CLEAVAGE OF ORGANOPOLYSILANES BY LITHIUM ALUMINUM HYDRIDE. INTERMEDIATE (ORGANOSILYL)ALUMINUM COMPOUNDS

HENRY GILMAN AND RICHARD A. TOMASI*

Department of Chemistry, Iowa State University of Science and Technology, Ames. Iowa 50010 (U.S.A.) (Received July 8th, 1968)

SUMMARY

The reductive cleavage of silicon-silicon bonds in phenyl-substituted organopolysilanes by lithium aluminum hydride has been shown to occur in refluxing tetrahydrofuran. The reaction of hexaphenyldisilane was studied in greater detail, and the results show that the cleavage produces intermediate(organosilyl)aluminum compounds. The formation of the latter from reactions of triphenylsilyllithium with some aluminum compounds was also investigated.

INTRODUCTION

The reductive cleavage of aryl-substituted silicon-silicon bonds in organopolysilanes by alkali metals has been accomplished under a wide variety of conditions**. In aprotic solvents, the reactions afford organosilylmetallic compounds, according to the following equation:

$$R_3 SiSiR_3 + 2M \rightarrow 2R_3 SiM \tag{1}$$

Only very low yields of trialkylsilylmetallic compounds have resulted from fission of 1,1,1-trialkyl-2,2,2-triphenyldisilanes. Thus, trimethylsilylpotassium was obtained utilizing sodium/potassium alloy in ether², and triethylsilyllithium was formed by the lithium cleavage of 1,1,1-triethyl-2,2,2-triphenyldisilane in tetrahydrofuran^{3,4}. In contrast to the reactions of aryl-substituted compounds, numerous attempts to effect similar cleavages of hexamethyl- and hexaethyldisilane have proved unsuccessful^{2,3,5-9}. Thus, it is apparent that aryl groups activate the silicon-silicon bond toward cleavage and/or stabilize the resulting organosilylmetallic. However, the cleavage of octamethyltrisilane and decamethyltetrasilane with sodium/potassium alloy in tetrahydrofuran has been reported¹⁰. Derivatization of the resulting silylmetallic species with chlorotrimethylsilane gave a variety of linear and cyclic permethylated polysilanes. Evidently, the presence of silicon-silicon bonds offers some stability to non-arylsubstituted silyl-metallics and/or facilitates cleavage.

A similar effect has also been observed in the reductive cleavage of siliconsilicon bonds by hydrogen over a "copper chromite" catalyst, with the phenyl-sub-

^{*} Present address: Chemistry Department, University of Tulsa, Tulsa, Oklahoma, U.S.A.

^{**} For comprehensive reviews on cleavages by alkali metals see ref. 1.

stituted compounds undergoing hydrogenolysis according to equation (2), while hexaethyldisilane was not affected under comparable conditions¹¹.

$$R_3 SiSiR_3 + H_2 \rightarrow 2R_3 SiH$$
⁽²⁾

Hexaphenyldisilane has been shown to be unreactive towards lithium aluminum hydride in ether¹². However, it has been noted that silicon-silicon bond cleavage occurs upon the addition of hexachlorodisilane to lithium aluminum hydride in tetrahydrofuran or in di-n-butyl ether, which afforded exclusively silane $(SiH_4)^{13}$, whereas disilane is obtained when diethyl ether was used as the solvent¹⁴. Also, cleavage of disilane by potassium hydride in ethylene glycol dimethyl ether yielded silane and silylpotassium¹⁵, as illustrated by equation (3). Similarly, the fission of

$$x H_3 SiSiH_3 + KH \rightarrow x SiH_4 + \left[\frac{1}{x}(SiH_2)_x\right] \xrightarrow{KH} KSiH_3$$
 (3)

silicon-silicon bonds in mono- and dialkyldisilanes by potassium hydride has been observed, accompanied by redistribution of the alkyl groups¹⁶.

