Recent developments in the chemistry of stable doubly bonded germanium compounds

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ABSTRACT

During the last 10 years, several compounds of the type Ge=X (X = C, Ge, N, P, S) have been isolated as monomers. The stabilization of such derivatives, which are generally highly polymerizable, was achieved by using very bulky groups both on the germanium atom and on the heteroelement X. Conjugation (particularly in a few germenes and germaimines) and intramolecular or intermolecular coordination with oxygen or nitrogen, also contributes, in some cases, to the stabilization. The X-ray analyses of such compounds show a significant bond shortening of the double bond (8-10%) relative to the corresponding single bond and a planar or nearly planar germanium. These doubly bonded germanium derivatives are usually thermally stable but must be handled in an inert atmosphere because of their high sensitivity to oxygen and moisture; they are extremely reactive, much more than the corresponding carbon analogues. Nearly quantitative additions on the double bond have been observed with electrophiles and nucleophiles, and various types of cycloadditions also occur. Except

in one case, a germylene behavior has not been observed, proving that such compounds retain their structural integrity in solution.

A. INTRODUCTION

A large part of the richness of organic chemistry results from the ability of carbon to form double and triple bonds with itself or with other elements. So it seemed very promising to try to obtain similar multiply bonded species with heavier elements of Group 14, such as silicon, germanium and tin, and consequently to develop new heterocompounds. However, all attempts over a long time have been unsuccessful, leading exclusively to oligomers. The first evidence for transient M=X species (M=Si, Ge, Sn; X=C, O, S, N, P, Si, Ge, Sn) was obtained in the 1960s and 1970s with their in situ trapping [la]. However, during the last 10 years, great progress has been made in the organometallic chemistry of Group 14 elements, with new synthetic routes to doubly bonded species and particularly with their stabilization due to substituents having large steric effects, and electronic effects in some cases.

Among organogermanium compounds, many doubly bonded molecules have now been stabilized, such as germenes >Ge=C<, digermenes >Ge=Ge<, germaimines >Ge=N-, germaphosphenes >Ge=P- and germathiones >Ge=S. Some reviews have summarized this field [1-4]. This paper describes the synthesis, the physicochemical studies and the reactivity of such doubly bonded germanium species. Only those reactions involving stable compounds are described. Transient doubly bonded species, which have been reviewed recently [4], are not reported.

B. GERMENES, >Ge=C<

(i) Synthesis

Three routes allow the synthesis of stable compounds with a germanium-carbon double bond (eqn. (1)):

- coupling between a germylene and a carbene (route a)
- dehydrohalogenation of a halogermane by lithio compounds (route b)
- addition of an organolithio compound to a fluorovinylgermane followed by elimination of LiF (route c).

(a) Route a

The reaction, reported by Berndt et al., between the electrophilic cryptocarbene 1 [5] and the stable germylenes 2 [6] and 3 [7] afforded the stable germenes 4 and 5 [8] (eqn. (2)). Compounds 4 and 5 are stable at room temperature in solution. The presence of the Ge=C double bond in 4 has been confirmed by X-ray diffraction (see Section B(ii)(a)). Moreover, 4 has been chemically characterized by addition of HCl across the Ge=C double bond [8] (eqn. (3)).

$$[Me_{3}Si)_{2}C \xrightarrow{tBu} \\ [Me_{3}Si)_{2}C \xrightarrow{t$$

$$4 + HC1 \longrightarrow (Me_3Si)_2C \xrightarrow{B} C -Ge \times N(SiMe_3)_2 \times HC1 \times N(SiMe_3)_2 \times HC1 \times N(SiMe_3)_2 \times HC1 \times$$

(b) Route b

In our laboratory, five stable germenes 6–10 have been obtained by dehydrohalogenation of chloro- or fluorogermanes by tert-butyllithium [9-11] (eqn. (4)). Owing to the high germanium-fluorine bond energy (113 kcal/mol) [12], substitution of germanium by fluorine rules out the Li/halogen exchanges frequently observed when germanium is substituted by other halogens; therefore the best result is generally obtained with fluorogermanes, but the reaction must be done in some cases with chlorogermanes, particularly when the starting fluorogermanes are insoluble in Et₂O [10]. The use of a bulky lithio compound such as 'BuLi is necessary to prevent direct alkylation of germanium. Addition of 'BuLi to the halogermanes was performed at –78°C and under such conditions, the yields of germene are usually about 80%. Like 4 and 5, germenes 6–10 are thermally stable but highly air- and moisture-sensitive derivatives obtained in the form of orange crystals.

 $\label{eq:reconstruction} R'R'': Mes_2 \ 6 \ [9] \ ; \\ (R_2CH)_2 \ 7 \ [10] \ ; \\ (Me_3Si)_2CH]_2 \ 8 \ [11] \ ; \\ Mes, \\ (Me_3Si)_2CH \ 9 \ [11]; \ X:F \ (Me_3Si)_2CH \ 9 \ [11]; \ Mes_1CH \ (Me_3Si)_2CH \ (Me_3Si)_2CH \ 9 \ [11]; \ Mes_1CH \ (Me_3Si)_2CH \ (Me_3Si)_2CH \ (Me_3Si)_2CH \ (Me_3Si)_2CH \ (Me_3Si)_2CH \ (Me_3Si)_2CH \ (Me_3Si)_2CH$

(c) Route c

This route was described for the first time by Jones [13], and then widely used by Auner for organosilicon compounds [14] to obtain transient silenes from vinylchlorosilanes. By using bulky mesityl groups on germanium, the stable dimesitylneopentylgermene could be obtained in 90% yield [15] (eqn. (5)). Compound 11, like the other germenes 6–10, must be handled in an inert atmosphere and appears thermally stable.

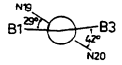


Fig 1. ORTEP view of 4 (reprinted from Angew. Chem. Int. Ed. Engl., 12 (1987) 221).

(ii) Physicochemical studies

(a) X-Ray structure determinations

X-Ray diffraction studies of 4 and 6 show short Ge—C distances (1.827 [8] and 1.801–1.806 Å [16], respectively), about 9–10% shorter than Ge—C single bonds, which are generally about 1.95–2.00 Å, depending on the degree of steric hindrance. Such shortening is typical of all doubly bonded main group elements [3,4,17–19]. The structure of 4 (Fig. 1) confirms the significance of the ylide resonance form B (eqn. (6)) as shown by short distances C(Ge)—B and an average twist angle at the Ge=C double bond of 36° [8].

$$Ge=C \qquad \qquad + Ge-C \qquad B$$

$$Ge=C \qquad \qquad + Ge-C \qquad B$$

$$Ge=C \qquad \qquad + Ge-C \qquad B$$

$$Ge=C \qquad \qquad + Ge-C \qquad Ge$$

$$Ge=C \qquad + Ge-C \qquad + Ge$$

$$Ge=C \qquad + Ge$$

$$Ge=C$$

Fig 2. ORTEP view of 6 (reprinted from Angew. Chem. Int. Ed. Engl., 27 (1988) 828).

In 6, a ylide form D could also be postulated since the negative charge on carbon can be delocalized on the fluorenylidene group (eqn. (6)). However, the X-ray structure of 6 [16] (Fig. 2) displays a planar germanium atom and a small twisting around the Ge-C bond (only 6°) consistent with structure C.

(b) Theoretical studies

The germanium-carbon double bond length (1.801-1.827 Å) is in good agreement with the sum of covalent radii (1.12+0.67=1.79 Å), with ab initio calculations using pseudopotentials (1.779 Å) at the SCF level, 1.812 Å with configuration interactions) [20], and with other calculations (1.767, 1.782 [22c]; 1.761, 1.784, 1.814 [22d]). Calculations have also predicted a rather important polarity of the Ge-C double bond [20] (eqn. (7)), but less important than in silenes, which is in complete agreement with the chemical behavior of Ge=C bonds.

The π -bond energies of Group 14 metal—carbon double bonds have been determined by various methods. The results show that π -bond energies of silenes (35–36 kcal/mol [21]) and germenes (31 kcal/mol [21] or 32.2–33 [22d]) are similar, greater than for stannenes (19 [21] or 20.9, 21 kcal/mol [22d]), and smaller by about 30 kcal/mol than for alkenes (64–68 kcal/mol [21]). Thus these results show that the order of ability to form π -bonds is C > Si ~ Ge > Sn.

Ab initio calculations showed that germanium is more reluctant to form doubly bonded compounds than silicon [22a]. For example, although much more stable than germylmethylene H_3Ge-CH [22b], the corresponding germene was predicted to be 11.4 [22c], 15 [20] and 24.2 kcal/mol [22a] less stable than methylgermylene H_3C-GeH ; however, the energy barrier for the isomerization was calculated to be about 36 [22b] or 33.1 kcal/mol [22c]. The dipole moment of 6 is estimated to be 4 Debyes [16] indicating that the Ge=C double bond is strongly polar. The great importance of the resonance structure $Ge^+=C^-R_2$ explains its high stabilization due to the delocalization of the excess negative charge into the fluorenylidene system.

(c) NMR and Raman spectra

Germenes are very stable in the crystalline state and in non-polar solvents such as pentane or benzene. In polar solvents such as ethers or amines, 6–10 form Lewis acid-base adducts as shown by ¹H NMR [9] (eqn. (8)). This type of complexation has already been observed with silenes >Si=C< [23a], silaimines >Si=N- [23b] and recently in germaimines (see Section D) in which the double bond is strongly polarized and the metal highly electrophilic, but never in symmetric molecules M=M (M = Si, Ge, Sn, P) [17–19]

or in metallaphosphenes >M'=P-(M'=Si, Ge, Sn) (see Section E) in which Group 14 metal is bonded to phosphorus, a less electronegative atom than carbon or nitrogen.

