# The Simplest Amino-borane $\mathbf{H}_{\mathbf{2}} \mathbf{B}=\mathbf{N H}_{\mathbf{2}}$ Trapped on a Rhodium Dimer: Pre-Catalysts for Amine-Borane Dehydropolymerization 

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#### Abstract

The u-amino-borane complexes $\left[R h_{2}\left(L^{R}\right)_{2}(\mu-H)\left(\mu-H_{2} B=N H R^{\prime}\right)\right]\left[B A r^{F}{ }_{4}\right] \quad\left(L^{R}=R_{2} P\left(\mathrm{CH}_{2}\right)_{3} P R_{2}\right.$; $\left.R=P h,{ }^{i} P r ; R^{\prime}=H, M e\right)$ form by addition of $H_{3} B \cdot N M e R^{\prime} H_{2}$ to $\left[R h\left(L^{R}\right)\left(\eta^{6}-C_{6} H_{5} F\right)\right]\left[B A r^{F}\right]$. DFT calculations demonstrate that the amino-borane interacts with the Rh centers through strong Rh-H and Rh-B interactions. Mechanistic investigations show that these dimers can form by a boronium-mediated route, and are pre-catalysts for amine-borane dehydropolymerization, suggesting a possible role for bimetallic motifs in catalysis.


Polyamino-boranes $\left(\left[\mathrm{H}_{2} \mathrm{BNRH}\right]_{n}\right)$ are potentially exciting new materials that are isoelectronic with technologically pervasive polyolefins, but are chemically distinct because of $(\delta-) \mathrm{HB}-\mathrm{NH}(\delta+)$ polarization. They are formed by the dehydropolymerization of amine-boranes $\left(\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NRH}_{2} ; \mathrm{R}=\right.$ H or Me, for example; Scheme 1A), ${ }^{[1]}$ and metal-catalyzed routes to polyamino-boranes offer the potential for fine control over molecular weight and polymer stereochemistry.


Scheme 1. A) Amine-borane dehydropolymerization; B) a suggested coordination/insertion mechanism, $\mathrm{P}=$ polymer chain; C ) examples of $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}$ coordinated to a metal center.

[^0]There is recent evidence that these processes occur at a metal center in which the catalyst needs to perform two roles: 1) formal dehydrogenation of amine-borane to form a latent source of amino-borane $\left(\mathrm{H}_{2} \mathrm{~B}=\mathrm{NRH}\right)$, and 2) subsequent $\mathrm{B}-\mathrm{N}$ bond formation. ${ }^{[2-6]}$ For some systems a coordination/ insertion mechanism is proposed, although the precise structure of the propagating species is currently unresolved (Scheme 1B). ${ }^{[3,5,6]}$ This is in contrast to olefin polymerization, in which the feedstock (for example, ethene or propene) is already unsaturated, and the active species and propagating mechanisms are well-defined. ${ }^{[7]}$ A clearer understanding of how the catalyst dehydrogenates amine-borane, traps intermediate amino-boranes, and promotes $\mathrm{B}-\mathrm{N}$ bond-formation, is central to harnessing the full potential of systems that ultimately deliver new well-defined $\mathrm{B}-\mathrm{N}$ polymeric materials on a useful scale.

Unlike ethene $\left(\mathrm{H}_{2} \mathrm{C}=\mathrm{CH}_{2}\right)$, which is stable under ambient conditions, the isoelectronic amino-borane $\left(\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}\right)$ has only been prepared in low temperature matrices and oligomerizes above $-150^{\circ} \mathrm{C} .^{[2,8]}$ Adding steric bulk to the nitrogen atom increases stability, so that, for example, $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NMeH}^{[9]}$ or $\mathrm{H}_{2} \mathrm{~B}=\mathrm{N}^{t} \mathrm{BuH}^{[10]}$ can be observed as transient species using in situ NMR spectroscopy before they also oligomerize. There are two examples where unstable $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}$ can be trapped by coordination to a single metal center. These originate after dehydrogenation of a putative $\sigma$-ammonia borane ${ }^{[11]}$ complex, forming $\mathrm{Ru}\left(\mathrm{PCy}_{3}\right)_{2}(\mathrm{H})_{2}\left(\eta^{2}-\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}\right) \mathbf{A}^{[12]}$ and (Cy-PSiP)-$\left.\mathrm{Ru}(\mathrm{H})\left(\eta^{2}-\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}\right) \mathbf{B}, \mathrm{Cy}-\mathrm{PSiP}=\kappa^{3}-\left(\mathrm{Cy}_{2} \mathrm{PC}_{6} \mathrm{H}_{4}\right)_{2} \mathrm{SiMe}\right) .{ }^{[13]}$

