[M@Sn_{14-x}Sb_x]^q− (M = La, Ce, or U; x = 6–8; q = 3, 4): Interaction of 4f or 5f Metal Ions with 5p Metal Atoms in Intermetalloid Clusters

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ABSTRACT: The impact of 4f metal ions Ln^{5+} (Ln = La or Ce) versus 5f metal ions U^{n+} (n = 3 or 4) on the compositions and distribution of 5p metal atoms in the cluster shells of endohedral species [M@Sn_{14-x}Sb_x]^q− (M = La, Ce, or U; x = 6–8; q = 3, 4) was studied by means of combined experimental and quantum chemical investigations. While all known f-block metal ion-centered endohedral clusters possessed combinations of larger main group metal atoms so far (Sn/Bi or Pb/Bi), resulting in mixtures of 13- and 14-atom cages, the 14-atom cages reported herein comprise exclusively Sn and Sb atoms and therefore are challenged in accommodating the large 4f and 5f ions. We show that the clusters form in reactions of (Sn,Sb)^2− anions with [Ln(C₅Me₅H)₅] or [U(C₅Me₅H)₅]Cl, and that salts of [La@Sn₈Sb₄]^{7−}, [La@Sn₈Sb₃]^{4−}, [U@Sn₆Sb₆]^{4−}, and [U@Sn₇Sb₃]^{−} can be isolated from them. The assignment of Sn versus Sb in the encapsulating cage follows a simple rule. Different central atoms cause only slight differences in this regard and with respect to distortions of the cluster shells. The reactions also yielded the salt of the new binary anion (Sn₈Sb₄)^2− that was recently predicted by quantum chemical studies.

INTRODUCTION

The generation and in-depth investigation of binary and ternary intermetalloid clusters, in which a metal atom of the d-block or f-block is encapsulated by a cage of p-block metal atoms, have been two of the primary topics of metal cluster research for approximately two decades. Ternary clusters in particular have been intensively studied by research groups worldwide because of the sheer endless variations of elemental combinations and structures, and the uncommon electronic properties resulting from their structural diversity. The comprehensive studies have revealed a large variety of bonding patterns, and clusters with combined d/p-block elemental combinations have meanwhile also been successfully used for catalytic reactions.

However, most studies in this field have focused on clusters with d-block metal atoms in the interstitial position, while studies of the chemistry and physics of clusters with endohedral lanthanide or actinide atoms have remained scarce. The first intermetalloid cluster comprising an f-block metal ion within a cage of main group metal atoms was reported in 2011, [Eu@Sn₆Bi₆]^{4−}. The corresponding [K(crypt-222)]^{+} (crypt-222: 4,7,13,16,21,24-hexaaza-[1,10-diazabicyclo[8.8.8]-hexacosane) salt exhibited mainly ionic interactions between the inner Eu^{2+} ions and the binary cage. This was also proven by magnetic measurements indicating an unchanged S = 7/2 ground state of the Eu^{2+} ion.

Studies following this first report comprised comprehensive investigations of compounds with binary cluster shells of either Sn and Bi, Pb and Bi, Ga and Bi, or In and Bi atoms. The isolation of compounds with 13-atom shells, [Sm@Ga₃H₁₂Bi₁₀]^{3−} (x = 0 or 1), [La@Pb₂₃Bi₄]^{7−} (Ln = La, Ce, Nd, Gd, Sm, or Tb; x/q = 9/4 or 10/3), [Ln@Sn₆Bi₆]^{4−} (Ln = La or Ce), [La@In₃Bi₃]^{4−}, and [U@Pb₄Bi₄]^{10−} indicated significantly lower yields for Sn/Bi and Ga/Bi elemental combinations. Larger cages in contrast, especially those comprising Pb and Bi atoms, are exceptionally well suited for accommodation of the large f-block metal ions. Quantum chemical studies were performed to further explore this observation, which confirmed that smaller cages can face steric problems.

