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Characterisation of Bis-methylgermanium and Bis-methylsilicon Carbodi-imides and their Reactivity with Protic Reagents

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Bismethylgermanium and bismethylsilicon carbodi-imides have been prepared in high yield by halide metathesis reactions with lead(II), silver(I), or silicon(IV) species. The new derivatives in the series of general formula $(Me_nH_{3-n}MN:)_2C$ (M = Si or Ge; n = 1, 2, or 3) have been characterised by ¹H n.m.r., i.r., and Raman spectroscopy as well as mass spectrometry and cleavage reactions. The utility of the germanium carbodi-imides as synthetic intermediates is shown by ready cleavage of the Ge-N bond by Group VI and other protic species leading to the formation of chalcogeno-germanes and related derivatives of the type $(Me_nH_{3-n}Ge)_2E$ (E = O, S, Se, or Te) and $Me_{n}H_{3-n}GeSR$ (R = Me, Bu^t, Ph, MeC(O), or -CH₂CH₂-). Comparative studies show that in the silicon carbodi-imides the Si-N bond is not susceptible to protolysis by the heavier chalcogenols.

NUMEROUS studies have demonstrated the utility of species containing the Si-N bond as synthetic intermediates in silane¹ and organosilicon² chemistry. Studies of the analogous Ge-N species have been confined largely to the fully substituted organogermanes.³

Several methods for the preparation of fully substituted organometallic carbodi-imides of the type R_3M . NCN·MR₃ (R = alkyl or aryl; M = Si, Ge, or Sn) are described in the literature.⁴⁻⁶ Early work by Pump and Wannagat⁴ showed that bistrimethylsilylcarbodi-imide, $(Me_3SiN:)_2C$, was formed in a variety of systems such as reaction of Na(R₃Si)₂N with COCl₂, Me₃SiCl with (LiMe₃-SiN)₂CO, or Me₃SiCl with AgCN₂. A modification to the silver salt procedure has recently appeared.⁵ Other workers report the formation of the germanium species (Et₃GeN:)₂C and (Ph₃GeN:)₂C in the reactions of the corresponding oxide or bromide respectively with cyanamide, H₂NCN, under forcing conditions.⁶ We find that Me₃SiCl and Me₃GeBr are readily converted into the carbodi-imides, (Me₃SiN:)₂C and (Me₃GeN:)₂C, by direct

heating with lead(II) cyanamide. However, these and the foregoing reactions are not suited for the formation of the thermally unstable hydridic species and, indeed, no reaction occurs between Me₂HSiCl and PbCN₂ at room temperature in the gas phase. In keeping with the well known heavy-metal salt exchange series 7 the metathetical reaction of the iodo-silanes and -germanes with lead(II) cyanamide leads to extremely satisfactory yields of the corresponding carbodi-imido-species, (Me_nH_{3-n}) MN:)₂C (M = Si or Ge; n = 0, 1, 2, or 3) [equation (1)].

$$2 \ge M-I + PbCN_2 \longrightarrow (\ge M-N:)_2C + PbI_2 (M = Si \text{ or } Ge) \quad (I)$$

The comparative reactions of the iodides with silver(I) cyanamide give significantly reduced yields with increased disproportionation as has been observed in other systems employing silver-salt exchange reactions.^{8,9}

Alternatively, the germanium carbodi-imides may be obtained from bistrimethylsilylcarbodi-imide by ex-

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⁶ A. Vostokov and Yu. I. Dergunov, Zhur. obshchei Khim., 1970, 40, 1666 and 1971, 41, 1647.

 ⁷ A. G. MacDiarmid, *Quart. Rev.*, 1956, 10, 208; C. Eaborn,
 ⁶ Organosilicon Compounds,' Butterworths, London, 1960, p. 147.
 ⁸ E. A. V. Ebsworth and M. J. Mays, *J. Chem. Soc.*, 1962, 4844;
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 ⁹ E. A. V. Ebsworth and M. J. Mays, *J. Chem. Soc.*, 1961, 4870; Snetwechim Acta 1962, 1061, 1062.

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change with the corresponding fluorogermanes.¹⁰ While the illustrative example (2) is a high yield reaction it is

$$(Me_{3}SiN:)_{2}C + 2MeH_{2}GeF \longrightarrow (MeH_{2}GeN:)_{2}C + 2Me_{3}SiF \quad (2)$$

less convenient because the fluoride must first be prepared from heavier halides. Thus the lead cyanamide route is the preferable one.