We now report that $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}$ can be trapped by a bimetallic $\left[\mathrm{Rh}_{2}\left(\mathrm{R}_{2} \mathrm{PCH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{PR}_{2}\right)_{2}\right]^{2+}$ fragment to give a novel bridging amino-borane bonding motif. We provide mechanistic evidence for formation of the complex from a monometallic precursor, and show that such dimeric aminoborane species may be important in dehydropolymerization pathways. This report builds upon previous observations that indirectly implicate bimetallic motifs during catalysis. ${ }^{[14-16]}$

Addition of a slight excess of $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}$ to a [ $\left.\mathrm{D}_{8}\right]$ THF solution of $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{Ph}}\right)\left(\eta^{6}-\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{~F}\right)\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right] \quad \mathbf{1}$ $\left(\mathrm{L}^{\mathrm{Ph}}=\mathrm{Ph}_{2} \mathrm{P}\left(\mathrm{CH}_{2}\right)_{3} \mathrm{PPh}_{2}, \quad \mathrm{Ar}{ }^{\mathrm{F}}=3,5-\left(\mathrm{CF}_{3}\right)_{2} \mathrm{C}_{6} \mathrm{H}_{3}\right) \quad$ resulted in the rapid formation of a bimetallic monocation, which was identified by NMR spectroscopy, electrospray ionization mass spectrometry (ESI-MS), and single-crystal X-ray diffraction, as $\left[\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{Ph}}\right)_{2}(\mu-\mathrm{H})\left(\mu-\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}\right)\right]\left[\mathrm{BAr}_{4}{ }_{4}\right]$ 3. One equivalent of the boronium ${ }^{[9,17-20]}$ cation [THF• $\left.\mathrm{BH}_{2} \cdot \mathrm{NH}_{3}\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]$ was also formed $\quad\left(\delta\left({ }^{(11} \mathrm{B}\right) \quad 0.5 \quad(\mathrm{t}), \quad J_{\mathrm{BH}}=108 \mathrm{~Hz} ; \quad\right.$ lit. ${ }^{[19]}$ $\left.\left[\mathrm{Et}_{2} \mathrm{O} \cdot \mathrm{BH}_{2} \cdot \mathrm{NH}_{3}\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right] \delta\left({ }^{11} \mathrm{~B}\right) 0.2, J_{\mathrm{BH}}=125 \mathrm{~Hz}\right)$.

In situ solution NMR data for $\mathbf{3}$ show a signal at $\delta\left({ }^{11} \mathrm{~B}\right)$ 51.5, a single ${ }^{31} \mathrm{P}$ environment $\left(\delta\left({ }^{31} \mathrm{P}\right) 18.2, J_{\mathrm{RhP}}=142 \mathrm{~Hz}\right)$, and a broad peak at $\delta\left({ }^{1} \mathrm{H}\right)-7.45$ (integral ca. 3 H relative to the
phenyl groups). ESI-MS shows a mono-cation at $m / z=1060.16$ (calcd 1060.16) with the correct isotope pattern. Crystallization (THF/pentane/ $-18^{\circ} \mathrm{C}$ ) gave a small number of crystals, for which a single-crystal X-ray diffraction study showed a $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}$ unit bridging a $\left\{\left(\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{Ph}}\right)_{2}(\mu-\mathrm{H})\right\}\right.$ unit (Supporting Information, Figure S21). However, insufficient material was obtained upon which to collect reliable NMR data. Complex $\mathbf{3}$ is unstable in solution at room temperature, decomposing after four hours to give a mixture in which $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{Ph}}\right)(\mathrm{THF})_{2}\right]\left[\mathrm{BAr}_{4}{ }_{4}\right] \mathbf{6}$ was present in approximately $30 \%$ yield. ${ }^{[21]}$ To put the structure and spectroscopic data on a firm footing, the equivalent reaction using the ${ }^{i} \mathrm{Pr}$ substituted chelating phosphine gave complex 4, $\left[\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{iPr}}\right)_{2}(\mu-\mathrm{H})\left(\mu-\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}\right)\right]\left[\mathrm{BAr}_{4}^{\mathrm{F}}\right]$, and 5 (Scheme 2).


Scheme 2. Formation of amino-borane coordinated dimers 3 and 4. $\left[\mathrm{BAr}_{4}\right]^{-}$anions are not shown.

