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Introduction

The terms "aromatic hydrocarbon" and "planar π system" can be used as synonyms for most representatives of these substance classes. Triptycenes are among the few aromatic hydrocarbons that enter the third dimension, which is one reason why they have fascinated the chemical community for decades. Triptycene is built upon a [2.2.2]-bicyclic system with three 1,2-phenylene blades to generate a D_{3h} -symmetric paddlewheel scaffold.^{1,2} Due to its exceptional rigidity, the triptycene framework has been used as a backbone for transition-metal ligands^{3,4} and as a building block for molecular motors.⁵ When incorporated into emitting polymers, triptycene units can sterically block the individual polymer chains so that the materials exhibit solution-like luminescence properties in the solid state, with applications in sensor technology.6 The threedimensional shape of a triptycene building block creates a free volume around the molecule (i.e., between the blades of the paddle-wheel structure), which can be designed such that it confers size selectivity in sensory responses,⁷ promotes the

Synthesis, bridgehead functionalization, and photoisomerization of 9,10-diboratatriptycene dianions[†]

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9,10-Diboratatriptycene salts $M_2[RB(\mu-C_6H_4)_3BR]$ (R = H, Me; M⁺ = Li⁺, K⁺, [*n*-Bu₄N]⁺) have been synthesized *via* [4 + 2] cycloaddition between doubly reduced 9,10-dihydro-9,10-diboraanthracenes $M_2[DBA]$ and benzyne, generated *in situ* from C_6H_5F and C_6H_5Li or LiN(*i*-Pr)₂. [HB(μ -C₆H₄)₃BH]²⁻ reacts with CH₂Cl₂ to form quantitatively the bridgehead-derivatized [ClB(μ -C₆H₄)₃BCl]²⁻, while twofold H⁻ abstraction with B(C_6F_5)₃ in the presence of SMe₂ leads cleanly to the diadduct (Me₂S)B(μ -C₆H₄)₃B(SMe₂). Photoisomerization of K₂[HB(μ -C₆H₄)₃BH] (THF, medium-pressure Hg lamp) provides facile access to diborabenzo[a]fluoranthenes, a little explored form of boron-doped polycyclic aromatic hydrocarbons. According to DFT calculations, the underlying reaction mechanism consists of three main steps: (i) photoinduced di- π -borate rearrangement, (ii) "walk reaction" of a BH unit, and (iii) boryl anion-like C–H activation.

desired alignment in oriented polymers,⁸ or creates porous organic materials.⁹ Of more academic interest – but no less fascinating – is the excited-state reactivity of triptycene and the resulting photorearrangement products (*i.e.*, a norcaradiene and a benzo[a]fluoranthene derivative).^{10,11}

A proper means of expanding the chemical space of triptycenes as well as their application potential is heteroatom doping, *i.e.*, the replacement of selected C atoms by other pblock elements. Our group has a long-standing interest in triptycenes featuring B atoms at their bridgehead positions. One example is the (BN)₂-triptycene A (Fig. 1). A-type structures are accessible by simple mixing of 9,10-dihydro-9,10diboraanthracenes (DBAs) with 1,2-diazenes, such as pyridazine. Since the reaction is based on the essentially barrierless formation of a double B-N adduct, A is obtained in very good yields of $\approx 80\%$.¹²⁻¹⁴ An organometallic (BN₂)₂-triptycene is the pyrazabole-bridged ansa-ferrocene B (Fig. 1).15 Recently, our group succeeded in abstracting one of its bridgehead H⁻ ligands and characterized the resulting free, cationic Lewis superacid $[\mathbf{B} - \mathbf{H}]^+$ by X-ray crystallography.¹⁶ Berionni and coworkers reported a series of boratatriptycenes C (Fig. 1) possessing various different atoms/groups "E" at the second bridgehead position ($E = CH, PH^+, S^+, Se^+$).^{17–20} A derivative of C combining E = CH and Ar = 4-t-Bu-C₆H₄ underwent selective protodeboronation of the exocyclic C-B bond upon treatment with bistriflimidic acid (HNTf₂); the resulting boratriptycene Lewis acid still engages in a dynamic association-dissociation equilibrium with the weakly coordinating $[NTf_2]^-$ anion.¹⁸ This finding underscores the potential for enhancing Lewis acidity