TABLE 1
Germenes

	Germenes	Synthetic route ^a	d(Ge=C) (Å)	¹³ C (ppm)	Ref.
4	$[(Me_3Si)_2N]_2G \rightleftharpoons C \bigvee_{\substack{B \\ \downarrow \\ LBu}} C(SiMe_3)_2$	a	1.827	115	8
5	tBu tBu N B Me ₂ Si Ge = C B C(Si Me ₃) ₂ tBu tBu	a		93	8
6	$Mes_2Ge=CR_2 (CR_2 : C)$	b	1.801 ^b 1.806		9,16
7	$(R_2CH)_2Ge=CR_2$	b			10
8	[(Me3Si)2CH]2Ge=CR2	b			11
9	${ m (Me_3Si)_2CH}_{ m Ge=CR_2}$	b			11
10	R_2 CH $Ge=CR_2$ t Bu	ь		79.8	10
11	$Mes_2Ge = CH - CH_2^tBu$	С		124.2	15

^a(a) Coupling reaction between a germylene and a carbene; (b) dehydrohalogenation of a halogermane; (c) addition of 'BuLi on a vinylfluorogermane, then elimination of LiF.

^bTwo crystallographic independent molecules.

Germenes have also been characterized by ¹³C NMR which shows chemical shifts between 79.8 and 124.2 ppm [8,10,15] (see Table 1) for the sp² carbon doubly bonded to germanium. Unfortunately it is not possible to characterize sp² germanium by NMR as ⁷³Ge NMR is efficient only for symmetric tetracoordinated germanium species. The Raman spectrum of 6 shows a band at 988 cm⁻¹ which is assigned to the Ge=C valence vibration [16], close to the calculated value of 1018 cm⁻¹ using MNDO [16]. Other recent calculations predicted values of 785, 827 and 904 cm⁻¹ [22d].

(iii) Reactivity

(a) Protic reagents. lithio compounds. disulfides. hydrides

Although bulky groups stabilize the Ge=C bond, 6-11 are extremely reactive towards various electrophiles and nucleophiles, and in cycloadditions. The most commonly used reagent to characterize germenes 6-11 is methanol which gives the corresponding methoxyadducts in nearly quantitative yield [9,10,11,15] (eqn. (9)). Other protic reagents react with germenes, for example HCl with 4 [8] and 11 [15], and thiols [9], hydrofluoric acid [9b] and imines, such as diphenylimine [24] with 6 (eqn. 10). Water gives two different products with 6 and 11, depending on the quantity of water. With an excess, germanols 12 are obtained whereas with 0.5 equiv., digermoxanes 13 are formed, due to the reaction of germanol with unreacted germene [15,25] (eqn. 11).

$$Ge=C \qquad \xrightarrow{MeOH} \qquad Ge-C \subset MeO \qquad H$$

$$Ge-C \subset MeO \qquad H \qquad (9)$$

A: Cl, F, EtS, Ph2C=N

Enolyzable ketones such as acetone also react with 6 as protic reagents, yielding the derivative 14 [25] by a classic germanotropic rearrangement [26] (eqn. (12)). These reactions are regiospecific with the more negative group bonded to germanium.

Lithio compounds give two types of reaction depending on steric hindrance. Methyllithium adds to the double bond of 6 to afford the corresponding germane [9b] after quenching with methyl iodide (eqn. (14)). The same type of addition to the Ge=C double bond is observed between tert-butyllithium and germene 11 [15] (eqn. (13)). A completely different reaction occurs between tert-butyllithium and germenes 6 and 8 which have a more pronounced steric hindrance around the double bond than 11. After quenching with methanol or methyl iodide, the germanes 17 and 18, respectively, are obtained. These reactions are best explained by a single-electron transfer mechanism involving the intermediates 15 and 16 (eqn. (14)).

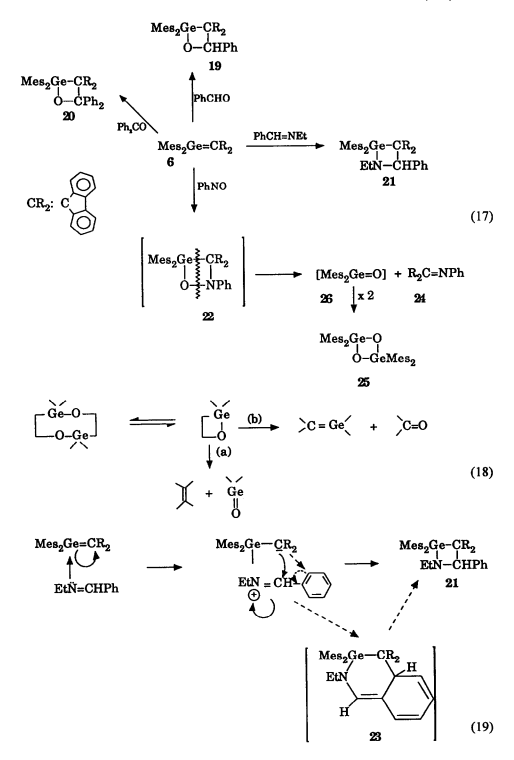
Disulfides such as dimethyldisulfide also add to the germanium-carbon double bond, probably according to a radical mechanism [9] (eqn. (15)). The double bond of 6 and 11 can be easily reduced by lithium aluminium hydride (eqn. (16)).

(b) [2+2] Cycloadditions: aldehydes and ketones, imines, nitroso compounds

Germenes, and particularly germene 6 which has been the most intensively studied, are very reactive in cycloadditions. Nearly quantitative [2+2] cycloadditions are observed with aldehydes [25], ketones [25], nitrosobenzene [24] at ambient temperature, and imines [24] after heating at 140° C in a sealed tube (eqn. (17)). Four-membered ring derivatives 19 and 20 are thermally very stable contrary to other heterocycles of this type which generally dimerize [27] or decompose by a [4] \rightarrow [2+2] process, with formation of germanone and alkene (route a), or by cycloreversion (route b) to give the starting material (eqn. (18)).

The process (b) has been observed by Wiberg in the four-membered ring compounds obtained by addition of benzophenone to the transient germene $Me_2Ge \approx C(SiMe_3)_2$ [28]. Thus, this germaoxetane is a convenient storage device for this germene. Heterocycles 19 and 20 are stable to oxidation and hydrolysis, even towards acetic acid. The chemical inertness of the Ge-O bond in these heterocycles is probably due to the large steric hindrance caused by mesityl groups on germanium, as clearly demonstrated by the X-ray structure.

Reaction of 6 with nitrosobenzene affords only the imine 24 and the 2,4-digerma-1,3-dioxetane 25, a dimer of the transient germanone 26 [24]. These products probably involve the preliminary formation of the four-membered ring heterocycle 22 followed by a classical $[4] \rightarrow [2+2]$ decomposition. Imines are much less reactive than carbonyl compounds towards germene 6 and a prolonged heating at 140° C in a scaled tube is necessary for the reaction [24]. As in the case of benzaldehyde and benzophenone, a [2+4] cycloaddition has never been observed although the preliminary formation of the six-membered heterocycle 23, followed by its isomerization to 21 is not excluded (eqn. (19))



(isomerization of a six-membered ring to a four-membered ring has been postulated by Wiberg in the formation of silaoxetanes from silenes and ketones [29]).

(c) [2+3] Cycloadditions: nitrones and diazo compounds

Facile [2+3] cycloadditions have been observed between 6 and N-(tert-butyl)- α -phenylnitrone and diazomethane leading to the five-membered heterocycles 27 [9a], 28 [30,31] and 29 [30] (eqn. (20)). The reaction with diazomethane is very dependent on experimental conditions. When a large excess of an ethereal solution of diazomethane, dried over potassium pellets, was added to a solution of 6 in Et₂O at -78° C, the only observable product was the germapyrazoline 29 [30]. When the reaction was carried out with a solution of distilled diazomethane (thus free of traces of KOH), the 4-germa-1-pyrazoline 28 was obtained [30,31]. The first steps of these reactions are probably the formation of the dipolar intermediates 30 or 31. Such a prototropy leading to 29 from 28 is well known and has already been observed in other five-membered rings containing the CH₂-N=N linkage [32]. The reverse addition of diazomethane to the Ge=C double bond, with nitrogen bonded to germanium, has never been observed.

Compound 28 was of particular interest as a potential precursor of three-membered ring germiranes 32 (eqn. (21)) which are still unknown, contrary to stable siliranes [33]. Heating 28 at 60°C does not afford the germirane 32, but only 9-methylenefluorene 34,

and oligomers (Mes_2Ge)_n according to a decomposition [3] \rightarrow [2+1] already observed with siliranes. The transient germylene has been trapped by methanol [30] or dimethylbutadiene [31]. Attempts to trap germirane have been unsuccessful probably because of its very short lifetime.

Other diazo compounds such as diazofluorene or diphenyldiazomethane give a completely different reaction with 6, leading to the digermadiazetidines 35 and fluorene R₂CH₂ exclusively [34]. A mechanistic study seems to prove the formation of germaimine 36 as an intermediate, followed by its dimerization (eqn. (22)). In the case of diazofluorene, the dimerization of germaimine 36 affords only the cyclodigermazane 35 with the R'₂CN in a cis position, which then quantitatively isomerizes to the trans isomer (as characterized by X-ray) after a week at room temperature in solution.

(d) [2+4] Cycloadditions: dienes, azobenzene, α -ethylenic aldehydes and ketones

[2+4] Cycloadditions have been observed between germene 6 and 2,3-dimethylbutadiene [9a], azobenzene [24] and various α -ethylenic aldehydes and ketones [35] (eqn. (23)). With 2,3-dimethylbutadiene, only the [2+4] cycloaddition occurs, whereas an ene-

reaction or a [2+2] cycloaddition with the C=C double bond have not been observed [9a].

Afthough the α -ethylenic aldehydes and ketones have different steric and electronic properties, as well as different charges on carbons 2 and 4 [36], exclusive [2+4] cycloaddition leading to the corresponding six-membered ring germaoxacyclohexenes 37-39 [35] have been observed. In the mass spectrum, the most important fragment is always (except for 37) the germylene Mes_2Ge^+ . This can arise from the germene $[Mes_2Ge=CR_2]^+$ as observed in most cases (fragmentation b), but more likely, from a $[6] \rightarrow [5+1]$ decomposi-

tion, since the $R_2C-C=CO$ moieties are also observed (fragmentation a). So it seems that the two types of fragmentation, a and b, occur in germaoxacyclohexenes 38, 39, whereas path c appears as a minor one since $[Mes_2Ge=O]^+$ is observed less frequently (eqn. (24)).

