

B–N, B–O, and B–CN Bond Formation via Palladium-Catalyzed Cross-Coupling of B-Bromo-Carboranes

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Supporting Information

ABSTRACT: Carboranes are boron-rich molecules that can be functionalized through metal-catalyzed crosscoupling. Here, for the first time, we report the use of bromo-carboranes in palladium-catalyzed cross-coupling for efficient B–N, B–O, and unprecedented B–CN bond formation. In many cases bromo-carboranes outperform the traditionally utilized iodo-carborane species. This marked difference in reactivity is leveraged to circumvent multistep functionalization by directly coupling small nucleophiles (-OH, -NH₂, and -CN) and multiple functional groups onto the boron-rich clusters.

I cosahedral carboranes are boron-rich molecular clusters that are often described as three-dimensional (3D) analogs to benzene.¹ Their unique delocalized 3D aromatic bonding, high stability, and potential for site-selective functionalization make them attractive building blocks for tunable pharmacophores, unique ligand scaffolds, and building blocks for materials applications.² Further development of these and other applications with carboranes requires efficient methods for cluster synthesis and functionalization, where ultimately each individual vertex can be specifically addressed.¹

Over the past 50 years, palladium-catalyzed cross-coupling has emerged as a powerful synthetic method for creating new molecules.³ In particular, the emergence of designer ligands (beyond PPh₃) for Pd-catalyzed cross-coupling dramatically expanded the scope of electrophile substrates beyond aryl iodides.^{4a} These new catalyst systems demonstrated a clear ability to cross-couple aryl-bromides and aryl-chlorides, thereby facilitating transformations of synthetically challenging substrates. Among existing ligand platforms, biaryl phosphine ligands significantly increased the efficacy of Pd-catalyzed C–C, C–N, and C–O bond formation.⁴

Despite these advances in catalyst design for aromatic substrates, effective methodologies for metal-catalyzed B-N, B-O and B-C cross-coupling in carboranes are lacking. In fact, only B-iodo-carboranes have been used in Pd-catalyzed crosscoupling thus far.⁵ Yet, analogy between carboranes and arenes provides a clear hypothesis that other B-functionalized electrophiles, beyond B-iodo-carboranes, may be competent cross-coupling partners. Here we report our discovery validating this hypothesis by demonstrating for the first time that B-bromo-carboranes can be efficient electrophiles for B–N, B–O, and B–CN bond formation in Pd-catalyzed cross-coupling. Furthermore, we show conditions where these B-bromo-carboranes are superior to the iodinated congeners enabling the synthesis of previously inaccessible B-substituted carboranes. This chemistry is furthermore attractive given the greater synthetic accessibility of B-bromo-carboranes compared to their iodo-based congeners (see SI).¹

Hawthorne and co-workers recently reported Pd-catalyzed amidation of 9-I-m-carborane (I-mCB) utilizing the biaryl phosphine ligand DavePhos (L1, Figure 1).^{5h} To test our hypothesis, we replaced I-mCB with the bromo-carborane congener, 9-Br-m-carborane (Br-mCB), as a substrate under the reported cross-coupling conditions. However, our initial attempts at cross-coupling trifluoroacetamide with Br-mCB proved unsuccessful. Rapid formation of Pd metal was observed without any consumption of Br-mCB. We postulated that the Pd(0) precursor (Pd_2dba_3 , dba = dibenzylideneacetone) was not efficiently forming the catalytically active species $[L1Pd^0]$. To resolve this issue, we employed a commercially available Pd(II) precatalyst (Figure 1B inset), which has been previously shown to dramatically improve catalytic activity across a large pool of aryl-based substrates and catalytic conditions. Importantly, this change tremendously improved the catalytic conversion of Br-mCB producing 1a in nearly quantitative conversion within 2 h (Figure 1A). This discovery demonstrates for the first time that one can efficiently activate a relatively inert B-Br bond in a carborane with electron-rich Pdbased species supported by a biaryl phosphine ligand (Figure 1B).

This example demonstrates the potential competence of **Br**-**mCB** toward cross-coupling (Figure 1B), which does not have any literature precedent. This advance was also appealing given that **Br-mCB** can be synthesized in a fraction of the time (1 h) that is required for the synthesis of **I-mCB** (1 day). We therefore investigated the scope of Pd-catalyzed cross-coupling

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Figure 1. (A) General amidation conditions (inset, GC-MS yield of 1a from **Br-mCB** and **I-mCB** using different palladium precursors). (B) Proposed catalytic cycle employing biaryl phosphine ligands (step i, oxidative addition; step ii, transmetalation; step iii, reductive elimination).

of **Br-mCB** with other nucleophiles utilizing biaryl-ligand containing precatalysts.

