0-Trialkyltin Hydroxylamines

of Grummit, Budewitz, and Chudd.³³ To 6.0 g (260 mg-atom) of magnesium powder (40 mesh) in ca. 125 ml of ether, freshly distilled from lithium aluminum hydride, in a flame-dried 300-ml roundbottomed flask equipped with magnetic stirrer, condenser, and No-Air stopper, was added at 0" ca. **0.5** ml of 1,2-dibromoethane. To the activated magnesium, 7.0 g (58 mmol) of freshly distilled allyl bromide was added at 0° over a 2-hr period. The solution was allowed to stir at 0" for an additional 5 **hr** and was transferred by **cannula** to a flamedried 200-ml storage bottle. The residual salts were washed with ca. 20 ml of ether, and the washings transferred to the storage bottle and mixed thoroughly. Titration with 0.105 *M* sec-butyl alcohol in xylene using $1,10$ -phenanthroline³⁴ as indicator showed the solution to be 0.36 *M.*

Tetra(trihaptoallyl)zirconium(N) *(5)* was prepared according to the procedure of Becconsall, Job, and O'Brien.' In a 100-ml roundbottomed flask was placed 0.79 g (3.9 mmol) of zirconium tetrachloride. To the salt, cooled to -60° , 40 ml of $0.30 M$ (12.0 mmol) allylmagnesium bromide was added over a 1.5 hr period. The mixture was stirred for 20 hr at external temperature of ca. **-70"** (Dry Ice-isopropyl alcohol) and filtered at $ca. -40^{\circ}$ through a Celite pad, which previously had been dried with $ca. 2$ ml of *n*-butyllithium. The solvent was removed under vacuum while maintaining the sample temperature between -40 and -30° .

methane, previously degassed by three freeze-thaw cycles, was distilled into the storage tube containing the red solid *5,* obtained by removal of ether under vacuum at **-78"** (Dry Ice-isopropyl alcohol). An aliquot of the resulting solution was transferred at -78° through a stainless steel cannula into an nmr tube, adapted with a 12/15 ball Preparation **of Nmr** Samples **of** *5. Cu.* 5 ml of fluorotrichloro-

Syntheses,"Collect. Vol. IV, Wiley, New York, **N. Y., 1963,** p **748. (33) 0.** Grummit, E. **P.** Budewitz, and **C.** C. Chudd, "Organic

165 (1967). (34) S. C. Watson and **J.** F. Eastham, *J.* Organometal. Chem., *9,* joint and joined to a vacuum stopcock, fitted with a **No-Air** stopper. TMS internal standard was added and the tube sealed under vacuum. The sample used showed no absorption in the nmr spectrum due to ether, but did show a broad impurity peak centered slightly upfield from the H₁ resonances of 5. The nature of this impurity was not explored. Extensive decomposition was observed if the sample was not stored below -20°

Trihaptoallylpalladium chloride dimer was prepared by Dr. D. L. Tibbets. The nmx sample used in this study was prepared by adding 75 mg (0.206 mmol) of the palladium complex and 64.5 mg (0.826 mmol) of dimethyl sulfoxide d_6 (2 equiv of DMSO d_6 /allyl group) to 0.5 ml of a 10% solution of benzene in deuteriochloroform.

Calculations were performed at the Massachusetts Institute of Technology Information Processing Center using the program EXCHSYS^{15,35} to compute eigenfunctions and line shapes of static spectra, K matrices, and interchange-broadened line shapes.

Registry No. Tetra(trihaptoallyl)zirconium(IV), 12090-34-5; chloro(trihaptoallyl)palladium(II) dimer, 12012-95-2.

Acknowledgment. The cooperation of D. L. Tibbets in obtaining the nmr spectra is appreciated. C. B. Powell, Jr., and J. **B.** Lisle assisted with the computer programming. Our colleague John Waugh corrected a number of our egregious misconceptions concerning the physical basis of mag netic resonance phenomena. Friedmanning.

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Bothner-By and S. M. Castellano, in D. **F.** deTar, "Computer **Pro-**grams for Chemistry," Vol. **I,** w. A. Benjamin, **New** York, N. **Y., 1968,** and local line shape programs, KMATRX and EXCNMR, written by J. B. Lisle, C. **P. Powell, J. K.** Krieger, and G. M. Whitesides. **(35)** EXCHSYS is based on the program **LAOCN3: A. A,**

Contribution from the Department of Chemistry, University of Nottingham, University Park, Nottingham NG7 2RD, England

