Lewis Acids

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Tris(*ortho*-carboranyl)borane: An Isolable, Halogen-Free, Lewis Superacid

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Dedicated in memory of the late Stephen A. Westcott, a great friend and mentor

Abstract: The synthesis of tris(*ortho*-carboranyl)borane (BoCb₃), a single site neutral Lewis superacid, in one pot from commercially available materials is achieved. The high fluoride ion affinity (FIA) confirms its classification as a Lewis superacid and the Gutmann-Beckett method as well as adducts with Lewis bases indicate stronger Lewis acidity over the widely used fluorinated aryl boranes. The electron withdrawing effect of *ortho*-carborane and lack of pi-delocalization of the LUMO rationalize the unusually high Lewis acidity. Catalytic studies indicate that BoCb₃ is a superior catalyst for promoting C–F bond functionalization reactions than tris(pentafluorophenyl)borane [B(C_6F_5)₃].

Boranes are useful Lewis acids in stoichiometric and catalytic reactions by taking advantage of the vacant porbital.^[1] Boron trihalides (BX₃; X=F, Cl, Br) are ubiquitous examples; however, their volatile nature and labile B-X bonds make them incompatible with many substrates that limits their utility. In this regard. tris(pentafluorophenyl)borane [B(C₆F₅)₃, Figure 1], became a standout borane due to excellent thermal stability and functional group tolerance which led to its widespread use as a Lewis acid reagent and catalyst.^[2] Although $B(C_6F_5)_3$ is a useful Lewis acid, its' fluoride ion affinity (FIA) is less than the defined benchmark Lewis superacid (LSA), SbF₅.^[3,4] In fact, reports on single-site LSA boranes are scarce and accessing new species with different acidity or steric profiles could lead to altered reactivity and selectivity

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Melbourne, Victoria 3086 (Australia) in stoichiometric or catalytic reactions. Although there have been efforts directed at isolating boron based Lewis superacids, most of the methods involve modifications on the substitution of the aryl group on triarylboranes.^[2b] This strategy was effective in achieving LSA for $B(p-CF_3-C_6F_4)_3$ while other fluoride loading approaches fell short of the LSA criterion.^[5] The increased FIA of $B(p-CF_3-C_6F_4)_3$ results in greater catalytic activity in C-H perfluoroalkylation and perfluoroarylation reactions compared to $B(C_6F_5)_{3.}^{[6]}$ Incorporating the boron center within an antiaromatic heterocycle results in compounds with high Lewis acidity based on Gutmann-Beckett acceptor numbers, or high FIAs, but their reactive B-C bonds limits application in synthesis.^{[7][8]} Fluorinated alkyloxy-boranes, boron pseudohalides $[B(OR^F)_3, R^F = -CF_3, -TeF_5, -SO_2CF_3]$, and perfluoroalkyl-boranes are known and believed to be LSAs; however, they are thermally unstable with the latter only generated as fleeting species.^[9,10] Chlorination approaches in triarylboranes have been extensively studied with little success.^[11] Gabbaï and co-workers applied a different approach, appending cationic substituents (trialkylammonium, sulfonium, and phosphonium) on aryl boranes as an alternative electron withdrawing group to generate a class of cationic Lewis acids with high FIAs.^[12] Recently, Berionni and co-workers achieved remarkable FIAs with rigid pyramidalized boranes tethered by carbon, phosphonium and sulfonium centers, although the free boranes were not isolable.^[13]

