

Sandwich Complexes

How to cite: Angew. Chem. Int. Ed. 2022, 61, e202206840 International Edition: doi.org/10.1002/anie.202206840 German Edition: doi.org/10.1002/ange.202206840

Stable Two-Legged Parent Piano-Stool and Mixed Diborabenzene-E₄ (E=P, As) Sandwich Complexes of Group 8

Maximilian Dietz, Merle Arrowsmith, Stephan Reichl, Leonardo I. Lugo-Fuentes, J. Oscar C. Jiménez-Halla, Manfred Scheer, and Holger Braunschweig*

Abstract: A cyclic alkyl(amino)carbene-stabilized 1,4diborabenzene (DBB) ligand enables the isolation of 18electron two-legged parent piano-stool Fe⁰ and Ru⁰ complexes, $[(\eta^6\text{-DBB})M(CO)_2]$, the ruthenium complex being the first of its kind to be structurally characterized. $[(\eta^6-DBB)Fe(CO)_2]$ reacts with E_4 (E=P, As) to yield mixed DBB-cyclo-E₄ sandwich complexes with planar E_4^{2-} ligands. Computational analyses confirm the strong electron-donating capacity of the DBB ligand and show that the E₄ ligand is bound by four equivalent Fe-P σ bonds.

Owing to their labile CO ligands low-valent half-sandwich transition metal carbonyl complexes, or parent piano-stool complexes, [1-3] are ideal precursors for a variety of halfsandwich and mixed ligand sandwich complexes used in organometallic chemistry and catalysis. [4-12] Two-legged parent piano-stool 18-electron complexes, $[(\eta^n - C_n R_n)M$ $(CO)_2$] (n=3-8), are relatively rare compared to their threeand four-legged counterparts. Often generated in situ by photolysis of tricarbonyl precursors, [13,14] they tend to dimerize via metal-metal bonding and/or CO bridging. [15-20] Bursten showed that the stability of these complexes is mainly governed by the relative energy of the HOMO versus the LUMO and HOMO-1.^[21] Their relative instability centered HOMO by the π -acidic aromatic ligand. [22,23] There are only a handful of zero-valent two-legged

often results from insufficient stabilization of the metal-

parent piano-stool group 8 complexes. Most of these are highly reducing anionic Fe⁰ and Ru⁰ cyclopentadienyl derivatives, often stabilized by additional CO--countercation interactions. [24-32] While $[(\eta^6$ -arene)M(CO)₂] (M=Fe, Ru, Os) complexes such as I-M, II and IIIR-X (Figure 1) have long been known, [33-36] only the iron complex III^{Me}-SiMe₃ has been structurally authenticated.^[36]

The substitution of endocyclic carbon atoms by more electropositive boron atoms in a benzene ring results in a removal of degeneracy of the highest filled molecular orbitals (MOs) and a destabilization of the MOs involving the boron 2p orbitals.^[37–42] This makes borabenzenes significantly more electron-donating than their isoelectronic organic counterparts. Our group has reported the synthesis and coordination chemistry of the doubly cyclic alkyl-(amino)carbene (CAAC)-stabilized 1,4-diborabenzene (DBB) 1 (Scheme 1).[43-45] Three-legged parent piano-stool group 6 DBB complexes (IV-M, Figure 1) displayed the lowest CO stretching frequencies yet observed for this class

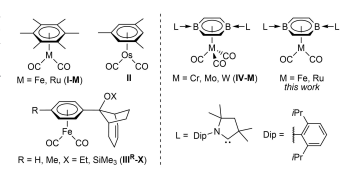


Figure 1. Known neutral two-legged parent piano-stool group 8 arene complexes (I-III) and parent piano-stool 1,4-diborabenzene (DBB) group 6 (IV) and group 8 complexes.

Scheme 1. Synthesis of two-legged parent piano-stool group 8 diborabenzene complexes.