The structure of these adducts is supported by their physicochemical data and also by their chemical reactivity. Addition of water or HF to 38 and 39 gives aldehydes and ketones, respectively, after cleavage of the Ge-O bond (eqn. (25)).

$$\begin{array}{c|c}
Ge-C \\
O \\
C=C
\end{array}$$

$$\begin{array}{c|c}
AH \\
HO \\
C=C
\end{array}$$

$$\begin{array}{c|c}
Ge-C \\
A \\
C \\
R
\end{array}$$

$$\begin{array}{c|c}
A:OH, F \\
C-C \\
R
\end{array}$$

$$\begin{array}{c|c}
R \\
H
\end{array}$$

$$\begin{array}{c|c}
Ge-C \\
C-C \\
R
\end{array}$$

$$\begin{array}{c|c}
R \\
H
\end{array}$$

$$\begin{array}{c|c}
Ge-C \\
C-C \\
R
\end{array}$$

$$\begin{array}{c|c}
A:OH, F \\
C-C \\
R
\end{array}$$

$$\begin{array}{c|c}
C+C \\
R$$

$$\begin{array}{c|c}
C+C \\
R
\end{array}$$

$$\begin{array}{c|c}
C+C \\
R$$

$$\begin{array}{c|c}
C+C \\
R$$

$$\begin{array}{c|c}
C+C \\
R
\end{array}$$

$$\begin{array}{c|c}
C+C \\
R$$

$$\begin{array}{c|c}
C+C \\
R$$

$$\begin{array}{c|c}
C+C \\
R$$

$$\begin{array}{c|c}
C+C \\
R$$

$$\begin{array}{c|c}
C+C \\
R
\end{array}$$

$$\begin{array}{c|c}
C+C \\
R$$

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C+C \\
R$$

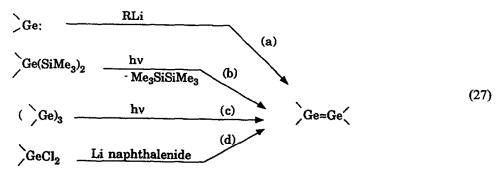
Azobenzene gives only a [2+4] cycloaddition involving one of the phenyl groups; the reaction occurs at room temperature with formation of 40 [24] (eqn. (23)). After a few days in solution, compound 41 was formed, the driving force of the rearrangement $40 \rightarrow 41$ being the re-aromatization. On the contrary, heating 40 leads to the four-membered ring 42. Thermolysis of 42 gives the two expected types of fragmentation [4] \rightarrow [2+2], with formation of starting material (a), of germainine 43 and the corresponding imine (b) (eqn. (26)).

C. DIGERMENES >Ge=Ge<

(i) Synthesis

Stable digermenes are obtained by four routes (eqn. (27)):

- reaction of aryl- or alkyllithio compounds with a stable germylene (route a);
- coupling of germylenes obtained by irradiation of bis-silylgermanes (route b);
- irradiation of cyclotrigermanes (route c);
- reaction of diaryldichlorogermanes with lithium naphthalenide (route d).



(a) Route a

Route (a) was used by Lappert to prepare digermene 45 [37], using bisyllithium, (Me₃Si)₂CHLi, with stable germylene 46 or a Grignard reagent with dichlorogermylene—dioxane complex 44. These reactions afforded 45 in good yield (eqn. (28)). Compound 45 was the first digermene to be prepared, but it retains its digermene structure only in the solid state, and in solution it behaves as two germylene species (see below). Digermene 47 [38] can also be synthesized in low yield (10%) by reaction of diiodogermylene, GeI₂, with 2 equiv. of 2,6-diisopropylphenyllithium, resulting in the formation of the transient germylene, Ar₂Ge, followed by its dimerization (eqn. (29)).

(b) Route b

Irradiation of bis(trimethylsilyl)germanes gives the corresponding germylenes which dimerize to digermenes [3,39-41]. This route allowed the synthesis of 48 [39], which was the first digermene to retain its Ge=Ge structure in solution, and of 49 [42a] (eqn. (30)). This route also affords other transient digermenes which dimerize at room

temperature because they are substituted by less bulky groups such as two 2,6-dimethylphenyl, two mesityl, a mesityl and a tert-butyl, a mesityl and a 2,6-diethylphenyl [3]. In the organosilicon field, the dimerization of a silylene obtained by photolysis of an acyclic trisilane was the first route to tetramesityldisilene [43].

(c) Route c

The photolysis of a cyclotrigermane was the first access to a transient digermene substituted by the 2,6-dimethylphenyl group [44]. However, this digermene could not be isolated because of its slow conversion to the starting cyclotrigermane. Other digermenes have been obtained by this route [3]. The only clean reaction is observed in the case of 48 (eqn. (31)) [45] from the corresponding cyclotrigermane [42b,45]. A similar route involving the irradiation of cyclotrisilanes or of a cyclotristannane was employed for the synthesis of disilenes [18] and a distannene [46].

(d) Route d

The treatment of diaryldichlorogermanes with lithium naphthalenide afforded the corresponding digermenes [38,47] (eqn. (32)). This route is particularly useful for the synthesis of digermenes bearing very large substituents, which cannot be obtained by route (c) since the synthesis of cyclotrigermanes is impossible due to the excessive steric hindrance.

(ii) Physicochemical studies

(a) X-Ray diffraction

The X-ray crystal structure has been determined for digermenes 45, 48 and 50. The Ge=Ge bond length is very different in the three derivatives: 2.213 Å in 48 [45,47], 2.30 Å in 50 [47] and 2.347 Å in 45 [48,51] (sum of covalent radii 2.24 Å) (see Table 2). These long bonds are probably due to an increase in the size of substituents. Lappert's digermene 45 (see Fig. 3), which has the longest Ge—Ge distance, retains its structural integrity only in the crystal [48] and reacts as two germylenes in solution. There is a rather good agreement with calculations which predicted bond lengths of 2.27–2.341 Å depending on the method used [22c,d,49b].

Digermenes 45 and 50 have a markedly *trans* conformation with a fold angle of 32 and 36°, respectively (eqn. (33)). Such fold angles correspond to calculations which predict values from 34 to 40° (depending on the basis set used) [3,48,49]. On the contrary, 48 (see Fig. 4), which contains two smaller groups on germanium than 45 and 50, is more planar, with a fold angle of only 12°. The structure of germene 45 has been described according to eqn. (33) [48].

TABLE 2
Digermenes

						
	Digermenes	Synthetic route ^a	UV nm ($\log \varepsilon$)	Fold angle (°)	d(Ge=Ge) (Å)	Ref.
45	$R_2Ge=GeR_2$ $(R = (Me_3Si)_2CH)$	a	302 (3.2) 414 (3.0)	32	2.35	37 48
47	$Ar_2Ge = GeAr_2$ $Ar = \bigcirc$	a,d	418 (4.6)	-	-	38
48	$Ar'_2Ge = GeAr'_2$ $Ar' = \bigcirc \bigcirc$	b,c	412 (3.9)	12	2.21	39 45 47
49	$Ar''_{2}Ge = GeAr''_{2}$ $Ar'' = $	b	305 (4.0) 418 (4.5)	-	-	42
50	Mes Ge=Ge Ar	c,d	-	36	2.30	47
51	Mes Ge=Ge Ar	c,d	412 (4.6)	-	-	47

^a(a) Reaction of lithio compound with a stable germylene; (b) coupling of germylenes obtained by irradiation of bis-silylgermanes; (c) irradiation of cyclotrigermanes; (d) reaction of dichlorogermanes with lithium naphthalenide.

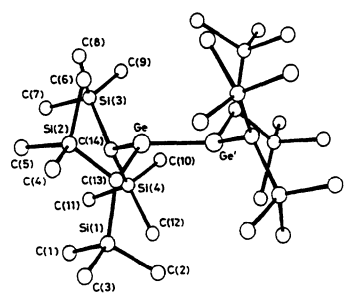


Fig. 3. Molecular structure of 45 (reprinted from J. Chem. Soc. Chem. Commun., (1984) 480).

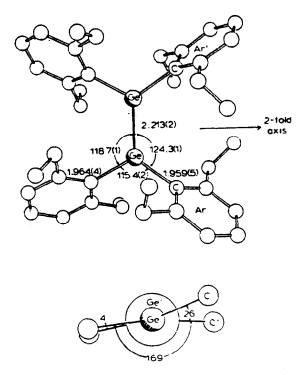
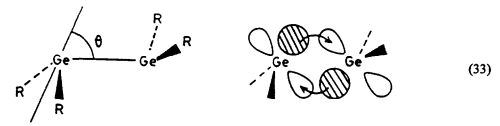


Fig. 4. Molecular structure of 48 (reprinted from Tetrahedron Lett., 25 (1984) 4191).

Note that the fold angles in digermenes are much greater than in disilenes which are generally more planar and, in some cases, completely planar with fold angles and twist angles of 0° [3]. The twist angles in digermenes 45, 47 and 51 are small, 0, 10 and 7°, respectively.



(b) UV spectra

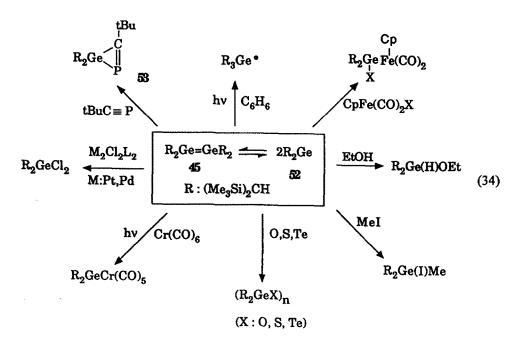
Like other doubly bonded main group species, digermenes are usually yellow. The absorptions of digermenes are blue-shifted compared to those of disilenes but considerably red-shifted from those of ethenes and are not affected by their conformation. The longest wavelength absorptions (between 412 and 418 nm) (see Table 2) have been attributed to a transition between modified π and π^* orbitals [3].

