Oxidation, Coordination, and Nickel-Mediated Deconstruction of a Highly Electron-Rich Diborane Analogue of 1,3,5-Hexatriene

Alexander Hermann*, Felipe Fantuzzi**, Merle Arrowsmith, Theresa Zorn, Ivo Krummenacher, Benedikt Ritschel, Krzysztof Radacki, Bernd Engels, and Holger Braunschweig*

Abstract: The reductive coupling of an N-heterocyclic carbene (NHC) stabilized dibromovinylborane yields a 1,2-divinyl-diborane, which, although isoelectronic to a 1,3,5-triene, displays no extended \( \pi \) conjugation because of twisting of the \( C_2B_2C \) chain. While this divinyl-diborane coordinates to copper(I) and platinum(0) in an \( \eta^6 \)-\( B_2 \) and \( \eta^4 \)-\( C_2B_2 \) fashion, respectively, it undergoes a complex rearrangement to an \( \eta^4 \)-1,3-diborane upon complexation with nickel(0).

Introduction

Linear conjugated alkenes owe their intrinsic stability to the delocalization of their \( \pi \) electrons. Found in many natural products and biologically relevant compounds,[1] they are also important building blocks in organic synthesis and materials chemistry. Conjugated trienes and higher oligoenes have attracted interest because of their photophysical properties, which enable applications in nonlinear optics and optical sensing.[2] In an industrial setting, conjugated dienes (e.g. butadiene, isoprene) are mainly used as monomers for the Ziegler–Natta synthesis of synthetic rubbers.[3] In organic chemistry, they are principally employed in 1,4-addition[4] and Diels–Alder reactions,[5] as well as numerous other transformations.[6] Many of these reactions are metal-catalyzed and involve transition-metal (TM) 1,3-diene complexes as key reaction intermediates. Solution and solid-state analyses of such complexes show that the diene ligand can be found either in the cis or trans conformation and switch between the \( \eta^2 \) and \( \eta^4 \) coordination modes (Figure 1a).[7] which may determine its subsequent reactivity with incoming substrates.

The substitution of one or more carbon atoms with more electronegative heteroatoms (e.g. N, O) has long been exploited to generate polar conjugated systems, which are employed in numerous organic reactions (e.g. Michael additions[8] and hetero-Diels–Alder reactions).[9] In contrast, however, the chemistry of conjugated heterodiienes or heterotrienes in which one or more carbon atoms have been substituted with a more electropositive element is virtually unexplored.

One of the focuses of our research is on the synthesis, reactivity, and metal coordination of compounds displaying boron–element[10] and boron–boron multiple bonds.[11–13] Among the latter, doubly base-stabilized diboranes, which are formally isoelectronic and isostructural to alkenes, have been relatively well studied since their first isolation by Robinson in 2007.[14] Unlike most alkenes, diboranes undergo 1,2-addition and [2+2] cycloaddition reactions without the need for a catalyst, owing to their high-lying HOMO and relatively low-lying LUMO.[11] While the B–B bond coordinates to coinage metals in an \( \eta^2 \) fashion reminiscent of metal–olefin \( \pi \) complexes,[12] it is also sufficiently electron-rich to bind to ZnII, CuII,[13] and even MgII centers,[17] which do not tend to form stable \( \pi \)-olefin complexes because of their limited capacity for \( \pi \) backdonation. DFT calculations have shown that the B–M interaction in these complexes is mostly electrostatic in nature (ca. 60–70%), with the electron-rich B–B bond donating into empty orbitals of the metal center, and no or little \( \pi \) backbonding from the metal to the diborane unit.[12,13,17] The only reactivity reported to date for such TM-diborane complexes is that of a PMe3-stabilized bis(9-
Results and Discussion

The NHC-stabilized dibromovinylboranes (IPr)BBR₂(C(R)=CHR) (IPr=1,3-dioxopropylimidazol-2-ylidene; R = R' = Me 1-Me, Ph 1-Ph; R = Me, R' = tBu 1-tBu) were synthesized by hydroboration of the corresponding R–CHR alkynes precursors with HBcat (Cat = catecholate),[23] followed by adduct formation with IPr and bromination with BBBr₂.[20] The reduction of 1-Me and 1-Ph with 2.5 equivalents KC₈ in benzene at room temperature resulted in relatively unselective reactions from which the only isolable crystalline products were the bis(IPr)-stabilized 1,4-bis(bromoboraneylidene)butanes 2-Me [δ(B) = 12.5 ppm, broad] and 2-Ph [δ(B) = 18.4 ppm, broad], resulting from radical C–C coupling of two vinylborane units at the β position. 1-Me and 1-Ph could be further reduced with 10 equivalents KC₈, albeit unselectively,[20] to the corresponding doubly IPr-stabilized 4,5-dihydro-1,2-diborinines 3-Me [δ(B) = 19.9 ppm, broad] and 3-Ph [δ(B) = 27.5 ppm, broad; Figure 2].[21]