Am Hubland, 97074 Würzburg (Germany) and

Institute for Sustainable Chemistry & Catalysis with Boron, Julius-Maximilians-Universität Würzburg

Am Hubland, 97074 Würzburg (Germany)

E-mail: h.braunschweig@uni-wuerzburg.de

S. Reichl, Prof. Dr. M. Scheer

Institute of Inorganic Chemistry, University of Regensburg Universitätsstraße 31, 93040 Regensburg (Germany)

L. I. Lugo-Fuentes, Prof. Dr. J. O. C. Jiménez-Halla Departamento de Química, División de Ciencias Naturales y Exactas, Unversidad de Guanajuato, Noria Alta S/N Col. Noria Alta, Guanajuato, C.P. 36050, Gto. (Mexico)

© 2022 The Authors. Angewandte Chemie International Edition published by Wiley-VCH GmbH. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

^[*] M. Dietz, Dr. M. Arrowsmith, Prof. Dr. H. Braunschweig Institute for Inorganic Chemistry, Julius-Maximilians-Universität Würzburg





of complex, reflecting the exceptionally strong electron-donating ability of $\mathbf{1}^{,[44]}$ Furthermore, $\mathbf{1}$ enabled the synthesis of the first heteroarene actinide complexes. [45] In this work we report the use of $\mathbf{1}$ as a stabilizing ligand for two-legged parent piano-stool group 8 complexes of the form $[(\eta^6\text{-DBB})M(CO)_2]$ (M=Fe, Ru) and the reduction of white phosphorus and yellow arsenic by the iron complex, yielding mixed-ligand $[(\eta^6\text{-DBB})M(\eta^4\text{-E}_4)]$ (E=P, As) sandwich complexes.

The two-legged piano-stool complexes 2-Fe and 2-Ru were synthesized by refluxing 1 with [Fe(CO)₅] or [Fe₂- $(CO)_9$] and $[Ru_3(CO)_{12}]$, respectively, in toluene (Scheme 1). The complexes were isolated as dark green and dark turquoise solids, respectively. The analogous reaction with $[Os_3(CO)_{12}]$ did not lead to the formation of **2-Os**. The ¹¹B NMR spectrum of 2-Fe and 2-Ru complexes shows a broad resonance around 4 ppm, significantly upfield from 1 (δ_{11B} = 24.8 ppm), [43] and similar to the group 6 complexes IV-M $(\delta_{11B} = 6-7 \text{ ppm}).^{[44]}$ The ¹H and ¹³C NMR DBB ring resonances are significantly upfield-shifted from 7.3 to ca. 4.7 ppm and from 150 to ca. 108 ppm, respectively, reflecting a reduction in aromaticity. The IR spectra showed several broad CO stretching bands in the 1822–1877 cm⁻¹ region for **2-Fe** and 1843–1978 cm⁻¹ for **2-Ru**, shifted to much lower wavenumbers than in the analogous arene complexes III^R-X $(v_{\rm CO} = 1877 - 1980 \text{ cm}^{-1})$ and I-Ru 1973 cm⁻¹). [34,36] This confirms the exceptionally strong backdonation from the [(DBB)M] fragment into the π^* orbitals of the CO ligands previously observed for **IV-M**. [44]

The UV-Vis spectra of **2-Fe** and **2-Ru** in benzene showed absorption maxima at 425 and 358 nm, respectively, as well as broad higher-wavelength absorption bands in the 570–710 nm region, and two additional low intensity bands at 453 and 494 nm for **2-Ru**. Calculations on complexes **IV-M** have shown that the absorption maximum around 400 nm corresponds to π - π * ligand transitions, [44] which undergo a hypsochromic shift upon descending the group, correlating with stronger binding to the metal. The broad bands at lower energy are likely metal-to-ligand charge transfer bands.

