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Redox-Active Carborane Clusters in Bond Activation Chemistry and Ligand Design

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Icosahedral *closo*-dodecaboranes have the ability to accept two electrons, opening into a dianionic *nido*-cluster. This transformation can be utilized to store electrons, drive bond activation, or alter coordination to metal cations. In this feature article, we present cases for each of these applications, wherein the redox activity of carborane facilitates the generation of unique products. We highlight the effects of exohedral substituents on reactivity and the stability of the products through conjugation between the cluster and exohedral substituents. Futher, the utilization of the redox properties and geometry of carborane clusters in the ligand design is detailed, both in the stabilization of low-valent complexes and in the tuning of ligand geometry.

Introduction

The activation of strong bonds has been the motivator of organometallic chemistry since the early development of transition metal catalysis. This reactivity largely stems from the combination of energetic and spatial features of filled and empty *d*-orbitals at a metal center, which influences its coordination environment and often leads to the ability to readily undergo two-electron cycling between oxidation states.¹

Currently, significant attention is focused on the conceptually similar metal-free redox bond activation transformations inspired by recent discoveries.^{2–19} It has been demonstrated that the electronic structure (i.e., energy levels of frontier orbitals) of main group compounds can be significantly altered by changes in the substitution patterns or by the introduction of geometric constraints. As one prominent example, cyclic (alkyl)(amino)carbenes (CAACs), which possess both higher-energy HOMOs and lower-energy LUMOs than Nheterocyclic carbenes (NHCs), resulting in significantly greater σ -donation and π -acceptance, participate in the activation of strong H-H, B-H, Si-H, and N-H bonds through oxidative addition at the carbene center.^{3,20,21} As another example, cycling between P(III) and P(V) states in geometrically constrained phosphorus compounds affords the possibility of oxidative addition and reductive elimination in a way that mimics the reactivity of transition metal complexes.^{6,22–24}

The redox transformations in main group systems can be centered at a single atom (carbon or phosphorus atoms in the examples above), or they can occur through the participation of larger molecular fragments. The focus of this feature article is on the utilization of the redox activity of icosahedral *closo*- dicarbadodecaboranes ($C_2B_{10}H_{12}$, further referred to simply as "carboranes"), which are remarkably robust neutral molecular clusters with ten boron and two carbon vertices.^{25,26} The electron delocalization within the cluster, which is related to the delocalized nature of planar aromatic rings, contributes to its high chemical stability.^{27–29} These clusters have attracted increasing attention due to new applications in luminescent materials, batteries, medicine, ligand design, and catalysis.^{30–41}

Neutral icosahedral $C_2B_{10}H_{12}$ cages are among the most studied examples of boron clusters. These carboranes are represented by three different isomers depending on the relative positions of the carbon atoms: *ortho-, meta-,* and *para*- $C_2B_{10}H_{12}$. During the past few decades, carborane chemistry has flourished due to its unique electronic, steric, and chemical properties. The unusually encumbered steric profile of the cluster is comparable to adamantane in van der Waals volume⁴² and is further increased due to the presence of five neighboring cluster atoms adjacent to an exohedral bond in the icosahedral molecule.

Similarly to aromatic hydrocarbons, which feature delocalized bonding, the 3-D aromaticity of carborane involves extensive delocalized molecular orbitals^{27–29,43,44}. The neutral carboranes possess a relatively low-energy LUMO that renders them reasonably strong electron acceptors amenable to electrochemical or chemical reduction.

The C–H bonds in carboranes are relatively acidic (pK_a is ca. 23), and as such can be deprotonated with strong bases, resulting in carbanions that can be utilized in reactions with organic or inorganic electrophiles, providing a major route for carbon vertex derivatization.^{45,46} Factors driving the selectivity of carborane deprotonation and functionalization have been extensively studied.^{47–49} The carbon atoms of *ortho*-C₂B₁₀H₁₂ form single bonds with exohedral substituents, and π -backdonation from those groups to the cluster can simultaneously increase the exohedral bond order and weaken the intracluster C–C bond.^{50–52} There have been theoretical

