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Synthesis, structure and bonding of bimetallic bridging borylene and boryl complexes

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ABSTRACT: New synthetic route for the synthesis of diruthenium boryl complexes has been established. Thermolysis of an *arachno*-ruthenaborane, [(Cp**Ru*)₂B₃H₈(CS₂H)] (**1**) (Cp* = η⁵-C₅Me₅) with phenyl acetylene, led to the formation of bridging boryl borylene complex as [(Cp**Ru*)₂(μ-HBS₂CH₂-κ²B:κ²S){μ-B(C₆H₄)C(CH₃)-κ²B:κ²C}] (**2**). In parallel to the formation of **2**, the reaction also yielded [(Cp**Ru*)(μ-H)BH{HC=C(H)Ph}{SC(H)S}] (**3a**) and [(Cp**Ru*)(μ-H)BH(PhC=CH₂){SC(H)S}] (**3b**). To understand the reaction pathways for the formation of **2**, we have thermolyzed **1** in toluene that afforded ruthenium bridging bis(boryl) complex, [(Cp**Ru*)₂(μ-HBS₂CH₂-κ²B:κ²S){μ,η²:η²-SBH}] (**4**) along with *nido*-ruthenathaborane [(Cp**Ru*)₂(Me)-(S₂B₂H₃)] (**5**). *Nido*-**5** is structurally and electronically similar with *nido*-[(Cp**Ru*)₂(S₂C₂Ph₂)] (Cp* = η⁵-C₅Me₄Et), which can be generated from the room temperature reaction of [(Cp**Ru*)₂(μ,η¹:η¹-S₂)(μ,η²:η²-S₂)] with diphenylacetylene. Thus, *nido*-**5** can be defined as a true mimic of organometallic cluster *nido*-[(Cp**Ru*)₂(S₂C₂Ph₂)]. Complex [(Cp**Ru*)₂(μ,η¹:η¹-S₂)(μ,η²:η²-S₂)] (**6**) the Cp* analogue of [(Cp**Ru*)₂(μ,η¹:η¹-S₂)(μ,η²:η²-S₂)] can be isolated from the reaction of Li[BH₂S₃] with [Cp**Ru*Cl₂]₂ along with a diruthenium boryl complex, [(Cp**Ru*)₂(μ,η¹:η¹-S₂)(μ-S₂BH-κ¹B:κ¹S:κ²S)] (**7**) in which the boryl unit (S₂BH) possesses no bulky heterocyclic ligand. Theoretical studies were performed to shed light on the bonding of these borylene and boryl complexes. The theoretical calculations reveal that the stability of these complexes is due to the strong interaction between the borylene and boryl unit with the ruthenium centers.

INTRODUCTION

Boron has enriched the coordination chemistry of both main group and the transition metals as it can form both electron precise (2c-2e) as well as electron deficient (3c-2e) bond/s with itself and other elements.¹ As a result, different types of boron-containing compounds having unusual geometry and diverse reactivity, were reported over the years.²⁻³ Other than practical applications in organic chemistry, boron containing compounds have received significant attention in experimental as well as theoretical chemistry. Discovery of hydroboration reaction by H. C. Brown has surged the development of organoborane reagents.^{4,5} Hydroboration of carbon-carbon multiple bonds using different metal catalysts allowed this field to grow further.⁶ Note that, the introduction of boryl moiety into C=C or C≡C bond using hydroboration reaction is one of the most effective ways to synthesize organoboronates with high regio, chemo, and stereoselectivity.⁷ As a result, valuable metal-based catalysts are developed that offer notable advancement in this field. The group 9 metal-based systems, in particular Rh and Ir complexes, are the most commonly employed catalysts used for such organic transformations.⁸

Although the reaction of alkyne/s with metallaborane/s was recognized in 1978, not many examples are known till date.^{9,10} Grimes *et al.* demonstrated the hydroboration reaction of [(CpCo)B₄H₈] with acetylene that led to the formation of metallacarborane cluster, [(CpCo)₂B₃H₇].^{9a} Subsequently, Fehlner *et al.* examined the reactivity of a series of open clusters with alkynes, for example, *nido*-[(Cp**Ru*)₂(μ-H)₂B₃H₇], *nido*-[(Cp**Rh*)₂B₃H₇] and *arachno*-

[(PMe₃)₂IrH(CO)B₈H₁₂].¹⁰ Interestingly, when the terminal or internal alkynes were treated with *nido*-[(Cp**Ru*)₂(μ-H)₂B₃H₇], this led to the formation of metallacarborane clusters.^{10a} In contrast, the Rh-analogue *nido*-[(Cp**Rh*)₂B₃H₇] acts as alkyne cyclotrimerization catalyst.^{10b} Subsequently, we have demonstrated the utility of *arachno*-[(Cp**Ru*CO)₂B₂H₆] as a catalyst for the cyclotrimerization of various terminal and internal alkynes.^{9c} The usefulness of various borate and borane species of transition metals have also been established that yielded metal-vinylborane and alkeneborane complexes.¹¹ As a result, we were interested to explore the reactivity of *arachno*-ruthenaborane [(Cp**Ru*)₂B₃H₈-(CS₂H)]¹², **1** with various alkynes. Herein, we describe the syntheses, structures and bonding of some unique ruthenium boryl borylene and boryl complexes.

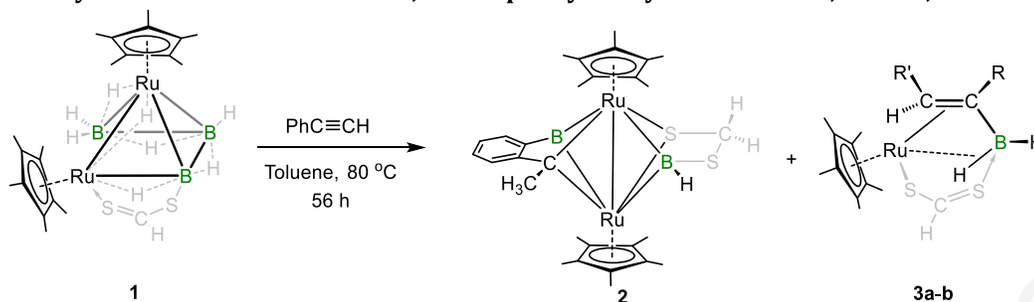
RESULTS AND DISCUSSION

Hydroboration of phenyl acetylene using *arachno*-[(Cp**Ru*)₂B₃H₈(CS₂H)], **1**

Thermolysis of *arachno*-**1** with phenyl acetylene over 56 hours led to the formation of air and moisture sensitive purple solid **2** along with known **3a-b**.^{11d} The reaction also yielded some unstable compounds in very low yields and as a result, the characterizations of these complexes were not possible. Complex **2** was isolated as 23% yield and was characterized by multi-nuclear NMR, IR spectroscopy, mass spectrometry and X-ray diffraction study. The ¹¹B{¹H} NMR spectrum of **2** showed a broad signal at δ = 90.4 ppm. The ¹H NMR spectrum of **2** revealed a chemical shift at δ = 1.36 ppm which is attributed to the methyl protons associated

with Cp* units. The presence of aryl protons and carbon atoms were confirmed from the ^1H chemical shifts at $\delta =$

Scheme 1. Reactivity of *arachno*-ruthenaborane, **1 with phenyl acetylene. **3a**: R = H, R' = Ph; **3b**: R = Ph, R' = H.**



6.52-7.57 ppm and ^{13}C chemical shifts at $\delta = 119.8$ - 128.7 ppm, respectively. Two singlets at $\delta = 4.10$ and 3.59 ppm, observed in the ^1H NMR spectrum of **2** have been assigned to methanedithiolate and another set of methyl protons respectively. The mass spectrometric data of **2** revealed isotopic distribution pattern at m/z 655.0649 corresponding to the molecular ion $\{[\text{M}]^+ + \text{H} - \text{B}_2\text{H}\}$.