The solid-state structures of **2-Fe** and **2-Ru** confirm the formation of the two-legged piano-stool complexes (Figure 2).^[46] While a number of neutral two-legged piano-stool Fe⁰ arene complexes have been structurally characterized, including doubly CO-,^[36] NHC-,^[47] silylene-,^[48] stannylene-,^[49] and phosphine-stabilized examples,^[50-52] **2-Ru** is, to our

Figure 2. Solid-state structures of 2-Fe (side view) and 2-Ru (top view). Atomic displacement ellipsoids at 50%. Ellipsoids of ligand periphery and CAAC hydrogen atoms omitted for clarity.

knowledge, the first structurally characterized Ru⁰ complex of this type. Unlike in 1 and complexes IV-M,[43,44] the DBB ring in 2-Fe and 2-Ru is slightly twisted out of planarity (B-C-C-B torsion angles: 3.0-5.4°). The CAAC ligands, rotated by 20.8–24.7° out of the C₄B₂ plane, adopt a trans conformation rather than the cis conformation observed in 1 and IV-M. The endocyclic C-C (avg. 1.41 Å) and exocyclic B-C bonds (avg. 1.57 Å) of the DBB ring are slightly longer than in 1 (1.38 and 1.54 Å, respectively), [43] reflecting a decrease in π delocalization within the DBB ring and in π backdonation to the CAAC ligands due to competition with the metal. Whereas the iron center in 2-Fe is equidistant from the two boron atoms (Fe1-B1 2.2075(17), Fe1-B2 2.2076(16) Å), the ruthenium atom in 2-Ru sits slightly offcenter (Ru1-B1 2.349(3), Ru1-B2 2.317(3) Å). The Fe---(DBB)_{centroid} distance (1.59 Å) and Fe-CO bond lengths (avg. 1.74 Å) in 2-Fe are identical to those in the arene analogue III^{Me}-SiMe₃.[36] The C-O bonds lengths of 2-Fe (1.171(2) and 1.166(2) Å), however, are slightly longer than in \mathbf{HI}^{Me} -SiMe₃ (1.151(2) and 1.159(2) Å), confirming the strong backdonation from the [(DBB)M] fragment suggested by the IR C-O stretching bands.

The cyclovoltammogram of **2-Fe** shows a reversible reduction at -2.46 V and a reversible oxidation at -1.13 V (vs Fc/Fc⁺ couple), which correspond to DBB-based redox events, as the free ligand itself shows two reversible redox events at -2.48 and -0.81 V.^[43] Further irreversible reduction and oxidation events are observed at -3.05 and -0.02 V, respectively. The cyclovoltammogram of **2-Ru** similarly shows four redox waves at -3.19, -2.53, -1.42 and -0.87 V, of which only the reduction at -2.53 V is reversible. Thus the identity of the metal center only influences the oxidation events, so that **2-Ru** is much less easily oxidized than **2-Fe**.

Irradiating a benzene suspension of **2-Fe** and white phosphorus for 4 days under a Hg/Xe UV lamp (185–2000 nm) resulted in quantitative conversion to the mixed sandwich complex **3-P**, isolated as a dark brown crystalline solid (Scheme 2). The heavier analogue **3-As** was obtained by refluxing **2-Fe** with yellow arsenic for 1 h in decalin, and isolated as a dark brown solid. The high reaction temperature of 190 °C did not lead to any substantial decomposition of the **2-M** precursors, thus demonstrating the stabilizing power of the DBB ligand. Despite the prolonged reaction time for the synthesis of **3-P** no reaction intermediates could

L
$$\rightarrow$$
 B \rightarrow L

Second B \rightarrow L

E = P: benzene, hv , 4 d

E = As: decalin, 190 °C, 1 h

E = P: benzene, hv , 4 d

E = P: benzene, hv , 4 d

E = P: benzene, hv , 4 d

Fe = P: benzene, hv , 4 d

Scheme 2. Reduction of P₄ and As₄ by 2-Fe.





be detected by ¹¹B or ³¹P NMR spectroscopy, the reaction proceeding cleanly from **2-Fe** to **3-P**. While P_4 potentially undergoes dissociation to P_2 under the photolytic conditions employed in the synthesis of **3-P**, ^[53] the latter is also formed under the thermal conditions employed in the synthesis of **3-As**, albeit less efficiently. Since the thermal dissociation of P_4 to P_2 and As_4 to As_2 require temperatures above 1100 K and 800 K, respectively, ^[54,55] we deem it unlikely that the formation of **3-E** proceeds via generation of E_2 fragments.

