Metal-Catalyzed and Metal-Free Nucleophilic

Substitution of 7-I-B₁₈H₂₁

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ABSTRACT

In this work, two pathways of reactivity are investigated to generate site-specific

substitutions at the B7 vertex of the luminescent boron cluster, anti-B₁₈H₂₂. First, a palladium-

catalyzed cross-coupling reaction utilizing the precursor 7-I-B₁₈H₂₁ and a series of model

nucleophiles was developed, ultimately producing several B-N and B-O substituted species.

Interestingly, the B-I bond in this cluster can also be substituted in an uncatalyzed fashion, leading to the formation of various B-N, B-O, and B-S products. This work highlights intricate differences corresponding to these two reaction pathways and analyzes the role of solvents and additives on the product distributions. As a result of our synthetic studies, seven new B₁₈-based clusters were synthesized, isolated, and characterized by mass spectrometry and Nuclear Magnetic Resonance (NMR) spectroscopy. The photoluminescent properties of two structurally similar ether and thioether products were further investigated, with both exhibiting blue fluorescence in solution at 298 K and long-lived green or yellow phosphorescence at 77 K. Overall, this work shows for the first time the ability to perform substitution of a boron-halogen bond with nucleophiles in a B₁₈-based cluster, resulting in the formation of photoluminescent molecules.

INTRODUCTION

Boron-halogen (B-X) bond substitution has proven to be a general and versatile transformation that permits precise placement of various substituents on different polyhedral boron cages. This reactivity has led to broad accessibility of new and interesting molecular architectures, particularly for *closo*-boranes such as carboranes and dodecaborate-based clusters. B-X bond substitution in these species is usually achieved through metal-mediated means, such as cross-coupling, that can procure B-C, B-N, B-O, B-P, and B-S bonds. Other methods include light-promoted homolytic cleavage of the boron-halogen bond leading to the formation of a B-centered radical, which can then undergo further substitution reaction. Another strategy for activating a boron-halogen bond centers on the oxidation of the halogen, resulting in the formation of an iodonium substituent that permits a more facile nucleophilic attack at the boron. Interestingly, in contrast to the classical alkyl C-X bonds, nucleophilic substitution of B-X cluster bonds is

generally intractable, with only a handful of reports to date.⁶ This can be generally ascribed to the increased strength of B-X bonds and the steric bulk of the cage that prevents nucleophilic attack on the side opposite the halide.⁷

Anti-B₁₈H₂₂ is a boron cluster which has gained recent attention as an inherently fluorescent boron hydride (λ_{em} = 410 nm, Φ = 0.97) that has potential for various photoluminescence applications.⁸ In order to tune the luminescent properties of the parent compound, selective derivatization methods for this cluster are needed. While there exist a number of routes that can place a halogen moiety onto various vertices of this boron hydride, surprisingly, there have been no reports so far exploring the post-functionalization of these boron-halogen bonds. Unlike closobased clusters for which boron-halogen substitution chemistry has been established, the open-face polyhedral cage configuration of anti-B₁₈H₂₂ presents several distinct synthetic challenges, including the presence of acidic bridging hydrogens (pK_a = 2.7 and 7.5)⁹ and reduced stability profile (Figure 1A). 8h In this work, we report our initial assessment for the feasibility of substituting a boron-iodine bond at the B7 vertex of the anti-B₁₈H₂₂ cluster. Specifically, we report the discovery of metal-catalyzed and metal-free pathways capable of producing B-N, B-O, and B-S bonds, ultimately leading to new B₁₈-based chromophores with tunable photoluminescent properties. Overall, this work represents the first example of B-X bond substitution in anti-B₁₈H₂₂ and provides a potential roadmap toward the development of a broad scope of B₁₈-based luminescent materials.

RESULTS AND DISCUSSION

To examine the feasibility of metal-catalyzed cross-coupling for halogenated $anti-B_{18}H_{22}$ clusters, we decided to employ the Buchwald-Hartwig amidation conditions previously reported

