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SILYL- AND GERMYLMERCURIALS IN ORGANIC SYNTHESIS. A NEW ROUTE TO O-SILYLATED AND O-GERMYLATED ENOLATES *

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Summary

The exchange reaction of α -mercurated ketones with bis(triethylsilyl)- (I) and bis(triethylgermyl)mercury (II) leads to the formation of the corresponding triethylsilyl and triethylgermyl enol ethers. O-Silylated and O-germylated enolates derived from ketones and aldehydes can be also prepared by treating mercurials I and II with appropriate α -bromo-carbonyl compounds. This new pathway also represents the best available method for preparing bromo-containing triethylsilyl and triethylgermyl enol ethers. NMR and IR spectral characteristic useful in the identification and characterization of these and related compounds are summarized.

Introduction

Silyl-, germyl-, and stannyl-mercurials are widely used in organic synthesis [1,2]. From this point of view, the exchange reactions between $(R_3M)_2Hg$ (M = Si, Ge, Sn) and mercurials of the type HgX_2 of interest. These reactions usually lead to unsymmetrical compounds R_3M —HgX or their demercurated products [3,4].

$$(R_3M)_2Hg + HgX_2 \stackrel{a}{\to} 2 R_3M - HgX \stackrel{b}{\to} 2 R_3MX + 2 Hg$$
 (1)

For example, the interaction of bis(triethylsilyl)mercury (I) with $Hg[C(N_2)COOEt]_2$ proceeds according to eq. 1a, b to give metallic mercury and ethyl triethylsilyldiazoacetate [5,6]. The compound $[(Me_3SiCH_2)_3Sn]_2Hg$ reacts similarly with HgX_2 , when X is CH_2COOMe , PhC = C, or $Co(CO)_4$ [4]. In

^{*} Dedicated to Professor G.A. Razuvaev on the occasion of his 85th birthday on August 23rd, 1980.

contrast, the reaction of bis(triethylgermyl)mercury (II) with fluorinated dialkylmercurials, HgR_2^f ($R_2^f = CF_3$, CH_2CF_3 , CF_2CF_3 etc.) proceeds via formation of the corresponding fluoroalkyl(triethylgermyl)mercurials, Et_3GeHgR^f (i.e., via the reaction depicted in eq. 1a) [7].

The formation of the fluoroalkyl germyl-mercury compounds was interpreted in terms of a four-centre mechanism of the type previously proposed for the exchange reactions between (CH₃)₂Hg and (CD₃)₂Hg [8a] or (Me₃Si)₂Hg * and (Me₃Si)₂Hg [8b].

A similar mechanism would account for the interaction of optically active bis(methyl-1-napthylphenylgermyl) mercury with Hg(CH₂COOMe)₂. The reaction proceeds according to eq. 1a, b and yields a product which retains the configuration at germanium [9].

It is possible that demercuration of intermediate R_3M —HgX involves the formation of ion pairs $R_3MHg^{\dagger}X^{-}$ and/or $R_3M^{\dagger}X^{-}$ [10]. In accord with this conclusion, addition of $(Et_3Si)_2Hg$ to an acetone solution of $Hg[C(N_2)COMe]_2$ causes a formation of metallic mercury and α -triethylsilyldiazoacetone together with unexpected diacetone alcohol [10d]. The most likely possibility is that exchange occurred to give, initially, intermediate $Et_3SiHgC(N_2)COMe$ which was subsequently demercurated to the ion pair. The anion $C(N_2)SOMe$ generated catalyzes the aldol-type condensation of acetone to give diacetone alcohol.

The reactions of mercurials I and II with α -mercurated ketones and α -bromoketones have now been investigated to ascertain whether silyl and germyl enol ethers can be prepared.

Results and discussion

The exchange reaction of α -mercurybisacetophenone with mercurials I and II in THF at 65°C results in a high yield of triethylsilyl enol ether (III) and its germanium analogue (IVa), respectively. Since the starting α -mercurated ketone has no enolic form [11] it may be assumed that reaction occurred with the initial formation of an unsymmetrical silyl- or germylmercury intermediate (V) containing a σ , π -conjugated bond system [12]. Further demercuration of intermediate V leads to ambident anions (VI). Subsequent recombination of the ion pairs gives III or IVa (Scheme 1).