The reduction of the elemental pnictogens was not achieved with **2-Ru** under similar conditions. The ¹¹B NMR shifts of 5.3 and 4.4 ppm for **3-P** and **3-As**, respectively, are similar to that of **2-Fe**. The ³¹P NMR spectrum of **3-P** displays a broad resonance at 61.4 ppm for the P₄ ring, downfield-shifted from that of Mézailles' related tris(phosphine) iron *cyclo*-P₄ complex (δ_{31P} =53.2 ppm, see complex **4** in Table 3). ^[56] The ⁷⁵As NMR shift of **3-As** could not be detected in the entire -1300–2000 ppm range. The UV-Vis spectra of **3-P** and **3-As** each show two major absorption bands in the 380–435 nm region and several broad low intensity bands in the 450–635 nm range.

X-ray crystallographic analyses of 3-P and 3-As confirmed the formation of the mixed sandwich complexes (Figure 3). Unlike in 2-Fe, the DBB ring is quasi-planar (B-C-C-B torsion angles < 1°). The Fe--(DBB)_{centroid} distance (1.55 Å) is slightly shorter than in **2-Fe**, suggesting a stronger interaction. The Fe···(E_4)_{centroid} distance in 3-P (1.77 Å) is only slightly shorter than in **3-As** (1.79 Å). The E₄ rings are parallel to the DBB ring and essentially squareplanar (Σ (E–E–E) \approx 360°). The E–E bonds in **3-P** (2.139(2)– 2.179(2) Å) and 3-As (2.383(1)-2.397(1) Å) are within the range of partial double bonds.^[57] Complexes with delocalized As4 rings are much rarer than their lighter cyclo-P4 analogues, both types including several double-decker iron and cobalt complexes. [58,59] The only other structurally characterized neutral end-deck cyclo-P4 iron and end-deck cyclo-As₄ complexes are [{κ³-(PhP(CH₂CH₂PCy₂)₂)}Fe(η⁴- P_4] (see complex **4**, Table 3)^[53] and $[(\eta^5-C_5Me_5)Nb(CO)_2(\eta^4-$ As₄)],^[60] respectively, both showing similar E₄ bonding parameters to 3-P and 3-As, respectively.

Given the strong π -acceptor properties of the DBB ligand, which favor its reduction to [DBB]^{2-,[44]} DFT calculations were carried out at the ω B97X–D/def2-svpp level to investigate the electronic structure of complexes **2**-

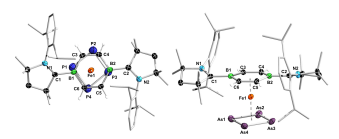


Figure 3. Crystallographically-determined solid-state structures of 3-P (top view) and 3-As (side view). Atomic displacement ellipsoids at 50%. Ellipsoids of ligand periphery and CAAC hydrogen atoms omitted for clarity.

M and **3-E** (see details in the Supporting Information). Bond dissociation energy (BDE) calculations for 2-M (M= Fe, Ru, Os) provided the lowest energy values for the neutral singlet fragments, DBB and M(CO)₂ (Table 1). The calculated net charge of the metal centers is close to neutral and effective oxidation state (EOS)^[61] calculations for 2-Fe and 2-Ru confirm their oxidation state of zero. The HOMO of 2-M is located at the metal carbonyl fragment whereas the LUMO is mainly located at the carbene moieties. The HOMO energy of 2-M decreases down the group (by only 3 kcal mol⁻¹), as do the HOMO-LUMO gaps (2-Fe 5.67 eV, 2-Os 5.50 eV). The reason why 2-Os is not formed lies in the instability of the Os(CO)2 fragment required for its formation. The $M \rightarrow CO \pi$ backdonation decreases from M =Fe to Os (Table S2), which in turn weakens the M-CO bonds, thus destabilizing the M(CO)₂ fragment.