We postulated that using an unconventional electron withdrawing group with significant steric protection could result in the isolation of a new class of trigonal planar Lewis acids. In the literature, carboranes^[14] offer both of these attributes although they have been minimally explored in this realm. $^{\left[15,16\right] }$ The $C_{2}B_{10}$ carborane cluster is exceptionally stable and the three-dimensional icosahedron presents a sphere-like steric profile in stark contrast to flat aryl groups. Within the C_2B_{10} carboranes, three isomers exist with each classified based on the relative positioning of the carbon atoms, ortho- (adjacent), meta- (one boron betwixt), and para- (carbon atoms on opposite sides of the icosahedron). Among these, the ortho-isomer (oCbH) has been documented as the most electron withdrawing if C-bound. This hypothesis is supported by a report of borole analogues featuring a bis(ortho-carborane) backbone in which the ocarborane engendered significantly higher Lewis acidity than the biphenyl variant.^[17] A notable advancement in this realm was the preparation of anthracene analogues by Ye

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Research Articles



Figure 1. Tris (pentafluorophenyl) borane, isolable single-site neutral boron Lewis superacids (LSAs) in the literature and feature complex, tris (*ortho*-carboranyl) borane (oCb = ortho-carborane, C₂B₁₀H₁₁).

and co-workers in which the boron centers are bridging two *ortho*-carboranes.^[18] In these species, LSA is achieved but the halides or azides present on the boron center are reactive and these fall into the category of dual site LSAs.^[19] In contrast to aryl or fluoroaryl boranes, the carborane cluster is likely unable to delocalize the LUMO, primarily a p-orbital on boron. We herein report the synthesis of BoCb₃ in one pot which represents an isolable, halogen-free,



Scheme 1. Synthesis of BoCb₃. [Isolation: After the reaction was complete, additional toluene was added and the reaction mixture filtered through a small pad of celite and washed with dichloromethane. The volatiles were stripped from the combined filtrate under vacuum and diethyl ether was added to make a suspension, which was filtered to give pure BoCb₃ as white solid.]



Figure 2. Solid state structure of BoCb₃. Ellipsoids depicted at the 50%

probability level and hydrogen atoms are omitted for clarity. Selected

bond lengths (Å) and angles (°): B(1)–C(1) 1.614(8), B(1)–C(3) 1.627(7), B(1)–C(5) 1.626(8); C(1)–B(1)–C(3) 119.8(5), C(1)–B(1)–C-

neutral, single-site Lewis superacid; and investigate its properties.

The lithiation of ortho-carborane with nBuLi and subsequent reaction with 0.33 equivalents of BCl₃ (1 M solution in hexanes) generated BoCb₃ in 29% yield after a work up (Scheme 1). When BBr3 was used instead of BCl3, an increased isolated yield of 35% was achieved. The structure was confirmed by a single-crystal X-ray diffraction study (Figure 2).^[20] The geometry is trigonal planar as all C-B-C bond angles are within error of 120° and the C-B bond lengths are slightly longer than the typical C(ocaroborane)-B single bond in boranes species [1.614(8)-1.627(7) Å c.f. \approx 1.58 Å], which is attributed to the bulk around the central B-atom. The downfield ¹¹B{¹H} resonance at 67.2 ppm is assigned to the central tricoordinate boron atom and peaks ranging from 7.4 to -12.4 ppm to the cluster atoms. The C–H protons are the most diagnostic in the ¹H NMR spectrum and appear as a singlet at 5.02 ppm while the corresponding carbon is observed in the ¹³C{¹H} NMR spectrum at 65.0 ppm and a broad peak at 69.3 ppm is assigned to the carbon bound to the central boron. Analytical purity was confirmed by microanalysis and the melting point exceeds 250 °C, indicating thermal stability.

To experimentally investigate the relative Lewis acidity, the corresponding acetonitrile $[CH_3CN \cdot BoCb_3]$ and benzaldehyde $[PhCHO \cdot BoCb_3]$ adducts were prepared (Scheme 2). In the ¹¹B NMR spectra, the peak for the central boron atom (67.2 ppm) shifts to the tetracoordinate region among the cluster boron peaks. The C–H resonance of $BoCb_3$ (5.02 ppm) in the ¹H NMR spectrum shifts upfield to 4.60, and 4.77 for CH₃CN $\cdot BoCb_3$ and PhCHO $\cdot BoCb_3$,



Scheme 2. Synthesis of nitrile- $[CH_3CN \cdot BoCb_3]$ and aldehyde- $[PhCHO \cdot BoCb_3]$ adducts of $BoCb_3$.

