

B-H Functionalization of Hydrogen-Rich [(Cp*V)(2) (B2H6)(2)]: Synthesis and Structures of [(Cp*V)(2)(B2X2)(2)H-8] (X = CI, SePh; Cp* = eta(5)-C5Me5)

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B-H Functionalization of Hydrogen-Rich [(Cp*V)₂(B₂H₆)₂]: Synthesis and Structures of [(Cp*V)₂(B₂X₂)₂H₈] (X = Cl or SePh; Cp* = η^5 -C₅Me₅)

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ABSTRACT: We have recently reported the perchlorinated diniobaborane species $[(Cp*Nb)_2(B_2H_4Cl_2)_2]$ from $[(Cp*Nb)_2(B_2H_6)_2]$ using CCl₄ as chlorinating agent. In an attempt to isolate the vanadium analogue, we have isolated $[(Cp*V)_2(B_2H_6)_2]$, **1** from the reaction of $(Cp*VCl_2)_3$ with $[LiBH_4\cdot THF]$ followed by thermolysis with excess $[BH_3\cdot THF]$. Subsequently, the thermolysis of **1** with CCl₄ for a prolonged period of time afforded perchlorinated divanadaborane $[(Cp*V)_2(B_2H_4Cl_2)_2]$, **2** along with the formation of bichlorinated divanadaborane $[(Cp*V)_2(B_2H_4Cl_2)(B_2H_5Cl)]$, **4.** Similarly, in order to functionalize the terminal B-H by {SePh} group, thermolysis of **1** was carried out with Ph₂Se₂ that yielded persubstituted divanadaborane $[(Cp*V)_2\{B_2H_4(SePh)_2\}_2]$, **5** in parallel to the formation of $[(Cp*V)_2\{B_4H_{11}(SePh)\}]$, **6.** Compound **5** is very fascinating in which all the terminal B-H hydrogens of **1** have been substituted by {SePh} ligands. All the compounds have been characterized by 1H , ^{11}B , ^{13}C NMR spectroscopy; mass spectrometry; IR spectroscopy and single crystal X-ray analysis. Density functional theory (DFT) and TD-DFT calculations provided further understanding regarding the electronic structures, bonding and electronic transitions of these persubstituted vanadaborane species.

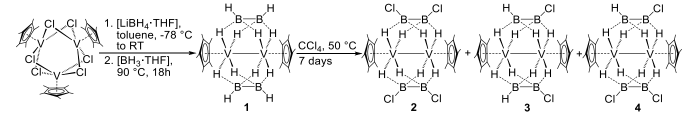
INTRODUCTION

In the expansion of rapidly growing polyhedral metallaborane^{1a-d} and metallacarborane^{1e-f} chemistry, functionalization of borane, metallaborane and carborane at B-H vertices became of interest because of their growing desirability in material science,² catalysis³ and drug design.⁴ However, compared to the widely explored carbon-functionalization⁵ in hydrocarbons, the boron-functionalization is relatively unexplored. Despite of the efforts devoted to the development of efficient routes for the functionalization of boranes⁷ and carboranes, 8 functionalization of metallaboranes met with little success.9 The icosahedral carboranes and boranes have been functionalized regularly because of their wide-spread applications as a basic component for Boron Neutron Capture Therapy (BNCT), 4a,10 weakly coordinating anion, 11 photoluminescent material, 12 radio imaging reagent, 13 tunable hybrid nanomolecules 14 etc. Reed and co-workers have shown that the hexahalogenated carboranes $[CB_{11}H_6X_6]^-(X = Cl, Br or I)$ play a significant role in stabilization of the coordinatively unsaturated cationic species, such as, the silylium cation (R_3Si^+) , 15 fullerene cation (C_{76}^+) , 16 hydronium ion (H₉O₄⁺), ¹⁷ and (tetraphenylporphyrinato)iron(III) ion [Fe-(tpp)⁺]. ¹⁸ Beginning with the pioneering contributions by Hawthorne ^{7a-b,8a} and Grimes, ^{6,8b-c} followed by Xie, ¹⁹ Yan, ^{19a,20} and others, 21,22 several synthetic methodologies have been established for the derivatization of boranes and carboranes. Recently, Spokoyny and co-workers have shown that perfunctionalization of borane clusters using pentafluroroaryl-terminated linkers yields size-tunable rigid nanomolecules that undergo conjugation with different types of thiols via S_NAr chemistry. 14

After the successful synthesis of perchlorinated rhenaborane [(Cp*Re)₂B₅Cl₅H₂],²³ search for new routes for the isolation of perhalogenated metallaboranes became of interest. One of the well-known approaches for the B-H functionalization of boranes or carboranes is the electrophilic substitution followed by cross coupling reactions.²⁴ Having a series of several open and *closo*-metallaboranes,²⁵⁻²⁷ we were interested in isolating perhalogenated compounds. Unfortunately, the above mentioned method was not successful for the generation of these species. Note that, recently we have reported various perhalogenated metalloboranes, albeit in lower yields, using PtBr₂, CCl₄, CHCl₃, MeI, BHCl₂·SMe₂ as halogenating agents. ²⁵⁻²⁷ recently, we have isolated perchlorinated [(Cp*Nb)₂(B₂H₄Cl₂)₂] using CCl₄ as chlorine source.²⁷ As a result, perchlorination of other group 5 metallaboranes became of interest. Herein, in this report, we illustrate the synthesis, structures and bonding of various persubstituted divanadaborane clusters.