The single crystal X-ray diffraction analysis was carried out on a suitable crystal of **2**, obtained by slow evaporation of hexane solution at 2°C . The solid-state X-ray structure of **2**, shown in Figure 1(a), shows the structure as $[(\text{Cp}^*\text{Ru})_2(\mu\text{-HBS}_2\text{CH}_2\text{-}\kappa^2\text{B}:\kappa^2\text{S})\{\mu\text{-B}(\text{C}_6\text{H}_4)\text{C}(\text{Me})\text{-}\kappa^2\text{B}:\kappa^2\text{C}\}]$, in which a boryl ligand, $\{\text{SCH}_2\text{SBH}\}$ and a borylene unit, $\{\text{B}(\text{C}_6\text{H}_4)\text{-C}(\text{Me})\}$ bridged two ruthenium centers in $\{\mu, \kappa^2:\kappa^2\}$ fashion. Although compound **2** has two boron environments, the $^{11}\text{B}\{^1\text{H}\}$ NMR spectrum shows only one signal at $\delta = 90.4$ ppm, which has been assigned to the borylene boron atom based on the similar chemical shift observed for transition metal borylene complexes along with the $^1\text{H}\text{-}^{11}\text{B}\{^1\text{H}\}$ HSQC NMR data and theoretical NMR calculation (Figure S5 and Table S2).¹³ Note that, we did not observe any signal corresponding to the boryl boron in $^{11}\text{B}\{^1\text{H}\}$ NMR spectrum. However, the DFT calculation predicted the ^{11}B chemical shift for the boryl boron for **2** at $\delta = 70.6$ ppm. This calculated value is comparable to the ^{11}B signals of the two boryl groups in **4** (65.9 & 50.9 ppm), but is very different from that of **7** (27.7 ppm). The $^{11}\text{B}\{^1\text{H}\}$ chemical shift at $\delta = 90.4$ ppm is due to the borylene boron which does not associate with any hydrogen. Therefore, any correlation between the $\delta_{\text{B}} = 90.4$ ppm and $\delta_{\text{H}} = 8.70$ ppm has not been observed in the $^1\text{H}\text{-}^{11}\text{B}\{^1\text{H}\}$ HSQC NMR spectrum. The $^1\text{H}\text{-}^{11}\text{B}\{^1\text{H}\}$ HSQC NMR spectrum is an indirect evidence that the ^{11}B chemical shift at $\delta_{\text{B}} = 90.4$ ppm is due to borylene boron.

The Ru-B1 distances (1.93(3) and 2.137 Å) are comparable to that of ruthenium μ -borylene complex $[(\text{Cp}^*\text{Ru})_2(\mu\text{-H})_2(\text{BC}_6\text{Me}_4\text{H})]$ ^{13a} (2.047(4) and 2.049(4) Å). In contrast, the Ru-B2 distances (2.174(15) and 2.195(15) Å) are reasonably shorter as compared to boryl complex $[(\text{Cp}^*\text{Ru})_2(\mu\text{-H})(\mu\text{-CO})(\mu\text{-BCat})]$,¹⁴ (2.26 Å) however, similar to that of $[(\text{Cp}^*\text{Ru})_2(\mu\text{-HBS}_2\text{CH}_2\text{-}\kappa^2\text{B}:\kappa^2\text{S})_2]$ ¹⁵ (2.197 Å). These dissimilar Ru-B bond lengths (Ru-B1 vs Ru-B2) indicate the presence of different boron centers in **2**. Complex **2** is a rare diruthenium complex that feature both the borylene and boryl unit. Gem-diborene species, $[(\text{Br})\text{B}\{(\text{C}_6\text{H}_4\text{CH}_2)\text{PCy}_2\}\text{-B}(\text{PMe}_3)]$ may be considered as one of such species in which both boryl and borylene units are present.¹⁶

It is reasonable to assume that four acute bite angles ($\angle\text{Ru1-B1-Ru}_1$ (74.21°), $\angle\text{Ru1-B2-Ru}_1$ ($68.5(5)^\circ$), $\angle\text{Ru1-S2-Ru}_1$ (61.04°) and $\angle\text{Ru1-C14-Ru}_1$ (74.88°)) bring the Ru centers to close proximity that resulted in a shorter Ru-Ru contact (2.4607(10) Å) as compared to that in *arachno*-**1** (2.9738(3) Å).¹² Note that, a Ru-Ru single bond in **2** is inferred according to the effective atomic number (EAN) rule. From the solid-state structure of **2**, it is very clear that the $\{\text{C}\equiv\text{CH}\}$ unit of phenyl acetylene underwent hydroboration to generate $\{\text{-C-CH}_3\}$ unit, in which the C14-C16 distance of 1.526(19) Å falls within the carbon-carbon single bond distance.^{11d} The B1-C11 bond distance of 1.575(19) Å is comparable to the unperturbed B-C bond distance, signifying the single bond character.¹⁵ Note that, structurally characterized known metal bridging-boryl complexes are very few since after the appearance of first X-ray structure of such complexes in 1990.^{1a} In that context, complex **2** is interesting as the structure is simple and does not require any bulky ligand attached to the boron atom to stabilize the system.

The chemistry of transition metal-boryl complexes has been practised to understand the electronic and steric properties of boryl ligands and their influence on the reaction processes. Further, the involvement of group 9 metal-boryl complexes in metal-catalysed hydroboration were established and recent progress is mostly centred on the utilisation of boryl systems in the catalysis of diboration and C-H functionalization processes.⁷ An outstanding growth in metal-boryl complexes has been achieved by catalysis based on precious heavy metals. Often, Rh- and Ir-based boryl complexes are used as catalysts and remain the most

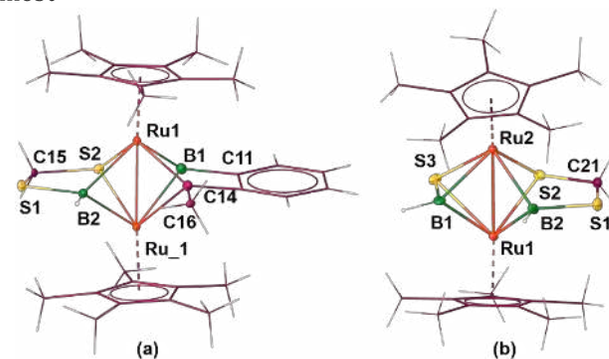


Figure 1. Molecular structure and labelling diagram of **2** (a) and **4** (b): Selected bond lengths (Å) and angles ($^\circ$) are: **2**: C16-C14 1.526(19), B1-C11 1.575(19), C15-S1 1.73(2), C15-S2

1.822(17), Ru_1-S2 2.401(3), Ru_1-B2 2.195(15), Ru1-B2 2.174(15), Ru1-B1 1.93(3), Ru_1-B1 2.137, S1-B2 1.852(17), Ru1-Ru_1 2.4607(10); Ru1-B2-Ru_1 68.5(5); **4**: Ru1-B1 2.111(13), Ru1-Ru2 2.6388(10), Ru1-B2 2.188(10), Ru2-B1 2.129(12), Ru2-B2 2.180(10), C21-S1 1.781(10), C21-S2 1.790(9), B1-S3 1.798(13), B2-S1 1.849(11); Ru2-S2-Ru1 67.94(7), Ru1-S3-Ru2 66.56(6), Ru1-B1-Ru2 77.0(4).