DISCUSSION AND RESULTS

As compared to other organopolysilanes, octaphenylcyclotetrasilane¹⁷ (I) is unusually reactive in silicon-silicon bond cleavage. Thus, it appeared likely that this compound, (I), might react with lithium aluminum hydride. After 168 h in refluxing ether, no evidence of reaction was indicated and octaphenylcyclotetrasilane was recovered in nearly quantitative amounts. However, after 20 h at room temperature in tetrahydrofuran (THF), no octaphenylcyclotetrasilane was recovered. The primary product of this reaction was decaphenylcyclopentasilane¹⁸ (II), isolated in 64% yield. It seems likely that the mechanism involved in this reaction is analogous to that proposed for this same cyclotetrasilane to cyclopentasilane transformation brought about by organosilyllithium compounds^{19,20} and other organometallic compounds^{19,21}. The proposed mechanism of this conversion is shown in the following sequence of reactions. The fact that decaphenylcyclopentasilane was recovered

in a 74% yield after similar treatment with lithium aluminum hydride is consistent with the accumulation of this compound in the reaction with octaphenylcyclotetrasilane. From these and other reactions of (I) and (II) with lithium aluminum hydride.

J. Organometal. Chem., 15 (1968) 43-56

further support of the mechanistic pathway was obtained by the isolation of products containing one, two, three, four and five diphenylsilylene units, corresponding to hydrolysis of intermediate organosilylmetallics such as (III), (IV) and others predicted from subsequent cyclization cleavage steps.

The reactions of lithium aluminum hydride with hexaphenyldisilane, 1,2dimethyl-1,1,2,2-tetraphenyldisilane, 1,1,2,2-tetramethyl-1,2-diphenyldisilane and hexaethyldisilane in refluxing tetrahydrofuran were also investigated. The general reaction is illustrated in equation (7). Under comparable conditions, the yields of

$$R_{3}Si-SiR_{3}+H^{-} \rightarrow R_{3}SiH+R_{3}Si^{-} \xrightarrow{H^{+}} 2R_{3}SiH$$
(7)

the corresponding products decreased, respectively, as the number of phenyl substituents decreased, with no triethylsilane being detected from hexaethyldisilane. 1,1,1-Trimethyl-2,2,2-triphenyldisilane also reacted, with trimethylsilane being obtained by distillation from the reaction mixture prior to hydrolysis, and triphenylsilane being isolated after hydrolysis. In addition, no reaction was observed with dodecamethylcyclohexasilane. The formation of organosilylmetallic species in these reactions is strongly suggested by the observation that the cleavages by lithium aluminum hydride closely follow the same general pattern of greater reactivity of aryl-substituted silicon-silicon bonds towards fission by alkali metals in the preparation of organosilylmetallic reagents¹⁻⁹.

The reaction of lithium aluminum hydride with hexaphenyldisilane was studied in some detail in order to get a better understanding of the cleavage reactions and, in particular, to investigate the nature of the proposed silylmetallic compounds, which could, conceivably, be associated with either lithium or aluminum.

To determine if appreciable amounts of triphenylsilyllithium were present, the reaction mixture obtained from the cleavage of hexaphenyldisilane was treated with excess chlorotrimethylsilane. Subsequent to acid hydrolysis, no 1,1,1-trimethyl-2,2,2-triphenyldisilane could be isolated, indicative of the absence of a silyllithium compound. Significantly, a 60% yield of triphenylsilane was obtained. Similar results were obtained upon repetition of this reaction on a little larger scale. In this case, the mixture was hydrolyzed in deuteriochloric acid. The overall yield of triphenylsilane in this instance was 78%. Of this, 31% was triphenylsilane-d, which established the presence of a silylmetallic compound other than lithium, namely, a (triphenylsilyl)aluminum compound. The following sequence of reactions is consistent with these findings. The fact that no 1,1,1-trimethyl-2,2,2-triphenyldisilane was isolated is apparently due to a lower reactivity of (triphenylsilyl)aluminum dichloride, (VI).

$$(C_{6}H_{5})_{6}Si_{2} + LiAlH_{4} \rightarrow (C_{6}H_{5})_{3}SiH + Li^{+}[(C_{6}H_{5})_{3}SiAlH_{3}]^{-}$$

$$(C_{6}H_{5})_{3}SiD \xrightarrow{D_{3}O^{+}} (C_{6}H_{5})_{3}SiAlCl_{2} \xrightarrow{3(CH_{3})_{3}Sic1} (V) \qquad (8)$$

$$(C_{6}H_{5})_{3}SiC1 \stackrel{1}{=} (VI)$$
No reaction

Based on these reactions, the actual yield of triphenylsilane-d, and therefore of silvaluminum compound (VI), was 62%.