This reaction was slower than that observed for $\mathrm{L}^{\mathrm{Ph}}$. Complex 4 can also be isolated in $78 \%$ yield as orange crystalline material using an alternative route (see below, Scheme 5). In the absence of $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}$, complex 4 is stable for at least two days in $\left[\mathrm{D}_{8}\right]$ THF solution. However, when formed in situ $\mathbf{4}$ decomposes over 24 hrs into a mixture of products, one of which can be characterized as $\left[\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{iPr}}\right)_{2}(\mathrm{H})_{2}(\mu-\mathrm{H})_{3}\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]{ }^{[22]}$ The room temperature solution NMR data obtained for $\mathbf{4}$ are very similar to those for $\mathbf{3}$ : $\delta\left({ }^{11} \mathrm{~B}\right) 51.1 ; ~ \delta\left({ }^{31} \mathrm{P}\right) 40.8, J_{\mathrm{RhP}}=142 \mathrm{~Hz} ; \delta\left({ }^{1} \mathrm{H}\right)-8.64(3 \mathrm{H}$, broad). Progressive cooling to 180 K splits the high field hydride resonance into two signals, in a 2:1 ratio; while two ${ }^{31} \mathrm{P}$ environments were also observed, suggesting a fluxional process at room temperature. An Eyring plot yields the activation data: $\Delta H^{+}=31.1 \pm 1.3 \mathrm{~kJ} \mathrm{~mol}^{-1}, \quad \Delta S^{\ddagger}=$ $-27 \pm 1 \mathrm{~J} \mathrm{~K}^{-1} \mathrm{~mol}^{-1}, \quad \Delta G(298 \mathrm{~K})^{\ddagger}=39.2 \pm 1.6 \mathrm{~kJ} \mathrm{~mol}^{-1} ;$ where the negative entropy of activation suggests an intramolecular process (Supporting Information, Figures S2-3).

The solid-state structure of complex 4 is shown in Figure 1 A . A dimeric $\mathrm{Rh}_{2}$ unit is accompanied by one $\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]^{-}$ anion, confirming that it is a mono-cation. Two $\left\{\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right)\right\}^{+}$ fragments are bridged by a hydride and a $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}$ unit. The $\mathrm{B}-\mathrm{N}$ distance $(1.377(6) \AA$ ) is consistent with a significant $\mathrm{B}-\mathrm{N} \pi$-interaction, and is similar to that measured in $\mathbf{A}$ (1.396(3) $\AA$ ) and B (1.359(8) $\AA)$, as well as the bridging borylene complex C (1.399(3) Å; Scheme 3). ${ }^{[23]}$ The Rh $\cdots$ B distances (2.070(5) and 2.055(5) $\AA$ ) are similar to those found in the amino-borane complexes $\mathbf{A}, \quad \mathbf{B}$, and $\left[\operatorname{Ir}\left(\mathrm{PCy}_{3}\right)_{2}(\mathrm{H})_{2}\left(\mathrm{H}_{2} \mathrm{~B}=\mathrm{NMe}_{2}\right)\right]\left[\mathrm{BAr}_{4}{ }^{\mathrm{F}}{ }^{[24]}\right.$ (spanning 1.956(2) to $2.140(13) \AA$ ), but significantly shorter than those measured in the bridging thexylborohydride complex $\mathbf{D}(2.330(3) \AA) .{ }^{[25]}$ The hydrogen atoms were located but refined using a riding model. Within the limits of X-ray diffraction the $\mathrm{B}-\mathrm{H}$

(B)
(C)



Figure 1. Solid-state structure of the cationic portion of complex 4. Displacement ellipsoids are shown at the $50 \%$ probability level. Selected bond distances ( $\AA$ ) and angles ( ${ }^{\circ}$ ): Rh1 $\ldots$ Rh2, $2.7874(4)$; Rh1-B1, 2.070(5); Rh2-B1, $2.055(5) ; \mathrm{B} 1-\mathrm{N} 11.377(6)$; P1-Rh1, 2.2550(10); P2-Rh1, 2.3063(10); Rh1-H1, 1.718; Rh2-H2, 1.723; $\Varangle$ plane ( N 1 B 1 H 1 H 2 )/plane ( N 1 B 1 Rh 1 Rh 2 ), 54.1; $\Varangle$ plane (Rh1P1P2)/ plane (Rh2P3P4), 100.2; $\Varangle\left(\mathrm{NH}_{2}\right) /\left(\mathrm{BH}_{2}\right) 24.3^{\circ}$.