frequently employed catalysts for such transformations.⁸ In particular, the metal-catalyzed borylation of arenes produces functionalized arenes with regioselectivity that is typically determined by steric factors. This process does not require a chelating substituent to trigger catalyst activity.⁷ Although complex **2** does not show any catalytic borylation, intramolecular borylation of the C-H bond of the arene ring is quite unique. In most of the cases, borylation of C-H bonds in arenes occurs by a different pathway in which the C-H bond is cleaved by orthometallation.⁷ In this regard, complex **2** is a notable example where the C-H borylation occurred. We believe that, the CCH₃ moiety is formed via insertion of the alkyne moiety into transiently generated Ru-H bonds. In order to trace the source of hydrogens in the methyl group of {B(C₆H₄)C(Me)} unit in **2** and to get an insight on the formation of this boryl complex, hydroboration of **1** with deuterium labelled phenyl acetylene was carried out. The ¹H and ²H NMR spectra evidently demonstrate that one of the hydrogens of methyl unit in the hydroborated complex (**2^D**) is replaced by deuterium (Scheme S1, Figures S22 and S23) which confirms the possible source of one of the hydrogens in {CCH₃} unit in **2** is the reactant alkyne. We also have tested the reactivity of **1** with other alkynes such as methyl propiolate, dimethyl acetylenedicarboxylate, cyclohexyl acetylene that led to either decomposition of the starting material or afforded no results. Although, we do not have any experimental evidences, looking at the structure of **2** it is reasonable to assume that **2** might have formed via insertion of the C≡C bond into transiently generated Ru-H bonds followed by the borylation of the arene unit.

To gain further insight into the bonding of the boryl borylene species **2**, computational studies were performed at the pbe1pbe/ def2-tzvp level of theory using DFT methods.¹⁷⁻²⁰ The optimized structure of **2** is in well agreement with its solid-state structure. The molecular orbital (MO) analysis of **2** reveals a large HOMO-LUMO gap of 3.234 eV. Based on MO analysis, it is evident that the HOMO of **2** shows the overlap of *p* orbitals of the boron atoms with the *d* orbitals of both Ru atoms (Figure 2(a)). Whereas, the HOMO-11 reveals the overlap of the donut like ring face of the *d_z²* orbitals of both Ru centers signifying strong bonding interaction between two Ru atoms (Figure 2(b)). Furthermore, the Wiberg bond index (WBI) of 0.3753 between Ru centers validates the Ru-Ru bonding interaction.²¹ The natural charge analysis also shows that the B atoms act as donor, whereas the Ru atoms act as acceptor (Table S1). In addition, the Laplacian electron density plot of **2** reveals the bonding scenario of the plane perpendicular to the Ru-Ru bond. As shown in Figure 2(c), an area of charge concentration along the C16-C14 bond is very prominent. The calculated C14-C16 bond length of 1.505 Å (exp. 1.534(17) Å) indicates the presence of a single bond. The WBI value of 1.0433 between C16 and C14 atoms

also indicates single bond interaction, which is further supported by NBO analysis (Figure S29).

Bimetallic bridging bis(boryl) complex, [(Cp*Ru)₂(μ-HBS₂CH₂-κ²B:κ²S){μ,η²:η²-SBH}], **4**

The formation of bridging boryl borylene species, **2** from the reaction of **1** with alkyne was somewhat unusual, specially the hydroboration of alkyne and C-H borylation of

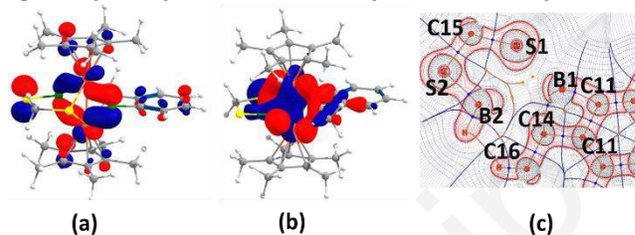


Figure 2. (a) HOMO and (b) HOMO-11 of the boryl borylene species **2**. (c) Laplacian electron density plot of the plane perpendicular to Ru-Ru bond; Solid red lines indicate areas of charge concentration ($\nabla^2\rho(r) < 0$), while dashed black lines show areas of charge depletion ($\nabla^2\rho(r) > 0$). Blue dots indicate bond critical points [BCPs].

the arene ring. From the solid-state X-ray structures of **2** and **1**, it is apparent that they are structurally related as the former (**2**) is a derivation from the latter (**1**) and both the species contain {(Cp*Ru)₂} and {S₂CBH} units. Also, we believe that there must be an intermediate which reacts with the alkyne to produce the hydroborated species **2**. Thus, we carried out the thermolysis of **1** in toluene with and without alkyne. The reaction without alkyne led to the formation of orange **4** and green **5** in 17 and 10% yields respectively. Both the complexes were separated by thin layer chromatography using hexane/CH₂Cl₂ (80:20) as eluents. They have been characterized by multi-nuclear NMR, IR spectroscopy, mass spectrometry and X-ray diffraction studies.

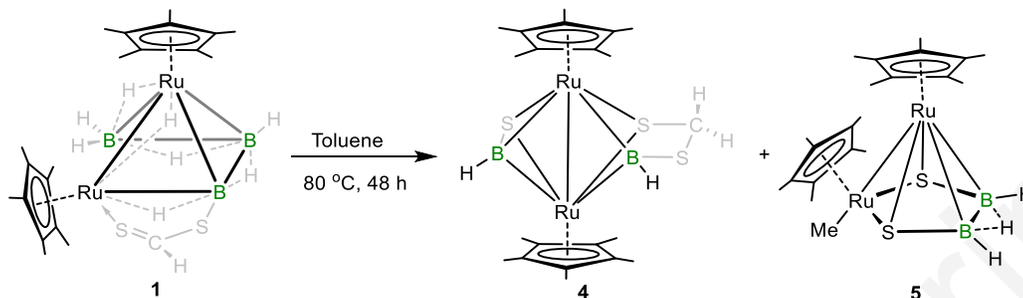
The ¹¹B{¹H} NMR spectrum of **4** showed two resonances at $\delta = 65.9$ and 50.9 ppm which are upfield shifted as compared to **2**. The presence of Cp* ligand in **4** was confirmed from the ¹H chemical shift at $\delta = 1.73$ ppm. This was further supported by the ¹³C{¹H} NMR spectrum. The ¹H NMR spectrum also showed a resonance at $\delta = 6.24$ ppm that is due to the presence of BH₂ proton/s. The mass spectrometric analysis of **4** showed an ionized peak at $m/z = 596.0003$ corresponding to {[M]⁺ + H - BH}. Suitable crystals for **4** were obtained by the slow evaporation of a hexane/CH₂Cl₂ (80:20) solution at 2 °C that allowed us to perform the X-ray diffraction study.

The solid-state X-ray structure of **4**, shown in Figure 1(b), reveals the structure of **4** as a bridging bis(boryl) complex of ruthenium. Two of the boryl units, {HBS} and {HBSCH₂S} are attached perpendicular to {Cp*Ru} units in (μ-κ²:κ²) fashion and the boron atoms reside on the same face unlike in **2**. The avg. B-S bond distance of 1.824 Å is in accord with the sum of the covalent radii of B and S however, shorter as compared to [(Cp*Ru)₂(μ-HBS₂CH₂-κ²B:κ²S)₂] (1.7885 Å).¹⁵ Similarly, the avg. C-S bond distance of 1.785 Å in methanedithiolate unit of **4** is within the single bond distance.^{22,23} The ¹H chemical shift at $\delta = 2.84$ ppm and a resonance at $\delta = 40.3$ ppm in ¹³C{¹H} NMR spectrum confirms the presence of methanedithiolate ligand in **4**. The Ru-Ru bond distance of 2.6388(10) Å in **4** is significantly shorter as compared to the ruthenium oxo-boryl species

$\{[Cp^*Ru(\mu-H)]_3(\mu_3-BO)(\mu_3-H)\}$ (avg. 2.7162 Å).²⁴ This shortening of bond length may be due to the presence of two bridging S atoms that bring the two ruthenium centers closer forming acute angles with them. The average Ru-B

bond distance of 2.152 Å is comparatively shorter as compared to $\{[Cp^*Ru(\mu-HBS_2CH_2-\kappa^2B:\kappa^2S)]_2\}$ ¹⁵ (2.185 Å). The shorter Ru-S bond lengths in $\{HBSCH_2S\}$ unit compared to

Scheme 2. Thermolysis of *arachno*-ruthenaborane, $[(Cp^*Ru)_2B_3H_8(CS_2H)]$, **1.**



that of $\{HBS\}$ unit in **4** suggests that the sulfur atom of former approaches the ruthenium centers more closely as compared to that of latter. Whereas, the boron atom of the $\{HBS\}$ unit approaches the ruthenium centers more closely as compared to $\{HBSCH_2S\}$ unit.

