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Spirocyclic germanes *via* transannular insertion reactions of vinyl germynes into Si–Si bonds†‡Małgorzata Walewska,^a Judith Baumgartner,^{id} *^a Christoph Marschner,^{id} *^a
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The reactions of two cyclic germylene phosphane adducts with monosubstituted acetylenes caused the formation of spirocyclic germanes, which is postulated to occur by double acetylene insertion into germylene attached bonds. Further insertion of the formed cyclic divinylgermylene into transannular Si–Si or Si–Ge bonds provides the spirocyclic germanes. Thermal treatment of two germacyclopropenes, formed by the reaction of the two cyclic germylene phosphane adducts with tolane, also produced spirocyclogermanes. The structures of the latter require, however, a more complicated mechanistic proposal.

Introduction

In addition to the chemistry of carbenes,^{1,2} that of analogous heavier congeners^{3–9} has also become increasingly popular in recent years. In addition to constituting an interesting class of new transition metal ligands,^{10–15} several of these compounds are able to participate in small molecule bond activation reactions.^{16–19}

Our own interest in this compound class concentrates on silylated tetrylenes.^{20–34} In contrast to the well-studied N-heterocyclic heavy carbene analogues, these are not stabilised by π -donation of electron density from the attached nitrogen lone pairs into the empty p-orbitals, which leads to a much higher reactivity. To prevent dimerization or rearrangement reactions the formation of phosphane adducts proved to be a viable concept since the interaction between the tetrylene center and the donor is strong enough to suppress its decomposition but is still weak enough to allow dissociation. This is important, since the introduction of stronger donor molecules such as small N-heterocyclic carbenes (NHC)

switches the reactivity pattern of the respective adduct from “free tetrylene” to “tetrylenoid”.⁶

In a recent communication we reported that the reaction of phenylacetylene with the PMe_3 adduct of bis[tris(trimethylsilyl)silyl]germylene (**1**) occurs by regioselective insertion into a Si–Ge bond, thus yielding a PMe_3 adduct of a vinylgermylene (**2**).³³ The reaction of **1** with tolane, however, gives the less unexpected germirene **3**³³ (Scheme 1). The formation of **2** is very promising as it clearly indicates that weak donor adducts of divalent germanium are energetically similar or even more stable than isomeric structures with tetravalent germanium. This is a prerequisite for the potential use of divalent germanium compounds as catalysts in catalytic cycles with the intermediate formation of tetravalent germanes.

In a subsequent study, we could show that the formation of **2** involves the intermediate formation of a germirene similar to **3**.³⁴ The reason that **3** itself does not rearrange to the isomeric vinylgermylene is due to the fact that the silyl shift onto the phenyl substituted sp^2 -hybridised carbon is sterically less favored. Heating compound **3** to 150 °C gives silagermacyclobutene **4**, likely occurring *via* the intermediate formation of a vinylgermylene, which stabilizes by insertion into a Si– SiMe_3 bond (Scheme 1).³⁴

A related germirene with two less sterically demanding ethyl substituents, obtained by the reaction of **1** with 3-hexyne, was shown to undergo a rearrangement to a structure analogous to **4** even at ambient temperature.³⁴

A number of related reactions were reported recently for silylenes and stannylenes. Rieger and co-workers reported the insertion of ethylene into the Si–Si(SiMe_3)₃ bond of a silylated silylene,³⁵ while Kato and Baceiredo described the insertion of an olefin into a Si–H bond of their stabilised silylene also involving the previous formation of a silirene.³⁶ Insertion of tolane into the Sn–B and Ge–B bonds of a stannylene and a

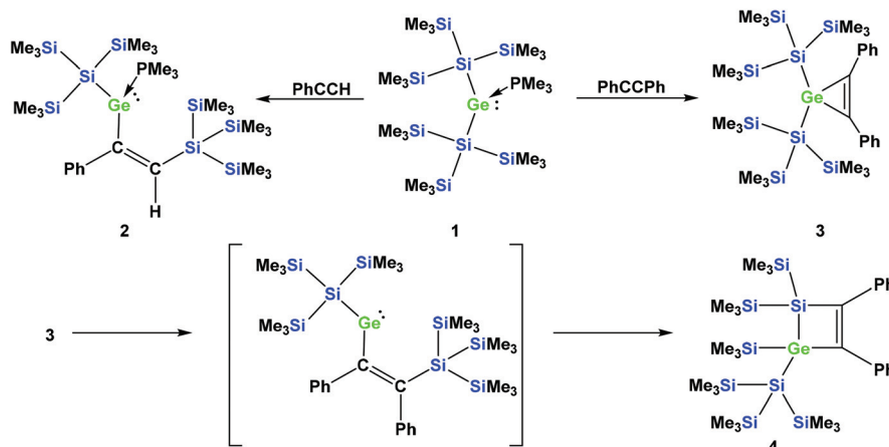
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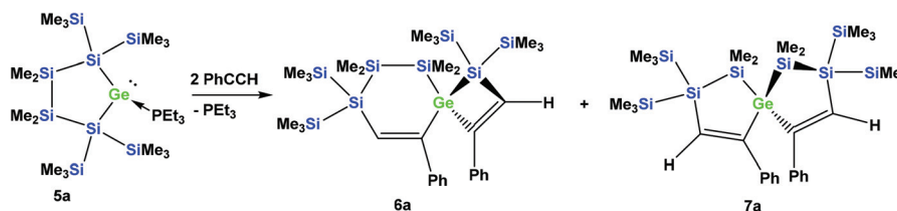
†Dedicated to Phil Power, a main-group chemist extraordinaire and a great person.

‡Electronic supplementary information (ESI) available: Tables containing the crystallographic information of compounds **6a**, **6b**, **7c**, **10a**, **13a**, and **13b** and the absolute and relative energies of compounds **6–9** and **10–16**; figures with relative energy diagrams and the correlation of calculated ²⁹Si NMR shifts; xyz-files of the calculated structures. CCDC 1450265, 1450269–1450273. For the ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c8dt00315g



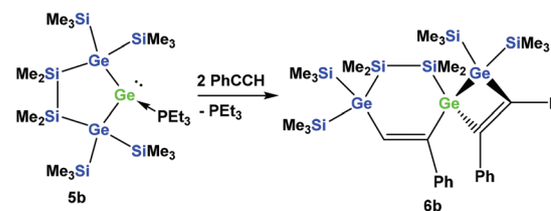


Scheme 1 Reactions of germylene adduct **1** with either phenylacetylene (top left) or toluene (top right). Thermally-induced rearrangement of germirene **3** to silgermacyclobutene **4** (bottom part).



Scheme 2 Reaction of cyclic germylene phosphane adduct **5a** with phenylacetylene.

germylene, respectively, was reported by Aldridge and co-workers.³⁷ An example where two germylens react with several different disubstituted alkynes to 1,2-digermacyclobut-3-enes was observed recently by Ketkov, Dostál and co-workers.¹⁹ A different approach to vinylgermylenes was communicated by Rivard and co-workers utilizing N-heterocyclic olefins.³⁸ Most recently, Sasamori and co-workers reported the use of a digermine for the catalytic regioselective cyclotrimerization of terminal alkynes. Their mechanistic explanation for this reaction also involves alkyne insertion into the bonds attached to germylens.³⁹



Scheme 3 Reaction of digermylated cyclic germylene phosphane adduct **5b** with phenylacetylene.

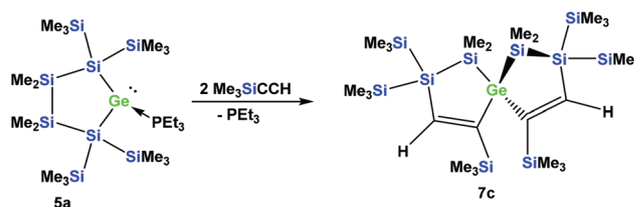
Results and discussion

Reactions of cyclic germylene adducts with two alkynes

In order to study whether the unexpected migratory insertion behavior of **1** is a general reactivity feature of silylated germylens, we further investigated the behavior of cyclic germylene adducts **5a** and **5b**.^{28,31} The reaction of the disilylated germylene **5a** with two equivalents of phenylacetylene was found to give a mixture of the two spirocyclic compounds **6a** and **7a** in an approximate ratio of 1 : 1 (Scheme 2).

