\(_{4}\) solid was obtained with a
high yield (−92%, 1.7 g) and stored under a dry nitrogen atmosphere in a glove box. Cr(CO)6 and Mo(CO)6 were purchased from Aldrich while Ag4Mes4 and Cu(PPh3)Cl were synthesized according to the literature with a yield of 55 and 60%, respectively58,59.

Synthesis of [K(2.2.2-crypt)]3[Cr2(CO)6Sn2Sb5] (1). K8SnSb4 (50 mg, 0.055 mmol) and 2.2.2-crypt (83 mg, 0.220 mmol) were dissolved in 2.5 mL en and stirred for 0.5 h to yield a dark brown solution. Cr(CO)6 (25 mg, 0.110 mmol) was added into the reaction mixture resulting in a red-brown suspension, and stirred for 2.5 h at 60 °C. The resulting solution was filtered through glass wool and transferred to a test tube, then carefully layered by toluene (3 mL) to allow for crystallization. After one week, dark red plate crystals of [K(2.2.2-crypt)]3[Cr2(CO)6Sn2Sb5] as a side product was also crystallized on the wall of the test tube.

Synthesis of [K(2.2.2-crypt)]3[Mo2(CO)6Sn2Sb5] (2). K8SnSb4 (50 mg, 0.055 mmol) and 2.2.2-crypt (83 mg, 0.220 mmol) were dissolved in 2.5 mL en and stirred for 0.5 h to yield a dark brown solution. Mo(CO)6 (33 mg, 0.110 mmol) was added into the reaction mixture resulting in a red-brown suspension, and stirred for 2.5 h at 60 °C. The resulting solution was filtered through glass wool and transferred to a test tube, then carefully layered by toluene (3 mL) to allow for crystallization. After one week, dark red plate crystals of [K(2.2.2-crypt)]3[Cr2(CO)6Sn2Sb5] as a side product was also crystallized on the wall of the test tube.

Synthesis of [K(2.2.2-crypt)]4[(AgSn2Sb5)2] (3). In vial 1, K8SnSb4 (50 mg, 0.055 mmol) and 2.2.2-crypt (83 mg, 0.220 mmol) were dissolved in 2.5 mL en and stirred for 0.5 h to yield a dark brown solution. In another vial, Ag4(Mes)4 (25 mg, 0.027 mmol) was dissolved in 0.5 mL tol resulting in a white suspension, and added into vial 1 dropwise. The reaction mixture turned into yellow-brown and stirred for 3.5 h at room temperature. The resulting solution was filtered through glass wool and transferred to a test tube, then carefully layered by toluene (3 mL) to allow for crystallization. After five months, brown thin plate crystals of [K(2.2.2-crypt)]4[(AgSn2Sb5)2] (3) were isolated in 12% (based on Ag4(Mes)4) together with red crystals [K(2.2.2-crypt)]2[Sn7Sb2] crystallized on the wall of the test tube.

Synthesis of [K(2.2.2-crypt)]4[(CuSn2Sb5)2] (4). In vial 1, K8SnSb4 (70 mg, 0.075 mmol) and 2.2.2-crypt (112 mg, 0.300 mmol) were dissolved in 2.5 mL en and stirred for 0.5 h to yield a dark green solution. Cu(PPh3)Cl (27 mg, 0.075 mmol) was added into the reaction mixture resulting in a yellow-brown suspension. After stirring for 3 h, the mixture was centrifuged for 10 min at 8000 r/min. The supernatant was filtered through glass wool and transferred to a test tube, then carefully layered by toluene (3.5 mL) to allow for crystallization. After one week, brown block crystals of [K(2.2.2-crypt)][(CuSn2Sb5)2] (4) was isolated in 15% (based on Cu(PPh3)Cl) together with red crystals [K(2.2.2-crypt)][Sn2Sb2].