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studies of the electronic and steric effects of exohedral substituents on this C–C bond length.^{53,54} The electronic interaction between exohedral donor groups and the carborane cluster can be very significant, as evidenced by the recently disclosed structure of the diamine derivative 1,2-(MesCH₂NH)₂-C₂B₁₀H₁₀, which features one of the longest reported C–C bonds (1.931(3) Å, cf. 1.624(8) Å for the parent unsubstituted cluster).⁵⁵ Natural bond order (NBO) analysis attributed this extreme bond elongation to the efficient negative hyperconjugation between the lone pairs of the nitrogen atoms and the antibonding σ^*_{C-C} orbital in the *ortho*-carborane cluster.⁵⁶

One feature, pertinent to this review, is the ability of the ortho-carborane, closo-{C₂B₁₀}, cluster to accept two electrons with conversion to the open *nido*- $\{C_2B_{10}\}$ isomer.^{57,58} The cyclic voltammograms of $\textit{ortho-}C_2B_{10}H_{12}$ and its derivatives often exhibit an unresolved two-electron reduction wave.59-61 For instance, $ortho-C_2B_{10}H_{12}$ has been reported to undergo reduction to a dianion at a potential of -2.9 V vs. Fc⁺/Fc.^{62,63} Significant structural rearrangements occur during cluster opening upon reduction, causing a quasi-reversible or irreversible appearance of the cyclic voltammograms.⁶⁴ The structure of open-cage isomers that form upon reduction is much less studied, with only a few examples of isolated reduced clusters.⁶⁴ Exohedral substituents have been documented to have significant effects on the reduction potentials of carborane derivatives.58 For example, the chemical reduction of unsubstituted $ortho-C_2B_{10}H_{12}$ with sodium metal produces the 7,9-C₂B₁₀H₁₂²⁻ isomer that features an open six-membered ring face, while our results presented below show that the chemical reduction of carboranyl diphosphines affords a different, C_{2} symmetric cage isomer (Fig. 1).^{57,65} In addition, the outcome of cluster reduction depends on the reagent used, as demonstrated by the reaction of $ortho-C_2B_{10}H_{12}$ and magnesium metal in the presence of dibromoethane, which, upon workup, cleanly produces a different 7,10-C₂B₁₀H₁₃⁻ cluster. 66,67



This feature article will focus on the recent discoveries by our group and others pertinent to the electronic structure of *ortho*- $C_2B_{10}H_{12}$ clusters and the utilization of their redox properties in the metal-free activation of strong bonds as well as in ligand design.

Bond Activations by Carboranyl Diphosphines

Conceptually, carboranyl diphosphines represent a molecular platform that merges two important features normally exhibited by transition metals with a relatively narrow HOMO–LUMO separation: the ability of the boron cluster to accept electrons and cycle through two-electron redox states, and the ability of the phosphine groups to donate electrons and provide binding sites. The resulting interplay between these two spatially separated fragments of the metal-free carboranyl diphosphine system opens a new reactivity manifold where coordination of a substrate and subsequent two-electron oxidative addition occur with the cleavage of substrate bonds.

The first example of the redox-active behavior of carboranyl diphosphines is the work of Pringle and co-workers.⁶⁸ The group synthesized a series of $1,2-\{C_2B_{10}\}$ diphosphines bearing various PR₂ groups at adjacent carbon atoms of the cluster, for example, ortho-1-P^tBu₂-2-PEt₂-C₂B₁₀H₁₀. It was noted that the ³¹P NMR spectrum of this compound in chloroform-d slowly changed upon standing for 28 days, with the appearance of two new singlets, one of which corresponded to the presence of a direct P-H bond. The crystal structure of the product revealed the cage-opening of the parent *closo*-{C₂B₁₀} cluster and its conversion to the *nido*-{C₂B₁₀} form. The more basic P^tBu₂ group was selectively protonated in the product, and the PEt₂ group was converted to a PEt₂Cl fragment. This compound could also be directly obtained in the reaction of the starting carboranyl diphosphine and ethereal HCl (Fig. 2). Interestingly, the reverse reaction occurred easily upon the addition of triethylamine to the nido-{C₂B₁₀} product. The overall outcome of this reaction is therefore a formation of the zwitterionic product containing two phosphonium cations and the dianionic nido-{C2B10} cluster. The open cluster geometry was different from the previously reported *nido*-{C₂B₁₀} anions and can be described as a derivative of a C_2 -symmetric 13-vertex docosahedron. Importantly, "regular" redox-inactive phosphines do not exhibit such reactivity with HCl, and instead become protonated at the phosphorus atom with the outer sphere halide counterion in the products.