(c) Energy of the Ge=Ge double bond

The synthesis of Z or E stereoisomers 50 and 51 [47] provides a means to determine the activation energy of the isomerization, which is 20.0 ± 0.3 kcal/mol for $E \rightarrow Z$ (50 \rightarrow 51) and 22.2 ± 0.3 kcal/mol for $Z \rightarrow E$ (51 \rightarrow 50) [47]. These values are in good agreement with calculations of the enthalpy of activation for the Ge=Ge bond isomerization, which are estimated at 25 [49e] and 25.4–28 [22d] kcal/mol. In disilenes, the values are about 4–7 kcal/mol higher [3]. The dissociation energy of 48 was estimated, by kinetic investigation, at an upper limit of 30 kcal/mol [3]. Calculations predict this dissociation energy to be between 30 and 45 kcal/mol for digermenes [48,49b,50].

(iii) Reactivity

Like all the doubly bonded main group species, the digermenes are extremely airand moisture-sensitive, but they can be stored for long periods in an argon or a nitrogen atmosphere. Although a large steric hindrance is necessary for their stabilization, they appear very reactive, showing double bond characteristics toward certain reagents. As noted previously, 45 retains its digermene structure only in the solid state. The reactivity of 45 involves the corresponding germylene 52 (eqn. (34)). Most of the reactions lead to tetravalent germanium compounds (e.g. with MeI, EtOH, $M_2Cl_2L_2$ (M = Pt, Pd), O, S, Te) [48]. Transition metal complexed germylenes are also obtained from 45 as well as germyl radicals R_3Ge by irradiation in benzene [51]. Recently, the first germaphosphirene 53 [52] was synthesized by reacting 45 \rightleftharpoons 52 with tert-butylphosphaalkyne (eqn. (34)). A similar behavior occurs for the tin analogue of 45 which appears as a distannene $R_2Sn=SnR_2$ in the solid state by X-ray diffraction and behaves as a stannylene R_2Sn in solution [54]. In sharp contrast, among silenes, the analogue of 45 retains its disilene structure $R_2Si=SiR_2$ ($R=(Me_3Si)_2CH$) in solution [55].



(a) Alcohols

The great difference between digermene 45 and other digermenes appears in their chemical behavior; for example, 48 gives the corresponding methoxydigermane with methanol (eqn. (35)), whereas digermene 45 displays the reaction of germylene shown previously in eqn. (34). This reaction proves that digermene 48, contrary to 45, retains its structural integrity in solution.

$$Ar'_{2}Ge=GeAr'_{2} \xrightarrow{MeOH} Ar'_{2}Ge-GeAr'_{2}$$

$$48 \qquad H OMe$$
(35)

(b) Oxidation

Digermenes 47 and 48 undergo oxygenation through various pathways [53] (eqn. (36)). Exposure of 47 or 48 to oxygen gives 54 quantitatively, which isomerizes to 57 on heating, whereas 54 gives 55 upon photolysis. This last derivative can also be obtained directly and quantitatively from digermenes by reaction with DMSO or N-methylmorpholine-N-oxide. The digermoxirane 56 also results from 47 or 48 by oxidation with N_2O at $60^{\circ}C$ [53] (eqn. (36)).

(c) Reaction with lithium naphthalenide

Addition of an excess of lithium naphthalenide to 47 produces the unexpected cleavage of a Ge-Ar bond to give the corresponding digermenyllithium 58 [38], the first compound of this type in Group 14 metal chemistry (eqn. (37)). A different reaction is observed with less hindered digermene 48 since, in this case, cleavage of the germanium-aryl bond does not occur. Instead, digermane 59 (due to the reduction of the Ge-Ge double bond), Ar'₂GeH₂, and unidentified polymers were obtained [3] (eqn. (38)).

(d) [2+1] Cycloadditions

Digermene 48 reacts with S [56], Se [56] and Te [57] to give the corresponding air-stable thia-, selena- or telluradigermiranes in good yields (eqn. (39)). These heterocycles show

temperature-dependent NMR spectra due to the slow rotation of the aryl groups caused by large steric hindrance [56]. Telluradigermirane exhibits thermochromism, being colorless at -196° C, pale yellow at room temperature, and orange at about 140° C [57]. The structures of 60a and 60b have been determined by X-ray crystallographic analyses [3], and show relatively short germanium-germanium bond distances (2.387 and 2.398 Å; cf. 2.44 Å in a standard Ge-Ge bond) and nearly planar arrangements of the two germanium atoms and of the four ipso carbon atoms of the aryl groups (the sum of angles on germanium is 357.3 and 356.7°). As short Ge-Ge bond distances are not characteristic of three-membered rings (cyclotrigermanes have a longer Ge-Ge bond than expected [44] [59]), these results can be interpreted in terms of Dewar's model of metal olefin bonding [60,61]. Thus, 60a and 60b should have some π -complex character (form A) (eqn. (40)). Similar structures have been reported in other three-membered heterocycles:

Ge-Ge-X
$$(X = CH_2, NPh; see Section C(iii)(f)$$

 $Si-Si-X$ $(X = O,S, CH_2, NSiMe_3)$
Ge-S-P $(see Section E)$

In compound 60c [57], the X-ray structure also shows that the ipso carbons of the aryl groups and the germanium atoms are almost coplanar (sum of angles on germanium 355.4 and 355.5°). But in contrast to 60a and 60b, the normal Ge-Ge bond length (2.435 Å) does not support the π -complex model. Similar [2+1] cycloadditions with sulfur and selenium have also been obtained from the metastable tetramesityldigermene Mes₂Ge=GeMes₂ [57,58].

(e) [2+2] Cycloadditions

Acetone and diphenylacetylene undergo [2+2] cycloadditions with digermene 48 to give the corresponding four-membered ring derivatives 61 and 62 [56] (eqn. (41)). In sharp contrast, acetone and acetophenone react only as protic reagents with germenes >Ge=C< and germaphosphenes >Ge=P-, which have polar bonds (see Sections (B(iii) and E(iii)).

$$Ar'_{2}Ge=GeAr'_{2}$$

$$Ar'_{2}Ge=GeAr'_{2}$$

$$O-CMe_{2}$$

$$Ar'_{2}Ge=GeAr'_{2}$$

$$PhC=CH$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$HC=CPh$$

$$62$$

(f) [2+3] Cycloadditions

Digermene 48 undergoes [2+3] cycloadditions in good yields with diazomethane and phenylazide [42a,56] (eqn. (42)). As for compounds 60a and 60b [3], short Ge—Ge distances and planarity around the germanium atoms are observed for 63 and 64 [42a], hence a π -complex can also be postulated for these cycloadducts.

$$Ar'_{2}Ge=GeAr'_{2}$$

$$Ar'_{2}Ge=GeAr'_{2}$$

$$PhN_{3}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

$$Ar'_{2}Ge-GeAr'_{2}$$

Photolysis of the two three-membered ring compounds 63 and 64 shows them to be good precursors of germylene Ar'₂Ge (which have been trapped by DMB and methanol) and of the corresponding germene and germaimine [42a] (eqn. (43)). However, only the germene could be confirmed by trapping with methanol. The germaimine probably polymerizes rapidly [42a] (eqn. (43)).

Interestingly [2+4] cycloadditions from stable digermenes (for example with dienes or α -ethylenic aldehydes and ketones) have never been observed. Diels-Alder reactions

with dienes have only been obtained from transient digermenes such as Et₂Ge=GeEt₂ [62]. Other types of cycloadditions have also been observed with transient digermenes (see [4] and refs. therein).

Until now, no stable germasilenes $R_2Ge=SiR_2$ or stannagermenes $R_2Sn=GeR_2$ have been synthesized. However, very recently, the transient germasilene $Mes_2Ge=SiMes_2$ has been obtained by Baines et al. by photolysis of hexamesityldigermasilirane and characterized by a trapping reaction with methanol and ²⁹Si NMR at -70°C [122]. At room temperature, this germasilene isomerizes to the corresponding silylgermylene Mes_3Si —GeMes [122], in accordance with calculations which predicted that silylgermylene is more stable than germasilene by 3–6 kcal/mol [50,122].

D. GERMAIMINES >Ge=N-

(i) Synthesis

Stable germaimines are obtained by five routes (eqn. (44)):

- coupling of germylenes with diazomalonic esters (route a);
- coupling of germylenes with azides (route b);
- photolysis of a germylazide (route c);
- reaction between germane and nitroso or nitroxy compounds (route d);
- dehydrochlorination (or fluorination) of a chloro- (or fluoro)germylamine by tert-butyllithium (route e).

$$Ge: + N_{2}C(COOR)_{2}$$

$$Ge: + RN_{3}$$

$$- N_{2}$$

$$Ge=N-$$

$$- \frac{c}{hv/- N_{2}} - GeN_{3}$$

$$- HX e$$

$$- \frac{d}{hv/- N_{2}} - GeN_{3}$$

$$- HX e$$

$$- Ge-N-$$

$$X H$$

$$GeH_{4} + RN=0$$

$$(44)$$

(a) Route a

The first stable germaimine 65 was obtained by Glidewell in the reaction between the electron-rich germylene 66 and the diazo-compound 67 [63,64] (eqn. (45)). Compound 65 exhibits high stability at room temperature in benzene solution, in the absence of air, since addition of EtOH 3 h or 23 h after reaction has minor effect on the yield of adduct. In contrast, when other diazo compounds containing an enolizable function such as $RCH_2COC(N_2)R'$ (R = H; R' = COOEt, $SO_2C_6H_4$ -p-Me, COPh; $R,R' = CMe_2CH_2C(O)$ —) are mixed with germylene 66, the resulting germaimines are not stable [64].

$$[(Me_{3}Si)_{2}N]_{2}Ge: + N_{2}C(COOMe)_{2} \longrightarrow [(Me_{3}Si)_{2}N]_{2}Ge=N-N=C(COOMe)_{2}$$

$$66 \qquad 67 \qquad 65$$

$$(45)$$

(b) Route b

Reaction of stable germylenes with azides substituted by bulky groups is the best route to stable compounds containing a germanium-nitrogen double bond. Seven stable germaimines 68, 69 [65] 70-72 [66], 73 [67] and 74 [68] have been obtained by this route in good yields (eqn. (46)) (see Table 3). Although stable germylenes were generally used [65-67], this reaction also occurs with transient dimesitylgermylene generated in situ [68].

$$[Mes_{2}Ge] \longrightarrow MesN_{3}$$

$$Me \xrightarrow{I} V$$

$$Si \xrightarrow{N} V$$

$$Me \xrightarrow{I} V$$

$$Si \xrightarrow{N} V$$

$$Me \xrightarrow{I} V$$

$$Et_{3}SiN_{3} \qquad Ar:Mes, Ar':Mes$$

$$Ar':Mes$$

$$Ar:Mes, Ar':Mes$$

In these reactions, the first step is probably the formation of the intermediate 75 [68] followed by elimination of nitrogen. However such an intermediate has never been observed (eqn. (47)). Germaimines 65, 68-73 (see Table 3) are stable in solution at room temperature, but 74 must be stabilized by complexation with trimethylamine. When 74 was generated without amine in a benzene or cyclohexane solution, it oligomerized rapidly and could not be chemically characterized.

$$Ge + RN-N \equiv NI \longrightarrow Ge-N-N \equiv NI \longrightarrow Ge=NR+N_{2}$$

$$R \xrightarrow{75}$$

$$Ge=NR+N_{2}$$

$$(47)$$

(c) Route c

Compound 74 has been obtained through a Curtius type rearrangement by irradiation of Mes₃GeN₃ [68,69] (eqn. (48)).