Structural Studies in Main-Group Chemistry. IV.' 0-Trialkyltin Hydroxylamines2

PHILIP *G.* HARRISON

Received November *15, 1972*

The organotin hydroxylamine derivatives $Me_3SnONEt_2$, $R_3SnONHCOPh$ ($R = Me$, *n*-Pr), and $R_3SnONPhCOPh$ ($R = Me$, *n*-Pr, Ph) and the organosilicon and organolead analogs $Me_3SiONPhCOPh$ and Ph₃PbONPhCOPh have been synthesized. are inert to hydrolysis save $Me₃SDONEt₁$, which readily reverts to its protic precursors in air, and monomeric in solution (osmometry). Attempts to prepare Ph,SnONHCOPh from triphenyltin hydroxide result in the formation of tetraphenyltin

in high yield, and from triphenyltin chloride and excess triethylamine the ionic species $(NEt₃H⁺)(Ph₃SnONCPhO$ is produced. The trimethyltin homolog of the latter is prepared by dissolution of Me,SnONHCOPh in triethylamine. The structures of the derivatives in both solution and solid phases are discussed in terms of their infrared, nuclear magnetic resonance, tin-1 19m Mossbauer, and mass spectra. The stability of the tin and lead derivatives is due to the increase of the coordination number at the metal atom via intramolecular coordination of the carbonyl group to the metal. The mass spectra for Me,SnONHCOPh and Me,SnONPhCOPh indicate the presence of associated, most probably weakly associated dimer, species in the solid. High-resolution mass measurements have confirmed the presence of the SnOSn backbone in the more abundant ditin fragments and also the composition of monotin fragments, among which are some which arise from R_3 SnONPhCOPh $(R = Me, n$ -
h have been synthesized. All
, and monomeric in solution
the formation of tetraphenylti
 $H^*(Ph_3$ SnONCPhO⁻
HCOPh in triethylamine. The
infrared, nuclear magnetic res-
is due to the increase of th

novel ring contraction reactions involving the elimination of oxygen or phenylnitrene from MONPhCPh=O heterocyclic species. The major fragmentation processes of Me,SiONPhCOPh have been corroborated by the observation of metastable ions and involve mainly the formation of phenyl-containing ions in which the charge may be effectively delocalized.

Introduction

metalloid derivatives containing the M-0-N linkage continues to arouse much interest.^{3,4} Oxime derivatives of lithium.³ The synthesis and properties of main-group metal and

(1) Part **111: P.** G. Harrison and **J. J.** Zuckerman, *J.* Orgunomefal. Chem., **in** press.

(2) A preliminary communication has appeared: **P.** G. Harrison, *J.* Organometal. Chem., **38,** C5 **(1972).**

(3) P. G. Harrison and **J. J.** Zuckerman, Inorg. Chem., **9, 175 (1970),** and references therein. **(4) P. J.** Harrison and **J. J.** Zuckerman, Inorg. *Nucl.* Chem. Lett.,

6, 5 (1970), and references therein.

silicon, germanium, tin, $3,4$ lead, arsenic, and antimony⁴ have previously been synthesized by us, while attempts to prepare similar phosphorus derivatives led to the formation of phosphorylamines *via* an Arbuzov rearrangement.4 The *0-* **(trialkylstannyl)oximes3** are volatile monomeric liquids except for cyclohexanone **0-(trimethylstannyl)oxime,** for which infrared, tin-I 19m Mossbauer, and high-resolution mass spectral data indicated association *via* distannoxane

SnOSnO ring formation in the solid. Such association is however rather weak since only monomeric species could be

detected in benzene solution, unlike the strongly associated 1,3-disubstituted distannoxanes where dimeric units are preserved on dissolution.⁵ Because of their different electronic and steric requirements, we have found it of interest to synthesize organotin derivatives of substituted hydroxylamines, some initial investigations of which are reported in this paper.

Results **and** Discussion

The organotin derivatives of N,N-diethyl-, N-benzoyl-, and **N-phenyl-N-benzoylhydroxylamines** were prepared by the azeotropic removal of water from the appropriate hydroxylamine and the organotin oxide or hydroxide

$$
{}^{1}/{}_{2}(\mathrm{R}_{3}\mathrm{Sn})_{2}\mathrm{O}
$$
\n
$$
\mathrm{R}_{3}\mathrm{SnOH}
$$
\n
$$
{}^{1}/{}_{2}(\mathrm{R}_{3}\mathrm{Sn})_{2}\mathrm{O}
$$
\n
$$
{}^{1}/{}_{3}\mathrm{SnOH}
$$
\n
$$
{}^{1}/{}_{3}\mathrm{R}_{3}\mathrm{O}(\mathrm{R}^{1})_{1}\mathrm{R}^{2}
$$
\n
$$
{}^{1}/{}_{3}\mathrm{R}_{3}\mathrm{O}(\mathrm{R}^{1})_{1}\mathrm{R}^{2} = \mathrm{Et}
$$
\n
$$
{}^{1}/{}_{3}\mathrm{R}_{3}\mathrm{O}(\mathrm{R}^{1})_{1}\mathrm{R}^{2} = \mathrm{Et}
$$
\n
$$
{}^{1}/{}_{3}\mathrm{R}_{3}\mathrm{O}(\mathrm{R}^{1})_{1}\mathrm{R}^{2} = \mathrm{L}\mathrm{O}(\mathrm{R})_{1}\mathrm{R}^{2} = \mathrm{C}\mathrm{O}(\mathrm{R})_{1}\mathrm{R}^{2} = \mathrm{C}\mathrm{O}(\mathrm{R
$$

0-(Triphenyllead)-N-phenyl-N-benzoylhydroxylamine (11) was similarly synthesized from triphenyllead hydroxide

$$
Ph3PbOH + HONPhCOPh \xrightarrow{-H2O} Ph3PbONPhCOPh
$$
II (2)