While repeated attempts at isolating analytically pure samples of 2-R and 3-R (R = Me, Ph) failed because of coscrystallization with other unidentified reaction products, the solid-state structures of 2-Ph (see Figure S52 in the Supporting Information), 2-Me, and 3-Ph (Figure 2) were unequivocally determined by X-ray diffraction analyses. These confirmed the presence of the B-C bonds [2-Me B1-C1 1.422(2), B2-C4 1.426(2); 2-Ph B1-C1 1.439(3), B2-C4 1.436(3); 3-Ph B1-C1 1.468(4) Å],[22] the newly formed C–C single bonds [2-Me C2-C3 1.551(2); 2-Ph C2-C3 1.558(2); 3-Ph C2-C2' 1.530(4) Å], and the endocyclic B–B bond in 3-Ph [1.694(5) Å]. Unfortunately, all attempts to synthesize these compounds more selectively failed.

In contrast, the reduction of 1-tBu with 4 equivalents KC₈ in benzene at room temperature led to the selective formation of the red-colored divinylborene 4, which was isolated in 72% yield as a brown solid (Scheme 1). The presence of the sterically demanding β-tert-butyl substituent prevents the recombination of the intermediate β-carbon-centered radical, favoring instead further reduction of the boron center and boron–boron bond formation. The diborene 4 presents a broad ¹¹B NMR resonance at δ = 25.1 ppm, in the range typical for NHC-stabilized diborenes.[31] The ¹H NMR spectrum displays a characteristic 2H quartet at δ = 4.80 ppm (J = 1.2 Hz) for the vinylic protons coupling to the tert-butyl protons and correlating by HSQC to a ¹³C[¹H] NMR resonance at δ = 132.4 ppm.

The solid-state structure of 4 is shown in Figure 3a. The B–B bond length of 1.601(2) Å lies in the upper range of B–B bonds[31] and is similar to that found in doubly IPr-stabilized di(2-thienyl)[23] or ferrocenediyl-bridged diborenes,[24] for

For 3-Ph: 10 KC₈, CdHex, rt

2 R = Me 1-Me, Ph 1-Ph

2.5 KC₈

CDHex, rt

allyl radical recombination

Figure 2. Reduction of 1-Me and 1-Ph with postulated radical mechanism and crystallographically derived molecular structures of 2-Me and 3-Ph. Thermal ellipsoids set at 50% probability.[20] Thermal ellipsoids of ligand periphery and hydrogen atoms omitted for clarity.
example. The diborene core is slightly distorted from planarity, with (C1,B1,B2,C21) and (C8,B1,B2,C28) torsion angles of 177.06(16) and 164.82(14)°, respectively. The vinyl groups display localized B–C single [1.580(2) and 1.589(2) Å] and C=C double bonds [1.354(2) and 1.354(2) Å], and a Z configuration of the alkyl substituents. Figure 3b shows that the vinyl units are not coplanar with the mean plane of the diborene, and are rotated by α = 37.9° and β = 30.7°. This conformation may arise from steric repulsion between the NHC iPr and the vinyl α-methyl groups. As a result, there is no π conjugation possible between the C=C and B=B bonds. This lack of π conjugation is supported by DFT calculations at the B3LYP/6–311G** level, which show that the π-bonding molecular orbitals of 4 are largely localized: the HOMO on the B=B bond, the HOMO-2 on the two C=C bonds and the HOMO-3 slightly delocalized over each of the C,B fragments (Figure 3c). This localization contrasts with the extensive π delocalization observed in the related and entirely planar 1,3,5-hexatriene molecule (Figure 3c). It is noteworthy that calculations on an analogue of 4 in which the methyl groups at the α-vinyl positions have been replaced by protons, the compound 4H, yield a quasi-planar C,B,C core with significantly shortened B–C bonds (1.566 Å), which displays a greater extent of π conjugation than 4 (Figure 3d). Consequently, the lack of planarity and π conjugation in 4 can be ascribed mainly to the steric at the α-vinyl positions. The UV-vis spectrum of 4 displays two absorption maxima at 453 and 573 nm, which account for the brown color of the compound. TDDFT calculations at the same level of theory provide a good match for the absorption maxima (439 and 582 nm) and show that the absorption at 573 nm results from a HOMO — LUMO transition (89%), while that at 453 nm is related to a HOMO — LUMO + 1 transition (89%).