However, the reaction of two equivalents of phenylacetylene with the analogous digermylated germylene adduct **5b** gave only a single product, namely the asymmetric spirocyclic **6b** (Scheme 3).

The fact that subtle differences are essential for the reactivity in these reactions was shown by the reaction of **5a** with two

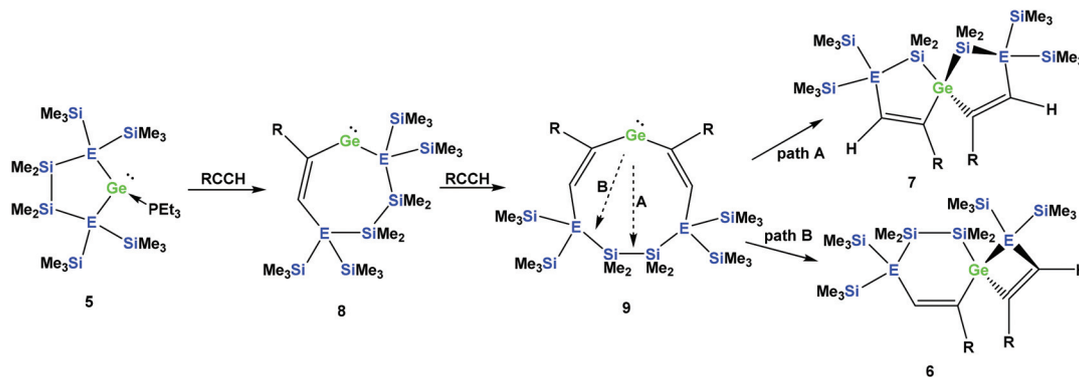


Scheme 4 Reaction of cyclic germylene phosphane adduct **5a** with trimethylsilylacetylene.

equivalents of Me_3SiCCH , which also gave only a single product, but in this case the symmetric spirocyclic compound **7c** was formed (Scheme 4).

It seems reasonable to assume that all these reactions proceed in a very similar manner. The first steps are likely to





Scheme 5 Assumed reaction mechanism of germylene adducts **5a** and **5b** with monosubstituted acetylenes to spirocyclic compounds (a: E = Si and R = Ph; b: E = Ge and R = Ph).

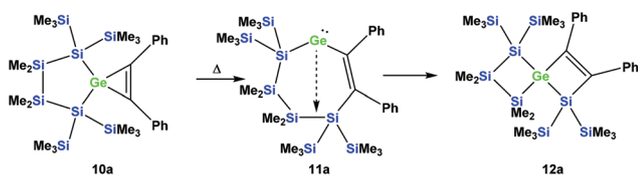
be double insertion reactions of the acetylene units into the Si–Ge or Ge–Ge bonds of **5a** and **5b** to give cyclic divinylated germylenes **9a**, **9b**, and **9c** (Scheme 5). The regiochemistry of these steps is the same as that described above for acyclic germylene adduct **1**.³² The resulting nine-membered ring germylenes are likely to be conformationally rather flexible. The subsequent intramolecular insertion of the released free germylenes **9** can occur either into transannular Me₂Si–SiMe₂ or Me₂Si–E(SiMe₃)₂ (E = Si, Ge) bonds (Scheme 5).

Attempts to run the reactions of **5a** and **5b** with only one equivalent of phenylacetylene led to more complicated mixtures. In the case of **5b** the ²⁹Si NMR spectrum showed the presence of spirocycle **6b** derived from **9b** and the starting material **5b**. In addition, signals which likely belong to a [3.3]-spirocycle derived from the mono-inserted germylene **8b**

(Scheme 5) were observed. This suggests that germylenes **5b** and **8b** are of similar reactivity and compete for phenylacetylene.

Depending on into which bond the germylene insertion occurs, either symmetric [4.4]-spirocyclic compounds like **7a** or **7c** or [5.3]-spirocyclic ones like **6a** or **6b** are formed (Schemes 2–5). The initial structural assignment was carried out on the basis of single crystal XRD analyses of **6a**, **6b**, and **7c**. Compounds **6a** and **7a** were obtained as a mixture and the signals in the ²⁹Si spectrum are not easily assignable. However, the spirocyclic compound **7c**, which is very similar to **7a** was formed selectively and displays a rather simple ²⁹Si NMR spectrum with five signals (Table 1). Apart from the trimethylsilyl vinyl signal at –6.9 ppm, the two geminal trimethylsilyl groups were observed at –12.1 and –12.7 ppm, and the dimethylsilyl unit was found at –28.2 ppm with the remaining Si(SiMe₃)₂ signals located at –66.4 ppm. Assuming that the set of very similar signals (–12.1, –12.5, –27.3, and –66.9 ppm) represent compound **7a**, which were then subtracted from the spectrum of the mixture of **6a** and **7a**, the remaining signals can be assigned to be those of **6a** (–9.6, –10.9, –11.4, –11.8, –35.1, –46.1, –46.8, and –86.6 ppm).

The four signals between –9.6 and –11.8 ppm are associated with the trimethylsilyl groups and the signal at –86.6 ppm belongs to the Si(SiMe₃)₂ in the six-membered



Scheme 6 Expected thermally-induced reaction pathway of **10a** to give [3.3]-spirocyclic vinylgermane **12a**.

Table 1 ²⁹Si NMR spectroscopic data of the acyclic germylene adducts and reaction products

Compound	²⁹ Si E(SiMe ₃)	²⁹ Si SiMe ₂	²⁹ Si Si(SiMe ₃) ₂	²⁹ Si Other SiMe ₃
5a ³¹	–4.3/–8.5	–22.3	–126.0	
5b ³¹	–2.0/–4.1	–16.9	n.a.	
6a	–9.6/–10.9/–11.4/–11.8	–35.1 (GeSiMe ₂)/–46.1 (SiSiMe ₂)	–46.8 (GeSiC)/–86.6 (Me ₂ SiSiC)	
7a	–12.1/–12.5	–27.3	–66.9	
6b	–3.3/–3.7/–4.7/–5.0	–35.5/–37.3	n.a.	
7c	–12.1/–12.7	–28.2	–66.4	–6.9
10a ³¹	–7.3	–30.3	–120.1	
10b	–1.5	–23.4	n.a.	
12a	–2.9/–7.6/–9.3/–12.5	–22.6/–35.3	–60.1 (CSi(SiMe ₃) ₂)/–119.1 (SiSi(SiMe ₃) ₂)	
13a	–8.6/–8.7/–9.5/–9.6	9.9, 3.8	–94.9 (GeSi(SiMe ₃) ₂)/–108.2 (SiSi(SiMe ₃) ₂)	
13b	–2.5/–2.9/–3.2/–4.5	3.5 (GeSiC)/18.4 (GeSiGe)	n.a.	



ring. The other $\text{Si}(\text{SiMe}_3)_2$ unit is assigned to the signal at -46.8 ppm, which shows the rather typical strong down-field shift in cyclotetrasilane rings.^{40,41} The two remaining signals at -35.1 and -46.1 ppm are those of the SiMe_2 groups in the six-membered ring, with the one at -35.1 being attached to the central germanium atom, which usually leads to a down-field shifted signal compared to a related unit attached to a silicon atom. This assignment of the signals of **6a** is fully consistent with what we observe for compound **6b**, which is isostructural to **6a**.