However attempts to prepare $Ph_3SnONHCOPh$ by the same method only resulted in the formation of tetraphenyltin in high yield, presumably by a disproportionation reaction,

although no pure diphenyltin derivative could be isolated
\nPh₃SnOH + HONHCOPh
$$
\xrightarrow{-H_2O}
$$
 [Ph₃SnONHCOPh] \longrightarrow
\nPh₄Sn + [Ph₂Sn(ONHCOPh)₂] (3)

The reaction of the hydroxylamine with triphenyltin chloride in the presence of triethylamine yielded a product, the triethylammonium salt of the tin-containing complex anion, IIIa, only when a large excess of amine was used

$$
\mathsf{Ph}_3\mathsf{SnCl} + \mathsf{HONHCOPh} \xrightarrow{\mathsf{excess Net}_3} \mathsf{NEt}_3 \mathsf{H}^+ \left[\mathsf{Ph}_3\mathsf{Sn} \begin{matrix} O-N \\ O-Ph \\ O-Ph \end{matrix} \right] (4)
$$

Dissolution of Me₃SnONHCOPh in triethylamine yielded the trimethyltin homolog, IIIb

Since the Lewis acidity of the trimethylsilyl moiety is much lower than that of the heavier group IV metals, the *0* trimethylsilyl derivative of **N-phenyl-N-benzoylhydroxyl**amine (IV) was synthesized for comparative purposes. Because of the inertness of the silicon-oxygen bonds of hexamethyldisiloxane, this compound was synthesized by the protolysis of the silicon-nitrogen bonds of the analogous disilazane

$$
^{1}/_{2}
$$
(Me₃Si)₂NH + HONPhCOPh $\frac{-NH_{3}}{N}$ Me₃SiONPhCOPh (6)

(5) R. Okawara and M. Wada, *Advan. Organometal. Chem., 5,* **15 1** (1967) .

Like O -(trimethylstannyl)acetoxime,³ the O -trimethyltin derivative of **N,N-diethylhydroxylamine** is a mobile liquid, which hydrolyzes readily in air to its component protic reagents, characteristic of the tin-oxygen bonds of fourcoordinate tin compounds. The feature of highest mass in the mass spectrum (Table I) is the polyisotopic parent ion at m/e 252. Other monotin-containing fragments produced by the fission of tin-carbon and tin-oxygen bonds are observed at lower mass. Corroboration of the unassociated nature of the compound comes from the nmr $2J(117,119Sn-C-1H)$ coupling constant, which appears to be sensitive to the coordination number at tin for methyltin derivatives. The value of this parameter for the trimethyltin derivatives of N , N -diethylhydroxylamine, acetoxime, 3 and cyclohexanone oxime³ all lie close to that for tetramethyltin (\sim 54 Hz) which is unable to associate.

The organotin derivatives of the N -benzoylhydroxylamines are. in contrast, extremely stable to moisture. Colligative measurements (osmometry) show that the derivatives of *N***phenyl-N-benzoylhydroxylamine** are monomeric in dilute chloroform solution. Similar measurements for 0-(trimethyl. **stanny1)-N-benzoylhydroxylamine** indicated some association. Evidence for association via hydrogen bonding of the nitrogenous proton comes from infrared dilution studies on the latter compound and its tripropyltin analog. The neat compounds exhibit a very broad N-H stretching frequency at $ca. 3200 \text{ cm}^{-1}$. On dissolution in dichloromethane, a second band of much narrower line width appears to higher energy at ca. 3410-3430 cm⁻¹, until, at concentrations of $\approx 1\%$, only the higher frequency band remains.

Information concerning the colligative nature in the solid phase is available from the mass spectra of $Me₃SnONHCOPh$, Me₃SnONPhCOPh, and Ph₃SnONPhCOPh, for which relevant data are listed in Tables 11-IV, respectively. In addition to monotin-containing fragments derived from fission of the monomer units, all three compounds exhibit several ditincontaining fragments. Furthermore, in the spectra of the two trimethyltin derivatives, there occur monotin fragments of mass in excess of the monomer parent ion. In contrast, the spectra of the O -trimethylsilyl (Table V) and O -triphenyllead (Table VI) compounds show only ions resulting from the fragmentation of a monomer species.

Recently we have completed a determination of the crystal structure of $Ph_3SnONPhCOPh.$ ⁶ The compound possesses a trigonal-bipyramidal arrangement of groups about tin, with two equatorial and one axial phenyl groups (Figure 1). The hydroxylamine residue is covalently bound at an equatorial site, and the carbonyl group coordinates *intra*molecularly to the tin atom via the remaining axial site. This is consistent with the lowering of the infrared carbonyl stretching frequency from 1620 cm^{-1} in the parent hydroxylamine to 1540 cm^{-1} in the O-triphenyltin derivative. Similar shifts to lower energy for this mode are observed for the other tin and lead derivatives of the N-benzoylhydroxylamines in both the solid and solution. In the O -trimethylsilyl derivative, however, a shift to *higher* energy (1660 cm^{-1}) is observed. This value is perhaps more truely representative of a "free," uncoordinated carbonyl group in this type of compound, since inter- or intramolecular hydrogen bonding of the type

