Closely related yet different: a borylene and its dimer are non-interconvertible but connected through reactivity†

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The self-stabilizing, tetrameric cyanoborylene [(cAAC)B(CN)]₄ (I, cAAC = 1-(2,6-diisopropylphenyl)-3,3,5,5-tetramethylpyrroloidin-2-ylidene) and its diborene relative, [(cAAC)CNB≡B(CN)(cAAC)] (II), both react with disulfides and diselenides to yield the corresponding cAAC-supported cyanoboron bis(chalcogenides). Furthermore, reactions of I or II with elemental sulfur and selenium in various stoichiometries provided access to a variety of cAAC-stabilized cyanoboron–chalcogen heterocycles, including a unique dithiaborirane, a diboraseselenirane, 1,3-dichalcogena-2,4-diboretanes, 1,3,4-trichalcogena-2,5-diborolanes and a rare six-membered 1,2,4,5-tetrathia-3,6-diborirane. Stepwise addition reactions and solution stability studies provided insights into the mechanism of these reactions and the subtle differences in reactivity observed between I and II.

Introduction

The dimerisation of carbenes [R₂C:] to form alkenes (Fig. 1A) is a fundamental reaction pattern in carbene chemistry and the basis of the so-called Wanzlick equilibrium. A similar dimerisation process has also been observed with heavier analogues of carbenes. However, despite the existence of isoelectronic group 13 species (LRL:), no carbene-analogous dimerisation (Fig. 1B) has been observed for group 13 species. Singly base-stabilised [LRL:] borylenes belong to the family of monovalent boron species discovered and recently reviewed by Bertrand et al. Perhaps the most definitive and well-defined example of this class of compounds is the linear borylene [(cAAC)CNB(SiMe₃)₂] (cAAC = 2-(2,6-diisopropylphenyl)-3,3-dimethyl-2-azaspiro[4,5]decan-1-ylidene) reported by Stephan and Bertrand. Over the past few years, however, our group has demonstrated that a number of doubly base-stabilized borylene species ([LL′RB:]), either through photodecarbonylation or the base-mediated deaggregation of tetrameric cyanoborylene [(cAAC)B(CN)]₄, is a tetramer (and synthetic equivalent) of the borylene [(cAAC)(CN)B:], while diborene II is a formal dimer of the same borylene. Having failed thus far to

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realise the interconversion of I and II through photolysis and/or heating, we were eager to test their respective reactivity with suitable reagents in order to define any differences or similarities, and hopefully gain definitive proof of their ability (or inability) to interconvert. Herein we present the reactivity of I and II with elemental chalcogens and dichalcogenides, based on our recently-reported reactions of these chalcogen reagents with B–B multiply-bonded species.12,13 Our results indicate that, although extensive similarities exist between borylene I and its formal dimer II, and they may therefore be viewed as close relatives, subtle differences in reactivity confirm that no Wanzlick-type equilibrium exists between the two.

Results and discussion

Synthetic studies and characterisation

The reaction of tetrameric borylene I with four equivalents of diorganyldichalcogenides, E2R2 (E = S, Se, R = Ph; E = Se, R = Me) in benzene proceeded selectively to the corresponding cAAC-supported cyanoboron dichalcogenides, [[cAAC]B(CN)(ER)2] (ER = SPh 1, SeMe 2, SePh 3), which were isolated in moderate to good yields (Scheme 2A). Whereas the reaction with diphenyl disulfide required prolonged heating at 80 °C to proceed to completion, reactions with diselenides proceeded at room temperature, the reaction with Se2Me2 being significantly faster than that with Se2Ph2. Compounds 1–3 showed 11B NMR resonances typical of sp3 hybridized boranes, with the bis(selenides) 2 (−18.4 ppm) and 3 (−14.4 and −15.8 ppm) exhibiting shifts significantly downfield from the bis(sulfide) 1 (−9.6 ppm). Furthermore, at room temperature the bis(phenylselenide) 3 displayed highly broadened 1H NMR ligand resonances as well as two distinct, very broad 11B NMR resonances (−14.4 and −15.8 ppm), which coalesced upon heating to 70 °C, indicating hindered rotation, presumably owing to steric interactions between the bulky cAAC substituents and the large phenylselenide ligands.