for B-iodo-carboranes. ¹⁰ Monoiodinated 7-I-B₁₈H₂₁ (1) was selected as the borane precursor due to its straightforward and efficient synthesis, with 67% and 75% isolated yield in its initial report and in this study, respectively. 8c Additionally, 2,2,2-trifluoroacetamide (I) was used as a coupling partner due to its inability to undergo β-hydride elimination and its low pK_a resulting in facile deprotonation during the transmetallation step. 11 The reaction progress and products formed were monitored in situ by electrospray ionization mass spectrometry in the negative mode (ESI(-)MS). While the initially selected conditions did not result in significant conversion of the starting material as observed by ESI(-)MS (Figure 1C, Entry 1), a mass consistent with product 2 was observed when the ligand and catalyst loading were increased from 10 mol% and 5 mol% to 50 mol% and 25 mol%, respectively (Figure 1C, Entry 2). However, degradation to a non-cluster species was also indicated by a significant amount of a water-soluble solid that manifested as a sharp singlet at ~13 ppm in the ¹¹B NMR spectrum. The formation of this unidentified byproduct was observed to a lesser extent when 1,4-dioxane was used as a solvent, however only B₁₈H₂₂ was observed by ESI(-)MS, suggesting a competing reduction pathway (Figure 1C, Entry 3). We previously employed palladium-based precatalysts in combination with biaryl phosphine ligands, which proved superior in cross-coupling efficiency for B-halo-carboranes compared to the conventional Pd₂dba₃ (tris(dibenzylideneacetone)dipalladium(0)) precursor (Figure 1B).¹² Improved conversion to product 2 in 1,4-dioxane was thereby achieved with a RuPhos/RuPhos Pd G4 precatalyst system (L2/[L2Pd], Figure 1B). Further optimization revealed that L2/[L2Pd] loading could be reduced to 25 mol%/10 mol% while maintaining nearly complete conversion (Figure 1C, Entry 4). With these optimized conditions we were able to purify the product mixture and isolate 2 as an analytically pure solid in 24% yield. The significant yield reduction is attributed to both the competing reduction pathway that produces B₁₈H₂₂ and the acidified silica gel column

chromatography conditions, which results in loss of the product material. 6c,13 Nevertheless, the reaction conditions also proved effective with a K'BuO substrate (II) to yield the monosubstituted boron cluster product, 3 (Figure 1C, Entry 5). This B-O cross-coupling can also be conducted in a microwave reactor at 100°C for 45 minutes (Figure 1C, Entry 6). In addition to appending the alkyl 'BuO substituent, we found that this reaction is suitable for the bulky aromatic substrate potassium 2,6-dimethylphenolate to yield 4. The metal-mediated nature of these reactions is supported by Figure S1.

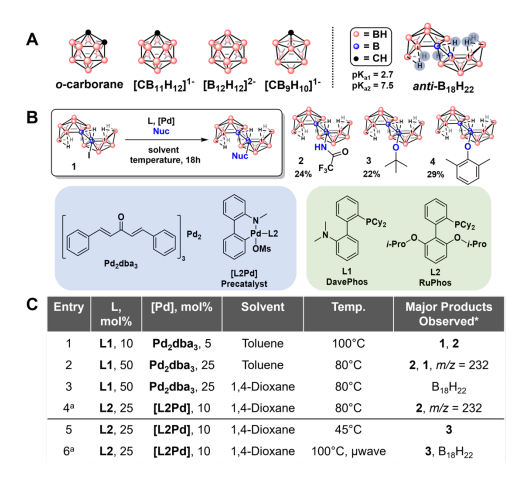


Figure 1. A) Select *closo*-boranes that can undergo metal-catalyzed cross-coupling vs. the *nido anti*-B₁₈H₂₂ cluster. B) General cross-coupling conditions, ligands and catalysts used, and resulting products **2-4** with isolated yields. C) Screening conditions employed to produce products **2** and **3**. See SI for general screening conditions procedure. ^aDone on a 30mg scale. *In order of decreasing intensity by ESI(-)MS.

With the isolated B-N and B-O products in hand, we turned to heteronuclear NMR spectroscopy for further characterization of compounds 2-4; each were subject to ¹³C, ¹H{¹¹B}, ¹¹B NMR and ¹H{¹¹B}-¹¹B{¹H} Heteronuclear Multiple Quantum Coherence (HMQC) analysis (Figures S9-13). Taking 3 as an example, the presence of the *tert*-butyl group is supported by characteristic resonances in the ¹³C NMR spectrum (-29.95 ppm and-78.24 ppm) and the singlet at 1.48 ppm in the ¹H{¹¹B} NMR spectrum. Also present in the ¹H{¹¹B} NMR spectrum are resonances corresponding to the 15 terminal and 6 bridging hydrogen nuclei on the boron cage, which is consistent with a monosubstituted product. The 18 boron nuclei are represented in the ¹¹B{¹H} NMR spectrum, and analysis of the corresponding ¹¹B NMR spectrum shows that all but three of these resonances exhibit doublet splitting, indicating they are bound to a terminal hydrogen. The three singlets at 8.3, 0.7, and -1.1 ppm should correspond to vertices B5, B6, and the newly substituted B7 (see Figure 2 for the numbering scheme). Integration of the boron resonances indicate that while 18 boron nuclei are present, several share similar chemical shifts, which makes definitive assignment of the boron resonances between 5 and -5 ppm more challenging. Therefore, a HMQC study of the ¹¹B{¹H} and ¹H{¹¹B} NMR spectra was conducted to assign the boron resonances and to confirm the overall structure of the molecule (Figure 2). The bridging hydrogen resonances in the HMQC spectrum provide an ideal starting point for the analysis of the 2D spectrum because they correspond to two separate boron nuclei. For example, the identification of the B8-B9 bridging hydrogen H8-9 resonance at -2.90 ppm^{10c} leads to the assignment of the resonances at -3.1 ppm and 2.6 ppm to the boron nuclei B8 and B9, respectively. In addition, the B9 resonance corresponds to a proton resonance at -0.26 ppm, which then correlates to two different boron resonances. This must signify the B9-B10 bridging hydrogen, which in turn leads to the assignment of the resonance at 9.6 ppm to the B10 nucleus. Continuing this process leads to