RESULTS AND DISCUSSION

Synthesis and reactivity of divanadaborane [(Cp*V)₂ (B₂H₆)₂], 1. Reaction of [(Cp*VCl₂)₃] with 6 equivalents of [LiBH₄·THF] followed by thermolysis with excess [BH₃·THF] led to the formation of blue 1 in 31 % yield (Scheme 1). Compound 1 was characterized by mass spectrometric data; 1 H, 11 B, 13 C NMR spectroscopy and IR spectroscopy. The 11 B NMR spectrum showed a resonance at δ = -2.9 ppm. The 1 H NMR spectrum shows the presence of a single Cp* environment. In addition, 1 H NMR spectrum exhibited a chemical shift at -10.12 ppm corresponds to V-*H*-B. Both the 1 H and 11 B NMR spectra showed similarity with those of



Scheme 1. Synthesis of divanadaborane (1) and chlorinated divanadaboranes (2-4).

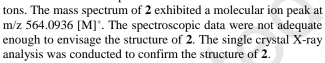
 $[(CpV)_2(B_2H_6)_2].^{28}$ The mass spectrometric data suggests the composition of 1 as $C_{20}H_{42}B_4V_2.$

In order to confirm the spectroscopic assignments of $\mathbf{1}$, the single crystal X-ray structure analysis was carried out. As depicted in Figure 1(a), the X-ray analysis revealed compound $\mathbf{1}$ as the the Cp* analogue of $[(CpV)_2(B_2H_6)_2]^{28}$ where two tetrahedral V_2B_2 units are fused by common V-V edge. The V-V, avg. V-B and avg. B-B bond lengths of 2.782, 2.304 and 1.758 Å respectively for $\mathbf{1}$ are practically analogous to that of $[(CpV)_2(B_2H_6)_2]$.

As shown in Scheme 1, reaction of $[(Cp*V)_2 (B_2H_6)_2]$, 1 with CCl_4 at 50 °C for one week led to the formation of violet

solid **2**. Room temperature ¹¹B NMR spectrum of **2** reveals a single resonance, appearing at $\delta = 10.1$ ppm. The ¹H NMR spectrum of **2** suggests the presence of one kind of Cp* environment, appeared at $\delta = 2.18$ ppm. In addition, the ¹H NMR spectrum of **2** shows one chemical shift at $\delta = -8.19$ ppm, indicating the existence of V–*H*–B pro-

B1 B2 V2 B4 B3

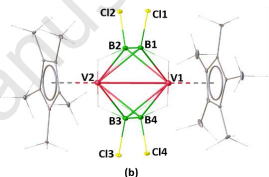


The X-ray diffraction study exposed the molecular formula of ${\bf 2}$ as $[(Cp^*V)_2(B_2H_4Cl_2)_2]$. The solid-state X-ray structure, shown in Figure 1(b), clearly shows that all the terminal B-H hydrogens are functionalized by Cl atoms. The V-V bond distance of ${\bf 2}$ (2.8389(6) Å) is slightly longer as compared to ${\bf 1}$ (2.7820 (9) Å). Whereas, the average V-B as well as B-B bond distance is comparable with that of ${\bf 1}$ (Table 1).

As shown in Scheme 1, in parallel to the formation of persubstituted divanadaborane 2, the prolonged thermolysis reaction of 1 with CCl₄ also yielded bichlorinated divanadaborane [(Cp*V)₂(B₂H₅Cl)₂], **3** and trichlorinated divanadaborane $[(Cp*V)_2(B_2H_4Cl_2)(B_2H_5Cl)]$, 4. The mass spectra show the molecular ion peaks of 3 and 4 at m/z 496.1801 and 530.1416 for [M]⁺, suggesting the molecular formula of C₂₀H₄₀Cl₂V₂B₄ and C₂₀H₃₉Cl₃V₂B₄ respectively. The ¹³C and ¹H NMR spectra reveals single Cp* environments both for 3 and 4. The ¹¹B NMR spectrum of 3 displays two boron resonances at $\delta = 9.0$ and -0.5 ppm. The ¹H NMR spectrum of 3 reveals distinct signals at δ = -8.66 and -9.65 ppm in 4:4 ratio for two dissimilar types of bridging V-H-B hydrogens. Whereas, the ¹¹B NMR spectrum of **4**, exhibits chemical shifts at $\delta = 10.3$ and -1.6 ppm. The ¹H NMR spectrum of 4 displays peaks at $\delta = -8.67$ and -9.80 ppm in 2:6 ratio that matches to bridging V-H-B hydrogens. The IR