While 4 was indefinitely stable under inert conditions in the solid state and in benzene solution up to 80°C, it decomposed entirely within minutes in polar solvents (THF, 3,5-difluorobenzene), thereby preventing the acquisition of cyclic voltammetry data. Its reducing power was, however, confirmed by one-electron oxidation with [C,F][BF₄] (ArF = 3,5-trifluoromethylphenyl), which yielded the red-purple radical cation [4][BF₄]· (Scheme 2a). Like other [(NHC)B,R₂]⁺ radical cations with [BF₄]⁻ as the counteranion,[25] the EPR spectrum of [4][BF₄]· in 3,5-difluorobenzene showed a broad signal, for which a simulation provided a hyperfine coupling parameter of α(B) = ca. 1.7 G (Figure 4, left). We have shown that the doubly iPr-stabilized 1,2-diisopropyl diborene is sufficiently reducing to react with 1-mesityl-2,3,4,5-tetraphenylborole (Eₚ= = −1.69 V, mesityl −

**Figure 3.** a) Crystallographically derived molecular structure of 4.[24] Thermal ellipsoids at 50% probability. Thermal ellipsoids of ligand periphery and hydrogen atoms omitted for clarity. Selected bond lengths [Å] and angles [°]: B1–B2 1.601(2), B1–C1 1.600(2), B1–C8 1.581(2), B2–C21 1.589(2), B2–C28 1.584(2), C1–C2 1.354(2), C21–C22 1.354(2), Σα,α 359.57(14), Σβ,α 359.89(14), torsion (C1,B1,B2,C21) 177.06(16), (C8,B1,B2,C28) 164.82(14). b) Truncated view of 4 along the B₂ plane. Angles between mean planes: α = 37.9°, β = 30.7°. Plots of the π bonding frontier molecular orbitals of 1,3,5-hexatriene (c), 4⁺ (d), an analogue of 4 in which the methyl groups at the α-vinyl positions have been truncated, and 4 (e) at the B3LYP/6–311G** level of theory.
2,4,6-Me(C$_6$H$_4$)$_2$Cl[27] to form the radical cation/radical anion pair [[(IPr)$_2$B$_2$Pr$_2$]$^+$][MesBC$_6$H$_5$]$^-$. Similarly, 4 reacted with the borole 5 to yield the radical cation/radical anion pair [4]$^+$[5]$^-$ (Scheme 2b), which displays a broad EPR signal consisting of the overlap of the radical cation and radical anion resonances (Figure 4, right). Given the success of this single-electron transfer reaction, we can conclude that the redox potential of 4 must be lower than that of 5, that is, $-1.69$ V. The lowest redox potential of a diborene measured to date remains that of [(IPr)$_2$B$_2$Pr$_2$] at $-1.95$ V[20d]. Since the oxidation potential of conjugated alkenes to radical cations occurs generally above $1$ V[23] we can conclude that the oxidation of 4 occurs exclusively at the diborone core and not at the vinyl moieties.

Like other diborones, 4 formed a $\pi$-diborone complex with CuCl, 4-Cu (Scheme 3a)[12a–c]. The bright-yellow compound displays a broad $^1$B NMR resonance at $\delta=19.5$ ppm, slightly upfield with respect to 4 [$\delta(^1$B) = 25.1 ppm] in accordance with other known coinage metal diborone complexes[12]. Similarly, the $^1$H NMR quartet of the vinylic protons (2H) is upfield-shifted from $\delta = 4.80$ ppm in 4 to $\delta = 4.29$ ppm in 4-Cu. The solid-state structure of 4-Cu (Figure 6) is similar to those reported for other $\pi$-diborone copper complexes,[12a–c] with a slight elongation of the B–B bond [1.627(4) Å] compared to that in 4 [1.601(2) Å] and increased distortion of the diborone core away from planarity, $\Sigma_4 = 375.8(3)$, $\Sigma_4 = 375.7(3)$, torsion angles (C1,B1,B2,C2) $-166.6(3)$, (C8,B1,B2,C8) $156.1(3)^\circ$.