To date, the only f-block metal-centered clusters with a purely 5p-block metal environment have been [Ln@Sn₁₂]^{3−} clusters (Ln = La, Y, Ho, Er, or Lu), which are structurally related to their 5f/Bi analogues [An@Bi₁₂]^{q−} (An/q = Th/4 or U/3). However, these clusters do not exhibit closed cluster shells but rather a doughnut-like 12-atom polycycle surrounding the inner ions. We were therefore interested in exploring the possibility of enclosing 4f or 5f metal ions within closed shells of smaller p-block metal atoms and for this chose the Sn/Sb combination for the first time.

Here, we report the success of this study for different synthetic approaches, namely, reactions of (a) K₄SnSb₄ and...
Preparation of \([\text{Bu}_3\text{P}](\text{LaSnSb})\) \(2\). \(\text{K}_2\text{SnSb}\) \((51 \text{ mg}, 0.16 \text{ mmol})\), \([\text{Bu}_3\text{P}]\text{Br}\) \((109 \text{ mg}, 0.35 \text{ mmol})\), and \([\text{La}(\text{C}_5\text{H}_7\text{Me})_3]\) \((27 \text{ mg}, 0.054 \text{ mmol})\) were combined in a Schlenk tube and dissolved in 3 \(\text{mL}\) of \(\text{en}\). After the mixture had been stirred for 10 min at room temperature, the dark red solution was sonicated for 1 h. Six milliliters of toluene was added, and the solution was stirred for 19 h. Then, it was filtered through a 0.45 \(\mu\text{m}\) PTFE syringe filter. The black solution was stirred for an additional 3 h at 60 °C, filtered again through a 0.45 \(\mu\text{m}\) PTFE syringe filter, layered with 8 \(\text{mL}\) of toluene, and finally stored at 5 °C. After 2 months, small black, block-like crystals (yield of <5%) were present at the bottom of the Schlenk tube.

Preparation of \([\text{Bu}_3\text{P}](\text{CeSnSb})\) \(3\). \(\text{K}_2\text{SnSb}\) \((51 \text{ mg}, 0.16 \text{ mmol})\), \([\text{Bu}_3\text{P}]\text{Br}\) \((109 \text{ mg}, 0.35 \text{ mmol})\), and \([\text{Ce}(\text{C}_5\text{H}_7\text{Me})_3]\) \((27 \text{ mg}, 0.054 \text{ mmol})\) were combined in a Schlenk tube and dissolved in 4 \(\text{mL}\) of \(\text{en}\). After the mixture had been stirred for 10 min at room temperature, the dark red solution was sonicated for 1 h. Six milliliters of toluene was added, and the solution was stirred for 20 h. Then, it was filtered through a 0.45 \(\mu\text{m}\) PTFE syringe filter. The black solution was stirred for an additional 3 h at 60 °C, filtered again through a 0.45 \(\mu\text{m}\) PTFE syringe filter, layered with 8 \(\text{mL}\) of toluene, and stored at 5 °C. After 3 months, large black, block-like crystals (yield of <5%) were present at the bottom of the Schlenk tube.

### EXPERIMENTAL SECTION

#### General

All syntheses were performed in the absence of air and moisture using standard Schlenk or glovebox techniques. Solvents and DMF were distilled from CaH\(_2\) and stored over 3 Å molecular sieves. Toluene (tol, Acros Organics, 99%) was distilled from a sodium–potassium alloy and stored over 3 Å molecular sieves. Crypt-222 and \([\text{Bu}_3\text{P}]\text{Br}\) were dried in vacuum for at least 18 h. \(\text{K}_2\text{SnSb}\) was prepared by stoichiometric fusion of the elements at 950 °C for 48 h in a niobium tube, sealed within an evacuated silica ampule according to the literature. \([\text{K}([\text{crypt}-222])\text{[Sn(Sb)]}_{2}\text{en}]\) was precipitated from a 2:1 m solution of crypt-222 and \(\text{K}_2\text{SnSb}\) upon addition of 5 equiv of toluene. \(\text{K}_2\text{SnSb}\) was prepared according to the literature. The complexes \([\text{Ln}(\text{C}_5\text{H}_7\text{Me})_3])\) \((\text{Ln} = \text{La} or \text{Ce})\) and \([\text{U}(\text{C}_5\text{H}_7\text{Me})_3]\text{Cl}\) were synthesized as reported previously.\(^{17,19}\)