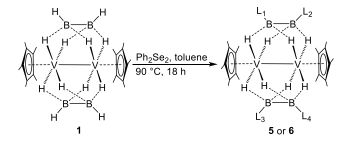
spectra of **3** and **4** further confirmed the existence of terminal B-H hydrogens. In order to deduce the solid-state X-ray structures of **3** and **4**, the single crystal X-ray analysis was undertaken. The solid state X-ray structure of **3**, shown in Figure S3, is a bichlorinated divanadaborane $[(Cp*V)_2(B_2H_5Cl)_2]$. Whereas, X-ray analysis of compound **4** showed a trichlorinated divanadaborane $[(Cp*V)_2(B_4H_9Cl_3)]$ (Figure S4). The V-V bond distances of **3** and **4** are slightly longer as compared to **1**, whereas the avg. B-B and avg.V-B bond lengths are slightly shorter.

Figure 1. Molecular structures and labeling of **1**(a) and **2**(b). Selected bond lengths [Å] and angles [deg] of **1**: V1-V2 2.7820(9),



V1-B1 2.300(4), B2-V1 2.295(4), B3-V1 2.301(4), B4-V1 2.307(4), B1-V2 2.307(4), V2-B2 2.315(4), V2-B3 2.299(4), V2-B4 2.308(4), B1-B2 1.752(7), B3-B4 1.764(6); B2-V1-V2 53.20(11), B2-V1-B1 44.81(16), B2-B1-V1 67.4(2), B2-V1-B1 44.81(16), B1-V1-B4 88.98(16); **2**: V1-V2 2.8389(6), V1-B1 2.302(3), V1-B2 2.302(3), V1-B3 2.306(3), C11-B1 1.844(3), C12-B2 1.844(3) B2-B1 1.755(5), B3-B4 1.756(5); B2-V1-V2 52.06(8), B1-V2-B2 44.64(12), B2-B1-V1 67.61(16), B2-V1-B1 44.80(12), B1-V1-B4 86.86(12), V1-B2-C12 140.52(18), V2-B2-C12 143.26(19).

Reactivity of [(Cp*V)2(B2H6)2] with Ph2Se2. Klumpp and co-workers have introduced R_2S_2 ligands (R = aryl or alkyl) as a source of 'S' and 'SR'.29 Recently, Himmel and co-workers have synthesized diborane(4) species $[R'SB(\mu-hpp)]_2$ (R' = Bz)or Ph) from the reaction of $[HB(\mu-hpp)]_2$ and Bz_2S_2 or Ph_2S_2 .³⁰ We have explored the reactivity of diaryl disulfide, diselenide and ditelluride ligands for the synthesis of various metallaheteroborane clusters. Recently, we have shown that the reaction of divanadaborane $[(CpV)_2(B_2H_6)_2]$ with Bz_2Se_2 or Ph_2S_2 yielded $[(CpV)_2\{B_2H_5(BzSe)\}(B_2H_6)],$ $[(CpV)_2\{B_2H_4(PhS)_2\}\{B_2H_5(PhS)\}],$ $[(CpV)_2\{B_2H_5(PhS)\}_2]$, albeit in poor yields.³¹ In an effort to isolate the persubstituted species, we have carried out the reaction of 1 with Ph₂Se₂ at 90 °C. It led to the formation of green solid **5**.



Scheme 2. Synthesis of 5 and 6 (5: $L_1 = L_2 = L_3 = L_4 = SePh$; 6: $L_1 = L_2 = L_3 = H$, $L_4 = SePh$).

The mass spectrum of **5** displays peak at m/z 1048.0567 for [M]⁺. The ¹H NMR spectrum shows single resonance at δ = 1.97 ppm that proposes one kind of Cp* environment. The ¹³C NMR spectrum further confirms the existence one Cp* environment. The ¹¹B NMR spectrum of **5** shows a single chemical shift at δ = 1.9 ppm. The ¹H NMR spectrum of **5** reveals signal for one type of V-*H*-B bridging hydrogens appeared at δ = -7.89 ppm. The IR spectrum of **5** confirmed the presence of V-*H*-B hydrogens. Although all these spectroscopic data suggest a highly symmetric structure of **5**, these data were not enough to predict the structure of **5**. As a result, single crystal X-ray analysis was undertaken on a suitable crystal.