Similar conclusions may be drawn from cleavage reactions which were car-

bonated prior to hydrolysis. In the case of hexaphenyldisilane, triphenylsilane was obtained in a 62% yield. From the reaction involving 1,1,1-trimethyl-2,2,2-triphenyldisilane mentioned earlier, which was also treated with carbon dioxide prior to hydrolysis, triphenylsilane was obtained in a 63.9% yield. Although no triphenylsilanecarboxylic acid was isolated from these reactions, some oxygen-containing organosilicon compounds were obtained, which could have arisen from decarbonylation of the acid²². However, these products could also arise from basic hydrolysis of triphenylsilane by moisture condensed on the cold, carbonation mixture prior to hydrolysis. These reactions may be illustrated by equation (9).

$$(V) + CO_2 \rightarrow (C_6H_5)_3 \text{SiAl}(OR)_2 \xrightarrow{H^+} (C_6H_5)_3 \text{SiH}$$
(9)
(VII)

(OR represents hydride reduction products of CO₂)

An alternate synthesis of lithium trihydro(triphenylsilyl)aluminate, (V), was carried out by the addition of triphenylsilyllithium to lithium aluminum hydride. The reaction, as illustrated by the following equation, is analogous to that reported for the reaction of metal alkyls with lithium aluminum hydride²³.

$$(C_6H_5)_3SiLi + LiAlH_4 \rightarrow (V) + LiH$$
(10)

Using a 1:1 mole ratio of reactants, half of the resulting mixture was carbonated and, subsequent to hydrolysis, triphenylsilane was isolated in a 78.4% yield. The remaining portion of the reaction mixture was treated with excess chlorotrimethylsilane and yielded 68% of triphenylsilane after acidification. As previously noted, both of these routes show that triphenylsilyllithium was no longer present in appreciable amounts, and indicates the formation of (V).

An attempt to prepare lithium dihydrobis(triphenylsilyl)aluminate by the reaction of two molar equivalents of triphenylsilyllithium with lithium aluminum hydride was apparently unsuccessful. The yield of triphenylsilane subsequent to carbonation and acid hydrolysis was only 43%. In a similar experiment, using lithium aluminum deuteride, only a 34.6% yield of a mixture of triphenylsilane and triphenylsilane-*d* was obtained upon acid hydrolysis alone. Based on the triphenylsilyllithium used, the yield of triphenylsilane-*d* was 4.3%, the formation of which will be discussed subsequently. The fate of the second equivalent of triphenylsilyllithium used in these reactions was not established.

The hydrolysis reaction of the (triphenylsilyl)aluminum compounds formed in these experiments does not differentiate between (V) and a possible equilibrium product, (VIII), as shown in the following equation. Since it has been shown that lithium tetraphenylaluminate has an initial reactivity greater than triphenylalu-

$$Li[(C_6H_5)_3SiAlH_3] \rightarrow LiH + (C_6H_5)_3SiAlH_2$$
(11)
(VIII)

minum²⁴, and further, that octaphenylcyclotetrasilane is cleaved by phenyllithium but not by the less reactive phenylmagnesium bromide under similar conditions¹⁹, it appears more likely that (V) is the intermediate in the reactions, as it must be of sufficient reactivity to cleave silicon-silicon bonds. This same cleavage of octaphenylcyclotetrasilane, catenation, and cyclization was also brought about by treatment with (V), formed by the complete reaction of hexaphenyldisilane with a one-molar equivalent of lithium aluminum hydride. In addition, octaphenylcyclotetrasilane reacted similarly with the reaction mixture obtained from the addition of triphenylsilyllithium to aluminum hydride, another synthetic approach to (V), as illustrated in equation (12). Although this method was not thoroughly investigated, the results