(6) T. **J.** King and P. G. Harrison, *J. Chem. SOC., Chem. Commun.,* **815 (1972).**

Table I. Mass Spectral Data for the Tin-Containing
Fragments of Me₃SnONEt₁

Mass	Rel intens	Assignment	
252	17.6	$Me3$ SnONEt ₂ ⁺	
237	11.0	Me ₂ SnONEt ₂ ⁺	
207	3.5	SnONEt, ⁺	
184	8.3	$C_{s}H_{s}Sn^{+}$?	
164	100	$Me2Sn+$	
149	9.7	$Me, Sn+$	
134	16.9	MeSn ⁺	
119	4.9	Sn^{+}	

Table **11.** Mass Spectra Data for the Mono- **and** Ditin Fragments of Me,SnONHCOPh

 a Mass measurement: Found: 329.9242. Calcd: Me, Sn₂O, 329.9179; Me,Sn,O,N, 329.8689. b Mass measurement: Found: 285.9837. Calcd: 285.9869.

is likely to occur in the parent hydroxylamine, lowering the carbonyl stretching frequency to that observed in both the solid and the chloroform solution (1620 cm^{-1}) .

Since Ph₃SnONPhCOPh is unequivocally monomeric in the solid phase, we attribute the observation of ditin fragments in the mass spectrum to the decomposition of the solid under electron impact at the temperature of the inlet (140°) . The observed ditin fragments at *m/e* 716, 639, and 562 are all derived from $Ph_3SnOSnPh_3$, which may be a product of the thermal decomposition. Such reactions of this and related compounds are the subject of further investigation in this laboratory. In the cases of the two trimethyltin derivatives, however, no decomposition was observed at the inlet ternperature used. Moreover, a greater number of fragments attributable to the fission of dimer species were observed for these compounds than for $Ph_3SnONPhCOPh$, including some indicative of the loss of only methyl groups from a dimer species.

It seems likely that, whereas $Ph₃SnONPhCOPh$ is monomeric in both the crystal and solution phases, the trimethyl-

 $a-h$ The following footnotes contain found and calculated highresolution mass measurements, respectively. a 329.9203, 329.9179. b 346.0219, 346.0253. **C** 329.9736, 329.9729. *d* 314.0757, 314.0419. **e** 287.9859,287.9920. *f* 270.9779, 270.9780. **g** 240.9356, 240.9311. *h* 226.9884,226.9882. *i* Intensity measurement not possible because of overlapping monotin and ditin fragments. **I'** Not assigned.

Table **IV.** Mass Spectral Data for Mono- and Ditin Fragments for Ph,SnONPhCOPh

Mass	Rel intens	Assignment		
Ditin Fragments				
716	16.4	Ph,SnOSnPh,†		
639	100	Ph, SnOSnPh, +		
621	7.6	α		
562	b	Ph ₂ SnOSnPh ₂ +		
281	6.0			
Monotin Fragments				
562	1.3	$Ph3$ SnONPhCOPh ⁺		
546	0.9	Ph ₃ SnOCPhNPh ⁺		
518	0.3	Ph ₃ SnNPh ₂ +		
485	100	Ph ₂ SnONPhCOPh ⁺		
469	4.9	Ph_2 SnOCPhNPh ⁺		
426	0.7	$Ph_3Sn(c-Hex)^+$ ^c		
394	10.5	$Ph_3SnO_2C^*$		
350	64.4	Ph, Sn^+		
331	3.1	SnONPhCOPh ⁺		
290	5.6	Ph ₂ SnOH ⁺		
273	9.0	$Ph, Sn+$		
240	3.5	$PhSnO2C+$		
212	1.3	PhSnO ⁺		
196	44.1	PhSn*		
136	6.1	SnOH+		
119	25.5	Sn*		

a Not assigned. *b* Intensity measurement not possible because of overlapping mono- and ditin fragments. c c-Hex = cyclohexadienyl.

tin derivatives are associated in the solid. The intermolecular forces must however be fairly weak since monomeric species, which presumably also have the trigonal-bipyramidal structure, are present in solution. In the absence of additional structural information, however, it is difficult to assess

Rel intens	Assignment
8.7	$(P + 1)^+$
33.7	Me ₃ SiONPhCOPh ⁺
13.7	$(P - Me + 1)^+$
33.7	Me, SiONPhCOPh ⁺
8.1	
5.0	PhNCOPh ⁺
6.9	
10.0	Me,SiONPhCO+
10.0	Me,SiONPh ⁺
3.8	
3.4	Me, SiOPh ⁺
4.6	SiONPh ⁺
7.2	$PhN=C=O^+$
31.1	PhCOH ⁺
93.6	PhCO ⁺
6.9	PhO ⁺
16.2	PhN ⁺
14.3	$C_6H_6^+$
100	$C_sH_s^+$
8.1	$C_{\alpha}H_{\alpha}^{+}$
33.7	
3.8	$Me2SiO+$
24.3	
10.6	
6.9	MeSiO ⁺
23.1	

Table **VI.** Mass Spectral Data for the Lead-Containing Fragments of Ph,PbONPhCOPh

the degree of association fn the solid. Dimerization, which is implied by the mass spectral data, would lead to possible formulations such as A, **B,** or C, while association may take place via the formation of linear chains, as in D, involving

Figure **1.** The structure of **0-(triphenylstanny1)-N-phenyl-N**benzoylhydroxylamine.

planar trimethyltin moieties, frequently encountered among trimethyltin derivatives in the solid state.⁷ This last structure may reasonably excluded however, since for this geometry only one infrared active tin-carbon stretching frequency is expected, contrary to observation (*vide infra*).