The solid-state X-ray structure of **5**, shown in Figure 2, is $[(Cp*V)_2\{B_2H_4(SePh)_2\}_2]$, where all the 4 terminal B-H hydrogens are substituted by {SePh} ligand. The V-V bond distance of 2.859 Å is comparatively longer in comparison to its parent divanadaborane **1**. However, the avg. B-B bond distance for **5** (1.741 Å) is slightly shorter to that of **1** and **2**. The avg. B-Se bond distance of **5** (2.033 Å) is analogous to other reported dimetallaselenaborane clusters having B-Se bonds.³¹

As shown in Scheme 2, in parallel to the formation of **5** the thermolysis reaction of **1** with Ph₂Se₂ afforded green solid **6**. The mass spectrum of compound **6** displays molecular ion peak at 584.2027 corresponds to the molecular formula of C₂₆H₄₆B₄SeV₂. The ¹H and ¹³C NMR spectra show resonances for one type of Cp* environment. The ¹¹B NMR spectrum of **6** displays two signals at δ = 0.3 and -6.5 ppm. The ¹H NMR spec-

trum shows distinct signals for two different types of bridging V-H-B protons at δ = -8.30 and -9.95 ppm. The single crystal X-ray analysis was undertaken to determine the solid-state structure of **6**. The single crystal X-ray study revealed compound **6** as monosubstituted divanadaborane in which one of the terminal B-H hydrogens is substituted by {SePh} ligand. The V-V bond distance of compound **6** is shorter compared to that of persubstituted compound **5**.

As all the divanadaboranes 1-6 have the identical V_2B_4 core, a structural comparison of them became of interest (Table 1). Compounds 1-6 can be viewed as 44 cluster valence electron (cve) species, where two V_2B_2 tetrahedral is fused by one V-V bond. Although, the substituent atoms which form exo-polyhedral bonds with the boron atoms do not affect the skeletal bonding, because of their electronegativity and size difference with

boron and hydrogen, they affect the geometrical parameters.³² On stepwise substitution of hydrogens in **1** by electronegative chlorine atoms, the V-V bond length increases and when all the terminal B-H hydrogens are substituted by Cl atoms, the V-V distance became longer. Likewise, due to the large size of {SePh} ligand, the V-V bond distance increases in **5** and the B-B bond distance decreases upon substitution. On substitution of the hydrogen atoms at B-H in **1** by chlorine, the ¹¹B chemical shifts gradually shifted to down field. However, a minor ¹¹B chemical shift is observed when the substitution was occurred by {SePh} group. These observations may be attributed due to higher electron pulling ability of Cl atoms compared to {SePh} ligands.

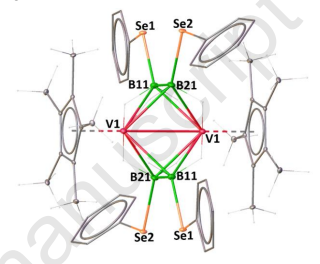


Figure 2. Molecular structure and labeling diagram of **5**. Selected bond lengths [Å] and angles [deg] of **5**: V1-V1 2.8595(10), V1-B11 2.331(3), V1-B21 2.333(3), B11-Se1 2.029(3), B11-B21 1.741(5); V1-V1-B11 52.28(8), B21-V1-B21 104.22(10), B21-V1-B11 87.56(12), B21-V1-B11 44.00(11), V1-B21-V1 75.78(10).

Table 1. Selected structural parameters and spectroscopic data of divanadaboranes.

Av = Average, d = distance

Compounds	Av <i>d</i> _{B-B} [Å]	dv-v [Å]	δ(V-H-B) ¹ H NMR (ppm)	¹¹ B{ ¹ H} NMR (ppm)
$[(CpV)_2(B_2H_6)_2]^{28}$	1.73	2.787	-9.7	1.7
$[(Cp*V)_2(B_2H_6)_2], 1$	1.758	2.782	-10.12	-2.9
$[(Cp*V)_2(B_2H_4Cl_2)_2], 2$	1.755	2.838	-8.19	10.1
$[(Cp*V)_2(B_2H_5Cl)_2], 3$	1.746	2.794	-8.66, -9.65	9.0, -0.5
$[(Cp*V)_2(B_2H_4Cl_2)(B_2H_5Cl)], 4$	1.740	2.810	-8.67, -9.80	10.3, -1.6
$[(Cp*V)_2\{B_2H_4(SePh)_2\}_2], 5$	1.741	2.859	-7.89	1.9
$[(Cp*V)_2(B_2H_6)\{B_2H_5(SePh)\}], {\bf 6}$	1.752	2.799	-8.3, -9.95	0.3, -6.5

Electronic structure analysis. To get some insight into the electronic structures and bonding of 1-6, theoretical calculations on the ground state of DFT at bp86/6-31g* level were carried out. Structural parameters of the optimized structures are almost consistent with the experimental bond lengths (Table S1). The molecular orbital (MO) analysis of 1-4 shows that the HOMOs of these divanadaboranes are centred on both vanadium atoms (d-orbitals). On the other hand, the HOMO of 5 is

localized at the d orbitals of the vanadium centers and the p orbitals of two Se atoms. Likewise, the HOMO of **6** is centred on both vanadium atoms (d-orbital) and one Se atom (p-orbital).