the identification of the bridging hydrogens H5-7' and H6-7 *via* their resonances at -0.66 and 0.98 ppm, respectively. From this, one can assign the B5 (0.7 ppm), B6 (8.3 ppm), and B7 (-1.1 ppm) nuclei to the initially observed singlets in the ¹¹B NMR spectrum. The significant downfield shift of the B7 resonance compared to the B7' resonance and the absence of a B7 terminal hydrogen resonance is consistent with the B7-substituted nature of the cluster.

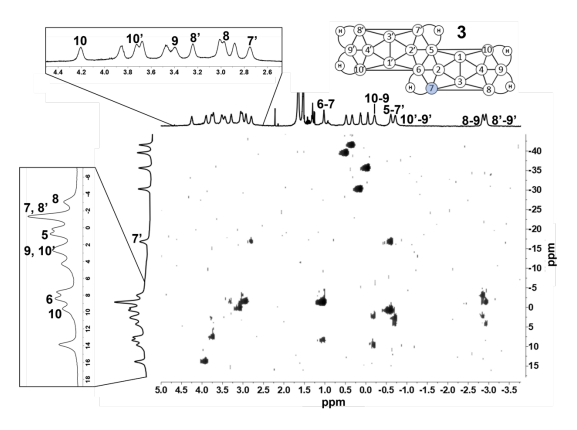


Figure 2. ¹H{¹¹B}-¹¹B{¹H} HMQC of **3** with select boron and hydrogen resonances labeled according to the numbering scheme at the top (substituted B7 vertex is highlighted in blue).

As we turned our attention to expanding the scope of cross-coupling to forge B-S bonds, we noticed product masses in the ESI(-)MS of the crude reaction mixtures that were not observed in previous reactions. For example, when the cross-coupling conditions were applied to potassium 2,6-dimethylthiophenolate (III), a mass corresponding to the desired B-S substituted cluster product plus a dioxane solvent molecule (m/z = 440.38) was the major species by ESI(-)MS.

Similar results were observed when the reaction was conducted in THF (m/z = 424.38) (Figure S20). Upon isolation of this THF-containing compound, analysis by 1D and 2D 11 B NMR spectroscopy indicated a monosubstituted cluster pattern, with the functionalized B7 resonance at $^{-3.83}$ ppm. The 1 H $\{^{11}$ B $\}$ NMR spectrum contained resonances corresponding to 21 cluster hydrogens ($^{-3}$ to 5 ppm) and the thioether group (7.11 and 2.57 ppm), but the distinctive THF resonances at 1.85 and 3.76 ppm were not observed. Instead, triplets at 3.97 and 2.73 ppm and multiplets at 1.82 and 1.71 ppm were present, suggesting the presence of a linear alkyl ether motif rather than an intact ethereal ring (Figure S21). This observation was further supported by 13 C and 13 C APT NMR spectroscopic experiments (Figures S24-26), which depict carbon resonances for both the thioether (143.20, 133.78, 128.25, 128.23, 22.8 ppm) and alkyl ether (70.33, 35.21, 30.49, and 36.56 ppm). In total, these data led us to conclude that the compound consisted of a B7-O bound, ring-opened THF molecule that resides between the electrophilic boron cluster and nucleophilic substrate (5).