Angew. Chem. Int. Ed. 2022, 61, e202212073 (2 of 7)

C(5) 120.3(5), C(3)-B(1)-C(5) 119.9 (5).

respectively. Their identities were further confirmed by single crystal X-ray diffraction experiments (Figure 3) with the central boron atoms adopting distorted tetrahedral geometries. The shorter B-donor atom (O, and N) bonds in the BoCb₃-adducts compared to the corresponding $B(C_6F_5)_3$ -complexes indicate higher Lewis acidity of $BoCb_3$ $[CH_3CN \cdot BoCb_3: B-N 1.5679(17) \text{ Å } c.f. CH_3CN \cdot B(C_6F_5)_3:$ B-N 1.616(3) Å; PhCHO·BoCb₃: B-O 1.551(2) Å, c.f. PhCHO·B(C_6F_5)₃: B–O 1.6108(8) Å].^[21] Although the structure of the PhCHO \cdot BoCb₃ was confirmed in the solid state, we observed a dissociation equilibrium in solution which could not be resolved in the ¹H NMR spectrum at 25 °C. The ¹H NMR spectra of BoCb₃ with gradient PhCHO concentrations at 25 °C and variable temperature studies of 1:1 and 1:1.4 molar mixtures of BoCb₃ and PhCHO confirm the dissociation equilibria (see Supporting Information Figures S21-S23). The CN stretching frequency is often used as a parameter to evaluate Lewis acidity. However, the CN stretching frequencies of $CH_3CN \cdot BoCb_3$ (2363 cm⁻¹) and $CH_3CN \cdot B(C_6F_5)_3$ (2367 cm⁻¹) are within experimental error. To find the relative Lewis acidity between BoCb₃ versus $B(C_6F_5)_3$, a competition study was performed to assess the relative binding affinity of acetonitrile. When equimolar $BoCb_3$, $B(C_6F_5)_3$, and CH_3CN were mixed in $CDCl_3$, $CH_3CN \cdot BoCb_3$ and $CH_3CN \cdot B(C_6F_5)_3$ were observed in a $\approx\!2\!:\!1$ ratio indicating higher CH_3CN affinity of BoCb3 over $B(C_6F_5)_3\!\!\!\!^{[22]}$

The Gutmann-Beckett method was applied to evaluate the Lewis acidity of BoCb₃ by synthesizing Et₃PO·BoCb₃ (Table 1).^[23] The difference in ³¹P chemical shift ($\Delta\delta^1$) of Et₃PO·BoCb₃ and Et₃PO is 34.1 ppm, 27.5 ppm, and 30.1 ppm in C₆D₆, CDCl₃, and CD₂Cl₂, respectively (Table 1).^[24] These differences in chemical shifts are higher than the results obtained for B(C₆F₅)₃ ($\Delta\delta^2$ =29.7 ppm in C₆D₆, 23.5 ppm in CDCl₃ and 26.1 ppm in CD₂Cl₂) and those reported for B(*p*-CF₃-C₆F₄)₃ ($\Delta\delta^2$ =31.9 ppm in C₆D₆ and $\Delta\delta^2$ =29.0 ppm in CD₂Cl₂).^[5] This indicates that BoCb₃ is more Lewis acidic than B(C₆F₅)₃ and B(*p*-CF₃-C₆F₄)₃ based on the Gutmann-Beckett method. Also, the shorter B–O bond in Et₃PO·BoCb₃ [B–O 1.500(2) Å] over Et₃PO· B(C₆F₅)₃ [B–O 1.533(3) Å] imply a stronger bond in the BoCb₃ adduct (Figure 3).^[25]