Although it seemed that the boryl species **4** is the intermediate for the formation of **2**, reaction of **4** with phenyl acetylene yielded no traces of **2**. Nonetheless, the availability of this boryl species **4** allowed us to compare its bonding and electronic structure with other diruthenaborane species with the aid of computational studies. The MO analysis reveals that the HOMO-LUMO gap of **4** (4.27 eV) is reasonably larger as compared to that of **2**. Further, the Ru-Ru bonding interaction of **4** is depicted in HOMO-3 where the donut like ring face of the d_{z^2} orbitals are overlapped (Figure 3(a)). The Ru-Ru bonding interaction is further verified by WBI (0.3754) and Laplacian electron density plot (Figure 3(b)). The covalent character of Ru1-Ru2 bond has also been reflected by the positive value of electron density ($\rho_b = 0.0681$) and a negative value of the energy density [$H(r) = -0.0205$] at bcp. Although the natural charges at both the Ru centers (acceptor) of **4** is comparable with that of **2**, the natural charges at B atom of the borylene unit of **2** is significantly higher as compared to boryl borons in **2** and **4** (Table S1).

Along with the formation of **4**, compound **5** was isolated as green solid. The 1H NMR spectrum of **5** showed two chemical shifts at $\delta = 1.68$ and 1.60 ppm due to the methyl protons of Cp^* ligands. Further, the presence of boron was confirmed by ^{11}B chemical shift, appeared at $\delta = 13.5$ ppm. The mass spectrometric analysis revealed an isotropic distribution pattern at m/z 566.0492 $\{[M]^+ + H - BH\}$.

The X-ray diffraction studies, performed on a single crystal of **5**, obtained from slow evaporation of hexane/ CH_2Cl_2 (70:30) solution of **5** at 2 °C, revealed its identity as *nido*- $[(Cp^*Ru)_2(Me)(S_2B_2H_3)]$. The solid-state X-ray structure of *nido*-**5**, shown in Figure 4(a), clearly

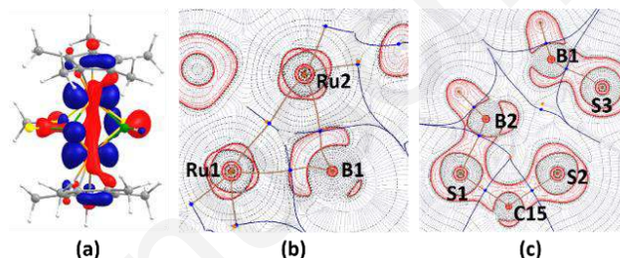


Figure 3. (a) HOMO-3 of **4**; (b) and (c) Laplacian electron density plot of **4** of plane (Ru1-B1-Ru2) and plane perpendicular to Ru-Ru bond, respectively.

displays a $\{S_2B_2H_3\}$ unit, attached to two $\{Cp^*Ru\}$ units. The Ru-Ru bond distance of 2.878(2) Å is considerably shorter as compared to ruthenaborane species, $[(Cp^*Ru)_2(B_3H_9)]$ ²⁵ (2.9689(17) Å). The B-S bond distance of 1.87(2) Å is in agreement with the sum of the covalent radii of B and S atoms.¹⁵ The molecular structure of **5** can be considered as a dimetallaheteroborane analogue of *nido*- $[B_6H_{10}]$, in which one apical BH and one basal BH atom is subrogated by $\{Cp^*Ru\}$ and $\{Cp^*RuMe\}$ fragments respectively and other two basal BH units are subrogated by two S atoms. The core of **5** is similar with ruthenaborane, $[(Cp^*Ru)_2(\mu-H)_2(S_2B_2H_4)]$ ²⁶. Note that one methyl group is attached to one of the Ru atoms in **5** and the chemical shift for this methyl protons appeared at $\delta = -0.20$ ppm in 1H NMR spectrum. The bridging hydrogens as well as terminal hydrogens associated with boron atoms were located crystallographically. They were further confirmed by 1H NMR that shows resonances at $\delta = 3.50$, 3.12 and -4.44 ppm, which are attributed to two BH_t and one B-H-B proton/s respectively. If one considers that $\{Cp^*Ru\}$, $\{B_2H_3\}$, (μ_3-S) and (Me) unit contribute 1, 5, 4 and 1 electrons to the cluster bonding framework respectively,²⁷ then **5** possesses eight skeletal electron pairs (SEP), one SEP lesser than $[(Cp^*Ru)_2(\mu-H)_2(S_2B_2H_4)]$ ²⁶. Therefore, *nido*-**5** is considered as electronically unsaturated species and may be potential to react with electron rich species. As shown in Figure 4(b), *nido*-**5** is analogous to *nido*- $[B_6H_{10}]$ and structurally similar with organometallic analogue $[(Cp^*Ru)_2S_2C_2Ph_2]$.²⁸ The $\{B_2H_3\}$ unit of *nido*-**5** provides 5 skeleton electrons for

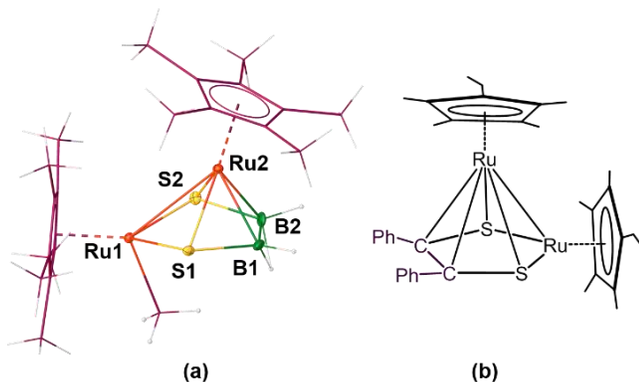
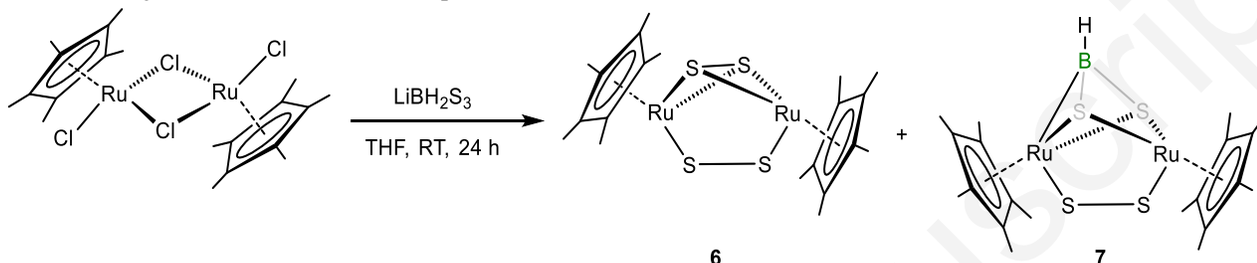


Figure 4. (a) Molecular structure and labelling diagram of **5**: Selected bond lengths (Å) and angles (°) are: Ru1-B1 2.111(13), Ru1-B2 2.188(10), Ru2-B1 2.129(12), Ru2-B2 2.180(10), Ru1-Ru2 2.6388(10), Ru1-S2 2.365(3), C21-S1 1.781(10), C21-S2 1.790(9), B1-S3 1.798(13), B2-S1 1.849(11); S1-C21-S2 113.4(5), Ru1-B1-Ru2 77.0(4). (b) schematic representation of $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$.