Theoretical methods. All structures were optimized at the DFT level using the PBE0 hybrid density functional and def2-TZVP basis set60. The Gaussian 16 code61 was used for the optimization procedures. The same level was used for chemical bonding analysis including the adaptive natural density partitioning (AdNDP)58,62 analysis and electron localization function (ELF)63 analysis. The AdNDP 2.0 code was used to perform AdNDP analysis, which is based on the general ideas of the NBO analysis64. The topology analysis of ELF was performed with the MultiWFN program65. The ChemCraft 1.8 software was used to visualize chemical bonding patterns and geometries of investigated compounds. The isostructural schemes of [Sn2Sb2]3− were explored via an unbiased Coalescence Kick (CK) algorithm66,67. 2000 sample structures were optimized at PBE0/LANL2DZ65 level. The isomeric structures of [K8SnSb4]− were explored at the same level with 4000 sample structures. The seven lowest structures for both CK calculations were reoptimized using different functionals and def2-TZVP basis set (Supplementary Tables 6, 7). The frequency calculations were performed using the harmonic approximation, and all reported structures were confirmed to be minima due to the absence of imaginary frequencies. To assess the aromaticity and anti-aromaticity with a quantitative parameter, NICS calculations were performed at PBE0/def2-TZVP level.

The nucleus-independent shielding tensors (n)52,66,67 were calculated within the GIAO formalism using the ADF2019 code68, employing the OPBE69,70 functional and an all-electron STO-TZ2P basis set, placed in a three-dimensional grid in order to evaluate the induced field (Biind), upon an external magnetic field (Bext), related via Biind = −σijBjext52,66,67. For convenience, the i and j suffixes are related to the x, y, and z-axes of the molecule-fixed Cartesian system (i, j = x, y, z). The values of Biind are given in ppm in relation to Bext. Relativistic effects were considered through the ZORA Hamiltonian74. Solvation effects were considered via the COSMO module as implemented in the ADF code by using ethylenediamine as a solvent.

X-ray diffraction. Suitable single crystals of compound 1′, 2′, 3′, and 4′ were selected for X-ray diffraction analyses. Crystallographic data were collected on Rigaku XtaLAB Pro MM007 DW diffractometer with graphite monochromated Cu Ka radiation (λ = 1.54184 Å). Structures were solved using direct methods and then refined using SHELXL-2014 and Olex2 to convergence57,77, in which all the non-hydrogen atoms were refined anisotropically during the final cycles and all hydrogen atoms of the organic molecule were placed by geometrical considerations. The disorder existed in crystal molecules of compound 3′ and heavy atoms of cluster 3 and 4 were solved by the Split SAME process in Olex 2. The disordered three solute solvent molecules of compound 1′ and 2′ were squeezed by the Solvent...
Electrospray ionization mass spectrometry (ESI-MS) investigations. Negative ion mode ESI-MS of the DMF solutions of crystals of 1 and 2 were measured on an LTQ linear ion trap spectrometer by Agilent Technologies ESI-TOF-MS (6320). The spray voltage was 5.48 kV and the capillary temperature was kept at 300 °C. The capillary voltage was 30 V. The samples were made up inside a glovebox under a nitrogen atmosphere and rapidly transferred to the spectrometer in an airtight syringe by direct infusion with a Harvard syringe pump at 0.2 mL/min.

Energy dispersive X-ray (EDX) spectroscopic analysis. EDX analysis on the compounds of 1, 2, 3, and 4 was performed using a scanning electron microscope (FE-SEM, JEOL JSM-7800F, Japan). Data acquisition was performed with an acceleration voltage of 15 kV and an accumulation time of 60 s.

Data availability

The data that support the findings of this study are available from the corresponding author on reasonable request. The X-ray crystallographic of compounds 1, 2, 3, and 4 reported in this study have been deposited at the Cambridge Crystallographic Data Centre (CCDC) under deposition numbers 2080455 and 2080457–2080459. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

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Z.M.S. conceived the project and designed the experiments. Y.H.X. performed the synthesis. N.Y.T., I.A.P., A.M.C., and A.I.B. performed the quantum chemical calculations and analyzed the data. I.Q. performed the single-crystal X-ray diffraction and analyzed the data. All authors co-wrote the manuscript.

Competing interests
The authors declare no competing interests.

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