In contrast, the reaction of Pt(nbe)$_1$ (nbe = norbornene) with 4 yielded the pink-colored 1,2-diborabutaadiene complex 4-Pt (Scheme 3b). The solid-state structure of 4-Pt shows that the divinylidiborone has displaced two of the nbe ligands and coordinates to platinum through $\pi$ interactions with both the B–B [B1-Pt1 2.248(4), B2-Pt1 2.343(4) Å] and one of the C–C bonds [C1-Pt1 2.227(3), C2-Pt1 2.236(3) Å; Figure 5], which is rotated in the direction of the metal center to give a cis-$\eta^1$ configuration. This cis-$\eta^1$ binding mode contrasts with the bonding of Pt$^6$ to conjugated olefins, which is usually limited to cis-$\eta^4$. The B1-Pt bond [1.601(2) Å] is shorter than those of 4–Cu [2.0023, 2.0025, 2.0023 Å, C1–C2 1.354(2) Å], while the B1–C1 bond [1.572–5 Å] is shorter than those of 4–Cu [2.0023 Å].

DFT calculations show that the HOMO of 4-Pt is a $\pi$ orbital mostly localized on the B–B bond donating into an empty $d$ orbital at the platinum center, with only a small contribution of the C–C bond and a node at the B1–C1 bond (Figure 5a). The HOMO-1 consists mainly of the vinylic bonding component localized on the B1–C1 bond, as already suggested by its shortened bond length in the solid-state structure (Figure 6). The nature of the Pt–C$_2$B$_2$ bonding was further analyzed by energy decomposition analysis combined

![Figure 4](image-url). Experimental (black) and simulated (red) EPR spectra of [4]$^+$[BAr]$^-$ in o-difluorobenzene (left) and [4]$^+$[5]$^-$ in hexanes (right) at room temperature. Simulated parameters are as follows:

- [4]$^+$[BAr]$^-$: $g_{oa}=2.0023$, $\alpha(B) = 0.17 G$ (peak-to-peak width $= 7 G$).
- [4]$^+$[5]$^-$: $g_{oa}=2.0025$, $\alpha(B) = 3.5 G$.

The B1–C1 and C1–C2 bonds [1.418(5) Å] are significantly longer than those of 4 [B1–B2 1.601(2), C1–C2 1.354(2) Å], while the B1–C1 bond [1.572–5 Å] is shorter than those of 4–Cu [1.600(2) Å], which suggests some amount of $\pi$ delocalization on the platinum-bound B2-B1-C1-C2 unit.

![Figure 5](image-url). a) Plot of the HOMO and HOMO-1 of 4-Pt at the B3LYP/TZV2P level of theory. b) Plot of deformation densities ($\Delta\rho_f$), at the same level of theory, of the orbital interactions of the C$_2$B$_2$ fragment $\pi$-donating to the Pt$^6$ center (left) and the Pt center $\pi$-backdonating to the C$_2$B$_2$ fragment (right). The $\lambda_{ij}$ values correspond to the eigenvalues of the complementary eigenfunctions ($\psi_+ \cdot \psi_-$) in the NOCV representation, while $\Delta\rho_{\text{orb}}$ is the $k$ orbital interaction energy (kcal mol$^{-1}$), with the percentage contribution to the total orbital interaction energy ($\Delta\rho_{\text{orb}}$) shown within parentheses. The electron density flows from yellow to purple.
with the natural orbitals for chemical valence theory (EDA-NOCV).

The results suggest that the bonding in 4-Pt is dominated by electrostatics (65.0%), with non-negligible orbital interaction contributions (35.0%). These arise from a combination of equal amounts of the C2B π-symmetrized fragment orbital (SFO), mostly centered on the B–B bond, donating into an empty platinum d SFO, and the platinum d π-donating into an empty π* SFO of the C2B fragment, with a strong B1–C1 bonding component (Figure 5b). This bonding picture is also reflected in the calculated Mayer bond orders (MBOs) of 4-Pt and the metal-free optimized cis-η*-like structure of 4, namely cis-4. While the bond order of the B–B and C1–C2 bonds decrease from 1.50 and 1.75, respectively, in cis-4 to 1.13 and 1.28, respectively, in 4-Pt, only a very small increase from 0.87 in cis-4 to 0.89 in 4-Pt is observed in the MBO of B1–C1.