$$(C_6H_5)_3SiLi + AlH_3 \rightarrow Li[(C_6H_5)_3SiAlH_3]$$
(12)

indicate the formation of (V). The first step in the reaction of (V) with octaphenylcyclotetrasilane could lead to the formation of either (IX) or (X). No products

$$(C_{6}H_{5})_{8}Si_{4} + (V) \rightarrow (C_{6}H_{5})_{2}Si - Si(C_{6}H_{5})_{2} - Si - (C_{6}H_{5})_{3} \text{ or} \\ (C_{6}H_{5})_{2}Si - Si(C_{6}H_{5})_{2} - AIH_{3}Li \\ (IX) \\ (C_{6}H_{5})_{2}Si - Si(C_{6}H_{5})_{2} - H \\ (C_{6}H_{5})_{2}Si - Si(C_{6}H_{5})_{2} - H \\ (C_{6}H_{5})_{2}Si - Si(C_{6}H_{5})_{2} - AIH_{2}[(C_{6}H_{5})_{3}Si]Li \\ (X)$$

containing the triphenylsilyl group bound to silicon could be isolated from the resulting complex mixture of products, which would have established (IX) as the intermediate. Negative evidence suggesting that (IX) is the predominant species was obtained by the isolation of only 50% of triphenylsilane from the reaction involving the preparation of (V) by hydride reduction of hexaphenyldisilane, and only 30% from the experiment wherein triphenylsilyllithium had been added to aluminum hydride. Also, the results of the reactions of triphenylsilyllithium with lithium aluminum hydride indicated that compounds of the type $Li[AIH_2(SiR_3)_2]$ are not readily formed, which supports (IX) as the intermediate, rather than (X).

With regard to the existence and nature of (VI), the reaction of triphenylsilyllithium with aluminum chloride was also investigated briefly. This reaction had been carried out in an earlier study, but the formation of an (organosilyl)aluminum compound was not ascertained²⁵. The addition of triphenylsilyllithium to aluminum chloride in a 1:1 molar ratio resulted in an almost colorless, clear solution. Color Test I²⁶ was essentially negative, indicating the absence of appreciable silyllithium reagent. Hydrolysis in deuteriochloric acid resulted in a 55% yield of a mixture of triphenylsilane-d and triphenylsilane in which the ratio of the deuterated to nondeuterated product was 1:1. Combination of triphenylsilyllithium and aluminum chloride in a 2:1 molar ratio resulted in a mixture which gave a marginally positive Color Test I^{26} . Subsequent to treatment with deuteriochloric acid, a 63% yield of a mixture of triphenylsilane-d and triphenylsilane was isolated, and analysis showed a 1:3 ratio, respectively. When a 3:1 molar ratio of silylmetallic reagent to aluminum chloride was treated similarly, the ratio of triphenylsilane-d to triphenylsilane was 1:10. These results indicate that only the mono-substituted product, (VI), was formed in appreciable quantities, and that the bis- and tris(triphenylsilyl)aluminum chlorides, if formed at all, decomposed to afford triphenylsilane prior to hydrolysis. The fact that deuterated product was obtained does not establish the formation of di- and trisubstituted products in these latter two reactions, since a mono-substituted silvlaluminum compound could arise in these cases.

The amount of non-deuterated triphenylsilane formed in the reaction using one equivalent of triphenylsilyllithium does not necessarily reflect an appreciable instability of (triphenylsilyl)aluminum dichloride, since a portion of the silyllithium compound could have interacted with the (triphenylsilyl)aluminum compound initially formed, resulting in triphenylsilane as in the other two cases.

Attempts to prepare(triphenylsily)aluminum compounds by cleavage of hexaphenyldisilane with aluminum metal in THF or by reaction of chlorotriphenylsilane with aluminum in the same solvent at reflux or at elevated temperatures under pressure were unsuccessful. In the latter cases, none of the easily isolable coupling product, hexaphenyldisilane, was obtained and there was not evidence of triphenylsilane subsequent to hydrolysis.