Scheme 3. Limiting valence bond descriptions for complex 4, and examples of bridging hydridoborate and borylene complexes. $[R h]=\left\{\operatorname{Rh}\left(L^{\mathrm{iPr}}\right)\right\}$, charge not shown.
distances suggest lengthened, but unbroken bonds (for example, $1.360 \AA$ ). The $\mathrm{NH}_{2}$ group is slightly twisted with respect to the $\mathrm{BH}_{2}$ group ( $24.3^{\circ}$; Figure 1 B ). The whole $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}$ fragment lies $54.1^{\circ}$ from the $\mathrm{Rh}-\mathrm{Rh}$ vector so as to accommodate appropriate overlap between the $\mathrm{B}-\mathrm{H}$ bonds and the two rhodium centers. These are best described as being two distorted square planes (for example, P1/P2/H3/ H1) twisted with respect to one another by $102^{\circ}$ (Figure 1C). This motif, which is similar to that observed for $\mathbf{D}$, is fully consistent with the low temperature NMR data, and are recreated well in the DFT calculated structure (Supporting Information, Figures S24-26). Each metal center in $\mathbf{4}$ is best described as $\mathrm{Rh}^{\mathrm{I}}$, with no $\mathrm{M}-\mathrm{M}$ bond. ${ }^{[26]}$ The end-on $\left\{\mathrm{Rh}_{2}\left(\mu-\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}\right)\right\}$ binding mode contrasts with $\mathrm{H}_{2} \mathrm{C}=\mathrm{CH}_{2}$ that bridges two metal centers symmetrically using both
carbon atoms, in either $\mu-\eta^{2}: \eta^{2}$ or $\mu-\eta^{1}: \eta^{1}$ bonding modes, ${ }^{[27,28]}$ highlighting the differences between these isosteres. ${ }^{[29]}$

Surprisingly, the amino-borane in $\mathbf{4}$ is quite strongly bound. It is only slowly displaced by excess acetonitrile (7\% in 50 min ) to give a mixture of species, one of which is $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right)(\mathrm{NCMe})_{2}\right]\left[\mathrm{BAr}_{4}\right] \cdot{ }^{\mathrm{F}} \cdot{ }^{[22]}$ No reaction occurs with toluene, which might be expected to form a $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right)\left(\eta^{6}-\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{Me}\right)\right]^{+}$complex if a monomeric $\left\{\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right)\right\}^{+}$ fragment were accessible. ${ }^{[30]}$ Addition of cyclohexene, shown to be a probe for free $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2},{ }^{[2]}$ gave no reaction. In contrast, $\mathrm{H}_{2}$ rapidly reacts with $\mathbf{4}$ to form $\left[\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{iPr}}\right)_{2}(\mathrm{H})_{2}(\mu-\mathrm{H})_{3}\right]\left[\mathrm{BAr}_{4}{ }_{4}\right] .{ }^{[22]}$

There are two limiting forms for the structure of 4 (and quasi-isostructural 3): 1) a bridging amino-borane at two $\mathrm{Rh}^{\mathrm{I}}$ centers, or 2 ) a bridging borylene dihydride $\left(\mathrm{Rh}^{\mathrm{III}}\right)$, Scheme 3 . The observed $\delta\left({ }^{11} \mathrm{~B}\right)$ chemical shift of 51 ppm is more consistent with the former as amino-boranes bound to one metal center show chemical shifts around $40-50 \mathrm{ppm},{ }^{[12,13,24,31]}$ while bridging borylenes ${ }^{[32]}$ are generally observed between 90 and 100 ppm. ${ }^{[23,33]}$

To probe the bonding of the amino-borane ligand in $\mathbf{4}$, DFT calculations were used as the basis for a Quantum Theory of Atoms in Molecules (QTAIM) analysis of the total electron density. The results are presented in Figure 2 A , along with selected bond critical point (BCP) metrics. Figure 2 B provides comparative BCP data for the bridging borylene complex $\mathbf{C}$, the hydridoborate complex $\mathbf{D}$, and $\left[\left(\mathrm{PPh}_{3}\right)_{2} \mathrm{Rh}(\mathrm{H})(\mu-\mathrm{H})(\mu-\mathrm{Cl})_{2} \mathrm{Rh}(\mathrm{H})\left(\mathrm{PPh}_{3}\right)_{2}\right]^{+}, \quad \mathbf{E}, \quad$ a welldefined $\mathrm{Rh}^{\text {III }}$ dimer with both terminal and bridging hydrides. ${ }^{[34]}$ Average data are presented for all complexes where appropriate, although the discussion will focus on the bonding around a single rhodium center (Rh1).