In solution, the 11B NMR spectrum of 4-Pt showed a broad resonance at δ = 10.6 ppm, which is strongly upfield-shifted from that of 4 [δ(C2B) = 25.1 ppm] and 4-Cu [δ(C2B) = 19.5 ppm], presumably owing to the strong π backdonation of the Pt* center. The room-temperature 1H NMR spectrum showed very broad signals and those for the vinylic protons were undetectable. At low temperature (−90 to −40°C) at least four different conformers are visible with vinylic proton resonances around δ = 5 ppm. These conformers could be rapidly exchanging cis/trans-η*-C2B2-Pt and η*-C2B2-Pt conformers, in which the C1=C2 and the C21=C28 bonds are alternately bound to the Pt center, similarly to the bonding motifs found in 1,3-diene complexes (Figure 1a). Moreover, at temperatures above 40°C 4-Pt decomposed rapidly in solution. An attempt to stabilize 4-Pt by replacing the remaining nbe ligand with iPr resulted in complete release of free 4, as observed by 11B and 1H NMR spectroscopic analyses (Scheme 3c; see Figures S34 and S35).

Unlike its reactions with CuI and Pt(nbe), the reaction of 4 with Ni(COD)2, (COD = 1,5-cyclooctadiene) did not result in simple coordination to the metal center. Instead a complex rearrangement of the B–B unit and one vinyl group took place, resulting in the formation of the NiC2B4 complex 6 [δNi(COD)2 = 13.3 ppm] as the major reaction product (Scheme 4). Unlike for 4-Pt, the addition of iPr to 6 did not result in the liberation of the diborete ligand and no reaction was observed.

The X-ray crystallographically derived structure of 6 (Figure 6) shows the nickel center bound to all four atoms of a 1-vinyl-1,3-diborete ligand, which displays a butterfly structure with the carbon atoms located at the tips of the wings, a puckering angle of 40.2° and a B–B distance of 1.890(2) Å. Furthermore, the β-vinyl hydrogen H2 has migrated from C2 to B1 [H1–H2 1.106(17) Å] and the two iPr ligands have migrated from boron to nickel, displacing the COD ligands. The B–C bond lengths are all relatively similar [1.5430(19) to 1.5584(19) Å] and shorter by about 0.05 Å compared to typical B–C bonds, suggesting some π delocalization over the C2B4 ring. This delocalization is also confirmed by the 13C NMR resonances of the C2B4 ring, which appear in the

![Figure 6. Crystallographically derived molecular structures of 4-Cu (the non-disordered one of the two distinct molecules present in the asymmetric unit), 4-Pt, and 6](image_url)
To assess the electronic situation of 6, the nature of bonding was examined by EDA-NOCV. Two distinct scenarios were assessed: a) The donor-acceptor interaction of a Ni\(^{II}\) fragment with a neutral 2π-electron 1,3-diborete ligand, and b) the interaction of a Ni\(^{II}\) center and a diatomic 4π-electron \([C_2B_4]^{2-}\) ligand. The calculations reveal that, irrespective of the choice of fragments, the main bonding contribution arises from σ interactions between Ni and the carbon atoms of the C\(_2\)B\(_4\) ring (Figure 7). The scenario involving Ni\(^{II}\) and a neutral C\(_2\)B\(_4\), 1,3-diborete, however, yields a lower absolute value of the total orbital interaction energy, \(\Delta E_{\text{orb}}\) (Figure 7; see Table S5),[33] which indicates a more appropriate choice of fragments. These data suggest that the bonding in 6 is best described as the result of the Ni\(^{II}\) fragment donating into an empty π\(^*\) SFO of the neutral diborete ligand located at the carbon centers. This donor–acceptor interaction accounts for more than 80% of \(\Delta E_{\text{orb}}\), thereby suggesting that the valence electrons of the C\(_2\)B\(_4\) ring are bystanders. The calculated MBOs of roughly unity for all endocyclic B–C bonds in 6 suggest delocalization of the two π electrons over the C\(_2\)B\(_4\) ring despite the lack of planarity. Furthermore, the MBO of only 0.25 for B1–B2 confirms the absence of B–B bonding.