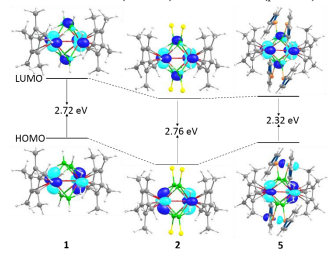


Figure 3. Comparison of frontier molecular orbitals of 1, 2 and 5.

Further, the MO analysis of **1-6** reveals that the energy gaps between HOMO-LUMO increase in the order of 6 < 5 < 1 < 3 < 4< 2 (Figure 3 and Table S2). These results have been further confirmed with the calculated natural charges on the vanadium atoms (qV), boron atoms (qB) and substituent atoms (qE) (Table S2). Upon persubstitution by chlorine, the natural charges on boron atoms (qB) become less negative and the natural valence populations around boron atoms decreases. This may be due to the electron pulling effect of chlorine atoms. Thus, the above results satisfyingly support the downfield ¹¹B chemical shifts of the boron atoms attached to chlorine. The HOMO-1 of divanadaboranes 1-4 are built of d orbitals of two vanadium atoms which are delocalized over two vanadium centres to form V-V bond. In case of 5, the HOMO-4 is localised on p orbitals of the Se atoms and d orbitals of two V atoms to form V-V bond. Further, the Laplacian electron density plots for 1, 2 and 5 show bond critical points (bcp) and bond paths between two vanadium atoms that indicate the presence of V-V bonds (Figure S38 and Table S3).

UV-vis absorption spectra of 1, 2 and 5. In order to investigate the effect on electronic transition upon B-H functionalization by electronegative Cl atoms and {SePh} ligands, UV-vis absorption studies became of interest. The UV-vis absorption studies of 1, 2 and 5 were performed using CH₂Cl₂ solution. As shown in Figure 4, compound 1 displays absorption peak at λ = 579 nm. The perchlorinated compound 2 shows two absorption peaks at λ = 410 and 565 nm. Whereas, the persubstituted compound 5 shows three absorption bands at λ = 432, 508 and 630 nm. In a comparison of the absorption spectra of 1, 2 and 5, a gradual blue shift (5 < 1 < 2) has been observed in the region of 650-550 nm.

To confirm these absorptions and get some insight into the electronic transition, we have carried out the time dependent-DFT calculation (Figures 4, S35-S37 and Tables S3-S5). In case of compound 1, the absorption nearby 579 nm may be assigned to the electronic transition that corresponds to HOMO to LUMO. As HOMO of 1 is centred on d-orbitals of two vanadium and LUMO is localised partially on the d-orbitals of vanadium and B-B bonds, according to TD-DFT calculation the

absorption neighbouring 579 nm may be assigned to an intramolecular MLCT transition. The absorption band at 565 nm for 2 corresponds to MLCT, whereas an absorption shoulder near 410 nm seems to be an intramolecular LMCT transition. The intense absorption bands at 630 nm for 5 appears due to the charge transfer from HOMO to LUMO and HOMO-2 to LUMO.

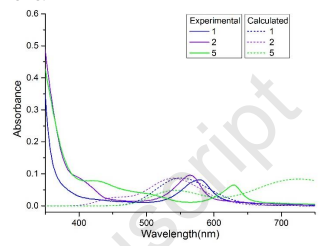


Figure 4. Experimental and calculated UV-vis spectrum of 1, 2 and 5

CONCLUSION

In summary, we have established a very simple and facile method for the perchlorination of $[(Cp*V)_2(B_2H_6)_2]$, 1 using CCl₄ as chlorine source. Further, we have synthesised persubstituted divanadaborane in which the hydrogens at B-H vertices have been substituted by {SePh} ligands. Upon perchlorination of 1, the UV-vis absorption spectrum shows a blue shift, while the persubstitution by {SePh} yielded a red shift of the absorption bands.