NMR spectroscopic data of reaction mixtures of diborene II and Ph4S2 or Ph4Se2 showed the monoboron bis(chalcogenides) 1 and 3 to be sole products of these reactions, independent of the stoichiometry used (Scheme 2B). These reactions proceeded much faster and at lower temperatures than the corresponding additions of dichalcogenides to the tetrameric borylene I. Although no intermediates were observed under these reaction conditions, it is likely that the formation of 1 and 3 from II proceeds via the successive 1,2-addition of the E–E σ-bond first across the B–B double bond and, subsequently, across the remaining B–B single bond. Although both compounds I and II reacted with Ph5Te2 at high temperatures, these reactions were rather unselective and did not yield any tractable products.

Compounds 1–3 readily crystallized from THF at room temperature, 1 as a colorless crystalline solid, and 2 and 3 as yellow crystals.‡ Fig. 2 shows the crystallographically determined structures of the bis(phenylselenides) 1 and 3, which crystallize in isomorphous unit cells. The B–C(CAAC) bond
length in disulfide 1 (1.6297(19) Å) is slightly elongated compared to that in diselenide 3 (1.597(9) Å). The B–C$_{CN}$ bond lengths (1: 1.584(2); 3: 1.580(9)) are similar to those found in other cAAC-supported cyanoboranes (1.574(5)–1.589(3) Å). While the B–E bond lengths (B–S: 1.9248(15), 1.9578(16) Å; B–Se: 2.109(7), 2.056(8) Å) are within the typical range for Lewis-base-stabilized boron organochalcogenides, such as Mader and Norman’s pyridine and phosphine adducts of 2,2′-bienzo[def]1,3,2]dithiaboroles (B–S: 1.899(5)–1.930(2) Å) or Wrackmeyer’s base-stabilized boron 1,2-diselenato-ortho-carboranes (B–Se: 2.031(6)–2.065(5) Å). While structurally unspectacular, species 1–3 represent the first examples of boron chalcogenides synthesized by the atom efficient insertion of a borylene into the E–E bond of a diorganodichalcogenide.

Stirring of a dark red suspension of tetrameric borylene I with elemental sulfur in a 1:1 boron-to-sulfur ratio in benzene for 5 d at room temperature resulted in a yellow suspension, which upon filtration and slow evaporation yielded compound 4 as a yellow crystalline solid (Scheme 3). Compound 4 displayed a $^{11}$B NMR singlet at −17.9 ppm and a single set of $^{1}$H NMR resonances for the cAAC ligand, suggesting a highly symmetrical species. Despite repeated recrystallization attempts, attempts, crystalline samples of 4 always contained 5–10% of another species displaying a higher field $^{11}$B NMR resonance at −9 ppm (vide infra). The analogous reaction with elemental selenium at 60 °C for 3 d similarly provided compound 5 as an orange crystalline solid in 70% yield. Compound 5 was isolated as a single species with an $^{11}$B NMR singlet at −33.5 ppm, shifted ca. 25 ppm upfield from that of 4, and a highly shielded, broad $^{77}$Se($^1$H) NMR resonance at −143.1 ppm. In solution at room temperature 5 partially isomerized to a second species presenting an $^{11}$B NMR shift at −31.8 ppm and slightly shifted $^1$H and $^{13}$C NMR resonances. High resolution mass spectrometry experiments performed on 4 and 5 provided molecular masses consistent with dimeric compounds of the formula [(cAAC)B(CN)E]$_2$ (E = S, Se, 5).