Fig. 1 Chemical reduction of *ortho*-carborane (top) and carboranyl diphosphines (bottom), highlighting the different $\textit{nido}-\{C_2B_{10}\}$ geometries.



Fig. 2 Cage-opening reactivity of a carboranyl diphosphine with HCl.

The Hey-Hawkins group reported a related carboranyl diphosphetane that displays similar oxidative addition-type reactivity in the presence of Mel.⁶⁹ The synthesized carboranyl diphosphetane undergoes P-P bond activation and cluster opening upon reduction with lithium metal. The reaction of the resulting Li intermediate with excess MeI induces the methylation of both phosphorus centers, generating a zwitterionic bis-phosphonium-nido-carborane (Fig. 3). An attempt to obtain this open cluster compound though the direct reaction of the neutral diphosphine 1,2-(P^tBuMe)₂-C₂B₁₀H₁₀ with excess MeI resulted in the observed formation of the same product along with significant unidentified side products. It can be surmised that in order to form the diphosphonium zwitterion, the iodide counterions need to be oxidized by the cluster, generating I_2 and thus highlighting the key role of the carborane in this redox-active system.



Fig. 3 Reaction of a carboranyl diphosphetane with Mel.

These early examples, along with the findings of intramolecular bond activation in biscarboranyl phosphines (see the section below) inspired us to devise the system that would target metal-free reactivity driven by the redox rearrangement of the carborane cluster.

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The targeted metal-free reactivity in main-group systems is based on the lowering in energy of the LUMO with the simultaneous maintenance or increase in energy of the HOMO. For carboranyl diphosphines, the HOMO can be envisioned to be centered on the phosphorus atoms, and therefore, the increase in the electron-donating abilities of the phosphine groups can be achieved by the utilization of alkyl substituents, such as tert-butyl or isopropyl groups. The LUMO is likely to be localized within the cluster; however, its energy is also governed by the conjugation with exohedral substituents. As mentioned in the introduction, the negative hyperconjugation between the lone pairs of the phosphine groups and the antibonding σ^*_{c-c} orbital within the ortho-carborane cluster can significantly increase the LUMO level, weakening the intracluster C–C bond and facilitating its opening upon reduction. Furthermore, the phosphine groups increase the steric encumbrance, thus contributing to the destabilization and lengthening of the intracluster C-C bond. Thus, in our work, we aimed to maximize both the electron-donating ability and the steric hindrance of phosphine substituents on the ortho-carborane cluster.

A carborane bearing two electron-rich and sterically crowded phosphines, 1-P^tBu₂-2-PⁱPr₂-C₂B₁₀H₁₀ (1), was obtained by stepwise deprotonation of the C-H bonds of ortho-C₂B₁₀H₁₂ and reactions with P^tBu₂Cl and PⁱPr₂Cl, respectively (Fig. 4). The ³¹P NMR spectrum of **1** contains two doublets with a large coupling constant ${}^{3}J_{PP}$ = 90 Hz, indicating a strong through-bond interaction between phosphorus nuclei. The presence of two inequivalent phosphines in the molecule and their strong ${}^{3}J_{PP}$ coupling in the neutral closo- cluster was expected to be advantageous in exploring the redox reactivity of this compound. As shown below, the transformation of the closocluster to nido- form upon reduction proceeds with the cleavage of the intracluster C–C bond and leads to the lack of appreciable coupling between the two inequivalent phosphorus nuclei, giving rise to a pair of singlets in ³¹P NMR spectra. Single crystal X-ray diffraction confirmed the expected connectivity and revealed an intracluster carbon-carbon bond length of 1.832(2) Å, which is drastically longer than that of the unsubstituted ortho-C₂B₁₀H₁₂ cluster (1.624(8) Å)⁷⁰ and is the longest reported C–C distance for any carboranyl diphosphine.⁷¹ This elongation was attributed to the aforementioned combination of the steric hindrance of the substituents as well as the electronic effects of the lone pairs of the electron-rich phosphine groups on the electron-accepting carborane cluster.