SCHEME 1
$$(\text{Et}_3\text{M})_2\text{Hg} + \text{Hg}(\text{CH}_2\text{COPh})_2 \longrightarrow 2\text{Et}_3\text{M} - \text{Hg} - \text{CH}_2 - \text{C} - \text{Ph}$$

$$(\text{I}, \mathbb{I})$$

$$(\mathbb{V})$$

$$- 2\text{Hg}$$

$$2\text{CH}_2 = \text{C} - \text{Ph}$$

$$(\mathbb{I}, \text{M} = \text{Si}; \mathbb{V}\text{a}, \text{M} = \text{Ge})$$

$$(\mathbb{V})$$

An alternative reaction scheme which involves demercuration of intermediate V (M = Si) to lead to α -triethylsilylacetophenone, Et₃SiCH₂COPh, is less probable. According to Lutsenko, Baukov et al. [13], the above ketosilane is quite stable and undergoes rearrangement to siloxyalkene III under severe conditions (160°C). It has been established [14] that the rearrangement Me₂SiCH₂COPh \rightarrow Me₃SiOC(Ph)=CH₂ occurs via an associative mechanism which involves a five-coordinate silicon intermediate. In the case of the reaction with compound II, along with the appropriate germyl enol IVa, α -triethylgermylacetophenone (IVb) is formed in a ratio of 86 : 14. Moreover, we have found (see below) that the O-isomer IVa slowly isomerizes to the C-isomer IVb at room temperature. This reaction is probably reversible (i.e IVa \rightleftharpoons IVb). A similar germanotropic equilibrium for the reversible isomarization of trimethylgermylated acetophenone and acetone has been reported by Lutsenko and his coworkers [15,16]. In both cases the equilibrium is strongly shifted to the C-isomer.

The reaction of equimolar quantities of α -mercurybiscyclohexanone and mercurial I or II under the same conditions yields metallic mercury and 1-triethylsiloxy-1-cyclohexene (VII) or 1-triethylgermoxy-1-cyclohexene (VIII), respectively. The results obtained are in agreement with the mechanism depicted in Scheme 1. The mechanism of the formation of VII and VIII is thought to involve initial formation of the ambident anion. This anion offers two potential sites for association with the counter-ion (e.g. Et_3Si^+). However, no formation of C-silylated or C-germylated isomers could be detected, which demonstrated the selectivity of this reaction.

Several methods are available for the synthesis of compound VII. One of them is based on the reaction of chlorotriethylsilane with an ambident enolate anion of cyclohexanone [17].

It is known [18] that mercury bisacetoacetic ester has the mercury atom bonded to the central carbon atoms of both ethyl acetoacetate groupings. However, the exchange reactions of this compound with I and II in THF lead to the ethyl ester of β -triethylsiloxycrotonic acid (IX) and its germanium analogue (X), respectively. The formation of the observed products could be rationalized by the mechanism shown in eq. 3, including decomposition of unsymmetrical mercurials to a silyl (or germyl) cation and ambident anion (XI), followed by simultaneous association of the counter-ions (cf. Scheme 1).

$$Et_{3}M \xrightarrow{H} G \xrightarrow{CH} C - Me \xrightarrow{-Hg} \left[Et_{3}M^{+} \quad Me - C \xrightarrow{CH} - C - OEt\right] \xrightarrow{OMEt_{3}} MeC = CHCOOEt \quad (3)$$

$$COOEt \qquad (XI) \qquad (IX, M = Si; X, M = Ge)$$

It should be noted that the O-silylated product IX was previously prepared by the reaction of chlorotriethylsilane with sodium acetoacetic ester [19]. In this reaction the ambident anion XI derived from proton abstraction also offers three potential sites for electrophilic attack. The structure and reactivity of enolate anion XI has been discussed previously [20,21].

According to ¹H NMR data (Table 2) products IX and X are mixtures of *cis* and *trans* isomers. For IX, the assignment of signals to *cis* and *trans* isomers is

carried out by comparison of the measured chemical shifts of the vinyl proton with the value calculated using an additive scheme [22]. The assignment corresponds to that [23] for $R_3Si^*OC(Me)=CHCO_2Et$ where $R_3Si^*=Me(1-C_{10}H_7)$ -PhSi. For cis and trans isomers of X the order of assignment of $CH_3C=C$ and C=CH proton signals is in agreement with and argues against the order found for product IX, respectively. With association of the counter-ions (cf. eq. 3) one cannot but take into account the alternative structure $MeC(O)CH=C(OEt)-OSiEt_3$ (XII). However, this structure may be excluded from consideration since the calculated vinyl proton shifts for trans and cis isomer of XII (4.25 and 4.08 ppm, respectively) differ by only 0.9 ppm from the measured values.

