Review

Silver and Copper Complexes with closo-Polyhedral Borane, Carborane and Metallacarborane Anions: Synthesis and X-ray Structure †

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† Dedicated to the memory of Dr. Natalia Votinova, a boron chemist and our colleague and friend.


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1. Introduction

From the days of alchemists, the main goal of chemistry has been understanding the behavior of molecules and their construction from the constituent atoms. In recent years, this goal has been extended beyond atomic and molecular chemistry into the realm of supramolecular chemistry controlled mainly by weak bonding. In contrast to molecular chemistry, which is predominantly based upon the covalent bonding of atoms, supramolecular chemistry is based upon intermolecular interactions, i.e., on the association of two or more building blocks, which are held together by intermolecular bonds. In molecular chemistry, strong association forces such as covalent and ionic bonds are used to assemble atoms into discrete molecules and hold them together. In contrast, the forces used to organize and hold together supramolecular assemblies are weaker noncovalent interactions (typically on the order of 1–5 kcal/mol), such as hydrogen bonding, polar attractions, van der Waals forces, hydrophilic-hydrophobic interactions and charge transfer interactions [1,2].

Among the systems that are held together by weak interactions are compounds containing reactive electrophilic p-block cations and transition metal complex cations. As a rule, such cations can be stabilized by so-called “non-coordinating anions”, the term which was first coined to describe anions such as ClO4−, SO3CF3−, BF4−, PF6−, AsF6− and SbF6−. These anions usually are non-coordinating in aqueous solution, however with development of routine X-ray crystallography it has become
evident that in many cases a “non-coordinating” complex anion can in fact easily coordinates to a metal in the solid state. Accordingly, the term “weakly coordinating anions” was proposed for the coordination for these anions. Due to capability of such anions to stabilize reactive electrophilic cations, their use as counter anions is of significant interest in both synthesis and catalysis. Owing to the importance of weakly coordinating anions in fundamental and applied chemistry, there is growing interest in synthesis of “the least coordinating anions”. Polyhedral boranes and carboranes ([B\textsubscript{10}H\textsubscript{10}]\textsuperscript{2-}, [B\textsubscript{12}H\textsubscript{12}]\textsuperscript{2-}, [1-CB\textsubscript{9}H\textsubscript{10}]\textsuperscript{-}, [CB\textsubscript{11}H\textsubscript{12}]\textsuperscript{-}, [3,3'-Co(1,2-C\textsubscript{2}B\textsubscript{9}H\textsubscript{11})\textsubscript{2}]\textsuperscript{-} (Scheme 1)) and especially their halogen derivatives are usually considered as weakly coordinating anions that are able to stabilize highly reactive electrophilic \textit{p}-block cations and transition metal complex cations [3–6].

![Scheme 1. Anionic polyhedral boron hydrides discussed in this review.](image)

One of the most common ways to obtain compounds with stabilized reactive cations are metathesis reactions of relatively weak bonded halide complexes with univalent salts of weakly coordinating anions M[X]. In many cases, Ag\textsuperscript{+} is the best cation to abstract halide from substrate and the silver salt metathesis is a time-honored method of halide ion abstraction. Its origin dates back to the early days of coordination chemistry when silver nitrate was established as the standard testing reagent to differentiate free and complexed chloride. More recently, the availability of numerous silver salts of weakly coordinating anions, Ag[X] (X = ClO\textsubscript{4}\textsuperscript{-}, SbF\textsubscript{6}\textsuperscript{-}, etc.), have exceeded silver salt metathesis as the widely used method of halide ion abstraction. The polyhedral borane and carborane anions owing to their large size and relative low charge can be considered soft bases. On the other hand, Ag\textsuperscript{+} and Cu\textsuperscript{+} cations are typical soft acids that should result in formation of great number of complexes of various structural types [7].

In coordination chemistry, silver(I) is able to adopt a wide variety of coordination geometries with coordination numbers varying from 2 to 8. This variety is mainly due to the lack of stereoechemical preference that arises from the d\textsuperscript{10} configuration. Furthermore, the weak nature of the silver-ligand bond means that in the solid state, weak interactions and crystal packing forces have a greater influence on structure than may be expected in more robust metal–ligand systems [8–10]. Copper(I) has the same d\textsuperscript{10} configuration and its coordination chemistry in some cases resembles coordination chemistry of silver(I) [11]. The availability of a number of silver and copper salts makes it easy to analyze the effects counter-anions have on the structure of supramolecular networks, such as their ability to coordinate to the metal center. Therefore, synthesis and X-ray structure studies of silver and copper salts of new types of weakly coordinating anions are of great importance. Several review papers concerning different aspects of chemistry of silver and copper complexes of polyhedral boranes and carboranes were published during the recent years [12–14].

Herein, we present a review of chemistry of silver and copper salts and complexes with polyhedral borane and carborane anions with a special emphasis on their solid state structure.
2. Copper and Silver Complexes with the closo-Decaborate anion \([\text{B}_{10}\text{H}_{10}]^{2-}\)

The closo-decaborate anion \([\text{B}_{10}\text{H}_{10}]^{2-}\) was the first representative of the closo-borane family, which was reported by Hawthorne and Pitochelli in 1959 [15]. The \textit{closo-}decaborate anion has structure of bicapped square antiprism with two different types of boron atoms (five-coordinated apical \(B_a\) and six-coordinated equatorial \(B_e\)) that makes some aspects of its chemistry unique and having no analogues in the chemistry of icosahedral boron hydrides. One of these peculiarities is the possibility to form various coordination isomers. Despite rather rapid development of its chemistry [16,17], the coordination chemistry of the \textit{closo-}decaborate anion was poorly understood until the last fifteen years, when a number of studies which cast light on this subject were published.

Copper(I) \textit{closo-}decaborate \([\text{Cu}_2[\text{B}_{10}\text{H}_{10}]]\) was the first crystallographically characterized salt with the \textit{closo-}decaborate anion [18]. The complex was prepared by reaction of \(\text{K}_2[\text{B}_{10}\text{H}_{10}]\) with copper(II) acetate in water. According to single crystal X-ray diffraction data, copper atoms are arranged near the centers of opposite polyhedral faces providing each boron cage to be surrounded by four copper atoms with the \(\text{Cu} \ldots \text{B}\) distances varying from 2.06 to 2.33 Å (Figure 1) [18]. Based on the X-ray data, the formation of three-centered two-electron Cu-H-B bonds between copper and the boron cage was suggested. This suggestion was supported by the IR spectrum wherein two absorption bands in the B-H stretching region are observed: one of them (with maximum near 2450 cm\(^{-1}\)) was assigned to non-coordinated BH groups and the other (with maximum at 2300 cm\(^{-1}\)) was assigned to the BH groups involved in the Cu-H-B interactions [19].

![Figure 1](image-url). The coordination sphere of the \([\text{B}_{10}\text{H}_{10}]^{2-}\) anion (a) and crystal packing (b) in the structure of \([\text{Cu}_2[\text{B}_{10}\text{H}_{10}]]\)\(_n\). Reproduced with permission from reference [18]. Copyright © (1962) AIP Publishing LLC.

The similar silver(I) salt \(\text{Ag}_2[\text{B}_{10}\text{H}_{10}]\) can be prepared by metathetical reactions in aqueous solution [20,21]. Its structure was not determined, however the existence of Ag-H-B interactions in the solid state was supported by the IR spectroscopy data [21].

In addition to the three-dimensional polymeric binary complexes \([\text{M}_2[\text{B}_{10}\text{H}_{10}]]\), the \textit{closo-}decaborate anion is able to form one-dimensional chain polymers of the \(\text{Cat}[\text{M}\text{B}_{10}\text{H}_{10}]\) type both with \(\text{Cu}^+\) and \(\text{Ag}^+\) cations. The complexes are formed at equimolar amounts of \(\text{Cat}_2[\text{B}_{10}\text{H}_{10}]\) and copper(II) and silver(I) salts, whereas in the case of two-fold excess of metal salts or in the presence of small cations (e.g., \(\text{NH}_4^+\)) formation of binary complexes \([\text{M}_2[\text{B}_{10}\text{H}_{10}]]\) takes place. It is worth noting that copper(I) complexes \(\text{Cat}[\text{Cu}\text{B}_{10}\text{H}_{10}]\) are formed from copper(II) salts due to reductive properties of the decahydro-\textit{closo-}decaborate anion. As a result, copper(I) complexes \(\text{Cat}[\text{Cu}\text{B}_{10}\text{H}_{10}]\) (where \(\text{Cat}^+ = \text{K}^+, \text{Cs}^+, \text{R}_2\text{NH}_2^+, \text{R}_3\text{NH}^+, \text{R}_4\text{N}^+\) (R = Me, Et, Bu), \(\text{Ph}_4\text{P}^+, \text{NaphCH}_2\text{PPh}_3^+, \text{Ph}_4\text{As}^+\)) [22–25] and
silver(I) complexes Cat[Agl[B10H10]] (where Cat = Cs+, R2NH2+, R3NH+, R4N+ (R = Me, Et, Pr, Bu)) [21] were prepared. IR spectra of all these compounds demonstrate characteristic bonds of M-H-B stretching at 2150–2400 cm⁻¹ due to interactions of Cu⁺ and Ag⁺ cations with the closo-decaborate anion.

In the copper(I) complex [Cs(Cu[B10H10])]ₙ the Cu⁺ cation is in distorted tetrahedral arrangement formed by two bridging closo-decaborate anions which are coordinated through Bₐ⁻Bₑ edges. The Cu…B₁ distances (2.159(6) Å) are significantly shorter than the Cu…Bₑ distances (2.228–2.287(6) Å). The binding of Cu⁺ by closo-decaborate edges of different types results in formation of zigzag chains running along the c axis (Figure 2a) [25,26]. In the silver(I) complex [Cs(Ag[B10H10])]ₙ the closo-decaborate anion is coordinated to copper cations through two Bₐ⁻Bₑ edges with a shared vertex (Ag…B distances of 2.645(4) and 2.593(6) Å) to form zigzag chains. The chains are connected into layers owing to additional contacts with BH groups of the neighboring [B10H10]²⁻ anions (Ag…B distances of 3.183(6) and 3.337(6) Å) (Figure 2b) [21].

Figure 2. Fragments of polymeric chains in [Cs(Cu[B10H10])]ₙ (a) and [Cs(Ag[B10H10])]ₙ (b).

In the structure of [Me₂NH₂(Cu[B10H10])]ₙ [25,26], the copper cation has a distorted tetrahedral environment formed by two bridging closo-decaborate anions which are coordinated through Bₐ⁻Bₑ edges resulting in the formation of straight [(Cu[B10H10])ₙ] chains that run along the b axis. In both independent chains, the polyhedra that coordinate the Cu atoms by the B(1)-B(2) and B(7)-B(10) edges alternate with those coordinating Cu atoms by the B(1)-B(2) and B(8)-B(10) edges. The Cu…Bₑ and Cu…Bₐ bond lengths substantially overlap (2.145–2.285(9) and 2.130–2.268(9) Å, respectively). The symmetrically related chains are packed into layers parallel to the bc plane. The layers alternate along the a axis. The [Me₂NH₂]⁺ cations are located between the anionic layers and form the NH…HB dihydrogen bonds with the polyhedral anions (Figure 3) [26].

Figure 3. Cont.
Comparison of structures of \{(\text{Et}_3\text{NH})(\text{M}[\text{B}_{10}\text{H}_{10}])\}_n\) (\text{M} = \text{Cu} \[22\] or \text{Ag} \[21\]) revealed that different polymeric chains are formed in these two complexes. In the copper(I) complex, the polymeric chain is formed by coordination of the boron cage to a copper atom via \(\text{B}_a-\text{B}_e\) and \(\text{B}_e-\text{B}_e'\) edges (\(\text{Cu} \cdots \text{B}_a 2.172\ \text{Å}, \text{Cu} \cdots \text{B}_e 2.213(4)–2.234(4)\ \text{Å}\) (Figure 4a). The \(\text{Et}_3\text{NH}^+\) cations form the \(\text{NH} \cdots \text{HB}\) dihydrogen bonds with the polyhedral anions (\(\text{NH} \cdots \text{HB} 2.20(6)–2.92(6)\ \text{Å}\)). In the silver(I) complex the metal atom is coordinated by two \textit{closo}-decaborate anions via \(\text{B}_a-\text{B}_e\) edges and one \textit{closo}-decaborate anion via \(\text{B}_e-\text{B}_e'\) edge (\(\text{Ag} \cdots \text{B} 2.561(5)–2.907(5)\ \text{Å}\)) (Figure 4b). Similar to the copper(I) complex, the \(\text{Et}_3\text{NH}^+\) cations in \{(\text{Et}_3\text{NH})(\text{Ag}[\text{B}_{10}\text{H}_{10}])\}_n\) form the \(\text{NH} \cdots \text{HB}\) dihydrogen bonds with \textit{closo}-decaborate anions.

\begin{figure}[ht]
\centering
\includegraphics[width=\textwidth]{figure3.png}
\caption{Fragment of X-ray structure (a) and crystal packing (b) of \{(\text{Me}_2\text{NH}_2)(\text{Cu}[\text{B}_{10}\text{H}_{10}])\}_n. Reproduced with permission from reference \[26\]. Copyright © (2003) Springer.}
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The trigonal-planar environment of copper cation is formed by acetonitrile nitrogen atoms. Two elongated contacts with the BH groups supplements the Cu\(^+\) coordination to trigonal bipyramidal (Cu…B(H), 3.319(7) Å; Cu…H(B), 2.40(7) Å) (Figure 6) [28].

In the structure of [(Ph\(_4\)P)(Cu[B\(_{10}\)H\(_{10}\)])\(_n\)] the copper cation is coordinated by two closo-decaborate anions through single B\(_e\)-H groups (Cu…B 2.160 and 2.187 Å) [27] forming infinite polymeric chain (Figure 5).

The unusual complex [{Cu(MeCN)\(_3\)}(Ag[B\(_{10}\)H\(_{10}\)])\(_n\)] was prepared by the reaction of Cu\(_2\)[B\(_{10}\)H\(_{10}\)] and Ag\(_2\)[B\(_{10}\)H\(_{10}\)] in acetonitrile in the presence of trifluoroacetic acid. The silver cation is coordinated by two closo-decaborate anions through B\(_e\)-Be edges (Ag…B\(_a\) and Ag…B\(_e\) bond lengths are 2.656(8) and 2.547(7) Å, respectively) and one closo-decaborate anion through B\(_e\)-B\(_e'\) edge (Ag…B\(_e\) 2.758(9) Å). The trigonal-planar environment of copper cation is formed by acetonitrile nitrogen atoms. Two elongated contacts with the BH groups supplements the Cu\(^+\) coordination to trigonal bipyramidal (Cu…B(H), 3.319(7) Å; Cu…H(B), 2.40(7) Å) (Figure 6) [28].
As it was demonstrated above, all the \([\text{Cat}([\text{M}[\text{B}_{10}\text{H}_{10}])]_n\) complexes in the solid state form polymeric chains with cations located between the chains. The discrete complex \((\text{Et}_3\text{NH})\{(\text{Cu}[1-\text{B}_{10}\text{H}_9\text{N}_2]_2)\) was prepared by the reaction of the charge-compensated diazonium derivative of the \(\text{closo}\)-decaborate anion \((\text{Et}_3\text{NH})[1-\text{B}_{10}\text{H}_9\text{N}_2]^-\) with \(\text{CuCl}\). The copper cation in this complex is coordinated by two \([1-\text{B}_{10}\text{H}_9\text{N}_2]^-\) anions through \(\text{B}(6)-\text{B}(10)\) edges that are the most distant from the substituent. The \(\text{Cu} \cdots \text{B}(6)\) and \(\text{Cu} \cdots \text{B}(10)\) distances are 2.184(9) and 2.168(8) Å, the \(\text{Cu} \cdots \text{H}(6)\) and \(\text{Cu} \cdots \text{H}(10)\) distances are 1.96 and 1.99 Å (Figure 7) [29].

Complex \([\text{Cu}^{II}_4(\text{OH})_4(2,2'\text{-Bipy})_4][\text{Cu}^{IV}_4\text{B}_{10}\text{H}_{10}]_3\) \(\cdot 4\text{(MeCN)}\) containing discrete \([\text{Cu}^{IV}_2\text{B}_{10}\text{H}_{10}]_3]^{4-}\) anions and \([\text{Cu}^{II}_4(\text{OH})_4(\text{Bipy})_4]^{4+}\) cations was prepared by the reaction of \(\text{Cu}_2\text{B}_{10}\text{H}_{10}\) with \(2,2'\text{-bipyridine}\) in refluxing acetonitrile in air [30,31]. In the anionic part three \(\text{closo}\)-decaborate anions are connected by two \(\text{Cu}^+\) cations into a linear \(\{\text{Cu}^{II}_2\text{B}_{10}\text{H}_{10}\}_3\}^{1-}\) oligomer (\(\text{Cu} \cdots \text{B}\) and \(\text{Cu} \cdots \text{H}(\text{B})\) distances are 2.139–2.205(8) and 1.69–2.01 Å, respectively). The \([\text{Cu}_4(\text{OH})_4(\text{Bipy})_4]^{4+}\) cation has a double-decker structure. Each of the \(\text{Cu}^{2+}\) ions is bound to a bidentate \(2,2'\text{-bipyridine}\) molecule and two or three bridging \(\text{OH}\) groups, which form a square (the \(\text{Cu}(1)\) and \(\text{Cu}(4)\) atoms) or a tetragonal pyramid (the \(\text{Cu}(2)\) and \(\text{Cu}(3)\) atoms). The \(\text{Cu}-\text{N}\) bond lengths are within 1.973–2.014(5) Å. The \(\text{Cu}-\text{O}\) bond lengths in the squares formed by the \(\text{Cu}(1)\) and \(\text{Cu}(4)\) atoms and in the bases of the pyramids...
formed by the Cu(2) and Cu(3) atoms are within 1.909–1.956 and 1.921–1.986 Å, respectively. The apical O atoms are at Cu-O distances of 2.342(4) and 2.384(4) Å from the Cu(2) and Cu(3) atoms, respectively. The BH groups of the terminal polyhedral anions of the oligomer complete the coordination of the Cu(1) and Cu(4) atoms to a square pyramid (4 + 1) (Cu . . . B(H), 3.089(7) and 3.175(7) Å; Cu . . . H(B), 2.17 and 2.33 Å) and that of the Cu(2) and Cu(3) atoms, to a square bipyramid (4 + 1 + 1) (Cu . . . B(H), 3.485(8) and 3.364(7) Å; Cu . . . H(B), 2.91 and 2.54 Å). Weak Cu . . . B(H) interactions connect the \([\text{Cu}_{4}(\text{OH})_{4}(\text{Bipy})_{4}]^{4+}\) complex cations and \([\text{Cu}_{2}[\text{B}_{10}\text{H}_{10}]_{3}]^{4-}\) complex anions into chains. The terminal \([\text{B}_{10}\text{H}_{10}]^{2-}\) anions of the oligomeric anion link the Cu$^{2+}$ and Cu$^{+}$ atoms forming Cu$^{2+}$···B$_{10}$H$_{10}$···Cu$^{+}$ bridges (Figure 8).

![Figure 8. Fragment of X-ray structure of \([\text{Cu}_{II4}(\text{OH})_{4}(\text{Bipy})_{4}]\text{[Cu}_{I2}[\text{B}_{10}\text{H}_{10}]_{3}]\cdot4(\text{MeCN})\). Reproduced with permission from reference [30]. Copyright © (2011) Springer.](image)

Solvent coordinated complexes of transition metals are widely regarded as useful starting materials in synthetic chemistry since their weakly coordinating solvent ligands, e.g., acetonitrile and tetrahydrofuran, can be easily replaced by more strongly coordinating ligands. In this way, they can be also applied as building blocks for inorganic and organometallic macromolecules.

Recrystallization of \([\text{Cu}_{2}[\text{B}_{10}\text{H}_{10}]]_{n}\) from acetonitrile solution in the presence of trifluoroacetic acid gave complex \({((\text{MeCN})_{2}\text{Cu})_{2}[\text{B}_{10}\text{H}_{10}])}\) with two \([(\text{MeCN})_{2}\text{Cu}]^{+}\) fragments coordinated to B(1)-B(2) and B(3)-B(6) edges [28] (Figure 9).
Complex \{[(\text{MeCN})_3\text{Ag}](\text{Ag}[2-\text{B}_{10}\text{H}_9\text{F}])\}_2\} was prepared by the treatment of (\text{Ph}_4\text{P})_2[2-\text{B}_{10}\text{H}_9\text{F}] with silver trifluoroacetate in 9:1 dichloromethane-toluene solution followed by crystallization from acetonitrile. The structure of the \{[(\text{MeCN})_3\text{Ag}](\text{Ag}[2-\text{B}_{10}\text{H}_9\text{F}])\}_2\} complex consists of dimeric \{\text{Ag}[2-\text{B}_{10}\text{H}_9\text{F}]\}_2^{2−}\) units weakly interacting with two nearly planar \{(\text{MeCN})_3\text{Ag}\}^+\) cations (Ag . . . B distances are 2.520–2.644(5) and 2.02–2.36(5) \(\text{Å}\), respectively) [32]. There is no interaction between the fluorine substituent in boron cage and either silver cation (Figure 10) [14].

Complex \{(\text{MeCN})_2\text{Cu})_2[\text{B}_{10}\text{H}_{10}]\}\). Complex \{(\text{MeCN})_2\text{Cu})_2[\text{B}_{10}\text{H}_{10}]\}\) was prepared by crystallization of \(\text{Ag}_2[\text{B}_{10}\text{H}_{10}]\) from \(\text{N},\text{N}\)-dimethylformamide [33]. The compound contains four types of silver atoms. The environment of the Ag(1) and Ag(4) atoms are formed by the O atoms of two DMF molecules and two BH groups of two closo-decaborate anions (Ag . . . B distances 2.555(10) and 2.646(12) \(\text{Å}\)). Two other BH groups form additional elongated bonds (Ag . . . B distances 2.891(11) and 2.929(10) \(\text{Å}\)). Although the environment of Ag(1) and Ag(4) have the identical composition, they differ in the structure (Figure 11a,b). The Ag(1)
atom forms short Ag...B contacts (2.613(11) and 2.696(11) Å) with one closo-decaborate anion and long Ag...B contacts (2.974(10) and 3.086(12) Å) with other one anion. The Ag(1) atom is coordinated by the B₃-B₄ and B₅-B₆ edges of the closo-decaborate anion, whereas the Ag(4) atom is coordinated only by B₃-B₆ edges. The environment of the Ag(2) and Ag(3) atoms includes only closo-decaborate anions (Figure 11c,d).

![Diagram](image)

Figure 11. The crystal environment of silver cations in structure of [Ag₂(DMF)[B₁₀H₁₀]ₙ]: (a) Ag(1); (b) Ag(4); (c) Ag(2); and (d) Ag(3). Reproduced with permission from reference [33]. Copyright © (2008) Springer.