Structural information concerning the solid may also be gained from tin-1 19m Mossbauer spectral data, which are collected in Table VII, together with those for some related compounds. The quadrupole splittings for the model compounds $Me₃SnOSiPh₃$, $(Ph₃Sn)₂O$, and $Ph₃SnOSiPh₃$, where the bulkiness of the groups prevents any deviation from fourcoordination at tin, have a value of *cu.* 2 mmjsec, close to that expected from point charge considerations of electronic imbalances in the σ framework.⁸ Similarly, Me₃SnOH and trimethyl- and triphenyltin acetates, which have the fivecoordinate trans- R_3SnX_2 structure, have quadrupole splittings \geq 3 mm/sec. Triphenyltin oxinate, which is presumed to have the cis - R_3SnX_2 structure, again has a splitting of the same order as the four-coordinate R_3SnX compounds. Assignments of a four-coordinate structure to $Me₃SnONEt₂$ and the five-coordinate cis -R₃SnX₂ to Ph₃SnONPhCOPh and

the related $Ph_3SnON=CPhO^-$ anion are consistent with previous structural assignments made on the basis of Mossbauer data. The increased splittings for $Me₃SnONPhCOPh$ (2.36) mm/sec), $Pr₃SnONPhCOPh$ (2.65 mm/sec), and $Me₃SnONH-$ COPh (2.74 mm/sec) indicate an increasing perturbation from the $cis-R_3SnX_2$ geometry involving additional, albeit weak, coordination as in structures B or C. In $Me₃SnONH-$ COPh, the association will be further reinforced by intermolecular hydrogen bonding. (See Figure 2 for a tin- 1 19m Mossbauer spectrum of Ph₃SnONPhCOPh.)

Although structures A and C may not be ruled out, a structure involving dimerization via the formation of distannoxane rings, which are prevalent in organotin chemistry,⁵ would be most likely to dissociate into monomeric species. In $Ph₃Sn-$ ONPhCOPh, the bulkiness of the phenyl rings bound to tin prevent any close intermolecular approach. and the smallest *intermolecular* Sn. $\cdot \cdot$ O distances are in excess of 6.5 Å.⁶ However, replacement by much smaller methyl groups would not disallow close approach for distannoxane ring formation, in which $Sn-O(ring) = 2.2 \text{ Å}^5$ (although much longer Sn-O distances have been observed), and the formation of dimeric species would depend on the Lewis acidity of the tin atom in that particular environment. In structure B (and C) the coordination number at tin would be raised to 6, higher than

(7) B. Y.-K. Ho and **J. J.** Zuckerman, *Organometal. Chem. Rev., Sect. A,* in **press.**

(8) R. **V.** Parish and R. **H.** Platt, *Znorg. Chim. Acta,* **4,** *65* **(1970).**

a P. J. Smith, *Organometal Chem. Rev., Sect. A, 5,* **373 (1970).**

that usually accepted for coordination saturation at tin in trialkyltin compounds.

Similar coordination-dimerization equilibria in solution have been observed for 1,3-disubstituted distannoxanes⁹ and the dialkyltin alkoxide halides,¹⁰ the latter having been elegantly demonstrated by the tin-1 19 nmr chemical shifts of the two species present in the solution.

Although the solid-state structure of the two trimethyltin derivatives is not unequivocal, it seems certain that in solution they have the trigonal-bipyramidal structure, which will possess inequivalent tin-carbon bond distances. In $Ph₃Sn-$ ONPhCOPh the two equatorial tin-carbon bond distances are 2.13 and 2.14 **A,** while the axial distance is 2.19 **A.** The proton nmr of the trimethyltin derivatives might, therefore, be expected to exhibit two distinct resonances due to the inequivalent methyl groups. One signal, together with a single set of tin-117,119 satellites, is observed in both cases. The value of $2J(117,119Sn-C-1H)$ of methyltin compounds, *via* a Fermi contact mechanism, is a guide to the s-electron density directed along the tin-carbon bond. We have noted before that the magnitude of this coupling constant parallels the tin-carbon bond distance, its value rising as the bond distance decreases.¹¹ In the compounds under discussion here, the value of $2J(Sn-H)$ is \sim 54 Hz, a value expected for fourcoordination at tin, and much lower than that observed for trans five-coordinate Me_3SnX_2 derivatives (~70 Hz). The observed proton nmr spectra are consistent with a model involving stereochemical nonrigidity of the groups about tin, the value of the coupling constant being such that one cannot distinguish between four-coordinate tetrahedral and five-coordinate cis trigonal-bipyramidal trimethyltin compounds. This may be rationalized since the latter may be regarded as rather distorted tetrahedral with smaller CSnC bond angles.