As may be seen from the data presented in Table 1, α -bromocarbonyl compounds derived from enolizable ketones and aldehydes react with mercurials I or II, giving the corresponding silyl or germyl enol ethers in high yields [24,25]. For example, α -bromoacetophenone reacts with I under mild conditions to give enol ether III along with metallic mercury and bromotriethylsilane. However, a similar reaction of α -bromoacetophenone with mercurial II affords both O- and C-germylated products, IVa and IVb, in a ratio of 20:80. The fact that the mixture predominantly contains the thermodynamically more stable isomer IVb is not surprising. This may be explained by the presence in the reaction mixture of bromotriethylgermane which catalyzes the process of reversible isomerization IVa \rightleftharpoons IVb (cf. [15,16]).

Thus, the results (Table 1) may be best explained by Scheme 2.

SCHEME 2
$$(Et_3M)_2Hg + BrCH_2COR \xrightarrow{(a)} Et_3MBr \qquad Et_3M-Hg-CH_2-C-R$$

$$(I,II)$$

$$(b) -Hg$$

$$Et_3MCH_2COR \xrightarrow{CH_2=C-R} (C) \left[Et_3M^+ CH_2^-C-R\right]$$

$$(IV b,XIV b) \qquad (II,M=Si,R=C_6H_5; IV a,b,M=Ge,R=C_6H_5; IV a,b,M=Ge,R=P-BrC_6H_4; IV a,b,M=Ge,R=P-BrC_6H_4; IV a,b,M=Ge,R=C_6F_5; IV a,b,M=Ge,R=C_6F_5; IV a,B=Ge,R=C_6F_5; IV a,B=Ge,R=C_6F_6; IV a,B=Ge,R=C_6F_6; IV a,B=Ge,R=C_6F_6F_6; IV a,B=Ge,R=C_6F_6; IV a,B=Ge,R=C_6F_6; IV a,B=Ge,R=C_6F_6F_6; IV a,B=Ge,R=C_6F_6; IV a,B=Ge,R=C_6F_6F_6; IV a,B=Ge,R=C_6F_6F_6; IV a,B=Ge,R=C_6F_6F_6; IV a,B=Ge,R=C_6F_6F_6F_6F_6F_6F_6F_$$

The mechanism of the formation of silyl and germyl enol ethers is thought to involve initial formation of unsymmetrical silyl- or germylmercurials via a four-centre exchange reaction [step a]. This step is similar to that postulated for thermal reaction of (Me₃Si)₂Hg with arylhalides [26,27], pentafluorobromo-

ANALYTICAL AND OTHER DATA FOR SILYL AND GERMYL ENOL ETHERS AND RELATED COMPOUNDS TABLE 1

Compound	Method h	Yield	B.p. (°C)	п _D	Analysis, fou	Analysis, found (calcd.) (%)	
		(%)	(mmHg)		υ	н	Sl/Ge
$\operatorname{Et}_3\operatorname{SiOC}(\operatorname{Ph})=\operatorname{CH}_2^d$ (III)	н с	67	87-89 (2)	1,5065			
Et3GeOC(Ph)=CH2 (IVa)	N -	83 79	81-83(1) $105-106(1-2)$	1.6075 1.5270	60,02	7.93	25.55
Et3GeCH2COPh (IVb)					(60.28)	(4.96)	(26,02)
E1350	-	74	97-99 (12)	1.4630	67,81	10,96	12,92
					(09'/0)	(11.39)	(13,22)
Et ₃ GeO一(文語)	1	72	70-71 (1)	1.4770	55,96	9,42	27.97
)					(56.10)	(9.41)	(28.25)
$Et_3SIOC(Me) = CHCO_2Et(IX)$	7	83	76-78 (1.5)	1,4580	58,88	66'6	11,20
					(58.97)	(06'6)	(11,48)
$Et_3GeOC(Me)=CHCO_2Et(X)$, ,	77	84-85 (1.5)	1.4760	50.18	8,33	25.20
	,	i			(49.89)	(8,37)	(25,13)
$Et_3SIOC(C_6H_4br_p)=CH_2$ (XIII)	23	90	$92-94 (2.8 \times 10^{-2})$	1,5315	53.94	6.49	8.90
7-1119/ HOLY "d' 11 0/0000 fd					(53.67)	(6.76)	(8.96)
Et-George (AIV)	62	77	$107-108 (2.1 \times 10^{-2})$	1,5528	46.65	5,89	20.22
Ergenerizon Gridary (Arty E)					(46.99)	(6.92)	(20.29)
$Et_3SIOC(C_6F_5)=CH_2$ (XV)	2	. 98	71-72 (1)	1,5498	51.88	5.23	8.45
	п			1.4480	(51.84)	(5.28)	(8,66)
$Et_3GeOC(C_6F_5)=CH_2(XVI)$	6 1 ,	89	80-82 (1)	1,4659	45.86	4.50	19.46
}	-			1.4681	(45.59)	(4.64)	(19.68)
	8	96	$60-62 (4 \times 10^{-2})$	1,4688	72.23	11.30	10.68
OSIEt ₃ (XVIII)					(72.11)	(11.35)	(10.54)
7							
>							
	c	S	61000	0		,	
OGELT ₃ (XVIII)	N	N D	$81-83 (1.1 \times 10^{-2})$	1.4852	61.29 (61.79)	9.60 (9.72)	23,44 (23,34)