Scheme 3. Synthesis of disulfide complexes of ruthenium



cluster bonding, whereas $\{\text{C}_2\text{Ph}_2\}$ moiety of $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$ provides 6 skeleton electrons towards the cluster bonding. The missing one skeleton electron of *nido-5* is achieved from the exo-polyhedral CH_3 group attached to one of the Ru centers and thus becomes electronically similar with $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$. As shown in Figure 5, the HOMO-9 of *nido-5* reveals the bonding orbitals of Ru- CH_3 bond. From the NBO analysis and Laplacian electron density plot, a high electron density is observed along the B-B and B-H-B (3c-2e) bonds of *nido-5* and the C-C bond of $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$ (Figures S30-S33). Further, both theoretical and experimental structural parameters of *nido-5* and $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$ clusters are comparable. Thus, *nido-5* can be defined as a true mimic of organometallic cluster *nido*- $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$. Note that, besides *nido-5*, there are several structurally characterized metallaboranes known that mimic classic organometallic clusters.^{2b}

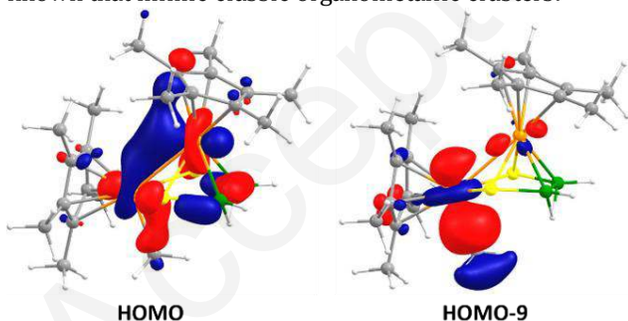


Figure 5. Selected frontier molecular orbitals of *nido-5*.

Diruthenium boryl complex, **7**

Ruthenium disulfide complex $[(\text{Cp}^*\text{Ru})_2(\mu, \eta^1: \eta^1\text{-S}_2)(\mu, \eta^2: \eta^2\text{-S}_2)]$,²⁸ made by Rauchfuss *et al.* in 1999, is a good precursor to synthesize *nido*- $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$ when reacted with phenyl acetylene. Interestingly, the Cp^* analogue of this disulfide species, $[(\text{Cp}^*\text{Ru})_2(\mu, \eta^1: \eta^1\text{-S}_2)(\mu, \eta^2: \eta^2\text{-S}_2)]$, **6** can also be isolated from the room temperature reaction of $\text{Li}[\text{BH}_2\text{S}_3]$ and $[\text{Cp}^*\text{RuCl}_2]_2$. Along with the formation of **6**, the reaction also yielded another

green complex **7**. Both the complexes **6** and **7** were separated using thin layer chromatography with 33 and 19% yields. Complex **6** was characterized in comparison of its spectroscopic data with $[(\text{Cp}^*\text{Ru})_2(\mu, \eta^1: \eta^1\text{-S}_2)(\mu, \eta^2: \eta^2\text{-S}_2)]$,²⁸ and an X-ray diffraction analysis shows **6** as the Cp^* analogue of $[(\text{Cp}^*\text{Ru})_2(\mu, \eta^1: \eta^1\text{-S}_2)(\mu, \eta^2: \eta^2\text{-S}_2)]$.²⁸ The HOMO of **6** displays the donating property through the *p*-orbitals of sulfur atoms and the *d*-orbitals of Ru centers (Figure S27).

Along with the formation of disulfide species **6**, complex **7** was isolated. The $^{11}\text{B}\{^1\text{H}\}$ NMR spectrum shows a signal at $\delta = 27.7$ ppm that suggests the presence of one boron environment. The chemical shift at $\delta = 1.78$ ppm in ^1H NMR spectrum is due to the presence of Cp^* protons. Presence of terminal hydrogen associated with boron atom was confirmed by ^1H NMR spectroscopy ($\delta = 6.72$ ppm). The mass spectrometry (ESI⁺ mode) showed an intense isotopic distribution pattern at m/z 612.9529 $[\text{M}]^+$. Nonetheless, all the spectroscopic data along with mass spectrometric data were not adequate enough to predict the molecular structure of **7**. Therefore, the single crystal X-ray diffraction analysis on a suitable crystal of **7** was necessary to reveal its identity.

The solid-state X-ray structure of **7**, shown in Figure 6, depicts the structure as diruthenium boryl species, $[(\text{Cp}^*\text{Ru})_2(\mu, \eta^1: \eta^1\text{-S}_2)(\mu\text{-S}_2\text{BH}\text{-}\kappa^1\text{B}\text{-}\kappa^2\text{S}\text{-}\kappa^2\text{S}')]_2$, in which one boryl ligand $\{\text{SBHS}\}$ is bridged between two $\{\text{Cp}^*\text{Ru}\}$ units in $(\mu\text{-}\kappa^1: \kappa^2: \kappa^2)$ fashion (Table 1).^{14,15,29} Both the ruthenium centers (Ru1 and Ru2) in **7** are in pseudo-octahedral environments. The boryl ligand binds with both the ruthenium centers orthogonally with the Ru-S-Ru angles of 94.78(2) and 94.42(2)°. The Ru-B bond length of 2.284(3) Å is significantly longer as compared to ruthenium boryl complex $[\text{RuH}(\text{CO})(\text{PPh}_3)_2\{\text{B}(\text{NCH}_2\text{PPh}_2)_2\text{C}_6\text{H}_4\}]$ ^{29c} (2.077(6) Å). The avg. Ru-S bond distance of 2.3751 Å in **7** is within the single bond distance as observed in ruthenium dithioformate complex $[\{\text{Cp}^*\text{Ru}(\mu, \eta^3\text{-SCHS})\}_2]$ ¹⁵ (2.3733 Å). The S-S bond distance of 1.9985(9) Å is significantly shorter

than typical disulfide bond (2.05 Å). The B-S bond distances in **7** (1.809(3) and 1.826(3) Å) are within the range of single bond distance as observed in other transition metal-boron species.^{15,22} This type of ruthenium complex stabilized by simple boryl ligand is unique and not many examples of this kind are known.