To support the experimental observations on the high Lewis acidity of B_0Cb_3 we undertook a variety of theoretical analyses against benchmark strong Lewis acidic boranes, specifically $B(p-CF_3-C_6F_4)_3$ and $B(C_6F_5)_3$ (Table 2). Considered parameters include fluoride ion affinity (FIA), hydride ion affinity (HIA), binding energy to the Lewis bases NH₃ and CH₃CN, Natural Population Analysis Charges, relative LUMO energies, and finally, the Global Electrophilicity



Figure 3. Solid state structure of adducts CH₃CN·BoCb₃, PhCHO·BoCb₃, and Et₃PO·BoCb₃ (left to right). Ellipsoids depicted at the 50% probability level, hydrogen atoms, and solvates are omitted for clarity. Selected bond lengths (Å) and angles (°) of CH₃CN·BoCb₃: B(1)–N(1) 1.5679(17), N(1)–C(7) 1.1380(18), B(1)–C(1) 1.714(2), B(1)–C(3) 1.7118(19), B(1)–C(5) 1.7143(19), B(1)–N(1)–C(7) 176.44(13); PhCHO·BoCb₃: B(1)–O(1) 1.551(2), O(1)–C(7) 1.254(3), B(1)–C(1) 1.714(3), B(1)–C(3) 1.718(3), B(1)–C(5) 1.706(3); B(1)–O(1)–C(7) 134.57(18), O(1)–C(7)–C(8) 123.2(2); Et₃PO·BoCb₃: B(1)–O(1) 1.500(2), O(1)–P(1) 1.5344(11), B(1)–C(1) 1.730(2), B(1)–C(3) 1.732(2), B(1)–C(5) 1.730(2); B(1)–O(1)–P(1) 177.85(11).

Table 1: Determination of the Lewis a	cidity of BoCb ₃ using t	he Gutmann–Beckett method. ^{[ª}
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		E	$BR_{3} \xrightarrow{1.00 \text{ equiv } Et_{3}PO} Et_{3}PO \cdot BR_{3}$ $C_{6}D_{6}/CDCl_{3}, 23 \text{ °C} \qquad Et_{3}PO \cdot BR_{3}$			
			$R = oCb, C_6F_5$			
Solv.	Et₃PO [δ ³¹ P]	$Et_3PO \cdot BoCb_3$ [$\delta^{31}P$]	$\begin{array}{c} Et_3PO\cdotB(C_6F_5)_3\\ [\delta^{31}P] \end{array}$	$Et_3PO \cdot BoCb_3 - Et_3PO$ $[\Delta \delta^1]$	$Et_3PO \cdot B(C_6F_5)_3 - Et_3PO$ $[\Delta \delta^2]$	
C ₆ D ₆	45.7	79.6	75.4	34.1	29.7	
CDCl ₃ CD ₂ Cl ₂	52.3 51.0	79.8 81.1	75.8 77.1	27.5 30.1	23.5 26.1	

Angew. Chem. Int. Ed. 2022, 61, e202212073 (3 of 7)

able 2: Calculated thermochemical and electronic values	(FIA=fluoride ion affinity	, HIA=hydride ion affinity,	GEI = global electrophilicity index).
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Compound	FIA (Krossing) [Δ <i>H</i> , kJ mol ⁻¹]	HIA (Krossing) [Δ <i>H</i> , kJ mol ⁻¹]	NH₃ affinity [ΔG, kJ mol ^{−1}]	CH₃CN affinity [∆G, kJ mol ^{−1}]	LUMO [eV]	GEI
BoCb ₃	605	622	149	89	-3.99	4.22
$B(C_6F_5)_3$	452	484	97	33	-3.50	3.78
$B(p-CF_3-C_6F_4)_3$	501	537	110	51	-4.09	4.79
SbF₅	493	-	-	-	-	-
BF ₃	-	299	-	-	-	-