In order to understand the reluctance of **2-Ru** to react with P_4 and As_4 , reaction energies were calculated for the formation of **(DBB)ME**₄ from **2-M** via the dissociation of both CO ligands. Table 2 shows that while the overall energies of formation of **(DBB)RuE**₄ are similar to those of **3-E**, the dissociation energy for both CO ligands from **2-Ru** $(\Delta G = 119.6 \text{ kcal mol}^{-1})$ is a prohibitive 45 kcal mol⁻¹ higher than from **2-Fe** $(\Delta G = 74.5 \text{ kcal mol}^{-1})$, thus preventing the reaction.

The electronic nature of **3-P** and **3-As** was assessed and compared with complex **4**, a simplified version of the known Fe^{II} complex $[\{\kappa^3-(PhP(CH_2CH_2PCy_2)_2)\}Fe(\eta^4-P_4)],^{[56]}$ in

Table 1: Frontier MOs, BDEs (corrected by BSSE) and HOMO-LUMO gaps of **2-M** at the ω B97X-D/def2-svpp level.

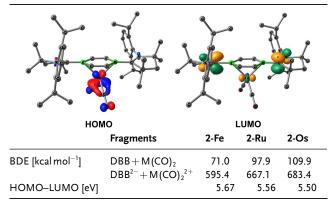


Table 2: Calculated reaction energies for the reduction of E_4 by **2-M** at the $\omega B97X-D/def2$ -svpp level.

Reaction	Е	ΔG [kcal mol $^{-1}$]
2-Fe →(DBB)Fe+2 CO		74.5
(DBB) $Fe + E_4 \rightarrow 3-E$	Р	-124.3
	As	-100.9
2-Fe + E_4 → 3-E + 2 CO	Р	-49.8
	As	-26.4
2-Ru →(DBB)Ru+2 CO		119.6
$(DBB)Ru + E_4 \rightarrow (DBB)RuP_4$	Р	-167.7
	As	-146.1
$2\text{-}\mathbf{R}\mathbf{u} + E_4 \!\!\rightarrow\! (DBB) Ru P_4 \!+\! 2 CO$	Р	-48.1
	As	-26.4

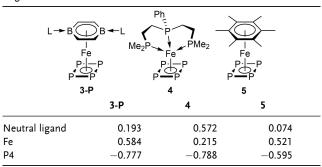




which the cyclohexyl groups were replaced by methyl groups, and the model arene-P4 sandwich complex 5 (Table 3). Calculated net charges at the metal centers and P₄ ligands and EOS calculations confirm that all four compounds are Fe^{II} complexes with dianionic E₄ ligands (see Supporting Information). The main differences between 3-P, 4 and 5 are seen in the natural bond orbital (NBO) charges summed by fragments.^[62] The NBO charge of the DBB fragment of 3-P is less positive (+0.193) than that of the tris(phosphine) fragment of 4 (+0.572), indicating that the latter is more electron-donating. This is in agreement with the downfield-shift of the ³¹P NMR resonance of 3-P (δ_{31P} = 61.4 ppm) compared to [{ κ^3 -(PhP(CH₂CH₂PCy₂)₂)}Fe- (η^4-P_4)] $(\delta_{31P}=53.2 \text{ ppm})$. [56] In turn, the DBB ligand of **3-P** is more electron-donating than the hexamethylbenzene ligand of 5 (+0.072), as expected upon exchanging two carbon atoms with boron.

The bonding within **3-P** and **3-As** was investigated using the NBO approach. Calculations suggest that these complexes have three Fe–E σ bonds (E=P, As) and one lone pair on E, which is strongly delocalized towards the vacant Fe s orbital (n(P) $\rightarrow s$ (Fe)), as revealed by the interaction energies (E(2)) of 76.1 and 78.8 kcal mol⁻¹ for **3-P** and **3-As** respectively (Figure 4, see Supporting Information for **3-As** and for IBO calculations). Due to greater charge transfer

Table 3: NBO charges of complexes 3-P, 4 and 5 summed by fragments.