Formation of polymeric complexes with the closo-decaborate anion can be avoided using ligands that can saturate effectively the coordination sphere of the metal atom. Taking in the account that tetrahedral environment is the most preferred both for copper(I) and for silver(I) complexes, it can be expected that the closo-decaborate anion should form discrete complexes of {(L₂M)₂[B₁₀H₁₀]} type (where M = Cu, Ag). Indeed, a large number of such complexes were synthesized and structurally characterized. The [L₂M]⁺ fragments have two free coordination places and are coordinated to the [B₁₀H₁₀]²⁻ anion through two BH groups of one edge in chelate-like fashion. Due to geometrical features of the closo-decaborate anion there are a total of 25 possible isomers for the {(L₂M)₂[B₁₀H₁₀]} complexes, 17 of which are chiral. Naturally, not all of these coordination modes can be realized due to the reaction ability of the boron clusters, steric hindrances of ligands, and other factors. Nevertheless, a number of these positional isomers were already obtained and structurally characterized.

Synthesis of the first complex of this type {((Ph₃P)₂Cu)₂[B₁₀H₁₀]} by the reaction of (Et₃NH)₂[B₁₀H₁₀] with [(Ph₃P)₃CuCl] in chloroform was described by Gill and Lippard in 1975 [34]. The copper atoms are in a quasi-tetrahedral environment formed by two phosphorus atoms of the Ph₃P ligands and two BH groups of the boron polyhedron. The closo-decaborate anion coordinates...
metal atoms by B₃-B₄ edges at different apical vertices of the boron polyhedron. Two enantiomers of \[((\text{Ph}_3\text{P})_2\text{Cu})_2[B_{10}H_{10}]\] were found in the crystal. In one enantiomer, coordination proceeds via B(1)-B(2) and B(6)-B(10) edges, whereas the other enantiomer—via B(1)-B(2) and B(9)-B(10) edges. The same compound was isolated by interaction of Cat₂[B₁₀H₁₀] (Cat⁺ = Bu₄N⁺, Ph₄P⁺) with \[((\text{Ph}_3\text{P})_3\text{CuCl})\] in acetonitrile [35]. Structure of the 1–2, 9–10 enantiomer of \[((\text{Ph}_3\text{P})_2\text{Cu})_2[B_{10}H_{10}]\] is presented in Figure 12a. The Cu . . . B bonds vary in the range from 2.260(9) to 2.317(9) Å, the Cu . . . HB distances range from 1.78 to 2.05 Å, the Cu-H-B angles are 109–122°. The existence of the Cu-H-B interactions in the solid state is supported by the presence of the Cu-H-B stretching band in the IR spectrum of \[((\text{Ph}_3\text{P})_2\text{Cu})_2[B_{10}H_{10}]\] at 2150–2400 cm⁻¹ [36]. The molecular conductance measurements indicate that the \[((\text{Ph}_3\text{P})_2\text{Cu})_2[B_{10}H_{10}]\] complex does not dissociate in dichloromethane solution [34].

Similar silver(I) complex 1–2, 6(9)–10 \[((\text{Ph}_3\text{P})_2\text{Ag})_2[B_{10}H_{10}]\] was prepared by the reaction of (Et₃NH)₂[B₁₀H₁₀] with \[((\text{Ph}_3\text{P})_3\text{Ag})\text{NO}_3\] in acetonitrile (Figure 12b). The geometrical parameters of MHB bond are: Ag . . . B and Ag . . . HB bonds are 2.561(4)–2.617(4) and 1.97–2.29 Å, respectively; the Ag–H–B angles are 90.0–105.3°. The Ag(1) . . . B bonds with the B-a-Be' edge (2.577(3) and 2.519(3) Å) are shorter than the Ag(2) . . . B

![Figure 12. Molecular structure of 1–2, 9–10 \[((\text{Ph}_3\text{P})_2\text{M})_2[B_{10}H_{10}]\]: M = Cu (a); M = Ag (b); schematic representation of cocrystallized 1–2, 6–10 and 1–2, 9–10 enantiomers (hereinafter view from the top of the boron cage) (c). Phenyl cycles are omitted for clarity. Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.](image-url)

Another positional isomer of \[((\text{Ph}_3\text{P})_2\text{Ag})_2[B_{10}H_{10}]\] was prepared by the reaction of (Et₃NH)₂[B₁₀H₁₀] with \[((\text{Ph}_3\text{P})_3\text{Ag})\text{NO}_3\] in the acetonitrile in the presence of trifluoroacetic acid. In the obtained \[((\text{Ph}_3\text{P})_2\text{Ag})_2[B_{10}H_{10}]\] complex (Figure 13), the \[B_{10}H_{10}^-\] anion is coordinated to silver atoms by B₃-B₄ edges at different apical vertices of the boron polyhedron. Two enantiomers of \[((\text{Ph}_3\text{P})_2\text{M})_2[B_{10}H_{10}]\] are shown in Figure 12c.
bonds with the $B_a$-$B_e$ edge (2.653(3) and 2.799(3) Å). The Ag...H distances range from 2.06 to 2.34 Å, the Ag-H-B angles are 90.0–105.3°. It should be noted that the boron cage in the complex is distorted: the $B(2)$–$B(3)$ bond is elongated and equal to 1.947(10), the $B(4)$–$B(5)$ bond is 1.895(10), respectively, the Ag-H-B angles vary from 118 to 120°. The reactions of $\text{Ag}_2[B_{10}H_{10}]$ with complex $[(\text{Ph}_3\text{P})_3\text{Ag}]\text{NO}_3$ in acetonitrile results in 1–2, 1–4 positional isomers, the Ag-H-B angles are 90.0–105.3°. When the protonated form of the closo-decaborate anion was used in complexation reactions, complexes with coordination of the both metal atoms by $B_a$-$B_e$ edges near the same apical vertex were obtained. The reaction of $(\text{Ph}_3\text{P})[B_{10}H_{11}]$ with $[(\text{Ph}_3\text{P})_3\text{Ag}]\text{NO}_3$ in acetonitrile results in 1–2, 1–4 $[((\text{Ph}_3\text{P})_2\text{Ag})_2[B_{10}H_{10}]]$ [35]. The Ag...B and Ag...H distances are 2.642(3)–2.783(4) and 2.12–2.55 Å, respectively, the Ag-H-B angles vary from 118 to 120°. It should be noted that the boron cage in the complex is distorted: the B(2)–B(3) bond is elongated and equal to 1.947(10), the B(4)–B(5) bond is shortened to 1.543(9) Å (Figure 14).

![Figure 13](image1.png)

**Figure 13.** Molecular structure of 1–2, 5–8 $[((\text{Ph}_3\text{P})_2\text{Ag})_2[B_{10}H_{10}]]$ (a) and schematic representation of cocrystallized 1–2, 3–7 and 1–2, 5–8 enantiomers (b). Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.

![Figure 14](image2.png)

**Figure 14.** Molecular structures of 1–2, 1–4 $[((\text{Ph}_3\text{P})_2\text{Ag})_2[B_{10}H_{10}]]$ complex (a) and schematic representation of the 1–2, 1–4 positional isomer (b). Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.
The reactions of Ag₂[B₁₀H₁₀] with complex [(Ph₃P)₃Ag]NO₃ or free triphenylphosphine in refluxing acetonitrile unexpectedly afforded polymeric complex [(Ph₃P)₃Ag₂[B₁₀H₁₀]]. In the crystal each closo-decaborate anion is coordinated by four [(Ph₃P)Ag]⁺ fragments to form polymeric chain (Figure 15) [38].

![Figure 15. Fragment of the X-ray structure of [(Ph₃P)₃Ag₂[B₁₀H₁₀]]. Reproduced with permission from reference [38]. Copyright © (2010) Springer.](image)

The unusual 1–2, 1–3 coordination isomer [(dmapm)Cu₂[B₁₀H₁₀]] (where dmapm is 1,1-bis[di-(o-N,N-dimethylanilinyl)phosphino]methane) was prepared by the reaction of distanna-closo-decaborate (Et₄N₂)₂Sn₂[B₁₀H₁₀] with dinuclear complex [Cu₂(MeCN)₂(dmapm)][BF₄]₂ [39]. The closo-decaborate anion coordinates two copper(I) atoms via 1–2 and 1–3 apical edges forming a relative short Cu . . . Cu bond (2.6631(5) Å). The Cu(1) . . . B(2) and Cu(2) . . . B(3) distances are 2.214(3) and 2.201(3) Å, respectively, and Cu(1) . . . B(1) and Cu(2) . . . B(1) distances are 2.227(3) and 2.541(3) Å, respectively. The close positions of metal atoms (near the 1–2 and 1–3 edges) can be caused by structure of the starting copper(I) complex and general tendency of the closo-borate anion to form complexes first with less coordinated apical vertex (Figure 16).

![Figure 16. Molecular structure of [(dmapm)Cu₂[B₁₀H₁₀]] complex (a) and schematic representation of 1–2, 1–3 positional isomer (b). Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.](image)

Reaction of Cu₂[B₁₀H₁₀] with phenanthridine (5NPhen) in acetonitrile resulted in formation of a mixture of 1–2, 1–4 and 1–2, 7(8)-10 complexes [(5NPhen)₂Cu₂[B₁₀H₁₀]] [40]. In the 1–2, 1–4 isomer both copper atoms are bonded with apical edges near the same boron vertex, the Cu-B bonds with the shared apical boron atom are elongated (2.316(3) and 2.313(3) Å) as compared to the Cu-B bonds between copper atoms and equatorial boron atoms (2.215(4) and 2.245(4) Å) (Figure 17). In the 1–2,
7(8)–10 isomer the Cu-B bonds with apical boron atoms of both edges are shorter that those with equatorial edges (Cu(1) . . . B 2.219(4) and 2.244(4) Å; Cu(2) . . . B 2.253(4) and 2.280(4) Å) (Figure 18a). The similar reaction of Cu$_2$[B$_{10}$H$_{10}$] with di(pyrid-2-yl)amine (BPa) in acetonitrile at 0 °C under N$_2$ atmosphere gives the 1–2, 7(8)–10 complex {[(BPa)Cu]$_2$[B$_{10}$H$_{10}$]} with the same coordination mode of the closo-decaborate anion [41] (Figure 18b). Positional isomers found in both complexes are schematically presented in Figure 18c.

Figure 17. Structure of 1–2, 1–4 isomer of complex {((5NPhen)$_2$Cu)$_2$[B$_{10}$H$_{10}$]}. Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.

Figure 18. Structures of 1–2, 7–10 enantiomers of {((5NPhen)$_2$Cu)$_2$[B$_{10}$H$_{10}$]} (a) and {[(BPa)Cu]$_2$[B$_{10}$H$_{10}$]} (b) and schematic representation of 1–2, 7(8)–10 enantiomers (c). Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.
The listed above positional isomers belong to complexes with the closo-decaborate anion that coordinates metal atoms by two edges. However, a few examples of the facial coordination of the closo-decaborate anion were revealed. The reactions of Cu$_2$[B$_{10}$H$_{10}$] with chelate ligands 2,2'-bipyridine (2,2'-Bipy) and 1,10-phenanthroline (Phen) in acetonitrile result in formation of copper(I) complexes [((L)Cu)$_2$[B$_{10}$H$_{10}$]] with facial and mixed edge-facial coordination mode of metal atoms by the closo-decaborate anion [42]. In the complex [((2,2'-Bipy)Cu)$_2$[B$_{10}$H$_{10}$]] the boron cluster coordinates metal atoms by apical B(1)-B(2) edge and the opposite B(6)-B(7)-B(10) face (or the corresponding B(8)-B(9)-B(10) face in other enantiomer). The Cu . . . B bonds fall in the range from 2.156(7) to 2.256(7) Å (Figure 19).

![Figure 19](image_url)

**Figure 19.** Structure of [((2,2'-Bipy)Cu)$_2$[B$_{10}$H$_{10}$]] (a) and schematic representation of the cocristallized enantiomers (b). Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.

The [((Phen)Cu)$_2$[B$_{10}$H$_{10}$]] complex is centrosymmetrical with crystallographically equivalent copper atoms [43]. The closo-decaborate anion coordinates both copper atoms asymmetrically by two opposite apical faces B(1)-B(2)-B(3) and B(7)-B(8)-B(10) (or B(8)-B(9)-B(10)). The Cu . . . B bonds with each boron face are 2.269(5), 2.119(5), and 2.497(6) Å (Figure 20).
The metal coordination mode in the closo-decaborate complexes depends on steric effects of ligands and their basicity, whereas the formation of specific positional isomers is strongly governed by the reaction conditions. A more detailed discussion of this subject was presented in the recent review [14].

It should be noted that complexation reactions proceeding under redox conditions could be accompanied with substitution reactions in the closo-decaborate anion. For example, the reaction of Cu$_2$[B$_{10}$H$_{10}$] and 2,2'-Bipy in acetonitrile gives not only the above mentioned complex {((2,2'-Bipy))Cu$_2$[B$_{10}$H$_{10}$]} but also a complex with substituted closo-decaborate anion {[(MeCN)$_2$Cu[2-B$_{10}$H$_9$(N-2,2'-Bipy)]} [44]. In this complex the substituted anion [2-B$_{10}$H$_9$(N-2,2'-Bipy)]$^-$ acts as bidentate ligand. The tetrahedral environment of the Cu atom is formed by two N atoms of acetonitrile molecules (Cu-N 1.995(3) and 1.976(3) Å), the N atom of the pendant pyridine (Cu-N 2.028(2) Å), and the B(1)-H(1) apical group of the cluster anion (Cu...B 2.601(3) Å, Cu...H 1.85(3)Å, Cu-H-B angle 123(2)°) (Figure 21).
The Cu-N bond lengths fall in the range 1.997(2)–2.082(2) Å. The endocyclic N-Cu-N bond angles are approximately equal (82.06(10)–2.31(9)°), whereas the exocyclic angles around two Cu atoms vary in the ranges 105.12(10)–156.58(10)° and 107.88(9)–146.71(10)°. The complex [CuI(Phen)2]+, respectively, and [B10H10]2− anions. The coordination polyhedra of the CuII and CuI complexes consist of cationic CuI and CuII complexes [CuII(Phen)3]2+ and [CuI(Phen)2]+, respectively, and [B10H10]2− anions, and solvate MeCN molecules. Both independent Cu atoms have a severely distorted tetrahedral coordination. The Cu-N bond lengths are approximately equal (82.06(10)–2.31(9)°), whereas the exocyclic angles around two Cu atoms vary in the ranges 105.12(10)–156.58(10)° and 107.88(9)–146.71(10)°. The complex [CuI(Phen)2][B10H10]2MeCN is built of [Cu(Phen)2]+ cations, [B10H10]2− anions, and solvate MeCN molecules. Both independent Cu atoms have a severely distorted tetrahedral coordination.

On the other hand, under definite conditions the reactions can result in the complete dissociation of the starting complex and formation of complexes without Cu...HB interactions. For example, the reaction of Cu2[B10H10] and 1,10-phenanthroline in refluxing acetonitrile gives complexes [CuI(Phen)2]2[B10H10]·MeCN and [CuI(Phen)2][CuII(Phen)3][B10H10]2 [43]. The complex [CuI(Phen)2][B10H10]·MeCN is built of [Cu(Phen)2]+ cations, [B10H10]2− anions, and solvate MeCN molecules. Both independent Cu atoms have a severely distorted tetrahedral coordination.

The Cu-N bond lengths fall in the range 1.997(2)–2.082(2) Å. The endocyclic N-Cu-N bond angles are approximately equal (82.06(10)–2.31(9)°), whereas the exocyclic angles around two Cu atoms vary in the ranges 105.12(10)–156.58(10)° and 107.88(9)–146.71(10)°. The complex [CuI(Phen)2][CuII(Phen)3][B10H10]2 consist of cationic CuI and CuII complexes [CuII(Phen)3]2+ and [CuI(Phen)2]+, respectively, and [B10H10]2− anions. The coordination polyhedra of the CuII and CuI atoms are a distorted octahedron and a tetrahedron, respectively. The Cu-N bond lengths are 2.026(6)–2.330(6) Å for CuII and 2.022(7)–2.056(7) Å for CuI complex cations (Figure 22) [43].

![Figure 21. X-ray structure of {(MeCN)2Cu[2-B10H9(N-2,2'-Bipy)]}. Reproduced with permission from reference [44]. Copyright © (2011) Springer.](image)

![Figure 22. Cont.](image)
The last complex contains both Cu\textsuperscript{I} and Cu\textsuperscript{II} cationic complexes. Another example of heterovalent copper(I,II) complex is [Cu\textsuperscript{II}\textsubscript{4}(OH)\textsubscript{4}(Bipy)\textsubscript{4}][Cu\textsuperscript{I}\textsubscript{2}[B\textsubscript{10}H\textsubscript{10}\textsuperscript{3}]·4(MeCN) (See Figure 8). It should be noted that although copper(II) is easily reduced to copper(I) with decaborate anion in solutions, upon crystallization of complexes containing azaheterocyclic ligands in air a reverse process, namely the oxidation of copper(I) to copper(II), often occurs.

The reaction of CuSO\textsubscript{4}·5H\textsubscript{2}O with an excess of 1,10-phenanthroline in DMF results in qualitative precipitation of copper(II) complex [Cu(Phen)]\textsubscript{2}[B\textsubscript{10}H\textsubscript{10}]. The crystal of [Cu(Phen)]\textsubscript{2}[B\textsubscript{10}H\textsubscript{10}] is built of \[\text{Cu(Phen)}\textsubscript{2}\]\textsuperscript{2+} cations and the \[\text{B}_{10}\text{H}_{10}\]\textsuperscript{2}− anions. The copper cation is in distorted plane square coordination (Cu-N bonds are 1.9962(12) Å, N-Cu-N angles are 83.04(7), 101.29(7) and 157.73(7)°). The \[\text{Cu(Phen)}\textsubscript{2}\]\textsuperscript{2+} cations and the \[\text{B}_{10}\text{H}_{10}\]\textsuperscript{2}− anions alternate in the \textit{a} direction (Figure 23) [43].

![Figure 22](image1.png)

**Figure 22.** X-ray crystal structures of [Cu\textsuperscript{I}(Phen)\textsubscript{2}][B\textsubscript{10}H\textsubscript{10}]·MeCN (a) and [Cu\textsuperscript{I}(Phen)\textsubscript{2}][Cu\textsuperscript{II}(Phen)\textsubscript{3}][B\textsubscript{10}H\textsubscript{10}]\textsubscript{2} (b). Reproduced with permission from reference [43]. Copyright © (2015) Elsevier.

![Figure 23](image2.png)

**Figure 23.** Fragment of X-ray structure of [[Cu(Phen)\textsubscript{2}][B\textsubscript{10}H\textsubscript{10}]]. Reproduced with permission from reference [43]. Copyright © (2015) Elsevier.
Reaction of Cu$_2$[B$_{10}$H$_{10}$] with 1,10-phenanthroline in acetonitrile followed by addition of DMF or DMSO afforded anti,anti- and anti,syn-isomers of complex [Cu$_2$(Phen)$_4$(μ-CO$_3$)][B$_{10}$H$_{10}$] with bridging CO$_3$ group [45]. These complexes were structurally characterized as solvates [Cu$_2$(Phen)$_4$(μ-CO$_3$)][B$_{10}$H$_{10}$]-2.5(DMSO)-2(H$_2$O) and [Cu$_2$(Phen)$_4$(μ-CO$_3$)][B$_{10}$H$_{10}$]-4(DMF) (Figure 24). In both structures the CO$_3^{2−}$ anion acts as a bridge between two [Cu(Phen)$_2$]$^{2+}$ fragments. The basic difference between both isomers is the spatial orientation of the {Cu(Phen)$_2$}$^{2+}$ fragments: anti,syn in the DMSO solvate and anti,anti in the DMF solvate. As a result, the Cu . . . Cu distance in the dimer with the anti,syn configuration (4.4409(14) Å) is considerably shorter that in the dimers with the anti,anti configuration (5.2886(7) Å) (Figure 24) [14].

![Figure 24](image-url). Structure of the [[(Phen)$_2$Cu(μ-CO$_3$)Cu(Phen)$_2$]$^{2+}$ cations in complexes anti,syn-[Cu$_2$(Phen)$_4$(μ-CO$_3$)][B$_{10}$H$_{10}$]-2.5(DMSO)-2(H$_2$O) (a) and anti,anti-[Cu$_2$(Phen)$_4$(μ-CO$_3$)][B$_{10}$H$_{10}$]-4(DMF) (b). Reproduced with permission from reference [14]. Copyright © (2016) Elsevier.

The anti,anti-complex [Cu$_2$(Phen)$_4$(μ-CO$_3$)][B$_{10}$H$_{10}$]-3.5(DMF)-1.25(H$_2$O) was also prepared by directed reaction of (Et$_3$NH)$_2$[B$_{10}$H$_{10}$] with previously prepared copper(II) complex [Cu$_2$(Phen)$_4$(μ-CO$_3$)]Cl$_2$ with the anti,anti configuration of the CO$_3$ group in DMF [43].

The reaction of Cu$_2$[B$_{10}$H$_{10}$] with 2,2′-bipyridine in an acetonitrile/DMSO mixture produces tetranuclear hydroxo bridged copper(II) cluster [[Cu$_4$(2,2′-Bipy)$_4$(OH)$_4$][B$_{10}$H$_{10}$]$_2$-2(DMSO)] [46]. The crystals are composed of tetranuclear [Cu$_4$(2,2′-Bipy)$_4$(OH)$_4$]$^{3+}$ cations, [B$_{10}$H$_{10}$]$^{2−}$ anions and DMSO molecules. The cation has a centrosymmetric double-decker structure of the Z type (open cubane-like complex). The asymmetric unit of the complex consists of two $(2,2′$-Bipy$)_4$Cu$^{2+}$ fragments connected to each other by OH$^−$ groups. The Cu(1) and Cu(2) atoms have a flattened environment of two
nitrogen atoms and two oxygen atoms. The coordination of the Cu(1) and Cu(2) atoms is completed to 4 + 1 by the H(2B) atom of the polyhedral anion and the O(2') atom of the second part of the complex, respectively. The axial Cu(2)-O(2') bond (2.332(1) Å) is elongated as compared with the equatorial Cu-O bonds (1.922-1.946(1) Å). The Cu(1)-H(2B) bond length is 2.22(2) Å. The Cu(1) and Cu(2) atoms form long axial contacts (Cu(1) . . . N(4') 3.123 Å; Cu(2) . . . H(1B) 2.86(2) Å), each of which completes the polyhedron to an asymmetrically elongated tetragonal bipyramid. The O(2)-H(2) hydroxyl group forms a hydrogen bond with the DMSO molecule (O(2) . . . O(3) 2.676(2) Å), and the O(1)-H(1) group is involved in the formation of dihydrogen bond with the B(3)-H(3) group of the anion (H(1) . . . H(3B) 2.16(3) Å). The DMSO molecule is additionally bonded to the complex cation by the C(1)-H . . . O(3) hydrogen bond. In the complex cation, there are π-π stacking interactions between overlapping 2,2'-Bipy molecules (Figure 25). The same tetranuclear hydroxo-bridged cations were found in complex {[CuII4(OH)4(Bipy)4][CuI2[B10H10]3·4(MeCN)} (See Figure 8).