Index (GEI) recently reported by Stephan and co-workers. $\ensuremath{^{[26]}}$

Calculations for gas phase fluoride and hydride affinities are beset with issues in obtaining values consistent with experiment due to the small size and hardness of the anions and also the lack of a consistent referencing system. Krossing and co-workers presented an isodesmic reaction benchmarked to accurate fluoride and hydride affinities for $[(CH_3)_3Si]^+$ combined with inexpensive calculations using BP86/SVP for the single point calculations for Lewis acids.^[27] Gas phase values for direct addition of $F^{\scriptscriptstyle -}$ and $H^{\scriptscriptstyle -}$ using B3LYP-D3/def2SVP are presented in the Supporting Information. Antimony pentafluoride has been established as the threshold LSA, with a calculated fluoride ion affinity (FIA) using Krossing's method of 493 kJ mol^{-1.[27]} Using Krossing's method benchmarked to their result for $B(C_6F_5)_3$ at 452 kJ mol⁻¹, we verified our calculations to reproduce their result for SbF₅ at 493 kJ mol⁻¹. The calculated FIA of $BoCb_3$ is 605 kJ mol⁻¹ and $B(p-CF_3-C_6F_4)_3$ is 501 kJ/mol, indicating that BoCb₃ has a significantly higher FIA than the other two boranes and SbF5, classifying it as a LSA. To validate the theoretically determined high FIA, BoCb₃ was reacted with [nBu₄N][SbF₆] aiming to abstract a fluoride ion from SbF₆⁻. The 1:1 reaction of BoCb₃ and [nBu₄N][SbF₆] at room temperature in CDCl₃ resulted in consumption of $BoCb_3$ and SbF_6^- within 30 min indicating fluoride ion abstraction based on ¹H, ¹⁹F{¹H}, and ¹¹B{¹H} NMR spectroscopy.^[22]

For hydride ion affinities, using Krossing's method, anchoring to their results for $B(C_6F_5)_3$ (543 kJ mol⁻¹) and BF_3 (299 kJ mol⁻¹), again $BoCb_3$ has the largest HIA of the considered Lewis acids at 622 kJ mol⁻¹. Calculating affinities for neutral Lewis bases (B3LYP-D3/def2SVP, gas phase), $BoCb_3$ also has a larger affinity with a ΔG for the dissociation of NH₃ being 149 kJ mol⁻¹, compared to 110 and 97 kJ mol⁻¹ for B(p-CF₃-C₆F₄)₃ and B(C₆F₅)₃, respectively. For acetonitrile, the ΔG for dissociation was calculated at $89 \text{ kJ} \text{mol}^{-1}$ for $BoCb_3$, $33 \text{ kJ} \text{mol}^{-1}$ for $B(C_6F_5)_3$, and 51 kJ mol⁻¹ for B(*p*-CF₃-C₆F₄)₃, qualitatively consistent with the experimental observation that BoCb₃ outcompetes $B(C_6F_5)_3$ for acetonitrile. For all affinity calculations, the results indicate BoCb₃ is the strongest Lewis acid, followed by $B(p-CF_3-C_6F_4)_3$, then $B(C_6F_5)_3$. However, the LUMO of $B(p-CF_3-C_6F_4)_3$, calculated using B3LYP-def2TZVP to give results directly comparable to the literature,^[26] is 0.1 eV lower in energy than BoCb₃, which also has a correspondingly larger Global Electrophilicity Index at 4.79 compared to 4.22 for BoCb₃. The discrepancy between the orbital energy difference in $B(p-CF_3-C_6F_4)_3$ and $BoCb_3$ versus the thermochemical Lewis acidity predictions are rationalized by a qualitative examination of the LUMOs themselves, depicted in Figure 4. The LUMO for BoCb₃ is entirely localized on the boron atom, whereas the LUMO for B(p- $CF_3-C_6F_4$ [and also $B(C_6F_5)_3$] is partially delocalized into the arene rings. Increased localization of the LUMO of BoCb₃ is also reflected in the Natural Population Charges on the boron centers, where the central boron BoCb₃ carries a positive charge of +1.83. The central boron atoms in B(p- CF_3 - C_6F_4)₃ and $B(C_6F_5)_3$ carry positive charges of +0.81 and +0.78, respectively, that is likely due to the phenyl rings ability to donate some electron density to the central atom.