The mass spectra of the compounds merit further discussion. In the spectrum of $Me₃SnONPhCOPh$, nine ditin fragments could be distinguished. That at *m/e* 239 was shown by a high-resolution mass measurement *not* to con. tain nitrogen, but to possess the SnOSn backbone. The fragments of higher mass may be interpreted similarly, with organic residues pendant from the oxygen, although six membered SnONSnON rings may not be ruled out entirely. The major fragmentation processes of the monomer involve the fission of the tin-oxygen and tin-carbon bonds, resulting

(10) A. C. Chapman, A. *G.* **Davies, P.** *G.* **Harrison, and W. McFarlane,** *J. Chem. SOC. C,* **821 (1970).**

(1 1) J. Buckle, M. K. Das, and P. *G.* **Harrison,** *Inorg. Chim.* **Acta, 6, 17 (1972).**

Figure 2. The tin-119m Mossbauer spectrum of Ph₃SnONPhCOPh run at 77° K *vs.* a Ba^{119m}SnO₃ source.

in the formation of $Me_nSnOCPhNPh⁺$ and $Me_nSn⁺$ (n = 0-3) ions, **A** high-resolution mass measurement has confirmed the composition of the SnOCPhNPhO' ion. Similar fragmentation processes were observed for the O-trimethyltin oximes, however in the present case the spectrum exhibits several monotin-containing fragments between *m/e* 226 and 345 of appreciable abundance which must arise *via* alternative fragmentation paths. The constitution of these fragments has been unequivocally determined by high-resolution mass measurements, although in the absence of metastable peaks their origin is not certain.

The fragment of highest mass at *m/e* 345 corresponds to loss of oxygen from the Me₂SnONPhCOPh⁺ ion. This may be readily rationalized by a ring contraction reaction involving the loss of oxygen from the tin-containing heterocyclic ring

The fragments at m/e 240 and 270 correspond to SnO_2CPh^+ and $Me₂SnO₂CPh⁺$, respectively, and probably originate *via* a similar ring contraction reaction involving the loss of the nitrene species, PhN:

The fragment at m/e 226 is, surprisingly, the Me₂PhSn⁺ ion. Although conceivably arising from the decomposition of the dimethyltin fragments at *m/e* 361 or 287, or by the elimination of phenyl isocyanate at *m/e* 345, its formation, necessarily *via* a 1,3-phenyl migration, is more probably by the decar-

⁽⁹⁾ D. L. Alleston, A. *G.* **Davies, M. H. Hancock, and R. F. M. White,** *J. Chem. Sac.,* **5469 (1963).**

Figure 3. Major fragmentation processes of Me₃SiONPhCOPh. Calculated values for metastable ions are given in parentheses.

boxylation of the $Me₂SnO₂CPh⁺$ ion, a process well known in main-group organometallic chemistry¹²

$$
Me2SnO2CPh+ \to Me2SnPh+ + CO2
$$
 (9)

The fragment at m/e 287 corresponds to the $Me₂SnO₂N₂HPh⁺$ ion, not the $Me₃SnO₂NPh⁺$ ion, and presumably originates from the decomposition of the dimer. The mass measurement on the fragment at *m/e* 313 was not satisfactory but indicated its probable constitution as the substituted carbodiimide $Sn(Ph)N=C=NPh^+$ ion.

In general the mass spectra of $Me₃SnONHCOPh$, $Ph₃Sn-$ ONPhCOPh, and Ph₃PbONPhCOPh exhibited similar fragmentation pathways, including the ring contraction reactions involving the elimination of oxygen and phenylnitrene.

Whereas the tin and lead derivatives give rise to only weak parent ions, that of the trimethylsilyl derivative IV is one of the strongest silicon-containing features of the spectrum. The major fragmentation processes, most of which have been confirmed by the observation of metastable ions, are shown in Figure 3. Cleavage of the silicon-oxygen bond is, as expected because of its high bond dissociation energy, not

(12) *G.* **B.** Deacon, *Organometal. Chem. Rev., Sect. A, 5, 355* **(1970).**

observed, and it is readily apparent that, on fragmentation of the hydroxylamine residue, the charge is retained by aromatic species where delocalization is possible.

Principal infrared bands for the benzoylhydroxylamines and their derivatives are listed in Table VIII. For the trimethylsilyl derivative, the silicon-oxygen and the antisymmetric and symmetric silicon-carbon stretching vibrations are located at 768, 690, and 640 cm^{-1} , respectively. The similar bands for the two trimethyltin derivatives are to be expected in the region $ca. 500-600 \text{ cm}^{-1}$. In this region, both compounds exhibit three bands. For a trigonal-bipyramidal (or a distorted octahedral) structure, group theoretical considerations predict three infrared-active metal-carbon stretching frequencies; hence a total of four bands are to be expected in this region. Figure 4 shows the infrared spectra of Me₃SnONPhCOPh in the region 450-600 cm⁻¹ in the solid and in *ea.* **5%** dichloromethane solution. The band at 508 cm^{-1} of the solid is clearly resolved into two components, at 507 and 497 cm^{-1} , on dissolution. No similar splitting could however be observed for $Me₃SnONHCOPh$. The coordinate tin-oxygen stretching vibration is extremely variable in position and is not assigned in this study. We have previously assigned this vibration for a number of phos-