Figure 25. X-ray structure of {[Cu4(Bipy)4(OH)4][B10H10]2·2(DMSO)}. Reproduced with permission from reference [46]. Copyright © (2012) Springer.

Reaction of Cu2[B10H10] with di(pyrid-2-yl)amine in acetonitrile at ambient temperature results in hydroxo-bridged copper(II) complex {[Cu2(Bpa)2(OH)2][B10H10]} [41]. The crystals are composed of the dimeric cations [Cu2(Bpa)2(OH)2]2+ and the closo-decaborate anions. A distorted square planar environment of copper atoms is formed by two nitrogen atoms of di(pyrid-2-yl)amine and oxygen atoms of two bridging OH− groups with the Cu(1) . . . Cu(1') distance of 2.9180(14) Å. The [B10H10]2− anion is located on the two-fold rotation axis that passes through the apical vertices B(1) and B(6). The H(2) and H(4) atoms of the anion form Cu . . . HB contacts of 2.73 and 2.63 Å with copper atoms of the complex cation thus completing copper atom coordination to 4 + 2 and connecting the cations and the anions into polymeric chains (Figure 26) [41].
The similar reaction at –20 °C gave a mixture of \([[(\text{Cu}^{II}_2\text{(Bpa)}_2\text{(OH)}_2)]_2[\text{Cu}^{I}_2\text{B}_{10}\text{H}_{10}]_3\cdot n(\text{MeCN})], [\text{Cu}^{II}\text{(Bpa)}(\text{CO}_3)]_2\cdot \text{H}_2\text{O}]\) and \([\text{Cu}^{II}_2(\text{Bpa})_2(\text{MeCN})_2][2-\text{B}_{10}\text{H}_9(\text{N-Bpa})]_2\cdot 2(\text{H}_2\text{O})\) complexes. The crystal of \([\text{Cu}^{II}_2(\text{Bpa})_2(\text{MeCN})_2][2-\text{B}_{10}\text{H}_9(\text{N-Bpa})]_2\cdot 2(\text{H}_2\text{O})\) consists of the \([\text{Cu}^{II}_2(\text{Bpa})_2(\text{MeCN})_2]^{2+}\) cations and the \([2-\text{B}_{10}\text{H}_9(\text{N-Bpa})]^{-}\) anions, where molecule of di(pyrid-2-yl)amine is attached through nitrogen atom of the pyridine ring to the B(2) position of the boron cluster anion (Figure 27) [41].
The complexation between the closo-decaborate anion and cyclic copper(I) and silver(I) 3,5-bis(trifluoromethyl)pyrazolates \([3,5-(\text{CF}_3)_2\text{Pz}]\text{M}\)\(_3\) (M = Cu, Ag) (Scheme 2) in CH\(_2\text{Cl}_2\) was studied by IR spectroscopy. Formation of two types of complexes, 1:1 and 1:2, was revealed and their stability constants were determined [47].

![Scheme 2. Copper(I) and silver(I) 3,5-bis(trifluoromethyl)pyrazolates \([3,5-(\text{CF}_3)_2\text{Pz}]\text{M}\)\(_3\) (M = Cu, Ag).](image)

Silver ammonia complex with decachloro-closo-decaborate anion \([\text{Ag(NH}_3)_2]_2[\text{B}\text{10Cl}_{10}]\) was prepared by reaction of \((\text{Et}_3\text{NH})_2[\text{B}\text{10Cl}_{10}]\) with AgNO\(_3\) in aqueous ammonia. According to X-ray diffraction, the crystal is built of the \([\text{Ag(NH}_3)_2]^+\) cations and \([\text{B}\text{10Cl}_{10}]^{2-}\) anions. The Ag-N distances are equal to 2.128(3) and 2.140(3) Å, the NAgN angle is almost linear (178.9°). There are rather short N–H...Cl (<2.95 Å) and Ag...Cl (<3.5 Å) contacts between the cations and anions (Figure 28) [48].

![Figure 28. The crystal environment of the \([\text{B}\text{10Cl}_{10}]^{2-}\) anions in structure of \([\text{Ag(NH}_3)_2]_2[\text{B}\text{10Cl}_{10}]\). Reproduced with permission from reference [48]. Copyright © (2016) Elsevier.](image)

An interesting case presents a silver complex with the dimeric icosaborate anion \(\text{trans-}[\text{B}_{20}\text{H}_{18}]^{2-}\) which can be considered as product of mild oxidation of the closo-decaborate anion \([\text{B}10\text{H}_{10}]^{2-}\) [49]. Complex \(((\text{Ph}_3\text{P})_3\text{Ag})_2[\text{trans-}\text{B}_{20}\text{H}_{18}]\) was prepared by the reaction of \((\text{Et}_3\text{NH})_2[\text{trans-}\text{B}_{20}\text{H}_{18}]\) with \(((\text{Ph}_3\text{P})_3\text{Ag})\text{NO}_3\) in N,N-dimethylformamide. In the obtained complex each silver atom is coordinated by the apical BH group of the dimeric boron anion and three molecules of triphenylphosphine. The Ag...B distance is 3.382(3) Å and the AgHB angle is 161.8° that reflects much weaker cation-anion bonding in comparison with the parent closo-decaborate anion (Figure 29) [50].
The Ag...B distances increase to 3.517(5) and 3.394(8) Å, respectively (Figure 29). The isomerization prepared [20,52,53]. Their X-ray structures were not determined, however the existence of M-H-B interactions in the solid state was supported by the IR spectroscopy data.

3. Copper and Silver Complexes with the closo-dodecaborate anion \([\text{B}_{12}\text{H}_{12}]^{2-}\)

The closo-dodecaborate anion \([\text{B}_{12}\text{H}_{12}]^{2-}\) is the most studied representative of the closo-borane family [51]. The \([\text{B}_{12}\text{H}_{12}]^{2-}\) anion has close to sphere icosahedral structure that causes decrease of its coordination ability. Nevertheless, a large number of its copper(I) and silver(I) complexes has been described.

The binary copper(I) and silver(I) salts of \([\text{B}_{12}\text{H}_{12}]^{2-}\) were prepared [50]. Complex \([[\text{(MeCN)}_2\text{Cu}]_2[\text{B}_{12}\text{H}_{12}]\]] was prepared by the reaction of the acid \((\text{H}_3\text{O})_2[\text{B}_{12}\text{H}_{12}]\) with \(\text{Cu}_2\text{O}\) followed by recrystallization from acetonitrile. In the structure of \([[\text{(MeCN)}_2\text{Cu}]_2[\text{B}_{12}\text{H}_{12}]\]] two

Figure 29. X-ray structures of \([[\text{(Ph}_3\text{P)}_2\text{Ag}]_2[\text{trans-B}_{20}\text{H}_{18}]\]] (a) and two cocry stallized isomers of \([[\text{(Ph}_3\text{P)}_2\text{Ag}]_2[\text{iso-B}_{20}\text{H}_{18}]\]] (b and c) (phenyl cycles are omitted for clarity). Reproduced from reference [50].

The UV radiation of a crystal of \([[\text{(Ph}_3\text{P)}_2\text{Ag}]_2[\text{trans-B}_{20}\text{H}_{18}]\]] results in its single-crystal-to-single-crystal isomerization to \([[\text{(Ph}_3\text{P)}_2\text{Ag}]_2[\text{iso-B}_{20}\text{H}_{18}]\]] without the crystal degradation with only minor (0.75%) contraction of the unit cell. There are two linkage isomers co-crystallize in the final crystal of \([[\text{(Ph}_3\text{P)}_2\text{Ag}]_2[\text{iso-B}_{20}\text{H}_{18}]\]]: in one of them, the BHAg angle is practically linear (172°), while in other isomer with facial coordination its value (116°) is typical for polydentate coordination of silver atom. The Ag...B distances increase to 3.517(5) and 3.394(8) Å, respectively (Figure 29). The isomerization is reversible, and heating \([[\text{(Ph}_3\text{P)}_2\text{Ag}]_2[\text{iso-B}_{20}\text{H}_{18}]\]] at 150°C produces \([[\text{(Ph}_3\text{P)}_2\text{Ag}]_2[\text{trans-B}_{20}\text{H}_{18}]\]] the iso-isomer was converted completely to the described trans-isomer [50].
{(MeCN)\textsubscript{2}Cu}\textsuperscript{+} fragments are coordinated to opposite edges of the closo-dodecaborate anion. The Cu…B bonds are 2.288(3) and 2.351(4) Å, Cu…H(B) bonds are 1.86 and 1.95 Å (Figure 30) [54].

![Figure 30. X-ray structure of \{((MeCN)\textsubscript{2}Cu)\textsubscript{2}[B\textsubscript{12}H\textsubscript{12}]\}. Reproduced with permission from reference [54]. Copyright © (2008) Springer.](image)

Complex \{((MeCN)Ag)\textsubscript{2}[B\textsubscript{12}H\textsubscript{12}]\}_n which was prepared by crystallization of Ag\textsubscript{2}[B\textsubscript{12}H\textsubscript{12}] from acetonitrile has a 3D-polymeric framework structure with two crystallographically different Ag cations. The Ag(1) cation is coordinated via Ag…(BH)\textsubscript{2} bridges by two closo-dodecaborate anions and one acetonitrile molecule forming slightly distorted square pyramid environments. The Ag(2) cation is coordinated via single Ag…BH bridges by three acetonitrile molecule forming tetrahedral arrangement. The Ag…B and Ag…H bonds are 2.582(3)–2.873(3) and 2.02(3)–226(4) Å, respectively (Figure 31) [55].

![Figure 31. Coordination environment of Ag(1) atoms (a), Ag(2) atoms (b) and the closo-dodecaborate anion (c) in the structure of \{(MeCN)Ag\textsubscript{2}[B\textsubscript{12}H\textsubscript{12}]\}_n. Reproduced with permission from reference 55. Copyright © (2005) John Wiley & Sons.](image)

Similar to complexes of the closo-decaborate anion, a series of complexes \{Cat(Cu[B\textsubscript{12}H\textsubscript{12}])\} (Cat\textsuperscript{+} = Cs, R\textsubscript{3}NH\textsuperscript{+} and R\textsubscript{4}N\textsuperscript{+} (R = Me, Et, Pr, Bu), Ph\textsubscript{3}P\textsuperscript{+}, Ph\textsubscript{4}As\textsuperscript{+}) was prepared by reaction of
Cat$_2$[B$_{12}$H$_{12}$] with CuSO$_4$ or CuCl$_2$ in aqueous solution in the presence of reducing agent (SO$_2$ or Na$_2$SO$_3$) [23,54]. In the IR spectra of the Cat[Ag[B$_{12}$H$_{12}$]] complexes along with the B-H stretching band at 2450–2540 cm$^{-1}$ a new band appears at 2220–2165 cm$^{-1}$, that can be assigned to Cu-H-B stretching by analogy with similar complexes of the closo-decaborate anion.

The similar silver(I) complexes [Cat(Ag[B$_{12}$H$_{12}$])] (Cat$^+$ = Cs, R$_3$NH$^+$ and R$_4$N$^+$ (R = Me, Et, Pr, Bu), Et$_3$NBz$^+$, Ph$_4$P$^+$, Ph$_4$As$^+$, Ph$_3$PCH$_2$Naph$^+$) were prepared by the reaction of Cat$_2$[B$_{12}$H$_{12}$] with AgNO$_3$, in aqueous solution [53,55]. Structures of complexes [Cat(Ag[B$_{12}$H$_{12}$])]$_n$ (Cat$^+$ = Et$_3$NBz$^+$ and Ph$_3$PCH$_2$Naph$^+$) were determined by single crystal X-ray diffraction.

The structure of complex (Ph$_3$PCH$_2$Naph)(Ag[B$_{12}$H$_{12}$])$_n$ is built of the {Ag[B$_{12}$H$_{12}$]}$_n^{-}$ chains and the [Ph$_3$PCH$_2$Naph]$^+$ cations. Each silver cation is coordinated by two closo-decaborate anions, to which silver is bound by Ag . . . (BH)$_2$ bonds (Ag . . . HB distances are 2.474(6)–2.580(6) Å). There are two types of closo-decaborate anions: the first one (A1) coordinate silver via B(1)–B(2) and B(9)-B(12) edges, whereas the second one (A2) coordinates silver via B(1)-B(2) and B(7)-B(8) edges. The centrosymmetric A1 anion provides linearity of the chain fragments of the chain, whereas the A2 anion provides a bend of the silver anionic chain (Figure 32) [53].

![Figure 32. Fragment of polymeric anionic chain in structure of (Ph$_3$PCH$_2$Naph)(Ag[B$_{12}$H$_{12}$])$_n$. Reproduced with permission from reference [53]. Copyright © (2008) Springer.](image1)

The structure of complex ([Et$_3$NBz](Ag[B$_{12}$H$_{12}$]))$_n$ is characterized by double anionic chain structure of [Ag[B$_{12}$H$_{12}$]]$^-$. Each silver cation is coordinated by three icosahedral cages, to which silver is bound by Ag . . . (BH)$_2$ bonds. Each closo-decaborate anion cage is coordinated by three silver cations. The Ag . . . B and Ag-H bonds fall in the ranges 263.8(2)–292.0(2) and 2.18(2)–2.40(2) Å, respectively (Figure 33) [55].

![Figure 33. Fragment of structure of double polymeric anionic chain in ([Et$_3$NBz](Ag[B$_{12}$H$_{12}$]))$_n$. Reproduced with permission from reference [55]. Copyright © (2005) John Wiley & Sons.](image2)
Complex \((\text{Ph}_3\text{PCH}_2\text{Naph})\)\(((\text{MeCN})\text{Ag})\text{[B}_{12}\text{H}_{12}]\)\)\()_n\) was obtained by crystallization of \(((\text{Et}_3\text{NBz})\text{[B}_{12}\text{H}_{12}]\)\)\()_n\) from acetonitrile. The crystals are built of the silver anionic chains and the \([\text{Ph}_3\text{PCH}_2\text{Naph}]^+\) cations. The silver chain contains one centrosymmetric \(\text{Ag} \cdots \text{B} \cdots \text{Ag}\) fragment similar to that found in the acetonitrile-free complex (see Figure 30, the A1 anion) and one additional fragment, in which silver atom is coordinated by two closo-dodecaborate anions and an acetonitrile molecule. Thus, the environment of silver \(\text{Ag}(1)\) and \(\text{Ag}(2)\) is formed by two edges of two symmetrically unequivalent icosahedra (A3 and A4) (Figure 34) [53].

**Figure 34.** Fragment of polymeric anionic chain in structure of \(((\text{Ph}_3\text{PCH}_2\text{Naph})\)\(((\text{MeCN})\text{Ag})\text{[B}_{12}\text{H}_{12}]\)\)\()_n\). Reproduced with permission from reference [53]. Copyright © (2008) Springer.

Complex \(((\text{Et}_3\text{NBz})\text{[B}_{12}\text{H}_{12}]\)\)\()_n\) was prepared by addition of aqueous \(\text{AgNO}_3\) to a solution of \(((\text{Et}_3\text{NBz})\text{[B}_{12}\text{H}_{12}]\)\)\()_n\) in acetonitrile. The asymmetric unit contains three silver atoms, two closo-dodecaborate anions, two acetonitrile molecules and the organic cation. Two silver atoms \(\text{Ag}(1)\) and \(\text{Ag}(2)\) are coordinated by two icosahedral cages via \(\text{Ag} \cdots \text{B} \cdots \text{Ag}\) bridges and acetonitrile molecule, giving a five-fold arrangement. The \(\text{Ag}(3)\) atom is coordinated by two closo-dodecaborate anions giving a distorted tetrahedron. The \(\text{Ag} \cdots \text{B}\) bonds fall in the range of 2.426(3)–2.987(3) Å (Figure 35) [55].

**Figure 35.** Fragment of X-ray structure of \(((\text{Et}_3\text{NBz})\text{[B}_{12}\text{H}_{12}]\)\)\()_n\). Reproduced with permission from reference [55]. Copyright © (2005) John Wiley & Sons.

Complex with bis(triphenylphosphine)gold(I) cation \(((\text{Ph}_3\text{P})\text{[B}_{12}\text{H}_{12}]\)\)\()_n\) was prepared by reaction of \(\text{Ag}_2\text{[B}_{12}\text{H}_{12}]\) with \(\text{[Ph}_3\text{P})\text{AuCl}\) in acetonitrile/benzene. The structure of the complex 

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consists of polymeric anionic chains \(\text{Ag}[\text{B}_{12}\text{H}_{12}]\)^+ which are extended along axis c and \([\text{Ph}_3\text{P}_2\text{Au}]^+\) complex cations. The silver atom is coordinated by the B(1)-B(2) edge of one anion and the B(9)-B(10)-B(12) face of the other anion. The Ag…B bond lengths vary from 2.498(11) to 2.738(11) Å. In the \([\text{Ph}_3\text{P}_2\text{Au}]^+\) cation the gold atom has a nearly linear coordination (Au-P bond lengths are 2.338(2) and 2.341(2) Å, P-Au-P angle is 170.91(7)°) (Figure 36) [56].

Silver dodecachloro-closo-dodecaborate \(\text{Ag}_2[\text{B}_{12}\text{Cl}_{12}]_n\) was obtained by the metathesis reaction of \(\text{Cs}_2[\text{B}_{12}\text{Cl}_{12}]\) with an aqueous solution of AgNO\(_3\) [57] or by reaction of an aqueous solution of the free acid \((\text{H}_3\text{O})_2[\text{B}_{12}\text{Cl}_{12}]\) with Ag\(_2\)O [57,58]. In the crystal each Ag\(^+\) cation is coordinated by six chlorine atoms from the edges of three \([\text{B}_{12}\text{Cl}_{12}]^\cdot\) anions providing a distorted octahedral environment (Ag…Cl distances 2.83–2.85 Å) (Figure 37) [57].

![Figure 36. Fragment of anionic polymeric chain (a) and crystal packing of \([\text{Ph}_3\text{P}_2\text{Au}][\text{Ag}[\text{B}_{12}\text{H}_{12}]])_n\) (b). Reproduced with permission from reference [56]. Copyright © (2011) Springer.](image-url)
Complex \(((\text{Me}_2\text{CO})\text{Ag})_2[B_{12}\text{Br}_{12}]\)_n was prepared by the metathesis reaction of \((\text{Et}_3\text{NBz})_2[B_{12}\text{Br}_{12}]\) with AgOTf in acetone [55]. In the structure, the Ag\(^+\) cations are coordinated in a square-pyramidal way by four bromine atoms from two \([B_{12}\text{Br}_{12}]^{2-}\) anions at the basis and by acetone molecule at the apex. Each dodecabromo-closo-dodecaborate anion is coordinated by four \(((\text{Me}_2\text{CO})\text{Ag})^+\) cations. The Ag…Br bonds fall in the range of 2.731–2.906 Å, the Ag-O bond is 2.344 Å (Figure 38).

Lowering the anion charge from 2- to 1- by an introduction of a charge-compensating group decreases its coordination ability. Complex \([\text{Ag(MeCN)}_4][B_{12}\text{Cl}_{11}\text{NH}_3] \cdot 3(\text{MeCN})\) prepared by crystallization from acetonitrile consists of the discrete \([\text{Ag(MeCN)}_4]^+\) cations, the \([B_{12}\text{Cl}_{11}\text{NH}_3]^-\) anions and acetonitrile solvate molecules (Figure 39) [59].
Several complexes of closo-dodecaborate anion with triphenylphosphine and di(2-pyridyl)amine ligands were prepared. In the structure of \{(Ph\textsubscript{3}P)\textsubscript{2}Cu\}[B\textsubscript{12}H\textsubscript{12}]\) two \{(Ph\textsubscript{3}P)\textsubscript{2}Cu\}\textsuperscript{+} fragments coordinated to opposite edges of the closo-dodecaborate anion. The Cu…B bonds are 2.392(18) and 2.387(17) Å, Cu…H(B) bonds are 1.82(13) and 1.88(15) Å (Figure 40) [60].

The polymeric complex \{((BPa)Ag)\textsubscript{2}[B\textsubscript{12}H\textsubscript{12}]\}_n was prepared by the reaction of Cs(Ag[B\textsubscript{12}H\textsubscript{12}]) or Ag\textsubscript{2}[B\textsubscript{12}H\textsubscript{12}] with di(2-pyridyl)amine in acetonitrile. The complex is built of centrosymmetric binuclear \{(BPa)Ag[B\textsubscript{12}H\textsubscript{12}]Ag(BPa)\} units connected into chains running along the \textit{a} axis. The silver cation is coordinated by the nitrogen atoms of the chelating di(2-pyridyl)amine ligand (Ag-N bonds are 2.288(3) and 2.320(3) Å), the B(1)H group of the closo-dodecaborate anion (Ag…B and Ag…H bonds are 2.544(4) and 1.87 Å, respectively, Ag-H-B angle is 114°), and the H(4) atom of the neighboring anion. The interaction between the neighboring complexes is apparently weak: Ag…H distance is 2.14 Å, the Ag-H-B angle is open to 143°, and Ag…B distance of 3.108(4) Å is too long to be considered a bond. In the crystal di(2-pyridyl)amine molecules pack into stacks with the interplanar spacing of ~3.50 Å (Figure 41) [61].

Figure 39. X-ray structure of [Ag(MeCN)\textsubscript{4}][B\textsubscript{12}Cl\textsubscript{11}NH\textsubscript{3}·3(MeCN). Reproduced with permission from reference [59]. Copyright © (2015) John Wiley & Sons.

Figure 40. X-ray structure of complex \{(Ph\textsubscript{3}P)\textsubscript{2}Cu\}[B\textsubscript{12}H\textsubscript{12}]\}.
The reaction of Cs(Ag[B_{12}H_{12}]) or Ag_{2}[B_{12}H_{12}] with di(2-pyridyl)amine in DMF produces discrete complex \(((\text{Bpa})\text{Ag})_2[B_{12}H_{12}]: 2(\text{DMF})\). The \(((\text{Bpa})\text{Ag})_2[B_{12}H_{12}]\) fragment is centrosymmetric. The silver atoms has a trigonal coordination formed by two nitrogen atoms of the chelating di(2-pyridyl)amine ligand (Ag-N, 2.2745(15) Å) and the BH group of the closo-dodecaborate anion (Ag-B, 2.60(2) Å). In the crystal two DMF molecules are attached to amino groups of di(2-pyridyl)amine ligands by N-H….O hydrogen bonds (Figure 42) [61].