TABLE 1 (continued)

15,23 (15,06) 30,31 (31,44) 9.26 (9.58) 21.21 (21.49) 11.07 (11.18) 24.71 (24.54) 10.97 (11.30) 25.71 (24.78) Si/Ge Analysis, found (calcd.) (%) 7.40 (7.46) 21.11 (11.90) 9.98 (9.60) 6.36 8,45 (8.59) 7.78 (7.62) I 65.01 (64.45) 52.52 (52.02) 49.19 (49.14) 42.94 (42.46) 43.43 (43.03) 35,99 (36,55) 72.43 (72.52) 61.49 (61.50) ပ 1,4358 1,4918 1,4920 1,4770 1,4731 1.6180 1,4551 1,5380 8,5 $62-64 (2.5 \times 10^{-2})$ $81-83 (2.5 \times 10^{-2})$ 48-50 (1) 61 - 52(1)44-45 (1) 48-49 (1) B.p. (°C) (mmHg) Yield (%) 95 c 9 9 G 22 69 93 79 84 38 Method h c) c4 Br (E) (XXIV) Br (E) (XXV) H(Z) (XIX) Ph Me Ph(E) (XXI)
Et₃SiOCH=CMe₂ h (XXII) H(Z) (XX) Et3GeOCH=CMe2 (XXIII) C=CH₂ (XXVII) C=CH₂ (XXVI) Compound Et3Ge0 Et 3GeO Me₃C Et3GeO, Me3C Et 3SiO Et 3SIO BrCH₂ Et₃SiO

10,64 (10,59)		22,97 (23,43)				
8.15 (7.98)		6.79 (6.83)				
45.16 (45.28)		38.71 (38.77)				
1,4810		1,4970		1,4674	1,4617	1,4609
47—48 (1)		50—52 (1)		93—94 (1) 102—104 (1)	108—109 (1)	106–107 (1)
80		89		78 84	67	70
Ø		લ		લ લ	લ	82
E_{t_3SlO} $C=C$ B_rCH_2 E_{t_3SlO}	C=CH ₂ (XXVIII) Mechbr Et3GeO H	BrCH ₂ Me (E) (XXXI)	Et ₃ GeO C=CH ₂ (XXIX)	MeChbr' Et3GeCH(CN)CO2Et ^d (XXXII) Et3GeCH(CO2Et)2 ^c (XXXIII) Et3GO	$c = cHCN^f(XXXIV)$	Et ₃ SiO $C = CHCO_2Et^g(XXXV)$ EtO

^a Lit. [13]: b.p. 86–88°C/1.5: $^{10}_{D}$ 1.5062. ^b Lit. [40a]; b.p. 89°C/25 mm; $^{10}_{D}$ 1.4337. ^c GLC was used in determining yields and isolating pure samples for analysis and spectroscopy. ^d Lit. [37]: b.p. 94–95°C/1 mm; $^{10}_{D}$ 1.4650. ^e Lit. [38]; b.p. 154°C/14 mm; $^{10}_{D}$ 1.4570. ^f Lit. [37]: b.p. 110°C/1 mm; $^{10}_{D}$ 1.4600. ^g Lit. [37]: b.p. 108°C/1—2 mm; $^{10}_{D}$ 1.4590. ^h 1, reaction with $^{10}_{C}$ reaction with $^{10}_{C}$ carbonyl compounds.