Preparation of \([\text{K}([\text{crypt}-222])\text{[La@Sn(Sb)]}_{2}\text{en}]\) \((103 \text{ mg}, 0.075 \text{ mmol})\). \([\text{K}([\text{crypt}-222])\text{[La(Sn)]}_{2}\text{en}]\) \((27 \text{ mg}, 0.054 \text{ mmol})\) were combined in a Schlenk tube and dissolved in 2.5 \(\text{mL}\) of DMF. After the mixture had been stirred for 18 h at 60 °C, a dark brown powder precipitated from the dark red solution upon addition of 24 \(\text{mL}\) of toluene. The precipitate was washed three times with 5 \(\text{mL}\) of toluene. Then, it was redissolved in 2 \(\text{mL}\) of a 1:1 DMF/\(\text{en}\) mixture and filtered through a 0.45 \(\mu\text{m}\) PTFE syringe filter. The black solution was layered with 6 \(\text{mL}\) of toluene and finally stored at 5 °C. After 2 months, small black, block-like crystals (yield of <5%) were present at the bottom of the Schlenk tube.

Another possibility is the reaction of \(\text{K}_2\text{SnSb}\) \((115 \text{ mg}, 0.125 \text{ mmol})\), crypt-222 \((380 \text{ mg}, 1.00 \text{ mmol})\), and \([\text{La}(\text{C}_5\text{H}_7\text{Me})_3])\) \((82 \text{ mg}, 0.163 \text{ mmol})\) in a mixture of 4 \(\text{mL}\) of \(\text{en}\) and 2 \(\text{mL}\) of toluene for 4.5 h at room temperature with stirring. The dark brown reaction mixture was filtered through a standard glass frit. The resulting reddish brown solution was layered with 4 \(\text{mL}\) of toluene. After 3 months, small black, block-like crystals (yield of <5%) were present in the wall of the Schlenk tube.

Single-Crystal X-ray Diffraction. Crystals suitable for X-ray diffraction analyses were investigated with a Bruker D8 Quest diffractometer at 100 K using Mo K\(_\alpha\) radiation \((\lambda = 0.7073 \AA; 1–3)\) or with a STOE Stadi Var diffractometer at 100 K using Cu K\(_\alpha\) radiation \((\lambda = 1.54186 \AA; 4 \& 5)\). Upon multiscan absorption correction (APEX II for 1–3 and STOE LANa for 4 and 5), the structure solution was performed by dual-space methods, followed by full-matrix least-squares refinement against F\(_2\), using SHELXTL, SHELXTL\(_8/3\), SHELXL\(_{11}\), SHELXL\(_{18/3}\), and OLEX2 software.\(^{10–23}\) Tables S4–S8 summarize the crystallographic data and
refinement results. Supplementary structural images are shown in Figures S17–S26.

Quantum Chemical Investigations. Density functional theory (DFT) calculations were performed with TURBOMOLE28 using the TPSS functional29 and employing def2-TZVP basis sets30 for Sn, Sb, Pb, and Bi, together with Dirac–Fock effective core potentials.31 For La, Ce, and U, Wood–Boring type effective core potentials were used12,32 together with segmented contracted basis sets from the same references with slight modifications in the g function space, available as “def2-TZVP” bases from ref 35. The conductor-like screening model36 was used for charge compensation.

RESULTS AND DISCUSSION

Figure 1 shows the molecular structures of the cluster anions within compounds 1–4. Compound 1 crystallizes in the monoclinic space group P2₁/n with four formula units in the unit cell. Compounds 2 and 3 crystallize in monoclinic space group C2/c with one formula unit in the unit cell. Compound 4 crystallizes in tetragonal space group P4₁/bc with one formula unit in the unit cell (see Tables S4–S8 and Figures S17–S26).