In the reactions involving the proposed (triphenylsilyl)aluminum compounds, the formation of some triphenylsilane, from sources other than hydrolysis, and hydride was detected. This became evident when the ratio of SiH to SiD in the triphenylsilane obtained from the cleavage by lithium aluminum deuteride followed by hydrolysis in hydrochloric acid was not the theoretical 1:1, but was of the order 1.5:1 of deuterated to non-deuterated product*. Nearly the reverse ratio was observed in the cleavage of hexaphenyldisilane by lithium aluminum hydride, followed by treatment with excess chlorotrimethylsilane and hydrolysis in deuteriochloric acid. The variances from the theoretical ratio may be explained by the fact that hydride/deuteride exchange was demonstrated between triphenylsilane and lithium aluminum deuteride in a separate experiment. Thus, in the cleavage of hexaphenyldisilane by deuteride ion, a larger than the expected amount of deuterated product would result if a portion of the silylaluminum complex were to decompose and afford triphenylsilane which could subsequently exchange with deuteride before hydrolysis. In the cleavage by lithium aluminum hydride, the triphenylsilane resulting from both sources would be constant upon hydrolysis in deuteriochloric acid, with only the remaining silylaluminum compound being deuterated. Similar reactions should explain the formation of the deuterated product from the reaction of triphenylsilvllithium with lithium aluminum deuteride. The mode of decomposition of the silvaluminum compounds was not determined, but may involve the formation of triphenylsilyl radicals and abstraction of hydrogen from solvent. A similar phenomenon has been observed in the reactions of triphenylsilyllithium with certain metallic halides, which has been proposed to result in unstable silylmetallic compounds²⁵. Furthermore, recent studies have indicated that some silylmetallic derivatives of mercury²⁷, antimony, and bismuth²⁸ undergo decomposition affording silicon hydrides. However, lithium(triphenylsily)}triphenylborate, prepared by the reaction between triphenylsilvllithium and triphenylboron, apparently has appreciable stability²⁹.

In the reactions of hexaphenyldisilane with lithium aluminum hydride in tetrahydrofuran, there was always obtained a small amount of diphenylsilane, showing that some cleavage of silicon-phenyl bonds occurred. This was also observed upon treatment of triphenylsilane with lithium aluminum hydride under similar conditions.

^{*} A significant exchange of hydrogen between triphenylsilane and the aqueous media is precluded by the observation that triphenylsilane-*d*, prepared by the reduction of the chloride with lithium aluminum deuteride, followed by hydrolysis in hydrochloric acid, did not contain silicon-hydrogen absorption bands in its IR spectrum³⁵.

In addition to recovered starting material in this case, there was obtained about 3% of diphenylsilane and a lesser amount of tetraphenylsilane. The formation of these products may be illustrated by the following equation:

$$(C_6H_5)_3SiH + LiAlH_4 \rightarrow (C_6H_5)_2SiH_2 + LiAlH_3(C_6H_5) \xrightarrow{(C_6H_5)_3SiH} \rightarrow (C_6H_5)_4Si + LiAlH_4$$

Upon similar treatment, tetraphenylsilane was recovered in a 96% yield, and there was no evidence of products containing silicon-hydrogen groups. Recently, it has been reported that sodium hydride similarly cleaves silicon-carbon bonds when heated in cyclohexane at 200° under high pressure for 10 h³⁰. Similarly, it has been demonstrated that silicon-phenyl bonds readily undergo hydrogenolysis when refluxed in dioxane in the presence of W-7 Raney nickel catalyst³¹.