TABLE 2 SOME IR AND ^1H NMR DATA FOR SILYL AND GERMYL ENOL ETHERS AND RELATED COMPOUNDS

No	$\frac{IR (cm^{-1})}{C=C}$	¹ H NMR, δ (ppm) J (Hz)						
	(C=O)	=CH cis a	=CH trans a	$=C-CH_n$ n=1, 2, 3	CH ₂	—CH ₃	⁽ⁿ⁾ J(HH)	
III	1617	4.72 d	4.27 d				1.8(2)	
IVa l	1628	4.58 d	4.01 d				1.8(2)	
IVb ^ʃ	(1662)			2.64 s				
VII	1667	4.73 t		1.96 m	1.55 m		2.8(3)	
VIII	1660	4.42 t		1.89 m	1.53 m		2.8 (3)	
IX	1625	5.00 s	4.94 s	2.22 s 1.85 s	4.01 k	1.22 t	7.5(3)	
X	1602	4.73 s	4.89 s	2.20 s 1.79 s	3.98 k	1.29 t	7.5(3)	
XIII	1615	4.76 d	4.34 d				1.8(2)	
XIVa l	1627	4.59 d	4.06 d				1.8(2)	
XIVb J	(1659)			2.59 s				
xv	1630	4.68 d	4.38 d, t				2.0(2)	
XVI	1610	4.47 d	4.27 d, t				2.0(2)	
XVII	1620	4.53 d		2.14 m	1.79 m 1.45 m	0.86 c 0.65 c	3.5(3)	
XVIII	1610	4.27 d		2.09 m	1.77 m 1.42 m	0.84 c 0.69 c	3.4(3)	
XIX	1657		5.29 k	1.91 d			0.9(4)	
XX l	1640		5.23 k	1.86 d			0.9(4)	
XXI J	1640	5.47 k		1.96 d			0.8(4)	
XXII	1685	5.95 m		1.55 d			1.0(4)	
				1.51 d			1.2(4)	
XXIII	1674	6.00 m		1.47 d			1.2(4)	
				1.52 d			1.0(4)	
XXIV	1618	5.52 s				1.09 s		
XXV	1587 1602	5.15 s				1.06 s		
XXVI	1629	4.35 d	4.17 m	3.74 m			1.3(2) 0.6(4)	
XXVII	1610	4.13 d	3.89 d, t	3.65 d			1.3(2) 0.6(4)	
	1638		•					
XXX 1	1648	4.52 q		3.60 q		1.43 d, t	6.6(3) 0.7(5)	
XXVIII	Ī	4.15 d	4.02 q	4.28 q		1.57 d	1.5(2) 6.6(3) 0.5(4)	
XXXI	1650 1606	4.70 q	•	3.68 q		1.47 d, t	6.8(3) 0.7(5)	
xxix)		4.12 d	3.82 q	4.42 q		1.68 d	1.5(2) 6.8(3) 0.5(4)	

^a The cis and trans position of the H atom relative to OMEt₃ is related to the E- and Z-forms, respectively.

benzene and bromotrifluoroethylene [28]. Other steps of Scheme 2 have been discussed above.

In these reactions, the reactivity of α -bromoacetophenones is enhanced by a decrease in the electron-donating effect of the aromatic ring in the order: $BrCH_2COC_6H_5 < BrCH_2COC_6H_4Br-p < BrCH_2COC_6F_5$. Thus, with $BrCH_2COC_6H_5$ the reaction proceeds under more severe conditions (100°C) than those required for $BrCH_2COC_6F_5$ which reacts with mercurials I and II in hexane at -78° C.

We have prepared the product IVa by a photochemical reaction of α -bromo-acetophenone with II in hexane at 30°C. Under these conditions the reaction

is complete within ca. 15 min to yield a pure O-isomer. It has been shown that the above isomer, when kept in an evacuated ampoule (2 months at ca 20°C) isomerizes to product IVb. The overall yield (without isolation of IVa) is 20%.

Since the difficulty synthesized silyl enol ether of camphor was formed in good yield under some of the reaction conditions [29–34], 3-bromocamphor was treated with mercurials I and II in benzene to demonstrate the synthetic value of this reaction. The results are shown in Table 1. The yields of triethylsilyl enol ether (XVII) and its germanium analogue (XVIII) represent isolated yields of distilled material. A similar reaction with 1-phenyl-1-bromopropan-2-one proceeds under mild conditions in a highly stereoselective fashion, leading predominantly to the Z-isomer.

$$(Et_{3}M)_{2}Hg + MeCCHBrPh \xrightarrow{ca\ 20^{\circ}C} \xrightarrow{H} + Me \\ \xrightarrow{Et_{3}MO} C = C \xrightarrow{H} + Et_{3}MO C = C \xrightarrow{Ph}$$

$$(XIX, M = Si; \quad (XXI, M = Ge)$$

$$XX, M = Ge)$$

$$(4)$$

A special experiment has shown that the presence of the E-isomer (XXI) in the reaction mixture (isomer ratio is 90 : 10 from the ¹H NMR data) cannot be explained in terms of an isomerization XX \rightarrow XXI. The reaction of 2-methyl-2-bromopropanal with I and II also affords silyl (XXII) and gernyl (XXIII) enolethers, respectively.

