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Cesium Carbonate Mediated C-H Functionalization of Perhalogenated 12-vertex Carborane Anions

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C-H functionalization of undecahalogenated carborane anions, $[HCB_{11}X_{11}^{-1}]$ (X = Cl, Br, I), is performed with Cs₂CO₃ in acetonitrile. We show that the requisite Cl, Br and I carborane dianions can all be efficiently accessed with Cs₂CO₃. The utilization of Cs₂CO₃ eliminates the complications associated with competing E2 elimination reactions providing an efficient, more functional group tolerant, and broader scope than previously reported. The ensuing functionalized cages provide potential synthons for constructing advanced materials and other molecular architectures for various applications.

Weakly-coordinating anions (WCAs) are highly useful tools that allow the full expression of their countercations reactivity profiles.¹ Their applications range from catalysis,²⁻⁵ isolation of reactive intermediates,^{6, 7} and materials relevant to batteries and other devices.² Ideally, WCAs function as inert charge carriers that do not undergo undesirable side reactions. Closo $carborane^{8,\ 9}\ anions^{2,\ 7,\ 10\text{-}12}$ are a class of WCAs, which are carbon containing analogues of polyhedral boranes. The 12vertex carborane anions [RCB₁₁X₁₁¹⁻], and derivatives of the isoelectronic $[B_{12}H_{12}^{2-}]$ dianion, are the most stable,⁷ and they display unmatched chemical, electrochemical,^{2, 11, 13} and thermally stability. These anions can be readily modified at the B-vertices with electrophiles, such as halogens, which render them more weakly coordinating and robust.¹⁰ Additionally, deprotonation of the parent 12-vertex carborane anion [HCB₁₁H₁₁¹⁻] carborane with very strong bases (i.e. *n*-BuLi or metal hydrides),¹⁰ leads to the formation of the isolable^{14, 15} [CB₁₁H₁₁²⁻] dianions that are nucleophilic but also very basic at carbon.¹⁰ This basicity often leads to side reactions with the THF solvent, as well as elimination reactions, instead of clean nucleophilic substitution. As first demonstrated by Strauss,¹⁶ perhalogenation of the carborane cage, specifically with fluorine, leads to dramatically enhanced acidity of the C-H vertex, enabling it to be deprotonated with relatively weak bases, such as hydroxide. However, the broad utility of this fluorinated anion is tempered by its facile cage degradation in aqueous hydroxide solution and its hazardous synthesis with F_2/HF . It should be noted that Vinas and Teixidor have been able to achieve some interesting functionalization reactions of halogenated ortho-and nido carboranes in aqueous media.¹⁷⁻²⁰

A more accessible and base stable cluster is the $[HCB_{11}CI_{11}^{11}]$ anion $1CI_{11}$ (Figure 1).²¹ Several years ago Ozerov and coworkers reported¹⁷ that $1CI_{11}$ could be deprotonated with the strong base KOt-Bu and subsequently functionalized with simple alkyl electrophiles. This procedure is a 2-step/1-pot process utilizing *t*-butoxide (3.5 eq) in *t*-butanol solvent, with subsequent addition of an alkyl iodide (7eq). Both the order of addition and excess base/electrophile are important to fully drive the reaction to completion, as both the base and alkyl iodide are consumed in undesirable E2 elimination reactions. This is perhaps the reason only terminal straight chain alkyl iodides were the sole substrates reported in this manuscript.

Here we report our findings on the C-H functionalization of $[HCB_{11}CI_{11}^{1-}]$ with the relatively weak base Cs_2CO_3 . We show that it is possible to efficiently achieve C-alkylation of the cage with diverse alkyl groups containing functionality, which makes the ensuing compounds potential building blocks for advanced materials and other downstream applications. Importantly we also show that such reactions are also possible with $[HCB_{11}Br_{11}^{1-}]$ and $[HCB_{11}I_{11}^{1-}]$.



Figure 1. Synthesis of C-functionalized $[RCB_{11}X_{11}^{1-}]$ anions $3RX_{11}$ with Cs_2CO_3 .

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⁺ Footnotes relating to the title and/or authors should appear here.

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While investigating the possibility of preparing functionalized carboranes with enhanced solubility in alkane solvents for silylium catalysis,² Nelson's team discovered that Cs_2CO_3 is sufficiently basic to generate and functionalize [HCB₁₁Cl₁₁¹⁻] (Figure 1). Given Lavallo's interest in functionalized clusters for ligand design and battery applications, we decided to develop this chemistry, but first probe the formation of the C-nucleophilic dianion [CB₁₁Cl₁₁²⁻] **2Cl₁₁** (Figure 1).