To further probe the scope of B–N bond formation using Br-mCB, we evaluated several conditions and substrates for Pdcatalyzed amination. Using morpholine as a substrate (2a, Figure 2), we evaluated the cross-coupling efficiency of three precatalysts featuring L1, SPhos (L2), and XPhos (L3) ligands (see SI). For this transformation, L2 afforded complete consumption of Br-mCB and a high amount of B-N coupling product 2a as determined by GC-MS analysis. Evaluation of various bases indicated the superior performance of K^tBuO for forming 2a. Importantly, Br-mCB showed superior crosscoupling efficiency compared to I-mCB for the formation of 2a (Figure 2A). Using these optimized conditions, cross-coupling of Br-mCB proceeds with primary, secondary, aromatic, and heterocyclic amines in nearly quantitative conversion affording the corresponding B-N compounds (2b-2e, Figure 2B and SI).

In general, cross-coupling using unprotected nitrogen-rich heterocyclic substrates is known to be challenging.^{6c} Amination of halocarboranes has only been shown on the 2-I-*p*-carborane, which is a significantly more reactive substrate than **Br-mCB**.⁷ The cross-coupling methodology we developed addresses this issue for the first time in the context of *m*-carborane chemistry since, to the best of our knowledge, **2e** represents the first product resulting from the direct cross-coupling of an unprotected five-membered heterocycle with a B-halo-*m*-carborane.

The versatility of **Br-mCB** as a cross-coupling partner can be further seen from its efficient reaction with challenging nucleophiles. For example, **Br-mCB** cross-couples with ammonia producing 2c (Figure 2B), whereas previously 2ccould only be prepared by lengthy hydrolysis of 1a.^{Sh}



Figure 2. (A) General amination and alkoxylation conditions (inset, GC-MS yield of 2a obtained from Br-mCB and I-mCB). (B) Amination scope using Br-mCB and X-ray crystal structure confirming B–N bond formation. (C) Alkoxylation scope using Br-mCB and X-ray crystal structure confirming B–O bond formation (ellipsoids at 50% probability and H atoms omitted for clarity). GC-MS yields, and isolated yields in parentheses. $*K^{T}BuO$ used as a base except for: 2e, anhydrous K₃PO₄; 2f, 1 M aqueous K₃PO₄; 2g, NaOCH₃.

Importantly, our method represents the first example of a direct cross-coupling leading to 2c and is enabled by the previously unrecognized reactivity of **Br-mCB** when using biaryl phosphine supported Pd-based catalysts.

During the course of our amination studies, we observed B–OH coupling with **Br-mCB** (2f, see SI) when nonanhydrous bases were used. This is remarkable, given that the only example of a Pd-catalyzed carborane B–O bond formation was reported on 2-I-*p*-carborane. Importantly, the I-mCB congener was previously deemed too unreactive.^{8a}

Based on these observations, we developed a new crosscoupling protocol enabling the direct coupling of water, methanol, trifluoroethanol, and 3,5-dimethylphenol with BrmCB (2f-2i, Figure 2C).

This constitutes the first reported Pd-catalyzed crosscoupling leading to a B–O bond formation with *m*-carborane substrates. Significantly, a control reaction where **I-mCB** was used as a substrate led to a significantly lower conversion to 2i (Figure 2A). This Pd-catalyzed route is also superior to the exisiting method for forming related B–O compounds utilizing carborane B-halonium salts.^{8b} Additionally, 2f can be readily converted to 2g by deprotonation with NaH and followed by treatment with MeI, demonstrating the added synthetic utility of 2f.

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The versatility of **Br-mCB** cross-coupling with small nucleophiles led us to investigate B–CN bond formation. Cyanide is known to be a difficult cross-coupling partner in metal catalysis due to its propensity toward binding to catalytically active species, resulting in their deactivation.⁹ Recently several groups reported efficient protocols for cyanation of aromatic substrates using K₄[Fe(CN)₆] as a mild cyanide source.^{9b,d} Pd-catalyzed cyanation of **Br-mCB** using K₄[Fe(CN)₆] with an L3-based precatalyst led to the formation of 9-CN-*m*-carborane in a nearly quantitive conversion (**3a**, Figure 3A). This example represents the first



Figure 3. (A) Cyanation protocol; GC-MS yield of 3a obtained from **Br-mCB** and **I-mCB** (isolated yield in parentheses). (B) X-ray crystal structure, ¹¹B and ¹³C{¹H} NMR spectra of 3a (ellipsoids at 50% probability and H atoms omitted for clarity).

direct cyanation of a halogenated derivative of dicarba-*closo*dodecaborane. Importantly, cross-coupling activity of the ImCB species under these conditions is dramatically diminished compared to Br-mCB (Figure 3A).