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Fig. 4 Synthesis and displacement ellipsoid plot (50% probability) of 1. Hydrogen atoms of alkyl groups are omitted for clarity.

Theoretical calculations demonstrated that the HOMO and HOMO–1 of **1** are localized at the lone pairs on the phosphorus atoms, while the major components of the LUMO were the antibonding σ^*_{C-C} orbital and the bonding π_{C-P} orbitals, which supports the hypothesis that **1** is able to both efficiently donate electrons through the phosphine groups and accept electrons in the cluster, resulting in the cage opening and an increase in C–P bond order. Thus, the carboranyl diphosphine **1** can be considered as an ambiphilic molecule with spatially separated but conjugated electron-donating and electron-accepting regions.

We probed the redox behavior of **1** both chemically and electrochemically. Cyclic voltammograms of **1** exhibited quasiirreversible overlapping reduction events with the position and peak shapes being solvent-dependent. Chemical reduction of **1** with sodium metal in THF resulted in the clean formation of a dianionic open cluster diphosphine **1**^{2–}, which was confirmed by NMR spectroscopy. Additionally, the structure of the reduced form of a congener of **1**, 1,2-(PⁱPr₂)₂-C₂B₁₀H₁₀, was determined by X-ray crystallography. Its crystal structure revealed the *C*₂symmetric dianionic *nido*-{C₂B₁₀} cluster, isostructural to those reported in the Pringle and Hey-Hawkins works mentioned above.

This molecular design proved advantageous for the target metal-free bond activation reactivity. We found that **1** undergoes oxidative addition-type reactions with a rather broad range of substrates beyond just stronger electrophiles such as HCl and Mel. Reactions of **1** with main group hydrides, alcohols, and electron-deficient terminal alkynes resulted in generally similar oxidative addition-type *nido*-{C₂B₁₀} products – that is, the carborane cluster experienced a two-electron reduction with concurrent rupture of the intracluster C–C bond, and the phosphorus centers were converted to phosphonium cations via protonation or addition of substrate fragments.

For example, the reaction of **1** and tributylstannane proceeded cleanly to afford a single product according to ${}^{31}P{}^{1}H$ NMR spectroscopy. The most noticeable change in the NMR spectrum was the disappearance of the two doublet

signals from the inequivalent phosphine groups in 1 and the appearance of two new singlets. Furthermore, the protoncoupled ³¹P NMR spectrum of the product contained two doublets (¹J_{PH} of 432 and 437 Hz) corresponding to direct P–H bonds. The two-electron reduction of the closo-cluster to the nido-form and the simultaneous conversion of the phosphine groups into protonated phosphonium cations was accompanied by the formation of the tin-tin bond in the dimeric species Bu₃Sn–SnBu₃, the presence of which was confirmed with ¹¹⁹Sn NMR spectroscopy. The crystal structure determination corroborated the spectral data and revealed the formation of a zwitterionic product featuring the open dianionic *nido*- $\{C_2B_{10}\}$ cluster with its separated carbon atoms and protonated phosphonium centers (2) (Fig. 5). The open cluster exhibited the same C_2 -symmetric geometry as that in the structure 1^{2-} . The exohedral bond lengths in 2 (1.747(1) Å and 1.751(1) Å), however, were significantly shorter that those in parent ${\bf 1}$ (1.878(1) Å and 1.915(1) Å), which indicates their partial ylide character and corroborates the DFT calculation results described above.