Compared to **6a**, for **6b** the four trimethylsilyl signals are shifted some 6 ppm down-field (Table 1), while the SiMe_2 signals appear at -35.5 and -37.3 ppm, reflecting the fact that in contrast to the situation of **6a**, both groups are attached to a germanium atom.

The striking dependence of the product formation on subtle differences in the reactants that are shown in Schemes 2–4 prompted us to investigate these reactions in detail using DFT calculations at the B3LYP-D3 level.⁴² Based on our previous investigations,³³ we supposed that cyclic bis-vinylgermylenes **9** (Scheme 5 and Fig. 1) are key compounds for the formation of the two different germanium spiro-cycles **7** (*via* path A, Fig. 1) and **6** (*via* path B). In both cases, the intramolecular

insertion reaction of the dicoordinated germanium atom into the remote Si–Si or Si–E (E = Si, Ge) in compounds **9** was calculated to be a strongly exergonic process (see Fig. 1).

As expected, the formation of the less strained [4.4]-spirocyclic compounds **7** is in each case thermodynamically favored over the [5.3]-spirocyclic isomers **6** ($\Delta\Delta G$ (**7a/6a**) = 52 kJ mol^{-1}), ($\Delta\Delta G$ (**7b/6b**) = 55 kJ mol^{-1}), and ($\Delta\Delta G$ (**7c/6c**) = 65 kJ mol^{-1}) (Fig. 1). Therefore, the computational results indicated that the generation of [5.3]-spirocycles **6** is only possible under kinetic control. We found that both types of insertion reactions proceeded in one concerted step *via* the transition states TS(**9/7**) and TS(**9/6**) (Fig. 1 and Table 2).

The calculated barriers ΔG^\ddagger for both processes are between 58 and 71 kJ mol^{-1} , in qualitative agreement with the reactions that proceed under ambient conditions (see Fig. 1). In view of the high barriers for the back reaction (155 – 168 kJ mol^{-1} for **6** \rightarrow **9**, and 211 – 222 kJ mol^{-1} for **7** \rightarrow **9**, Fig. 1), the establishment of a thermodynamic equilibrium between the three isomers **9**–**6** is unlikely. For the phenyl-substituted germylene **9a** the energy difference $\Delta\Delta G^\ddagger$ between both competing reaction paths is merely 3 kJ mol^{-1} in favor of the formation of the [5.3]-spirocycle **6a**. Based on a basic Boltzmann treatment this would suggest the formation of a product mixture of the two

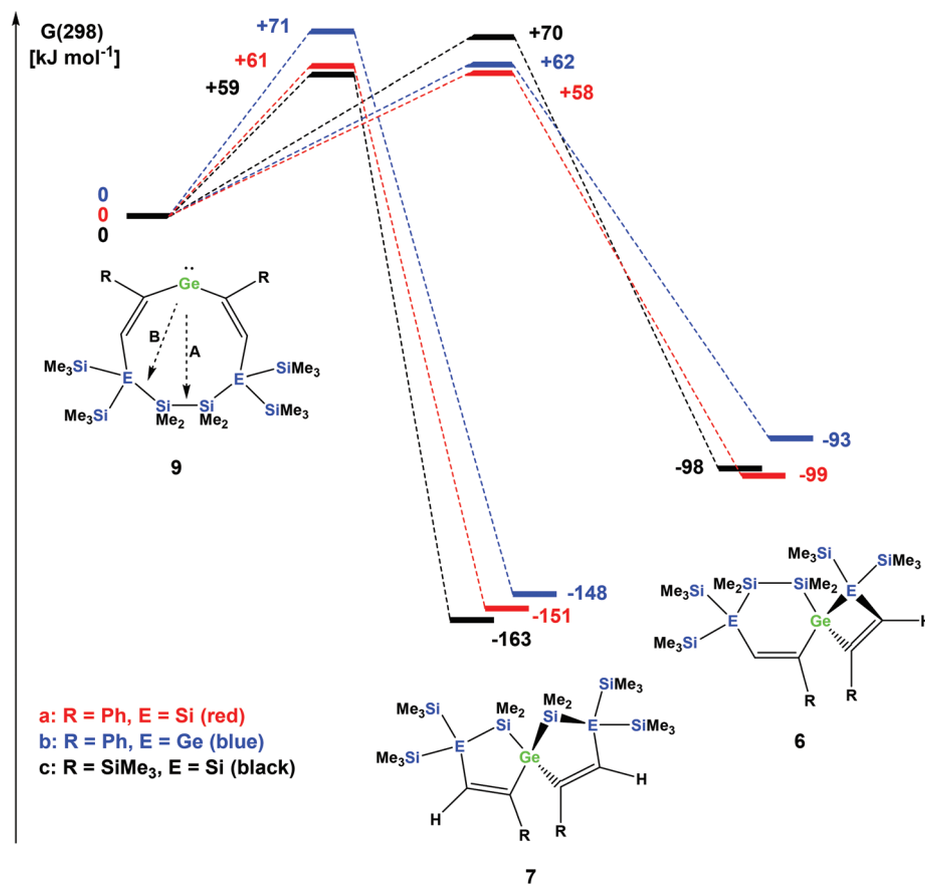


Fig. 1 Calculated reaction coordinates for the intramolecular bond insertion reaction of germylenes **9**; A: insertion into the Si–Si bond to give [4.4] germanium spirocycles **7**; B: insertion into the Si–E bond to give [5.3] germanium spirocycles **6** (free energy differences at 298 K $G(298)$, at B3LYP-D3/6-311G(d,p)(Si,C,H); def2tzvp(Ge)).



Table 2 Calculated energies E of compounds **6** and **7** and the connecting transition states relative to the corresponding starting germynes **9** (in kJ mol^{-1} , ΔG at 298 K is given in parenthesis)

Compound	TS germylene 9/spiro [5.5] 7	Spiro [5.5] 7	TS germylene 9/spiro [6.4] 6	Spiro [6.4] 6
R = Ph, E = Si	+54 (+61)	-152 (-151)	+61 (+58)	-101 (-99)
R = Ph, E = Ge	+59 (+71)	-150 (-148)	+57 (+62)	-103 (-93)
R = SiMe ₃ , E = Si	+52 (+59)	-165 (-163)	+64 (+70)	-104 (-98)

isomers **6a** and **7a** of approximately 23 : 77. This is in qualitative agreement with the experimental results that the reaction of germylene **5a** with phenylacetylene provides a mixture of the isomers **6a** and **7a** (Scheme 2). For the reaction of germylene **5b** with phenylacetylene the bis-vinylgermylene **9b** is the most likely intermediate. The intramolecular insertion of the dicoordinated germanium atom into the weaker Ge–Si (path B, Scheme 5) is favored over the insertion into the Si–Si bond (path A) by $\Delta\Delta G^\ddagger = 9 \text{ kJ mol}^{-1}$. Under the conditions of kinetic control at room temperature, this energy difference suggests a nearly exclusive formation of the less stable [5.3] germanium spirocycle **6b** (ratio **7b** : **6b** = 3 : 97), in perfect agreement with the experimental results. The situation is different for the reaction of germylene **5a** with trimethylsilylacetylene (Scheme 4). In the case of the corresponding cyclic bis-vinylgermylene **9c**, we found for the generation of the more stable [4.4]-spirocycle **7c** a barrier which is by $\Delta\Delta G^\ddagger = 11 \text{ kJ mol}^{-1}$ lower than that predicted for the competing reaction channel to give the [5.3] isomer **6c** (Fig. 1). Therefore, our computations suggest the almost exclusive formation of the more stable isomer **7c** (ratio **7c** : **6c** = 99 : 1) in accordance with the experiment.

Rearrangement reactions of spirocyclic germacyclopropenes

Cyclic germylene adducts **5a** and **5b** can be reacted with toluene and the expected spirocyclic germacyclopropenes **10a**³⁴ and **10b** are obtained. Further attempts to react **5a** and **5b** with bis(trimethylsilyl)acetylene, however, did not give the expected germacyclopropenes.