EXPERIMENTAL SECTION

General procedures and instrumentation. Syntheses of all these compounds were performed underneath of argon pressure using glovebox and Schlenk techniques. Solvents were distilled underneath of N2 atmosphere using common methods. [LiBH4·THF], [BH3·THF], CCl4 and Ph₂Se₂ (Aldrich) were used as purchased. All substances (Cp*VCl₂)₃³³ and [Bu₄N(B₃H₈)]³⁴, reference for ¹¹B{¹H} NMR spectra were prepared following the literature. On silica-gel TLC plates supported by dialuminium (Merck) thin layer chromatography was carried out. On a Bruker 500 MHz FT-NMR spectrometer all the NMR spectra were recorded. For ^{11}B NMR a sealed tube contains [Bu₄N(B₃H₈)] ($\delta =$ -30.07 ppm) in C₆D₆ was taken as an external reference. ¹¹B{¹H} spectra were processed with a backward linear prediction algorithm to eliminate the broad ¹¹B background signal of the NMR tube. ³⁵ For ¹H NMR spectra residual solvent protons were employed as reference ($\delta = 7.26$ ppm, CDCl₃; $\delta = 7.16$ ppm, C₆D₆). For ¹³C{¹H} NMR spectra residual solvent carbons were employed as a reference (δ = 77.1 ppm, CDCl₃; δ = 128.6 ppm, C₆D₆). On a Thermo Scientific (Evolution 300) UV-VIS spectrometer UV-vis absorption spectra were recorded. All the infrared spectra were recorded on a FT/IR-1400 (JASCO) spectrometer. Using 6545 Qtof LC/MS and Qtof Micro YA263 HRMS instruments all the mass spectra were recorded.

Synthesis of 1. In an air and moisture free Schlenk tube, (Cp*VCl₂)₃ (0.50 g, 0.65 mmol) was dissolved in toluene (25 ml) and chilled to -78 °C. To the reaction mixture [LiBH₄·THF] (1.95 ml, 3.90 mmol) was added slowly using a syringe, and slowly brought to room temperature by stirring for 1 h. Using a syringe BH₃·THF (3.9 ml, 3.9 mmol) was

added, followed by heating at 90 $^{\circ}$ C for 18 h. After evaporation of the solvents under vacuum, it was extracted using hexane/CH₂Cl₂ and celite. After exclusion of the solvent, it was exposed to chromatography on TLC plates and eluted with a CH₂Cl₂/hexane (v/v 30:70) mixture which produced blue [(Cp*V)₂B₄H₁₂] (0.13 g, 31%).

1: MS (ESI⁺): Calculated for [C₂₀H₄₂V₂B₄]⁺: 428.2550, found: 428.2530. 11 B{ 1 H} NMR (160 MHz, 22 $^{\circ}$ C, CDCl₃): δ = -2.9 (4B) ppm. 1 H NMR (500 MHz, 22 $^{\circ}$ C, CDCl₃): δ = 2.06 (s, 30H, C₅Me₅), 2.11 (br., 4H, BH_t), -10.12 (s, 8H, V-*H*-B) ppm. 13 C{ 1 H} NMR (125 MHz, 22 $^{\circ}$ C, CDCl₃): δ = 113.08 (s, C₅Me₅), 13.01 (s, C₅Me₅) ppm. IR (CH₂Cl₂, cm⁻¹): $\bar{\nu}$ = 2404.8 (BH_t).

Synthesis of 2-4. Inside a moisture free Schlenk tube, 1 (0.20g, 0.47 mmol) in CCl₄ (25ml) was stirred at 50 °C for 7 days. Conversion of 1 to chloro-substituted products were monitored using TLC. After elimination of the solvent, it was exposed to chromatography on TLC plates. Elution using a CH₂Cl₂/hexane (v/v 30:70) mixture afforded violet coloured compounds 2 (0.03g, 11%), 3 (0.04g, 18%) and 4 (0.05g, 20%). Note that, this reaction produced compound 2 as exclusive product in one day time under reflux condition. We have not observed any chlorination at B-H vertices while we record the NMR. However, keeping it in CDCl₃ for prolonged time, led to partial chlorination.

2: MS (ESI⁺): Calculated for [C₂₀H₃₈B₄Cl₄V₂]⁺: 564.1003, found: 564.0936. ¹¹B{¹H} NMR (160 MHz, 22 °C, CDCl₃): δ = 10.1 (4B) ppm. ¹H NMR (500 MHz, 22 °C, CDCl₃): δ = 2.18 (s, 30H, C₅Me₅), -8.19 (s, 8H, B-*H*-V) ppm. ¹³C{¹H} NMR (125 MHz, 22 °C, CDCl₃): δ = 117.22 (s, C₅Me₅), 12.31 (s, C₅Me₅) ppm.

3: MS (ESI⁺): Calculated for $[C_{20}H_{40}B_4Cl_2V_2]^+$: 496.1777, found: 496.1801. ${}^{11}B\{{}^{1}H\}$ NMR (160 MHz, 22 °C, CDCl₃): δ = 9.0 (2B), -0.5 (2B) ppm. ${}^{1}H$ NMR (500 MHz, 22 °C, CDCl₃): δ = 2.11 (s, 30H, C₅Me₅), 2.58 (br., 2H, BHt), -8.66 (s, 4H, V-t-B), -9.65 (s, 4H, B-t-V) ppm. ${}^{13}C\{{}^{1}H\}$ NMR (125 MHz, 22 °C, CDCl₃): δ = 115.10 (s, t-SMe₅), 12.67 (s, t-SMe₅) ppm. IR (CH₂Cl₂): \bar{v} = 2433.7 cm⁻¹ (B-t-Ht).