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Fig. 1 Boron-containing triptycenes. Left: the DBA-pyridazine adduct A and the 1.1'-ferrocenediyl-bridged pyrazabole B as BN derivatives of triptycene. Right: Berionni's BC-triptycenes C featuring one B atom at a bridgehead position (E = CH, n = 1; $E = PH^+$, S^+ , Se^+ , n = 0; Ar = aryl) and the dibora(ta)triptycenes D presented in this work (R = H, Me, n = 2; $R = SMe_2$, n = 0). DBA = 9,10-dihydro-9,10-diboraanthracene.

by designing a rigid organic framework in a way to impose a structural constraint on the BC_3 core that forces it out of planarity (see also the Lewis superacidity of $[\mathbf{B} - \mathbf{H}]^+$ mentioned above¹⁶).¹⁷

From the application examples of triptycenes, it is obvious that the degree of repulsive or attractive interactions between the paddle-wheels' blades and their immediate environments play an important role. Since these interactions will be strongly influenced by the overall charge of the triptycene scaffold, we are herein targeting dianionic diboratatriptycenes **D** (Fig. 1; **R** = **H**, Me; n = 2). We will show that H⁻ abstraction *in situ* releases a Lewis acid with potentially cooperating B atoms²¹ and allows convenient derivatization of both bridgehead positions. We finally show that the photorearrangement of **D** (**R** = **H**; n = 2) provides access to the little explored compound class of diborabenzo[*a*]fluoranthenes.²²

Results and discussion

Synthesis of 9,10-diboratatriptycenes D

We tested two conceptually different ways to prepare our **D**-type target compounds: (a) adduct formation on neutral DBA and (b) [4 + 2] cycloaddition on doubly reduced $[DBA]^{2-}$. The first approach (a) is inspired by the facile synthesis of **A** and based on the assumption that pyridazine can, in principle, be replaced by the isoelectronic benzene-1,2-diide to assemble the tricyclic scaffold of **D** *via* a twofold B–C-adduct formation. Unfortunately, the available literature on 1,2-dimetalated benzenes indicates a rather limited synthetic value of these highly reactive species.²³ We therefore resorted to a stepwise reaction protocol, starting from *in situ*-prepared *o*-iodophenyllithium and **1** (-110 °C, THF/Et₂O (1:1); Scheme 1).²⁴ Cyclization of the obtained B(sp²)–B(sp³) intermediate Li[4] was subsequently



Scheme 1 Synthesis of 9,10-diboratatriptycenes by either sequential B–C-adduct formation (left) or [4 + 2] cycloaddition of *in situ*-generated benzynes with M₂[DBA] salts (right). (i) 1,2-C₆H₄I₂, *t*-BuLi, THF/Et₂O (1:1), -110 °C \rightarrow room temperature, overnight, 21% yield; (ii) 1,2-C₂H₄Br₂ (6 mol%), THF, room temperature, 3 d; (iii) THF, room temperature, 1 h, quant. conv. by NMR; (iv) C₆H₅F (1-2 equiv.), base (LiN(*i*-Pr)₂ or C₆H₅Li), THF, -78 °C \rightarrow room temperature, overnight to 5 d, Li₂[5]: 50-75% yield, K₂[6]: 41% yield, K₂[7]: few crystals.

accomplished by stirring with Mg⁰ (room temperature, THF). Since the NMR-spectroscopic investigation of the crude product still showed the formation of some unwanted side products that later proved difficult to remove, the B-C-adduct approach was eventually abandoned in favor of the Umpolung strategy (b): given that doubly reduced DBAs easily undergo [4 + 2]cycloaddition reactions with (hetero)olefins or internal alkynes,²⁵ the reaction of DBA salts M₂[1]-M₂[3] with in situgenerated benzyne should likewise generate diboratatriptycenes $M_2[5]-M_2[7]$ (Scheme 1). For an initial proof-ofprinciple experiment, we selected Li₂[1], because we have had good experience with cycloadditions on this compound in the past.²⁵ LiN(*i*-Pr)₂ in THF was added with stirring at -78 °C to a mixture of freshly prepared Li₂[1] (ref. 26 and 27) and C₆H₅F in THF (Scheme 1). After warming to room temperature overnight, the reaction mixture had lost the characteristic red color of Li₂[1] and turned to pale yellow. ¹¹B NMR spectroscopy showed the selective formation of one new species; its chemical shift value of $\delta(^{11}B) = -13.2$ ppm testified to the presence of tetracoordinated B atoms.²⁸ An ¹H NMR spectrum, recorded on the crude product after a solvent change to THF- d_8 , indicated an essentially quantitative conversion: a signal at 0.56 ppm (6H)