As was described above for the temperature-induced rearrangement of the germacyclopropene **3**,³⁴ compound **10a** was also subjected to 100 °C to facilitate the insertion step of the olefin group into the Ge–Si bond. We expected that upon the formation of the seven-membered cyclic germylene **11a**, the insertion into a transannular Si–Si bond would give the [3.3]-spirocyclic compound **12a** with the two bis(trimethylsilyl)silylene units in different rings (Scheme 6).

According to the ²⁹Si NMR spectroscopic analysis of the reaction solution, two products were formed with two signal sets of eight symmetry non-equivalent ²⁹Si signals each (Fig. 2). Although it was not possible to separate the mixture of crystalline products, we were nevertheless able to obtain a crystal of one of the main products which was subjected to a single crystal XRD study. According to this analysis a spirocyclic product was obtained again.

However, the product was not the expected **12a** following the pathway as outlined above (Scheme 6). Despite the fact that the crystal structure solution showed indeed the for-

mation of a [3.3]-spirocyclic vinylgermane (**13a**), it is not the expected but an isomeric one. Compound **13a** features both bis(trimethylsilyl)silylene and one dimethylsilylene units in one ring and another dimethylsilylene and 1,2-diphenylethylene in the other one (Scheme 7). However, analysis of the ²⁹Si{¹H} NMR spectrum of the mixture suggests that in addition to **13a** another product is present in which the SiMe₂ units are part of a five-membered ring. Dimethylsilylene units are quite sensitive to the ring-size and their ²⁹Si chemical shift is typically around -10 ppm when incorporated into a four-membered ring and can be found further up-field in larger rings.

Also subjecting digermylgermacyclopropene **10b**, easily available by the reaction of **5b** with toluene, to a thermally-induced rearrangement gave only a single product (**13b**), which is isosteric to **13a** and features the two additional germanium atoms bearing two trimethylsilyl groups each (Scheme 8). Scheme 9 (top part) shows a mechanistic proposal for the formation of **13a** and **13b**. We assume that indeed the reactions first follow the steps outlined above: (1) insertion of the 1,2-diphenylacetylene unit into the Ge–E bond to give **11a** and **11b** and (2) germylene insertion into the transannular Si–E bond to give **12a** for the silicon case and **12b** for E = Ge (Scheme 9). At the stage of **12a** and **12b** we assume that a cycloreversion reaction leads to the formation of the isomeric seven-membered cyclic vinylgermylenes **14a** and **14b** (Scheme 9). Also, if now a reversal of the germacyclopropene-vinylgermylene rearrangement occurs, the resulting germacyclopropenes (**15a** and **15b**) have the possibility to shift the acetylene unit to the Ge–SiMe₂ bond, this way forming germynes **16a** and **16b** (Scheme 9). If the latter molecules (**16a/16b**) now insert into the transannular Me₂Si–SiMe₂ bonds the observed spirocyclic germanes **13a** and **13b** are formed (Scheme 9, top part). If the insertion of the germylene (**11a** or **14a**) occurs instead of the endocyclic Si–Si bond into an exocyclic Si–SiMe₃ bond either **17a** or **18a** would be formed (Scheme 9, bottom part). As both compounds feature five-membered rings consistent with the ²⁹Si NMR resonances in Fig. 2 they are likely candidates for the unknown compound as shown in Scheme 7. This assignment is further supported by ²⁹Si NMR chemical shift calculations. For a test set that consists of the isolated and well-characterised compounds **6a**, **7a**, **10a**, **13a** and **13b**, we found that the GIAO/M06L/6-311G(2d,p)⁴² method allows a reliable prediction of the ²⁹Si NMR chemical shifts for this family of oligosilanyl compounds (see ESI, Fig. S3†). The characteristic ²⁹Si NMR chemical shifts that were predicted for the [3.3]-spirocyclic compound **12a** deviate significantly from the experimental data of the unknown com-



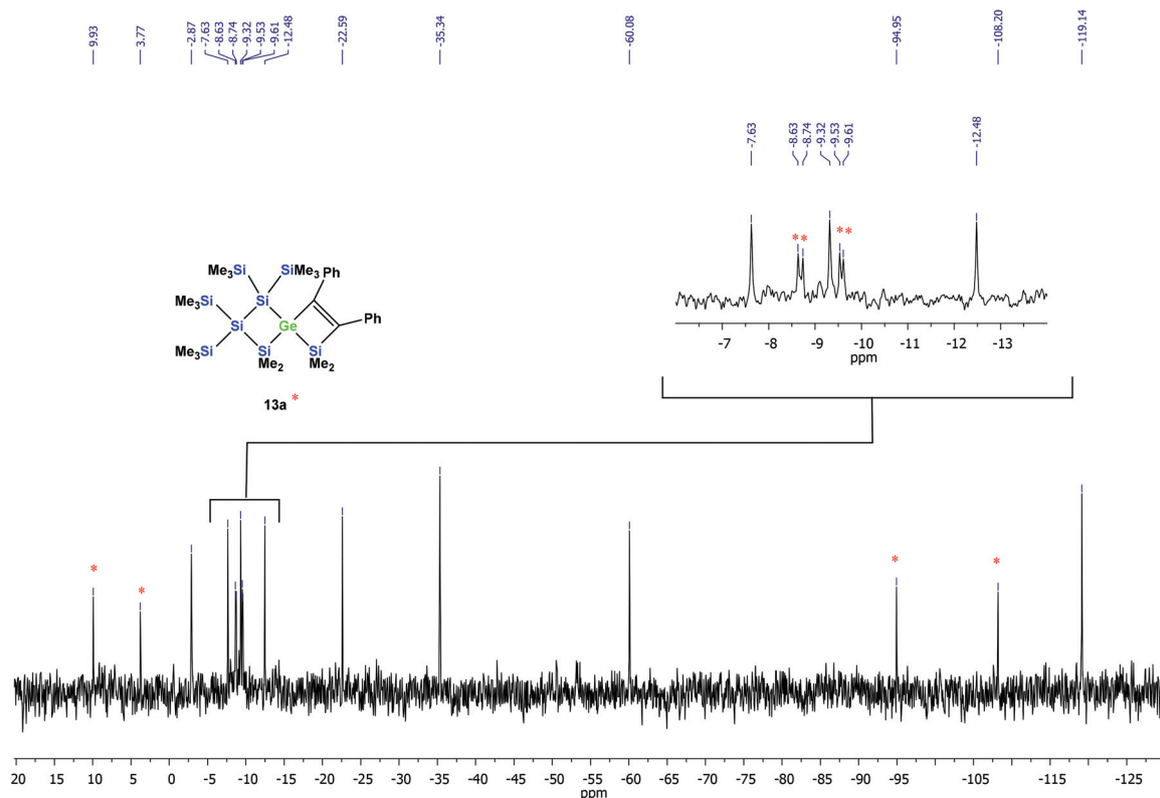
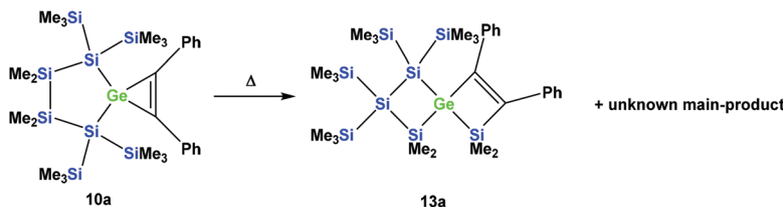
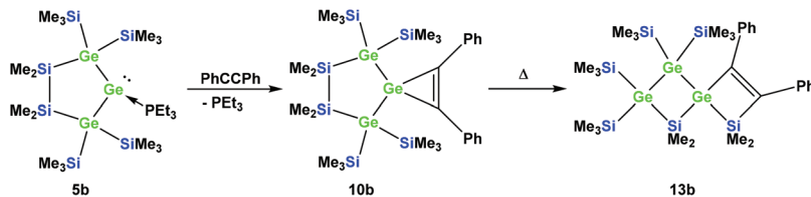


Fig. 2 $^{29}\text{Si}\{^1\text{H}\}$ INEPT NMR spectrum of the thermally promoted reaction of compound **10a**. The peaks assigned to compound **13a** are marked with an asterisk.