It is interesting to compare the reactions of the organometallic carbodi-imides to their organic analogues RNCNR. In the hydrolysis of an organic carbodiimide¹¹ the central carbon atom provides the electropositive centre for nucleophilic attack so that an urea results with no fission of a C-N bond. By contrast, in the germanium or silicon derivatives, the metal atom provides the focus for nucleophilic attack with cleavage

and arene-thiols [equation (4)] providing convenient
syntheses for the previously reported
$$H_3GeSMe^{12}$$
 and
 H_3GeSPh^{13} as well as the new hydride derivatives,
 H_3GeSBu^t , $H_3GeSC(O)Me$, $(H_3GeSCH_2^{-})_2$, and
 $Me_nH_{3-n}GeSR$ [equation (4)].

$$(H_{3}GeN:)_{2}C + RSH \longrightarrow H_{3}GeSR + \frac{1}{2}(H_{2}NCN)_{2} \quad (4)$$

$$R = Me, Bu^{t}, MeC(O), -CH_{2}CH_{2}-, \text{ or } Ph$$

$$also H_{3}Ge \equiv Me_{n}H_{3-n}Ge$$

Comparative experiments indicate that the SiNCNSi system is more resistant than is the GeNCNGe linkage to nucleophilic attack by the heavier chalcogenols, SH, SeH, and TeH. Proponents of the importance of (p-d) π bonding in the Si-N system,¹⁴ relative to Ge-N, would no doubt accept this as further evidence of the 'extra stability of the Si-N bond or of the decreased polarity of

TABLE 1

Protolysis reactions of (R₃MN.)₂C species with chalcogenols

 $(R_3MN)_2C + H_2E \longrightarrow (R_3M)_2E + \frac{1}{2}(H_2NCN)_2$

				(I)	-	(11)						
	$\mathbf{E} = \mathbf{O}$		E = S		E = Se		E = Te					
	(1)	(II)	Yield	(I)	(II)	Yield	$\overline{(I)}$	(II)	Yield	(I)	(II)	Yield
$R_{3}M$	(mmol)	(mmol)	(%)	(mmol)	(mmol)	(%)	(mmol)	(mmol)	(%)	(mmol)	(mmol)	(%)
MeH ₂ Ge	0.75	0.68	90	1.45	1.38	95	1.42	1.18	83	1.62	1.01	62
Me "H Ge	1.41	1.16	82	1.50	1.32	88	1.52	1.34	88	1.50	1.18	79
Me ₃ Ge	1.24	1.07	86	0.89	0.74	83	1.55	1.21	78	1.09	0.97	89
MeaSi	2.40	2.30	96	No	o reaction	1	No	reaction	a	Ne	o reaction	L
H₃Ğe	2.36	1.91	98 ^b	0.24	0.22	92	1.25	1.05	84 °			

⁶ BuSeH is reported to react with (Me₃SiN:)₂C (100 °C, 1 h, benzene) to give Me₃SiSeBu (70%), Yu I. Dergunov, I. A. Vostokov, and V. T. Bychkov, *Zhur. obshchci Khim.*, 1972, 42, 371, 380. ^b Ref. 10. ^c S. Cradock, E. A. V. Ebsworth, and D. W. H. Rankin, J. Chem. Soc. (A), 1969, 1628.

TABLE 2

Protolysis reactions of (R₃MN:)₂C species with various thiols

 $(R_3MN:)_2C + 2R'SH \longrightarrow 2R_3MSR^3 + \frac{1}{2}(H_2NCN)_2$

		(1)	(11)			
	$\mathbf{R} = \mathbf{M}\mathbf{e}$	$R = Bu^t$	R = Ph	$\mathbf{R} = -\mathbf{C}\mathbf{H_2}\mathbf{C}\mathbf{H_2} -$	R = MeC(O)	
	(I) (II) Yield	(I) (II) Yield	(I) (II) Yield	(I) (II) Yield	(I) (II) Yield	
$R_{3}M$	(mmol) (mmol) (%)	(mmol) (mmol) (%)	(mmol) (mmol) $\binom{0}{0}$	(mmol) $(mmol)$ $(%)$	(mmol) (mmol) (%)	
H ₃ Ge	0.45 0.72 81	1.41 2.53 90	0.60 0.98 80	0.86 0.42 49	1.05 1.64 78	
MeH.Ge	$1.42 \ 2.15 \ 75$	1.25 2.13 85	1.41 2.20 78	1.40 1.15 82	1.30 2.31 89	
Me "H Ge	1.42 2.38 84	0.95 1.49 78	1.12 1.77 79	1.02 0.70 68	1.12 1.95 87	
Me Ge	1.08 1.69 78	1.50 2.60 87	0.99 1.41 71	0.99 0.83 83	0.99 1.53 77	
Me₃Si	No reaction	2.00 1.56 39	2.70 1.83 34	No reaction	1.90 1.59 42	

of the M-N bond and formation of $(H_2NCN)_n$ and the germyl or silvl chalcogenide.