Figure 4. Infrared spectra in the region 450-600 cm⁻¹ of Me,SnONPhCOPh in the solid state (upper) and in dichloromethane solution (lower).

phine and arsine oxide complexes of the stannic halides in the range $338-482$ cm⁻¹.¹³

Experimental Section

All compounds with moisture-sensitive tin-oxygen and siliconnitrogen bonds were handled under an atmosphere of dry nitrogen or argon. Infrared measurements were performed using a Perkin-Elmer 521 spectrophotometer calibrated with polystyrene film and water vapor. Mass spectra were recorded at 70 eV on an AEI MS-9 instrument. The polyisotopic tin and lead containing fragments, because of their distinctive isotope abundances,¹⁴ are easily recognized. The m/e values listed in Tables I-VI are based on the isotopes tin-1 19 and lead-207 for convenience. Where high-resolution mass measurements have been performed, the particular isotopic peak used is self-evident. Proton nmr spectra were recorded at 100 MHz using a Varian HA-100 instrument. Osmometric molecular weight determinations were carried out in chloroform solution on a Mechrolab instrument. Tin-119m Mossbauer spectra were recorded at 77°K *us.* a Ba119mSn0, source, and the data were refined by least squares to give the best lorenztian fits.

1. *O*-(Trimethylstannyl)-N,N-diethylhydroxylamine, Ia. Trimethyltin hydroxide (8.0 g, 44.2 mmol) and N,N-diethylhydroxylamine $(4.0 \text{ g}, 45.0 \text{ mmol})$ in benzene (50 ml) were refluxed in a flask equipped with a Dean and Stark water separator. Water came over immediately and the reaction was complete within 10 min. On cooling, the solvent was removed *in vacuo* leaving the product as a mobile, colorless oil, bp 30-31' (0.2 mm). *Anal.* Calcd for C₇H₁₉NOSn: C, 33.4; H, 7.6; N, 5.6. Found: C, 33.2; H, 7.7; N, 5.4. Nmr data (CH₂Cl₂ solution): τ 9.68 (Me₃Sn), ²J(^{117,119}Sn-¹H) = 55.5 Hz (unresolved); triplet τ 8.97, quartet τ 7.42 (N-Et), $J(H-H)$ = *ea.* 7 Hz.

2. O-(Trimethylstannyl)-N-phenyl-N-benzoylhydroxylamine, Id. Trimethyltin hydroxide (1.81 g, 10.0 mmol) and N-phenyl-Nbenzoylhydroxylamine (2.1 3 g, 10.0 mmol) were similarly azeotropically dehydrated. Recrystallization from cyclohexane yielded colorless needle crystals of **0-(trimethylstanny1)-N-phenyl-N-benzoyl**hydroxylamine, mp 120-122", sub1 pt 120" (0.1 mm). *Anal.* Calcd for $C_{16}H_{19}NO_2Sn$: C, 51.1; H, 5.1; N, 3.7 (mol wt 376). Found: C, 51.7; H, 5.1; N, 3.7 (mol wt 338 (0.022 M)). Nmr data (CH₂Cl₂)

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(14) D. **B.** Chambers, **F.** Glockling, and M. Weston, *J. Chem.* **SOC.** *A,* **1759 (1967).**

solution): τ 9.58 (Me₃Sn), ²J(^{117,119}Sn-¹H) = 54.1, 56.7 Hz.

3. O-(Tri-n-propylstannyl)-N~phenyl-N-benzoylhydroxylamine, Ie, was similarly prepared as a golden viscous oil from bis(tripropy1tin) oxide (2.56 g, 5.0 mmol) and the hydroxylamine (2.13 g, 10.0 mmol). The compound could not be distilled *in uaeuo* but decomposed at temperatures in excess of *ca.* 150° . *Anal.* Calcd for $C_{22}H_{31}NO_2Sr$ C, 57.4, H, 6.8; N, 3.1. Found: C, 58.0; H, 6.5; N, 3.0.

4. O-(Triphenylstannyl)-N-phenyl-N-benzoylhydroxylamine, If, was obtained from triphenyltin hydroxide (3.67 g, 10.0 mmol) and the hydroxylamine (2.13 g, 10.0 mmol) and recrystallized from benzene as colorless needle crystals, mp 115.5-1 16.5". *Anal.* Calcd for $C_{31}H_{25}NO_2Sn$: C, 66.2; H, 4.5; N, 2.5 (mol wt 562). Found: C, $65.\overline{9}$; H, 4.7 ; N, 2.4 (mol wt 561 (0.013 M)).

5. **O-(Triphenyllead)-N-phenyl-N~benzoylhydroxylamine,** 11. Triphenyllead hydroxide (4.55 g, 10.0 mmol) and the hydroxylamine (2.13 g, 10.0 mmol) were azeotropically dehydrated in boiling benzene (50 ml). Removal of the solvent *in vucuo* gave a golden viscous oil, which could be recrystallized as an oil from hexane. Pumping at high vacuum for 24 hr produced a golden yellow crystalline solid, mp 44-46°. *Anal.* Calcd for C₃₁H₂₅NO₂Pb: C, 57.2; H, 3.9; N, 2.2. Found: C, 57.3; H, 4.0; N , 2.2.