Scheme 7 Thermal rearrangement of the [4.2]-spirocyclic germacyclopropene **10a** to the [3.3]-spirocyclic vinylgermane **13a** and an unknown product.



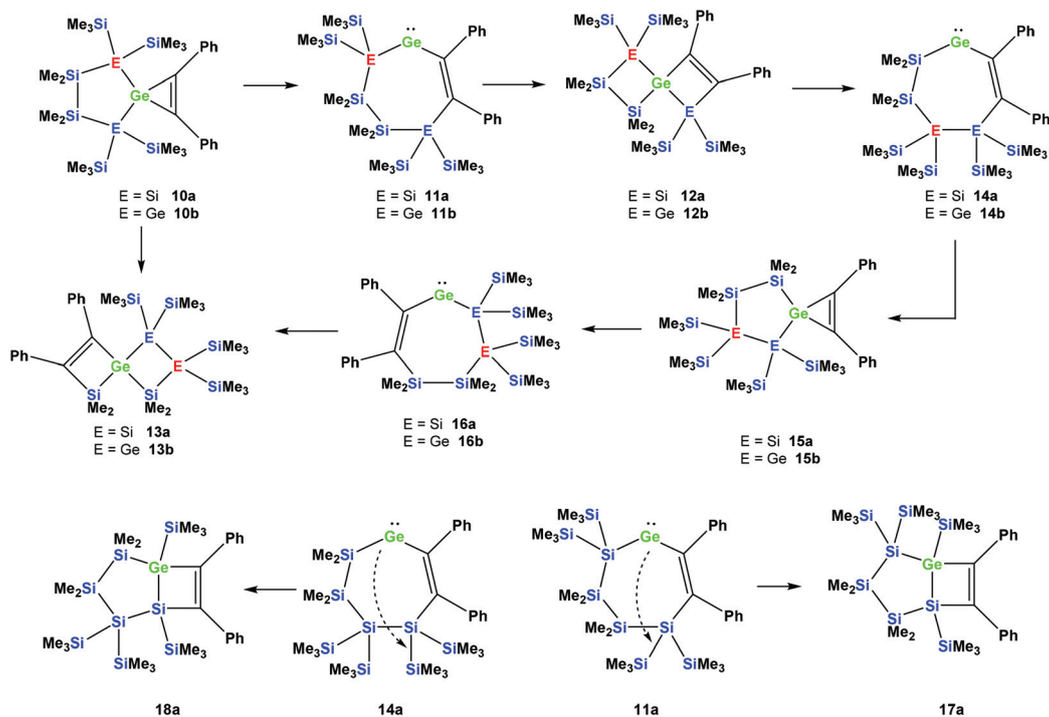
Scheme 8 Reaction of **5b** with toluene gave the expected [4.2]-spirocyclic germacyclopropene **10b**, which upon heating rearranged into the [3.3]-spirocyclic vinylgermane **13b**.

compound. The predicted ^{29}Si NMR chemical shifts for the annulated bicycles **17a** and **18a**, however, agree well with the experimental data (see Fig. S3† in the ESI).

Recently, we have reported a structure very similar to **17a**, which was formed in the reaction of compound **5a** with 1,4-bis

(trimethylsilyl)butadiyne.³¹ We interpreted the formation of this product as a cycloaddition reaction of an alkyne across a cyclic silagermene.³¹ Given the evidence of the chemistry described in the current paper, we might be inclined to revise this initial mechanistic explanation.





Scheme 9 Mechanistic rationale for the formation of **13a** and **13b** from **10a** and **10b**, respectively (top part), as well as for the formation of compounds with five-membered rings such as compounds **17a** and **18a** (bottom part).

Quantum mechanical calculations at the B3LYP-D3⁴² level of theory support the mechanistic proposal for the formation of **17** or **18**. For both cases, E = Si and E = Ge, the energy diagrams are very similar (see Fig. S1 and S2 in the ESI[†]): All spirocycles (**10a**, **10b**, **12a**, **12b**, **15a**, **15b**, **13a**, and **13b**) and the ring-annulated compounds **17a** and **18a** are lower in energy and well separated from the seven-membered ring germylenes (**11a**, **11b**, **14a**, **14b**, **16a**, and **16b**). The reason for this energetic preference is clearly connected to the formation of two additional bonds in the bicyclic compounds involving tetra-coordinated germanium. It is, however, noteworthy that for both investigated potential energy surfaces, the energy difference between Ge(II) and Ge(IV) compounds is small enough to allow an easy thermal switch between both coordination modes.

The ratio and the different constitutions of the obtained products are probably determined by the subtle interplay between the kinetic and thermodynamic factors. Although a thorough investigation of all barriers involved in the rearrangements suggested in Scheme 9 is beyond the scope of the present investigation, it seems reasonable to assume that thermodynamic factors such as ring strain and bond strengths are important for the product formation. The [2.4]-spirocyclic compounds **10** and **15** that involve unsaturated three-membered rings are clearly destabilized compared to the four-membered [3.3]-spirocycles **12** and **13**. In addition, for the pair **10a/13a** an estimate based on the calculated strengths of the Si–Si, Si–Ge, Si–C and Ge–C bonds⁴³ favors compounds **13** by 11 kJ mol^{−1}.

Interestingly, when dispersion interactions are neglected by using only the B3LYP functional, the seven-membered rings

(**11a**, **11b**, **14a**, **14b**, **16a**, and **16b**) lose notable relative stability ($\Delta\Delta G = 20\text{--}50$ kJ mol^{−1}), whereas the relative energies of the spirocyclic compounds (**10a**, **10b**, **12a**, **12b**, **15a**, **15b**, **13a**, and **13b**) only show little changes of $\Delta\Delta G = 1\text{--}8$ kJ mol^{−1} (see Fig. S1 and S2 in the ESI[†]). This fact indicates the importance

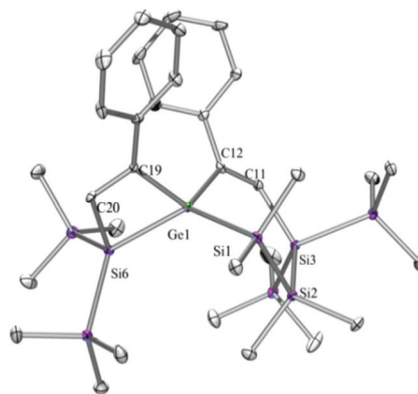


Fig. 3 Molecular structure of **6a** (thermal ellipsoid plot drawn at the 30% probability level). All hydrogen atoms are omitted for clarity (bond lengths in Å and angles in degrees). Ge(1)–C(12) 1.987(5), Ge(1)–C(19) 2.001(5), Ge(1)–Si(1) 2.3899(15), Ge(1)–Si(6) 2.4276(15), Si(1)–C(1) 1.882(5), Si(1)–Si(2) 2.3271(19), Si(3)–C(11) 1.889(5), Si(3)–Si(4) 2.3441(19), C(11)–C(12) 1.342(7), C(19)–C(20) 1.346(7), C(12)–Ge(1)–C(19) 116.7(2), C(12)–Ge(1)–Si(1) 103.51(14), C(19)–Ge(1)–Si(1) 111.31(14), C(12)–Ge(1)–Si(6) 120.67(14), C(19)–Ge(1)–Si(6) 72.70(14), Si(1)–Ge(1)–Si(6) 128.48(5), C(12)–C(11)–Si(3) 134.7(4), C(11)–C(12)–Ge(1) 123.5(4), C(20)–C(19)–Ge(1) 103.6(3), C(19)–C(20)–Si(6) 108.6(4).



of stabilizing dispersion contributions during this rearrangement, without which the barriers would be too high to be overcome under the experimental conditions.