and two resonances at 7.24 (br, 6H) and 6.44 ppm (m, 6H) are assignable to two chemically equivalent CH₃ groups and three chemically equivalent C₆H₄ rings, respectively. Also the ¹³C{¹H} NMR spectra supported the proposed D_{3h} -symmetric molecular structure of [5]²⁻ in solution. Purification of Li₂[5] through crystallization from THF gave yields of 50–75% (attempts at Xray analysis were not successful). Next, we moved on from DBA **1** to DBA **2** to also get a hold of the parent diboratatriptycene. K₂[6] was prepared in a similar manner as Li₂[5] and isolated in 41% yield (δ (¹¹B) = -8.3 ppm, d, ¹*J*(B,H) = 87 Hz); the BH resonance was found at 3.36 ppm in the ¹H{¹¹B} NMR spectrum).²⁹ To stress the steric limits of the [4 + 2] cycloaddition, we finally treated dimesitylated K₂[3] with benzyne and still obtained K₂[7], albeit in negligible yield of only a few crystals.²⁴

Crystals of [K(dme)][K(dme)₂][6] suitable for X-ray diffraction (XRD) were obtained from a gas-phase diffusion experiment (n-1)hexane/1,2-dimethoxyethane (DME); Fig. 2). In [K(dme)]- $[K(dme)_2][6]$, the diboratatriptycene dianion $[6]^{2-}$ acts as chelating ligand toward both K^+ ions (via the π -electron clouds of its 1,2-phenylene rings). The coordination sphere of $K(1)^+$ is completed by two dme ligands; $K(2)^+$ carries only one dme ligand and is further bonded to two phenylene rings of an adjacent $[6]^{2-}$ moiety, thereby creating a coordination polymer in which all three pockets of the paddle-wheel anion are occupied by K^+ ions (Fig. S61⁺). The six B-C-bond lengths of C_1 symmetric (in the crystal lattice) $[6]^{2-}$ range from 1.620(3) to 1.643(3) Å (avg. 1.631 Å) and are comparable to those in the structurally unconstrained, acyclic analogue [HBPh₃]⁻ (1.628(6) to 1.633(6) Å, avg. 1.631 Å).³⁰ Of the three dihedral angles 107.9°, 124.8°, and 127.3° between phenylene rings, the smallest is associated with the pocket hosting the doubly η^6 -coordinated $K(1)^{+}$.

Activation of the B–H bonds of the 9,10-diboratatriptycene $[6]^{2-}$

Having the diboratatriptycene salt [K(dme)][K(dme)₂][6] in hand, we next addressed the question of how to activate its B–H bonds to derivatize the bridgehead positions and/or generate the free Lewis acid *in situ*. The neutral ditopic Lewis acid **S1** (Scheme 2) should be among the strongest neutral B-based



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Scheme 2 Reactivity of 9,10-diboratatriptycene $[6]^{2-}$. (i) (1) THF, room temperature, 1 h; (2) extraction into CH₂Cl₂, 93% yield; (ii) 50 °C, 18 h, 89% yield; (iii) CD₂Cl₂, room temperature, 5 min, **9**: quant. conv. by NMR.