The reactions of **1** with HBpin, NH₃BH₃, or BH₃(SMe₂) produce the same open cluster 2, albeit in lower yields (Fig. 5). Careful monitoring of the reaction progress between 1 and BH₃(SMe₂) by ³¹P NMR spectroscopy revealed the initial formation of two major open cluster products both containing P–H and P–B bonds, as evidenced by the presence of two pairs of sharp singlets and broad 1:1:1:1 quartets in the ³¹P{¹H} NMR spectra of the reaction mixture. We were able to isolate and determine the crystal structure of one of these products, which featured the same open dianionic *nido*-{C₂B₁₀} cluster geometry along with a protonated H–P^tBu₂ group and an (SMe₂)BH₂–ⁱPr₂ phospinoboryl group. This apparent product of the borane B-H bond activation by carboranyl diphosphine 1 eventually converts into 2 upon standing in solution. The reaction of 1 and ammonia borane required elevated temperatures to proceed and, along with the formation of 2, was accompanied by the appearance of signals corresponding to oligomeric N-B containing products in the ¹¹B NMR spectrum.

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Fig. 5 Activation of main group hydrides by **1** (top) and displacement ellipsoid plot (50% probability) of **2** (bottom). Hydrogen atoms of alkyl groups are omitted for clarity.

Similar reactivity was observed in the activation of alcohols by **1**, producing P^tBu₂(OR)-PⁱPr₂(H)-*nido*-C₂B₁₀H₁₀ (**3a** R = Me, **3b** R = OⁱPr). The reaction proceeded in the presence of an acid, suggesting that the protonation of one of the phosphine groups is an initial step. Interestingly, in the final products **3a/3b**, protonation occurs at the PⁱPr₂ group, and the more basic P^tBu₂ group is connected to the alkoxide fragment. The cluster in **3a/3b** adopts the same open dianionic *nido*-{C₂B₁₀} geometry as in **2**.

A closely related example of bond activation by redox-active **1** was discovered in its reactions with alkynes.⁷² We found that electron-deficient terminal alkynes were activated by our system in a process that is driven by cluster reduction and formation of the *nido*-{ C_2B_{10} } cage, yet the products **4a/4b** did not follow the same reactivity pattern as the case described above (**Fig. 6**). Instead, the final outcome of the reaction is the regioselective activation of the B–H bond of the boron atom adjacent to both carbon atoms in the cluster. As a result, a boron–carbon bond is formed, and the alkyne fragment connects to the phosphorus atom of the P^tBu₂ group, completing a five-membered C–P–C(H)=C(Ar)–B cycle. The C–H

bond of the terminal alkyne is retained, and the activated B–H bond of the cluster protonates the P^iPr_2 group, as confirmed by deuterium labeling experiments. The transformation of the formally hydridic B–H bond into a proton provides two electrons for cluster reduction and opening. This cascading sequence of elementary steps creates a unique boron-containing cationic phosphacycle fused to the anionic *nido*-{C₂B₁₀} cluster. Importantly, the presence of the two phosphine groups is important for this outcome, as carboranyl monophosphines do not exhibit B–H bond activation reactivity. ^{73,74}



Fig. 6 Activation of electron-deficient terminal alkynes by 1.

To reiterate, the unusual reactivity of carboranyl diphosphine **1** described herein relies on the synergy between the electron-donating phosphine groups and the electron-accepting carborane cluster. In these novel transformations, the neutral closo-{C₂B₁₀} cluster accepts two electrons and becomes an open dianionic nido-{C₂B₁₀} cluster, while the two phosphine groups are converted into phosphonium cations. The resulting zwitterionic products exhibit ylide-type phosphorus–cluster bonding. Thus, the overall reactivity may be classified as the oxidative addition of substrates (i.e., main group hydrides, alcohols, and electron-deficient terminal alkynes) to a carboranyl diphosphine.

Prior to our work with carboranyl diphosphine **1**, we observed intriguing reactivity while attempting to synthesize 1,1'-bis-*ortho*-carboranyl diphosphines. The structure of 1,1'-bis-*ortho*-carborane ("biscarborane"), similarly to biphenyl, features two carborane clusters joined together by a single carbon–carbon bond. In an attempt to obtain biscarboranyl diphosphines by deprotonation of two C–H bonds of biscarborane and a subsequent reaction with phosphine chlorides, we instead observed unprecedented regioselective cluster B–H activation at room temperature.⁷⁵ In retrospect, these transformations described below correspond to the promotion of intramolecular bond activation by a redox-active cluster, and are conceptually the same as the aforementioned examples of the reactivity of **1**.