The mixed-ligand complex \(((\text{BPa})(\text{Ph}_3\text{P})\text{Ag})_2[B_{12}H_{12}]\) was prepared by reaction of Cs(Ag[B_{12}H_{12}]) with triphenylphosphine and di(2-pyridyl)amine in a DMF/acetonitrile/benzene mixture. Crystals of this compound are built of centrosymmetric binuclear complexes \(((\text{BPa})(\text{Ph}_3\text{P})\text{Ag}(B_{12}H_{12}))\text{Ag}(\text{Bpa})(\text{PPh}_3))\). The pyridine nitrogen atoms of di(2-pyridyl)amine and the phosphorus atom of triphenylphosphine form strong bonds with the silver cation (Ag-N, 2.356(2) and 2.369(3) Å; Ag-P, 2.3963(7) Å). The trigonal coordination of silver is complemented by two weak bonds with the BH groups of the closo-dodecaborate anion (Ag(1)….B(1), 2.831(3); Ag(1)….H(1), 2.32(3) Å; Ag(1)….B(2), 2.886(3);...
Ag(1) ... H(2), 2.43(3) Å (Figure 43). Although the Ag ... B distances in this complex are much longer than in [(BPa)Ag][B₁₂H₁₂] \_2(DMF) and [(BPa)Ag][B₁₂H₁₂]n (2.413-2.544 Å), the Ag-H-B stretching band at 2389 cm\(^{-1}\) in the IR spectrum unambiguously indicates the existence of Ag-H-B interactions between the silver atom and the closo-dodecaborate anion [61].

![X-ray structure of complex \{((BPa)(Ph₃P)Ag)₂[B₁₂H₁₂]\}. Reproduced with permission from reference [61]. Copyright © (2016) Elsevier.](image)

Complex \((\text{HAgu})_2[\text{Cu(Agu)}_2][\text{B}_{12}\text{H}_{12}]\) was prepared by reaction of \(\text{CuSO}_4\) with \((\text{HAgu})_2[\text{B}_{12}\text{H}_{12}]\) in aqueous solution. The structure is built of \([\text{Cu(Agu)}_2]^{2+}\) and \([\text{(HAgu)}_2]^{2+}\) cations and the \([\text{B}_{12}\text{H}_{12}]^{2-}\) anions. The square planar environment of copper atom is formed by two flat aminoguanidine molecules, which close five-membered chelate cycles (Cu-N, 1.9702(13) and 2.0261(10) Å; N-Cu-N angle, 81.53(4)°). In addition, copper atom forms two weak bonds with hydrogen atoms of two closo-dodecaborate anions (Cu ... H, 2.805(14) Å; B-H-Cu angle, 132.5(10)°). In the cells where copper positions are vacant, pairs of HAgu\(^+\) cations are connected by the intermolecular N(1)-H(13) ... N(4) hydrogen bonds into centrosymmetric dimers. The N(4)H atom is also involved in intermolecular dihydrogen bond with the B(3)H atom (Figure 44) [62].

The complexation between the closo-dodecaborate anion and cyclic copper(1) and silver(1) 3,5-bis(trifluoromethyl)pyrazolates \([3,5-(\text{CF}_3)_2\text{Pz}]\text{M}\) \(\text{(M} = \text{Cu, Ag)}\) in CH₂Cl₂ was studied by IR spectroscopy. Formation of two types of complexes, 1:1 and 1:2, was revealed and their stability constants were determined. It was demonstrated that closo-dodecaborate anion forms less stable complexes than the closo-decaborate anion [47].

![Representation of complexes](image)

(a)

Figure 44. Cont.
4. Silver Complexes with the 1-carba-closo-decaborate anion \([1-\text{CB}_9\text{H}_{10}]^-\)

Synthesis of the 1-carba-closo-decaborate anion \([1-\text{CB}_9\text{H}_{10}]^-\) anion was reported for the first time almost fifty years ago in the late 1960s [63,64], its coordination chemistry has been studied much less than chemistry of other weakly coordinated anions due to the absence of practical methods of its synthesis until 2000-s [65]. The situation was changed when convenient approach to synthesis of the parent anion and its C-aryl derivatives via reaction of decaborane with aldehydes in alkali media was elaborated [66–72].

The silver salt of the parent 1-carba-closo-decaborate anion, Ag\([1-\text{CB}_9\text{H}_{10}]\), can be prepared by metathesis of the cesium salt with \(\text{AgNO}_3\) in aqueous solution followed by crystallization from toluene [73]. In the crystal the coordination geometry of silver approximates linear two-coordination except that the \([1-\text{CB}_9\text{H}_{10}]^-\) anions provide \(\eta^2\)-like coordination via the antipodal to carbon B(6)-B(10) and B(8)-B(10) edges to form infinite polymeric chain. The silver atom is closer to B(6) (2.496(3) Å) than to B(10) (2.578(2) Å); however, this probably does not indicate site nucleophilicity on the decaborane cage, but rather reflects the fact that the B(10)-H bond is shared by two silver atoms (Figure 45) [73].

Figure 44. X-ray structure of \([\text{HAgu}]_2[\text{Cu(Agu)}_2][\text{B}_{12}\text{H}_{12}]_2\]: fragment of polymeric chain (a) and crystal packing (projection along the x axis) (b). Reproduced with permission from reference [62].

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The silver salt of 6,7,8,9,10-pentabromo-1-carba-closo-decaborate anion, Ag\([1-\text{CB}_9\text{H}_5-6,7,8,9,10-\text{Br}_5]\), was prepared by the metathesis of Cs\([1-\text{CB}_9\text{H}_5-6,7,8,9,10-\text{Br}_5]\) with \(\text{AgNO}_3\) from aqueous solution [73].

Figure 45. Fragment of X-ray structure of \([\text{Ag}[1-\text{CB}_9\text{H}_{10}]]_n\) (only hydrogen atoms bonded to silver are shown). Reproduced with permission from reference [73]. Copyright © (1994) American Chemical Society.
The toluene solvate \([\text{Ag}(\eta^2\text{-toluene})[1\text{-CB}_9\text{H}_5\text{-6,7,8,9,10-Br}_5]]_n\) was obtained by crystallization from toluene/\(n\)-hexane. The coordination sphere of silver is formed by \(\eta^2\)-toluene, one bidentate carborane anion, and one monodentate carborane anion. The anions act as bridging ligands to a zigzag linear chain of silver ions. The \(\eta^2\) coordination of toluene has Ag-C distances of about 2.62 Å. The coordination of the anion is notable on two counts. Firstly, the bridging disposition is quite unsymmetrical, making it “belong” more to one silver atom than another. This is true of both denticity and bond length. The bidentate binding is characterized by Ag-Br bond lengths of \(\sim 2.78\) Å, whereas the monodentate interaction is weaker at \(2.842(6)\) Å. Secondly, in the bidentate interaction, the Ag-Br(10) and Ag-Br(6) bond lengths are practically indistinguishable \(2.783(6)\) and \(2.775(6)\) Å, respectively (Figure 46) [73].

![Figure 46. Fragment of X-ray structure of \([\text{Ag}(\eta^2\text{-toluene})[1\text{-CB}_9\text{H}_5\text{-6,7,8,9,10-Br}_5]]_n\). Reprinted with permission from reference [73]. Copyright © (1994) American Chemical Society.](image)

The silver salts of C-substituted perchloro and perbromo derivatives of 1-carba-closo-decaborate anion, \(\text{Ag}[1\text{-Bn-1-CB}_3\text{Cl}_6]\) and \(\text{Ag}[1\text{-H}_2\text{N-1-CB}_3\text{Br}_9]\), were prepared by treatment of the corresponding sodium salts with \(\text{AgNO}_3\) in aqueous solution [74]. The silver salt of the perbromo derivative of 1-carba-closo-decaborate anion, \(\text{Ag}[1\text{-HCB}_9\text{Br}_9]\), was prepared by the metathesis of \(\text{Cs}[1\text{-HCB}_9\text{Br}_9]\) with \(\text{AgNO}_3\) from aqueous solution and crystallized from benzene/THF solution to give \([\text{Ag}(\eta^2\text{-benzene})[1\text{-HCB}_9\text{Br}_9]]_n\) [74]. The \(\text{[1-HCB}_9\text{Br}_9]^-\) anions act as bridging ligands in a one-dimensional coordination polymer chain, an increasingly familiar motif in silver carborane structures. The silver atom is in a trigonal arrangement of one \(\eta^2\)-benzene and two monodentate bridging carborane anions. It is noteworthy that the \(\text{[1-HCB}_9\text{Br}_9]^-\) anion coordinates to the silver atom through the Br(6) atom rather than Br(10) antipodal to the carbon atom. The average Ag-Br distance of \(2.778\) Å is close to those found in \([\text{Ag}(\eta^2\text{-toluene})[1\text{-CB}_9\text{H}_5\text{-6,7,8,9,10-Br}_5]]_n\). The asymmetric \(\eta^2\) benzene coordination has Ag-C distances of \(2.629(8)\) and \(2.649(7)\) Å (Figure 47) [74].

![Figure 47. Fragment of X-ray structure of \([\text{Ag}(\eta^2\text{-benzene})[1\text{-HCB}_9\text{Br}_9]]_n\). Reprinted with permission from reference [74]. Copyright © (2000) American Chemical Society.](image)
At the same time crystals of $[(\eta^2\text{-benzene})(\eta^1\text{-benzene})\text{Ag}_3[1\text{-HCB}_9\text{Br}_9][\text{NO}_3](\text{THF})]_n$ were obtained. The X-ray analysis reveals that this complex is a one-dimensional coordination polymer with both carborane and nitrate anions acting as bridging ligands. Each Ag atom in asymmetric unit has a unique coordination environment although they are all five-coordinated. Ag(1) is in a propeller-like arrangement having one $\eta^2$-benzene and two bidentate bridging $[1\text{-HCB}_9\text{Br}_9]^-$ anions. Ag(2) is bound to one bidentate $[1\text{-HCB}_9\text{Br}_9]^-$ anion, one bidentate NO$_3^-$ anion, and the oxygen atom of the coordinated THF molecule in a distorted square-pyramidal geometry. Ag(3) is in a highly distorted square-pyramidal arrangement having one $\eta^1$-benzene, one bidentate $[1\text{-HCB}_9\text{Br}_9]^-$ anion, and one bidentate NO$_3^-$ anion. It is noteworthy that a bromine atom from the upper tetragonal belt can also coordinate to an Ag$^+$ ion (Figure 48) [74].

**Figure 48.** Fragment of X-ray structure of $[(\eta^2\text{-benzene})(\eta^1\text{-benzene})\text{Ag}_3[1\text{-HCB}_9\text{Br}_9]^2[\text{NO}_3](\text{THF})]_n$. Reprinted with permission from reference [74]. Copyright © (2000) American Chemical Society.

Synthesis of silver salts of some other derivatives of the 1-carba-closo-decaborate anion (Ag[1-Ph-1-CB$_9$H$_9$]) [75], Ag[1-PhCH$_2$-1-CB$_9$Cl$_9$] [74], Ag[1-NH$_2$-1-CB$_9$Br$_9$] [74], Ag[1-Ph-1-CB$_9$H$_5$-6-I] [76], Ag[1-Ph-1-CB$_9$H$_4$-6,7,8,9,10-I$_5$] [75]) was described as well.

While the relatively labile chloride ligand in complexes MCl(CO)(PPh$_3$)$_2$ (M = Ir, Rh) can be readily abstracted with silver perchlorate or triflate, the reactions with Ag[1-CB$_9$H$_5$-6,7,8,9,10-Br$_5$] result in M→Ag donor-acceptor metal-metal bonded adducts (PPh$_3$)$_2$(CO)CIM·Ag[1-CB$_9$H$_5$-6,7,8,9,10-Br$_5$] that were isolated in the solid state [73]. The iridium complex was characterized by single crystal X-ray diffraction to adopt the expected square pyramid silver-iridium Lewis acid-base structure. The $[1\text{-CB}_9\text{H}_5^-6,7,8,9,10\text{-Br}_5]$ anion is coordinated to silver atom through three bromine atoms realizing the $\eta^3$-coordination mode. The Ag-Br distances range from 2.721(8) to 2.973(6) Å (Figure 49) [73].

**Figure 49.** X-ray structure of $[[\text{PPh}_3)_2$(CO)ClIr]Ag[1-CB$_9$H$_5$-6,7,8,9,10-Br$_5$]]. Reprinted with permission from reference [73]. Copyright © (1994) American Chemical Society.
Complex \([\text{[Ag(4,4'-Bipy)][1-Ph-1-CB_9H_5]·MeCN}_n]\) was prepared by reaction of Ag[1-Ph-1-CB_9H_5] with 4,4'-bipyridine in acetonitrile [75]. The Ag(I) has two coordinate bonds to two 4,4'-Bipy ligands in a near-linear fashion, with an angle N(1)-Ag-(1)-N(2) of 177.42(15)° and with Ag-N distances of Ag(1)-N(1) 2.132(4) and Ag(1)-N(2) 2.144(5) Å. Each 4,4'-bipyridine ligand bridges two silver atoms to give an infinite linear coordination chain. The chains group together in pairs, with the Ag(I) center of one chain positioned directly above the center of the C(4)-C(4') bond of the 4,4'-bipyridine of the second chain at a closest Ag … C distance of 3.585 Å. There are face-to-face π-stacking interactions between the arene rings of each chain at a centroid separation of 3.793 Å. The carborane anions do not form any close interactions with other molecular components. Both [1-Ph-1-CB_9H_5]^- anion and MeCN solvent molecules are arranged in a fashion that creates unidirectional channels of rhomboid cross section that run along the y-direction and contain the [Ag(4,4'-Bipy)]^+ coordination chains (Figure 50).

**Figure 50.** The crystal packing in X-ray structure of \([\text{[Ag(4,4'-Bipy)][1-Ph-1-CB_9H_5]·MeCN}_n]\).

Similar complex with iodinated carborane anion \([\text{Ag(4,4'-Bipy)[1-Ph-1-CB_9H_4-6,7,8,9,10-I_5]·MeCN}_n}\) was prepared by the reaction of Ag[1-Ph-1-CB_9H_4-6,7,8,9,10-I_5] with 4,4'-bipyridine in acetonitrile [75]. In contrast to the previous complex, the silver atom here has a distorted tetrahedral geometry with two coordinate bonds to two 4,4'-Bipy ligands with an angle N(1)-Ag(1)-N(2) of 127.42(11)° and two symmetrically equivalent Ag-N bonds of 2.271(4) Å. The coordination sphere is completed by two iodine atoms of the chelating \([\text{1-Ph-1-CB_9H_4-6,7,8,9,10-I_5}]^-\) anion, which coordinates to the metal center at Ag-I distances of Ag(1)-I(1) 2.985(5) and Ag(1)-I(2) 2.8642(10)Å. The [Ag(4,4'-Bipy)]^+ coordination chain is puckered to create a zigzag chain. At each silver center along the chain the coordinated [1-Ph-1-CB_9H_4-6,7,8,9,10-I_5]^- anions point in alternating directions (Figure 51) [75].

Complex \([\text{Ag(bppz)[1-Ph-1-CB_9H_5]·0.5MeCN}_n]\) was prepared by reaction of Ag[1-Ph-1-CB_9H_5] with 2,3-bis(2-pyridyl)pyrazine in acetonitrile [75]. The silver atom has a distorted six-coordinate geometry, arising from two bidentate chelating interactions with two symmetry-related bppz ligands with Ag-N distances in the range of 2.325(2) and 2.593(2) Å and from two Ag … H-B interactions at distances of 2.291 Å (Ag(1) … H(103)) and 2.471 Å (Ag(1) … H(101)), both involving the same [1-Ph-1-CB_9H_5]^- anion. The 2,3-bis(2-pyridyl)pyrazine ligand acts as a bis-chelating bridging ligand between the Ag(I) centers and thereby propagates a coordination chain that runs parallel with the crystallographic c-axis (Figure 52a) [75]. The coordinated anions are all on one side of the chain but show two alternating orientations along the chain. The chains stack with an identical orientation in the x-direction, such that the phenyl groups of the anions of one chain slot into the grooves created by the pyridine rings of the bppz ligands of an adjacent chain. The orientation of the chains alternates along the y-direction (Figure 52b) [75].
**Figure 51.** Fragment of X-ray structure of \([\text{Ag}(4,4'-\text{Bipy})[1-\text{Ph}-1-\text{CB}_{9}\text{H}_{4}-6,7,8,9,10-\text{I}_{5}]]_{n}\). Reprinted with permission from ref. 75. Copyright © (2007) American Chemical Society.

**Figure 52.** Fragment of X-ray structure (a) and crystal packing (b) in \([\text{Ag}(\text{bppz})[1-\text{Ph}-1-\text{CB}_{9}\text{H}_{9}]]_{n}\). Reprinted with permission from reference [75]. Copyright © (2007) American Chemical Society.
In contrast to the other complexes with the 1-carba-closo-decaborate anion reported here, the reaction of Ag[1-Ph-1-CB$_9$H$_8$-6-Br] with 2,3-bis(2-pyridyl)pyrazine in acetonitrile produces discrete dimeric complex [Ag(bppz)[1-Ph-1-CB$_9$H$_8$-6-Br]]$_2$ [75] rather than polymeric one. The silver atom has an irregular tetrahedral geometry: Ag(1) is coordinated with two bppz ligands, to form a chelating interaction to one ligand [Ag-N distances of Ag(1)-N(1) 2.332(10) and Ag(1)-N(3) 2.372(9) Å], with the third coordination bond occurring to the pyridyl nitrogen atom of a second bppz ligand (Ag(1)-N(2) 2.268(9) Å). The coordination environment of silver is completed with the bromine atom of the [1-Ph-1-CB$_9$H$_8$-6-Br]$^-$ (Ag(1)-Br(1) 2.9382(11) Å) (Figure 53).

Complex [Ag(DABCO)[1-Ph-1-CB$_9$H$_8$]]$_n$ was prepared by reaction of Ag[1-Ph-1-CB$_9$H$_8$] with 1,4-diazabicyclo[2.2.2]octane in acetonitrile [77]. The silver atom has a distorted tetrahedral geometry with two coordinate bonds to two crystallographic equivalent DABCO ligands with an angle N(1)-Ag(1)-N(2) of 123.22(1)° and Ag-N bonds of 2.2750(18) and 2.2713(18) Å for Ag(1)-N(1) and Ag(1)-N(2), respectively. Furthermore, it interacts with two BH groups of the [1-Ph-1-CB$_9$H$_8$-6-Br]$^-$ anion, the Ag...H-B interaction with equatorial BH group being much stronger (1.8623(4) Å) than the Ag...H-B interaction with apical BH group (2.3529(6) Å) which can be considered as an agnostic interaction. The 1D coordination network is puckered to create the [Ag(DABCO)]$^+$ zigzag chain. At each silver center along the chain the coordinated [1-Ph-1-CB$_9$H$_8$]$^-$ anions point in alternating directions (Figure 54) [77]. These coordination-chain structural features (zigzag chain with the adjacent coordinative anions pointing in opposite directions) are considerably similar to that reported for the complex [Ag(4,4'-Bipy)[1-Ph-1-CB$_9$H$_4$-6,7,8,9,10-I$_5$]-MeCN]$_n$ (see above).

**Figure 53.** X-ray structure of [Ag(bppz)[1-Ph-1-CB$_9$H$_8$-6-Br]]$_2$.

**Figure 54.** Fragment of X-ray structure of [Ag(DABCO)[1-Ph-1-CB$_9$H$_8$]]$_n$. Reprinted with permission from reference [77]. Copyright © (2013) American Chemical Society.
Similar complex with monoiodinated carborane \([\text{Ag(DABCO)}][\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])_n\) was prepared by reaction of \([\text{Ag}[\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}]]\) with 1,4-diazabicyclo[2.2.2]octane in acetonitrile as well \([77]\). The silver atom establishes coordination bonds with two nitrogen atoms from two crystallographically equivalent DABCO ligands (Ag(1)-N(1) and Ag(1)-N(2) bonds of 2.263(2) and 2.260(2) Å, respectively), and the iodine atom of the \([\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])^-\) anion (Ag(1)-I(1) distance of 2.9155(4) Å). The Ag(1)-N and Ag(1)-I coordinative interactions originate the formation of 1D coordination chains with the carborane anions incorporated in the chains. As was observed in the previous case, the coordination chains run along the a axis with some zigzag arrangement. However, in contrast with the previous chain, all the carbaborane anions point in the same direction (Figure 55) \([77]\).

**Figure 55.** Fragment of X-ray structure of \([\text{Ag(DABCO)}][\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])_n\). Reprinted with permission from reference \([77]\). Copyright © (2013) American Chemical Society.

Cyclotrimeratrylene (CTV) and its analogues (Scheme 3) represent a class of shape-persistent C_3-symmetric molecular hosts derived from the tribenzo[a,d,g]cyclononatriene scaffold. CTV is able to form crystalline clathrate complexes with various small organic molecules, in which the solvent guest may be contained in the molecular cavity of cyclotrimeratrylene. CTV and its analogues have also demonstrated an affinity for globular, electron-poor guests, such as fullerenes and carboranes, which form “ball-in-socket” superstructures. Therefore, study of interactions of cyclotrimeratrylene and its analogues with weakly-coordinating carboranes and metallacarboranes is very exciting \([78]\).

Star-burst tetrahedron complex \([\text{Ag}_4(\text{tris(isonic)CTG})_4(\text{MeCN})][\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])_2\cdot 12(\text{MeCN})\) was obtained by the reaction of \([\text{Ag}[\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}]]\) with tris(isonicotinoyl) cyclotrimeratrylene (tris(isonic)CTG) in acetonitrile \([76]\). The two carborane anions coordinate to two external vertices of the tetrahedron, giving a \([\text{Ag}_4\text{L}_4(\text{MeCN})_2][\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])^{2+}\) prism. Each of the four silver centers in the tetrahedron have a slightly different coordination environment. All are coordinated by three pyridyl arms from three different ligands. Two Ag(I) centers are irregular tetrahedral with terminal ligands external to the prism: either the MeCN ligand, or the \([\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])^-\) anion that coordinates through its iodine atom at Ag-I distance of 2.9305(9) Å. The coordinated \([\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])^-\) anion is oriented such that its phenyl ring accepts an edge-to-face π-stacking interaction with a pyridyl ring of one of the ligands with a CH...πcentroid distance of 2.60 Å. One of the other Ag(I) centers forms a weak Ag...H-Beq interaction (2.332 Å) to another \([\text{1-Ph-1-CB}_9\text{H}_8\text{-6-I}])^-\) anion, again external to the prism. The final Ag(I) has an irregular trigonal-planar geometry but with an additional weak coordination to a guest MeCN molecule located inside the prism (Figure 56) \([76]\).
Scheme 3. Cyclotriveratrylene (CTV) and its analogues, tris(isonicotinoyl)cyclotriguaiacylene (tris(isonic)-CTG), tris(3-pyridylmethylamino)cyclotriguaiacylene (tris(3-pyamino)CTG), tris(4-pyridylmethylamino)-cyclotriguaiacylene (tris(4-pyamino)CTG), tris(isonicotinoyl)-tris(propoxy)-cyclotriguaiacylene (tris(isonic)-CTG-Pr), tris(2-quinolylmethyl)-tris(propoxy)-cyclotriguaiacylene (tris(2-quinamino) CTG-Pr).