In 4 , the $\{\mathrm{Rh} 1 / \mathrm{B} 1 / \mathrm{H} 1\}$ moiety displays bond paths between all three centers, and these enclose a ring critical point (RCP). Thus, 4 has direct Rh1-B1 and Rh1-H1 bonding interactions, while the $\mathrm{B} 1-\mathrm{H} 1$ bond is also intact. Comparison with the Rh1-B1 interaction in $\mathbf{C}$ provides similar $\rho(r)$ and $H(r)$ values, but highlights a much reduced bond ellipticity $(\varepsilon)$ of 0.08 ; this low value indicates dominant $\sigma$-bond character, whereas the value of 0.47 in 4 reflects the asymmetry introduced by the B1-H1 unit. In $\mathbf{D}$, the absence of $\mathrm{Rh}-\mathrm{B}$ BCPs confirms a lack of any direct Rh-B interaction, and this also reduces the average ellipticity of the Rh1-H1 and $\mathrm{B} 1-\mathrm{H} 1$ bonds. Also noticeable are the higher values of $p(r)$ and $H(r)$ for the terminal $\mathrm{B} 1-\mathrm{H} 4$ bond in $\mathbf{D}$ compared to the bridging $\mathrm{B}-\mathrm{H}$ bonds in both that structure and, in particular, $\mathbf{4}$, all of which is consistent with a weakening of the latter. For $\mathbf{E}$, the Rh1-H1 BCP has larger values for $\rho(r)$ and $H(r)$ than the Rh1-H1 BCP in 4, as well as a minimal $\varepsilon$ value. These data indicate a terminal $\mathrm{Rh}-\mathrm{H} \sigma$-bond and stress the differences in bridging character of H 1 and H 2 in 4. BCP data for the Rh1-H3-Rh2 bonds in $\mathbf{4 , D}$, and $\mathbf{E}$ are very similar, suggesting that this moiety varies little across these three systems.

Taken together, the QTAIM analyses suggest that $\mathbf{4}$ is best described as a $\mu$-amino-borane $\mathrm{Rh}^{\mathrm{I}}$ species; a $\mu$-borylene hydride $\mathrm{Rh}^{\text {III }}$ formulism can certainly be ruled out in light of the intact $\mathrm{B} 1-\mathrm{H} 1 / \mathrm{B} 1-\mathrm{H} 2$ bonds and the lack of $\mathrm{Rh} 1-\mathrm{H} 1 /$ $\mathrm{Rh} 2-\mathrm{H} 2$ terminal hydride character. The $\mu$-amino-borane


Figure 2. A) Contour plot of the electron density of the central part of 4 presented in the $\{\mathrm{Rh} 1 \mathrm{~B} 1 \mathrm{Rh} 2\}$ plane with projected stationary points, bond paths, bond critical points (BCP; green), and ring critical points (RCP; red) ; the associated table shows selected BCP metrics (a.u.; average data for indicated bonds) and computed $\delta\left({ }^{11} \mathrm{~B}\right)$ chemical shifts. B) Calculated BCP metrics (a.u.; average data for indicated bonds) for comparator complexes $\mathbf{C}$ (including the computed ${ }^{11} \mathrm{~B}$ chemical shift), $\mathbf{D}$ and $\mathbf{E}(\rho(r)=$ electron density, $\nabla \rho(r)=$ Laplacian of electron density, $\varepsilon=$ bond ellipticity, $H(r)=$ local energy density). All geometries are based on the crystallographically determined heavy atom positions with hydrogen atoms optimized with the BP86 functional. For a full summary of parameters see Figures S24-27 and associated Tables in the Supporting Information.
ligand in 4 interacts with the rhodium centers through stretched $\mathrm{B}-\mathrm{H}$ bonds that engage in strong $\mathrm{Rh}-\mathrm{H}$ and $\mathrm{Rh}-\mathrm{B}$ interactions. Further support for this assertion comes from the computed $\delta\left({ }^{11} \mathrm{~B}\right)$ chemical shifts (Figure 2) and the Pipek-Mezey localized orbitals, where a strong bonding interaction spanning all three $\mathrm{Rh} 1, \mathrm{~B} 1$, and H 1 centers was identified (see Figure 3).

The mechanism of the room temperature fluxional process observed for 4 was also probed with DFT calculations and a single transition state was found to account for this process (Scheme 4). This is accessed by cleavage of one (blue) $\mathrm{B}-\mathrm{H}$ bond to give a transition state structure featuring two $\mathrm{Rh}-\mathrm{H}-\mathrm{Rh}$ bridging hydrides; movement of the original (red) $\mathrm{Rh}-\mathrm{H}-\mathrm{Rh}$ hydride into a $\mathrm{Rh}-\mathrm{H}-\mathrm{B}$ bridging position then completes the exchange ( $\mathbf{4}^{\prime}$ ). Repeating this process from $\mathbf{4}^{\prime}$ exchanges a second $\mathrm{B}-\mathrm{H}$ hydrogen (black) into the $\mathrm{Rh}-\mathrm{H}-\mathrm{Rh}$ bridging position $\left(4^{\prime \prime}\right)$. The computed free energy of activation is $55.2 \mathrm{~kJ} \mathrm{~mol}^{-1}$, somewhat higher than the experimental value ( $39.2 \pm 1.6 \mathrm{~kJ} \mathrm{~mol}^{-1}$ ) but still consistent with facile room temperature exchange.