In addition to the reactions of LiAlH₄ in THF reported herein, some organopolysilanes were treated with the following reducing systems: LiAlH₄ in ether and in pyridine; NaH and LiH in THF; NaH/AlCl₃ in THF; LiAlH₄/2 AlCl₃ in THF; NaBH₄ in dioxane; Zn(Hg)/HCl in a mixture of dioxane and toluene; Na/Hg in acetic acid; HI/P (red) in refluxing xylene; Al(iso-OC₃H₇)₃ in benzene and in THF; Mg in THF: Mg/MgI₂ in THF; and Al in THF. In the case of sodium hydride in THF, octaphenylcyclotetrasilane was converted into decaphenylcyclopentasilane in high yield after 20 h at reflux. Lithium hydride, after 72 h at reflux in THF, resulted in only a 3% yield of the cyclic pentasilane. None of the other treatments gave evidence of reductive cleavages. These reactions are summarized in Tables 5 and 6.

EXPERIMENTAL

All reactions were run in oven-dried glassware under atmospheres of oxygenfree, dry nitrogen. The tetrahydrofuran (THF) was dried and purified by refluxing over sodium for at least 24 h, followed by distillation into lithium aluminum hydride

CLEAVAGE OF ORGANOPOLYSILANES BY LITHIUM ALUMINUM HYDRIDE IN THF									
Compound		Mole ratio compd.: LiAlH4	Reflux (h)	Yield of $(C_6H_5)_{2n}Si_nH_2^a$ (%)				Yield of	
	(g)			$\overline{n=1}$	n=2	n=3	n=4	n=5	(C ₆ H ₅)10 ^{Si5^e} (%)
$(C_6H_5)_8Si_1^b$	10	4:1	c	d	ď		3		64
(-0++3/80+4	10	4:1	c	đ	đ		6		62
	7.3	1:1	1.5	đ	đ		16.5	8.2	16.5
(C6H5)10Si5"	10.9	4:1	0.2 ^r	đ	đ		6		74
(-05)105	10.9	1:1	41	đ	10		7		9
	10.9	1:1	1.5	đ	2	1	5	2	68
	10.9	1:2	44	16 ⁹	34		10		
(C ₆ H ₅) ₈ Si ₄ Cl ₂ ^a	9.1	1:1	24	12	26	14.4	14		
$(C_6H_5)_8Si_4H_2^a$	2	1:1	2	7.5 ^g	10	30	5		

TABLE 1

^a Linear polysilanes. ^b Octaphenylcyclotetrasilane. ^c The reaction mixture was stirred at room temperature for 20 h. ^d No attempt was made to isolate these materials. ^c Decaphenylcyclopentasilane. ^f Refluxed for 12 min after stirring at room temperature for 18 h. ^g Obtained on distillation under reduced pressure.

and redistillation from the hydride immediately before use. All temperatures recorded are uncorrected. Products were identified by comparison of physical properties and infrared spectra with known samples. Mixed-melting point determinations were carried out on solid products.

Organopolysilanes and LiAlH₄

The reagents, reaction conditions, and results are listed in Table 1. In general, ca. 100 ml of THF was used in each case. The resulting reaction mixtures were hydrolyzed by cautiously pouring onto excess dilute acid. Subsequent to the usual work-up, products were isolated either by distillation under reduced pressure or by crystallization. The yields reported in Table 1 are for pure products, many of which were obtained only after repeated fractional recrystallizations.

Organodisilanes and LiAlH₄

The organodisilanes listed in Table 2 were treated with $LiAlH_4$ in 1:1 mole ratios using *ca.* 100 ml of THF in each run. Subsequent to hydrolysis in excess acid and the customary work-up, the products were isolated by distillation. The reaction conditions and results are shown in Table 2.