This was confirmed by X-ray crystallographic analyses, the results of which are displayed in Fig. 3. Compounds 4 and 5 crystallized in near-identical triclinic unit cells as centrosymmetric species presenting planar 1,3-dithia- and 1,3-diselenato-2,4-diborethen cores, respectively. The B$_2$S$_2$ ring in 4 is an approximate square with four almost identical B–S bonds (1.939(3) and 1.940(3) Å) and near-perpendicular B–S–B and S–B–S angles (85.21(12) and 94.80(12)$^\circ$, respectively), whereas the planar B$_2$Se$_2$ ring in 5 displays two slightly different B–Se bond lengths (2.069(4) and 2.100(4) Å) as well as near-perpendicular B–Se–B and Se–B–Se angles (85.13(17) and 94.87(17)$^\circ$, respectively). There are only a couple of structurally characterized 1,3,2,4-dichalcogenadiborethenes in the literature, all displaying sp$^2$ hybridized boron centers stabilized by π-donating amino substituents. Due to the lower coordination number at boron, the B–E bonds in Nöth’s 2,2,6,6-tetramethylpiperidine-supported B$_2$S$_2$ and B$_2$Se$_2$ heterocycles are ca. 0.08–0.10 Å shorter than in 4 and 5, while the B–E and E–E angles are ca. 3° narrower and wider, respectively.

For both compounds the plane consisting of the cyanoboron moiety and the π-framework of the cAAC ligand forms a ca. 80° angle with the B$_2$E$_2$ plane, with the respective ligands lying in trans-conformation with respect to the B$_2$E$_2$ core. While the solid-state structure of 5 shows a single conformational isomer, it remains unclear whether the observation of two isomers in solution is the result of a 180° rotation of one of the cAAC ligands around the B–C$_{cAAC}$ bond or the existence of a cis-
conformer, in which both cyano ligands occupy a cis-arrangement with respect to the heterocyclic core. The B–C_{cAAC} bond lengths (4: 1.639(4), 5: 1.606(5) Å) suggest that the cAAC ligand functions as a pure σ-donor ligand to the sp³ borane, unlike in borylene I, where a significant contribution of π-backbonding from the electron-rich B(I) center shortens the bond (ca. 1.47 Å). It is noteworthy that both the B–C_{cAAC} and B–C_{CN} bond lengths are significantly longer in 4 than in 5 (B–C_{CN}: 4 1.597(4), 5 1.577(6) Å).

Whereas 4 proved indefinitely stable in solution at room temperature, the ¹¹B NMR spectrum of an analytically pure sample of 5 in CD₂Cl₂ left at room temperature for 3 d showed the partial disappearance (<10%) of 5 concomitant with the appearance of a new boron-containing species at −22.8 ppm (Scheme 4A). A few crystals of this species could be isolated by reocrystallization from diethyl ether and analyzed by mass spectroscopy, revealing a compound of the formula [(cAAC)₂-B₂(CN)₂]Se [6]. X-ray crystallographic analysis of this compound showed that 6 is indeed a bis(cAAC)-stabilized 2,3-dicyano-2,3,1-diboraselenirane (Fig. 4). The structure of 6 is similar to that of the bis(NHC)-stabilized 2,3-dithienyl-2,3,1-diboraselenirane obtained by our group upon reaction of one equivalent of selenium with the corresponding diborene precursor,¹³ and likewise displays a trans-arrangement of the cyano and cAAC ligands with respect to the B₂Se core. Attempts to generate 6 from I using a 2 : 1 boron-to-selenium ratio failed, resulting instead in 50% conversion to the diselenadiboretane 5, with the remaining 50% of I left unreacted (Scheme 5). It thus appears that 6 is not accessible directly from borylene I but can be generated in solution at room temperature through loss of one selenium atom from 5, concomitant with B–B bond formation. This was also confirmed by closer inspection of the mass spectrum of compound 5 which showed mass patterns corresponding to 6.