Rapid and non-reversible reactions occur with weak acid donors (Tables 1 and 2). With hydrogen chalcogenides, H₂E, protolytic cleavage leads to the corresponding germyl-Group VI derivatives and dicyanodiamide [equation (3)].

c.g.
$$(MeH_2GeN)_2C + H_2E \longrightarrow$$

 $(MeH_2Ge)_2E + \frac{1}{2}(H_2NCN)_2$ (3)
 $(E = O, S, Se, or Te)$

Cleavage also occurs with a wide variety of alkane-¹⁰ S. Cradock and E. A. V. Ebsworth, J. Chem. Soc. (A), 1968, 1423. ¹¹ F. Kurzev and K. Daurachi-Zadeh, Chem. Rev., 1967, 67,

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 ¹² J. T. Wang and C. H. Van Dyke, Chem. Comm., 1967, 612.

the bond making the silicon atom less open to nucleoattack. Unfortunately, supportive thermophilic chemical data, particularly for germyl derivatives, is scant and inconsistent. It seems unlikely that the supposed relative weakness of the Ge-N bond could completely account for the quantitative cleavage of Ge-N to form Ge-O, -S, -Se, and -Te bonds. However, the ready cleavage of Si-N by H₂O and ROH is probably associated with the formation of the strong Si-O bond.¹⁵

More important than these speculations is that we have

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¹⁴ E. A. V. Ebsworth in 'Organometallic Compounds of the Group IV Elements,' ed. A. G. MacDiarmid, Dekker, New York,

¹⁶ T. Tanaka, J. Inorg. Nuclear Chem., 1960, 13, 225; A. E. Beezer and C. T. Mortimer, J. Chem. Soc. (A), 1966, 514.

established that the Ge-N linkage in germanium carbodiimides makes them excellent and versatile intermediates for the formation of other Ge-M bonds, a property we are utilising in current researches.

EXPERIMENTAL

Manipulations and spectroscopic measurements were as described previously.16 Silyl and germyl halides were prepared by standard methods,¹⁷⁻¹⁹ and their purity established spectroscopically. Ag2CN2 Was obtained from AgNO₃ and CaCN₂ in basic solution.²⁰ Hydrogen sulphide, selenide, and telluride were prepared by hydrolysis of aluminum chalcogenides.²¹ Other materials were reagent grade of commercial origin.

Preparation of the Carbodi-imides, $(Me_nH_{3-n}MN^*)_2C$ (M = Si or Ge; n = 0, 1, 2, or 3). (i) Reactions of $Me_nH_{3-n}MX$ with $PbCN_2$ and Ag_2CN_2 . The gaseous iodo-silane or -germane (ca. 5 mmol) was passed at 25 °C over the anhydrous cyanamide (ca. 25 g) held in a column on glass wool. Exothermic reactions ensued and after two or three doublepasses the ¹H n.m.r. spectra of the colourless, sparingly volatile, material showed no iodide resonances. After fractional condensation at -23 °C the yields of the silyland germyl-carbodi-imides were frequently in the region of 90% in the PbCN₂ reactions with little evidence for disproportionation: e.g. GeH₃I (0.50 mmol) gave (H₃GeN.)₂C (0.21 mmol, 84%; v.p. ca. 0.6 mmHg at 0 °C; m.p. 10 °C, δ GeH 4.94 p.p.m.) and SiH_aI (2.89 mmol) gave (H_aSiN.)₂C (1.25 mmol, 87%; v.p. 18 mmHg at 0 °C, δ SiH 4.45 p.p.m.). Using Ag_2CN_2 yields of only 40-60% could be realised with considerable formation of involatile residues (shown to contain polymeric H₂NCN) and parent hydrides. The corresponding hydrido-chlorides or-bromides showed little or no reactivity in the gas phase. However, the high thermal stability of Me₃SiCl (11.20 mmol) and Me₃GeBr (11.30 mmol) permitted direct heating (ca. 100 °C, 5 h) with the cyanamide salt (ca. 20 g) in a sealed ampoule when again satisfactory conversion into (Me₃SiN.)₂C (5.03 mmol, 90%) and (Me₃-GeN.)₂C (5.08 mmol, 90%) occurred.

(ii) Reaction of $Me_nH_{3-n}GeF$ with bis(trimethylsilyl)carbodi-imide. Typically (Me₃SiN:)₂C (0.597 g, 3.21 mmol) was distilled into a reaction vessel (10 ml) held at -196 °C containing a slight excess of MeH₂GeF (0.833 g, 7.67 mmol). The mixture was allowed to react (room temp., 15 min) and then vacuum fractionated. A pure sample of (MeH₂GeN:)₂C (0.636 g, 2.90 mmol; 90.3%) was retained in a trap at -23 °C; traces of (Me₃SiN:)₂C (ca. 0.3 mmol) in a trap at -45 °C; small amounts of MeH₂GeF (ca. 1.8 mmol; identified by i.r. and n.m.r. spectra ¹⁷) in a trap at -95 °C; and Me₃SiF (ca. 5.8 mmol; identified by i.r.²² and n.m.r.²³ spectra) in a -196 °C following trap. By similar procedures Me₂HGeF (8.80 mmol) and Me₃GeF (9.12 mmol) reacted with (Me₃SiN:)₂C (ca. 4 mmol) to give (Me₂HGeN:)₂C