To understand the bonding situation in **7**, the computational studies have been performed at the pbe1pbe/def2-tzvp level of theory. The HOMO-LUMO

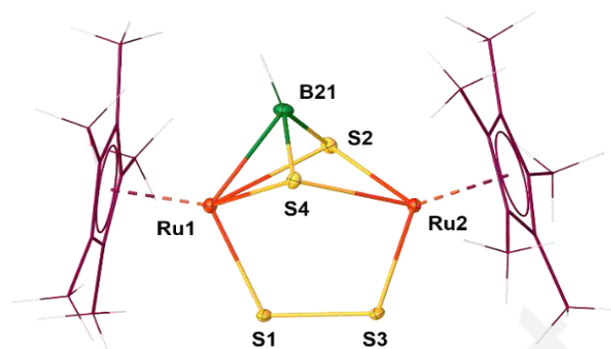
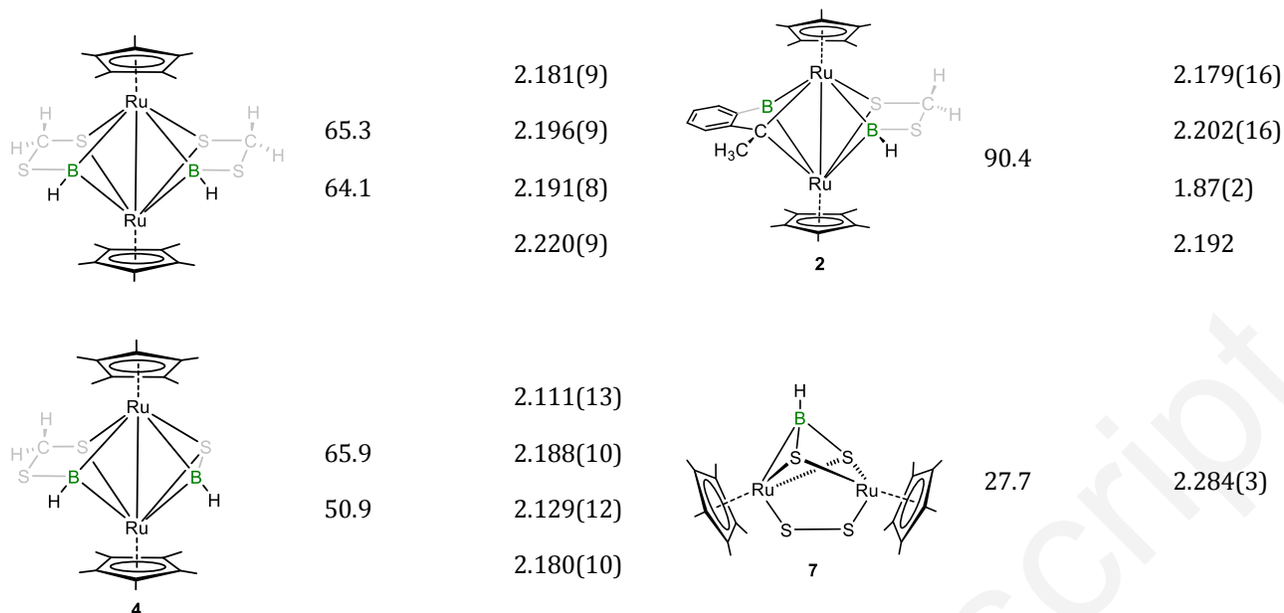


Figure 6. Molecular structure and labelling diagram of **7**. Selected bond lengths (Å) and angles (°) are: Ru1-S1 2.2599(6), Ru1-B21 2.284(3), Ru1-S4 2.4655(6), Ru2-S3 2.1916(6), Ru2-S2 2.4341(6), S1-S3 1.9985(9), B21-S2 1.809(3), B21-S4 1.826(3); S2-B21-S4 105.76(16), S2-B21-Ru1 73.28(10).

Table 1. Selected structural parameters and ¹¹B NMR chemical shifts of ruthenium boryl complexes^{14,15,29}

Boryl complexes	¹¹ B NMR (ppm)	d(M-B) (Å)	Complexes	¹¹ B NMR (ppm)	d(M-B) (Å)
	47.0	2.075(4)		60.3	2.115(2)
	-28.9	2.243(8)		38.2	2.173(3)
	61.3	2.077(6)		44.8	2.022(4)
	53.7	2.230(4) 2.290(4)		43.9	2.307(6)
	110.5	2.137(2)		43.4	2.0424(19)



energy gap for **7** of 2.72 eV is quite smaller as compared to other boryl species **2** and **4**. The HOMO-1, HOMO-12 and HOMO-14 of **7** reveal the bonding scenario of the boryl unit with Ru and S centers (Figures 7a and S28). The Laplacian electron density plot along the Ru1-B21-S4 plane of **7** shows high charge concentrations (Figure 7(b)). In addition, the BCPs between Ru-S, Ru-B and B-S bonds clearly indicate strong bonding interactions (Figure 7(b)). The natural charge at B atom of the boryl unit in **7** is negative, whereas for **2** and **4** the values are positive (Table S1). The negative natural charge of B atom in **7** may be arrived due to the highly positive natural charges of other four sulfur atoms.

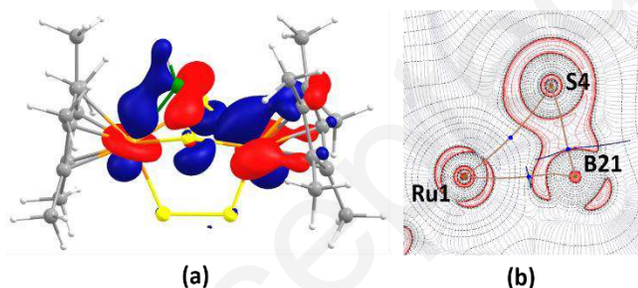


Figure 7. (a) HOMO-1 of **7**. (b) Laplacian electron density plot of **7** along Ru1-B21-S4 plane.

The availability of various structural data for species **2**, **4** and **7** allowed us to perform a structural comparison with known ruthenium boryl species as listed in Table 1.^{14,15,29} In general, the metal center influences the structural parameters of the attached ligands. Therefore, comparing the bond distances between the metal/s and ligand/s in a complex a general conclusion on the role of metal-ligand interaction can be drawn. For example, the Ru-B bond distances in **2** (2.1108 Å) and **4** (2.152 Å) are within the Ru-B single bond distance similar to ruthenium-boryl species, such as $[(\text{Cp}^*\text{Ru}(\mu\text{-HBS}_2\text{CH}_2\text{-}\kappa^2\text{B}:\kappa^2\text{S}))_2]$ ¹⁵ (2.1955 Å) and $[(\text{SiMe}_3)_3(\text{PMe}_3)(\text{CO})_3\text{RuB}(\text{Cl})(\text{C}_6\text{HMe}_4)]$ ^{29h} (2.137(2) Å) (Table 1). However, the Ru-B bond distance of 2.284(3) Å in

7 is considerably longer as compared to **2** and **4**. The bridging {SCH₂S} ligand in **2** and **4** may be accountable to keep the boryl units in close proximity with the metal centres.

The ¹¹B NMR is an important tool for the elucidation of structures of boron complexes.³⁰ Thus, to get some insight into the variations in the attachment and bonding of the ligands in these boryl species, we have analyzed the ¹¹B chemical shift values of the species, listed in Table 1. The ¹¹B chemical shift values of the bridging boryl complex **4** are comparable with that of bridging bis(boryl) complex $[(\text{Cp}^*\text{Ru})_2(\mu\text{-HBS}_2\text{CH}_2\text{-}\kappa^2\text{B}:\kappa^2\text{S})_2]$.¹⁵ This may be due to the presence of a similar type of boryl ligand {HBS₂CH₂}. On the other hand, the ¹¹B chemical shift for boryl ligand { $\mu,\eta^2:\eta^2$ -SBH} in **4** appeared in the region of 50-65 ppm. Further, the downfield ¹¹B chemical shift values for **2** and **4** as compared to **7** may be due to the fact that the boron atom in **7** is connected with two sulfur atoms.

CONCLUSION

In this feature article, we have established some new synthetic routes for different types of bridging borylene and boryl species of ruthenium. Further, we have established the reactivity of *arachno-1* with terminal alkyne that generated new type of bridging boryl borylene complex, which is stabilized at the diruthenium template. During the course of formation of bridging boryl borylene complex, the C-H bond of arene ring underwent borylation, which is very unusual. Based on the skeleton electron counts and the structures, *nido-5* may be considered as analogous to the *nido*-[B₆H₁₀] and $[(\text{Cp}^*\text{Ru})_2\text{S}_2\text{C}_2\text{Ph}_2]$ and it's a true mimic of organometallic counterpart, $[(\text{Cp}^*\text{Ru})_2(\text{S}_2\text{C}_2\text{Ph}_2)]$. Investigations to evaluate the scope of synthesis of boryl complexes concerning early transition metals are underway.