Instead of the intended diphosphine, zwitterionic *closo*- $\{C_2B_{10}\}$ -*nido*- $\{C_2B_{10}\}$ clusters featuring an open *nido*- $\{C_2B_{10}\}$ cluster and two phosphonium centers were isolated (**Fig. 7**). Both phosphorus atoms are connected to the carbon atoms of the biscarborane cage; therefore, one can assume that the installation of PR₂ groups to the cluster carbons indeed occurred in the initial stages of the reaction. However, the

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proximity of the phosphine substituents to the neighboring cluster evidently resulted in a transition state that corresponded to the close B–H···P contacts. The ensuing B–H bond activation is promoted by the *closo*-{C₂B₁₀} cluster accepting two electrons from the formally hydridic B–H bond and releasing a proton. The cluster is thus converted to the open dianionic *nido*-{C₂B₁₀} form, the P–B bond is formed, and the proton eventually migrates to the remaining neutral phosphine substituent, resulting in a protonated phosphonium cation. Importantly, the geometry of the *nido*-{C₂B₁₀} cluster is the same as for the zwitterionic products derived from 1 that are described above. Thus, the observed intramolecular B–H bond activation and the formation of a five membered C–C–B–P–C cycle is akin to the reactivity of 1 with alkynes.

Furthermore, our group has reported the ability of the zwitterionic compound $closo-(C_2B_{10}H_{10})(P\{N^iPr_2\}_2)-nido (C_2B_{10}H_9)(PH\{N^iPr_2\}_2)$ to activate the O-H bonds of water at room temperature, resulting in the partial removal of one boron and one carbon vertex from the $nido-\{C_2B_{10}\}$ cluster and conversion to the nido-{CB9} cage. This series of transformations culminates in the addition of H and OH fragments to separate sites of the molecule (Fig. 7).76 While deboronation of carboranes is generally considered irreversible and requires external boron sources to reform the cluster, this reaction is reversible upon formal dehydration of the unusual boronic acid hydride (C)B(H)(OH) fragment at elevated temperatures, releasing water and rearranging to the starting $nido{\{C_2B_{10}\}}$ cage.



Fig. 7 Intramolecular B–H bond activation with biscarboranyl diphosphines.

Redox-Active Carboranes in the Formation and Stabilization of Low-Valent Main Group Centers

In addition to facilitating the activation of strong bonds, carboranes in the pre-formed open nido-{C₂B₁₀} structure have

also been employed in facilitating the formation and stabilization of unique main group centers. One such candidate for this design is a carborane cluster decorated with two intramolecularly donor-stabilized N-heterocyclic silylene (NHSi)⁷⁷ groups, **5**, which is isolobal to carboranyl diphosphines and is therefore able to accept and release electrons (**Fig. 8**).



Fig. 8 Synthesis and displacement ellipsoid plot (50% probability) of 5. Hydrogen atoms of alkyl and aryl groups are omitted for clarity.

Recently, Xie and co-workers have demonstrated a new method of direct borylene insertion into a B–H bond by employing **5** as a ligand.⁷⁸ The monobromoborylene **6** features a nucleophilic, electron-rich boron(I) center, stabilized by carborane-connected, strongly donating silylene groups. Upon reacting with sodium naphthalenide, **6** was shown to undergo the unprecedented insertion of a borylene fragment into a carborane cluster B–H bond, forming a new B–B bond. This unique reactivity is likely promoted by the electron-accepting propensity of the carborane cluster and the nucleophilicity of the borylene.

Mechanistic studies showed that an unstable carborane radical anion forms as a result of a one-electron reduction, promoting an intramolecular single-electron transfer from the boron(I) center to the carborane cluster. As a consequence, the more stable dianionic *nido*- $\{C_2B_{10}\}$ cage and a boron-centered radical cation form (**Fig. 9**). During the reduction of the carborane cage, carbon–silicon bonds shorten, indicating a partial ylide character, analogous to the zwitterionic carboranyl diphosphines described in the previous section. Further reduction with sodium abstracts the bromide and creates an active borylene species, which then inserts into a B–H bond of the reduced cluster, creating a B–B bond. The capability of the carborane cluster to accept two electrons is crucial for this reactivity.