Finally, in order to demonstrate the synthetic potential of these new reagents, we have used the method described above to effect a simple synthesis of bromo(triethylsiloxy)alkenes (i.e., triethylsilyl bromoenol ethers) and their germanium analogues. Two methods for the synthesis of such silyl enol ethers are known.

A recent report [35] describes that *vic*-bromo(trimethylsiloxy)alkenes can be prepared regioselectively from the appropriate silyl enol ethers via bromination at low temperature followed by dehydrobromination of the 1,2-dibromoalkyl trimethylsilyl ether obtained. Another possible pathway involves treatment of 2-methyl-2-bromobutan-3-one with lithium diisopropylamide and subsequent reaction of the lithiated product with Me₃SiCl [36].

We have examined the reaction of mercurials I and II with α,α - and α,α' -dibromo carbonyl compounds. For example, of great interest was the success of such a reaction with Br₂CHCOt-Bu which did indeed give the expected (E)-3,3-dimethyl-2-triethylsiloxy-1-bromo-1-butene (XXIV) and its germanium analogue (XXV), respectively.

$$(Et_{3}M)_{2}Hg + Br_{2}CHCt-Bu \xrightarrow{-Hg,-Et_{3}MBr} Et_{3}MO C=C \xrightarrow{Br} (E)$$

$$(XXIV, M = Si; XXV, M = Ge)$$
(5)

With α,α' -dibromo ketones, enol ethers with the bromine atom bound to the

 sp^3 carbon atom are formed.

$$RCHBrCCH_{2}Br + (Et_{3}M)_{2}Hg \xrightarrow{-Hg, -Et_{3}MBr} RCH_{2}Br + (Et_{3}M)_{2}Hg \xrightarrow{-Hg, -Et_{3}MBr} RCH_{3} +$$

We have also studied the reactions of germylmercurial II with bromocyanoacetic and bromomalonic esters. These reactions were carried out in THF at room temperature. These conditions lead to the formation of the known C-germylated products (XXXII and XXXIII) [cf. 37.38]. Only traces of the corresponding O-germylated isomers were detected in the reaction mixtures.

$$(Et_{3}Ge)_{2}Hg + BrCH(R)COOEt \xrightarrow{-Hg, -Et_{3}MBr} Et_{3}GeCH(R)COOEt$$

$$(XXXII, R = CN;$$

$$XXXIII, R = COOEt)$$
(7)

In contrast, the reactions of the same bromoesters with silylmercurial I in THF afford predominantly, if not exclusively, O-silylated products. The yields of 2-ethoxy-2-triethylsiloxy-2-propenenitrile (XXXIV) and the ethyl ester of the same acid (XXXV) are 57 and 70%, respectively. Products XXXIV and XXXV were identified by comparison of IR and NMR spectra with previously described [37] samples.

Experimental

All reactions were carried out in evacuated sealed ampoules following the technique described in ref. 39. The IR spectra were recorded on a UR-20 spectrometer. NMR spectra were obtained on a Tesla BS 487C (80 MHz) instrument with hexamethyldisiloxane as internal standard. Typical experiments are given below.

3-Triethylgermoxy-1-cyclohexene (VIII)

A mixture of 3.8 g (7.3 mmol) II, 2.90 g (7.3 mmol) of Hg[CHCO(CH₂)₃CH₂]₂ and 20 ml of THF was heated in an ampoule for 6 h at 65°C, the characteristic yellow colour of the germylmercirial having disappeared. Metallic mercury (2.93 g, ca. 100%) was precipitated. Distillation of the organic layer gave 2.70 g (72%) of compound VIII (see Table 1).

Photochemical reaction of the mercurial II with α -bromoacetophenone A mixture of 3.46 g (17.4 mmol) of α -bromoacetophenone, 9.10 g (17.4

mmol) of II and 30 ml of hexane was irradiated in a molydenic glass ampoule at -30° C for 20 min. (DPT-375 lamp; 10 cm distance from the light source). The mixture was decanted from mercury (3.44 g, 99%) and the hexane and volatile products removed by recondensation in vacuo. Distillation of the residue gave 2.74 g (63%) of 1-phenyl-1-triethylgermoxyethylene (IVa) b.p. $103-105^{\circ}$ C/1 mm Hg, n_D^{20} 1.5260. (Found: C, 60.01; H, 7.91; Ge, 25.53. $C_{14}H_{22}$ GeO calcd.: C, 60.29; H, 7.95; Ge, 26.02%). IR (cm⁻¹): ν (C=C) 1628. NMR (CCl₄) (δ , ppm): 4.59 (d, 1 H, =CH₂), 4.05 (d, 1 H, =CH₂). No C-isomer. IVb was detected in the reaction mixture. The volatiles were analyzed by GLC. Bromotriethylgermane (88%) was identified.