¹⁶ J. E. Drake and R. T. Hemmings, Canad. J. Chem., 1973, 51,

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 ¹⁷ J. E. Drake, R. T. Hemmings, and C. Riddle, J. Chem. Soc. (A), 1970, 3359; G. K. Barker, J. E. Drake, R. T. Hemmings, and B. Rapp, J. Chem. Soc. (A), 1971, 3291; Spectrochim. Acta, 1972, 28A, 1113.
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(3.44 mmol, 89.4%) and $(Me_3GeN.)_2C$ (3.35 mmol; 88.9%)respectively.

Characterisation of the Carbodi-imides.-(i) ¹H N.m.r. spectra. All the carbodi-imides give first-order spectra (Table 3) consistent with free rotation about the C-M bonds. The chemical shifts (δ p.p.m.) of M-H resonances in the $(Me_nH_{3-n}MN)_2C$ species are comparable to those of related compounds containing an M-N bond, viz. (H₃Ge)₃N 4.9; H₃Ge-N₃ 5.1; -NCO 5.05, -NCS 5.18; (H₃Si)₃N 4.44; H₃Si-N₃ 4.49; -NCO 4.42, -NCS 4.46 δ.^{24, 25} Marked dilution shifts (e.g. ca. 0.5 p.p.m. from neat liquid to 5% CCl₄ solution) suggest association. This is supported by line-broadening in the more concentrated solutions. Broadening as a result of interactions involving the ¹⁴N quadrupole should be concentration independent.

(ii) The vibrational spectra of $(Me_nH_{3-n}MN!)_2C$ (M = Si or Ge). These spectra are displayed in Tables 3-5. Prin-

TABLE 3

The ¹H n.m.r. parameters * of $(Me_nH_{3-n}MN)_2C$ (M = Si or Ge) and $(Me_{*}H_{*}, Ge)_{*}E$ (E = O. S. Se, or Te) species

	(/2- (-	0, 0, 00, 0	(10) Speek
Compound	δ(Me)	δ(MH)	JHH ^{vic}	Јсн
(MeH ₂ SiN:) ₂ C ^a	0.36	4.61	3.75	125.5
(Me ₂ HSiN:) ₂ C ^b	0.25	4.74	3.30	123.4
(Me ₃ SiN:) ₂ C ^c	0.14			118.6
(MeH ₂ GeN:) ₂ C	0.58	4.99	3.00	129.6
(Me ₂ HGeN:) ₂ C	0.51	5.18	2.63	129.0
(Me ₃ GeN:) ₂ C	0.43			128.1
(MeH ₂ Ge) ₂ O	0.59	5.28	2.91	129.1
(Me ₂ HGe) ₂ O	0.40	5.40	2.43	128.2
(Me ₃ Ge) ₂ O ^d	0.29			125.9
(MeH ₂ Ge) ₂ S	0.66	4.87	3.30	129.4
(Me ₂ HGe) ₂ S	0.54	4.93	2.91	129.0
(Me ₃ Ge) ₂ S ^d	0.51			127.5
(MeH ₂ Ge) ₂ Se	0.77	4.55	3.38	129.6
(Me ₂ HGe) ₂ Se	0.69	4.73	2.96	127.5
(Me ₃ Ge) ₂ Se ^d	0.58			127.1
(MeH ₂ Ge) ₂ Te	0.93	4.12	3.53	132.6
(Me ₂ HGe) ₂ Te •	0.80	4.65	3.37	131.7
(Me ₃ Ge) ₂ Te ^f	0.71			129.1

* The spectra were recorded at ambient temperature in CCl solution (ca. 5% v/v). Chemical shifts ($\delta \pm 0.02$ p.p.m.) are in p.p.m. to low field of Me₄Si as internal standard. Deviations for coupling constants are $J(\text{HH}) \pm 0.05$ Hz, $J(^{13}\text{CH}) \pm 0.2$ Hz. Deviations

• $J(^{29}SiH)$, 218 Hz. • $J(^{29}SiH)$, 217 Hz. • $J_{\rm SiH}^{\rm gem}$, 6.9 Hz. • Compare with $\delta(Me_3Ge)_2E$ (CCl₄ solution) of 0.31(0); 0.53(S); 0.66(Se) p.p.m. in H. Schmidbaur and I. Ruidisch, *Inorg. Chem.*, 1964, **3**, 599. • Recorded in cyclohexane (5% v/v). ^J Agrees with approximate value of δ 0.74 p.p.m. (C₆H₆ solution) in H. Schumann, R. Mohtachemi, H. J. Kroth, and U. Frank, *Chem. Ber.*, 1973, **106**, 2049; $J_{\rm HTe}^{\rm vic}$ 5.5 Hz.