INDLIG	TABLE	2
--------	-------	---

CLEAVAGE OF ORGANODISILANES BY LITHIUM ALUMINUM HYDRIDE IN THF

Compound		Reflux	Treatment	Products		
	(g)	(h)			(%)	
(C ₆ H ₅) ₆ Si ₂ ^a	10.4	72	Acid hy- drolysis	(C ₆ H ₅) ₃ SiH	60	
[(CH ₃)(C ₆ H ₅) ₂ Si] ₂	6.0	67	Acid hy- drolysis	(CH ₃)(C ₆ H ₅) ₂ SiH ^b	40	
[(CH ₃) ₂ (C ₆ H ₅)Si] ₂	10.0	65	Acid hy- drolysis	(CH ₃) <u>2(</u> C ₆ H ₅)SiH (impure) ^e ; [(CH ₃)2(C ₆ H ₅)Si]2	23 41.5	
(CH ₃) ₃ SiSi(C ₆ H ₅) ₃	6.0	61	Carbon- ation, acid hydrolysis	$(CH_3)_3(C_6H_5)_3$ $(C_6H_5)_3$ SiH; $(C_6H_5)_3$ SiOH	50 63.9 14	
$(C_2H_5)_6Si_2$	12.5	6567	Acid hy- drolysis	$(C_{2}H_{5})_{3}SiH$ $(C_{2}H_{5})_{6}Si_{2}$	0° 72.8°	

^a A representative sample; see Table 3 for more complete information. ^b B.p. 107–110° (2.3 mm), n_D^{20} 1.5717; reported³⁶ n_D^{20} 1.5717. ^c Collected in three fraction, b.p. 50–70° (50–70 mm), with n_D^{20} values ranging from 1.4928–1.4988; reported³⁷ n_D^{20} 1.4988. ^d Collected prior to hydrolysis by means of a trap cooled to -65° , which was attached to the end of the reflux condenser. Identification was made by its IR spectrum as solutions in carbon disulfide and carbon tetrachloride. ^e The results of two runs. The solvent was carefully distilled through an 18″ spinning band column, rated at 23 theoretical plates. No evidence of triethylsilane was detected during the distillation or in the IR spectra of any of the distillation fractions.

Hexaphenyldisilane and $LiAlH_4$

The amounts of reagents, reaction conditions, subsequent treatments, and results are given in Table 3. In general, the reactions were carried out by simply mixing a 1:1 mole ratio of hexaphenyldisilane and LiAlH₄ in *ca*. 100 ml of THF and bringing to reflux the recorded length of time. The reaction mixtures usually took on a

CLEAVAGE OF ORGANOPOLYSILANES BY LIAIHA

(C ₆ H ₅) ₆ Si ₂ (g)	Reflux (h)	Treatment	Yield of (C ₆ H ₅)₃SiH ^b	• Other products ^e		
			(%)		(%)	
5.2	48	(CH ₃) ₃ SiCI (excess); acid hydrolysis	60			
10.4	72	Acid hydrolysis	65.6	(C6H5),SiH2	3	
10.4	24	Carbonation ^d ; acid hydrolysis	62.0 ^e	$(C_6H_5)_6Si_2$	10	
10.4	60	Acid hydrolysis	60.1	$(C_{6}H_{5})_{2}SiH_{2}$	2	
6.2	24	(LiAlD _{\downarrow} was used) hydrolyzed in H ₃ O ⁺	31 ¹ (62) ⁹	(C ₆ H ₅) ₃ SiD	44.3 ¹ (88.6) ^g	
10.4	96	reflux with excess	47 ^{e,f} (94) ^g	(C ₆ H ₅) ₃ SiD	31 ^{e.f} (62) ^g	
		(CH ₃) ₃ SiCl; hydro- lized in D ₃ O ⁺		$(C_6H_5)_6Si_2$	36.5	
5.2	26	$(C_6H_5)_8Si_4^{+}$, 7.2 g stirred at room temp. (36 h), acid hydrolysis	52	$(C_6H_5)_8Si_4^h$ $(C_6H_5)_{10}Si_5^i$ polysilanes ^k	45.2 55 ^j	