acids, as judged by its computed F⁻-ion affinity FIA_{soly}(S1) of 384 kJ mol^{-1} (cf. FIA_{solv}(B(C₆F₅)₃) = 315 kJ mol^{-1} ; solv = CH₂Cl₂; for FIAs of further B2-triptycenes and a more detailed discussion see the ESI[†]).¹⁶ As the dme ligands present in the complex cations could potentially become B-coordinating and thereby interfere with the planned reactivity studies, we first performed a cation exchange with $[n-Bu_4N]$ Cl in THF. $[n-Bu_4N]_2$ [6] precipitates together with KCl from the reaction mixture and was separated from the latter by extraction into CH_2Cl_2 (93% yield). Care must be taken to remove the solvent from the product in time after workup to avoid B-H/B-Cl exchange.31 On the other hand, the reaction between $[n-Bu_4N]_2[6]$ and CH_2Cl_2 can be used for the straightforward synthesis of bridgehead-chlorinated [n- $Bu_4N_2[8]$ (14 d at room temperature or 18 h at 50 °C; 89% yield; Scheme 2). In an attempt to release the free monotopic Lewis acid, $[n-Bu_4N]_2[6]$ was treated with 1 equiv. of $B(C_6F_5)_3$ in CD_2Cl_2 . NMR spectroscopy showed that $B(C_6F_5)_3$ had been quantitatively transformed to $[n-Bu_4N][HB(C_6F_5)_3]$, even though some unconsumed starting material [n-Bu₄N]₂[6] remained in the mixture. Several non-identified species and small quantities of a precipitate had formed. When the amount of $B(C_6F_5)_3$ added was increased to 2 equiv., the H⁻ scavenger was still fully converted, phenylene resonances were no longer detectable in the ¹H NMR spectrum, and substantial amounts of precipitate were found in the NMR tube. A similar result was obtained when $[Ph_3C][B(C_6F_5)_4]$ was used instead of $B(C_6F_5)_3$. The precipitate could not be re-dissolved by addition of SMe2. The situation changed when the double H⁻ abstraction was carried

[K(dme)][K(dme)₂][6]

out with $B(C_6F_5)_3$ in the presence of SMe₂, which led to the clean formation of the diboratriptycene-SMe2 diadduct 9 (Scheme 2). Based on the computed FIAs of relevant B2-triptycenes, we propose that the formation of 9 proceeds in a stepwise manner without intermediate formation of S1.24 X-ray quality crystals of $9 \cdot CH_2Cl_2$ were grown from CH_2Cl_2 (Fig. 2).

The ¹¹B NMR spectrum of **9** is characterized by a signal at -1.4 ppm, in the region of tetracoordinated B nuclei.²⁸ The ¹H/¹³C resonances of the SMe₂ ligands appear at 3.07/21.1 ppm (CD₂Cl₂), significantly downfield-shifted from those of free SMe₂ (2.09/18.4 ppm) and even Me₂S-B(C₆F₅)₃ (2.17/20.8 ppm;³²) CDCl₃), thus indicating a strong S-B-adduct bond. This conclusion is further confirmed by the short S-B-bond lengths of 9 (1.9710(19), 1.9775(18) Å; cf. $Me_2S-B(C_6F_5)_3$: S-B = 2.091(5)Å (ref. 33)).

To summarize, we assume that after H^- abstraction on $[6]^{2-}$, the corresponding free Lewis acid S1 is not stable but rather forms the insoluble solid by rearrangement or ring-opening polymerization. Although the structure of the solid remains to be confirmed, a plausible candidate is a polymer composed of 1,2-phenylene-bridged DBAs (Fig. S15[†]). The formation of a precipitate is suppressed if SMe₂ is present from the beginning to stabilize the diboratriptycene.

Photoisomerization of the 9,10-diboratatriptycene $[6]^{2-}$

In contrast to the thermal rearrangement of free neutral diboratriptycene, photoisomerization of the diboratatriptycene salt $K_2[6]$ results in the well-defined, soluble product $K_2[10]$ (Scheme 3). $K_2[10]$ can be viewed as a molecule in which a DBA moiety shares one B atom and one phenylene ring with a 9H-9borafluorene unit.

Irradiation of a THF solution of $K_2[6]$ with a mediumpressure Hg lamp in a quartz vessel for 1 h led to a color change from colorless to yellow.34 NMR spectroscopy on a THF d_8 solution revealed the highly selective formation of K₂[10]. Its ^{11}B NMR spectrum shows a doublet ($\delta(^{11}\text{B})$ = -13.7 ppm, ${}^{1}J(B,H) = 73$ Hz) and a pseudo-triplet ($\delta({}^{11}B) = -16.7$ ppm, ${}^{1}J(B,H) = 78$ Hz), assignable to a BH- and a BH₂ group, respectively. The corresponding BH_n resonances were found in the ¹H{¹¹B} NMR spectrum at 2.65 (BH) and 2.52/2.40 ppm (BH₂). The ¹³C{¹H} NMR spectrum supports the proposed C_1 symmetric molecular structure, since we observe 18 signals belonging to three different six-membered rings (according to ¹H-¹³C-HSQC and -HMBC experiments). Of those signals, five are severely broadened due to ${}^{1}I(B,C)$ coupling and the