The steric profiles and buried volume with respect to the boron centers for B_0Cb_3 and $B(C_6F_5)_3$ were compared using the SambVca 2.1 routine^[28] on the calculated optimized structures. The z-axis was defined as perpendicular to the BR₃ trigonal plane. It was found that the carborane substituents result in greater steric demand at B with a buried volume of 92.2 % compared to a buried volume of 83.5 % for $B(C_6F_5)_3$.^[22]



Figure 4. Depictions of the LUMO for $BoCb_3$, $B(C_6F_5)_3$, and $B(p-CF_3-C_6F_4)_3$ [from left to right, B3LYP/def2TZVP calculations]

Angew. Chem. Int. Ed. 2022, 61, e202212073 (4 of 7)

Next, we evaluated BoCb₃ as a Lewis acid in catalytic transformations. The abnormally high fluoride ion affinity encouraged us to explore it as a catalyst in C-F bond activation reactions.^[29] The catalytic functionalization of C-F bonds is challenging given that C-F is the strongest carbon single bond. To check the efficacy of BoCb₃, we attempted a model hydrodefluorination^[30] reaction of 1fluoroadamantane (1-F-Ad) and triethylsilane. To our delight, when one equivalent of 1-F-Ad was treated with 1.1 equivalents of HSiEt₃ in the presence of 0.5 mol% BoCb₃ in CDCl₃ at 23°C for 10 minutes, the reduction product, adamantane (Ad) was obtained in quantitative yield (>97 % yield by NMR spectroscopy) along with FSiEt₃ as the side product (Table 3, entry 1). Interestingly, when 0.5 mol% $B(C_6F_5)_3$ was employed (entry 2), adamantane (Ad), was only generated in 60% yield based on NMR spectroscopy, along with unreacted starting materials (Table 3, entry 1). Longer reaction times did not change the outcome. However, Stephan and co-workers showed that $B(C_6F_5)_3$ can undergo the same transformation but in a tenfold increase in catalyst loading (5 mol%).^[31] It is known that B(C₆F₅)₃ reacts with $HSiEt_3$ upon heating to give $HB(C_6F_5)_2$ (Piers' borane)^[32] but surprisingly, BoCb₃ is unreactive with HSiEt₃ ruling out the possibility of HBoCb₂ being the active catalyst.^[33] When BoCb₃ was reacted with 1-F-Ad, both starting materials were consumed resulting in a complex mixture based on ¹H, ¹⁹F{¹H}, and ¹¹B{¹H} NMR spectroscopy. This suggests that fluoride abstraction is the first step in the mechanism of the catalytic transformation.^[22]

Friedel–Crafts reactions of 1-F-Ad were also investigated. When 1-F-Ad was reacted with benzene in the presence of 1 mol% BoCb₃, 1-Ph-Ad was obtained in 90% isolated yield after 30 min (Table 3, entry 3). Interestingly, when $B(C_6F_5)_3$ was employed in the same transformation, we did not observe any product formation even after 4 h (entry 3 and 4). Moran and co-workers reported a $H_2O \cdot B(C_6F_5)_3$ -catalyzed Friedel–Crafts reaction of 1-F-Ad but highly electron rich arenes and specific solvent combinations were required to enable the reaction.^[34] The analogous transformation with toluene and *m*-xylene provides the corresponding products 1-Tol-Ad (entry 5) in 72 % isolated yield (20:3 *p*-:*m*- isomeric ratio), and 1-Xyl-Ad selectively (entry 6) in 92 % yield. The improved outcome in the reactions of $BoCb_3$ over $B(C_6F_5)_3$ is attributed to its higher FIA and Lewis acidity. The catalytic hydrosilylation reaction^[2e] of benzaldehyde with triethylsilane (1:1 PhCHO and HSiEt₃ in presence of 5 mol% $BoCb_3$ in CDCl₃) formed a mixture of the hydrosilylation product PhCH₂OSiEt₃ and dialkyl ether (PhCH₂)₂O in 52 % and 36 % yield by NMR spectroscopy at room temperature after 24 h.^[22]