Understanding how bimetallic species such as $\mathbf{3}$ and $\mathbf{4}$ are formed, and subsequently react, is important for delineating


Figure 3. Pipek-Mezey localized orbital, highlighting the bonding interaction of the $\mathrm{B} 1-\mathrm{H} 1$ bond with center Rh1 (see Supporting Information, Figure S28, for details and related orbitals spanning the $\{\mathrm{Rh} 2 \mathrm{~B} 1 \mathrm{H} 2\}$ and $\{\mathrm{Rh} 1 \mathrm{H} 3 \mathrm{Rh} 2\}$ moieties).


Scheme 4. Proposed fluxional process occurring in 4 (and 3). Hydrogen atoms shown by filled circles. See Supporting Information for DFT calculated geometries and energies.
their role in amine-borane dehydrocoupling. The single equivalent of boronium [THF• $\left.\mathrm{BH}_{2} \cdot \mathrm{NH}_{3}\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]$ (5) formed indicates that a hydride abstraction route may be operating, as recently outlined by Conejero and co-workers for the dehydrocoupling of $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMe}_{2} \mathrm{H}$ by cationic $\{\mathrm{Pt}-\mathrm{NHC}\}^{+}$ catalysts, ${ }^{[17]}$ as well as that occurring in cationic $\mathrm{Ru} / \mathrm{Ir}$-systems ${ }^{[35]}$ or with $\left.\mathrm{B}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)\right)_{3}{ }^{[19]}$ We reasoned that a similar process would yield 5 by B-H activation ${ }^{[16]}$ and subsequent attack by THF (Scheme 5), alongside $\left\{\operatorname{Rh}\left(\mathrm{L}^{\mathrm{R}}\right) \mathrm{H}\right\}$


Scheme 5. Mechanism of formation of $\mathbf{3}$ and $\mathbf{4}$ by boronium protonation of neutral dimer $\mathbf{H}$. $(\mathrm{S})=\mathrm{THF}$ or $\mathrm{Et}_{2} \mathrm{O}$. $\left[\mathrm{BAr}_{4}\right]^{-}$anions are not shown.
that would dimerize to give neutral $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{R}}\right) \mathrm{H}\right]_{2}$ (for example, complex $\mathbf{H}$ ). Subsequent protonation ${ }^{[17]}$ by boronium 5 and elimination of $\mathrm{H}_{2}$ would give $\mathrm{H}_{2} \mathrm{~B}=\mathrm{NH}_{2}$ trapped on a rhodium dimer. To test this hypothesis, addition of $\mathbf{5}$ to the neutral dimer is required. $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{Ph}}\right) \mathrm{H}\right]_{2}$ is unknown, and our attempts to prepare it have not been successful. $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right) \mathrm{H}\right]_{2}$ is a known complex, first prepared by Fryzuk in 1989, ${ }^{[36]}$ and addition of one equivalent of the known boronium salt
$\left[\mathrm{Et}_{2} \mathrm{O} \cdot \mathrm{BH}_{2} \cdot \mathrm{NH}_{3}\right]\left[\mathrm{BAr}_{4}{ }_{4}\right]^{[19]}$ to $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right) \mathrm{H}\right]_{2}$ in $\mathrm{Et}_{2} \mathrm{O}$ solvent, resulted in the immediate formation of $\mathbf{4}$ and gas evolution $\left(\mathrm{H}_{2}\right)$, which is consistent with the mechanism shown.