The ability to append multiple functional groups is crucial to developing carboranes for new and existing materials.^{2,10,11} While polyfunctionalization of arene-based electrophiles via cross-coupling is well-established, similar methods for carboranes are rare.^{5,10} Our methodology can be applied toward disubstitution cross-coupling chemistry. Specifically, 9,10-Br₂-*m*-carborane (**4a**) can be functionalized with two bulky 3,5-dimethylphenolate substituents (**4c**, Figure 4). Interestingly, under B–OH cross-coupling conditions (*vide supra*), **4a** undergoes exclusive monosubstitution to produce **4d**.

In addition, given the pronounced orthogonal reactivity of B–Br versus B–I bonds in cross-coupling, our methodology can be used to heterofunctionalize mixed halo-carborane substrates. We leveraged the selectivity of $PdCl_2(PPh_3)_2$ for B–I bond functionalization to produce 9-Br-10-Et-*m*-carborane (4e) from 9-Br-10-I-*m*-carborane (4b, Figure 4 and SI).

Selective Pd-catalyzed cross-coupling of the B–Br moiety in 4e with L2-containing precatalyst yields the heterofunctionalized 9-O- $(3,5-Me_2C_6H_3)$ -10-Et-*m*-carborane (4f). This transformation represents the first metal-catalyzed B-heterofunctionalization of dicarba-*closo*-dodecaborane via cross-coupling demonstrating that B-Br-carboranes offer an additional pathway for multifunctionalization. These experiments also suggest that our methodology is amenable to sterically encumbered carborane-based electrophiles.

Ortho-carboranes are the most challenging substrates in cross-coupling methodologies, since these species undergo



Figure 4. Difunctionalization conditions and X-ray crystal structure confirming B–O bond formation. X-ray crystal structure (ellipsoids at 50% probability and H atoms omitted for clarity); see SI for detailed conditions. GC-MS yields, and isolated yields in parentheses.

facile deboronation in the presence of nucleophiles.¹² Our conditions are sufficiently mild and enable the cross-coupling of 3-Br-*o*-carborane (**Br-oCB**, see SI for details) with amine and alcohol substrates that are not strongly nucleophilic (5a-5b, Figure 5). Using 3-Br-*o*-carborane in this case is preferred, given its higher conversion efficiency and ease of preparation compared to the 3-I-*o*-carborane analogue.



Figure 5. Alkoxylation and amination of *ortho*-carboranes using **BroCB** (ellipsoids at 50% probability and H atoms omitted for clarity). GC-MS yields, and isolated yields in parentheses.

In summary, we discovered that B-bromo-m-carboranes undergo efficient Pd-catalyzed B-N, B-O, and B-CN crosscoupling enabled by precatalysts featuring electron-rich biaryl phosphine ligands. The higher reactivity of Br-mCB likely stems from faster transmetalation (Figure 1B, step II) due to a weaker Pd-Br bond compared to Pd-I congener. This is consistent with previously observed trends in palladiumcatalyzed transformations using aryl halide electrophiles and Pd-based catalysts supported by bulky electron-rich phosphine ligands.^{13,14} The use of B-bromo-carboranes allows direct access to previously unknown B-functionalizations of these clusters. In addition, judicious use of Pd-catalyst systems with either iodo- or bromo-functionalized carborane was used to access unprecedented heterofunctionalized species. This approach is also amenable to o-carborane, which is the most challenging carborane substrate. Notably, this cross-coupling chemistry is complementary to the recently developed efforts in directed B-H functionalization strategies¹⁵ and, if successfully combined, may provide unprecedented densely functionalized carborane species.¹⁶ Further expansion of this methodology to other cross-coupling chemistry¹⁷ along with a full mechanistic investigation¹⁸ is currently underway in our laboratory.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.6b05505.

Full procedures and other characterization data (PDF) Crystallographic data (CIF)

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Notes

The authors declare no competing financial interest.

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