Figure 56. X-ray structure of the \{Ag_4L_4(MeCN)_2[1-Ph-1-CB_9H_8-6-I]_2\}^{2+} cation. Reproduced from reference [76] with permission from The Royal Society of Chemistry.
5. Silver Complexes with the carba-closo-dodecaborate anion [CB$_{11}$H$_{12}$]$^-$

The 1-carba-closo-dodecaborate anion [CB$_{11}$H$_{12}$]$^-$ was reported for the first time together with the [1-CB$_9$H$_{10}$]$^-$ anion [63,64], however its chemistry received much more extensive development due to its better availability, especially after 2000, when a convenient method of its synthesis avoiding usage of highly toxic and expensive decaborane was proposed [79]. A powerful impetus to the development of chemistry of the [CB$_{11}$H$_{12}$]$^-$ anion was given by the possibility of its use as weakly coordinating anion, that resulted in the synthesis of huge amounts of its derivatives [80,81]. The silver salt of the parent carba-closo-dodecaborate anion, Ag[CB$_{11}$H$_{12}$], was prepared by metathesis of the cesium salt with AgNO$_3$ in aqueous solution crystallized from benzene/hexane solution to give [Ag(η$^1$-benzene)[CB$_{11}$H$_{12}$]-C$_6$H$_6$]$_n$ [82]. The silver atom has a formal coordination number of three, bonding to two carborane anions and one benzene molecule. The interatomic distances Ag...H(B$^5$') and Ag...H(B$^9$) are 1.968 and 1.974 Å, respectively (Ag...B($^5$') and Ag...B($^9$) distances are 2.581(6) and 2.682(5) Å). The benzene molecule is coordinated to the silver atom in a η$^3$-fashion with short Ag-C distance of 2.400(7) Å (Figure 57). The carborane anion acts as a bridging ligand to give an alternating cation-anion chain which is maintained in one dimension throughout the lattice. Noteworthy that, according to IR spectral data, the Ag...HB interactions preserve in the benzene solution (2380 cm$^{-1}$) [82]. As it was revealed by $^{11}$B NMR spectroscopy, the same is true for the solution of Ag[CB$_{11}$H$_{12}$] in dichloromethane [83].

![Figure 57. Fragment of X-ray structure of {Ag($^1$-benzene)[CB$_{11}$H$_{12}$]-C$_6$H$_6$}$_n$. Reprinted with permission from reference [82]. Copyright © (1985) American Chemical Society.](image)

As it was mentioned above, one of the most common ways to obtain compounds with stabilized reactive cations are metathesis reactions of weak or sometimes even covalently bonded halide complexes with silver salts of weakly coordinating anions. In particular, triaryltrim(IV) cation containing O,C,O-coordinating pincer ligand [1-[2,6-(MeO)$_2$C$_6$H$_3$]SnPh$_2$]$_+$[CB$_{11}$H$_{12}$]$^-$ was prepared by the treatment of the corresponding triaryltrim chloride with the equimolar amount of Ag[CB$_{11}$H$_{12}$] in tetrahydrofuran. However, a change in the stoichiometry of the starting compounds (2:3) and solvent (CH$_2$Cl$_2$) resulted in the complex with new complex anion [1-(2,6-(MeO)$_2$C$_6$H$_3$]SnPh$_2$][Ag[CB$_{11}$H$_{12}$]$_3$]. The complex anion consists of the central Ag$^+$ ion, which is surrounded by three [CB$_{11}$H$_{12}$]$^-$ anionic carborane ligands. Each of them is bonded to the Ag$^+$ center via bridging Ag...H(B12) (opposite to the cage carbon) and Ag...H(B8) (adjacent to B12) bonds, which results in a 6-fold coordination of Ag$^+$ by the B-H bonds. The coordination polyhedron around the silver atom approximates a distorted octahedral shape. The Ag...H distances are 2.2388–2.2583 Å (Figure 58). The IR spectrum of the complex contains two bands at 2551 and 2347 cm$^{-1}$, which are diagnostic for ν(BH) and ν(AgHB) stretching, respectively [84].
An even more complex anion was found in the complex $[\text{Ag(IMes)}_2]^2[\text{Ag}_2\{\text{CB}_{11}\text{H}_{12}\}_4]$ formed by the treatment of 1,3-dimesitylimidazol-2-ylidene (IMes) with Ag[CB$_{11}$H$_{12}$] in toluene. The complex anion contains two silver atoms bridged by two [CB$_{11}$H$_{12}$]$^-$ anions in addition to one terminally bound carborane on each metal. The coordination geometry around each silver atom is approximately trigonal prismatic, with each metal center involved in four shorter and two longer interactions with the carborane BH groups. The shorter Ag-H-B interactions vary in distance ranging from 1.96 to 2.23 Å (Ag…H) and 2.586(7) to 2.79(2) Å (Ag…B). The longer interactions are in the range 2.84(2)-2.94(2) Å (Ag…B) (Figure 59) [83].

Figure 58. X-ray structure of $[1-{2,6-(\text{MeO})_2\text{C}_6\text{H}_3}\text{SnPh}_2][\text{Ag}\{\text{CB}_{11}\text{H}_{12}\}_3]$. Reprinted with permission from reference [84]. Copyright © (2006) American Chemical Society.

Figure 59. Cont.
Similar to Ag[1-CB9H5-6,7,8,9,10-Br5], the reaction of Ag[CB11H12] with iridium chloride complex IrCl(CO)(PPh3)2 in toluene results in the square pyramidal Ir\(\rightarrow\)Ag donor-acceptor metal-metal bonded complex \((\text{PPh}_3)_2\text{CO} \text{Ir} \cdot \text{Ag[CB11H12]}\) \([85,86]\). The \([\text{CB11H12}]^-\) anion is coordinated to silver atom through the antipodal to the carbon atom BH group (the Ag…H(B) and Ag…B distances are 1.901 and 2.528 Å, respectively) (Figure 60) \([85]\). The metathesis proceeds only in donor solvents, such as acetone, giving cationic species with coordinated solvent \([86]\).

The long-lived (up to 1 week at room temperature) intermediate complexes \((\text{Cp})\text{Fe(CO)}_2\cdot \text{Ag[CB11H12]}\) with \{\((\text{Cp})\text{Fe(CO)}_2\)\}[\text{CB11H12}]\} as the ultimate product \([86]\) were obtained by reacting Ag[CB11H12] with iron halide complexes \((\text{Cp})\text{Fe(CO)}_2X\) (X = Cl, Br, I) in toluene \([86]\). The similar reaction with \((\text{Cp})\text{Mo(CO)}_3\) in dichloromethane also proceeds through formation of stable dimeric \{\((\text{Cp})\text{Mo(CO)}_3\)\}[\text{Ag[CB11H12]}\}_2 intermediate which was characterized by single crystal X-ray diffraction (Figure 61) \([87,88]\).

![Figure 59. X-ray structure of cationic (a) and anionic (b) parts of \([\text{Ag(IMes)}_2\)\][\text{Ag}_2\text{[CB11H12]}_4\]. Reprinted with permission from reference \([83]\). Copyright © (2002) American Chemical Society.](image1)

![Figure 60. X-ray structure X-ray structure of \{\((\text{PPh}_3)_2\text{CO} \text{Ir} \cdot \text{Ag[CB11H12]}\}\}. Reprinted with permission from reference \([85]\). Copyright © (1987) American Chemical Society.](image2)
The silver salt of C-alkyl substituted carba-closo-dodecaborate anion Ag[1-R-1-CB\(_{11}\)]\(_{11}\) \(\{R = \text{Me} \ [89], \ Et \ [90]\}\) were prepared as well. The crystal structure of [Ag[\(\eta^2\)-benzene][1-Me-1-CB\(_{11}\)]\(_{11}\)]\(_{n}\) resembles structure of the corresponding salt of the parent anion with the silver atom bonding to two carborane anions and one benzene molecule. The interatomic Ag ... Ag(9A) distances are 2.613(1) and 2.613(1) Å, respectively. The benzene molecule is coordinated to the silver atom in asymmetric \(\eta^2\)-fashion with Ag-C distances of 2.460(1) and 2.698(1) Å (Figure 62) \[89\].

In the crystal structure of [Ag[1-1-Et-1-CB\(_{11}\)]\(_{11}\)]\(_{n}\) the silver atom is coordinated to three monocarborane anions. The coordination occurs through BH groups, with the first anion coordinating via BH group in position 7 (Ag(1) ... H(B7) is 2.078 Å), the second anion coordinating via BH group in position 10 (Ag(1) ... H(B10) is 1.943 Å), and with the third anion coordinating via two adjacent BH groups in positions 8 and 9 (Ag(1) ... H(B8) is 2.352 Å and Ag(1) ... H(B9) is 2.301 Å). These results in slightly distorted tetrahedral geometry around the Ag\(^{+}\) cation. The extended solid state structure is characterized by layers that pack parallel to the \(bc\) plane of the unit cell (Figure 63) \[90\].

The silver salt of 12-fluoro-1-carba-closo-dodecaborate anion, Ag[1-CB\(_{11}\)H\(_{11}\)-12-F], was prepared by cation exchange from the corresponding cesium salt. The crystals of the silver salt of 12-fluoro-1-carba-closo-dodecaborate anion [Ag[\(\eta^2\)-benzene]\(_2\)[1-CB\(_{11}\)H\(_{11}\)-12-F]]\(_n\) were obtained by crystallization from benzene. The silver atom has a formal coordination number of four, bonding to two carborane anions and two benzene molecules. Both [1-CB\(_{11}\)H\(_{11}\)-12-F]\(^{-}\) anions are coordinated to the silver atom via BH groups (Ag ... H(B) distances of 2.18 and 2.28 Å), the benzene molecules...
are coordinated to the silver atom in a $\eta^2$-fashion (Figure 64) [91]. Significantly, there is no interaction between the silver ion and the fluorine atom.

Figure 63. The crystal packing in X-ray structure of $\{\text{Ag}[1-\text{Et-1-CB}_{11}\text{H}_{11}]\}_n$.

Figure 64. Fragment of X-ray structure of $\{\text{Ag}(\eta^2\text{-benzene})_2[1\text{-CB}_{11}\text{H}_{11}-12\text{-F}]\}_n$. Reprinted with permission from reference [91]. Copyright © (1995) American Chemical Society.

The silver salt of 12-bromo-1-carba-closo-dodecaborate anion, Ag[1-CB$_{11}$H$_{11}$-12-Br], was prepared using the standard metathesis approach [92]. The crystals $\{\text{Ag}(\eta^1\text{-benzene}) [1\text{-CB}_{11}\text{H}_{11}-12\text{-Br}]\}_n$ were obtained by crystallization from benzene. The silver atom has a formal coordination number of four, bonding to three carborane anions and one benzene molecule. Two $[1\text{-CB}_{11}\text{H}_{11}-12\text{-Br}]^-$ anions are coordinated to the silver atom via BH groups (Ag . . . H(B) distances of 2.20 and 2.27 Å) and the third anion is coordinated by the bromine atom (Ag . . . Br(B) distance of 2.642 Å), the benzene molecule is coordinated to the silver atom in a $\eta^1$-fashion with Ag-C distance of 2.485 Å (Figure 65) [92].
The reaction with \((\text{Cp})\text{Mo}(\text{CO})_3\text{I}\) with \(\text{Ag}[\text{CB}_{11}\text{H}_{11}-12-\text{Br}]\) in dichloromethane produces dimeric complex \(\{[(\text{Cp})\text{Mo}(\text{CO})_3\text{I}]\text{Ag}[\text{CB}_{11}\text{H}_{11}-12-\text{Br}]\}_2\) which was characterized by single crystal X-ray diffraction. Its structure contains a centrosymmetric planar \(\{\text{Ag}_2\text{I}_2\}\) core appended with two \([\text{CpMo}(\text{CO})_3]\) fragments and two bidentate carborane anions. Each carborane anion is coordinated to a silver atom through one \(\text{Ag-Br}\) (2.6456(8) Å) and one \(\text{Ag-HB}\) (\(\text{Ag} \ldots \text{B} 3.159(3)\) Å, \(\text{Ag} \ldots \text{H} 2.43(2)\) Å) bonds. The two metal fragments are orientated \textit{trans} with respect to one another and the central \(\{\text{Ag}_2\text{I}_2\}\) core (Figure 66) [93].

The silver salts of 6,7,8,9,10,11,12-hexahalogeno-1-carba-closo-dodecaborate anions, \(\text{Ag}[1-\text{CB}_{11}\text{H}_6-6,7,8,9,10,11,12-\text{X}_6]\) (\(X = \text{Cl} [94], \text{Br} [86], \text{I} [94]\)), were prepared from the treatment of the corresponding cesium salts with \(\text{AgNO}_3\) in aqueous solution. The crystals of \(\{\text{Ag}(\eta^2-para-xylene)[1-\text{CB}_{11}\text{H}_6-6,7,8,9,10,11,12-\text{Cl}_6]\}_n\) were obtained by crystallization from 1,4-xylene. The silver atom is in the five-coordinated propeller arrangement built of one \(\eta^2-p\)-xylene and two bidentate bridging carborane anions. The \(\text{Ag} \ldots \text{Cl}\) distances range from 2.640(1) to 2.926(1) Å. The \(p\)-xylene is coordinated in asymmetric \(\eta^2\) fashion with \(\text{Ag-C}\) bonds of 2.506(3) and 2.481(4) Å (Figure 67) [94].
The crystals of \(\text{Ag}[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{Br}_{6}]\)_n were obtained by crystallization from toluene/hexane in the presence of \(i\)-Pr_2SiHCl. The silver atom is in distorted octahedral arrangement with two tridentate bridging carborane anions. The bromine atom attached to B(12) is shared by two silver ions. The average Ag…Br distance is 2.862(2) Å (Figure 67) [95]. In the crystals of \(\text{Ag}[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{I}_{6}]\cdot0.5(\text{benzene})\)_n the silver atom is complexed by iodine atoms to one anion in bidentate fashion and to another in tridentate fashion adopting a square-pyramidal arrangement (Figure 67) [95].

![Figure 67. Fragments of X-ray structure of \(\text{Ag}(\eta^2-\text{para-xylene})[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{Cl}_{6}]\)_n (a), \(\text{Ag}[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{Br}_{6}]\)_n (b) and \(\text{Ag}[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{I}_{6}]\cdot0.5(\text{benzene})\)_n (c). Reproduced from reference [95] with permission from The Royal Society of Chemistry.](image)

The discrete \(\text{Ag}[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{Cl}_{6}]\)\(^-\) anions with six-coordinated silver atom is in an distorted octahedral arrangement (Figure 68) were found in the X-ray structure of the azafullerene carbocation salt \([\text{C}_{59}\text{N}]\lfloor\text{Ag}[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{Cl}_{6}]\rfloor_2\) 3(\(o\)-dichlorobenzene) which was prepared by reaction of \((\text{C}_{59}\text{N})_2\) dimer with \([\text{HBPC}·\text{Ag}[\text{1-CB}_{11}\text{H}_{6}-6,7,8,9,10,11,12-\text{Cl}_{6}]\rfloor_2\) (HBPC = hexabromo(phenyl)carbazole) in \(o\)-dichlorobenzene [96].
Figure 68. Structure of the \(\text{[Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Cl}_{6}]]}^{-}\) anion in \([C_{60}N]\text{[Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Cl}_{6}]]}\cdot3(\text{o-dichlorobenzene})\). Reprinted with permission from reference [96]. Copyright © (2003) American Chemical Society.

The similar discrete \(\text{[Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}]]}^{-}\) anions (Figure 69) were found in the X-ray structure of the \([\text{Fe(TPP)}][\text{Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}]]_{2}]\cdot4(p\text{-xylene})\) complex which was prepared by the treatment of \([\text{Fe(TPP)}]\text{Br}\) with \(\text{Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}]}\) in \(p\text{-xylene}\) at 100 °C, subsequent removal of the resulting AgBr precipitate, and vapor diffusion of \(n\text{-hexane}\) [97].

Figure 69. Structure of the \(\text{[Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}]}^{-}\) anion in complex \([\text{Fe(TPP)}][\text{Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}]]_{2}]\cdot4(p\text{-xylene})\). Reproduced with permission from reference [97]. Copyright © (1994) John Wiley & Sons, Inc.

Similar to Ag\([\text{CB}_{11}\text{H}_{12}]\), the reaction of Ag\([1\text{-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}}]\) with iridium chloride complex IrCl(O)\((\text{PPh}_{3})_{2}\) in fluorobenzene results in the donor-acceptor metal-metal bonded complex \((\text{PPh}_{3})_{2}(\text{CO})\text{ClIr}\cdot\text{Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}]_{2}}\) identified by IR spectroscopy [86].

The similar reaction with \((\text{Cp})\text{Mo(CO)_{3}I}\) in dichloromethane also proceeds through formation of stable dimeric \([[(\text{Cp})\text{Mo(CO)_{3}I}\text{Ag[1-CB}_{11}\text{H}_{6-6,7,8,9,10,11,12-\text{Br}_{6}]]_{2}]\) intermediate which was characterized by single crystal X-ray diffraction. Its structure bears close similarities with that found in \([[(\text{Cp})\text{Mo(CO)_{3}I}\text{Ag[1-CB}_{11}\text{H}_{12}]]_{2}]\), with a centrosymmetric planar \(\text{[Ag_{2}I_{2}}]\) core appended with two \((\text{Cp})\text{Mo(CO)_{3}I}\) fragments and two bidentate carborane anions. The two metal fragments are orientated \(\text{trans}\) with respect to one another and the central \(\text{[Ag_{2}I_{2}}]\) core (Figure 70) [88].
The silver salts of C-methyl hexahalogen carba-closo-dodecaborate anions Ag[1-Me-1-CB_{11}H_{5-6,7,8,9,10,11,12-X_6}] (X = Cl, Br, I) were prepared using standard metathesis methodology. The crystal structure of \{Ag(\eta^1\text{-mesitylene})[1-Me-1-CB_{11}H_{5-6,7,8,9,10,11,12-Cl_6}]\}_n resembles structure of \{Ag(\eta^2\text{-p-xylene})[1-CB_{11}H_{5-6,7,8,9,10,11,12-Cl_6}]\}_n with five-coordinated silver atom in a distorted square pyramidal arrangement of one \eta^1\text{-mesitylene} and two bidentate bridging [1-Me-CB_{11}H_{5-6,7,8,9,10,11,12-Cl_6}]^{-} anions. The Ag . . . Cl distances range from 2.683(2) to 2.874(2) Å (Figure 71) [89]. The solid state structure of \{Ag[1-Me-1-CB_{11}H_{5-6,7,8,9,10,11,12-Br_6}]\}_n is almost identical to that of \{Ag(1-CB_{11}H_{5-6,7,8,9,10,11,12-Br_6})\}_n with the average Ag . . . Br distance of 2.858(1) Å (Figure 71) [89]. In the solid state structure of \{Ag[1-Me-1-CB_{11}H_{5-6,7,8,9,10,11,12-I_6}]\}_n the silver atom is complexed by four iodine atoms from two bidentate bridging [1-Me-1-CB_{11}H_{5-6,7,8,9,10,11,12-I_6}]^{-} anions in a distorted tetrahedral arrangement with the Ag . . . I distances varying from 2.798(2) to 2.872(2) Å (Figure 71) [89].
The silver salts of the perchloro and perbromo derivatives of 1-carba-closo-dodecaborate anion, Ag[1-HCB\(_{11}\)Cl\(_{11}\)] and Ag[1-HCB\(_{11}\)Br\(_{11}\)], were prepared by treatment of the corresponding sodium salt with AgNO\(_3\) in aqueous solution. Single crystals of {[(\(\eta^1\)\(\eta^1\)-p-xylene)\(_2\)Ag[1-HCB\(_{11}\)Cl\(_{11}\)]]·1/2(p-xylene) were grown from p-xylene. The complex has a monomeric structure with four-coordinated silver atom in a propeller arrangement of two \(\eta^2\)-p-xylene molecules and a single bidentate [1-HCB\(_{11}\)Cl\(_{11}\)]\(^-\) anion (Figure 72a) \cite{99}. Single crystals of [(\(\eta^1\)-mesitylene)(MeCN)Ag[1-HCB\(_{11}\)Br\(_{11}\)] were grown from a mesitylene-acetonitrile solution. The silver atom has a distorted tetrahedral arrangement consisting of one \(\eta^1\)-bound mesitylene molecule, one acetonitrile molecule and two bromine atoms of the bidentate [1-HCB\(_{11}\)Br\(_{11}\)]\(^-\) anion (Ag…Br bonds are 2.772(1) and 2.776(1) Å) (Figure 72b) \cite{99}.

Silver salts of mixed halocarborane anions Ag[1-HCB\(_{11}\)-2,3,4,5,6-Br\(_5\)-7,8,9,10,11,12-Cl\(_6\)], Ag[1-HCB\(_{11}\)-2,3,4,5,6-I\(_5\)-7,8,9,10,11,12-Cl\(_6\)], Ag[1-HCB\(_{11}\)-2,3,4,5,6-Cl\(_5\)-7,8,9,10,11,12-Br\(_6\)] and Ag[1-HCB\(_{11}\)-2,3,4,5,6-Br\(_5\)-7,8,9,10,11,12-Cl\(_6\)] were prepared by treatment of the corresponding sodium salts with AgNO\(_3\) in aqueous solution. Single crystals of {[(\(\eta^2\)-mesitylene)\(_2\)Ag[1-HCB\(_{11}\)-2,3,4,5,6-Br\(_5\)-7,8,9,10,11,12-Cl\(_6\)]·mesitylene and {[(\(\eta^1\)-mesitylene) Ag[1-HCB\(_{11}\)-2,3,4,5,6-Cl\(_5\)-7,8,9,10,11,12-Br\(_6\)]·mesitylene}\(_n\) were grown from mesitylene and mesitylene/acetone solutions, respectively. Similar to structure of {[(\(\eta^2\)-p-xylene)\(_2\)Ag[1-HCB\(_{11}\)Cl\(_{11}\)]·1/2(p-xylene), [(\(\eta^2\)-mesitylene)\(_2\)Ag[1-HCB\(_{11}\)-2,3,4,5,6-Br\(_5\)-7,8,9,10,11,12-Cl\(_6\)]·mesitylene has monomeric structure with four-coordinated silver atom in a propeller arrangement of two \(\eta^2\)-mesitylene molecules and a single bidentate [1-HCB\(_{11}\)Br\(_5\)Cl\(_6\)]\(^-\) anion which is coordinated through its Cl(8) and Cl(12) atoms (the Ag…Cl(8) and Ag…Cl(12) distances are 2.986(2) and 2.889(2) Å, respectively) (Figure 73) \cite{99}. It is noteworthy that no interaction between Ag\(^+\) and any one of the bromine atoms from the upper belt of the cage is observed, although Ag\(^+\) usually prefers to bind to a bromine atom rather than a chlorine atom according to the hard-soft acid-base principle.
These results may suggest that the charge distribution on the icosahedral cage plays a predominant role in the coordination chemistry of carborane anions. The four-coordinated silver atom is in a distorted tetrahedral arrangement of one \( \eta^1 \)-mesitylene molecule, two bromine atoms from a bridging \([1-\text{HCB}_{11}\text{Cl}_{11}]^{-}\) anion and one bromine atom from another bridging \([1-\text{HCB}_{11}\text{Br}_{11}]^{-}\) anion (the Ag...Br distances of 2.750(2)–2.873(1) Å) (Figure 74) [99]. There are no interactions between Ag\(^+\) and any one of the five chlorine atoms from the upper belt of the carborane cage, which is understandable because Br\(^-\) is a softer base than Cl\(^-\) and the lower belt atoms bear more negative charge than the upper belt ones. These results may suggest that the charge distribution on the icosahedral cage plays a predominant role in the coordination chemistry of carborane anions.