The complex 6 may also be viewed as a 22 electron C\(_2\)B\(_4\)Ni close-cluster according to the Wade–Mingos rules and is the smallest nickel-carborane cluster reported to date. The average bond lengths within the C\(_2\)B\(_4\)Ni fragment [Ni-C\(_{\text{org}}\) 1.99; Ni-B\(_{\text{org}}\) 2.25; B–B 1.890(2); B–C\(_{\text{org}}\) 1.55 Å] are within the range of other nickel carborane clusters.[34]

Considering the number of strong bonds broken (one C= C bond and the B–B bond, one C–H and two B–C bonds, as well as four Ni–C\(_{\text{org}}\) π interactions) and reformed (three B–C, one B–H, two Ni–B, and four Ni–C bonds) during the formation of 6, the reaction is surprisingly selective.[35] We therefore decided to undertake a computational analysis of the mechanism of formation of 6 at two different levels of theory, the results of which are shown in Figure 8 (see the Supporting Information for details). We propose that in the first step, Ni coordinates to the divinyl diborete in an analogous manner to Pt, yielding the slightly favorable intermediate 4-Ni. The next step, which is the rate-determining one, consists of the migration of the first NH ligand to the Ni center and liberation of one molecule of COD. This step is followed by an intramolecular [2+2] cycloaddition of the alkene moiety to the B–B bond, starting from intermediate (4-Ni)b and leading to the 1,2-diborete complex (4-Ni)c. While a handful of cycloaddition reactions of alkynes to B–B multiple bonds accompanied by C\(_2\)B\(_4\) rearrangements have been reported,[36] this is the first example of cycloaddition of an alkene to a B–B multiple bond. The rearrangement of (4-Ni)c to its 1,3-diborete isomer (4-Ni)d may be expected: extensive experimental studies in the 1980s[36,37] and later computational investigations[38] have shown that in the absence of electronic stabilization 1,2-diboretes rearrange to their thermodynamically more stable 1,3-isomers. The final formation of 6 by migration of the second NH to Ni and of H2 from C2 to B1 is calculated to be highly favorable from a thermodynamic point of view (\(AG = \approx -28.2\) kcal mol\(^{-1}\) at the B3LYP-D3(BJ)/def2-TZVPP level), and the barrier heights obtained are consistent with a reaction temperature of 80 °C.

**Conclusion**

The synthesis of 4 from the reductive coupling of two NHCl-stabilized (dibromo)vinylboranes was only rendered possible by suppressing β-carbon radical recombination through the introduction of a sterically hindering tert-butyl group in this position. While formally isoelectronic to a 1,3,5-hexatriene, experimental and theoretical data show that 4 does not display any delocalization of π electron density over the C\(_2\)B\(_2\)C\(_2\) core. Calculations show that this lack of delocalization is mainly a result of the steric of the methyl groups at the α-vinyl positions preventing planarization.

We have shown that the coordination mode of such a 3,4-dibora-1,3,5-hexatriene is strongly dependent on the nature of the metal used, unexpectedly resulting in three different outcomes with three different late transition metals. Whereas with CuClL, 4 forms a typical π-diborete complex, it coordinates to Pt\(^{4+}\) in a fashion reminiscent of 1,3-dienes by forming a cis-γ-vinyl diborete complex, the coordination of which is fluxional in solution. EDA-NOCV calculations show that, despite a stronger degree of planarization in the metal-bound C\(_2\)B\(_2\) unit, there still is little delocalization of the π electron density: π donation to platinum occurs mostly from the B–B double bond, while the Pt center π-backdonates into the empty π\(^*\) orbital of the C\(_2\)B\(_2\) ligand. In contrast, coordination of the vinyl diborete unit to a Ni\(^{II}\) complex induces a complex rearrangement into an η\(^1\),1,3-diborete complex, which proceeds by a novel metal-templated cycloaddition of the alkene moiety to the adjacent diborete.

This study demonstrates once more that the replacement of a C=C bond by an iso-electronic, yet much more electron-rich B–B bond considerably alters the chemistry of the resulting olefin analogue, opening up new avenues for
reactivity. Furthermore, the hitherto undocumented coordination of B=B bonds to group 10 metals known for their catalytic performance in olefin functionalization is promising for future applications in catalytic diborene functionalization reactions.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: carbenes · conjugation · density-functional calculations · rearrangements · structure elucidation

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See Supporting Information for synthetic details and solid-state structures of (IbPr)BCat(C(R)≡CHR) and (IbPr)Br[Cr(C≡CHR)] precursors.
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