To our surprise, further control experiments that excluded the palladium-based precatalyst and ligand from the reaction led to significant formation of **5** when the mixture was analyzed by ESI(-)MS *in situ*. This observation ultimately suggested that nucleophilic substitution of 7-I-B₁₈H₂₁ can potentially proceed *via* a separate mechanism that does not involve oxidative addition of Pd(0) into a B-I bond. This is significant considering that a metal-free route toward B₁₈H₂₂ functionalization could alleviate some of the limitations observed in the metal-catalyzed reactions, specifically the competing reduction process that produces B₁₈H₂₂ and the slow reductive elimination step that necessitates sterically bulky substrates. ¹⁶ To explore this opportunity further, we examined the conditions under which products similar to **5** could form for various substrates (Figure 3A). When individual substrates **I-V** were combined with **1** in THF at 80-100°C, the ESI(-

)MS spectrum of each reaction mixture exhibited only the mass corresponding to its respective alkyl ether product derived from THF ring-opening. The only exception was the K'BuO (II) reaction mixture, which also contained a mass corresponding to the direct nucleophilic substitution product 3. To determine if interconversion between these two compounds was possible, the reaction mixture was further refluxed for 18 hours but showed no change in the product distribution as observed by ESI(-)MS. This finding led us to consider two possible pathways of reactivity for this reaction (Figure 3B). The Lewis acidic B₁₈-based cluster can interact with THF^{10h} to weaken the B-I bond, providing sufficiently nucleophilic substrates such as II the opportunity to attack the boron atom (Figure 3B(a)). Weaker nucleophiles, however, will react at the THF carbon to yield the ring-opened product (Figure 3B(b)). To probe the conditions under which each pathway could be potentially favored, we reacted 1 with II in THF at room temperature. This provided nearly full conversion to 3 as observed by ESI(-)MS, suggesting that the ring-opened product is favored at elevated temperatures. These results, however, were exclusive to 3; only the THF-derived products were observed by ESI(-)MS for the other substrates, even when the reactions were conducted at -78°C. While sulfur-based reagents III and IV are more nucleophilic than their oxygen counterparts, the thermodynamic preference for B-O bond formation compared to B-S congener renders the formation of products 6 and 7 less favorable than the corresponding THF-derived products. ¹⁷ In an effort to overcome this, we found that incorporation of 18-crown-6 in the reaction mixture increases conversion to products 3 and 7 as observed by ESI(-) MS, possibly due to the increased Lewis basicity of the substrate caused by sequestration of the cation.¹⁷ The remaining substrates I, III, and V however, still required a more inert solvent to avoid the ring-opening pathway altogether (Figure 3A, pink). As such, when I, III, and V are reacted with 1 in toluene there is no conversion observed by ESI(-)MS, yet including 18-crown-6 in the reaction mixture

ultimately produces the desired compounds **2**, **6**, and **8**. Overall, the ability for substrates **I**, **III** and **V** to undergo nucleophilic substitution is aided not only by the crown ether but also toluene, which has a destabilizing effect on the nucleophile by solvating it to a lesser degree than THF. Compounds **2**, **3**, and **5-8** can be easily isolated from the corresponding reaction mixtures. Structural formulations of these products were confirmed by ESI(-)MS and ¹H{¹¹B}, ¹¹B{¹H}, ¹¹B, ¹³C, ¹⁹F, HMQC NMR spectroscopy. Interestingly, neither the reactions in THF nor toluene led to significant formation of **4**, potentially due to its weak nucleophilicity. Cross-coupling therefore remains the most viable route for producing **4** and other structures unsuitable for nucleophilic substitution. Still, the ability for 7-I-B₁₈H₂₁ to undergo metal-free nucleophilic substitution at the B-I bond is remarkable given the comparatively inert nature of boron-halogen bonds in other polyhedral borane clusters.

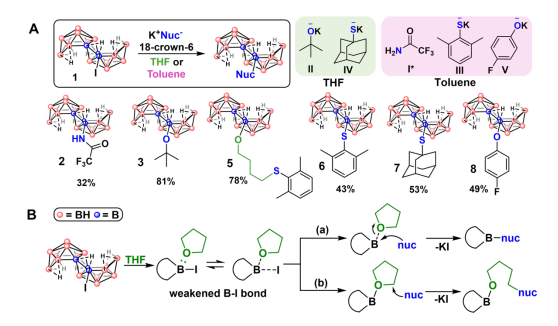


Figure 3. A) Reaction for nucleophilic substitution, substrates employed, and resulting products. Substrates can undergo substitution in THF (green) or toluene (pink). B) Pathways of reactivity for nucleophiles reacting with **1** in THF solution. *K₃PO₄ is included in the reaction mixture.