As indicated in the Introduction, the observation of salts that comprise only one of the described cluster sizes, such as those observed in 1–4, is relatively rare and has so far been widely restricted to compounds with 13-atom cages, which comprised the smallest Ln³⁺ ions or group 13/15 shells.16,19,20 In contrast to the more common mixtures of 13- and 14-atom cages, which are observed for intermetallic clusters with Sn/Bi or Pb/Bi shells encapsulating endohedral lanthanide ions (see above),15,18 the clusters reported herein form exclusively 14-atom cages, which we ascribe to the smaller atomic radii of Sn and Sb. A similar steric reason caused the exclusive crystallization of the only other salt of an exclusive 14-atom cluster anion with an elemental combination of Eu³⁺/Sn/Bi, because of the larger ionic radius of the Eu²⁺ ion formed by in situ reduction,35 and the crystallization of two isostructural clusters with other elemental combinations comprising much smaller ions and atoms throughout, [M@Ge₄As₄]³⁻ (M = Nb or Ta).37

The 14-atom cage that is observed in these clusters represents a deltahedral polyhedron with nine faces (six pentagonal ones and three square ones).38–41 It can be constructed by cutting the three equatorial corners of a trigonal bipyramid, which indicates its high symmetry, as a rationale for its observation instead of a deltahedral 14-atom Frank–Kasper polyhedron, for instance. Due to its high, almost spherical symmetry (disregarding the occupation of the corners by different types of atoms), this topology apparently enjoys exceptional thermodynamic stability. However, although they are isostructural and isoelectronic (with 70 = Sn valence electrons, where n = 14),13,42 the cluster shells of compounds 1–4 exhibit different ratios of the Sn and Sb atoms, which is required to adopt the overall charge of the cluster anion to the preferred amount of three [K(crypt-222)]⁺ counterions or four [K(crypt-222)]⁺ or [“Bu₃P”]⁺ counterions, respectively, for crystallization. As the interstitial cations possess (formal) charges of 3+ (La or Ce) or 4+ (U), different amounts of formally Sn²⁻ or Sb³⁻ (pseudo)group 15 atoms are thus required in the 14-atom shell to afford a total cluster charge (q) of 3− or 4−. For La³⁺ or Ce³⁺, a Sn:Sb composition of 6:8 or 7:7 corresponds to the observed number of counterions in 1–3, as confirmed by μ-XPS measurements (Table S1 and Figures S1–S3). The uranium cluster in 4 exhibits a 1:1 mixture of cluster anions of both charges, and hence of clusters with a 7:7 and an 8:6 composition of Sn and Sb atoms, also in agreement with elemental analysis (Table S1 and Figure S4). Given these different compositions of the cluster shells, we were interested in seeing whether there are certain patterns in the distribution of the two types of 5p metal atoms at the 14 atomic positions.

The stability of this cluster architecture and the occupation of the corners with either Sn and Sb atoms when the interstitial ion is La³+, Ce³⁺, or U⁴⁺ were thus investigated by means of DFT (for computational details, see the Experimental Section). For a first impression of the potential energy surface, we calculated single-point energies for all distributions of Sn and Sb in (LaSn₅Sb₇)³⁻ that are symmetry-distinct in a D₁₅-symmetric topology of identical atoms. As listed in Table S10, the four most favorable isomers are found within an energy range of 10 kJ/mol, while the fifth is 17 kJ/mol above the most energetically favorable cluster. The energetically following 237 isomers are densely distributed between 28 and 162 kJ/mol, and the five most disfavored are between 168 and 185 kJ/mol. Common for the Sn/Sb distributions of the four energetically most favorable isomers is the occupation of the top and bottom positions (1 and 14, respectively, in Figure 1a) by Sn, and the occupation of at least two positions in each of the three equatorial rings by Sb, which leads to the maximum possible number of heteroatomic bonds, 16. Next, we treated the anions in 1–4 and, for comparison, also the corresponding clusters with a Pb/Bi shell (note that [La@Pb₅Bi₄]⁻) was computed earlier and served to predict the experimentally found species35). All of them have 42 electrons in the valence p orbitals of the 14 shell atoms that form 21 two-electron, two-center bonds, which is evident from a Boys localization procedure. For the targeted search of the best p element distributions, we applied a first-order perturbation theory in the nuclear charge to [M@Tt/Pn]⁻ (Tt/Pn = Sn/Sb or Pb/Bi; M = La/4, Ce/4, or U/3), [M@Sn₅Sb₇]³⁻ (M = La or Ce), and [U@Sn₅Sb₇]³⁻ (see Figure 2).