6. **O-(Trimethylsilyl)-N-phenyl-N-benzoylhydroxylamine,** IV. **N-Phenyl-N-benzoylhydroxylamine** (2.13 g, 10.0 mmol) and hexamethyldisilazane (0.80 g, 5.0 mmol) were refluxed in dry tetrahydrofuran (50 ml) until the evolution of ammonia had stopped (1 hr). The solvent was removed and distillation of the residue yielded the *compound* as a golden oil, bp 108' (0.05 mm). *Anal.* Calcd for $C_{16}H_{19}NO_2Si$: C, 67.1; H, 6.6; N, 4.9. Found: C, 66.7; H, 6.5; N, 5.1.

7. 0-(Trimethylstanny1)-N-benzoylhydroxylamine, Ib. Trimethyltin hydroxide (1.81 g, 10.0 mmol) and N-benzoylhydroxylamine (1.37 g, 10.0 mmol) were azeotropically dehydrated in boiling benzene. On cooling, the hot solution yielded O (trimethylstannyl)- N -benzoylhydroxylamine as a white powder, mp 255 $^{\circ}$ dec. The compound decomposed on attempted sublimation at 160" (0.01 mm). *Anal.* Calcd for C₁₀H₁₅NO₂Sn: C, 40.0; H, 5.0; N, 4.7 (mol wt 299). Found: C, 40.7; H, 5.0; N, 4.7 (mol wt 394 (0.015 *M*)). Nmr data (CH₂Cl₂ solution): τ 9.62 (*Me*₃Sn), ²*J*(1171119Sn-1H) = 54.3, 56.7 Nmr data (CH₂Cl₂ solution): τ 9.62 ($Me₃Sn$), $2J(117,119Sn-1H) =$
54.3, 56.7 Hz. Recrystallization from triethylamine yielded

 $(NEt₃H⁺)(Me₃SnONCPhO⁻)$, mp \sim 200[°] dec. *Anal.* Calcd for $C_{16}H_{30}N_2O_3Sn$: C, 47.9; H, 7.5; N, 7.0. Found: C, 48.4; H, 7.2; N, 6.8.

8. **O.(Tri-n-propylstanny1)-N-benzoylhydroxylamine,** IC, was similarly obtained by the azeotropic dehydration of the hydroxylamine $(1.37 g, 10.0 mmol)$ and bis(tripropyltin) oxide $(2.56 g, 5.0$ mmol). The compound was a colorless viscous oil which could not be purified by distillation. *Anal.* Calcd for C₁₆H₂₇NO₂Sn: C, 50.0; H, 7.1; N, 3.7. Found: C, 49.5; H, 6.5; N, 2.8.

9. Attempted Syntheses **of 0-(Triphenylstanny1)-N-benzoyl**hydroxylamine. (a) Triphenyltin hydroxide (3.67 g, 10.0 mmol) and N-benzoylhydroxylamine (1.37 g, 10.0 mmol) were azeotropically dehydrated in benzene (50 ml). On cooling the hot mixture, a mass of white needle crystals were obtained, which were identified as tetraphenyltin by comparison with a authentic sample.

(b) An exothermic reaction took place when triethylamine (6.1 g, 60.0 mmol) was added to triphenyltin chloride (3.85 g, 10.0 mmol) and N -benzoylhydroxylamine (1.37 g, 10.0 mmol) in methanol (150 ml). After *ea.* 2 hr much solid had crystallized out of solution and was filtered off and dried; mp *ea.* 200" dec. *Anal.* Calcd for $C_{31}H_{36}N_2O_2Sn$: C, 63.4; H, 6.2; N, 4.8. Found: C, 63.1; H, 5.9; N, 4.7.

Registry **No.** *0-(* Trime thylstanny1)-N,N- diethylhydroxylamine, 37127-23-4; trimethyltin hydroxide, 56-24-6; *N,N*diethylhydroxylamine, 37 10-84-7; 0-(trimethylstanny1)-N**phenyl-N-benzoylhydroxylamine,** 3953 1-79-8; N-phenyl-Nbenzoylhydroxylamine, 304-88-1; O -(tri-*n*-propylstannyl)-**N-phenyl-N-benzoylhydroxylamine,** 39046-68-9; bis(tripropyltin) oxide, 1067-29-4; **0-(triphenylstanny1)-N-phenyl-**N-benzoylhydroxylamine, 39046-69-0; triphenyltin hydroxide, 76-87-9; **0-(tripheny1lead)-N-phenyl-N-benzo** ylhy droxylamine, 39532-11-1; *O*-(trimethylsilyl)-*N*-phenyl-*N*-benzoylhydroxylamine, 3953 1-82-3; triphenyllead hydroxide, 894- 08-6; hexamethyldisilazane, 999-97-3; O-(trimethylstannyl)-N-benzoylhydroxylamine, 3953 1-72- 1 ; N-benzoylhydroxylamine, 495-18-1; *O*-(tri-*n*-propylstannyl)-*N*-benzoylhydroxylamine, 39531-73-2.