cipal features can be assigned by comparison with the spectra of related but simpler molecules.9,10,16-18 The N= C=N asymmetric stretch is placed in the i.r. spectra at ca. $2 250 \text{ cm}^{-1}$ for the silanes and $2 150 \text{ cm}^{-1}$ for the germanes. Not surprisingly, in view of the effective, local centro-

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The i.r. and Raman spectra (cm⁻¹) of the bismethylgermanium carbodi-imides *

	(MeH	I ₂ GeN:) ₂ C	$(Me_2HGeN.)_2C$ $(Me_3Ge$		e ₃ GeN.) ₂ C	
Tentative assignment	I.r. (gas)	Raman (liq)	I.r. (liq)	Raman (liq)	I.r. (liq)	Raman (liq)
(a)	2.997m	2 992m. dp	2 990m	2 991m. dp	2.995m	2.985m, dp
CH_3 stretch $\begin{cases} (a) \\ (s) \end{cases}$	2 928w	2 919vs. D	2 916w	2 918vs, p	2 918m	2 915vs, p
N=C=N stretch (a)	2 1495	N.o.	2 142vs "	N.o.	2 130vs	N.o.
Ge-H stretch	2 086s	2 083vs. D	2 073vs	2 063vs, p		
N=C=N stretch (s)		1 418m. p ^b		1 412m, p ^b		1409m, p ^b
	1 405vw	, 1	1 417w	· •	1 408m	-
$CH_3 def. \left\{ \begin{array}{c} \langle x \rangle \\ \langle x \rangle \end{array} \right\}$	1 259w	1249m, p	1 249m	1 248m, p	1 252m	1 250m, p
GeH.def (sc)	882m	877m, p				
((a'')	845m	· · · 040-1	832s	850w, dp	825s	833 vw
CH_rock (a')	802s	<i>ca.</i> 840511	816s	-		
(a', a'')			ca. 760sh		ca. 750sh	763vw
GeH. GeH def.	728m	731m, dp	712m	705m, p		
2, 2,				636m, dp		
Skeletal bend	675m	675m, dp	661 m		670m	663 sh
$C = C$ = t = t = t $\int (a'')$		-	619s	603m, dp °	617s	615m, dp
Get stretch $\left(a' \right)'$	612m	611vs, p	593m	579vs, p	572m	576vs, p
GeH, rock	474w	478 m, dp				
Call stants (a)	545m		530s	<i>ca</i> . 500sh	527s	ca. 480sh
Gen stretch l(s)	398w?	421s, p	425w?	420s, p		430s, p
Claster I and a doc		227m, p		225m		$240 \mathrm{sh}$
Skeletal modes		194m, dp		184s, dp		189s
		· -				163ms

* Spectra recorded at room temperature.

m = medium, s = strong, w = weak, v = very, sh = shoulder, p = polarised, dp = depolarised.

• Gas. • CH₃(a) is expected in this region. • Polarisation spectrum.

TABLE 5

The i.r. and Raman spectra (cm⁻¹) of the bismethylsilicon carbodi-imides * †

	(Me	H ₂ SiN:) ₂ C	(M	e ₂ HSiN:) ₂ C	ISiN:) ₂ C (Me ₃ SiN:)		
Tentative assignment	I.r. (gas)	Raman (liq)	I.r. (gas)	Raman (liq)	I.r. (liq)	Raman (liq)	
CH_3 (stretch $\begin{cases} (a) \\ (s) \end{cases}$	2 981m 2 924w	2 972m, dp 2 911vs, p	2 972m 2 911w	2 968m, dp 2 906vs, p	2 967m 2 903w	2 966m, dp 2 904vs, p	
N=C=N stretch (a)	2 258vs	N.o.	2 241vs	N.o.	2 205s	N.o.	
Si-H stretch	2 176vs	2 170vs, p 1 566m	2 149vs	2 147vs, p 1 552m			
N=C=N stretch (s) ^a	N.o.	1 504m	N.o.	1 518m 1 481m	N.o.	1 514m 1 453m	
CH, def	1 419vw	1 418w, dp	1 429w 1 381w	ca. 1 420w, br	1 406w	1 416m, dp	
(b)	1 267m	1 257m. p	1 262ms	1 259m, p	1 253s	1 262m, p	
SiH, def (sc)	964s	963m, p				· 1	
CH, rock (a')	915vs ca. 870sh	929w, đp 868w, dp	906s 884vs	907w, dp 883w	837s	845w, br	
(a', a'')			838s	843w	736s	760w. dp	
SiH ₂ ,SiH def	686w	707vs, p		750m			
	<i>ca</i> . 730m	726m, dp	627m	632m			
Skeletal bend	587m	N.o.	584w 739s	N.o. 702m dp	581m 698m	N.o. 699m dn	
SiC stretch { a /	760-	761vs p	1003	674vs p	640w	645vs p	
SiH ₂ rock	512w	515m, dp		01103, p	01010	01073, p	
SiN stretch $\begin{cases} (a) \\ (c) \end{cases}$	789vs	4036 5	775s	772w?	760s	480c D	
(s)		4935, p		273m	279m	283E, dp	
Skeletal modes		232m, p		246m 199s dp		208s, p 195sh	
	* Se	e footnote to Table 4	. + N.o. = no	t observed.			