72 K]2 K hν (i) K₂[10] K₂[6]

Scheme 3 Photoisomerization of K₂[6] to the 8,12b-dihydro-8,12bdiborabenzo[a]fluoranthene derivative K₂[10]. (i) Medium-pressure Hg lamp, THF, room temperature, 1 h, quant. conv. by NMR.

quadrupolar effect of the ^{10/11}B nuclei and therefore correspond to the five required B-bonded C atoms; deshielded, guaternary ¹³C atoms giving rise to NMR resonances at 151.1/144.3 ppm point toward the presence of a biphenyl fragment in the molecule. Moreover, the DFT-computed ¹³C shift values of [10]²⁻ agree well with the experimentally observed ones for $K_2[10]$ (avg. deviation: 2.0 ppm).²⁴ In order to gain more information on the mechanism underlying the photoisomerization of $[6]^{2-}$ to $[10]^{2-}$, we performed corresponding quantum-chemical calculations.

Assessment of the photoisomerization of K₂[6] by theory

All energies were calculated at the SMD(THF)³⁵/wB97XD³⁶/6-311++G** level of theory, with optimized structures at SMD(THF)/\u03c6B97XD/6-31+G**. Vertical excitation energies were calculated with the respective unrestricted functional (T_1 states) or by TD-DFT calculations (S_1 states). The calculations were performed on the bare dianionic species (Fig. 3a).37

We took the known photoisomerization pathway of carbonaceous triptycene 6^{C} and the di- π -borate rearrangement of the tetraarylborate K[11] as blueprints for our mechanistic studies (Fig. 3b and c).^{10,11,38} The first reaction step $[6]^{2-} \rightarrow [Int1]^{2-}$ involves C–C coupling between two aryl rings with simultaneous extrusion of one BH unit. In line with experiment, the associated energy barrier of 62.0 kcal mol^{-1} is prohibitively high for a reaction under ambient conditions (in the S_0 states). To account for the actually applied photochemical protocol, this crucial step was investigated in detail also for the excited T1 and S_1 states of the molecules involved.

First, we verified the basic premise, namely that the Hg lamp employed (emission range: 200-600 nm) does indeed provide the wavelength required to overcome the computed $S_0 \rightarrow S_1$ energy gap of $[6]^{2-}$ ($\Delta E_{S_0 \rightarrow S_1} = 5.068$ eV, corresponding to $\lambda_{exc} =$ 244.7 nm; $f_{\rm osc} = 0.1010$).

We scanned the T₁ and S₁ energy surface topographies along the reaction coordinate between $[6]^{2-}$ and $[Int1]^{2-}$ based on ground-state structures obtained from the [TS1]²⁻-IRC scan (Fig. 4; IRC = intrinsic reaction coordinate): the energy curves for both excited states have minima located downstream of the [TS1]²⁻ structure along the reaction coordinate. It is reasonable that regardless of a potential intersystem crossing - both these minima may provide access back to the ground state.³⁹ From that on, a barrierless relaxation leads to $[Int1]^{2-}$ at 36.9 kcal mol⁻¹ relative to $[6]^{2-}$. We note that boratanorcaradiene substructures, such as the one present in the key intermediate $[Int1]^{2-}$, have been characterized by XRD in the form of K[12] and Na[14] (Fig. 3c and d).38,40

All barriers of the following "walk reaction" on the S₀ surface are reachable at ambient conditions:⁴¹ [Int1]²⁻ undergoes a [1,5] sigmatropic shift of the boryl group to form [Int2]²⁻, which is 2.7 kcal mol⁻¹ more stable than [Int1]^{2–}. The corresponding barrier $[Int1]^{2-} \rightarrow [TS2]^{2-}$ amounts to $\Delta G^{\ddagger} = 17.6$ kcal mol⁻¹. A second, slightly endergonic [1,5]sigmatropic shift via [TS3]²⁻ gives $[Int3]^{2-}$ ($\Delta G^{\ddagger} = 19.7 \text{ kcal mol}^{-1}$, $\Delta G = 7.9 \text{ kcal mol}^{-1}$). In the last step, C–H activation from $[Int3]^{2-}$ leads to $[10]^{2-.42}$ The corresponding $[TS4]^{2-}$ lies 12.7 kcal mol⁻¹ above $[Int3]^{2-}$, so that the free-energy span to be covered from $[Int2]^{2-}$ to $[TS4]^{2-}$