TABLE 3
Germaimines

	Germaimines	Synthetic route ^a	d(Ge=N) (Å)	UV λ_{\max} (nm)	Ref.
65	[(Me3Si)2N]2Ge=N-N=C(COOMe)2	a	_	502 (C ₆ H ₆) [72] 504 (Et ₂ O) [72]	63,64
68	[(Me3Si)2N]2Ge=N-SiEt3	b	_	55. (25/25) [,2]	65
69	[(Me3Si)2N]2Ge=N-Si(OtBu)3	b	_		65
70	(Mes N), Ge=NMes	b	1.691(3)		66
71	(Mes N), Ge=N-Ar Ar:	b	-		66
72	Ar (Me,Si N),Ge=NAr	b	1.703(2)	δ ¹⁵ N (ratio 2:1): -295, -193	66
73	Bu tBu Si N Ge=NSiMe, Me Bu BuN N Ge=NSiMe, Me Bu	b	1.688(9)		67
74a	Mes,Ge=NMes ♠ NMe,	b c		Free 74: 309,459 [69]	68 68,69

TABLE 3 (continued)

Germaimines	Synthetic route ^a	<i>d(</i> Ge=N) (Å)	UV λ _{max} (nm)	Ref.
F ₃ C-N=GeH ₂	d	-	IR v(Ge=N) 1030 cm ⁻¹	70
$F_3C-N=Ge(H)ON(CF_3)_2$	d	_	1028 cm ⁻¹	70
	d	_	1070 cm^{-1}	70
$F_3C-NO-N=GeH_2$	đ	-		71
Mes,Ge=N O C S	e	-	325 nm 261 nm	77
Mes,Ge=N	е			77b
	F ₃ C-N=GeH ₂ F ₃ C-N=Ge(H)ON(CF ₃) ₂ F ₃ C-N=Ge[ON(CF ₃) ₂] ₂ F ₃ C-NO-N=GeH ₂ Mes ₃ Ge=N O MeO	F ₃ C-N=GeH ₂ F ₃ C-N=Ge(H)ON(CF ₃) ₂ F ₃ C-N=Ge[ON(CF ₃) ₂] ₂ F ₃ C-NO-N=GeH ₂ Mes ₃ Ge=N Mes ₄ Ge=N O C C C C C C C C C C C C	F ₃ C-N=GeH ₂ Gain and a counter of the following states of the following st	F ₃ C-N=GeH ₂ Groute ^a F ₃ C-N=GeH ₂ F ₃ C-N=Ge(H)ON(CF ₃) ₂ F ₃ C-N=Ge[ON(CF ₃) ₂] ₂ F ₃ C-NO-N=GeH ₂ Mes ₃ Ge=N O Mes ₄ Ge=N O Mes ₄ Ge=N O C Groupe Groute ^a A A IR r(Ge=N) 1030 cm ⁻¹ 1070 cm ⁻¹ 1070 cm ⁻¹ C 325 nm 261 nm

^aSynthetic routes: (a) coupling between a germylene and a diazomalonic ester; (b) coupling between a germylene and an azide; (c) photolysis of a germylazide; (d) reaction between a germane and a nitroso compound; (e) dehydrochlorination (or fluorination) of a chloro- (or fluoro-) germylamine by tert-butyllithium.

(d) Route d

A novel synthesis of CF₃-bearing germaimines 78–80 [70] and 81 [71] (see Table 3) has been reported in two preliminary publications which appeared 4 years ago (eqn. (49)). Although supporting analytical and spectroscopic data appear to be strong, no comment was made about the astounding resistance of 78 to hydrolysis, unlike all other germaimines. Yet hydrogen iodide added readily. Moreover, the surprisingly high proton chemical shift of the GeH₂ group was unexplained even though the value is higher than and inconsistent with those of all other organogermanium compounds. These results are

also astonishing since doubly bonded germanium species are usually stabilized by sterically demanding groups. It remains to be seen whether the reported synthesis can be extended to a wider range of germanes and organofluorine compounds. Interestingly, the CF₃ group appears to stabilize compounds having a phosphorus—carbon [73] or a phosphorus—nitrogen [84] double bond.

(e) Route e

A new germaimine 82 [77] that is stable in solution at room temperature, has been very recently prepared by dehydrochlorination or dehydrofluorination of the corresponding chloro- (or fluoro)germylamine in excellent yield (nearly 90%) (eqn. (50)). The minor product 82a (~10%) was obtained when the reaction was performed at room temperature. A similar route leads to stable germylamine 83 [77b] (eqn. (50)). This is the first time that dehydrohalogenation has been used in germaimine synthesis. But, whereas this method is very efficient for germenes >Ge=C< and germaphosphenes >Ge=P-, the dehydrohalogenation of halogermylamines afforded only cyclogermazanes [78a]. When the steric hindrance was important, it appeared very difficult [78b], as also observed in silicon chemistry. In the case of 82, elimination of halogen is probably easier because of

the nucleophilic assistance of oxygen [77]. Germaimine 82 is probably stabilized by conjugation with the thiophoate group, and also by an intramolecular coordination with oxygen of the ester groups as evident from the decrease in the $\nu(C=0)$ band in the IR spectrum and the bathochrome shift in the UV spectrum [77]. On this basis, resonance structures 82A \leftrightarrow 82B can be postulated. Moreover, mesityl groups on germanium appear to be necessary for stabilization since, with ethyl groups, the germaimine was not stable and instead only the corresponding cyclodigermazane was obtained [80].

(ii) Physicochemical studies

(a) X-Ray

Three germaimines have been structurally characterized by X-ray diffraction. The Ge=N double bond lengths: 1.691(3) Å in 70 [66], 1.703(2) Å in 72 [66] (see Fig. 5) and 1.688(9) Å in 73 [67] (see Fig. 6), are in excellent agreement with the calculated value for $H_2Ge=NH$ (1.695 Å at the SCF level and 1.727 Å at the CI level) [20]. The bond shortening relative to the standard single bond is about 7.8%. In 70 and 72, the C-N=Ge-N torsion angles (-1.5(5), 178.4(3)° and -1.5(5), 178.1(3)°, respectively) and the sum of angles at germanium (exactly 360°) are consistent with a normal $p\pi$ - $p\pi$ double bond.

In compound 73, the sum of angles at germanium is 357.7° [67], also very close to 360°. From X-ray and NMR data, it seems that it should be described by three resonance structures [67] (eqn. (51)). Apparently, 73 is stabilized by an intramolecular base-stabilization, a new concept which appears to be very efficient and does not affect the reactiv-

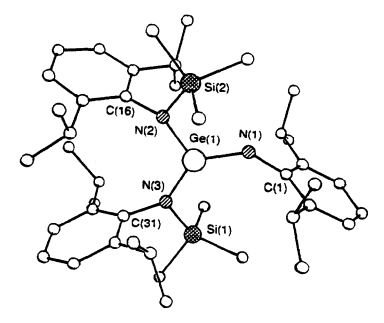


Fig. 5, Molecular structure of 72 (reprinted from J. Chem. Soc. Chem. Commun., (1991) 1123).

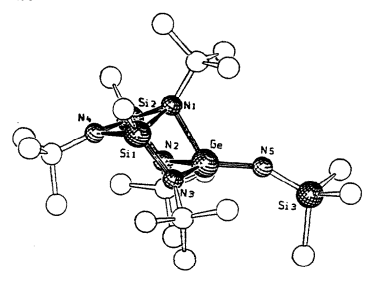


Fig. 6. Molecular structure of 73 (reprinted from Angew. Chem. Int. Ed. Engl., 29 (1990) 216).

ity of the germaimine. A similar stabilization by intramolecular coordination with a nitrogen occurs in silaimines >Si=N- [74] and silaphosphenes, >Si=P- [74], but in these cases, the Si=N(P) double bond is unreactive.

(b) Theoretical studies

Germaimine $H_2Ge=NH$ was predicted to be planar [20]. The calculated geometry and the Ge=N bond distances agree well with the experimental values [66,67]. The charge distribution shows a very polar bond (Ge, +0.54; N, -0.70 [20]) and a large dipole moment (2.99 D).