1-Pentafluorophenyl-1-triethylgermoxyethylene (XVI)

A solution of 2.81 g (10.4 mmol) of BrCH₂COC₆F₂ in 15 ml of hexane was degassed, frozen with liquid nitrogen and 5.61 g (10.8 mmol) of II was added. During the course of subsequent defrosting mercury was eliminated (2.04 g, 98%). The reaction was completed after 3–5 min. The organic layer was fractionated to obtain the desired product (2.71 g, 68%), b.p. 64–66° C/1 mm Hg, n_D^{20} 1.4658. (Found: C, 45.86; F, 25.71; Ge, 19.46. C₁₄H₁₇F₅GeO calcd.: C, 45.59; H, 4.64; F, 25.74; Ge, 19.68%). IR (cm⁻¹): ν (C₆F₅) 1495, 1520, 1650, ν (C=C) 1610. NMR (CCl₄) (δ , ppm): 4.47 (q, 1 H, =CH₂), 4.27 (d, t, 1 H, =CH₂, 2 J(HH) 2.0 Hz).

(E)-3,3-Dimethyl-2-triethylgermoxy-1-bromo-1-butene (XXV)

Bis(triethylgermyl)mercury (4.65 g, 8.9 mmol) was added to a solution of 2.31 g (8.9 mmol) of BrCHCOt-Bu in 7 ml of benzene. The exothermal reaction was complete after 10–15 min when the organic layer was decanted from the metallic mercury (1.71 g, 96%). Distillation of the organic layer gave 2.81 g (93%) of XXV, b.p. $51-52^{\circ}$ C/1 mm Hg, n_D^{20} 1.4918. (Found: C, 42.91; H, 7.40; Br, 23.59; Ge, 21.21. $C_{12}H_{25}$ BrGeO calcd.: C, 42.66; H. 7.46; Br, 23.65; Ge, 21.47%). (see Table 2). The volatiles were analyzed by GLC. Bromotriethylgermane (83%) was identified.

References

- 1 N.S. Vyazankin, G.A. Razuvaev and O.A. Kruglaya, Organometal. Chem. Syn., 1 (1971) 205.
- 2 W.P. Neumann and K. Reuter, J. Organometal. Chem. Libr., 7 (1979) 229.
- 3 O.A. Kruglaya, G.S. Kalinina, B.I. Petrov and N.S. Vyazankin, J. Organometal. Chem., 46 (1972) 51.
- 4 B.V. Fedot'ev, O.A. Kruglaya and N.S. Vyazankin. Izv. Akad. Nauk SSSR, Ser. Khim., (1974) 713.
- O.A. Kruglaya, I.B. Fedot'eva, B.V. Fedot'ev, I.D. Kalikhman, E.I. Brodskaya and N.S. Vyazankin, J. Organometal. Chem., 142 (1977) 155.
- 6 O.A. Kruglaya, I.B. Fedot'eva, B.V. Fedot'ev, N.S. Vyazankin and I.D. Kalikhman, Izv. Akad. Nauk SSSR, Ser. Khim., (1975) 2790.
- 7 B.I. Petrov, O.A. Kruglaya, N.S. Vyazankin, B.I. Martynov, S.R. Sterlin and B.L. Dyatkin, J. Organometal. Chem., 34 (1972) 299.
- (a) R.E. Dessy, F. Kaplan, G.R. Coe and R.M. Salinger, J. Amer. Chem. Soc., 85 (1963) 1191.
 (b) T.F. Schaaf and J.P. Oliver, J. Organometal. Chem., 32 (1971) 307.
- 9 V.M. Vodolazskaya, B.V. Fedot'ev, Yu.I. Baukov, O.A. Kruglaya and N.S. Vyazankin, J. Organometal. Chem., 63 (1973) C5.
- 10 (a) A.C. Lee, J. Organometal. Chem., 16 (1969) 321. (b) D.C. McWilliam and P.R. Wells, J. Organometal. Chem., 85 (1975) 335. (c) D.C. McWilliam and P.R. Wells, J. Organometal. Chem., 85 (1975) 347. (d) I.B. Fedot'eva, O.A. Kruglaya, I.D. Kalikhman and N.S. Vyazankin, Izv. Akad. Nauk SSSR. Ser. Khim., (1979) 2365.