A dimeric species similar to $\mathbf{3}$ was also formed when one equivalent of $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMeH}_{2}$ was added to $\mathbf{1}$ in THF solution. This was characterized by in situ NMR spectroscopy and ESI-MS as $\left[\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{Ph}}\right)_{2}(\mu-\mathrm{H})\left(\mu-\mathrm{H}_{2} \mathrm{~B}=\mathrm{NMeH}\right)\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right] \quad 8$ : $\delta\left({ }^{1} \mathrm{H}\right) \quad-6.84 ; \quad \delta\left({ }^{31} \mathrm{P}\left\{{ }^{1} \mathrm{H}\right\}\right) \quad 22.2, \quad 21.5 ; \quad \delta\left({ }^{11} \mathrm{~B}\right) \quad 50.6 .{ }^{[21]}$ [THF $\left.\cdot \mathrm{BH}_{2} \cdot \mathrm{NMeH}_{2}\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]$ was also formed $\left(\delta\left({ }^{11} \mathrm{~B}\right) 2.8(\mathrm{t})\right.$, $J_{\mathrm{HB}}=123 \mathrm{~Hz}$; lit. $\mathrm{Et}_{2} \mathrm{O}$ adduct $\delta\left({ }^{11} \mathrm{~B}, \mathrm{CD}_{2} \mathrm{Cl}_{2}\right) 1.7(\mathrm{t}), J_{\mathrm{HB}}=$ $\left.121 \mathrm{~Hz}^{[9]}\right)$. A more complex mixture of species was formed with $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMe}_{2} \mathrm{H}$, suggesting steric factors may be important in the formation of these aminoborane dimers, although a signal observed at $\delta\left({ }^{11} \mathrm{~B}\right) 52.7$ suggests dimer formation. Complexes 3, 4, and $\mathbf{8}$ presumably form via a $\sigma$-complex $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{R}}\right)\left(\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NRH}_{2}\right)\right]\left[\mathrm{BAr}_{4}\right], \mathrm{R}=\mathrm{H}(\mathbf{F}$ Scheme 5) or Me. In THF solution, using the $\mathrm{L}^{\mathrm{Ph}}$ ligand, these $\sigma$-complexes were not observed as boronium formation and subsequent formation of $\mathbf{3}$ is fast. For $L^{\mathrm{iPr}}$, an intermediate $\sigma$-complex could be observed on the way to $4,\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right)\left(\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}\right)\right]\left[\mathrm{BAr}_{4}^{\mathrm{F}}\right]$, presenting NMR data consistent with structure $\mathbf{F}$. ${ }^{[21]}$ Using $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMe}_{3}$ (in which the $\mathrm{N}-\mathrm{H}$ bonds are absent) $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right)\left(\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMe}_{3}\right)\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]$ (7) was isolated and structurally characterized, confirming the in situ NMR studies (Supporting Information, Figure S23). The rapid reaction of $\left[\mathrm{Et}_{2} \mathrm{O} \cdot \mathrm{BH}_{2} \cdot \mathrm{NH}_{3}\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]$ with $\left[\mathrm{Rh}\left(\mathrm{L}^{\mathrm{iPr}}\right) \mathrm{H}\right]_{2}$ to form 4 suggests protonation is not slow for this system; currently we cannot determine whether $\mathrm{B}-\mathrm{H}$ activation or boronium formation is the rate limiting process, although it is likely that either could be promoted by excess amine-borane via $\mathrm{N}-\mathrm{H} \cdots \mathrm{H}-\mathrm{B}$ interactions. ${ }^{[37]}$ Calculations on the $\{\mathrm{Pt}-\mathrm{NHC}\}^{+} / \mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMe}_{2} \mathrm{H}$ system suggest boronium formation is rate limiting. ${ }^{[17]}$

Complex 1 ( $0.5 \mathrm{~mol} \%$, THF, 3 hrs , open system) promoted the dehydrocoupling of $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}$ (1.2 equiv of $\mathrm{H}_{2}$ evolved by gas burette; Supporting Information, Figures S4-S7) to form oligomeric species such as $B$-(cyclotriboraza-nyl)amine-borane (BCTB), ${ }^{[3,38]}$ and insoluble polyaminoborane. ${ }^{[3]}$ With more soluble $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMeH}_{2}$, polymethyla-mino-borane was formed $\left[\mathrm{H}_{2} \mathrm{BNMeH}\right]_{n}$, which was isolated by precipitation from hexanes ( $M_{w}=30600 \mathrm{~g} \mathrm{~mol}^{-1}, ~ Đ=2.6$ ), alongside $\mathrm{H}_{2}$ (1.1 equiv, gas burette). Consistent with the rapid formation of dimers such as $\mathbf{8}$ in THF, no induction period was observed (as measured by $\mathrm{H}_{2}$ evolution) and similar TOF values were recorded (ca. $200 \mathrm{hr}^{-1}$ for 1 equiv $\mathrm{H}_{2}$ ), starting from monomeric $\mathbf{1}$ or in situ formed dimeric $\mathbf{8}$ (Scheme 6). ${ }^{[39]}$ Changing the solvent to non-nucleophilic $1,2-\mathrm{F}_{2} \mathrm{C}_{6} \mathrm{H}_{4}$, and using $\mathbf{1}$ or in situ generated $\mathbf{8}$ as a catalyst, did not present an induction period and also revealed a faster TOF (for $\mathbf{8}$, ca. $1000 \mathrm{hr}^{-1}$ with 1 equiv of $\mathrm{H}_{2}$ released). ${ }^{[40]}$ Sub-catalytic in situ experiments in this solvent ${ }^{[21]}$ show that dimer 8, $\left[\left(\mathrm{BH}_{2}\right)_{2} \mathrm{NMeH}(\mu-\mathrm{H})\right]$ and boronium $\left[\left(\mathrm{NH}_{2} \mathrm{Me}\right)_{2} \mathrm{BH}_{2}\right]\left[\mathrm{BAr}_{4}{ }^{\mathrm{F}}\right]$ are present; ${ }^{[44]}$ the latter is suggested to arise from $\mathrm{NMeH}_{2}$ formed from $\mathrm{B}-\mathrm{N}$ bond cleavage in $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMeH}_{2} \cdot{ }^{[17]}$ Thus, it is likely that similar active species are present in THF or $1,2-\mathrm{F}_{2} \mathrm{C}_{6} \mathrm{H}_{4}$. The lack of induction period is in direct contrast to xantphos-based rhodium catalysts, which show induction periods for $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMeH}_{2}$ dehydrocoupling in $\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{~F}^{[5,15]}$ suggesting that a different kinetics regime or mechanism is in operation.