Compound 65 slowly changes over a period of a few days into a second species. The similarities between the NMR spectra of the two compounds show that they are isomeric. The initial form is the transoid 65a, and the more stable the cisoid 65b [63,72] (eqn. (52)).

$$(Me_{3}Si)_{2}N \qquad (Me_{3}Si)_{2}N \qquad Ge=N \qquad (Me_{3}Si)_{2}N \qquad N$$

$$(Me_{3}Si)_{2}N \qquad N \qquad (S2)$$

$$(Me_{3}Si)_{2}N \qquad N \qquad (S2)$$

$$(Me_{3}Si)_{2}N \qquad N \qquad (S2)$$

These results have been confirmed by MNDO calculations on the simpler analogue [(H₃Si)₂N]₂GeNN(COOMe)₂ [72]. Structural data suggest that 65 should be represented as a hybrid of three forms, with C as the most important contributor [72] (eqn. (53)).

$$Ge=N-N=C \qquad \bigoplus Ge-N=N-C \qquad \bigoplus Ge-N-N=C \qquad (53)$$

$$A \qquad B \qquad C$$

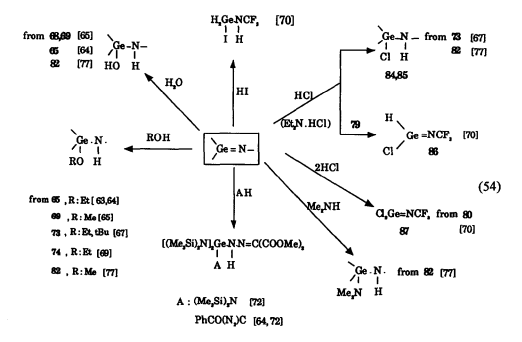
(c) IR, UV, NMR

The Ge=N stretching vibration has been observed experimentally at 1028 (79), 1300 (78) and 1070 cm⁻¹ (80) [70]. It was one of the most important physicochemical data responsible for the characterization of germaimines 78–80 [70]. These values are slightly higher than previously reported for $F_2Ge=NPh$ (970 cm⁻¹ [la]) and much higher than calculated (854 cm⁻¹ [20]). In the UV spectra, bands were observed at 502 nm (in Et₂O) and 504 nm (in C₆H₆) for 65 [72]. In the case of 74, two bands were observed at 459 nm (n $\rightarrow \pi^*$) and 309 nm ($\pi \rightarrow \pi^*$) [69]. These measurements were done in an argon matrix since 74 is only stable at room temperature when it is complexed by an amine. In the ¹⁵N NMR spectrum of 72, two signals were observed in the ratio 2:1 at -295 and -193 ppm, respectively, for the three nitrogen atoms [66].

(iii) Reactivity

(a) Protic reagents

Whereas the reactivity of transient germaimines is now well known [4 and refs. therein], the chemical behavior of stable germaimines is still relatively undeveloped. Most characterization reactions have been done with protic reagents (alcohols, water, mineral acids) or compounds with active hydrogen ((Me₃Si)₂NH, PhCOCH(N₂)) (eqn. (54)). The reactions displayed in eqn. (54) deserve some comment. They occur regiospecifically, with hydrogen always on nitrogen, and usually in excellent yield. In the case of hydrochloric acid, addition to the Ge=N double bond is normally observed in 73 and 82, giving 84 [67] and 85 [77], respectively. In contrast, a rather surprising reaction occurs between HCl and 79 or 80; cleavage of the Ge-O bond is observed, leading only to the corresponding chlorogermaimines 86 and 87 in 53 and 57% yield, respectively, together with an almost quantitative yield of bis(trifluoromethyl)hydroxylamine. These new germaimines are stable at -20°C. In some cases the labile Ge-N bond is cleaved by an excess of protic reagent, leading to dimesityldiethoxygermane (e.g. 74 with EtOH [69]) or germoxane ([(Me₃Si)₂N]₂GeO)_n (e.g. 65 with H₂O [64,72]). Similar results are obtained when EtOH is added to 65 3 h or 23 h after its synthesis; the yield decreases only from 83 to 75%, indicating a considerable lifetime at room temperature for this germaimine.



(b) Lithio compounds and methyliodide

Methyllithium and methyliodide add to the double bond of 73 with the electropositive part of the reactant bound to nitrogen and the electronegative part on the germanium atom as expected. Quenching 88 with ethanol, and addition of trimethylstannane to 89 yield the two regioisomers 90 and 91, respectively. Such reactions are interesting since they give two compounds which differ only in the position of the hydrogen atom and methyl group on germanium and nitrogen [75] (eqn. (55)).

$$Ge: tBuN N Ge$$

$$Si N Ge$$

$$Me tBu Si N Ge$$

$$Si N Ge$$

$$Me tBu N N Ge$$

$$Si N Me tBu Me tBu$$

$$Me tBu Me tBu$$

$$Me tBu Me tBu$$

Addition of methyliodide to the Ge=N double bond of 73 shows the high reactivity of such a derivative, although the germanium is complexed with nitrogen. Note that such addition of alkyl iodide to a stable doubly bonded germanium species has never been observed before.

(c) Cycloadditions

The free transient germaimine Mes₂Ge=NMes 74 can be characterized by trapping with benzaldehyde during the course of the irradiation of Mes₃GeN₃. But if benzaldehyde is added after the end of the irradiation, no pseudo-Wittig reaction occurs, showing that free 74 is not stable at room temperature [68]. When the irradiation was done in the presence of trimethylamine, the complexed germaimine 74a showed good stability, since addition of benzaldehyde or diphenylnitrone gave the expected pseudo-Wittig reaction even after the end of the irradiation [68] (eqn. (56)).

$$\begin{array}{c|c}
 & PhCHO \\
\hline
 & Mes_2Ge-NMes \\
\hline
 & O-CHPh \\
\hline
 & 92
\end{array}$$

$$\begin{array}{c|c}
 & Mes_2GeO)_2 + PhCH=NMes \\
\hline
 & PhCH=NPh \\
\hline
 & Mes_2GeO)_2 + PhCH=NMes \\
\hline
 & O-NPh \\
\hline
 & 93
\end{array}$$

$$\begin{array}{c|c}
 & Mes_2GeO)_2 + PhCH=NMes \\
\hline
 & O-NPh \\
\hline
 & PhN=NPh
\end{array}$$

$$\begin{array}{c|c}
 & Mes_2GeO)_2 + PhCH=NMes \\
\hline
 & O-NPh \\
\hline
 & PhN=NPh
\end{array}$$

$$\begin{array}{c|c}
 & Mes_2GeO)_2 + PhCH=NMes \\
\hline
 & O-NPh
\end{array}$$

These reactions involve the intermediate formation of four- and five-membered ring derivatives 92 and 93 which are not stable and give the classical decomposition of cycloadducts of germaimines [76]. A [2+4] cycloaddition has been observed between germaimine 82 and 3,5-di-tert-butyl-ortho-quinone, leading to cycloadduct 94 [77], then to the corresponding germadioxolane 95 according to a type of decomposition already observed in derivatives having similar structure [81] (eqn. (57)).

E. GERMAPHOSPHENES >Ge=P-

(i) Synthesis

Stable germaphosphenes are obtained in two ways (eqn. (58)):

dehydrohalogenation of halogermylphosphines by tert-butyllithium (route a); synthesis in one step by reacting dihalogermanes with dilithiophosphides (route b).

(a) Route a

The dehydrofluorination of the corresponding fluoro precursor by tert-butyllithium is the best route to germaphosphene 96 [82], 97, 98 [83] and 99 [83], which are all obtained nearly quantitatively (eqn. (59)). In dehydrohalogenation reactions, the best results are obtained with fluoro-germanium derivatives. Owing to the high germanium-fluorine bond energy (113 kcal/mol) [12], the Li/halogen exchange frequently observed when germanium is substituted by other halogens, does not occur. Moreover, bulky lithio compounds such as 'BuLi prevent direct alkylation of germanium. Note that silaphosphenes >Si=P- are usually obtained from chloro precursors [85-87], and rarely from fluoro precursors [88]. Germaphosphene 96 can also be prepared by dehydrochlorination of the corresponding chlorogermylphosphine by trimethylmethylenephosphorane [82] or DBU [82], but in low yields. These are only marginal routes to germaphosphenes (eqn. (60)).

(b) Route b

The germaphosphene 100 [89] can be synthesized in a one-step reaction between dimesityldifluorogermane and 2,4,6-tri-isopropylphenyldilithiophosphide, which is ob-

tained from the corresponding phosphine and 2 equiv. of butyllithium. This type of process, generally less clean than the two-step reaction, gave good results for 100 (eqn. (61)).

(ii) Physicochemical studies

(a) X-Ray diffraction

The presence of a double bond between germanium and phosphorus was proved unambiguously by an X-ray crystal structure determination. In the cases of 96 (2.138 Å) [91] (Fig. 7) and 98 (2.14 Å) [83] (Fig. 8), a shortening of about 8–9% compared to the Ge-P single bond (2.33–2.36 Å for a single bond [92]) was observed. This is typical for organometallic doubly bonded compounds. Moreover, 96 and 98 adopt a nearly planar conformation of the central atoms $C_2Ge=PC$ with the sum of angles on germanium practically 360°. The Ge=P bond length is in good agreement with calculated values by Barthelat [93] of 2.169 Å at the SCF level with configuration interactions, and 2.136 Å by ab initio calculations using pseudopotentials. These calculations also agree well with the sum of covalent radii (2.12 Å) of sp² hybridized germanium (1.12 Å) and sp² phosphorus (1.00 Å).

(b) Isomerization barrier

As mentioned above, dehydrofluorination of fluorogermylphosphine 101 afforded germaphosphenes 97 and 98 [83]. However, only the isomer with a *cis* arrangement of Mes and Ar groups (97) is observed immediately after the reaction, even when starting

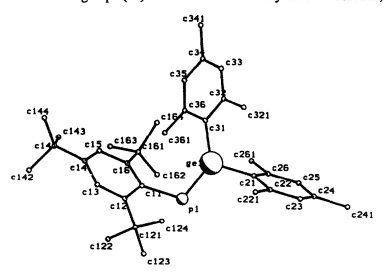


Fig. 7. Molecular view of 96 (reprinted from Organometallics, 7 (1988) 1010)

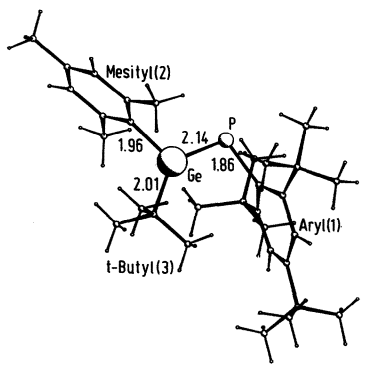


Fig. 8. Molecular view of 98 (reprinted from New J. Chem., 13 (1989) 389).

with a 60:40 mixture of the two diastereoisomers 101a/101b (eqn. (62)). This phenomenon can be explained by the preferential reaction of one of these two diastereoisomers, followed by a rapid thermodynamic equilibrium between them, probably due mainly to the high configurational instability of asymmetric fluorogermanes [90]. After 5 h at room temperature, complete isomerization of 97 to 98 was observed [83] (eqn. (62)). Both isomers were identified by NMR spectroscopy, and 98 also by X-ray diffraction (see Section (E(ii)(a)).