If the formation of spirocyclic compounds **13a** and **13b** really follows the mechanism outlined in Scheme 9, three different vinylgermylenes are involved. All attempts to capture one of these germylenes as a stable adduct by the addition of several different NHCs failed. The presence of NHCs did not change the outcome of the reaction at all.

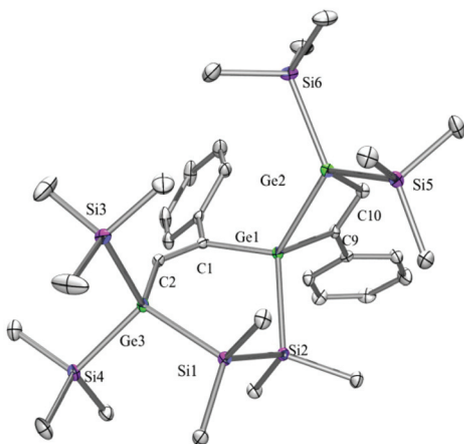


Fig. 4 Molecular structure of **6b** (thermal ellipsoid plot drawn at the 30% probability level). All hydrogen atoms are omitted for clarity (bond lengths in Å and angles in degrees). Ge(1)–C(1) 1.989(3), Ge(1)–C(9) 1.995(3), Ge(1)–Si(2) 2.3937(9), Ge(1)–Ge(2) 2.4741(6), Ge(2)–C(10) 1.976(3), Ge(2)–Si(6) 2.4025(9), Ge(3)–C(2) 1.973(3), Ge(3)–Si(3) 2.3809(9), Si(2)–C(20) 1.869(3), Si(2)–Si(1) 2.3275(12), C(1)–Ge(1)–C(9) 116.33(11), C(1)–Ge(1)–Si(2) 104.54(8), C(9)–Ge(1)–Si(2) 111.73(8), C(1)–Ge(1)–Ge(2) 119.71(7), C(9)–Ge(1)–Ge(2) 74.07(8), Si(2)–Ge(1)–Ge(2) 127.29(3), C(10)–Ge(2)–Si(5) 109.22(8), C(10)–Ge(2)–Si(6) 109.10(8).

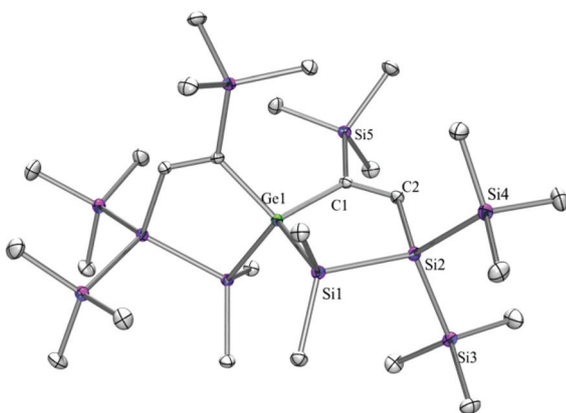


Fig. 5 Molecular structure of **7c** (thermal ellipsoid plot drawn at the 30% probability level). All hydrogen atoms are omitted for clarity (bond lengths in Å and angles in degrees). Ge(1)–C(1) 1.9870(11), Ge(1)–Si(1) 2.4017(5), Si(1)–C(3) 1.8805(13), Si(1)–Si(2) 2.3387(6), Si(2)–C(2) 1.8953(12), C(1)–C(2) 1.3518(17), C(1)–Ge(1)–Si(1) 99.81(4), C(1)–C(2)–Si(2) 126.64(9).

X-Ray crystallography

Compounds **6a** (Fig. 3), **6b** (Fig. 4), **7c** (Fig. 5), **13a** (Fig. 6), and **13b** (Fig. 7) all obtained by the conversion of the cyclic germylenes **5a** or **5b** were characterized by single crystal X-ray diffraction analysis. The structure of **10a** was determined previously.³¹ Although we obtained a fair number of structures of different germanium spirocyclic compounds (**6a**, **6b**, **7c**, **10a**, **13a**, and **13b**), they show no big differences regarding their structural features and do not appear to be very strained. The

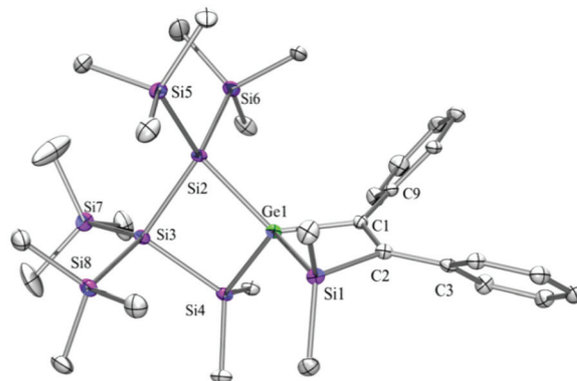


Fig. 6 Molecular structure of **13a** (thermal ellipsoid plot drawn at the 30% probability level). All hydrogen atoms are omitted for clarity (bond lengths in Å and angles in degrees). Ge(1)–C(1) 2.002(4), Ge(1)–Si(1) 2.3758(12), Ge(1)–Si(4) 2.3984(12), Ge(1)–Si(2) 2.4217(12), Si(1)–C(2) 1.892(4), Si(2)–Si(5) 2.3605(16), Si(2)–Si(3) 2.3818(15), Si(3)–Si(4) 2.3557(16), C(1)–C(2) 1.361(5), C(1)–C(9) 1.476(5), C(2)–C(3) 1.475(5), C(1)–Ge(1)–Si(1) 72.15(11), C(1)–Ge(1)–Si(4) 121.30(10), Si(1)–Ge(1)–Si(4) 119.68(4), C(1)–Ge(1)–Si(2) 129.02(11), Si(1)–Ge(1)–Si(2) 131.33(4), Si(4)–Ge(1)–Si(2) 88.17(4), C(2)–Si(1)–Ge(1) 77.59(11), Si(3)–Si(2)–Ge(1) 86.42(4), Si(4)–Si(3)–Si(2) 90.12(5), C(2)–C(1)–Ge(1) 105.2(3), C(1)–C(2)–Si(1) 104.7(3).

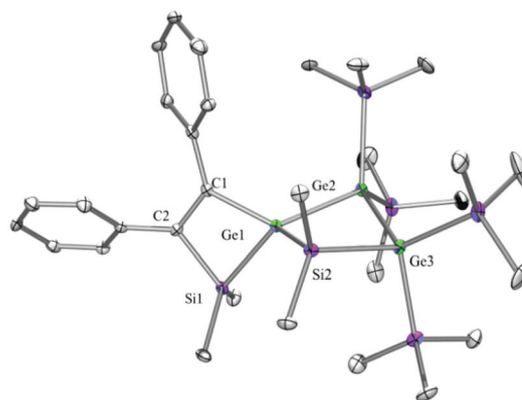


Fig. 7 Molecular structure of **13b** (thermal ellipsoid plot drawn at the 30% probability level). All hydrogen atoms are omitted for clarity (bond lengths in Å and angles in degrees). Ge(1)–C(1) 1.988(7), Ge(1)–Si(1) 2.376(2), Ge(1)–Ge(2) 2.4644(11), Ge(2)–Ge(3) 2.4758(11), Si(1)–C(2) 1.895(7), Si(2)–C(17) 1.873(7), C(1)–C(2) 1.380(10), C(1)–C(9) 1.471(10), C(1)–Ge(1)–Si(1) 72.7(2), C(1)–Ge(1)–Ge(2) 128.1(2), Si(1)–Ge(1)–Ge(2) 130.56(6), Ge(1)–Ge(2)–Ge(3) 85.47(4), C(2)–Si(1)–Ge(1) 77.5(2), C(2)–C(1)–Ge(1) 105.0(5), C(1)–C(2)–Si(1) 104.4(5).



longest Ge–Si distances were found as expected in the four-membered rings with 2.4276(15) Å in **6a** and 2.4217(12) Å in **13a** not differing much from the Ge–Ge distances of 2.4644(11) and 2.4758(11) Å in **13b**.