Conversely, compound 6 was the only product of the reaction of diborene II with Se (2 : 1 ratio of boron to selenium) at room temperature in C₆H₆, as shown by the appearance of the ¹¹B NMR shift around −22 ppm (Scheme 5). Isolated samples of 6 showed asymmetric ¹H NMR cAAC resonances due to the presence of the chiral centers in B1 and B2. Attempts to detect the ⁷⁷Se NMR resonance of 6 failed due to coupling to the two adjacent quadrupolar boron nuclei. Similarly, the reaction of diborene II with S (2 : 1 ratio of boron to sulfur) at room temperature in C₆H₆ yielded the corresponding 2,3-dicyano-1,2,3-thiadiborirane 7 (Scheme 5), which presented an intense orange coloration in solution and is, to our knowledge, the first example of a thiadiborirane. The ¹¹B NMR shift of 7, observed at

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Scheme 4  Reversible ring contraction of 1,3-diselena-2,4-diboretane 5.

Scheme 5  Direct synthesis of 1,2,3-chalcogenadiboriranes from II only.

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Fig. 4  Crystallographically determined solid-state structures of the major one (88%) of the two overlapping disordered molecules of 6 in the asymmetric unit (left)§ and 7 (right). Atomic displacement ellipsoids depicted at 50% probability level. Hydrogen atoms and atomic displacement ellipsoids of peripheral substituents omitted for clarity. Selected bond lengths (Å) and angles (°) for 6: B1–C1 1.585(5), B2–C31 1.599(6), B1–C21 1.597(4), B2–C51 1.571(6), B1–B2 1.757(6), B1–Se1 2.030(4), B2–Se1 2.027(4); B1–Se1–B2 111.31(18), Se1–B1–B2 64.27(18), Se1–B2–B1 64.4219. For 7: B1–C1 1.575(6), B2–C31 1.580(6), B1–C21 1.557(6), B2–C51 1.549(7), B1–B2 1.777(6), B1–S1 1.860(5), B2–S1 1.869(5); B1–S1–B2 56.9(2), S1–B1–B2 61.8(2), S1–B2–B1 63.2(2).
slightly wider B/C14 lent of elemental sulfur to thiadiborirane/C0 33.5 ppm (Scheme 4B). Similarly, the addition of one equivalent of elemental sulfur to thiadiborirane/C0 3.5 diborolane 8 as the major reaction product in 71% isolated yield (Scheme 7B). This suggests that 8 is formed directly from I and independently of 4, rather than by insertion of a sulfur atom into the 1,3-disulfa-2,4-diboretane ring. Isolated samples of 8 showed two distinct 11B NMR resonances at −8.5 and −9.1 ppm in a 55 : 45 ratio, ca. 9 ppm downfield from that of 4. The 3H NMR spectrum of 8 displayed two isomers in that same ratio, each presenting two sets of cAAC resonances in a 1 : 1 ratio. Furthermore, for each isomer, the isopropyl resonances of the Dip residue were split into two sets of asymmetric resonances. The ratio of the two isomers was temperature-independent, suggesting they may be non-exchanging diastereomers.

The analogous 1,2,4-triselena-3,5-diborolane 9 was successfully isolated as an orange-colored solid as the major product (83% yield) of the reaction of borylene I with selenium in a 2 : 3 boron-to-selenium ratio in benzene at 60 °C (Scheme 7B). Compound 9 presented an 11B NMR resonance at −12.3 ppm, ca. 20 ppm downfield from that of 5. Similarly to 8, the 3H NMR resonances of the isopropyl groups of the Dip residue were split into two sets, indicative of an asymmetric structure. Attempts to detect the 77Se{1H} NMR resonances of 9 failed due to strong broadening of the signal. Furthermore, while the BS5 analogue 7 proved stable in solution, compound 9 was observed to fully decompose to the B2Se2 heterocycle 5 in benzene solution over a period of 4 days at room temperature (Scheme 8).

Both 1,2,4-trichalcogena-3,5-diborolanes 8 and 9 could also be obtained by reacting diborene II with three molar equivalents of elemental sulfur or selenium, respectively, or from the heating at 60 °C, only starting material was recovered (Scheme 7A).