Given that this is the first report of ether and thioether-functionalized anti-B₁₈H₂₂, we wanted to benchmark the luminescent properties of these new derivatives. Specifically, the structurally similar compounds 4 and 6 were assessed to gauge the effects of oxygen vs. sulfur on cluster luminescence (Figure 4). In cyclohexane solution, the absorption of both compounds is largely in the UV region, with ($\varepsilon_{346} = 6600 \,\mathrm{M}^{-1}\mathrm{cm}^{-1}$ for 4 and $\varepsilon_{392} = 3300 \,\mathrm{M}^{-1}\mathrm{cm}^{-1}$ for 6). Emission spectra of these solutions show blue fluorescence at 427 nm and 460 nm in 4 and 6, respectively (Figure 4, S50, S52). Notably, this emission was exceptionally weak, with quantum yields (Φ) less than 0.01. However, when the measurements were repeated at 77 K in methylcyclohexane the luminescence was significantly shifted to the red ($\lambda_{em} = 530$ nm for 4 and $\lambda_{em} = 572$ nm for 6). The measured emission lifetimes (τ) at these wavelengths show long-lived, phosphorescent processes, with $\tau = 360$ ms for 4 and $\tau = 7.7$ ms for 6. (Figures S51, S53). Phosphorescence is typically seen in molecules containing heavy atoms, which lend the molecule spin-orbit coupling (SOC) needed to increase the efficiency of intersystem crossing (ISC) from the singlet excited state to the triplet excited state. Based on the weak fluorescence in these two compounds, it appears that SOC arising from oxygen and sulfur leads to efficient ISC. This phenomenon is consistent with TD-DFT calculations of 4 and 6, in which the valence orbitals participating in lowest electronic transitions (Figures S54-55) support El-Sayed's rules for efficient ISC. 19 The extraordinarily long phosphorescent lifetimes in 4 and 6 provide ample opportunity for competitive non-radiative decay to occur (e.g., through molecular motion), ultimately resulting in minimal luminescence at ambient conditions. At 77 K, however, the molecules are constrained in a frozen matrix that inhibits these non-radiative decay processes and thus permits strong phosphorescence. This data highlights the varying effects that substituents can have on the photophysical properties

of *anti*-B₁₈H₂₂ and shows that B-O and B-S connectivity alters the absorption, emission, and fluorescent behavior of the corresponding B₁₈-based cluster.

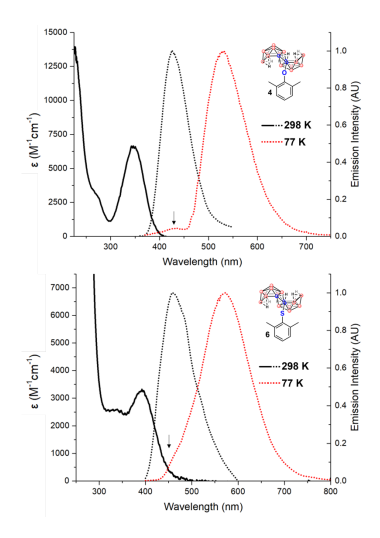


Figure 4. Absorption (solid line) and emission (dotted line) spectra of compounds **4** (top) and **6** (bottom). Emission spectra at 298 K and 77 K display fluorescence and phosphorescence, respectively. The arrows indicate fluorescent shoulders visible in the 77 K spectra. For **4**, $\lambda_{exc} = 340$ nm at 298 K, $\lambda_{exc} = 350$ nm at 77 K. For **6**, $\lambda_{exc} = 380$ nm.

In this work we detail two new and complementary strategies for generating nitrogen, oxygen, and sulfur-functionalized *anti*-B₁₈H₂₂ analogues from 7-I-B₁₈H₂₁. We first demonstrate that palladium-catalyzed cross-coupling can forge B-N and B-O bonds at the B7 vertex with sterically hindered substrates. To our surprise, we discovered that these connectivities could be produced through a metal-free nucleophilic substitution pathway, which also proved applicable for generating B-S functionalized clusters. In addition to unveiling this unique reactivity of monoiodinated B₁₈H₂₂, we demonstrate that the luminescence of the parent borane can be modulated through cage functionalization. Aside from a rare example of a metal-free nucleophilic substitution chemistry of a boron-halogen moiety in a boron cluster, this work broadens the scope of luminescent boron cluster-based molecules.¹⁸

ASSOCIATED CONTENT

Supporting Information.

The Supporting Information is available free of charge.

Detailed experimental and analytical procedures, characterization data, and DFT calculations (PDF)

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The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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TOC:



Synopsis: Nucleophilic substitution of the halogen moiety at 7-I- $B_{18}H_{21}$ cluster leading to a series of new B-N, B-O and B-S substituted clusters is reported. This work highlights several independent mechanistic pathways responsible for the B-I bond breaking events and showcases how the discovered substitution chemistry can affect the luminescent properties of the newly formed cluster products.