In contrast, a polymeric structure is observed in \((([\eta^1 \text{-mesitylene}]\text{Ag}[1-\text{HCB}_{11}-2,3,4,5,6-\text{Cl}_5-7,8,9,10,11,12-\text{Br}_6])_n\) mesitylene\(_n\). The four-coordinated silver atom is in a distorted tetrahedral arrangement of one \( \eta^1 \)-mesitylene molecule, two bromine atoms from a bridging \([1-\text{HCB}_{11}\text{Cl}_{11}\text{Br}_{6}]^{-}\) anion and one bromine atom from another bridging \([1-\text{HCB}_{11}\text{Br}_{11}]^{-}\) anion (the Ag...Br distances of 2.750(2)–2.873(1) Å) (Figure 74) [99]. There are no interactions between Ag\(^+\) and any one of the five chlorine atoms from the upper belt of the carborane cage, which is understandable because Br\(^-\) is a softer base than Cl\(^-\) and the lower belt atoms bear more negative charge than the upper belt ones. These results may suggest that the charge distribution on the icosahedral cage plays a predominant role in the coordination chemistry of carborane anions.

The silver salts of C-methyl perhalogen derivatives of the carba-closo-dodecaborate anion Ag\([1-\text{Me}-1\text{-CB}_{11}X_{11}]\) (X = Cl, Br) were prepared by treatment of the sodium salts with AgNO\(_3\) in...
aqueous solution [98]. Single crystals of [Ag[1-Me-1-CB11Cl11]]n were grown from benzene solution. As expected, the complex has polymeric structure. Each silver atom coordinates to six chlorine atoms from the three [1-Me-CB11Cl11]− anions in a distorted-octahedral arrangement at an average Ag . . . Cl distance of 2.882(1) Å, and each anion is surrounded by six silver atoms in a regular octahedral arrangement to form a three-dimensional coordination polymer (Figure 75) [99].

Figure 75. The coordination sphere of the Ag atom in [Ag[1-Me-1-CB11Cl11]]n (a) and the coordination sphere around the carborane anion in [Ag[1-Me-1-CB11Cl11]]n (b). Reprinted with permission from reference [99]. Copyright © (2000) American Chemical Society.

Single crystals of [Ag[1-Me-1-CB11Br11]]n were grown from a chlorobenzene solution. Like many other silver(I) salts of icoshedral carborane anions, it is a one-dimensional coordination polymer with the carborane anions acting as bridging ligands. The silver atom has six-coordinate geometry which is very similar to that found in [Ag[1-HCB11H5-7,8,9,10,11,12-Br6]]n with the metal atom bonding to three bromine atoms from each of two bridging [1-Me-CB11Br11]− anions (the average Ag . . . Br distance of 2.848(5) Å) (Figure 76) [98]. The methyl group is completely mixed with eleven bromine atoms of the cage anion. Therefore, it is not possible to distinguish the BBr vertex antipodal to carbon from other vertexes. Unlike in the solid state structure of [Ag[1-HCB11H5-7,8,9,10,11,12-Br6]]n, where the most basic bromine atom attached to B(12) antipodal to carbon is shared by two silver ions, there is no bromine atom shared by the silver ion in [Ag[1-Me-1-CB11Br11]]n, perhaps reflecting the steric effect and/or the fact that there is no significantly electron-rich bromine atom on the cage due to donor effect of the methyl group.

Figure 76. Fragment of X-ray structure of [Ag[1-Me-1-CB11Br11]]n. Reprinted with permission from reference [98]. Copyright © (1998) American Chemical Society.

Fractional crystallization of Ag[1-Me-1-CB11H11-nBrn] (n = 8–10) from a hot p-xylene solution containing a very small amount of acetonitrile gave X-ray-quality crystals of [(p-xylene)
Fractional crystallization of Ag\([1-\text{Me-CB}_{11}\text{H}_{11}-n\text{Br}_n]\) from a hot...

...anion Ag\([1-\text{Me-CB}_{11}\text{F}_{11}]\) was prepared by metathesis reaction of the corresponding cesium salt and Ag\(\text{NO}_3\)

...prepared by treatment of the corresponding sodium salt with Ag\(\text{NO}_3\) in aqueous solution. Single...

...Ag-C bonds of 2.514(5) and 2.485(5) Å (Figure 77) [98].

The silver salt of the C-butyl perchloro-carba-closo-dodecaborate anion Ag\([1-\text{Bu-1-CB}_{11}\text{Cl}_{11}]\) was prepared by treatment of the corresponding sodium salt with Ag\(\text{NO}_3\) in aqueous solution. Single crystals of \([\text{Ag}(\eta^2-\text{C}_6\text{H}_5\text{F})(\text{H}_2\text{O})][1-\text{Bu-1-CB}_{11}\text{Cl}_{11}]\) were grown from fluorobenzene solution. The silver atom is coordinated by \(\eta^2\text{-fluorobenzene}\) (the Ag-C bonds of 2.553(3) and 2.587(3) Å), water (the Ag-O bond of 2.317(2) Å), one bidentate (the Ag . . . Cl distances are 2.847(1) and 2.878(1) Å) and one monodentate (the Ag . . . Cl distance is 3.127(1) Å) bridging \([1-\text{Bu-1-CB}_{11}\text{Cl}_{11}]\) anions (Figure 78) [100].

Figure 77. Fragment of X-ray structure of \([\text{Ag}(\eta^2-\text{para-xylene})][1-\text{Me-CB}_{11}\text{Br}_{10}]\)\(_n\). Reprinted with permission from reference [98]. Copyright © (1998) American Chemical Society.

The silver salt of the C-butyl perchloro-carba-closo-dodecaborate anion Ag\([1-\text{Bu-1-CB}_{11}\text{Cl}_{11}]\) was prepared by treatment of the corresponding sodium salt with Ag\(\text{NO}_3\) in aqueous solution. Single crystals of \([\text{Ag}(\eta^2-\text{C}_6\text{H}_5\text{F})(\text{H}_2\text{O})][1-\text{Bu-1-CB}_{11}\text{Cl}_{11}]\)\(_n\) were grown from fluorobenzene solution. The silver atom is coordinated by \(\eta^2\text{-fluorobenzene}\) (the Ag-C bonds of 2.553(3) and 2.587(3) Å), water (the Ag-O bond of 2.317(2) Å), one bidentate (the Ag . . . Cl distances are 2.847(1) and 2.878(1) Å) and one monodentate (the Ag . . . Cl distance is 3.127(1) Å) bridging \([1-\text{Bu-1-CB}_{11}\text{Cl}_{11}]\) anions (Figure 78) [100].

Figure 78. Fragment of X-ray structure of \([\text{Ag}(\eta^2-\text{C}_6\text{H}_5\text{F})(\text{H}_2\text{O})][1-\text{Bu-1-CB}_{11}\text{Cl}_{11}]\)\(_n\). Reproduced from reference [100] with permission from The Royal Society of Chemistry.
AgNO₃ in refluxing benzene [101]. Single crystals of [Ag(CHPh₃)[1-Me-1-CB₁₁F₁₁]]ₙ were obtained from the reaction of Ag(C₆H₆)[1-Me-1-CB₁₁F₁₁] with Ph₃CCl in dichloromethane. The structure consists of layers of [Ag(CHPh₃)⁺]ₙ polycations with [1-Me-1-CB₁₁F₁₁]⁻ anions filling half of the trigonal prismatic holes defined by six Ag⁺ ions from successive [Ag(CHPh₃)⁺]ₙ layers. The [Ag(CHPh₃)⁺]ₙ triangular-net layers contain the first example of a rigorously trigonal planar Ag(arene)₃⁺ complex. In the [Ag(CHPh₃)⁺]ₙ layer three phenyl groups from three different CHPh₃ ligands in asymmetric η²-fashion coordinate to the Ag⁺ ion being in trigonal planar arrangement. There is no bonding between the Ag⁺ cations and the [1-Me-1-CB₁₁F₁₁]⁻ anions (Figure 79) [101].

Figure 79. The coordination sphere of the Ag atom (a) and crystal packing (b) in the structure of [Ag(CHPh₃)[1-Me-1-CB₁₁F₁₁]]ₙ. Reprinted with permission from reference [101]. Copyright © (2004) American Chemical Society.

The silver salt [Ag[1-Me-1-CB₁₁F₁₀-12-SiPh₃]·0.5(benzene)]ₙ was obtained by metathesis of the corresponding cesium salt and AgBF₄ followed by crystallization from benzene solution. The structure contains two types of silver cations in different arrangement. The coordination sphere of one of Ag⁺ cations is trigonal planar and consists of two η²-coordinated phenyl rings of two different [1-Me-1-CB₁₁F₁₀-12-SiPh₃]⁻ anions and one η¹-coordinated benzene molecule. The other Ag⁺ cation is coordinated in η¹-fashion by four phenyl groups of two different [1-Me-1-CB₁₁F₁₀-12-SiPh₃]⁻ anions forming the tetrahedral coordination sphere (Figure 80) [100]. Again, there are no Ag…F interactions between the Ag⁺ cations and the [1-Me-1-CB₁₁F₁₀-12-SiPh₃]⁻ anions.

Figure 80. The coordination sphere of Ag atoms in the structure of [Ag[1-Me-1-CB₁₁F₁₀-12-SiPh₃]·0.5(benzene)]ₙ. Reprinted with permission from reference [102]. Copyright © (2007) American Chemical Society.

Copper(I) complexes (CO)₂Cu[1-Bn-1-CB₁₁F₁₁] and [Cu(CO)₄][1-Et-1-CB₁₁F₁₁] were prepared by treating the corresponding silver salts with CuCl in dichloromethane under an CO atmosphere.
In the first complex the copper(I) dicarbonyl moiety is coordinated in asymmetrical \( \eta^2 \)-fashion by the phenyl ring of the \([1\text{-Bn-1-CB}_{11}\text{F}_{11}]^-\) anion (Figure 81a), whereas in the second complex there is no interactions between the \([\text{Cu(CO)}_4]^+\) cation and the \([1\text{-Et-1-CB}_{11}\text{F}_{11}]^-\) anion (Figure 81b) [103]. Again, there are no Ag \ldots F interactions between the Cu\(^+\) cations and the fluorinated carborane anions that can be explained taking in the account the hard-soft acid-base principle.

![Figure 81. X-ray structures of complexes (CO)\(_2\)Cu[1-Bn-1-CB\(_{11}\)F\(_{11}\)] (a) and [Cu(CO)\(_4\)][1-Et-1-CB\(_{11}\)F\(_{11}\)] (b). Reprinted with permission from reference [103]. Copyright © (1999) American Chemical Society.](image)

The silver salt of the permethylated carba-closo-dodecaborate anion \(\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]\) was prepared by metathesis reaction of the corresponding cesium salt and \(\text{AgNO}_3\) in water-\(\text{Et}_2\text{O}\). Single crystals of \(\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]_n\) were obtained by crystallization from dichloromethane/hexanes. One tridentate and three monodentate carborane anions affords a distorted octahedral coordination environment around the silver cation. Each carborane anion is closely associated with four silver cations. This results in a three-dimensional coordination polymer (Figure 82). The Ag \ldots C distances span the range 2.652(2)–3.089(2) Å [104].

![Figure 82. The coordination sphere of the Ag atom in \(\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]_n\) (a) and the coordination sphere around the carborane anion in \(\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]_n\) (b). Reprinted with permission from reference [104]. Copyright © (2004) American Chemical Society.](image)
In contrast to the silver salts of the parent carba-closo-dodecaborate anion and its halogenderivatives, the reaction of Ag[HCB$_{11}$HMe$_{11}$] with (Cp)Mo(CO)$_3$I in dichloromethane results in complex [{(Cp)Mo(CO)$_3$I$_2$}Ag$_2$][HCB$_{11}$HMe$_{11}$)$_2$], where each silver atom in the central [Ag$_2$I$_2$] core is surrounded by three {(Cp)Mo(CO)$_3$I} units, one of which bridges the two silver centres (Figure 83) [93].

Single crystals of {(MeCN)$_2$Ag[CB$_{11}$H$_{12}$]}$_n$ were grown by crystallization of Ag[CB$_{11}$H$_{12}$] from acetonitrile/trifluoroethanol solution. The silver atom is linearly coordinated by two crystallographically equivalent acetonitrile ligands at the Ag-N distance 2.172(5) Å and N-Ag-N angle of 157.6(2)$^\circ$. There are four additional interactions to BH groups of three surrounding [CB$_{11}$H$_{12}$]$^-$ anions at Ag...H-B distances 2.421 and 2.405 Å forming a distorted octahedral coordination environment (Figure 84a). Each [CB$_{11}$H$_{12}$]$^-$ anion forms Ag...H-B interactions with three silver centers, and every silver atom interacts with three [CB$_{11}$H$_{12}$]$^-$ anions forming a 2D network. These networks pack in the crystal lattice through interdigitation of the terminal acetonitrile ligands (Figure 84b) [105].

Crystals of [Ag(NCCH$_2$CN)$_2$[CB$_{11}$H$_{12}$]]$_2$ were grown by crystallization of Ag[CB$_{11}$H$_{12}$] from a malonitrile/ethanol solution. The complex is a discrete dimeric complex. There are two types of the malonitrile ligand. One type of the ligand bridges between two Ag$^+$ cations to form a cyclic dimer.
other coordinates in a terminal fashion with a pendant nitrile group that is not involved in coordination with adjacent dimers. The silver atom has a distorted tetrahedral geometry with coordination by three malonitrile nitrogen atoms and one Ag···H-B interaction (2.038 Å) (Figure 85) [105].

![Figure 85. The X-ray structure of \{Ag(NCCH\(_2\)CN)\(_2\)[CB\(_{11}\)H\(_{12}\)]\}_2.](image)

Single crystals of \{Ag\(_4\)(NC(CH\(_2\)\(_2\))CN)\(_5\)[CB\(_{11}\)H\(_{12}\)]\(_4\}\)_n were grown by crystallization of Ag[CB\(_{11}\)H\(_{12}\)] from succinonitrile/ethanol solution. The silver cations fall into two groups, either coordinated by four succinonitrile ligands (Ag\(_2\), Ag\(_3\), Ag\(_5\)) or by one succinonitrile ligand and two [CB\(_{11}\)H\(_{12}\)]\(^-\) anions (Ag\(_1\), Ag\(_4\), Ag\(_6\)). The first type of silver cations propagates a coordination chain. All have distorted tetrahedral geometry with Ag-N distances ranging from 2.194 to 2.350(9) Å and N-Ag-N angles ranging from 97.8 to 121.3(3)°. Silver cations of the other type are not involved in propagating the chain structure but are effectively terminal branches from the chain. Each is connected to the chain via a single bridging succinonitrile, and the other coordination sites are occupied by Ag···H-B interactions. For Ag\(_1\), the coordination environment is distorted tetrahedral with three Ag···H-B bonds of 1.971, 2.062, and 2.139 Å. The coordination environment around Ag\(_4\) is very similar to that of Ag\(_1\), while for Ag\(_6\) the environment shows two short Ag···H-B interactions and two long Ag···H-B interactions at around 2.5 Å (Figure 86) [105].

![Figure 86. The coordination sphere of the Ag atoms (a) and crystal packing (b) in the structure of \{Ag\(_4\)(NC(CH\(_2\)\(_2\))CN)\(_5\)[CB\(_{11}\)H\(_{12}\)]\(_4\}\)_n.](image)

Complex \{Ag(pyz)[CB\(_{11}\)H\(_{12}\)]\}_n was prepared by the reaction of Ag[CB\(_{11}\)H\(_{12}\)] with pyrazine in acetonitrile [106]. The silver cation is coordinated to two crystallographically equivalent pyrazine ligands at Ag-N distance of 2.203(3) Å in an almost linear fashion (N-Ag-N angle 174.76(14)°). There are also four Ag···H-B interactions between the silver cation and four [CB\(_{11}\)H\(_{12}\)]\(^-\) anions with Ag···H-B distances in the range 2.287–2.460 Å. Each pyrazine molecule bridges between two silver cations to form an infinite planar chain, which runs along the b-axis. Each carborane anion interacts with four
silver cations. On the whole, the coordination network has 3D pillared checkerboard arrangement (Figure 87).

Complex \([\text{Ag}(4,4\text{-Bipy})][\text{1-Ph-1-CB}_{11}\text{H}_{5-7,8,9,10,11,12-I_{6}}]\cdot\text{MeCN}]_{n}\) was prepared by reaction of \(\text{Ag}[\text{1-Ph-1-CB}_{11}\text{H}_{5-7,8,9,10,11,12-I_{6}}]\) with 4,4\text{-bipyridine} in acetonitrile [75]. Structure of the complex is similar to those found in complex \([\text{Ag}(4,4\text{-Bipy})][\text{1-Ph-1-CB}_{9}\text{H}_{4-6,7,8,9,10-I_{5}}]\cdot\text{MeCN}]_{n}\). Each silver atom is coordinated through two Ag-N bonds of two crystallographically distinct 4,4\text{-Bipy} ligands at distances of Ag(1)-N(1) 2.274(6) and Ag(1)-N(3) 2.242(7) Å and by a chelating interaction with the \([\text{1-Ph-1-CB}_{11}\text{H}_{5-I_{6}}]\)\text{`} anion resulting from two Ag-I coordination bonds. Each 4,4\text{-Bipy} ligand bridges between two silver cations giving a zigzag coordination chain where the coordinated \([\text{1-Ph-1-CB}_{11}\text{H}_{5-I_{6}}]\)\text{`} anion points in two alternating directions above or below the chain (Figure 88) [75].
Complexes $\{[\text{Ag}_2(\text{bppz})_3][\text{CB}_{11}\text{H}_{12}]_2\cdot\text{MeCN}]_n$ [106] and $\{[\text{Ag}_2(\text{bppz})_3][1-(4-\text{BrC}_6\text{H}_4)-1-\text{CB}_{11}\text{H}_{11}]_2\cdot1.5(\text{MeCN})]_n$ [75] were prepared by the corresponding silver salts with 2,3-bis(2-pyridyl)pyrazine in acetonitrile. Both complexes contain virtually identical $\{[\text{Ag}_2(\text{bppz})_3]^{2+}]_n$ coordination chains. Each silver cation has a distorted tetrahedral geometry and is coordinated by N-atoms belonging at three different bppz molecules with one chelating interaction per metal atom. Two types of ligands with different binding motifs can be distinguished. One bridges between two silver cations with a chelating interaction to one metal and a single bond through a pyridine nitrogen atom to the other. There are two such ligands, forming a double bridge between two adjacent silver cations. The other type of bppz ligand forms a single bridge between two silver cations and coordinates only through its pyridine nitrogen atoms (Figure 89).

![Figure 89. Fragment of the infinite $\{[\text{Ag}_2(\text{bppz})_3]^{2+}]_n$ chain of the complex $\{[\text{Ag}_2(\text{bppz})_3][1-(4-\text{BrC}_6\text{H}_4)-1-\text{CB}_{11}\text{H}_{11}]_2\cdot1.5(\text{MeCN})]_n$. Reprinted with permission from reference [75]. Copyright © (2013) American Chemical Society.](image)

Complex $\{\text{Ag}(\text{DABCO})[1-\text{Ph-1-CB}_{11}\text{H}_{5}-7,8,9,10,11,12-\text{I}_6]\}_n$ was prepared by reaction of Ag[1-Ph-1-CB$_{11}$H$_{5}$-7,8,9,10,11,12-I$_6$] with 1,4-diazabicyclo[2.2.2]octane in acetonitrile [77]. The silver atom has a slightly distorted tetrahedral geometry with two coordinate bonds to two crystallographic equivalent DABCO ligands with an angle N-Ag-N of 111.40(14)° and Ag-N bonds of 2.346(4) and 2.321(4) Å. Furthermore, it interacts with two iodine atoms of the [1-Ph-1-CB$_{11}$H$_5$I$_6$]$^-$ anion with an angle I-Ag-I of 95.41(3)° and Ag…I distances of 2.8276(9) and 2.817(1) Å. The $\{\text{Ag}(\text{DABCO})[1-\text{Ph-1-CB}_{11}\text{H}_{5}-7,8,9,10,11,12-\text{I}_6]\}_n$ coordination chains are comparable to those observed in $\{\text{Ag}(\text{DABCO})[1-\text{Ph-1-CB}_{11}\text{H}_{5}]\}_n$ (Figure 90) [77].

![Figure 90. Fragment of X-ray structure of $\{\text{Ag}(\text{DABCO})[1-\text{Ph-1-CB}_{11}\text{H}_{5}-7,8,9,10,11,12-\text{I}_6]\}_n$. Reprinted with permission from reference [77]. Copyright © (2013) American Chemical Society.](image)

Complex $\{(\text{Py})_2\text{Ag}[1-\text{HCB}_{11}\text{Br}_5\text{I}_6]\cdot\text{Py}\}$ was obtained by recrystallization of Ag[1-HCB$_{11}$Br$_5$I$_6$] from pyridine/acetonitrile. The four-coordinated silver atom is in a distorted-tetrahedral arrangement of two iodine atoms from the [1-HCB$_{11}$Br$_5$I$_6$]$^-$ anion (Ag-I bonds 2.894(1) and 2.835(1) Å) and two pyridine molecules (Ag-N bonds 2.258(11) and 2.267(4) Å). It is interesting to note that the third
pyridine molecule points toward the C-H cage proton with the N⋯C distance of 3.061(15) Å and C-H-N angle of 172(1)°, indicating there is a hydrogen bonding between pyridine molecule and the acidic C-H cage proton (Figure 91) [107].

Capsule-like complex {[Ag2(tris(pyamino)CTG)2(MeCN)2][CB11H12]2} was obtained by co-crystallization of Ag[CB11H12] with tris(3-pyridylmethylamino)cyclotriguaiacylene cavitand (tris(pyamino)CTG) from acetone. It structure is very close to those found in the complex with cobalt bis(dicarboxamide) anion which will be discussed below [108].