^a See text.

symmetry of the carbodi-imide structure ²⁶ this mode is absent in the Raman effect.²⁷ The N=C=N symmetric stretch is a polarised medium-intensity band at *ca*. 1 409— 1 418 cm⁻¹ in the Raman spectra of the germanes. In the silanes, a more complex pattern appears at 1 500—1 570 cm⁻¹ which seems to be a characteristic feature of silylcarbodi-imides ⁹ and so is assigned as the N=C=N symmetric

²⁶ J. D. Murdoch and D. W. H. Rankin, J.C.S. Chem. Comm., 1972, 748.

stretch. In $(H_3SiN:)_2C$, the feature was attributed to Fermi resonance with the overtone of the i.r.-active SiH_3 deformation mode.⁹ In $(MeH_2SiN:)_2C$ it is reasonable to assume that there is Fermi resonance between the N=C=N symmetric stretch and the overtone $(2 \times 761 \text{ cm}^{-1})$. For

²⁷ M. Davies and W. J. Jones, *Trans. Faraday Soc.*, 1958, 54, 1454; L. Kahovec and K. W. F. Kohlrausch, *Z. phys. Chem.*, 1937, **B37**, 421; G. D. Meakins and R. J. Moss, *J. Chem. Soc.*, 1957, 993.

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the other species, the rationale for the complex band-pattern is less obvious, but could involve the overtone of the Si-N asymmetric stretch, which is expected in the region 760- 790 cm^{-1} [788 cm⁻¹ in (H₃SiN:)₂C⁹]. For the germanium series, the assignment of the Ge-N asymmetric stretch to a strong i.r. band in the region 527-545 cm⁻¹ [547 cm⁻¹ in $(H_3GeN!)_2C$ ¹⁰ is clear as are the assignments of the Ge-C stretching modes (ca. 576-619 cm⁻¹) 17 and the GeH₂ and GeH deformation modes. The M-N symmetric stretches appear in the Raman spectra as strong polarised bands in the regions of 480-493 cm⁻¹ for the silanes [496 cm⁻¹ in $(H_3SiN.)_2C$]⁹ and of 420-430 cm⁻¹ for the germanes [410 cm⁻¹ in (H₃GeN.)₂C].¹⁰

(iii) Mass spectra. In all cases the molecular ion, viz., $(Me_nH_{3-n}MN^*)_2C^+$, is detected and provides confirmation of the molecular species. The low normalised abundance (i.e. ca. 1 to 20) of this ion is a common feature in the spectra of organo-silanes and -germanes.28,29 Ions arising from H and Me stripping appear to be most abundant although significant ion current in the germanes is carried by 'rearranged ' ions such as $\mathrm{Ge}_2\mathrm{N}^+$ or $Ge_{2^{2^{+}}}$ and ions from N-C bond cleavage, *i.e.* $Me_{n}H_{3-n}GeN^{+}$. This is in general agreement with the observations in the spectrum of (H₃GeN:)₂C but we have been unable to detect ions of the type MN₂C⁺ reported in the same study.¹⁰

(iv) Chemical characterisation (i.e. analysis for $Me_nH_{3-n}M^{-1}$ groups). This was achieved by protolytic cleavage of the M-N linkage with an excess of gaseous hydrogen halides leading to the isolation of stoicheiometric quantities of familiar halogeno-silanes and -germanes, $Me_nH_{3-n}MX$ (X = Cl, Br, or I), which were identified spectroscopically.17, 18, 30, 31 The extensive protolysis reactions with chalcogenols (see next section and Table 1) also provide indirect evidence for the stoicheiometric/monomeric nature of the carbodi-imides.