- 11 I.I. Kritskaya, Izv. Akad. Nauk SSSR, Ser. Khim., (1979) 1946.
- 12 A.N. Nesmeyanov, V.T. Aleksanyan, L.I. Denisovich, Yu.S. Nekrasov, E.I. Fedin, V.I. Khvostenko and I.I. Kritskaya, J. Organometal. Chem., 172 (1979) 133.
- 13 I.F. Lutsenko, Yu.I. Baukov, O.V. Dudukina, and E.P. Kramarova, J. Organometal. Chem., 11 (1968) 35.
- 14 H. Kwart, and W.E. Barnette, J. Amer. Chem. Soc., 99 (1977) 614.
- 15 A.N. Tvorogov, L.V. Goncharenko, I.Yu. Belavin, Yu.I. Baukov and I.F. Lutsenko, Zh. Obshch. Khim. 43 (1973) 441.
- 16 I.F. Lutsenko, Yu.I. Baukov, I.Yu. Belavin and A.N. Tvorogov, J. Organometal. Chem., 14 (1968) 229.
- 17 P.F. Hudrlik, J.M. Takacs, J. Org. Chem., 43 (1978) 3861; R. Bourhis and E. Frainnet, Bull. Soc. Chim. France, (1967) 3552; K. Rühlmann, A. Sitzki and C. Michael, Chem. Ber., 101 (1968) 285.
- 18 C. Glidewell, J. Organometal. Chem., 136 (1977) 7.
- 19 H. Gilman and R.N. Clark, J. Amer. Chem. Soc., 69 (1947) 967.
- 20 J.M. Lefour, P. Sarthoun, G. Bram, A. Loupy and J. Seyden-Penne, Tetrahedron Lett., (1978) 3831.
- 21 C. Cambillau, G. Bram, J. Corset, C. Riche and C. Pascard-Billy, Tetrahedron, 34 (1978) 2675.
- 22 U.E. Matter, C. Pascual, E. Pretsch, A. Pross, W. Simon and S. Sternhell, Tetrahedron, 25 (1969) 691.
- 23 I.K. Kusnezova, K. Rühlmann and K. Gründemann, J. Organometal. Chem., 47 (1973) 53.
- 24 D.V. Gendin, O.A. Kruglaya, I.D. Kalikhman and N.S. Vyazankin, Zh. Obshch. Khim., 49 (1979) 951.
- 25 D.V. Gendin, O.A. Kruglaya, I.D. Kalikhman, N.S. Vyazankin and G.G. Furin, Izv. Akad. Nauk SSSR, Ser. Khim., (1979) 2370.
- 26 C. Eaborn, R.A. Jackson and R.W. Walsingham, J. Chem. Soc. C, (1967) 2188.
- 27 R. Fields, R.N. Haszeldine and R.E. Hutton, J. Chem. Soc. C, (1967) 2559.
- 28 R. Fields, R.N. Haszeldine and P.J. Palmer, Tetrahedron Lett., (1971) 1879.
- 29 J.K. Rasmussen, Synthesis, (1977) 91.
- 30 J. Dedier, P. Gerval and E. Frainnet, J. Organometal. Chem., 185 (1980) 183.
- 31 G.C. Joshi and L.M. Pande, Synthesis, (1975) 450.
- 32 G. Simchen and W. Kober, Synthesis, (1976) 259.
- 33 R.D. Clark and C.H. Heathcock, J. Org. Chem., 41 (1976) 1396.
- 34 C.W. Jefford and C.G. Rimbault, J. Amer. Chem. Soc., 100 (1978) 6437.
- 35 M. Zembayashi, K. Tamao and M. Kumada, Synthesis, (1977) 422.
- 36 H. Sakurai, A. Schirahata and A. Hosomi, Angew. Chem., 91 (1979) 178.
- 37 L.I. Belousova, O.A. Kruglaya, V.V. Neretin, I.D. Kalikhman, N.A. Ivanova, A.A. Tatarinova and N.S. Vyazankin, Izv. Akad. Nauk SSSR, Ser. Khim., (1977) 2070.
- 38 J. Sadge and M. Rivière-Baudet, Recl. Trav. Chim. Pays-Bas, 94 (1975) 22.
- 39 N.S. Vyazankin, G.A. Razuvaev, O.A. Kruglaya and G.S. Semchikova, J. Organometal. Chem., 6 (1966) 474.
- 40 E. Frainnet and R. Bourhis, J. Organometal. Chem., 93 (1975) 309.