EXPERIMENTAL SECTION

I Experimental Details.

General procedures and instrumentation

All manipulations were conducted under an atmosphere of dry argon or in vacuo using standard Schlenk line or glove box techniques. Solvents (hexane, toluene, thf) were purified by distillation from appropriate drying agents (sodium/benzophenone) under dry argon prior to use. CDCl_3 was degassed by three freeze-pump-thaw cycles and stored over molecular sieves. $[(\text{Cp}^*\text{Ru})_2\text{B}_3\text{H}_8(\text{CS}_2\text{H})]^{12}$, **1** $[\text{Cp}^*\text{RuCl}_2]_2$,³¹ and $\text{Li}[\text{BH}_2\text{S}_3]$,³² were prepared according to literature methods, while phenyl acetylene was obtained commercially and used as received. Thin-layer chromatography was performed on 250 mm aluminum supported silica gel TLC plates. NMR spectra were recorded on 400 and 500 MHz Bruker FT-NMR spectrometers. The residual solvent protons were used as reference (δ , ppm, CDCl_3 , 7.26), while a sealed tube containing $[\text{Bu}_4\text{N}(\text{B}_3\text{H}_8)]$ in d_6 -benzene (δ , ppm, -30.07) was used as an external reference for the $^{11}\text{B}\{^1\text{H}\}$ NMR. For $^{13}\text{C}\{^1\text{H}\}$ NMR spectra, residual solvent carbon was used as a reference (δ , ppm, CDCl_3 , 77.1). Mass spectra were recorded on Qtof Micro YA263 HRMS and 6545 Qtof LC/MS instruments. The infrared spectra were recorded on an Agilent Cary 630 FTIR spectrometer in dichloromethane solvent.

I.1 Synthesis and Characterizations

Synthesis of 2 and 3a-b: In a flame-dried Schlenk tube, the red solution of **1** (0.05 g, 0.084 mmol) and one equivalent of phenylacetylene in toluene (20 mL) were stirred at 80 °C for 56 hours. The solvent was evaporated in vacuum; residue was extracted into hexane/thf (95:5 v/v) and passed through celite. After removal of solvent, the residue was subjected to chromatographic work-up using silica gel TLC plates. Elution with hexane yielded purple **2** (0.013 g, 23%) and orange solids **3a-b** (~18%).

2: MS (ESI⁺) calcd for $\text{C}_{29}\text{H}_{40}\text{Ru}_2\text{S}_2^+ \{[\text{M}]^+ + \text{H} - \text{B}_2\text{H}\} m/z$ 655.0680, found 655.0649; $^{11}\text{B}\{^1\text{H}\}$ NMR (160 MHz, CDCl_3 , 22 °C): δ = 90.4 (br, 1B); ^1H NMR (500 MHz, CDCl_3 , 22 °C): δ = 8.70 (br, 1H; *BH*), 7.57-6.52 (m, 4H; Ph), 4.10 (s, 2H; *CH_2S_2*), 3.59 (s, 3H; *CH_3*), 1.36 ppm (s, 30H; $2 \times \text{Cp}^*$); $^{13}\text{C}\{^1\text{H}\}$ NMR (125 MHz, CDCl_3 , 22 °C): δ = 143.1 (CCH₃), 128.7-119.8 (s, Ph), 91.5 (s, *C_5Me_5*), 51.5 (CH₂S₂), 37.3 (CCH₃), 9.5 ppm (s, *C_5Me_5*); IR (CH₂Cl₂) $\tilde{\nu}$ = 2411 cm⁻¹ (BH₃).

Synthesis of 4 and 5: In a flame-dried Schlenk tube, the red solution of **1** (0.05 g, 0.084 mmol) in toluene (20 mL) was stirred at 80 °C for 48 hours. The solvent was evaporated in vacuum; residue was extracted into hexane/CH₂Cl₂ (70:30 v/v) and passed through celite. After removal of solvent, the residue was subjected to chromatographic work-up using silica gel TLC plates. Elution with a hexane/CH₂Cl₂ (75:25 v/v) mixture yielded orange **4** (0.008 g, 17%) and green **5** (0.005 g, 10%).

4: MS (ESI⁺) calcd for $\text{C}_{21}\text{H}_{34}\text{Ru}_2\text{B}_3\text{S}^+ \{[\text{M}]^+ + \text{H} - \text{BH}\} m/z$ 596.0025, found 596.0003; $^{11}\text{B}\{^1\text{H}\}$ NMR (160 MHz, CDCl_3 , 22 °C): δ = 65.9 (br, 1B), 50.9 ppm (br, 1B); ^1H NMR (500 MHz, CDCl_3 , 22 °C): δ = 6.24 (br, 2H; *BH*), 2.84 (s, 2H; *CH_2S_2*), 1.73 ppm (s, 30H; $2 \times \text{Cp}^*$); $^{13}\text{C}\{^1\text{H}\}$ NMR (100 MHz, CDCl_3 , 22 °C): δ = 94.0 (s, *C_5Me_5*), 40.3 (s, CH₂S₂), 10.2 ppm (s, *C_5Me_5*); IR (CH₂Cl₂) $\tilde{\nu}$ = 2552, 2417 cm⁻¹ (BH₃).

5: MS (ESI⁺) calcd for $\text{C}_{21}\text{H}_{35}\text{Ru}_2\text{B}_2\text{S}_2^+ [\text{M} - \text{BH}]^+ m/z$ 566.0462, found 566.0492; $^{11}\text{B}\{^1\text{H}\}$ NMR (160 MHz, CDCl_3 , 22 °C): δ = 13.5 ppm (br, 2B); ^1H NMR (500 MHz, CDCl_3 , 22 °C): δ = 3.50, 3.12 (br, 2H; *BH*), 1.60, 1.68 (s, 30H; $2 \times \text{Cp}^*$), -0.20 (s, 3H, *CH_3*), -4.44 ppm (br, 1H; *B-H-B*); $^{13}\text{C}\{^1\text{H}\}$ NMR (100 MHz, CDCl_3 , 22 °C): δ = 96.6 (s, *C_5Me_5*), 19.4 (s, CH₃), 11.3, 10.6 ppm (s, *C_5Me_5*); IR (CH₂Cl₂) $\tilde{\nu}$ = 2513(BH₃), 2390 (BH₃) cm⁻¹.

Synthesis of 6 and 7: In a flame-dried Schlenk tube, a suspension of $[\text{Cp}^*\text{RuCl}_2]_2$ (0.05 g, 0.081 mmol) in 10 mL tetrahydrofuran at -78 °C was charged dropwise with a freshly prepared solution of $\text{Li}[\text{BH}_2\text{S}_3]$ in tetrahydrofuran (10 mL, 0.324 mmol, 0.0324 mol/L) over 15 min and kept under constant stirring for 24h at room temperature. The solvent was evaporated in vacuum; residue was extracted into hexane/CH₂Cl₂ (60:40 v/v) and passed through celite. After removal of solvent, the residue was subjected to chromatographic work-up using silica gel TLC plates. Elution with a hexane/CH₂Cl₂ (80:20 v/v) mixture yielded blue **6** (0.016 g, 33%) and green **7** (0.010 g, 19%).