The *cis-trans* isomerization barrier 97 = 98 (22.3 kcal/mol) has been determined by NMR spectroscopy. A similar value (20.3 kcal/mol) was obtained for the diphosphene ArP=PAr (Ar: 2,4,6-tri-tert-butylphenyl) [94]. The *cis* arrangement of Ar and 'Bu may seem surprising since tert-butyl is considered to be bulkier than mesityl [95]. However, free rotation of all groups is not possible when mesityl and Ar are *cis*.

(c) NMR and Raman spectra

Germaphosphenes display characteristic low field ³¹P NMR signals (between +145 and +173.6 ppm; see Table 4) [82,83,89]. A rather large temperature dependence has been observed for the δ^{31} P [83] with increasing temperature effecting a low field shift, up to 12 ppm, between -80°C and +80°C (see Table 4). This corresponds to a red shift in the electronic absorption spectrum. As a result, germaphosphenes display high thermochromism, yellow at -80°C, orange at room temperature, and orange-red at +80°C. Such a reversible phenomenon has previously been noted in other doubly bonded species like disilenes [17] and diphosphenes [19], and similar relationship between electronic absorption and ¹¹⁹Sn chemical shifts is observed in the series Ph₃Sn(Sn^tBu₂)_nSnPh₃ (n = 1-4) [96]. Germaphosphenes 96 and 98 have strong Raman emission bands at 503 and 501.5 cm⁻¹, respectively (see Table 4) [83]. A good correlation is observed with previous calculations: 481 cm⁻¹ [93] using pseudopotentials and configuration interactions.

(iii) Reactivity

(a) Protic reagents

Orange solutions of germaphosphenes 96-100 are extremely air- and moisture-sen-

TABLE 4
Germaphosphenes

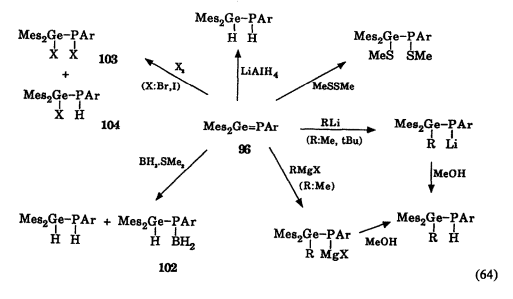
	Germaphosphenes	Synthetic route ^a	Raman (cm ⁻¹)	δ ³¹ P (ppm)	d(Ge=P) (Å)	Ref.
96	Mes ₂ Ge=PAr	a	503	+173.6 (+164.8: -80°C +178.2: +80°C)	2.139(3) [91]	82 91
97	Mes Ar	a	-	+169.2		83
98	tBu Ge=P Ar	С	501.5	+157.4 (+149.1: -80°C +161.0: +80°C)	2.14(4) ^b 2.14(3)	83
99	^t Bu ₂ Ge=PAr	a	_	+156.6 (+148.5: -80°C +159.8: +80°C)		83
100	Mes ₂ Ge=PIs	b	-	+146.8		89

^a(a) Dehydrofluorination by ^tBuLi of a fluorogermylphosphine; (b) one step process: reaction of difluorodimesitylgermane with a dilithiophosphide; (c) isomerization from another germaphosphene.

^bTwo crystallographic independent molecules.

sitive, but they can be stored without change for months in an inert atmosphere. They behave only as doubly bonded compounds, never as germylene R_2 Ge or phosphinidene RP should the germanium-phosphorus connection be very weak. Germaphosphene 96 reacts quantitatively with protic reagents or compounds with active hydrogen, such as phenylacetylene or trimethylmethylenephosphorane (eqn. (63)) [82,97-99]. These reactions are regiospecific, giving only secondary phosphanes. The regiochemistry is determined primarily by polar effects since the germanium-phosphorus bond is slightly polarized $Ge^{\delta+}=P^{\delta-}$. Addition of water or hydrofluoric acid to 98 gives a mixture of two diastereoisomers $Mes({}^tBu)(A)GeP(H)Ar$ (A=F or OH) due to a rapid thermodynamic equilibrium [83].

(b) Lithio compounds, Grignard reagents, hydrides, disulfides, halogens and carbon tetrachloride



Germaphosphene 96 is highly reactive towards nucleophiles. Methyllithium, tert-butyllithium and methylmagnesium iodide react rapidly at room temperature to give, after methanolysis, the corresponding secondary phosphanes [97,99] (eqn. (64)), and the reactions are all regiospecific. Lithium aluminium hydride reduces the germanium-phosphorus double bond [97,99], as do boranes such as BH₃·SMe₂. In this case, the adduct 102 was also observed in minor quantities [97] (eqn. (64)) depending on experimental conditions. Germaphosphene 96 reacts with dimethyldisulfide, probably by a

radical process [97,99]. Each reaction involves only the germanium-phosphorus double bond. The Ge-P single bond, which is usually cleaved by lithio compounds [100], Grignard reagents [101] and hydrides [102], is unaffected due to large steric hindrance by mesityl and tri-tert-butylphenyl groups.

Halogens react with 96 to produce the expected adducts 103 and, surprisingly, halogermylphosphines 104 as major products [97,99] (eqn. (64)). Two competitive mechanisms probably occur, an electrophilic addition leading to 103 and a radical process forming the radical Mes₂Ge(X)—P Ar which reacts further with halogen, or by hydrogen abstraction to produce either 103 or 104.

The reaction of 96 with carbon tetrachloride is particularly interesting because it allows the quantitative synthesis of a compound with a phosphorus—carbon double bond 105 from a compound having a germanium—phosphorus double bond [97] (eqn. (65)).

(c) [2+1] Cycloadditions

Reaction of an equivalent sulfur or selenium with germaphosphene 96 leads to stable three-membered heterocycles, the germathiaphosphirane 106 and the germaselenaphosphirane 107 [103] (eqn. (66)). The first step of the reaction is probably the formation of the germaphosphene sulfide (or selenide) 108 followed by its isomerization to the three-membered ring. A similar process has been demonstrated by Yoshifuji in the reaction of sulfur with a diphosphene [104] (eqn. (67)). The X-ray crystal structure of 106 is rather surprising since germanium, phosphorus and the three ipso carbons of the aromatic rings are nearly in a plane as in the starting germaphosphene. Moreover, the germanium atom is nearly planar (sum of angles 358.1°) with respect to the two mesityl groups and the P atom, and the germanium—phosphorus bond is shorter than a classical Ge—P single bond (2.316 Å versus 2.35–2.36 Å) [92]. Apparently the hybridization of germanium is between sp² and sp³ and the bonding intermediate between a normal three-membered ring and a π -complex (eqn. (68)).

Similar observations have been reported by West for disilaoxiranes

in which the Si-Si bonds are shorter than expected with quasi planar silicon atoms. This hypothesis is supported by a theoretical investigation of three-membered rings containing two silicon atoms [107]. The same phenomenon has been described for other three-membered ring derivatives (see Section C2).

$$ArP = PAr \xrightarrow{1/8 S_0} ArP = PAr \xrightarrow{\Delta \text{ or hv}} ArP = PAr$$

$$S$$

$$S$$

$$Ge = P - Ge = P - Ge$$

$$Ge = P - Ge$$

(d) [2+2] Cycloadditions

Germaphosphenes 96 and 100 react readily with benzaldehyde by [2+2] cycloaddition to give the corresponding four-membered ring heterocycles 109 [89] (eqn. (69)).

The first step of this reaction is probably the nucleophilic attack of oxygen on germanium, followed by cyclization. Although the Ge-P bond is only slightly polarized, a regioselective reaction takes place with oxygen bonded exclusively to germanium. Sixmembered ring heterocycles such as 110 are never obtained, contrary to [2+4] cycloadditions reported between benzophenone and some transient germenes Ge=C [28] or silenes Si=C [108]. The ³¹P NMR signal of 109 appears at low field (between +45 and +82 ppm depending on group R on phosphorus). Such a large deshielding has been explained by folding of the four-membered ring allowing an interaction between phosphorus and oxygen lone pairs [89]. In the mass spectrum of 109, the classical [2+2] decomposition of organometallic four-membered ring heterocycles has been observed, the most important route a (a/b 95:5) regenerating the starting material [89] (eqn. (70)).

(e) [2+3] Cycloadditions

Reactions of α -phenyl-N-tert-butylnitrone with germaphosphenes 96 and 100 afford the corresponding five-membered ring heterocycles 111 and 112 [89] (eqn. (71)). Whilst two diastereoisomers were obtained for 112, only one was observed for 111, probably due to the large steric hindrance imposed by the 2,4,6-tri-tert-butylphenyl group. An insertion with ring expansion, equivalent to a [2+3] cycloaddition, involving oxirane or thiirane, was the first method of characterization of transient germaphosphenes obtained by dechlorosilylation reactions between dichlorogermanes and disilylphosphines [109] (eqn. (72)).

$$R_{2}GeCl_{2} + PhP \xrightarrow{Si} \xrightarrow{Cl} [R_{2}Ge=PPh] \xrightarrow{X} R_{2}Ge-PPh \\ \xrightarrow{Me_{2}Si} \xrightarrow{Me_{2}Si} X: O, S$$

$$(71)$$

$$R_{2}Ge-PPh X$$

$$(72)$$

(f) [2+4] Cycloadditions

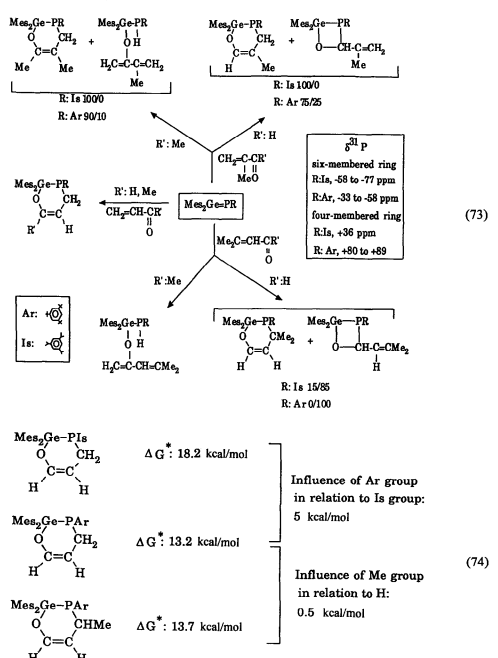
Germaphosphenes 96 and 100 react with α -ethylenic aldehydes and ketones to produce six-membered ring germaoxaphosphorinenes according to a [2+4] cycloaddition [110]. However, four-membered ring compounds ([2+2] cycloaddition) and open-chain products are also obtained with α or β methyl-substituted aldehydes and ketones, respectively (eqn. (73)). Hence, it seems that the outcome of the reaction strongly depends upon steric hindrance, both on the carbonyl compound and on the germaphosphene. The first step of these reactions is probably the nucleophilic attack of oxygen on the germanium centre while the attack of phosphorus occurs at the carbonyl carbon or the β carbon, depending on the substituents on these atoms.