In contrast, the room temperature reaction of tetrameric borylene I with elemental sulfur in a 2 : 3 boron-to-sulfur ratio yielded the corresponding yellow 1,2,4-trithia-3,5-diborolane 8 as the major reaction product in 71% isolated yield (Scheme 7B). This suggests that 8 is formed directly from I and independently of 4, rather than by insertion of a sulfur atom into the 1,3-disulfa-2,4-diboretane ring. Isolated samples of 8 showed two distinct 11B NMR resonances at −8.5 and −9.1 ppm in a 55 : 45 ratio, ca. 9 ppm downfield from that of 4. The 3H NMR spectrum of 8 displayed two isomers in that same ratio, each presenting two sets of cAAC resonances in a 1 : 1 ratio. Furthermore, for each isomer, the isopropyl resonances of the Dip residue were split into two sets of asymmetric resonances. The ratio of the two isomers was temperature-independent, suggesting they may be non-exchanging diastereomers.

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reaction of the corresponding B₂E heterocycles, 7 and 6, with two molar equivalents of sulfur or selenium, respectively (Scheme 7C).

Like their B₂E₂ counterparts, compounds 8 and 9 crystallized in isomorphous unit cells. For both compounds, the X-ray crystallographic structures (Fig. 5) show a five-membered central 1,2,4-trichalcogena-3,5-diborolane ring bearing one cyano and one cAAC ligand on each boron atom, arranged in a trans-configuration with respect to the B₂E₃ ring. While the structures thus represent a single (R,R,S,S) diastereomer, the observation of two non-exchanging isomers of 8 in solution suggests that the meso diastereomer of 8 may also be formed.

The formation of a single diastereomer of 9 may be attributable to the higher reaction temperature favoring the thermodynamic (R,R,S,S) diastereomer. The structure of 8 is reminiscent of that of the bis(NHC)-stabilized 3,5-dithienyl-1,2,4-trithia-3,5-diborolane obtained by the reductive insertion of three sulfur atoms into the B≡B double bond of a diborene precursor, with very similar B–S bond lengths and B–S–B and S–B–S angles.¹³

The 1,2,4-triselena-3,5-diborolane ring of 9 is reminiscent of that obtained by Tokitoh and co-workers upon irradiation of a boron bis(methylselenide) bound to a very bulky aryl ligand \( [\text{Tbt}_2\text{B}_2\text{Se}_3] \). Due to the sp³ hybridization of the cAAC-supported boron atoms in 9, however, the B-Se bonds (2.068(4), 2.086(4) Å) are slightly elongated and the Se–B–Se angle (109.4(2)°) is significantly more acute than in \( [\text{Tbt}_2\text{B}_2\text{Se}_3] \) (B–Se: 1.942(7), 1.926(8) Å; Se–B–Se: 118.8(4)°).²¹

NMR spectroscopic analysis of the crystallization filtrate of 1,2,4-trithia-3,5-diborolane 8 revealed the presence of another boron-containing species presenting an \(^{11}B\) NMR singlet at \(-11.2\) ppm and a single symmetrical cAAC ligand environment in its \(^{1}H\) NMR spectrum. Surmising that this may be a tetra-thiadiborinane resulting from a 1 : 4 boron-to-sulfur reaction, a scaled-up reaction with this stoichiometry was carried out (Scheme 9). The resulting orange suspension was filtered and the filtrate slowly evaporated to give compound 10 as a pale yellow solid in 53% isolated yield based on boron and its formulation confirmed by LC-MS.

Multiple recrystallizations of 10 from various solvents provided extremely thin, heavily twinned crystals, which diffracted too weakly to provide data suitable for structural discussion. However, the results of these X-ray crystallographic experiments provided conclusive proof of connectivity, confirming that 10 is indeed a bis[cAAC]-stabilized 3,6-dicyano-1,2,4,5-tetrasulfa-3,6-diborinane displaying a cis-arrangement of the cyano and cAAC ligands with respect to the central B₂S₄ ring, which displays a boat conformation (see Fig. S46 for the solid-state structure of 10). Attempts to insert a fourth sulfur atom into the isolated 1,2,4-trithia-3,5-diborolane 8 failed to yield 9, suggesting that, like 8, compound 9 forms directly from I, presumably by dimerization of a monomeric \([\text{cAAC}B(CN)\text{S}_2]\) intermediate. While 1,2,4,5-tetrasulfa-3,6-diborinanes have been postulated as minor reaction products by both Lappert and later Nöth, based solely on elemental analysis, mass spectrometry and IR spectroscopy data,²² compound 10 constitutes
the first structurally and NMR-spectroscopically characterized B$_2$S$_3$ heterocycle.