6: MS (ESI⁺) calcd for $\text{C}_{20}\text{H}_{30}\text{Ru}_2\text{S}_4^+ [\text{M}]^+ m/z$ 601.9327, found 601.9351; ^1H NMR (500 MHz, CDCl_3 , 22 °C): δ = 1.89 ppm (s, 30H; $2 \times \text{Cp}^*$); $^{13}\text{C}\{^1\text{H}\}$ NMR (100 MHz, CDCl_3 , 22 °C): δ = 95.3 (s, *C_5Me_5*), 10.9 ppm (s, *C_5Me_5*).

7: MS (ESI⁺) calcd for $\text{C}_{20}\text{H}_{31}\text{Ru}_2\text{S}_4\text{B}^+ [\text{M}]^+ m/z$ 612.9509, found 612.9529; $^{11}\text{B}\{^1\text{H}\}$ NMR (160 MHz, CDCl_3 , 22 °C): δ = 27.7 ppm (br, 1B); ^1H NMR (500 MHz, CDCl_3 , 22 °C): δ = 6.72 (br, 1H; *BH*), 1.78 ppm (s, 30H; $2 \times \text{Cp}^*$); $^{13}\text{C}\{^1\text{H}\}$ NMR (100 MHz, CDCl_3 , 22 °C): δ = 95.3 (s, *C_5Me_5*), 10.6 ppm (s, *C_5Me_5*); IR (CH₂Cl₂) $\tilde{\nu}$ = 2524 cm⁻¹ (BH₃).

I.2. X-ray Analysis Details

Suitable X-ray quality crystals of **2**, **4**, **5**, **6** and **7** were grown by slow diffusion of a hexane-CH₂Cl₂ solution. The crystal data for **2**, **4** and **5** were collected and integrated using a Bruker AXS Kappa APEX2 CCD diffractometer with graphite monochromated Mo-K α (λ = 0.71073 Å) radiation at 296(2) K. The crystal data for **6** and **7** were collected and integrated using a D8 VENTURE Bruker AXS diffractometer with graphite monochromated Mo-K α (λ = 0.71073 Å) radiation at 150(2) K. The structure of **2** was solved by heavy atom methods using SIR92 and refined using SHELXL-2018/3.^{33,34} The structure of **4** was solved by heavy atom methods using SHELXT-2014/5 and refined using SHELXL-2018/3.³⁴ The structure of **5** was solved by heavy atom methods using SHELXT-2014/5 and refined using SHELXL-2018/3.³⁴ The structure of **6** and **7** were solved by heavy atom methods using SHELXT-2015 and refined using SHELXL-2018/3.³⁴ The molecular structures were drawn using Olex² software.³⁵ The non-hydrogen atoms were refined with anisotropic displacement parameters. Crystallographic data have been deposited with the Cambridge Crystallographic Data Center as supplementary publication no CCDC-1990065 (**2**), CCDC-1990066 (**4**), CCDC-1990067 (**5**), CCDC-1990068 (**6**) and CCDC-1990069 (**7**). These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

Crystal data for **2**: $\text{C}_{29}\text{H}_{40}\text{B}_2\text{Ru}_2\text{S}_2$, M_r = 676.49, Tetragonal, space group $I4_1/a$, a = 13.1579(4) Å, b = 13.1579(4) Å, c = 32.8239(16) Å, α = 90°, β = 90°, γ = 90°, V = 5682.8(4) Å³, Z = 8, ρ_{calcd} = 1.581 g/cm³, μ = 1.227 mm⁻¹, $F(000)$ = 2752, R_i = 0.0470, wR_2 = 0.1030, 1972 independent reflections [$2\theta \leq 49.976^\circ$] and 192 parameters. Crystal data for **4**: $\text{C}_{21}\text{H}_{34}\text{B}_2\text{Ru}_2\text{S}_3$, M_r = 606.42, Triclinic, space group $P-1$, a = 13.9273(5) Å, b = 14.3620(7) Å, c = 14.7096(7) Å, α = 76.888(3)°, β = 64.981(2)°, γ = 64.953(2)°, V = 2411.37(19) Å³, Z = 4, ρ_{calcd} = 1.670 g/cm³, μ = 1.518 mm⁻¹, $F(000)$ = 1224, R_i = 0.0454, wR_2 = 0.0830, 4236 independent reflections [$2\theta \leq 49.998^\circ$] and 521 parameters.

Crystal data for **5**: $\text{C}_{21}\text{H}_{36}\text{B}_2\text{Ru}_2\text{S}_2$, M_r = 576.38, Monoclinic, space group $P2_1/c$, a = 14.329(4) Å, b = 9.1095(16) Å, c = 19.357(5) Å, α = 90°, β = 105.817(8)°, γ = 90°, V = 2431.0(10) Å³, Z = 4, ρ_{calcd} = 1.575 g/cm³, μ = 1.419 mm⁻¹, $F(000)$ = 1168, R_i = 0.0932, wR_2 = 0.2001, 5447 independent reflections [$2\theta \leq 44.204^\circ$] and 268 parameters.

Crystal data for **6**: $\text{C}_{20}\text{H}_{30}\text{Ru}_2\text{S}_4$, M_r = 600.82, Monoclinic, space group $P2_1/c$, a = 15.2538(14) Å, b = 10.4684(9) Å, c = 15.1172(13) Å, α = 90°, β = 107.016(3)°, γ = 90°, V = 2308.3(4) Å³, Z = 4, ρ_{calcd} = 1.729 g/cm³, μ = 1.673 mm⁻¹, $F(000)$ = 1208, R_i = 0.0222, wR_2 = 0.0532, 4829 independent reflections [$2\theta \leq 54.962^\circ$] and 245 parameters.

Crystal data for **7**: $\text{C}_{20}\text{H}_{31}\text{BRu}_2\text{S}_4$, M_r = 612.64, Monoclinic, space group $P2_1/n$, a = 7.9142(7) Å, b = 20.1634(19) Å, c = 15.2351(14) Å, α = 90°, β = 100.935(4)°, γ = 90°, V = 2387.0(4) Å³, Z = 4, ρ_{calcd} = 1.705 g/cm³, μ = 1.619 mm⁻¹, $F(000)$ = 1232, R_i = 0.0241, wR_2 = 0.0517, 4737 independent reflections [$2\theta \leq 54.968^\circ$] and 257 parameters.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: xx. It contains ^1H , $^{11}\text{B}\{^1\text{H}\}$, $^{13}\text{C}\{^1\text{H}\}$ NMR, mass spectra and X-ray analysis for complexes **2**,

4, 5, 6 and 7. This material is available free of charge via the internet at <http://pubs.acs.org>.

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. [§] S. K. and U. K. are equal second authors.

Notes

The authors declare no competing financial interest.

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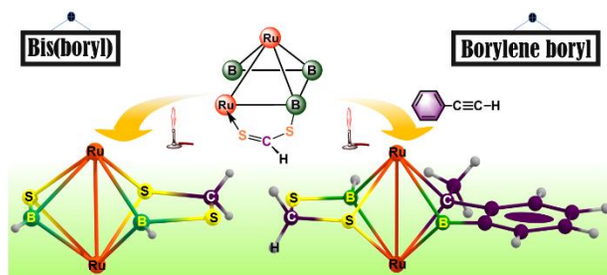
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New synthetic methods for ruthenium borylene and boryl complexes have been developed. Thermolysis of *arachno*-ruthenaborane, $[(\text{Cp}^*\text{Ru})_2\text{B}_3\text{H}_8(\text{CS}_2\text{H})]$ with phenyl acetylene led to the formation of ruthenium bridging boryl borylene complex, $[(\text{Cp}^*\text{Ru})_2(\mu\text{-HBS}_2\text{CH}_2\text{-}\kappa^2\text{B}:\kappa^2\text{S})\{\mu\text{-B}(\text{C}_6\text{H}_4)\text{C}(\text{Me})\text{-}\kappa^2\text{B}:\kappa^2\text{C}\}]$ via the hydroboration of alkyne (see picture).



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