Fig. 9 Construction of B-B bond assisted by cage reduction.

The ability of the carborane-linked bis-NHSi backbone **5** to reversibly accept and release electrons has also been applied to promote the formation and stabilization of low-valent compounds. Driess and co-workers reported the ability of **5** to chelate and stabilize a zero-valent silicon center, named "silylone".⁷⁹ Similar to the aforementioned borylene insertion into a cluster B–H bond, the mechanism depends on the electron-accepting ability of the carborane cluster to facilitate electron transfer. Further reduction of a low-valent Si(I)–Si(I) complex **8**, featuring two supporting dianionic *nido*-{C₂B₁₀} clusters (**Fig. 10**).



Fig. 10 Synthesis of silylone 7 via a dianionic *nido*-carboranyl bis-NHSi and the Si–Si homocoupling product 8.

Stabilization of a monovalent nitrogen was also realized in a similar manner by Driess and co-workers. (**Fig. 11**).⁸⁰ The reaction of **5** with adamantyl azide involves the reduction of the carborane cluster and the formation of silylene cations, generating zwitterionic product **9**.



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Fig. 11 Synthesis of nido-carboranyl bis-silylene-adamantyl azide adduct 9.

The reduction of **9** with KC₈ in the presence of 18-crown-6 results in one-electron reduction and the loss of adamantyl and dinitrogen fragments to form a new anionic compound **10** (Fig. **12**). The positive charge is localized at the nitrogen(I) center instead of the supporting silicon atoms, and the dinegative charge remains within the cluster, as indicated by the maintenance of a *nido*-{C₂B₁₀} geometry. The open cluster in **9** can be oxidized when reacted with AgOTf, which results in the formation of cationic compound **11**, with closure of the cluster, re-formation of the intracluster C–C bond, and loss of dinitrogen and adamantyl fragments. The compounds **10** and **11** coproportionate into a carborane-based radical species **12**.



Fig. 12 Synthesis of low-valent nitrogen complexes 10 and 11 from 9 and the carborane-based radical nitrogen(I) species 12 from 10 and/or 11.

Driess and co-workers recently reported the conceptually similar use of 5 in the synthesis of zero- and monovalent germanium species.⁸¹ One-electron reduction of the zerovalent germanium(0) complex 13 was shown to induce an intramolecular electron transfer, reducing the carborane cluster by two electrons (exemplified by the transition from closo-{ C_2B_{10} } to the nido-{ C_2B_{10} } structure) and oxidizing the germanium center to Ge(I). This one-electron reduction, as with silylone 7, facilitates Ge–Ge coupling to form a low-valent dimer complex 14. Meanwhile, one-electron oxidation bv ferrocenium forms a similar dicationic Ge(I)–Ge(I) complex 15 where both carborane clusters remain in the closo-form, thus not involving the redox properties of carborane for Ge(I) generation (Fig. 13).



Fig. 13 Synthesis of Ge(I)–Ge(I) complexes 14 and 15 from germylone 13.

The carboranyl bis-NHSi ligand **5** was also employed in reactions with alkynes.⁸² Treatment of **5** with either phenylacetylene or diphenylacetylene led to the formation of cyclic products **16a/16b** (**Fig. 14**), which features a C=C bridge between the two silicon centers and the open *nido*-{C₂B₁₀} cluster. The exohedral C–Si bonds in **16a** are rather short (1.769(2) Å), which is consistent with its zwitterionic formulation. The geometry of the open cluster is analogous to that of **8**, **9**, **10**, **12**, and **14**.



Fig. 14 Synthesis of the zwitterionic cyclic products 16a/16b.