As evidenced by ¹¹B-NMR spectroscopy reaction of $[HNMe_3^+][HCB_{11}Cl_{11}^{1-}]$ with two equivalents of Cs_2CO_3 (one eq is to destroy the HNMe3⁺ cation) in dry acetonitrile generates an equilibrium mixture (33:67) of the starting material and the dianion 2Cl₁₁, respectively (Figure 2). The equilibrium could be driven to essentially 100% dianion by increasing the amount of base to 4 equivalents. After confirming that a significant population of the dianion $\mathbf{2Cl}_{11}$ could be generated in solution with Cs₂CO₃ we were curious if related perhalogenated species $1Br_{11} \mbox{ and } 1I_{11} \mbox{ could undergo similar reactions, even though }$ they contained less electron withdrawing halides. Indeed, as indicated by ¹¹B-NMR we discovered that both 1Br₁₁ and 1I₁₁ have sufficiently acidic C-H vertices to undergo deprotonation with Cs₂CO₃. As one might expect because of electronegativity trends, under the same conditions (2eq base), 50:50 and 75:25 equilibrium mixtures of 1Br₁₁/2Br₁₁ and 1I₁₁/2I₁₁ formed, respectively (Figure 2). It should be noted we found that

hexahalogenated species $[H_6CB_{11}X_6^{1-}]$ (X= Cl, Br, or I) are not suitable substrates under these reaction conditions, which highlights the importance of perhalogenation to achieve deprotonation with the weak base Cs_2CO_3 .



Figure 2. HNMe₃⁺ salts of $1X_{11}$ reacted with 2 eq of Cs₂CO₃ yields equilibrium mixtures of $1X_{11}$ and $2X_{11}$.

The use of Cs₂CO₃ to generate the dianion was predicted to be adventitious for the functionalization of these cages, as E2 elimination can be avoided and other functional groups not compatible with K-OtBu could be utilized. We subsequently began to explore different classes of electrophiles that may be utilized to functionalize the cages to produce derivatives 3RX11 (Figure 3). Under optimized reaction conditions 1Cl₁₁ with ndecyl iodide (1.1 eq), Cs₂CO₃ (5 eq), dissolved in acetonitrile in open air, the reaction proceeds to completion with a 96% isolated yield of a in 18 hours at room temperature (Figure 3). Likewise, the bromo analogue 1Br11 was also functionalized to afford **b** in identical yield under the same conditions (Figure 3). However, while the iodo derivative could be similarly functionalized to produce c in comparable yield, it required the use of an inert atmosphere and dry acetonitrile for consistent results, likely due to adventitious water interfering with the formation of 2I₁₁. We were also able to successfully react 1CI₁₁ with more sterically hindered isopropyl iodide to afford **d** in 88% yield over 18 hours, at 85 °C (Figure 3). However, the more sterically demanding 1Br11 required 36 hours to completely form e, and the analogous reaction with 11,1 produced only 30% conversion to product **f** over one week.



Figure 3. Scope of cesium carbonate mediated C-H functionalization of $[HCB_{11}X_{11}^{1-}]$ cages **1.** Yields were calculated by weighing the purified salt products. * = conversion via NMR not isolated yield; reactions were performed at romm temperature unless indicated by ** (= 85 °C). Unlabeled vertices = B.

—After examining the scope of simple alkylations of compounds $1X_{11}$, we decided to further broaden the breadth of reactivity of $1Cl_{11}$. Utilizing the optimized procedure, we

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began investigating suitable substrates tolerant of the reaction conditions. Derivative g, featuring an epoxide moiety, was synthesized in 89% yield via nucleophilic addition to epichlorohydrin. Benzyl substituted cluster h was synthesized in 96% yield using vinylbenzyl chloride as the electrophile. The norbornene functionalized cage i was prepared in modest yield (64%) via reaction with the corresponding norbornyl tosylate, although the reaction required 5 days and heating to 85 °C. Ester derivative j was accessible in 95% yield via the reaction of $1Cl_{11}$ with ethyl bromoacetate. Crotyl substituted species k was also isolated in moderate yield (72%) using technical grade 75% ethyl bromocrotonate as the electrophile. Compound I, featuring a pendant nitrile group, was prepared with bromopropionitrile in 92% yield. As demonstrated by the formation of compound m, a cluster functionalized with an acetal group could also be produced in 90% yield, but required heating to 85 °C. Lastly we utilized dibromoethane as an electrophile, affording compound n in 93% yield. Notably, no evidence for side elimination reactions were observed.

The above communication illustrates a facile method to access dianionic carboranes $2X_{11}$ derived from $[HCB_{11}X_{11}^{1-}]$ $1X_{11}$. When the formation of these species is conducted in the presence of suitable electrophiles, efficient C-H functionalization of the cage is achieved. This novel methodology allows access to various hitherto unknown functionalized carborane building blocks that will likely be useful for catalyst and electrochemical materials design. We currently are investigating the utilization of these products to form single ion conducting polymers for electrochemical cells.

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Author Contributions

All authors contributed equally to the manuscript.

Conflicts of interest

There are no conflicts to declare

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