4: MS (ESI⁺): Calculated for [C₂₀H₃₉B₄Cl₃V₂]⁺: 530.1390, found: 530.1416. ¹¹B{¹H} NMR (160 MHz, 22 °C, CDCl₃): δ = 10.3 (1B), -1.6 (3B) ppm. ¹H NMR (500 MHz, 22 °C, CDCl₃): δ = 2.10 (s, 30H, C₅Me₅), 2.33 (br., 1H, B-H_t), -8.67 (s, 2H, B-H-V), -9.80 (s, 6H V-H-B) ppm. ¹³C{¹H} NMR (125 MHz, 22 °C, CDCl₃): δ = 114.25 (s, C₅Me₅), 12.84 (s, C₅Me₅) ppm. IR (CH₂Cl₂): $\bar{\nu}$ = 2433.7 cm⁻¹ (B-H_t).

Synthesis of 5 and 6. In a moisture free Schlenk tube, **1** (0.30g, 0.70 mmol) was dissolved in toluene (25 ml) followed by stirring with excess of Ph_2Se_2 (0.66g, 2.10 mmol) for 18 hrs at 90 °C. Conversion of **1** to substituted products were monitored using TLC. After exclusion of the solvent, on silica gel TLC plates the residue was exposed to chromatography and eluted using a CH_2Cl_2 /hexane (v/v 30:70) mixture which afforded green **5** (0.04g, 6%) and green **6** (0.06g, 16%).

5: MS (ESI⁺): Calculated for $[C_{44}H_{58}B_4Se_4V_2]^+$: 1048.0521, found: 1048.0567. ¹¹B{¹H} NMR (160 MHz, 22 °C, CDCl₃): δ = 1.9 (4B) ppm. ¹H NMR (500 MHz, 22 °C, CDCl₃): δ = 7.13-7.20 (C₆H₅), 1.97 (s, 30H, Cp* C₅Me₅), -7.89 (s, 8H, B-*H*-V) ppm. ¹³C{¹H} NMR (125 MHz, 22 °C, CDCl₃): δ = 137.38, 130.98, 128.76, 124.74 (C₆H₅), 117.44 (s, *C*₅Me₅), 11.73 (s, C₅*Me*₅) ppm.

6: MS (ESI⁺): Calculated for [C₂₆H₄₆B₄SeV₂]⁺: 584.2039, found: 584.2027. ¹¹B{¹H} NMR (160 MHz, 22 °C, CDCl₃): δ = 0.3 (1B), -6.5 (3B) ppm. ¹H NMR (500 MHz, 22 °C, CDCl₃): δ = 7.09-7.16 (C₆H₅), 2.16 (s, 30H, C₅Me₅), 2.47 (br, 3H, B-H_t), -8.30 (s, 2H, B-H-V), -9.95 (s, 6H, V-H-B) ppm. ¹³C{¹H} NMR (125 MHz, 22 °C, CDCl₃): δ = 138.31, 130.18, 127.66, 123.70 (C₆H₅), 114.30 (s, C₅Me₅), 12.73 (s, C₅Me₅) ppm. IR (CH₂Cl₂): $\bar{\nu}$ = 2403.8 cm⁻¹ (B-H_t).

X-ray crystal structure determinations. The X-ray crystal data of compounds **1** and **3-6** were collected at 150 K and integrated in a D8 VENTURE Bruker AXS diffractometer under graphite-monochromated MoK $_{\alpha}$ radiation ($\lambda = 0.71073$ Å). Using a Bruker APEX-II CCD diffractometer equipped with graphite-monochromated MoK $_{\alpha}$ radiation ($\lambda = 0.71073$ Å), X-ray crystal data of compound **2** was collected

at 296 K and integrated. Using SADABS³⁶ program multi-scan absorption correction has been done on the X-ray crystal data. All the structures were solved employing SHELXS-97 or SIR92 and refined with SHELXL-2014.³⁷ Using Olex2³⁸ all the molecular structures were drawn. CCDC 1940467 (1), 1940465 (2), 1940441(3), 1940442(4), 1940439(5) and 1940466 (6) contains crystallographic data. These data are available free of charge at www.ccdc.cam.ac.uk/data_request/cif.

Crystal data of 1. C₂₀H₄₂B₄V₂, Mr = 427.65 g.mol⁻¹, Monoclinic, P21/n, a = 11.3343(9) Å, b = 14.6762(10) Å, c = 14.5522(10) Å, $\alpha = 90^{\circ}$, $\beta = 109.887(2)^{\circ}$, $\gamma = 90^{\circ}$, V = 2276.3(3) Å³, $\rho_{calcd} = 1.248$ g.cm⁻³, Z = 4, $\mu = 0.823$ mm⁻¹, F(000) = 912, GOF = 1.061, $R_1 = 0.0866$, $wR_2 = 0.2490$, reflections collected 18801, independent reflections 5086 (20 $\leq 54.954^{\circ}$) and 281 parameters.