In summary, the halogen-free thermally stable borane, tris(ortho-carboranyl)borane, is accessed from three convenient reagents: ortho-carborane, nBuLi, and BBr3. Calculated fluoride ion affinities reveal the Lewis acidity greatly exceeds SbF₅ making it a rare example of a Lewis superacidic borane. Theoretical binding studies comparing BoCb₃ to the state-of-the-art fluoroarylborane Lewis acids, $B(C_6F_5)_3$ and $B(p-CF_3-C_6F_4)_3$, reveal fluoride and hydride ion affinities as well as Lewis base affinities of ammonia and acetonitrile are all the highest for BoCb₃. Experimentally, a competition study in solution with acetonitrile and one equivalent of $BoCb_3$ and $B(C_6F_5)_3$ corroborate preference for binding BoCb₃. In the solid state, X-ray diffraction data from benzaldehyde, acetonitrile and triethylphosphine oxide adducts all validate stronger binding to BoCb₃ than B(C₆F₅)₃. In solution, Gutmann-Beckett studies indicate the Lewis acidity of BoCb₃ exceeds the literature values for $B(C_6F_5)_3$ and $B(p-CF_3-C_6F_4)_3$. The high FIA can be taken advantage of in catalytic C-F bond activation reactions of unactivated alkyl fluorides by reduction with triethylsilane

Table 3: Reactions of 1-fluoroadamantane (0.1 mmol) in the presence of borane catalysts.

	F Nu E-Nu "conditions") + EF	
	1-F-Ad 1-Nu-Ad	d	
	H Ad 1-Ph-Ad 1-Tol-Ad (p-:m- isomer)	C CH ₃ 1-Xyl-Ad	
Entry	"Conditions"	Product (1-Nu-Ad)	Yield
1	1.1 equiv HSiEt₃, 0.5 mol% BoCb₃, CDCl₃ (1 mL), 23 °C, 10 min	Ad	$> 97\%^{[a]}$
2	1.1 equiv HSiEt ₃ , 0.5 mol% B(C ₆ F ₅) ₃ , CDCl ₃ (1 mL), 23 °C, 30 min	Ad	60 % ^[a]
3	1 mol% BoCb ₃ , C ₆ H ₆ (2 mL), 23 °C, 30 min	1-Ph-Ad	90 % ^[b]
4	1 mol% B(C ₆ F ₅) ₃ , C ₆ H ₆ (2 mL), 23 °C, 4 h	1-Ph-Ad	
5	1 mol% BoCb ₃ , C ₇ H ₈ (2 mL), 23 °C, 30 min	1-Tol-Ad	72 % ^[b]
6	1 mol% BoCb ₃ , <i>m</i> -Xylene (2 mL), 23 °C, 30 min	1-Xyl-Ad	92 % ^[b]

[a] Yields are determined by NMR spectroscopy using 1-bromo-4-fluorobenzene as an internal standard. [b] Isolated yields. [c] NR=No reaction.

Angew. Chem. Int. Ed. 2022, 61, e202212073 (5 of 7)

and C–C bond forming Friedel–Crafts reactions with arenes. In this disclosure of B_0Cb_3 , we merely touch on the potential of this reagent and catalyst. Given the powerful Lewis acidity and unusual steric profile, it offers unique synthetic opportunities in boron mediated transformations.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

The data that support the findings of this study are available in the supplementary material of this article.

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