Reactions of the Carbodi-imides with Selected Protic Reagents.—(i) Reactions of $(R_3MN)_2C$ (R = H or Me) with H_2O (E = O, S, Se, or Te). A general procedure was followed for all the volatile protic reagents. The carbodiimide (ca. 1-2 mmol) was distilled into a greaseless reaction vessel (150 ml) with a bulb which could be immersed in a low-temperature bath and a side arm which could be sealed. An excess of the H₂E species was distilled into the vessel with the bulb head at -196 °C. The mixture was allowed to attain room temperature, with quenching as the reaction became too vigorous (particularly for the thermally unstable H_2 Te). After typically 60 min, vacuum fractionation of the products was carried out. For the H2S, H2Se, and H2Te reactions examination of the products involatile at -45 °C showed no resonances attributable to unchanged carbodiimide whilst the excess of H₂E, volatile at this temperature, was identified in a -196 °C trap. For the H₂O reactions the excess of H_2O was retained in a -23 °C trap and spectroscopically pure oxides were collected in a -196 °C trap. An amorphous white material remaining in the reaction vessel was subsequently characterised by i.r. spectroscopy 32 as polymeric H₂NCN. Details of the reactions are given in Table 1 and the ¹H n.m.r. parameters of the new methylgermanium chalcogenides are collected in Table 3. General features in the Raman spectra of (Me₂HGe)₂E characteristic of $Me_nH_{3-n}Ge^-$ and Ge-E-Ge moieties appear as follows (cm^{-1}) : Ge-E stretching (2 bands) at 486m,p and 792s (i.r.) [O]; ca. 380vs,p and 409s (i.r.) [S]; 273vs,p and 282 s (i.r.) [Se]; 228vs,p [Te]; Ge-C stretching (2 bands) at ca. 585vs,p and ca. 610m,dp; Ge-H stretching at ca. 2 040-2 080vs,p; CH₃ stretching (2 bands) at ca. 2 915 vs,p and 2 985 m,dp; CH₃ def. (2 bands) at ca. 1 240m,p and 1 414 w,dp; Ge-E-Ge def. at 169m,p [O]; 95m,p [S]; 76m,p [Se], and 63m,p [Te].

Analysis for the $Me_nH_{3-n}Ge^-$ groups was carried out by cleavage of the germanium-chalcogen bond with an excess of gaseous HI, which produced stoicheiometric quantities of H_2E and the familiar iodogermanes, $Me_nH_{3-n}GeI$, which were identified spectroscopically.^{17,18} The 70 eV mass spectra of the $(Me_nH_{3-n}Ge)_2E$ species further support the formulation of the chalcogenides as well as providing molecular-weight confirmation and support for the a priori assignment of n.m.r. and vibrational spectra.

Further preliminary experiments 33 also show that the corresponding germyl chalcogenols $Me_nH_{3-n}GeEH$ (E = S, Se, or Te) are formed rapidly at room temperature when the chalcogenides are treated with an excess of H₂E in sealed tubes.

(ii) Reactions of (R₃MN:)₂C species with R'SH species. An exactly similar procedure to that described above was

TABLE 6

¹ H N.m.r.	parameters *	of Me	$e_n H_{3-n} MSF$	2 species	pre-
	pared ir	ı this	study †		

	δ	δ				
	(Me)	(GeH) ^b	J _{HH} vic	Jсн	δ(CH')	JCH'
IeH,GeSMe'	0.65	4.77	3.52	130.7	2.11	143.4
∕Ie₂H̃GeSMe′	0.56	4.86	3.24	129.5	2.04	142.8
Ae ₃ GeSMe′ ه	0.51			125.7	1.97	141.0
I3GeSCMe3'		4.35			1.42	125.5
MeH ₂ GeSCMe ₃ '	0.64	4.90	3.75	134.9	1.49	132.9
∕Ie₂HGeSCMe₃′	0.55	5.02	3.06	130.3	1.42	129.6
Ae ₃ GeSCMe ₃ ' °	0.50			127.5	1.40	126.3
Ie ₃ SiSCMe ₃ '	0.05			119.2	1.42	128.6
AeH ₂ GeSPh'	0.50	5.01	3.67	135.1	7.43	
∕Ie₂HGeSPh′	0.47	5.17	3.36	132.3	7.38	
Ae ₃ GeSPh' ^c	0.40			N.o.	7.16	
Ae ₃ SiSPh' ^b	0.25			123.0	7.15	
$H_3GeSCH'_2-)_2$		4.74			2.86	N.o.
$MeH_2GeSCH_2'-)_2$	0.71	4.89	3.53	133.8	2.79	N.o.
Me ₂ HGeSCH ₂ '-) ₂	0.61	5.04	3.45	132.7	2.72	N.o.
Me ₃ GeSCH ₂ '-)	0.50			N.o.	2.60	N.o.
I ₃ GeSC(O)Me'		4.62			2.49	127.2
$MeH_2GeSC(O)Me'$	0.78	4.92	3.42	133.8	2.47	135.0
Me ₂ HGeSC(O)Me'	0.68	5.03	3.18	131.7	2.41	132.0
Ie ₃ GeSC(O)Me'	0.60			128.1	2.35	128.8
Ae ₃ SiSC(O)Me'	0.39			120.8	2.54	131.1

* See footnote to Table 3. † N.o. = not observed.