The presence of the redox-active carborane cluster in the ligand backbone of **5** is integral to the formation of the aforementioned unique examples of low-valent Si, N, and Ge species. In most cases, the initial one-electron reduction of bissilylene **5** drives the two-electron reduction of the carborane cluster, governed by the comparative stability of the dianionic *nido*-{C₂B₁₀} cluster with respect to its radical monoanionic form; thus the second electron for the reduction of the cluster is sourced intramolecularly from the coordinated main group element center. For the germanium and silicon complexes, this results in the formation of Ge or Si radicals, which subsequently couple intermolecularly to form their respective dimers. For the

low-valent nitrogen complex, this one-electron intramolecular oxidation results in the fragmentation of the adamantyl azide moiety and the formation of the nitrogen(I) cation.

Redox-Active Carboranes as Supporting Ligands for Transition Metals

C–C Bond Activation with Metallacarboranes

The activation of C–C bonds often requires the involvement of transition metal catalysts. Among different classes of organic compounds, aromatic carbon–carbon bonds are some of the most stable and most challenging for activation. Welch and coworkers have reported an interesting example of aromatic C–C bond cleavage in a simple arene at room temperature by a metallacarborane complex.⁸³

The reduction of the biscarborane cluster with excess lithium metal and the subsequent reaction with $[(p-cymene)RuCl_2]_2$ failed to produce the intended C–C bridged bis-13-vertex metallacarborane. Instead, only one cluster of biscarborane was expanded with the metal vertex, producing an $\{RuC_2B_{10}\}$ cluster. Surprisingly, a triple-decker "fly-over bridge" bimetallic complex formed, with the rupture of the C–C bond in one of initial *p*-cymene ligands (**Fig. 15**). The results of DFT calculations suggest that the mechanism of the aromatic C–C bond activation involves an intramolecular electron transfer from the reduced pendant carborane cluster to the coordinated arene ligand. This discovery highlights the role of the carborane cluster as an electron reservoir for a subsequent redox process at a remote metal site.



Fig. 15 Synthesis of fly-over bridge complex 17.

Change of Ligand Coordination Mode

Ménard, Dobrovetsky, Hayton, and co-workers recently illustrated the novel use of a carborane-based ligand for the capture and release of uranium in the form of $UO_2^{2+.84}$ Reduction of neutral 1,2-(Ph₂PO)₂-1,2-*closo*-C₂B₁₀H₁₀ by two electrons leads the formation of the dianionic *nido*-{C₂B₁₀} cluster. This cage opening upon reduction leads to a significant

increase in the geometrical separation of the exohedral phosphine oxide binding sites, thus dramatically changing the bite angle of this ligand. This increased bite angle facilitates its coordination to the large uranyl cation. Once coordinated, the UO₂²⁺ fragment can be effectively released through oxidation, converting the cluster backbone of the ligand back to the closoform and decreasing the bite angle (Fig. 16). This reaction proceeds reversibly via the chemical or electrochemical reduction of the carborane compound, making this a controllable redox cycle. This development has substantial potential in the processing and separation of nuclear fuel waste. Both neutral and dianionic forms of the ligand are stable in the presence of water and thus the concept of geometric changes driven by the redox cycling of carboranyl ligands has been extended to the selective biphasic extraction and release of uranyl.85



Fig. 16 Capture and release of UO₂²⁺ by a redox-active carborane derivative.

Conclusions and Outlook

The findings summarized in this feature article put forward a new mode of metal-free bond activation reactions by a main group cluster systems. The 3D aromaticity and electronic properties of carboranes, as well as the influence of exohedral substituents, have been investigated and harnessed since the discovery of these intriguing clusters. The reversible redox transformations of carborane clusters have proven to be a tool for a diverse range of applications, as exemplified by the reactivity discussed in this article. The tunability of these cluster compounds through exohedral substitution will likely lead to new bond activation discoveries as even subtle changes in the substituents can have a profound impact on the reactivity. The electron-accepting ability of the icosahedral carborane cluster can be used to facilitate metal-free reactivity by inducing a transition from the neutral *closo*-{ C_2B_{10} } to the dianionic *nido*-{ C_2B_{10} } structure. Furthermore, the redox properties of the cluster could be utilized in an indirect way, as the geometric changes associated with the opening of the cluster upon reduction can be used to alter its coordination chemistry.

Conflicts of interest

There are no conflicts to declare

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