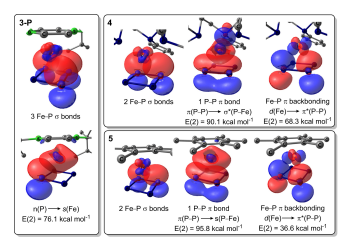


Figure 4. Fe $-P_4$ bonding NBOs of complexes 3-P, 4 and 5. Isosurface value: 0.05 a.u.

from the tris(phosphine) ligand to the metal and P_4 fragments in **4**, as well as the low C_s symmetry of the tris(phosphine) ligand, the NBOs of **4** show only two Fe–P σ bonds and a P–P π bond, which allows π backbonding from one metal d orbital into the P–P π^* orbital (E(2) = 68.3 kcal mol⁻¹). Complex **5** shows a P–P π bond and two Fe–E σ bonds, albeit with weaker π backbonding (E(2) = 36.6 kcal mol⁻¹) due to the lesser extent of charge transfer from the neutral ligand to the metal and P_4 fragments. This shows that the bonding situation between the metal center and the E_4 ligand is strongly affected by the electronics and symmetry of the neutral ligand.

To conclude, the highly electron-donating DBB ligand enables the stabilization of rare two-legged parent Fe^0 and Ru^0 piano-stool complexes, $[(\eta^6\text{-DBB})M(CO)_2]$ with high thermal stability. The ruthenium complex is the first of its kind to be crystallographically characterized. The reduction of white phosphorus or yellow arsenic by the iron complex yields the mixed sandwich complexes $[(\eta^6\text{-DBB})Fe(\eta^4\text{-E}_4)],$ which display square-planar $E_4^{\ 2^-}$ ligands. Calculations confirm these are Fe^{II} complexes and show that the $\mbox{cyclo-}E_4$ ligand is symmetrically bound by four σ bonds. Comparison with related $[LFe(\eta^4\text{-E}_4)]$ complexes shows that $Fe\mbox{-P}_4$ bonding is strongly influenced by the nature of the neutral ligand L.

Acknowledgements

Financial support from the Deutsche Forschungsgemeinschaft (DFG) is gratefully acknowledged. M.D. and S.R. thank the Studienstiftung des deutschen Volkes for a doctoral fellowship. L.I.L.-F. acknowledges CONACyT for financial support through his M.Sc. fellowship #1080463. Open Access funding enabled and organized by Projekt DEAL.

Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Keywords: 1,4-Diborabenzene · Bonding · Group 8 Metals · Parent Piano-Stool Complex · Pnictogen Reduction

^[1] H. Werner, Angew. Chem. Int. Ed. Engl. 1983, 22, 927–949; Angew. Chem. 1983, 95, 932–954.

^[2] P. Kubácek, R. Hoffmann, Z. Havlas, Organometallics 1982, 1, 180–188.

^[3] F. A. Cotton, J. R. Kolb, J. Organomet. Chem. 1976, 107, 113– 119

^[4] P. Piou, T. Rovis, Acc. Chem. Res. 2018, 51, 170-180.