Four- and six-membered ring derivatives are easily differentiated in ³¹P NMR spectra by their very different chemical shifts (-33 to -77 ppm for six-membered rings, +36 to +89 ppm for four-membered rings) [110].

The inversion barrier of phosphorus has been determined for three derivatives [110] (eqn. (74)). These low barriers are due to the substitution of phosphorus by the electropositive germanium (electronegativity 2.0) and particularly by the large steric effects of the substituted aromatic rings. Such studies have determined the influence of the aryl group bonded to phosphorus (Ar in relation to Is) and of the substitution on the 4-carbon of the carbonyl (Me in relation to H).

Interestingly an insertion with ring expansion, equivalent to a [2+4] cycloaddition, has previously been observed with germaphosphenes obtained by thermolysis of four-

membered rings. This was, for a long time, practically the only evidence for the formation of transient germaphosphenes [114] (eqn. (75)).



$$\begin{array}{c|c}
R_2Ge - PPh & 150^{\circ}C/10^{\circ 2} \text{mmHg} \\
& - CH_2 = CH_2
\end{array}
\qquad
\begin{bmatrix}
R_2Ge = PPh
\end{bmatrix}
\qquad
\begin{array}{c}
113 \\
R_2Ge
\end{array}
\qquad
PPh$$
P-GeR₂
PPh

(R: Me, Et)

Various types of cycloadditions have been observed between α -ethylenic aldehydes and ketones and other doubly bonded main group elements including exclusive [2+2] cycloadditions with disilenes [111], [2+4] with germenes [25], [2+2] and [2+4] with silenes [112]. With diphosphenes a cycloaddition involving one of the phosphorus atoms was observed to give a five-membered ring [113] (eqn. (76)).

(g) Thermolysis

Heating 96 at 140°C affords the stable germaphosphetene 114 [115] (eqn. (77)) probably by an intramolecular radical aromatic substitution. Compound 114 is the first heterocycle of this type to be prepared and has been structurally characterized. Although the four-membered ring is highly strained, it is thermally stable.

$$Mes_2Ge=PAr \xrightarrow{140^{\circ}C} Mes_2Ge \xrightarrow{P} \xrightarrow{solvent} Mes_2Ge-PH$$
96 114 (77)

Of the silaphosphenes, >Si=P-, germaphosphenes >Ge=P- and stannaphosphenes >Sn=P- series, the germaphosphenes have been the most extensively studied. This is

probably due to their straightforward synthesis allowing the isolation of pure crystalline derivatives. The germanium—phosphorus double bond appears very reactive in spite of the large steric hindrance necessary for its stabilization.

F. GERMATHIONES >Ge=S

(i) Synthesis

Stable germathiones are obtained by two routes (eqn. (78)):

coupling of a germylene with sulfur (route a);

decomposition of a germatetrathiolane (route b).

(a) Route a

The germathione 115 is obtained in high yield by a reaction between the stable germylene 116 and sulfur [116] (eqn. (79)). All previous attempts to isolate germathiones failed even with sterically demanding ligands on the germanium, and compound 115 is the first derivative with a base-stabilized double bond between sulfur and germanium. Stabilization of a doubly bonded compound of the type >Ge= M_{16} (M_{16} = O, S, Se) is of course much more difficult than the stabilization of compounds such as >Ge= M_{15} (M_{15} = N, P) or >Ge= M_{14} (M_{14} = C, Ge) since there is no substituent on the M_{16} element. Therefore complexation of germanium by a Lewis base is advantageous. A similar stabilization allowed the isolation of a stable compound with a silicon-sulfur double bond [117].

By contrast with sulfur, the reaction of germylene 116 with oxygen did not afford stable germanone >Ge=O, but only its dimer [116].

(b) Route b

Recently, Tokitoh obtained the new stable germathione 118 by thermolysis of the germatetrathiolane 117. Compound 118 is stabilized by the extremely bulky substituents on germanium (eqn. (80)) [121].

(ii) Physicochemical studies

(a) X-Ray diffraction

The geometry of germanium in 115 can be described as trigonal-planar (GeSN⁴N²) with an additional bond (GeN³) since the sum of angles on germanium deviates by only 5° from the ideal value of 360° [116]. The Ge-S bond length (2.063(3) Å) is the shortest germanium-sulfur bond observed, 0.2 Å shorter than that of a single bond [118] (Fig. 9). Theoretical calculations predict the Ge-S double bond length to be 2.02 Å in H₂Ge-S [119] (using pseudopotentials or MNDO methods) and 2.01 Å in Me₂Ge-S [120] (ab initio Hartree-Fock level calculations). On the basis of X-ray and NMR measurements, it seems that 115 can be formally described in terms of the two resonance structure 115a and 115b [67] (eqn. (81)).

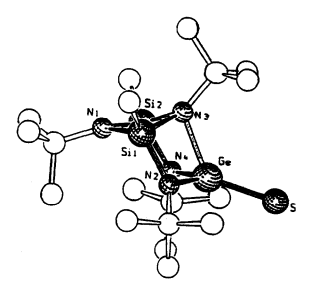


Fig. 9. Molecular view of 115 (reprinted from Angew. Chem. Int. Ed. Engl., 28 (1989) 1237).

TABLE 5
Germathiones

	Synthetic route ^a	d(Ge=S)(A)	Ref.
tBuN N Ge=S tBuN N Ge=S tBuN N Ge=S tBuN N Ge=S tBu Bu	a	2. 063 (3)	116
Ar Ar: $(R:(Me_3Si)_2)$	R.		121

^a(a) Reaction of a stable germylene with sulfur; (b) decomposition of a germatetrathiolane.

(b) IR spectra

The only IR spectrum of a germathione obtained to date is that for transient Me₂Ge=S trapped in an argon matrix at 18-20 K. The observed Ge=S stretching vibration (604.6 cm⁻¹) is in good agreement with calculated values (666 cm⁻¹ [119] and 586 cm⁻¹ [la]).

(iii) Reactivity

The reactivity of transient germathiones is well known (see [4] and refs. therein). But, for the recently prepared stable germathione 115, only the addition of methyl iodide to the germanium-sulfur double bond has been studied [75] (eqn. (82)). In the adduct 119, iodine is bound to germanium and the methyl group to sulfur, as expected from the bond polarity $Ge^{\delta +}=S^{\delta -}$. This reaction shows that the base-stabilized germathione 115 behaves as a normal unsaturated molecule.

$$\begin{array}{ccc} Ge=S & \xrightarrow{IMe} & Ge-S & \\ & IMe & \\ & & & \\ &$$

G. CONCLUSION

Great progress has been made in the field of organogermanium doubly bonded species in the 1980s and the beginning of the 1990s during which more than 30 stable germenes, digermenes, germaimines, germaphosphenes and germathiones have been synthesized and isolated. These compounds are stabilized by various means, the most important being the use of very bulky substituents which prevent their oligomerization. However, electronic effects, such as conjugation, also play a role, especially in germenes and germaimines. In the last case, the powerful—I effect of CF₃ attached to nitrogen probably allowed the extraordinary stabilization of germaimines substituted by small groups. The intermolecular or intramolecular base stabilization (by oxygen or nitrogen) also appears to be an effective way to stabilize germaimines.

Some doubly bonded germanium species have been structurally characterized by single crystal X-ray diffraction, showing germanium to be planar or nearly planar, with a significant bond shortening of about 8-10% relative to the corresponding single bond. Studies of the reactivity of these species confirm the presence of >Ge=X unsaturation as easy and nearly quantitative additions on the double bond are observed with many electrophiles and nucleophiles. Various types of cycloadditions ([2+1], [2+2], [2+3], [2+4]) are also possible with these species. The Ge=X derivatives appear extremely reactive, much more than the corresponding carbon analogues. Their reactivity as a germylene has never been observed (except in Lappert's digermene), but only their behavior as doubly bonded germanium, proving that they retain their structural integrity in solution.

Stable >Ge=X compounds are of particular interest because their reactivities can be thoroughly studied whereas studies of transient species are severely limited. The study of these new 'organometallic functions' is now in progress and appears very promising, particularly in organic and organometallic synthesis, including organometallic polymers. Although the chemistry of low-coordinated germanium species has developed rapidly during the last 10 years, many aspects remain to be examined, such as the synthesis and isolation of new types of doubly bonded germanium species including stable >Ge=B-, >Ge=Si<, >Ge=Sn<, >Ge=As- and >Ge=O. Another challenge is the preparation of triply-bonded germanium species -Ge=X, germa-allenic derivatives X=Ge=X, heterogermadienes and heterodigermadienes (for example: >Ge=X-X=X, X=Ge-X=X, >Ge=X-X=Ge<, X=Ge-Ge=X... with X=C, N, P) which, as yet, have neither been isolated nor clearly characterized by a trapping reaction.

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