Fig. 3 (a) Computed mechanism of the photoisomerization $[6]^{2-} \rightarrow [10]^{2-}$ and free energies of intermediates and transition states in their S₀ states. The energetically unfavorable and thus photochemically conducted step $[6]^{2-} \rightarrow [Int1]^{2-}$ is marked in red. (b) Literature-reported intermediates of the photoisomerization of triptycene 6^{C} and their calculated free energies (this work). (c) and (d) Known boratanorcaradienes K[12] and Na[14] obtained from photoreactions of tetraarylborates K[11] and Na[13], respectively. Carbon atoms marked with asterisks bear *t*-Bu substituents.



Fig. 4 Computed energy profiles of the photoinduced reaction $[6]^{2-} \rightarrow [Int1]^{2-}$ in the S₀ (gray), T₁ (red), and S₁ (blue) states. The dashed black line marks the position of $[TS1]^{2-}$ in the ground state; the blue arrows mark the electronic excitation and relaxation (the overall conclusion would remain the same if the relaxation occurred after a putative intersystem crossing from T₁).

is 20.6 kcal mol⁻¹.⁴³ The final product $[10]^{2^-}$ is essentially isoenergetic to the starting material $[6]^{2^-}$. Nevertheless, we consider a reverse reaction unlikely for the following reasons: (i) the energy barrier for the step $[Int1]^{2^-} \rightarrow [TS1]^{2^-}$ in the reverse reaction is significantly higher (25.1 kcal mol⁻¹; Fig. 3) than any barrier downstream $[Int1]^{2^-}$ for the forward reaction. (ii) Assuming that the barrier $[Int1]^{2^-} \rightarrow [TS1]^{2^-}$ could be overcome photochemically, the system would again be in the state described in Fig. 4 and should therefore preferentially fall back to $[Int1]^{2^-}$.

Conclusions

We achieved the synthesis of the parent dianionic 9,10diboratatriptycene $[6]^{2-}$ and its bridgehead derivatization. In the salt $[K(dme)][K(dme)_2][6]$, $[6]^{2-}$ acts as a chelating ligand toward both K⁺ ions, suggesting a possible future use as a trisbenzene ligand for mono- to trinuclear transition metal complexes.³ Abstraction of a bridgehead H⁻ substituent releases the free Lewis acid, which is sufficiently long-lived to be trapped by SMe₂; this transformation can be carried out at both

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bridgehead positions and eventually leads to the neutral boratriptycene-diadduct **9**. By using ditopic Lewis bases, this derivatization chemistry should provide an entry point to polymeric dibora(ta)triptycenes for materials science. The facile photoisomerization of $K_2[6]$ furnishes the diborabenzo[*a*]-fluoranthene derivative $K_2[10]$, which combines the two prominent structural motifs of 9,10-dihydro-9,10-diboraanthracene and 9*H*-9-borafluorene and belongs to a virtually unexplored class of B-doped polycyclic aromatic hydrocarbons.

Data availability

The datasets supporting this article have been uploaded as part of the ESI. \dagger

Author contributions

S. E. P. performed the experimental studies and characterized all new compounds. S. E. P. and J. G. performed the quantumchemical calculations. S. V. T., L. C., and M. We. assisted with the synthesis of compounds **9**, $[n-Bu_4N]_2[8]$, and $Li_2[5]$, respectively. M. B. performed the X-ray crystal structure analyses of the compounds $[K_2(dme)_3][3]$, $[Li(thf)_2][4]$, $[K(dme)]_2[6]$, and $[K(dme)][K(dme)_2][7]$. A. V. performed the X-ray crystal structure analysis of **9** \cdot CH₂Cl₂. H.-W. L., F. F., and M. Wa. supervised the project. The manuscript was written by S. E. P., J. G., and M. Wa. and edited by all co-authors.

Conflicts of interest

There are no conflicts to declare.

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