Communications



- [5] C. Johnson, M. Albrecht, Coord. Chem. Rev. 2017, 352, 1-14.
- [6] N. V. Shvydkiy, D. S. Perekalin, Coord. Chem. Rev. 2017, 349, 156–168.
- [7] P. Kumar, R. K. Gupta, D. S. Pandey, Chem. Soc. Rev. 2014, 43, 707–733.
- [8] J. Václavík, P. Šot, B. Vilhanová, J. Pecháček, M. Kuzma, P. Kačer, Molecules 2013, 18, 6804–6828.
- [9] W. E. Geiger, Coord. Chem. Rev. 2013, 257, 1459–1471.
- [10] M. Nishiura, Z. Hou, Bull. Chem. Soc. Jpn. 2010, 83, 595-608.
- [11] J. Liu, X. Wu, J. A. Iggo, J. Xiao, Coord. Chem. Rev. 2008, 252, 782–809
- [12] H. Werner, Organometallics 2005, 24, 1036-1049.
- [13] N. C. Baentiger, R. M. Flynn, N. L. Holy, Acta Crystallogr. Sect. B 1979, 35, 741–744.
- [14] L. R. Byers, L. F. Dahl, Inorg. Chem. 1980, 19, 277-284.
- [15] T. S. Janik, C. H. Lake, M. R. Churchill, Organometallics 1993, 12, 1682–1685.
- [16] C. P. Casey, H. Sakaba, P. N. Hazin, D. R. Powell, J. Am. Chem. Soc. 1991, 113, 8165–8166.
- [17] R. J. Klingler, W. M. Butler, M. D. Curtis, J. Am. Chem. Soc. 1978, 100, 5034–5039.
- [18] M. D. Curtis, W. M. Butler, J. Organomet. Chem. 1978, 155, 131–145.
- [19] R. D. Fischer, A. Vogler, K. J. Noack, Organomet. Chem. 1967, 7, 135–149.
- [20] F. A. Cotton, G. Yagupsky, Inorg. Chem. 1967, 6, 15–20.
- [21] B. E. Bursten, M. G. Gatter, K. I. Goldberg, *Polyhedron* 1990, 9, 2001–2011.
- [22] T. A. Albright, R. Hoffmann, Y.-C. Tse, T. D'Ottavio, J. Am. Chem. Soc. 1979, 101, 3812–3821.
- [23] D. L. Lichtenberger, D. C. Calabro, G. E. Kellogg, *Organo-metallics* 1984, 3, 1623–1630.
- [24] H. Braunschweig, R. D. Dewhurst, K. Ferkinghoff, Chem. Commun. 2016, 52, 183–185.
- [25] T. Pugh, N. F. Chilton, R. A. Layfield, Angew. Chem. Int. Ed. 2016, 55, 11082–11085; Angew. Chem. 2016, 128, 11248–11251.
- [26] K. F. Kalz, N. Kindermann, S.-Q. Xiang, A. Kronz, A. Lange, F. Meyer, *Organometallics* 2014, 33, 1475–1479.
- [27] I. Sänger, T. I. Kückmann, F. Dornhaus, M. Bolte, M. Wagner, H.-W. Lerner, *Dalton Trans.* 2012, 41, 6671–6676.
- [28] M. P. Blake, N. Kaltsoyannis, P. Mountford, J. Am. Chem. Soc. 2011, 133, 15358–15361.
- [29] B. T. Carter, M. P. Castellani, A. L. Rheingold, S. Hwang, S. E. Longacre, M. G. Richmond, *Organometallics* 2002, 21, 373– 270
- [30] J. Kühn, K. Ruck-Braun, J. Prakt. Chem. 1997, 339, 675-678.
- [31] E. Hey-Hawkins, H. G. von Schnering, *Z. Naturforsch.* **1991**, 46h 621–624
- [32] J. S. Plotkin, S. G. Shore, *Inorg. Chem.* **1981**, *20*, 284–287.
- [33] S. R. Weber, H. H. Brintzinger, J. Organomet. Chem. 1977, 127, 45–54.
- [34] K. Roder, H. Werner, Chem. Ber. 1989, 122, 833-840.
- [35] S. Stahl, H. Werner, Organometallics 1990, 9, 1876-1881.
- [36] J. Chen, J. Yin, Z. Fan, W. Xu, J. Chem. Soc. Dalton Trans. 1988, 2803–2808.
- [37] "Synthesis and Reactivity of Bora- and Boratabenzenes": I. Krummenacher, J. Schuster, H. Braunschweig *Patai's Chem. Funct. Groups* (Eds.: I. Marek, M. Gandelman), Wiley, Hoboken, 2019.
- [38] B. Su, R. Kinjo, Synthesis 2017, 49, 2985-3034.