Mechanistic considerations

With all these data in hand, it was now possible to reassess the viability of the mechanism proposed by Tokitoh and co-workers for the formation of [(Tbt)$_2$B$_2$Se$_3$] from [(Tbt)B(SeMe)$_2$]. In a first step, the authors postulated the formation of a “(Tbt)B” borylene intermediate upon irradiation of [(Tbt)B(SeMe)$_2$]. This monomeric borylene may then dimerize to an electron-deficient diborene [(Tbt)B–B(Tbt)] (Scheme 10, path A), which then undergoes the reductive insertion of three atoms of selenium atoms with full cleavage of the B=Se bond. For path B, the authors proposed a monomeric [(Tbt)B–Se] intermediate resulting from the reaction of “(Tbt)B:” with Se, which dimerizes to a 1,3-diselena-2,4-diboretane and finally inserts the third selenium atom. Regarding path A, numerous attempts on our part have failed to convert borylene I into its diborene relative, compound II, under thermal and/or photolytic conditions. The fact that the three-membered B$_2$E heterocycles 6 and 7 can only be accessed directly from diborene II, but not from borylene I, is further evidence that, despite extensive overlap of reactivity outcomes, I and II do not interconvert.

Our previous report on the deaggregation of tetramer I by Lewis bases showed that only a relatively small and strong Lewis base – the NHC 1,3,4,5-tetramethylimidazol-2-ylidene – is able to break up the tetramer and generate the mixed base-stabilised [(cAAC)(NHC)B(CN)] borylene. This leads us to conclude that, while acting in all appearance as a source of monomeric borylene “[B]:” [(B = cAAC)(CN)B, Scheme 10], such a species is not, in fact, ever present in solution. Reactivity with I instead occurs by association of the reactant with the tetramer itself, ultimately causing it to deaggregate into mononuclear intermediates and products. For reactions involving I, the generation of intermediate [(B)=E] (E = S, Se) monomers following path B therefore seems the most likely. While the dimerization of [(cAAC)(CN)B=E] via bridging chalcogen atoms provides compounds 4 and 5, the latter do not insert another chalcogen atom to form 8 and 9, respectively, under the reaction conditions employed herein (Scheme 10). This means that the addition of the third chalcogen atom must occur prior to ring-closure, leading us to propose a monomeric dichalcogenaborirane [(B)E$_n$] intermediate, which upon reaction with [(B)E$_n$] yields the B$_2$E$_3$ heterocycles 8 and 9. Similarly, the B$_2$S$_4$ heterocycle 10 would result from the dimerization of [(B)S$_2$].

It is noteworthy that, while the selenium-based reactions were highly selective, those based on sulfur always yielded a mixture of products. For example, based on $^{11}$B NMR spectroscopic analysis of the final reaction mixture, the reaction of I with sulfur in a 2:3 boron-to-sulfur ratio yielded 8 in 70–85% selectivity at most, alongside the smaller and larger heterocycles 4 and 10, whereas the analogous reaction with selenium provided 9 in near-quantitative yield. The increased selectivity in the formation of 9 may be ascribed to the possibility of selenium de-insertion, which allows any B$_2$Se$_3$ heterocycle (5) formed in the course of the reaction to lose a Se atom, forming the diboraseleinirane 6, which can in turn be converted to 9 (Scheme 10). In contrast, any B$_2$S$_2$ (4) and B$_2$S$_4$ (10) heterocycles formed in the course of the reaction are inert towards sulfur de-insertion.

Scheme 10 Possible pathways to the boron–chalcogen heterocycles presented herein.