^a Compare with values of $\delta(H_3 \text{GeSR})$: 4.71 (R = Ph), C. Glidewell and D. W. H. Rankin J. Chem. Soc. (A), 1969, 753 C($_{6}H_{12}$); 4.48 (R = Me), J. T. Wang and C. H. Van Dyke, *Inorg. Chem.*, 1968, **7**, 1319. ^b See K. A. Hooton and A. L. Allred, *Inorg. Chem.*, 1965, **4**, 671. ^e E. W. Abel, D. A. Armitage, and D. B. Brady, *J. Organometallic Chem.*, 1966, **5**, 130.

followed for the volatile MeSH and ButSH. The thiogermanes could be transferred in a pure state to the side arm after pumping off the excess of thiol at -45 °C. Reaction

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²⁸ F. Glockling and J. R. C. Light, J. Chem. Soc. (A), 1968, 717; D. B. Chambers, F. Glockling, and J. R. C. Light, Quart. Rev., 1968, 22, 317.

 ²⁹ G. P. Van der Kelen, O. Volders, H. Van Onckelen, and Z. Eeckhaut, Z. anorg. Chem., 1965, 338, 106.
 ³⁰ A. L. Smith, J. Chem. Phys., 1953, 21, 1997; J. R. Durig and C. W. Hawley, *ibid.*, 1973, 58, 237; *ibid.*, 1973, 59, 1; H. Burger, Scattering Acta, 1968, 24A, 2015. Spectrochim. Acta, 1968, 24A, 2015.

times of the order of 60 min at room temperature gave essentially quantitative conversion. The less-volatile species, PhSH, HSCH₂CH₂SH , and MeC(O)SH, were handled volumetrically and syringed into the reaction vessel and degassed thoroughly before distilling in the carbodi-imides. For these reactions it was found advantageous to use a deficit of the thiol (see Table 2). Unchanged (R₃MN:)₂C species were pumped from the reaction vessel leaving involatile oily sulphides which could be transferred to the side arm and sealed for subsequent spectroscopic analysis (Table 6). The amorphous white material adhering to the walls of the reaction vessel was again shown to be $(H_2NCN)_n$. The very low reactivity of (Me₃SiN:)₂C towards the thiols was evident even when the mixtures were held at higher temperatures (ca. 60-100 °C). Preliminary experiments

more acidic alcohols R'OH, leading to Me_aSiOR' species which together with analogous derivatives of the hydridosilanes and -germanes are currently under study and will be presented at a later date. Thus Me₃SiOMe' [88%; δ (Me) 0.07 and $\delta({\rm Me'})$ 3.37 p.p.m], $Me_3{\rm SiOPh}$ [90%; $\delta({\rm Me})$ 0.25

show that, as expected, (Me₃SiN:)₂C reacts rapidly with the

 ³⁴ S. Cradock, *J. Chem. Soc.* (A), 1968, 1426.
 ³⁵ D. F. Van De Vondel, E. V. Van den Berghe, and G. P. Van der Kelen, J. Organometallic Chem., 1970, 23, 105.

p.p.m], Me_3Si(OCH_2)_2 [80%; $\delta(Me)$ 0.08 and $\delta(CH_2)$ 3.57 p.p.m.] and Me_3SiOC(O)Me' [92%; $\delta(Me)$ 0.26, $\delta(Me')$ and 1.99] were obtained from (Me₃SiN:)₂C with MeOH, PhOH, HOCH₂CH₂OH, and HOC(O)Me respectively.

The ¹H n.m.r. parameters of the new Me_nH_{3-n}GeSR' species are presented in Table 6. The i.r. and Raman spectra show features characteristic of both $Me_nH_{3_n}Ge^-$ and -SR' moities but we have not attempted any detailed analysis at this stage. By comparison with the spectra of $\rm H_{3}GeSMe~^{34}$ and $\rm H_{3}GeSPh~^{13}$ and other thiogermanes 35 the Ge-S stretching frequency has been assigned as follows in the Raman spectra of the liquids: viz. 386-396 cm⁻¹ (R = Me); $450-451 \text{ cm}^{-1}$ (R = Bu^t); $378-381 \text{ cm}^{-1}$ (R = Ph); 393-398 cm⁻¹ (R = $-CH_2CH_2$ -); 288-293 cm⁻¹ [R = MeC(O)]. The Ge-H stretching frequency appears surprisingly low in the range 2 040-2 088 cm⁻¹ for all species. The mass spectra of the more volatile $Me_nH_{3-n}GeSMe$ have also been obtained and show the expected molecular ions in low abundance consistent with their formulation.

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