Scheme 6. $\mathrm{H}_{2}$ evolution experiments using 1 or 8 , and $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMeH}_{2}$ ( $0.5 \mathrm{~mol} \%$ [Rh], 0.41 m amine-borane, THF, 298 K ). [ $\left.\mathrm{BAr}^{\mathrm{F}}\right]^{-}$anions are not shown.

Determination of the resting state in catalysis was hampered by the addition of excess amine-borane $\left(\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}\right.$ or $\left.\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMeH}_{2}\right)$ to the preformed dimeric species $\mathbf{3}$ or $\mathbf{4}$ in THF, resulting in a mixture of products that have been resistant to characterization. Turning to the pure and well-characterized dimer 4, initial rate measurements in a closed system ( $4 \mathrm{~mol} \%$ rhodium, THF) were more informative, and a first-order dependence for either $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}$ or $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NMeH}_{2}$, as well as catalyst 4 , were measured for the early pseudo zero-order phase of catalysis (Supporting Information, Figures S19 and 20). Such behavior is not consistent with a rapid dimer-monomer equilibrium for which an order of $[4]^{1 / 2}$ would be expected, ${ }^{[22,36,42]}$ a view supported by the stoichiometric reactions with acetonitrile or toluene (see above). Under these conditions complexes 2 or $\mathbf{4}$ do not promote full conversion of amine-borane (for 4, 70\% conversion of $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}$ after 10 hrs ). Informed by the subcatalytic experiments and $\mathrm{H}_{2}$ addition studies, we propose that $\left[\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{iPr}}\right)_{2}(\mathrm{H})_{2}(\mu-\mathrm{H})_{3}\right]\left[\mathrm{BAr}_{4}{ }^{\mathrm{F}}\right]^{[22]}$ is formed during catalysis. Consistent with this hypothesis, isolated $\left[\mathrm{Rh}_{2}\left(\mathrm{~L}^{\mathrm{iPr}}\right)_{2}(\mathrm{H})_{2}(\mu-\mathrm{H})_{3}\right]\left[\mathrm{BAr}^{\mathrm{F}}{ }_{4}\right]$ is a poorer catalyst for $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}$ dehydrocoupling in a sealed system ( $4 \mathrm{~mol} \%$ [Rh], $30 \%$ conversion after 10 hrs ) than both 2 and 4. Interestingly, degassing the closed system restarted catalysis, indicating that inhibition by the $\mathrm{H}_{2}$ formed during dehydrocoupling is partially reversible (Supporting Information, Figure S10). Co-promotion of dehydrocoupling by boronium is discounted, as these studies show that isolated $\mathbf{4}$ is an active pre-catalyst in its absence. Consistent with this statement, dehydrocoupling of $\mathrm{H}_{3} \mathrm{~B} \cdot \mathrm{NH}_{3}$ is not catalyzed by $\left[\mathrm{Et}_{2} \mathrm{O} \cdot \mathrm{BH}_{2} \cdot \mathrm{NH}_{3}\right]\left[\mathrm{BAr}^{\mathrm{F}}\right]$ under the conditions used here ( $0.5 \mathrm{~mol} \%$, THF, $298 \mathrm{~K}, 3 \mathrm{hrs}$ ). ${ }^{[19]}$ Overall, these observations do not let us discriminate between active catalysts derived from dimeric 4 (or 3) or monomeric species that result from irreversible, but fast, consumption of 4 (or $\mathbf{3}$ ), under the conditions of excess amine-borane. ${ }^{[43]}$

The ambiguity surrounding mono/bimetallic catalysis has parallels with xantphos-based amine-borane dehydropolymerization catalysts, where P-C activated phosphido-bridged species are formed that are also active catalysts, in contrast to
the amino-borane-bridged dimers observed here. ${ }^{[15]}$ Deconvoluting these systems under conditions of high amine-borane concentration is thus a significant challenge to address if precise control over the resulting polyamino-borane is to be achieved by metal/ligand design. Nevertheless, the observation of novel and unexpected bridging amino-borane complexes as the first-formed species, offers tantalizing clues as to the nature of the actual catalysts; and also suggests that boronium cations may play a more general role in amineborane dehydrocoupling than generally appreciated. ${ }^{[17,19]}$

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