Scheme 11 Selected reactions showing the convergent and divergent reactivity of the two boron(I) isomers I and II with dichalcogenides and elemental selenium.
insertion and will therefore remain as by-products. Finally, the fact that the four- and five-membered BₓE₂ and BₓE₃ heterocycles are inert towards chalcogen insertion, whereas the three-membered BₓE heterocycles may be converted to both the larger BₓE₂ and BₓE₃ heterocycles, suggests that ring-expansion only proceeds by insertion of a single chalcogen atom or an E₂ unit into any remaining B–B bonds.

Conclusions

This first comparative study on the reactivity of two boron(i) compounds of the same empirical formula, the tetrameric, self-stabilizing cyanoborylene I and its dicyanoborene relative II, has demonstrated that, while both compounds provide access to the same products in many cases, this occurs via different pathways as shown in Scheme 11.

In reactions with dichalcogenides, both I and II yielded the mononuclear cyanoboron bis(chalcogenides) as sole products (Scheme 11). In the case of I this presumably occurs by insertion of a monomeric borylene into the chalcogen–chalcogen bond, whereas for II a successive 1,2-addition mechanism across the B–B double bond, resulting in full B–B bond cleavage is most likely at work.

Reactions with elemental sulfur and selenium were found to be highly dependent on the stoichiometry used. Reactions of I or II employing a 1 : 1 or 2 : 3 boron-to-chalcogen ratio yielded the corresponding 1,3-dichalcogena-2,4-diborolanes or 1,2,4-trichalcogena-3,5-diborolanes, respectively. A unique 1,2,3-thiadiborirane could only be accessed from the reaction of diborene precursor II with sulfur in a 2 : 1 boron-to-sulfur ratio, whereas the corresponding diboraselenirane was accessible both directly from the analogous reaction with selenium, and indirectly by de-insertion of one selenium atom from the four-membered 1,3-diselena-2,4-diborolane (Scheme 11). In the case of sulfur a rare example of a 1,2,4,5-tetrasulfa-3,6-diborirane was isolated from the reaction of borylene I with sulfur in a 1 : 2 boron-to-sulfur ratio.

Careful stepwise addition of chalcogen equivalents to either I or II and stability studies of the resulting heterocycles also gave insight into several mechanistic aspects of these reactions:

(i) Borylene-based reactions do not proceed via a diborene intermediate but likely via monomeric borachalcone and dichalchogenoborirane intermediates;

(ii) Ring-expansion reactions can only proceed by insertion of chalcogens into existing B–B bonds;

(iii) Ring-contraction is possible in the case of boron–selenium heterocycles only by de-insertion of 5e atoms.

Although the subtle divergences in reactivity between I and II provide confirmation that no Wanzlick-type equilibrium exists between a putative monomeric form of I and its B–B bonded dimer, diborene II, the question remains as to whether this is a result of the extremely stable, tetrameric constitution of borylene I. To date, I and II represent the only existing borylene/diborene pair with the same empirical formula, however, recent advances in the synthesis of borylenes will hopefully enable a more definitive answer to the question of a possible interconversion between [LRB] borylenes and [LRB=BRL] diborenies. Beyond the interest of such an interconversion from a fundamental point of view, its undeniable potential for providing a new, more reliable route towards hitherto inaccessible diborenies should continue to stimulate research into this area.

Conflicts of interest

The authors declare no conflicts of interest.

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Notes and references

† Compound 2 repeatedly crystallized as extremely thin, overlapping yellow plates, which could not be separated. X-ray crystallographic analysis of these provided proof of connectivity but data were of insufficient quality for further structural discussions (see ESI, Fig. S44†).

‡ The solid-state structure of molecule 6 presented a two-fold mirror disorder in a 88 : 12 ratio around the central diboraselenirane core, with the mirror plane running approximately through the plane containing (N₁, Se₁, N₂). The disordered 2,3-dicyano-2,3-diboroselenirane cores of both parts were freely refined. Fig. S45† provides an overlay of the two parts.

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