The Ge–C and the C=C bond lengths are within a small range for all these compounds: Ge–C between 2.002(4) Å in **13a** and 1.958(2) Å in **10a** and the C=C between 1.380(10) Å in **13b** and 1.340(4) Å in **6b**. The two ring planes with the germanium atom at their intersection engage an angle of about 84° in spiro[4.2] **10a** (84.9°) and in spiro[5.3] **6a** (84.1°) and **6b** (84.7°) and about 88° in spiro[3.3] **13a** (88.1°) and **13b** (88.2°) and in spiro[4.4] **7c** (88.1°). The four-membered ring systems containing a double bond (**6a**, **6b**, and **13a**) are largely planar, whereas the second four-membered ring in **13a** is folded by some 30°. The five-membered rings in **7c** and **10a** engage the expected envelope and the six-membered rings in **6a** and **6b** a twisted boat conformation.

Conclusions

The reactions of the two cyclic germylene phosphane adducts **5a** and **5b** with phenylacetylene and trimethylsilylacetylene at ambient temperatures led to the formation of spirocyclic compounds **6a–7c**. In contrast to the acyclic case described earlier,^{33,34} no intermediate formation of the phosphane adducts of cyclic mono- or divinylgermylenes was observed. Instead, germylene insertion into the transannular bonds caused the formation of spirocyclic germanes. The result of DFT calculations indicated that the formation of the different spirocyclic compounds is controlled by the reaction kinetics of the intramolecular insertion of the germylene into the remote Si–Si bonds.

There is no clear energetic differentiation between the insertion into the two different endocyclic Si–Si bonds in germylene **9a** (Fig. 1), leading to the formation of the [4.4]-spirocyclic germsilane **7a** along with the [3.5]-spirocycle **6a**. For germylene **9b**, the insertion reaction into the weaker Ge–Si bond is clearly favored over insertion into the central Si–Si bond, and thus leads to the exclusive formation of the less stable [3.5]-spirocyclic compound **6b**. In contrast, α -trimethylsilyl substitution as in the intermediate germylene **9c** favors insertion into the central Si–Si bond leading to the more stable [4.4]-spirogermsilane **7c**.

Thermal treatment of germacyclopropenes **10a** and **10b**, which were obtained by the reaction of phosphane adducts **5a** and **5b**, produced spirocyclogermanes (**13a** and **13b**) and annulated bicyclic germanes (**17a** or **18a**). Therefore, the thermolysis of germacyclopropenes provides straightforward access to several classes of new bicyclic silagerma-heterocycles. It is, however, noteworthy that these reactions that involve a cascade of ring-opening and ring-closure reactions *via* germylene (Ge(II)) and germane (Ge(IV)) intermediates proceed in solution at temperatures below 150 °C. The results of the DFT calculations confirm that the bicyclic Ge(IV) products are more stable than the suggested cyclic germylene intermediates, but

the calculated energy differences between both families of compounds are relatively small (less than 120 kJ mol⁻¹). These thermal rearrangements provide examples that the introduction of destabilizing factors to Ge(IV) compounds such as ring strain allows reaching an energetic regime that favors reversible reductive elimination to Ge(II) compounds. In consequence, this would lead to model systems which allow a reversible Ge(II)/Ge(IV) interconversion based on vinylgermylenes and spiro-germacyclopropenes and -germacyclobutenes. To this point, we would like to emphasize that Sasamori and co-workers recently communicated a catalytic cyclotrimerization reaction of alkynes based on a vinylgermylene catalyst.³⁹

Experimental section

General remarks

All reactions involving air-sensitive compounds were carried out under an atmosphere of dry nitrogen or argon using either Schlenk techniques or a glove box. All solvents were dried using a column-based solvent purification system.⁴⁴ The chemicals were obtained from different suppliers and used without further purification. Compounds **5a**²⁸ and **5b**²⁸ were prepared following the reported procedures.

The ¹H (300 MHz), ¹³C (75.4 MHz), and ²⁹Si (59.3 MHz) NMR spectra were recorded on a Varian INOVA 300 spectrometer in C₆D₆ and are referenced to tetramethylsilane (TMS) for ¹H, ¹³C and ²⁹Si. To compensate for the low isotopic abundance of ²⁹Si the INEPT pulse sequence^{45,46} was used for the amplification of the signal. Elemental analysis was carried out using a Heraeus Vario Elementar instrument.

X-Ray structure determination

For X-ray structure analyses the crystals were mounted onto the tip of glass fibers, and data collection was performed with a BRUKER-AXS SMART APEX CCD diffractometer using graphite-monochromated Mo K α radiation (0.71073 Å). The data were reduced to F²_o and corrected for absorption effects with SAINT⁴⁷ and SADABS,^{48,49} respectively. The structures were solved by direct methods and refined by the full-matrix least-squares method (SHELXL97).⁵⁰ If not noted otherwise all non-hydrogen atoms were refined with anisotropic displacement parameters and all hydrogen atoms were located in calculated positions to correspond to the standard bond lengths and angles. All diagrams were drawn with 30% probability thermal ellipsoids and all hydrogen atoms were omitted for clarity.

Crystallographic data (excluding structural factors) for the structures of compounds **6a**, **6b**, **7c**, **10a**, **13a**, and **13b** reported in this paper have been deposited with the Cambridge Crystallographic Data Center as supplementary publication nos. CCDC 1450265 (**6a**), 1450272 (**6b**), 1450270 (**7c**), 1450269 (**10a**), 1450273 (**13a**), and 1450271 (**13b**).[†] Figures of the solid-state molecular structures were generated using Ortep-3 as implemented in WINGX⁵¹ and rendered using POV-Ray 3.6.⁵²



5,5,6,6-Tetramethyl-3,9-diphenyl-1,1,7,7-tetrakis(trimethylsilyl)-1,5,6,7-tetrasila-4-germaspiro[3.5]nona-2,8-diene (6a) and 1,1,6,6-tetramethyl-4,9-diphenyl-2,2,7,7-tetrakis(trimethylsilyl)-1,2,6,7-tetrasila-5-germaspiro[4.4]nona-3,8-diene (7a)

A solution (2 mL THF) of **5a** (130 mg, 0.20 mmol) and phenylacetylene (41 mg, 0.40 mmol) was stirred for 18 h at r.t. The solvent was removed under reduced pressure and the products were extracted with pentane (3 × 5 mL). The solvent was removed and an inseparable colorless mixture of **6a** and **7a** (131 mg) in an approximate ratio of 1 : 1 was obtained. NMR: **6a** δ in ppm: -9.6 (SiMe₃), -10.9 (SiMe₃), -11.4 (SiMe₃), -11.8 (SiMe₃), -27.3 (SiMe₂), -35.1 (GeSiMe₂), -46.1 (SiSiMe₂), -46.8 (GeSiC), -86.6 (Me₂SiSiC) and **7a** δ in ppm: -12.1 (SiMe₃), -12.5 (SiMe₃), -27.3 (SiMe₂) and -66.9 (CSiSiMe₂). Although it was not possible to separate **6a** and **7a**, the quantitative inspection of the crystal batch under the microscope revealed good crystals of **6a**, which were subjected to single crystal XRD analysis.