All obtained structures follow the distribution scheme formulated above. Furthermore, for the 7:7 composition, the most favorable distribution, (7:7)⁷, is the same for M = La and Ce and for Tt/Pn = Sn/Sb and Pb/Bi; the most favorable distribution for [U@Sn₅Sb₇]³⁻, (7:7)⁹, is very similar.
Consequently, the energy differences (for optimized structure parameters) between the two distributions are very small (see Table 1). For Ce and La, the most favorable isomers for a 6:8 composition are achieved by replacing the Sn atom at position 10 (see Figure 1b) in (7:7)I with Sb, and the most favorable distribution for [(U@Sn,Sb)x]4+ is achieved by replacing the Sb atom at position 12 (see Figure 1b) in (7:7)II with Sn. The same replacements for the almost isoenergetic isomers with higher energies with a 7:7 composition yield again almost isoenergetic isomers with compositions of 6:8 (La and Ce) and 8:6 (U). Thus, for 14-atom Tf/Pn cluster shells with 42 electrons in the valence p orbitals of the shell atoms, the distribution rule presented above, "Tf at top and bottom position, at least two Pn in each of the equatorial rings", is robust, regardless of the elemental composition, regardless of whether the f occupation of the central ion is 5f0(La), 5f1(Ce), or 5f2(U), and regardless of whether a lanthanide or an actinide resides in the center. As summarized in Table S11, the distances within the (Sn,Sb) cluster shells are subject to notable changes for different interstitial ions though, indicating the potential of these cages to "breathe" to a certain extent.

During the synthesis of compound 4, single crystals of compound 5 were isolated after shorter crystallization times (1 day instead of 2 weeks). It comprises binary [Sn,Sb]4− isomers (see Figure 3a,b), the architecture of which does not indicate a direct structural relationship with the cluster observed in 4. Hence, most likely, it is not an intermediate on the way toward the uranium-centered 14-atom cage in 4 but rather represents the product of a parallel pathway of cluster growth, which stops at a convenient charge for crystallization with [K(crypt-222)]+ counterions (2− here). The composition of the anion, again not available through X-ray diffraction alone, was confirmed by μ-XFS measurements (Table S1 and Figure S5).

Notably, exactly this [Sn,Sb]4− anion was recently predicted to exist by quantum chemical calculations in the context of another binary eight-atom anion comprising Sn and Sb atoms, ([Sn,Sb]3)−, which was obtained from K2SnSb and [4-Bu4P]+Br in the absence of a Lewis-acidic f-block metal ion.27 As observed for the f metal atom-centered clusters described above, the smaller [4-Bu4P]+ cation tends to crystallize with similarly sized, more highly charged anions than do the larger [K(crypt-222)]+ cations (e.g., 4− in compound 2 vs in 3− in compound 1). Electrospray ionization mass spectrometry (ESI-MS) measurements of crystals of 5 (Figures S12−S16) indicate significant rearrangement processes in solution that also afford an anionic {Sn,Sb} fragment (detected as a monoanion, as usual), which obviously does not crystallize in the presence of [K(crypt-222)]+ cations though. Both binary anions are isoelectronic and possess the same topologies of a 2n + 6 arachno type cage. The most energetically favorable isomers of both can be derived from a bicapped square antiprism, but the calculated structure with the lowest energy for (Sn,Sb)4− is slightly distorted as compared to the more Sn-rich analogue, indicating the subtle differences between these two types of neighboring atoms and the limits of the pseudoelement concept at this point. The energetically most favorable isomers of the two anions as calculated with DFT methods are shown in panels c and d of Figure 3, respectively.