Triphenylphosphine complex {(Ph3P)Ag[CB11H12]} was prepared by the reaction of Ag[CB11H12] with equimolar amount of PPh3 in dichloromethane, whereas the excess of phosphine results in the formation of dimeric complex {(Ph3P)2Ag[CB11H12]}2 [109,110]. In the complex {(Ph3P)Ag[CB11H12]} the {AgPPh3}+ fragment interacts with the carborane anion through the three BH units of triangular face that is the most distant from the carbon atom with Ag⋯B distances of 2.504(1)-2.619(2) Å (Figure 92a) [109,110]. In the complex {(Ph3P)2Ag[CB11H12]}2 each carborane unit bridges two silver centers. Each boron cage interacts with the two silver cations through one shorter (Ag(1)-H(12′) 2.17(2) Å, Ag(1)-B(12′) 2.892(2) Å) and one longer (Ag(1)-H(7) 2.51(2) Å, Ag(1)-B(7) 3.494(2) Å) Ag⋯H⋯B interactions (Figure 92b) [110]. Notably, according to 11B NMR spectral data, the Ag⋯H⋯B interactions are preserved in dichloromethane solution of {(Ph3P)Ag[CB11H12]} but not in {(Ph3P)2Ag[CB11H12]}2, which reflects the significantly weaker interactions in the dimeric complex [110].
In a similar way, complex \( \{(\text{Ph}_3\text{P})\text{Ag}[\text{HCB}_{11}\text{H}_5\text{-7,8,9,10,11,12-Br}_6]\}\) was prepared by the reaction of \( \text{Ag}[\text{HCB}_{11}\text{H}_5\text{-7,8,9,10,11,12-Br}_6]\) with the equimolar amount of triphenylphosphine in dichloromethane, whereas the excess of phosphine give complex \( \{(\text{Ph}_3\text{P})_2\text{Ag}[\text{HCB}_{11}\text{H}_5\text{-7,8,9,10,11,12-Br}_6]\}\) \([109,110]\). The silver center in \( \{(\text{Ph}_3\text{P})\text{Ag}[\text{HCB}_{11}\text{H}_5\text{Br}_6]\}\) is coordinated with the carborane anion through two shorter (2.6963(5) Å and 2.8710(5) Å) \(\text{Ag} \cdots \text{Br}\) interactions and one longer one (Ag(1)-B(12) 3.0953(5) Å). There is also a long interaction (3.4901(5) Å) between Ag(1) and Br(11') on a symmetry related cage in the lattice; this bromine atom approaches Ag(1) approximately trans to Br(12) (Br(12)-Ag(1)-Br(11') 157.15(1)°). Although this distance is long, the coordination motif around the silver suggests that it is significant in the solid state. This interaction completes the coordination sphere at the silver center (Figure 93) \([110]\).

Figure 92. X-ray structures of \( \{(\text{Ph}_3\text{P})\text{Ag}[\text{CB}_{11}\text{H}_{12}]\}\) (a) and \( \{(\text{Ph}_3\text{P})_2\text{Ag}[\text{CB}_{11}\text{H}_{12}]\}_2 \) (b). Reproduced with permission from reference \([110]\). Copyright © (2002) John Wiley & Sons.

Figure 93. Fragment of X-ray structure of \( \{(\text{Ph}_3\text{P})\text{Ag}[\text{HCB}_{11}\text{H}_5\text{-7,8,9,10,11,12-Br}_6]\}\). Reproduced with permission from reference \([110]\). Copyright © (2002) John Wiley & Sons.
In a similar way, a series of complexes \{(R_3P)Ag[HCB_{11}Me_{11}]\} (R = C_6H_5, C_6H_{11}, C_6H_3-3,5-Me_2) was prepared by reactions of Ag[HCB_{11}Me_{11}] with the corresponding phosphines in dichloromethane [104,111]. In the structure of \{(P_3P)Ag[HCB_{11}Me_{11}]\} there is a close contact between the silver cation and carborane anion through Ag...MeB interaction (Ag(1)-C(7) 2.544(2) Å). There are two more, albeit substantially longer, contacts from adjacent carborane anions in the lattice Ag(1)...C(9') 3.154(2) Å and Ag(1)...C(12'') 3.336(2) Å (Figure 94) [104,111]. Similarly in the structure of \{(Cy_3P)Ag[HCB_{11}Me_{11}]\} there is a single close Ag...MeB contact, this time from the antipodal B(12)Me vertex (Ag-C(12) 2.608(2) Å). In the lattice there are two other relatively close Ag-C distances resulting in an approximate C_{3v} motif around the central silver atom (Figure 95a) [104]. In the structure of \{((3,5-Me_2C_6H_3)P)Ag[HCB_{11}Me_{11}]\} the steric bulk of the 3,5-methyl groups has switched off any significant extra interactions from carborane anions adjacent in the lattice, there being only three close intermolecular Ag...MeB contacts from a single carborane anion. These come from three methyl groups all on the lower surface of the cage. Two of these contacts are relatively short (Ag...C(7) 2.785(10) Å and Ag-C(12) 2.554(10) Å), while the other is longer (Ag-C(8) 3.166(10) Å), however, all are well within the sum of the van der Waals radius of the methyl group and the ionic radius of Ag\(^+\) (3.29 Å) (Figure 95b) [104].
The reaction of \((\text{Ph}_3\text{P})\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]\) with triphenylphosphine results in \([(\text{Ph}_3\text{P})_2\text{Ag}] [\text{HCB}_{11}\text{Me}_{11}]\) which also can be obtained by the reaction of \(\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]\) with an excess of \(\text{Ph}_3\text{P}\) (Figure 96) \cite{104}. The attempt to recrystallize \([(\text{Ph}_3\text{P})\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]]\) from diethyl ether resulted in \([(\text{Et}_2\text{O})_2\text{Ag}({\text{PPh}_3})][\text{HCB}_{11}\text{Me}_{11}]\) \cite{104}.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{image.png}
\caption{X-ray structure of \([(\text{Ph}_3\text{P})_2\text{Ag}[\text{HCB}_{11}\text{Me}_{11}]].\) Reprinted with permission from reference \cite{104}. Copyright © (2004) American Chemical Society.}
\end{figure}

It is known that the carba-c\textit{closo}-dodecaborate anion has slightly acidic CH group but its acidity increases significantly on halogenation of the cage. This open a way to synthesis of carborane derivatives with covalent bonds C-M (M = Ag, Cu). The deprotonation of the cage C-H proton of the \([\text{HCB}_{11}\text{I}_3\text{Br}_6]\) anion with KOH followed by treatment with AgNO\(_3\) and recrystallization from acetonitrile gave complex \([(\text{(MeCN})_4\text{Ag}_3)(\text{Ag}[\text{HCB}_{11}\text{I}_3\text{Br}_6]_2)_2\cdot 2\text{(MeCN)}]_n\). The C-Ag(1)-C fragment is linear and Ag(1)-C distance is 2.142(7) Å. The interesting structural feature is the strong bonding interactions between the second silver and the upper belt iodine atoms. This is a brand new coordination motif that has not been observed before in the silver salts of any carborane monoanions. Each six-coordinated silver Ag(3) links two cages together, leading to the formation of a zig-zag coordination polymeric chain. The Ag...IB and Ag...BrB distances vary from 2.953(2) to 3.008(1) Å and from 2.954(1) to 3.030(2) Å, respectively (Figure 97) \cite{107}.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{image.png}
\caption{Fragment of X-ray structure of \([(\text{(MeCN})_4\text{Ag}_3)(\text{Ag}[\text{HCB}_{11}\text{I}_3\text{Br}_6]_2)_2\cdot 2\text{(MeCN)}]_n.\) Reproduced with permission from reference \cite{107}. Copyright © (2003) John Wiley & Sons.}
\end{figure}

Reaction of \((\text{Bu}_4\text{N})[\text{HCB}_{11}\text{F}_{11}]\) with \([\text{Cu}(\text{mesityl})]_n\) and \(\text{Bu}_4\text{NCl}\) in dichloromethane resulted in formation of \((\text{Bu}_4\text{N})_2[1\text{-ClCu-1-CB}_{11}\text{F}_{11}]\). The C-Cu-Cl fragment in the \([1\text{-ClCu-1-CB}_{11}\text{F}_{11}]^-\) anion is near linear (176.0(2)°) and the Cu-C and Cu-Cl bonds are 1.917(5) and 2.136(1) Å, respectively (Figure 98) \cite{112}.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{image.png}
\caption{Figure 98. X-ray structure of \([1\text{-ClCu-1-CB}_{11}\text{F}_{11}]^-\). Reproduced with permission from reference \cite{112}. Copyright © (2003) John Wiley & Sons.}
\end{figure}
6. Silver and Copper Complexes with Cobalt bis(dicarbollide) anion [3,3′-Co(1,2-C2B9H11)2]−

Cobalt bis(dicarbollide) [3,3′-Co(1,2-C2B9H11)2]− was prepared for the first time fifty years ago [113,114] and it still remains the subject of the increasing interest [115] due to good perspectives of its application in various fields starting from medicine [116–118] and finishing with nuclear waste treatment [119], and weakly coordinating anions is one of them.

The silver salt of cobalt bis(dicarbollide) Ag[3,3′-Co(1,2-C2B9H11)2] was prepared by metathesis of the cesium salt or acid form (H3O)[3,3′-Co(1,2-C2B9H11)2] with AgNO3 in aqueous solution [73,120]. Crystals of [Ag(η2-p-xylene)][3,3′-Co(1,2-C2B9H11)2]n were grown by crystallization of Ag[3,3′-Co(1,2-C2B9H11)2] from p-xylene [116]. The silver atom is bound to one p-xylene molecule in an η2-fashion (Ag-C bonds are 2.562(7) and 2.411(7) Å) and three B-H bonds from the three dicarbollide cages through H(8), H(8) and H(9A) atoms (the Ag…B distances are very long—2.705(6), 2.926(6) and 2.938(6) Å, however all Ag…H distances fail in a range 2.01–2.08 Å) in a distorted tetrahedral geometry. The location of the three coordinating BH groups involves that portion of the bis(dicarbollide) anion which is believed to be the most electron-rich. The bis(dicarbollide) anion has cisoid conformation, that can be stabilized by B(8)H…Ag…HB(8’) interactions. The tridentate [3,3′-Co(1,2-C2B9H11)2]− anion and Ag+ cations serve as alternating bridging ligands to give a zigzag cation-anion chain (Figure 99) [121].

Figure 99. Fragment of X-ray structure of [Ag(η2-p-xylene)[3,3′-Co(1,2-C2B9H11)2]]n.
The silver salt of 8,8′,9,9′,12,12′-hexabromo cobalt bis(dicarbollide) \( \text{Ag}[8,8′,9,9′,12,12′-\text{Br}_6-3,3′-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_10)_2] \) was prepared by metathesis of the corresponding cesium salt with AgNO\(_3\) in aqueous solution [73]. Complex \([\text{(MeCN)}_4\text{Ag}][8,8′,9,9′,12,12′-\text{Br}_6-3,3′-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_8)_2]\) were prepared by crystallization of Ag[8,8′,9,9′,12,12′-Br\(_6\)-3,3′-Co(1,2-\text{C}_2\text{B}_9\text{H}_8)_2] from an acetonitrile/p-xylene solution. The crystal composes of discrete \([\text{(MeCN)}\text{Ag}]^+\) cations and metallacarborane anions (Figure 100) [121].

![Figure 100. X-ray structure of \([\text{(MeCN)}_4\text{Ag}][8,8′,9,9′,12,12′-\text{Br}_6-3,3′-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_8)_2]\).](image)

An unusual complex \([\text{Ag}(\eta^2-p\text{-xylene})][8,8′-\text{μ-}\text{C}_6\text{H}_4-3,3′-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]\) was prepared by metathesis of the corresponding cesium salt with AgNO\(_3\) in aqueous solution followed by crystallization from p-xylene [121]. It is interesting to note that no interaction between the Ag\(^+\) and any B-H bonds from the cage is observed. The Ag atom is \(\eta^2\)-bound to each of two aromatic rings in an approximately linear geometry. The average distances from the Ag atom to two aromatic rings are about the same (2.45(2) and 2.42(2) Å), perhaps indicating that the \(\pi\)-donor ability of the p-xylene molecule is no different than that of the bridging aromatic ring in the \([8,8′-\text{μ-}\text{C}_6\text{H}_4-3,3′-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]^−\) anion. The two aromatic rings are not parallel to each other and they are separated by about 4.1 Å. There are two crystallographically independent molecules in the unit cell, which are associated to each other through a \(\pi\)-\(\pi\) interaction between the bridging aromatic ring and the p-xylene of the second complex (Figure 101) [121].

![Figure 101. X-ray structure of \([\text{Ag}(\eta^2-p\text{-xylene})][8,8′-\text{μ-}\text{C}_6\text{H}_4-3,3′-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]\).](image)

Complex \(\{[(\text{MeCN})_2\text{Ag}][3,3′-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\}_n\) was obtained by crystallization of Ag[3,3′-Co(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2] from an acetonitrile/2,2,2-trifluoroethanol solution. The silver cation is coordinated by two acetonitrile molecules arranged in a nearly linear fashion with a N-Ag-N angle of 167.26(16)\(^°\). The coordination sphere of silver is completed by weak interactions to B-H groups of
the \([3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]^-\) anion. Two BH groups from one metallacarborane anion, and one from a symmetry related another anion interact with the Ag\(^+\) at Ag…B distances ranging from 2.35 to 2.56 Å. The rather long distances indicate that these interactions are weak. The geometry around the Ag\(^+\) is distorted trigonal bipyramidal, with the three coordinating H atoms forming the equatorial plane. The bis(dicarbollide) anions have \textit{cisoid} conformation. Each \([3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]^-\) anion coordinates to two Ag\(^+\) cations through three BH groups, and a stepped cation-anion coordinate chain is formed (Figure 102) [122].

![Figure 102. X-ray structure of \((\text{MeCN})_2\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\)_n: coordination arrangement of silver cation (a) and crystal packing (b). Reprinted with permission from reference [122]. Copyright © (2003) American Chemical Society.](image)

Single crystals of \(((\text{NC}(\text{CH}_2)_3\text{CN})\text{Ag})[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]_n\) were grown by crystallization of Ag\([3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\) from ethanol in the presence of glutaronitrile (1:1). The silver cation has two glutaronitrile ligands at Ag-N distances 2.160(3) and 2.171(4) Å and two close agostic Ag…H interactions at 2.036 and 2.118 Å. The corresponding B…Ag distances are not close however at 2.90 and 3.05 Å. The glutaronitrile ligands deviate from linearity with an N-Ag-N angle of 153.32(13)°. The Ag…BH interactions come from a single \(\eta^2\)-coordinated cobalt bis(dicarbollide) anion and run roughly perpendicular to the N-Ag-N axis, to form a silver coordination environment that can be regarded as a distorted T-shape. The glutaronitrile ligand bridges between two symmetry-equivalent silver cations, and a single-stranded coordination chain is propagated (Figure 103) [105].

![Figure 103. X-ray structure of \((\text{NC}(\text{CH}_2)_3\text{CN})\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]_n\).](image)

Single crystals of \([\text{Ag}(\text{NC}(\text{CH}_2)_3\text{CN})_2][3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]_n\) were grown by crystallization of Ag\([3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\) from ethanol in the presence of glutaronitrile (1:2) [105]. The silver cations are coordinated by glutaronitrile ligands with distorted tetrahedral geometries. Each glutaronitrile ligand bridges between two silver cations to give a \([\text{Ag}(\text{NC}(\text{CH}_2)_3\text{CN})_2]^+\) coordination chain with a criss-crossed ladder topology. The ladder is of unusual topology for a 4-connected polymer, and the
Ag...Ag separations within the ladder are 6.95 Å along the sides and 4.52 Å along the zigzagging struts. The \([\text{Ag(NC(CH}_2)_3\text{CN})_2]^+\) coordination chains do not show any close interactions with the \([3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]^\text{−}\) anions, and the crystal packing shows isolated coordination chains and double columns of cobalt bis(dicarbollide) anions (Figure 104) [105].

**Figure 104.** Crystal packing in structure of \([\text{[Ag(NC(CH}_2)_3\text{CN})_2][3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]_n\). Reprinted with permission from reference [105]. Copyright © (2004) American Chemical Society.

Single crystals of \([(\text{NC(CH}_2)_4\text{CN})\text{Ag})[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]_n\) were grown by crystallization of \(\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\) from ethanol in the presence of adiponitrile (1:1) [105]. Likewise complex \([(\text{NC(CH}_2)_3\text{CN})\text{Ag})[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]_n\), each silver cation is coordinated by two adiponitrile ligands with a distorted linear geometry. The Ag-N distances range from 2.151(4) to 2.174(3) Å, and N-Ag-N angles are 148.60(16) and 157.33(14)°. There is one close Ag...H interaction to Ag(1) at a distance 2.160 Å and a further long interaction at Ag...H distance 2.49 Å. The geometry around Ag(1) is best described as distorted trigonal pyramidal. There are three long Ag...H interactions to Ag(2) at distances ranging from 2.31 to 2.57 Å, with overall distorted trigonal bipyramidal geometry (Figure 105) [105].

**Figure 105.** Fragment of X-ray structure of \([(\text{NC(CH}_2)_4\text{CN})\text{Ag})[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]_n\). Reprinted with permission from reference [105]. Copyright © (2004) American Chemical Society.

Complex \([(\text{NC(CH}_2)_3\text{CN})\text{Ag})[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2][\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]_n\) was prepared by crystallization of \(\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\) from acetone/trifluoroethanol in the presence of...
The two crystallographic independent silver cations Ag(1) and Ag(2a) exhibit a very distorted tetrahedral coordination. Ag(1) connects two N-atoms belonging to two crystallographic equivalent pimelonitrile ligands (Ag(1)-N(1) and Ag(1)-N(2i)) distances of 2.228(4) Å, and 2.208(4) Å, respectively, N(1)-Ag(1)-N(2i) angle of 124.530(2)° and further interacts with two BH groups of an adjacent cobalt bis(dicarbollide) anion at Ag . . . H distances of 2.012(1) Å (Ag(1) . . . H(18)-B(8)) and 2.073(1) Å (Ag(1) . . . H(28)-B(17)) to complete the distorted tetrahedral coordination environment. The short Ag(1) . . . H(18) and Ag(1) . . . H(28) distances indicate that these interactions exhibit strong character. Ag(2) interacts with four BH groups belonging to two neighbouring crystallographic equivalent [3,3'-Co(1,2-C2B9H11)2]− anions: Ag(2) . . . H(35a”) and Ag(2) . . . H(43”) at short Ag . . . H distances of 2.097(1) and 2.032(1) Å, respectively, and two weaker interactions for Ag(2) . . . H(45a) (2.247(1) Å) and Ag(2) . . . H(54a) (2.192(1) Å) (Figure 106) [123].

**Figure 106.** X-ray structure of {[(NC(CH2)6-CN)Ag][3,3'-Co(1,2-C2B9H11)2][Ag[3,3'-Co(1,2-C2B9H11)2]]}n; coordination arrangement of silver cation (a) and crystal packing (b). Reproduced from reference [123] with permission from The Royal Society of Chemistry.

Complex {Ag3(NC(CH2)6-CN)2[3,3'-Co(1,2-C2B9H11)2]}n was prepared by crystallization of Ag[3,3'-Co(1,2-C2B9H11)2] from ethanol in the presence of suberonitrile [123]. The structural unit is composed of three Ag⁺ cations, two bridging ligands, and three cobalt bis(dicarbollide) anions. Two crystallographic independent atoms, Ag(1) and Ag(2), reveal identical coordination environment, whereas the Ag(3) is distinct. Both Ag(1) and Ag(2) are four coordinated centers, bounded by two N-atoms of two crystallographic independent suberonitrile ligands and also two BH groups belonging to two distinct [3,3'-Co(1,2-C2B9H11)2]− anions, and mutually exhibiting distorted tetrahedral geometry. In Ag(1) center the Ag-N bond lengths are 2.228(4) Å (Ag(1)-N(1)) and 2.208(4) Å (Ag(1)-N(4iii)) and Ag . . . H distances are found to be 1.9817(7) Å (Ag(1) . . . H(52iv)) and 1.9728 Å (Ag(1) . . . H(81)), suggesting the existence of two strong interactions. The Ag-N distances in the Ag(2) center are analogous to those observed for Ag(1) at 2.245(4) Å (Ag(2)-N(3)) and 2.198(4) Å (Ag(2)-N(2)). However, while the two Ag . . . H-B interactions involving Ag(1) are similar, those of the Ag(2) center are significantly distinct: Ag(2) . . . H(63) is a stronger interaction and Ag(2) . . . H(71v) has a considerable weaker nature, as evidently indicated by Ag . . . H distances of 1.877(1) and 2.314(1) Å, respectively. The Ag(3) coordination environment consists exclusively of six Ag-H-B interactions with the BH groups belonging to three crystallographic independent cobalt bis(dicarbollide) anions, leading to a center with highly distorted octahedral geometry (Figure 107) [123].
Complex \( [(Ag(pyrazine)(MeCN))_2][3,3'-Co(1,2-C_2B_9H_11)_2]_n \) was prepared by crystallization of \( Ag[3,3'-Co(1,2-C_2B_9H_11)_2] \) with pyrazine in acetonitrile [106]. The silver cation is in a distorted tetrahedral geometry, coordinating to two acetonitrile molecules and to two pyrazine ligands. (Ag-N bonds range from 2.266(2) to 2.471(3) Å). Each pyrazine ligand is bridging, and a coordination chain is formed which runs along the \( c \)-axis. The chains occur in pairs with a second chain running parallel but offset with the first, so that silver cations of one chain lies over the center of the pyrazine ring of the other one with an Ag...centroid distance of 3.732 Å. The coordinated acetonitrile molecules are directed away from the chain and lie parallel to acetonitrile molecules from two adjacent chains. The orientation of the acetonitrile molecules of adjacent coordination chains creates a “box-like” arrangement and the cobalt bis(dicarbollide) anions form rows and lie in the channels created by this packing arrangement. Each \( [3,3'-Co(1,2-C_2B_9H_11)_2]^- \) anion is in close proximity to neighboring anions with a closest contact of 2.271 Å between the BH...HC groups, indicating the presence of weak dihydrogen interactions between the metallacarborane anions (Figure 108) [106].

![Figure 107. X-ray structure of \( [Ag_3(NC(CH_2)_6CN)_2][3,3'-Co(1,2-C_2B_9H_11)_2]_n \): coordination arrangement of silver cation (a) and crystal packing (b). Reproduced from reference [123] with permission from The Royal Society of Chemistry.](image)
Two polymorphs of complex \([(\text{Ag}(4,4'-\text{Bipy})(\text{MeCN}))][3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\)\(_n\) was prepared by crystallization of \(\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\) with 4,4'-bipyridine in acetonitrile [106]. The silver coordination environment is essentially the same for both polymorphic forms, with silver cation coordinated by an acetonitrile and by two 4,4'-bipyridine molecules in a T-shape arrangement. In both polymorphs, the silver atom is in close proximity to a neighboring cobalt bis(dicarbollide) anion with Ag...H distances of 2.733 and 2.696 Å, too long to be considered effective interactions. The bridging 4,4'-bipyridine ligands form coordination chains that occur in infinite π-stacked columns. Each chain shows the coordinated acetonitrile molecules alternating in opposite directions. A box-like channels are generated that contains metallacarborane anions (Figure 109) [106].