5,5,6,6-Tetramethyl-3,9-diphenyl-1,1,7,7-tetrakis(trimethylsilyl)-5,6-disila-1,4,7-trigermaspiro[3.5]nona-2,8-diene (6b)

A solution (2 mL THF) of **5b** (37 mg, 0.05 mmol) and phenylacetylene (10 mg, 0.10 mmol) was stirred for 2 h at r.t. The solvent was removed under reduced pressure and the product was extracted with pentane (3 × 5 mL). The solvent was removed and crystallization from pentane at -35 °C gave colorless crystalline **6b** (38 mg, 90%). Mp = 101–102 °C. NMR: δ in ppm: ¹H: 7.78 (s, 1H, C-H), 7.58 (d, 2H, o-H), 7.41 (s, 1H, C-H), 7.07–7.00 (m, 4H, Ph), 6.95–6.85 (m, 2H, Ph), 0.54 (s, 3H, SiMe₂), 0.52 (s, 3H, SiMe₂), 0.43 (s, 9H, SiMe₃), 0.41 (s, 3H, SiMe₂), 0.40 (s, 9H, SiMe₃), 0.38 (s, 3H, SiMe₂), 0.32 (s, 9H, SiMe₃), 0.31 (s, 9H, SiMe₃). ¹³C: 169.7, 162.5, 149.4, 146.6, 142.6, 140.1, 126.5, 126.1, 125.9, 2.6 (SiMe₃), 2.2 (SiMe₃), 1.8 (SiMe₃), 1.5 (SiMe₃), -2.0 (SiMe₂), -3.1 (SiMe₂), -3.8 (SiMe₂), -5.1 (SiMe₂), ²⁹Si: -3.3 (SiMe₃), -3.7 (SiMe₃), -4.7 (SiMe₃), -5.0 (SiMe₃), -35.5 (SiMe₂), -37.3 (SiMe₂). Anal. calcd for C₃₂H₆₀Ge₃Si₆ (831.25): C 46.24, H 7.28. Found: C 46.01, H 7.34.

1,1,6,6-Tetramethyl-2,2,4,7,7,9-hexakis(trimethylsilyl)-1,2,6,7-tetrasila-5-germaspiro[4.4]nona-3,8-diene (7c)

A mixture of **5a** (65 mg, 0.10 mmol) and trimethylsilylacetylene (20 mg, 0.20 mmol) was stirred in THF (2 mL) for 18 hours at r.t. The solvent was removed under reduced pressure and the residue was extracted with pentane (3 × 5 mL).

The solvent was evaporated to give **7c** as a white solid (63 mg, 86%). Crystallization from pentane at -35 °C gave colorless crystals suitable for X-ray diffraction analysis. Mp = 113–116 °C. NMR: δ in ppm: ¹H: 7.80 (s, 2H, CH), 0.55 (s, 12H, SiMe₂), 0.28 (s, 18H, SiMe₃), 0.21 (s, 18H, SiMe₃), 0.20 (s, 18H, SiMe₃). ¹³C: = 176.6, 158.2, 1.2 (SiMe₃), 1.1 (SiMe₃), 0.3 (SiMe₃), -0.3 (SiMe₂), -1.4 (SiMe₂). ²⁹Si: -6.9 (SiMe₃), -12.1 (SiMe₃), -12.7 (SiMe₃), -28.2 (SiMe₂), -66.4 (Si(SiMe₃)₃). Anal. calcd for C₂₆H₆₈GeSi₁₀ (734.31): C 42.53, H 9.33 Found: C 42.24, H 9.03.

5,5,6,6-Tetramethyl-1,2-diphenyl-4,4,7,7-tetrakis(trimethylsilyl)-5,6-disila-3,4,7-trigermaspiro[2.4]hept-1-ene (10b)

The same procedure as that for the preparation of **10a** using **5b** (74 mg, 0.10 mmol) and diphenylacetylene (18 mg, 0.10 mmol) was performed. Crystallization from pentane at -35 °C gave yellow crystalline **10b** (52 mg, 64%). Mp = 168–169 °C. NMR: δ in ppm: ¹H 7.82 (m, 4H); 7.19 (m, 4H), 7.00 (m, 2H, p-H), 0.46 (s, 12 H, SiMe₂), 0.28 (s, 36 H, SiMe₃). ¹³C: 147.7, 135.4, 131.6, 128.8, 128.2, 127.1, 3.0 (SiMe₃), -2.2 (SiMe₂). ²⁹Si: -1.5 (SiMe₃), -23.4 (SiMe₂). Anal. calcd for C₃₀H₅₈Ge₃Si₆ (805.21): C 44.98, H 6.59. Found: C 44.98, H 6.58.

1,1,5,5-Tetramethyl-6,7-diphenyl-2,2,3,3-tetrakis(trimethylsilyl)-1,2,3,5-tetrasila-4-germaspiro[3.3]hept-6-ene (13a) and an unknown by-product

A solution of **10a** (50 mg, 0.07 mmol) in toluene (1 mL) was heated up to 100 °C for 18 h. The solvent was removed under reduced pressure and the residue was extracted with pentane (3 × 5 mL). After the removal of the solvent, an inseparable mixture of colorless crystalline **13a** (minor component, identified by single crystal XRD) and an unknown compound was obtained in a ratio of about 1 : 2. Based on the signal intensities and calculated ²⁹Si NMR shifts we assign the following ²⁹Si NMR shifts to **13a**: δ in ppm: ²⁹Si: 9.9 (SiMe₂), 3.8 (SiMe₂), -8.6 (SiMe₃), -8.7 (SiMe₃), -9.5 (SiMe₃), -9.6 (SiMe₃), -94.9 (GeSi(SiMe₃)₂), -108.2 (SiSi(SiMe₃)₂). The remaining signals in the ²⁹Si spectrum: -2.9 (GeSiMe₃), -7.6 (SiMe₃), -9.3 (SiMe₃), -12.5 (SiMe₃), -22.6 (SiMe₂), -35.3 (SiMe₂), -60.1 (CSi(SiMe₃)₂), -119.1 (SiSi(SiMe₃)₂) suggest a compound with the SiMe₂ groups being part of a five-membered ring.

Although we were not able to separate the crystals, it was nevertheless possible to use a single crystal of **13a** for crystal structure analysis.

1,1,5,5-Tetramethyl-6,7-diphenyl-2,2,3,3-tetrakis(trimethylsilyl)-1,5-disila-2,3,4-trigermaspiro[3.3]hept-6-ene (13b)

A solution (C₆D₆ 1 mL) of **10b** (35 mg, 0.04 mmol) in a closed tube was heated up to 150 °C for 18 h. The solvent was removed under reduced pressure and the residue was extracted with pentane (3 × 5 mL). The extract was concentrated to 0.5 mL and stored at -35 °C. Colorless crystals (20 mg, 58%) of **13b** were obtained. Mp = 168–170 °C. NMR: δ in ppm: ¹H: 7.28 (m, 2H, o-H), 7.21–7.09 (m, 4H), 7.04–6.91 (m, 4 H), 0.94 (s, 3H, SiMe₂), 0.91 (s, 3H, SiMe₂), 0.64 (s, 3H, SiMe₂), 0.60 (s, 3H, SiMe₂), 0.44 (s, 18 H, 2 × SiMe₃), 0.34 (s, 9 H, SiMe₃), 0.21 (s, 9H, SiMe₃). ¹³C: 170.6, 156.9, 144.0, 141.0, three signals overlapping with C₆D₆, 126.6, 126.1, 126.0, 5.4 (SiMe₂), 3.7 (SiMe₃), 3.7 (SiMe₃), 3.6 (SiMe₃), 3.5 (SiMe₃), 1.9 (SiMe₂), 1.6 (SiMe₂), 0.2 (SiMe₂). ²⁹Si: 18.4 (SiMe₂), 3.5 (SiMe₂), -2.5 (SiMe₃), -2.9 (SiMe₃), -3.2 (SiMe₃), -4.5 (SiMe₃). Anal. calcd for C₃₀H₅₈Ge₃Si₆ (805.21): C 44.75, H 7.26. Found: C 44.88, H 7.17.

Conflicts of interest

The authors state that there are no conflicts to declare.



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