**CONCLUSION**

In summary, with this study, we improve our understanding of the family of intermetalloid clusters with 14-atom cages by presenting the first examples with purely 5p metal (Sn/Sb) cluster shells. As shown by quantum chemistry, interstitial Ln4+ or U4+ ions do not significantly affect the preferences of Sn and Sb atoms for occupying certain atomic positions in the cluster shell. Because of mainly ionic interactions, the relative stabilities of isomers with different Sn and Sb atom assignments are dominated by the stability of the respective main group cluster shells. However, small structural changes of the cluster shells are notable for the different interstitial...

**Figure 2.** Energetically most favorable isomers found by perturbation theory in the nuclear charge. Arabic numerals indicate the Sn:Sb ratio (6:8, 7:7, or 8:6), and Roman numerals refer to the cluster type, I or II. Differences are colored green. For relative energies, see Table 1.

**Table 1.** Relative Energies of the Energetically Most Favorable Isomers of [M@SnSbx−]x[−] [M = La, Ce, or U; x = 6−8 (see Figure 2)] in Kilojoules per Mole

<table>
<thead>
<tr>
<th>M</th>
<th>q</th>
<th>Sn/Sb</th>
<th>Pb/Bi</th>
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<tr>
<td></td>
<td></td>
<td>I</td>
<td>II</td>
</tr>
<tr>
<td></td>
<td>7:7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>La</td>
<td>−4</td>
<td>0.1</td>
<td>0.0</td>
</tr>
<tr>
<td>Ce</td>
<td>−4</td>
<td>0.2</td>
<td>0.0</td>
</tr>
<tr>
<td>U</td>
<td>−3</td>
<td>1.8</td>
<td>2.4</td>
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<tr>
<td></td>
<td>6:8</td>
<td></td>
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</tr>
<tr>
<td>La</td>
<td>−3</td>
<td>2.0</td>
<td>3.3</td>
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<tr>
<td>Ce</td>
<td>−3</td>
<td>2.1</td>
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<td>U</td>
<td>−4</td>
<td>2.0</td>
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**Figure 3.** (a and b) Molecular structures of the (Sn,Sb)x−−anion in S in two views (thermal ellipsoids at 50% probability), with selected interatomic distances: Sn/Sb1−Sn/Sb2/3/4, 2.8296(5)−2.8569(5) Å; Sn/Sb2/3/4−Sn/Sb1/2/3/4, 3.2517(6)−3.3262(1) Å; Sn/Sb2/3/4−Sn/Sb5/6/7/8, 2.8322(4)−3.0548(4) Å; Sn/Sb5/6/7/8−Sn/Sb5/6/7/8, 2.7861(5)−2.8468(5) Å. As in Figure 1, all atoms are given as two-color octants, with the dominant color indicating the atom type that is more likely on the respective atomic position according to the DFT studies. The numbers refer to the respective atom labels and are also given in the color of the more probable atom at this position. (c) Calculated structures of the energetically most favorable isomer of the binary anion (Sn,Sb)x−. (d) Calculated structure of the energetically most favorable isomer of the binary anion (Sn,Sb)x− for comparison.
cations. The isolation of a second compound from the reaction with the uranium complex comprising a novel binary anion, \((\text{Sn}_x\text{Sb}_y)^{2-}\), provides important insight into the variance of the energy hypersurface in the reaction space of Sn/Sb-based Zintl anions and its diversity in the presence or absence of Lewis-acidic metal ions. At the same time, it confirmed a previously made theoretical prediction about the existence of this anion. Future studies will aim to use this new anion as a starting point for new ternary clusters.

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**Notes**

The authors declare no competing financial interest.

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