- [39] J. Singh, Y. Wang, G. Raabe, Z. Naturforsch. 2010, 65a, 113– 122.
- [40] P. B. Karadakov, M. Ellis, J. Gerratt, D. L. Cooper, M. Raimondi, J. Quantum Chem. 1997, 63, 441–449.
- [41] C. W. Allen, D. E. Palmer, J. Chem. Educ. 1978, 55, 497–500.
- [42] R. Boese, N. Finke, T. Keil, P. Paetzold, G. Schmid, Z. Naturforsch. 1985, 40b, 1327–1332.
- [43] M. Arrowsmith, J. Böhnke, H. Braunschweig, M. A. Celik, C. Claes, W. Ewing, I. Krummenacher, K. Lubitz, C. Schneider, Angew. Chem. Int. Ed. 2016, 55, 11271–11275; Angew. Chem. 2016, 128, 11441–11445.
- [44] J. Böhnke, H. Braunschweig, J. O. C. Jiménez-Halla, I. Krummenacher, T. E. Stennett, J. Am. Chem. Soc. 2018, 140, 848–853.
- [45] V. Paprocki, P. Hrobárik, K. L. M. Harriman, M. S. Luff, T. Kupfer, M. Kaupp, M. Murugesu, H. Braunschweig, Angew. Chem. Int. Ed. 2020, 59, 13109–13115; Angew. Chem. 2020, 132, 13209–13216.
- [46] Deposition Numbers 2169216 (for 3-As), 2169217 (for 2-Ru), 2169218 (for 2-Fe), and 2169219 (for 3-P) contain the supplementary crystallographic data for this paper. These data are provided free of charge by the joint Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service.
- [47] B. Blom, G. Tan, S. Enthaler, S. Inoue, J. D. Epping, M. Driess, J. Am. Chem. Soc. 2013, 135, 18108–18120.
- [48] M.-P. Luecke, D. Porwal, A. Kostenko, Y.-P. Zhou, S. Yao, M. Keck, C. Limberg, M. Oestreich, M. Driess, *Dalton Trans*. 2017, 46, 16412–16418.
- [49] J. J. Schneider, N. Czap, D. Bläser, R. Boese, J. Ensling, P. Gütlich, C. Janiak, *Chem. Eur. J.* 2000, 6, 468–474.
- [50] S. L. Daifuku, J. L. Kneebone, B. E. R. Snyder, M. L. Neidig, J. Am. Chem. Soc. 2015, 137, 11432–11444.
- [51] A. Casitas, H. Krause, S. Lutz, R. Goddard, E. Bill, A. Fürstner, *Organometallics* **2018**, *37*, 729–739.
- [52] H. Kubo, M. Hirano, S. Komiya, J. Organomet. Chem. 1998, 556, 89–95.
- [53] G. Rathenau, Physica 1937, 4, 503-514.
- [54] H. W. Melville, S. C. Gray, Trans. Faraday Soc. 1936, 32, 271–285.
- [55] J. J. Murray, C. Pupp, R. F. Pottie, J. Chem. Phys. 1973, 58, 2569–2578.
- [56] A. Cavaillé, N. Saffon-Merceron, N. Nebra, M. Fustier-Boutignon, N. Mézailles, Angew. Chem. Int. Ed. 2018, 57, 1874–1878; Angew. Chem. 2018, 130, 1892–1896.
- [57] R. C. Fischer, P. P. Power, *Chem. Rev.* **2010**, *110*, 3877–3923.
- [58] L. Giusti, V. R. Landaeta, M. Vanni, J. A. Kelly, R. Wolf, M. Caporali, *Coord. Chem. Rev.* 2021, 441, 213927.
- [59] M. Seidl, G. Balázs, M. Scheer, Chem. Rev. 2019, 119, 8406– 8434
- [60] O. J. Scherer, J. Vondung, G. Wolmershauser, J. Organomet. Chem. 1989, 376, C35–C38.
- [61] E. Ramos-Cordoba, V. Postils, P. Salvador, J. Chem. Theory Comput. 2015, 11, 1501–1508.
- [62] G. Knizia, J. Chem. Theory Comput. 2013, 9, 4834-4843.

Manuscript received: May 10, 2022

Accepted manuscript online: July 4, 2022

Version of record online: July 27, 2022