Crystal data of 2. $C_{20}H_{38}B_4Cl_4V_2$, Mr = 565.42 g.mol⁻¹, Monoclinic, P21/n, a = 10.4129(4)Å, b = 17.7675(7)Å, c = 14.8537(5) Å, $a = 90^{\circ}$, $\beta = 98.7410(18)^{\circ}$, $\gamma = 90^{\circ}$, V = 2716.18(18) Å³, $\rho_{calcd} = 1.383$ g.cm⁻³, Z = 4, $\mu = 1.089$ mm⁻¹, F(000) = 1560, GOF = 1.028, $R_1 = 0.0389$, $wR_2 = 0.1075$, reflections collected 21357, independent reflections 4784 (20 $\leq 49.992^{\circ}$) and 313 parameters.

Crystal data of 3. C₂₀H₄₀B₄Cl₂V₂, $Mr = 496.18 \text{ g.mol}^{-1}$, Orthorhombic, Pnma, a = 20.5180(11) Å, b = 15.4472(9) Å, c = 7.6522(4) Å, $\alpha = \beta = \gamma = 90^{\circ}$, V = 2425.3(2) Å³, $\rho_{calcd} = 1.266 \text{ g.cm}^{-3}$, Z = 4, , $\mu = 0.884 \text{ mm}^{-1}$, F(000) = 976, GOF = 1.186, $R_1 = 0.0441$, $wR_2 = 0.0983$, reflections collected 17157, independent reflections 2862 ($2\theta \le 54.963^{\circ}$) and 160 parameters.

Crystal data of 4. C₂₀H₃₁B₄Cl₃V₂, $Mr = 522.92 \text{ g.mol}^{-1}$, monoclinic, P21, a = 7.9109(6) Å, b = 15.3964(9) Å, c = 10.8018(7) Å, $\alpha = \gamma = 90^{\circ}$, $\beta = 101.498(3)^{\circ}$, V = 1289.25(15) Å³, $\rho_{calcd} = 1.347 \text{ g.cm}^{-3}$, Z = 2, $\mu = 1.041 \text{ mm}^{-1}$, F(000) = 536, GOF = 1.080, $R_1 = 0.1035$, $wR_2 = 0.2840$, reflections collected 9083, independent reflections 5318 ($20 \le 54.964^{\circ}$) and 150 parameters.

Crystal data of 5. C₄₄H₅₈B₄Se₄V₂, Mr = 1047.86 g.mol⁻¹, Triclinic, P-1, a = 10.6356(14) Å, b = 10.7180(13) Å, c = 11.2446(15) Å, $\alpha = 106.352(4)$ °, $\beta = 113.721(4)$ °, $\gamma = 96.642(4)$ °, V = 1087.1(2) Å³, $\rho_{calcd} = 1.601$ g.cm⁻³, Z = 1, $\mu = 3.809$ mm⁻¹, F(000) = 524, GOF = 1.052, $R_1 = 0.0441$, $wR_2 = 0.1273$, reflections collected 21602, independent reflections 4991 ($20 \le 55.102$ °) and 261 parameters.

Crystal data of 6. C₂₆H₄₆B₄SeV₂, Mr = 582.71 g.mol⁻¹, Triclinic, P = 1, a = 8.1846(6) Å, b = 10.2047(7) Å, c = 18.8337(14) Å, $\alpha = 105.551(2)^{\circ}$, $\beta = 92.550(3)^{\circ}$, $\gamma = 107.010(3)^{\circ}$, V = 1436.18(18) Å³, $\rho_{calcd} = 1.347$ g.cm⁻³, Z = 2, $\mu = 1.931$ mm⁻¹, F(000) = 604, GOF = 1.029, $R_1 = 0.0290$, $wR_2 = 0.0717$, reflections collected 33324, independent reflections 6577 ($20 \le 54.968^{\circ}$) and 341 parameters.

Computational Details. Optimization of all the molecules are done using the Gaussian 09³⁹ program. The gradient-corrected bp86⁴⁰ functional along with 6-31g* basis set had been employed for the optimization. Starting from X-ray crystallographic coordinates the model compounds were fully optimized without any solvent effect in gaseous state. Frequency calculations were carried out at the same level of theory. With the natural bond orbital (NBO) partitioning scheme⁴¹ as employed in the Gaussian 09 programs natural bonding analyses were executed. Multiwfn V.3.6 package⁴² was used to carry out the QTAIM analysis. All the optimized structures and orbital pictures were created with the Gaussview⁴³ and Chemcraft⁴⁴ programs.

ASSOCIATED CONTENT

Supporting Information

Details of X-ray structures, spectroscopic data, DFT and TD-DFT. The supporting Information is available free of charge on the ACS publication website.

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Notes

The authors declare no competing financial interests.

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