The crystallization of \(\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\) from dimethylsulfoxide in the presence of 4,4'-bipyridine gave complex \([(\text{Ag}(4,4'-\text{Bipy})(\text{DMSO})_2)][3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2].\text{DMSO}\)\(_n\) [106]. The silver cation is coordinated by two 4,4'-bipyridine molecules (Ag-N bond length 2.187(8) and 2.161(8) Å) in an almost linear arrangement (N-Ag-N angle 165.4(3)°). The four-coordinated silver sphere is completed by two DMSO molecules. The \([\text{Ag}(4,4'-\text{Bipy})(\text{DMSO})_2]\)\(^{+}\) cationic fragments form coordination chains. The chains are arranged in pairs with the chain offset so that the silver cations are opposite the center of one of the arene rings of a ligand from the neighboring chain, at Ag... centroid distance of 3.874 Å. There are π...π staking interactions between the chais at a centroid separation of 3.576 Å. The extended structure reveals a herring-bone arrangement of the cobalt bis(dicarbollide) anions in the bc-plane (Figure 110).
Figure 109. X-ray structure of \( \{\text{Ag}(4,4'-\text{Bipy})(\text{MeCN})\}[-3,3'-\text{Co}(1,2-\text{C}_2\text{B}_{9}\text{H}_{11})_2]_n \): coordination arrangement of silver cation (a) and crystal packing (b).

Complex \( \{\text{Ag}_2(bppz)_3\}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_{9}\text{H}_{11})_2]_2 \cdot 3.5(\text{MeCN})_n \) was prepared by crystallization of \( \text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_{9}\text{H}_{11})_2] \) from acetonitrile in the presence of 2,3-bis(2-pyridyl)pyrazine [106]. The crystals contain the \( \{\text{Ag}_2(bppz)_3\}^{2+} \) coordination chain which is similar to those found in the related complexes with the carba-closo-dodecaborate anion (See Figure 89). In the extended structure, the chains are separated by snaking chains of cobalt bis(dicarbollide) anions. The anionic chains have anions in one of two orientations and show dihydrogen bonding with closest BH…HC distances of 1.947 and 2.312 Å. Both the chains of polymeric \( \{\text{Ag}_2(bppz)_3\}^{2+} \) cations and the cobaltacarborane anions run in a parallel fashion along the c-axis forming a square grid or checkerboard pattern (Figure 111).

Figure 110. Fragment of the X-ray structure of \( \{\text{Ag}(4,4'-\text{Bipy})(\text{DMSO})_2\}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_{9}\text{H}_{11})_2]_2 \cdot \text{DMSO}_n \).
Complex $\{\text{Ag(DABCO)}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\}_n$ was prepared by crystallization of $\text{Ag}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]$ from acetonitrile in the presence of 1,4-diazabicyclo[2.2.2]octane [77]. The silver atom coordinates to two nitrogen atoms belonging at two crystallographically equivalent DABCO ligands (Ag-N distances 2.1752(17) and 2.180(17) Å) and further interacts with three BH groups belonging at three equivalent cobalt bis(dicarbollide) anions (Ag…H distances 2.8889(2), 2.4465(2), and 2.8958(1) Å). With consideration of the Ag-N coordination bonds and all the Ag-H-B interactions, the coordination geometry around silver atom can be visualized as intermediate between a deformed trigonal bipyramid and a distorted square pyramid. The bridging nature of the DABCO ligand between the Ag centers lead to the formation of cationic linear polymer chains $\{\text{Ag(DABCO)}\}^+$, which run along the $a$ axis with the Ag…Ag intrachain distance of 6.9544(5) Å. These cationic coordination chains are intercalated by the metallacarborane anions, ultimately generating a zigzag arrangement of both the coordination chains and carborane anions (Figure 112) [77].

![Figure 111](https://example.com/figure111.png)

**Figure 111.** Crystal packing in X-ray structure of $\{\text{Ag}_2(\text{bppz})_3\}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2\cdot 3.5(\text{MeCN})]\}_n$.

![Figure 112](https://example.com/figure112.png)

**Figure 112.** X-ray structure of $\{\text{Ag(DABCO)}[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]\}_n$: coordination arrangement of silver cation (a) and crystal packing (b). Reprinted with permission from reference [77]. Copyright © (2013) American Chemical Society.
The supramolecular complexes \([\text{Ag(MeCN)}(\text{H}_2\text{O})(\text{CTV})_2][3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\) and \([\text{Ag(MeCN)}_3(\text{CTV})][3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\) were prepared by co-crystallization of \(\text{Ag}[3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\) and cyclotriveratrylene (CVT) from a acetonitrile/2,2,2-trifluoroethanol solution depending on the solvent ratio [122]. In the structure of \([\text{Ag(MeCN)}(\text{H}_2\text{O})(\text{CTV})_2][3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\) the silver cation is located within the CTV molecular cavity. The coordination sphere of silver atom is four-coordinate with distorted tetrahedral geometry. There is one aquo ligand at \(\text{Ag-O distance 2.369(15)} \text{ Å}\), an acetonitrile molecule at \(\text{Ag-N 2.038(13)} \text{ Å}\) and \(\text{Ag-C bonds to the C6 position of one veratrole ring within the CTV (Ag-C distance 2.6077(7) Å)}\) to and the symmetry equivalent C3 position of an adjacent ring. The aquo ligand is exo to the CTV cavity, while the acetonitrile ligand is inside the CTV cavity and points along the plane of the arene ring not involved in coordination to silver. The bis(CTV) capsule actually comprises a \([\text{Ag(H}_2\text{O)(MeCN)}(\text{CTV})]^+\) complex with a second molecule of CTV in an inverted orientation, associated with the first via host-guest interactions with the coordinated acetonitrile (Figure 113a). The capsules stack in the \(c\) direction, possibly through a weak hydrogen bond between the coordinated aquo ligand and dimethoxy group of neighboring capsules (O . . . O separation 3.43 Å). Capsules that are adjacent in the \(bc\) plane associate through \(\pi\)-\(\pi\) interactions between arene rings of the CTV molecules where the centroid separation is 3.696 Å. The cobalt bis(dicarbolide) anions are located in hydrophobic pockets and show no significant interactions with neighboring molecules (Figure 113b) [122].

![Figure 113](image)

**Figure 113.** X-ray structure of \([\text{Ag(MeCN)}(\text{H}_2\text{O})(\text{CTV})_2][3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\): coordination arrangement of silver cation (a) and crystal packing (b). Reprinted with permission from reference [122]. Copyright © (2003) American Chemical Society.

In the structure of \([\text{Ag(MeCN)}_3(\text{CTV})][3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\) 0.5MeCN the silver cation has an unusual coordination sphere, comprising three acetonitrile ligands and two methoxy O atoms from the CTV. The Ag-N bond lengths are 2.184(2), 2.206(2), and 2.466(2) Å, and the MeCN ligands are arranged in a distorted T-shape. The coordination sphere of silver is completed by coordination to both O atoms of one of the veratrole moieties within the CTV at long Ag-O distances 2.685(2) and 2.834(2) Å to form a five-membered chelate ring. The coordination geometry around the silver atom may be described as highly distorted trigonal bipyramidal with MeCN ligands in the axial positions, and the dimethoxy unit and other MeCN comprising the equatorial plane. One of the coordinated acetonitrile molecules occupies the CTV molecular cavity to form a host-guest interaction. The guest MeCN is parallel to the coordinating veratrole ring of the CTV. In the crystal layers of the \([\text{Ag(MeCN)}_3\text{CTV}]^+\) cations separated by layers of the \([3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\) anions. The \([\text{Ag(MeCN)}_3\text{CTV}]^+\) cations form pairs with the roughly linear MeCN-Ag-NMe links nearly parallel to each other and with the CTV hosts in opposite orientations. Each \([3,3'\text{-Co(1,2-C}_2\text{B}_9\text{H}_{11})_2]\) anion is at hydrogen bonding distances to one CTV arene ring at C-H . . . \(\pi\) distance 2.78 Å, but do not show other close contacts to other molecular components (Figure 114) [122].
The [Ag(MeCN)$_4$](CTV)$[3,3'$-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$]·0.25MeCN were prepared by co-crystallization of Ag[3,3'-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$] and cyclotriveratrylene from acetonitrile [122]. The silver atom forms tetrahedral [Ag(MeCN)$_4$]$^+$ complex ion, which is a guest molecule with one coordinated MeCN ligand forming a host-guest interaction with the CTV. The guest MeCN extends into the CTV molecular cavity and approaches over one of the -CH$_2$- links of the CTV. The cobalt bis(dicarbollide) anion forms close interactions with both the [Ag(MeCN)$_4$]$^+$ cation and CTV molecule. There are three C-H...O hydrogen bonds between the [3,3'-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$]$^-$ anion and the methoxy groups of a single veratrole moiety of CTV (2.52, 2.59, 2.79 Å). The fourth C-H group of the [3,3'-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$]$^-$ anion is in close contact with a N atom of a coordinated MeCN at C-H...N distance 2.51 Å, leading to the dimeric $[[[Ag(MeCN)$_4$]$^+$∩CTV]$[3,3'-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$]]_2$ assembly (Figure 115) [122].

Figure 114. X-ray structure of [Ag(MeCN)$_3$(CTV)]$[3,3'-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$]$·0.5$MeCN.

Figure 115. X-ray structure of $[[[Ag(MeCN)$_4$]$^+$∩CTV]$[3,3'-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$]]_2$. Reprinted with permission from reference [122]. Copyright © (2003) American Chemical Society.

Another supramolecular host-guest complex [Ag(tris(isonic)(CTG)$_2$]$[3,3'$-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$]$·9$(MeCN) was prepared by co-crystallization of Ag[3,3'-Co(1,2-C$_2$B$_9$H$_{11}$)$_2$] with tris-(isonicotinoyl) cyclotriguaiacylene (tris(isonic)(CTG)) from acetonitrile. The silver atom has a tetrahedral geometry
(Ag-N bond lengths 2.233(7)–2.448(7) Å) and is bound by four different ligand molecules. Each molecule of L acts as a bidentate bridge of two symmetry-related silver atoms to form a 1-D \{Ag(tris(isonic)CTG)\}_{2+}^n coordination polymer. The 1-D coordination polymers are interwoven through dimeric host-guest associations between the coordinated molecular host moieties of adjacent chains and π...π stacking interactions to give an extended 2-D network (Figure 116). The cobalt bis(dicarbollide) anions are within channels created by packing the interwoven 2-D sheets [124].

![Figure 116. Structure of cationic lattice in \([Ag(tris(isonic)CTG)]_2[3,3'-Co(1,2-C_2B_9H_{11})_2]·9(MeCN)\). Reproduced with permission from reference [124]. Copyright © (2004) American Chemical Society.](image)

Complex \([Ag_2(tris(3-pyamino)CTG)_2(MeCN)_2][3,3'-Co(1,2-C_2B_9H_{11})_2]^2·2(MeCN)\) was prepared by co-crystallization of \([Ag][3,3-Co(1,2-C_2B_9H_{11})_2]·9(MeCN)\) from a acetonitrile/acetone solution [108]. The structure is formed by the \([Ag_2(tris(3-pyamino)CTG)_2(MeCN)_2]^+\) cations, cobalt bis(dicarbollide) anions and MeCN solvent molecules. The cation \([Ag_2(tris(3-pyamino)CTG)_2(MeCN)_2]^2+\) is a dinuclear, dimeric species with a capsule-like structure. The centrosymmetric capsule forms through the head-to-head coordination of two ligands to two silver atoms. The silver cation is coordinated by a terminal acetonitrile ligand at an Ag-N distance of 2.321 Å, and three pyridyl groups at Ag-N distances of 2.327, 2.350, and 2.292 Å in a distorted tetrahedral geometry (Figure 117) [108].

![Figure 117. Structure of \([Ag_2(tris(3-pyamino)CTG)_2(MeCN)_2]^+\) cation. Reproduced with permission from reference [108]. Copyright © (2006) John Wiley & Sons.](image)

Polymeric complex \({\{[Ag(tris(4-pyamino)CTG)]NC(CH_2)_3CN]\cdot(NC(CH_2)_3CN)·[3,3'-Co(1,2-C_2B_9H_{11})_2]\cdot n(NC(CH_2)_3CN)\}_n\) was prepared by co-crystallization of \([Ag][3,3-Co(1,2-C_2B_9H_{11})_2]\)
with tris(4-pyridylmethylamino)cyclotriguaiacylene (tris(4-pyamino)CTG) from acetonitrile-2,2,2-
trifluoroethanol in the presence of glutaronitrile [108]. The silver cation has a tetrahedral geometry
coordinated by three pyridyl moieties from three different molecules of tris(4-pyamino)CTG (Ag-N
bond lengths of 2.343(6)–2.369(6) Å) and a monodentate capping glutaronitrile molecule (Ag-N bond
length of 2.25(1) Å). The three cavitands coordinating to each silver cation are orientated such that two
have the top face orientated in the same direction with the third directed in an opposing direction. The
cavitand is likewise coordinated to three different symmetry-related silver cations and acts as a host
for the second molecule of glutaronitrile. This results in 2D-network structure incorporating roughly
rectangular channels. These channels are largely occupied by the globular metallacarbaborane anions
and glutaronitrile molecules (Figure 118) [108].

Complex \([\text{Ag}[\text{tris}(\text{isonicotinoyl})\text{-tris(propoxy)}]\text{-cyclotriguaiacylene}}\) (tris(isonic)CTG-Pr) from solution in N,N-dimethylformamide [125]. In the crystal, the
Ag-N and Ag...H(B) bond lengths were measured at 2.264(4), 2.280(4) and 2.176 Å, respectively, alongside N-Ag-N bond angles of 127.48(14), 101.76(14) and
97.97(13°). It results in a 1D ladder-type motif, whereby the tris(isonic)CTG-Pr ligands are all inwardly
orientated to afford a quasi-cylindrical arrangement of head-to-head ligands (Figure 119) [125].

Complex \([\text{Ag}_2(\text{tris}(\text{2-quinolylmethyl})\text{-tris(propoxy)}]\text{-cyclotriguaiacylene}}\) (tris(2-quinolylmethyl)tris(propoxy)-
cyclotriguaiacylene (tris(isoquinolyl)CTG-Pr) from a solution in acetonitrile/nitromethane [125].
In the crystal each silver cation is in approximately linear arrangement and is coordinated by two
independent tris(2-quinolynoic)CTG-Pr ligands, with Ag-N bond lengths and N-Ag-N angles of
2.189(7)–2.197(7) Å and 174.3(3) and 176.2(3)°, respectively, that results in 1D coordination polymer.
There is some bonding contribution from a proximal ethereal oxygen, with O...Ag distance of
2.513(7) Å. The host-guest interactions between quinolyl arm and the shallow hydrophobic
cavity of the ligand core are stabilised by π...π interactions between the quinolyl moiety and the
[a,d,g]cyclononatriene core, with centroid separation of 3.809 Å. Neighbouring 1D chains aggregate
through π...π interactions, with aromatic centroid separations of 3.687 Å, and aggregation of propyl

Figure 118. The crystal packing in \([(\text{Ag}[\text{tris}(4-pyamino)CTG])\text{[NC(CH}_2)_3\text{CN}]\cap(NC(CH}_2)_3\text{CN})
[3,3'-\text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2] \cdot n(NC(CH}_2)_3\text{CN})]_n\). Reproduced with permission from reference [108].
moieties which affords small pockets that are filled with disordered solvent and cobalt bis(dicarbollide) anions (Figure 120) [125].

![Figure 119](image1.png)

**Figure 119.** Fragment of X-ray crystal structure (a) and crystal packing (b) in {[Ag(tris(isonic)CTG-Pr)[3,3'-Co(1,2-C2B9H11)2]]·2.5(DMF)·H2O)n. Reproduced from reference [125] with permission from The Royal Society of Chemistry.

![Figure 120](image2.png)

**Figure 120.** Structure of cationic lattice in [Ag2(tris(3-pyamino)CTG)2(MeCN)2]+ cation in {[Ag2(tris(2-quinamino)CTG-Pr)2][3,3'-Co(1,2-C2B9H11)2]2·1.5(MeNO2)}n. Reproduced from reference [125] with permission from The Royal Society of Chemistry.
Syntheses of two silver(I) complexes with cobalt bis(dicarbollide) based on the chelating phosphine ligand \([1,1'-(\text{Ph}_3\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]^-\) were described [126]. The complex \([(\text{Ph}_3\text{P})\text{Ag}[1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]]^-\) acetonitrile was prepared by the reaction of \((\text{Me}_4\text{N})[1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]^-\) with \([(\text{Ph}_3\text{P})\text{AgCl}]\) in ethanol, whereas the complex \([(\text{Me}_4\text{CO})\text{Ag}[1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]^-\) was prepared by the reaction of \((\text{Me}_4\text{N})[1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]^-\) with \(\text{AgClO}_4\) in ethanol/acetone. The silver cation in both complexes is tetrahedrally coordinated, with one long and three short contacts. In the first complex two of the short contacts are due to the two P atoms from the \([1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]^-\) ligand (Ag–P 2.5507(9) and 2.5470(10) Å), and the third is due to PPh\(_3\) (Ag–P 2.4713(10) Å); the fourth bond is due to the O atom of acetone molecule (Ag … O 2.906(3) Å). In the second complex two short contacts are due to the P atoms of the \([1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]^-\) ligand (Ag–P 2.4894(6) and 2.4933(6) Å), and the third is due to the O atom of acetone molecule (Ag–O 2.3895(17) Å), whereas the bond is due to Ag … H(12′) contact (2.17 Å), assembling the molecules to 1D-polymeric chain (Figure 121) [126].

![Figure 121. X-ray structures of \([(\text{Ph}_3\text{P})\text{Ag}[1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]]^-\) acetonitrile (a) and \([(\text{Me}_4\text{CO})\text{Ag}[1,1'-(\text{Ph}_2\text{P})_2\cdot 3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{10})_2]]^-\) (b). Reproduced with permission from reference [126]. Copyright © (2004) John Wiley & Sons.](image)

Unlike the silver complexes, copper complexes with cobalt bis(dicarbollide) anion are represented by only a few examples. A mixture of complexes \([(\text{Ph}_3\text{P})\text{Cu}[3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]\) and \([(\text{Ph}_3\text{P})_2\text{Cu}[3,3'\cdot \text{Co}(1,2-\text{C}_2\text{B}_9\text{H}_{11})_2]]\) was prepared by the reaction of Cs[3,3'–Co(1,2–C\(_2\)B\(_9\)H\(_{11}\))\(_2\)] with CuCl\(_2\) and triphenylphosphine or, alternatively, with complex \([(\text{Ph}_3\text{P})\text{CuCl}]_4\) in dichloromethane, whereas the reaction of Cs[3,3'–Co(1,2–C\(_2\)B\(_9\)H\(_{11}\))\(_2\)] with complex \([(\text{Ph}_3\text{P})\text{CuCl}]\) gives only the second complex \([(\text{Ph}_3\text{P})_2\text{Cu}[3,3'–\text{Co}(1,2–\text{C}_2\text{B}_9\text{H}_{11})_2]]\). The last one can be obtained by the treatment of the first complex \([(\text{Ph}_3\text{P})\text{Cu}[3,3'–\text{Co}(1,2–\text{C}_2\text{B}_9\text{H}_{11})_2]]\) with excess of PPh\(_3\). The copper(I) cation in the complex \([(\text{Ph}_3\text{P})\text{Cu}[3,3'–\text{Co}(1,2–\text{C}_2\text{B}_9\text{H}_{11})_2]]\) is in distorted tetrahedral arrangement formed by the P atom of triphenylphosphine ligand (Cu–P bond is 2.2264(2) Å) and three B(8)H, B(8′)H and B(4′)H groups of the cobalt bis(dicarbollide) anion (Cu … B and Cu … H distances are 2.2265(9)–2.3243(9) and 1.721(14)–2.014(14) Å, respectively) (Figure 122) [127,128].
with the hydride atoms in the case of bulky cobalt bis(dicarbollide) monoanion. The coordination takes place at the most electron-rich positions of boron polyhedra. Geometrical features of the structure of which contains two four-coordinated boron atoms, to very weak M…H interactions with the hydride atoms in the case of bulky cobalt bis(dicarbollide) monoanion. The coordination takes place at the most electron-rich positions of boron polyhedra.

Complexes of silver(I) and copper(I) with anionic polyhedral boron hydrides demonstrate a wide variety of structural types, relating to both the metal coordination environment and coordination modes of boron hydride anions. The latter can range from very strong coordination via the polyhedron triangular face including formation of 3c-2e MHB bonds in the case of the closo-dodecaborate dianion, the structure of which contains two four-coordinated boron atoms, to very weak M…H interactions with the hydride atoms in the case of bulky cobalt bis(dicarbollide) monoanion. The coordination takes place at the most electron-rich positions of boron polyhedra. Geometrical features of the closo-decaborate anion allow one to control positions of metal atoms around the boron polyhedron resulting in preparation of positional isomers \([\{L_2M\}_2[B_{10}H_{10}]\] varying reaction conditions. In general, 

The copper(II) complex \([Cu(Phen)_3][3,3'-Co(1,2-C_2B_9H_{11})_2]\) was prepared by the reaction of \([Cu(Phen)_3](OAc)_2\) with Cs[3,3'-Co(1,2-C_2B_9H_{11})_2] in aqueous solution (Figure 123) [129,130].

**Figure 122.** X-ray structures of \([\{(Ph_3P)Cu\}[3,3'-Co(1,2-C_2B_9H_{11})_2]\) (a) and \([\{(Ph_3P)\_2Cu\}[3,3'-Co(1,2-C_2B_9H_{11})_2]\) (b). Reproduced with permission from reference [128]. Copyright © (2015) John Wiley & Sons, Inc.

**Figure 123.** X-ray-structure of \([Cu(Phen)_3][3,3'-Co(1,2-C_2B_9H_{11})_2]\) in aqueous solution (Figure 123) [129,130].

7. Conclusions

Complexes of silver(I) and copper(I) with anionic polyhedral boron hydrides demonstrate a wide variety of structural types, relating to both the metal coordination environment and coordination modes of boron hydride anions. The latter can range from very strong coordination via the polyhedron triangular face including formation of 3c-2e MHB bonds in the case of the closo-decaborate dianion, the structure of which contains two four-coordinated boron atoms, to very weak M…H interactions with the hydride atoms in the case of bulky cobalt bis(dicarbollide) monoanion. The coordination takes place at the most electron-rich positions of boron polyhedra. Geometrical features of the closo-decaborate anion allow one to control positions of metal atoms around the boron polyhedron resulting in preparation of positional isomers \([\{L_2M\}_2[B_{10}H_{10}]\] varying reaction conditions. In general,
the coordination ability of the studied polyhedral boron hydride anions decreases in the series closo-decaborate \( [\text{B}_{10}H_{10}]^2^- \) > closo-dodecaborate \( [\text{B}_{12}H_{12}]^2^- \) > 1-carba-closo-decaborate \( [1\text{-C}\text{B}_{9}H_{10}]^- \) > carba-closo-dodecaborate \( [\text{C}\text{B}_{11}H_{12}]^- \) > cobalt bis(dicarbollide) \( [3,3'\text{-Co}(1,2\text{-C_{2}B_{9}H_{11}})]^- \). In the case of the closo-decaborate and closo-dodecaborate anions, the coordination ability can be decreased by an introduction of charge-compensating substituents which reduce the anion charge. The substitution of hydrogen atoms for halogens reduces the coordination ability of the polyhedral boron hydrides, which decreases in the halogen order I > Br > Cl > F, with no M...FB interactions being found.

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