TABLE 3

CLEAVAGE OF HEXAPHENYLDISILANE BY LITHIUM ALUMINUM HYDRIDE IN THF®

^a The reactions were carried out as described in the experimental part, with the mole ratio of reactants being 1:1 in each case. ^b Yield based on two moles of product from one mole of disilane. ^c In addition to the products listed, distillation residues yielded varying amounts of viscous oils, which resisted attempts to purify. The IR spectra indicated the presence of SiOH and SiOSi groups. ^d Carbonation was accomplished by pouring the reaction mixture onto a slurry of Dry-Ice and ether, and allowing to warm to room temperature before hydrolyzing. ^e The yield was based on unrecovered starting material. ^J See the experimental part for the analysis of the mixture of $(C_6H_5)_3$ SiH and $(C_6H_5)_3$ SiD. ^g The yield in parentheses is based on one mole of disilane forming one mole each of $(C_6H_5)_3$ SiH and $(C_6H_5)_3$ SiD. ^h Octaphenylcyclotetrasilane. ⁱ Decaphenylcyclopentasilane. ^j Yield based on unrecovered octaphenylcyclotetrasilane. ^k Two additional solids were isolated. One, 0.5 g, m.p. 140–150° was obtained as crystals before distillation; the second, 0.7 g, m.p. 67–80°, was recovered from the distillation residue. Attempts to purify these materials were unsuccessful. Their IR spectra indicated the presence of silicon–phenyl and silicon–hydrogen groups.

greyish-green coloration. In one case, the grey lithium aluminates present in the lithium aluminum hydride were removed by filtration of a THF solution through a sintered-glass funnel prior to mixing with hexaphenyldisilane. After 24 h of reflux, an almost clear, light-yellow solution resulted. Subsequent to the treatment as given in Table 3 and hydrolysis, the mixtures were filtered to remove any unreacted hexaphenyldisilane. Following the usual work-up, the products were distilled under reduced pressure. Usually, a small amount of diphenylsilane was obtained at about 60° (0.3 mm) which was identified by IR spectral analyses and by its characteristic odor. Triphenylsilane was normally collected at temperatures of *ca*. 125–130° (0.16 mm), and solidified on cooling, m.p. 44–46°.

Triphenylsilyllithium and aluminum compounds

Triphenylsilyllithium was prepared in THF by cleavage of hexaphenyldisilane by lithium³². Subsequent to decantation from the excess lithium, the solution was added dropwise to a stirred solution of the aluminum compounds in THF. The triphenylsilane was isolated by distillation under reduced pressure. The reactants, molar quantities, reaction conditions and results are shown in Table 4.

TA	BL	F	4
ເກ	.DL	L	-

(C6H5)3SiH Other products (C6H4)3SiLi^a Aluminum Treatment compound (mmoles) (%) (%) (mmoles) (C6H3)3SiOH 2.4 30^b LiAlH. Stirred at room temp. (3h), added 68 (CH₃)₃SiCl, (160 mmole), in ether, (30) reflux (1 h), acid hydrolysis 30* 78.4 LiAlH₁ Stirred at room temp. (3 h), carbonated, acid hydrolysis (30) 40 LiAlH₄ Stirred at room temp. (72 h), 43 carbonated, acid hydrolysis (20)(C₆H₅)₃SiD^c 4.4 10 LiAID. Stirred at room temp. (7 h), 30.2 (5) hydrolyzed in dil. HCl AIH.ª 30.8 (C₆H₅)₈Si₄[€] 11.7 20 Stirred at room temp. (1 h), followed by stirring with 48.8^f (20) $(C_6H_5)_{10}Si_5^{f}$ (C₆H₅)₈Si₄^e (20 mmole), at room temp. (30 h); acid hydrolysis AlCly Stirred at room temp. (15 min)^h, 28 $(C_6H_5)_3SiD$ 27 14 hydrolyzed in dil. DCl (14)AIC13 Hydrolyzed in dil. DCl after 50 (C6H3)3SiD 15 14 15 min (7) AICI₃" 2^j 21 Hydrolyzed in dil. DCl^j 20^j (C6H5)3SiD (7)

REACTIONS BETWEEN TRIPHENYLSILYLLITHIUM AND ALUMINUM COMPOUNDS

^a Based on the amount of hexaphenyldisilane used in the preparation. ^b Originally, 60 mmoles each of $(C_6H_5)_5$ SiLi and LiAlH₄ were combined together. The resulting mixture was divided in half and each portion treated individually as described in the first two experiments. ^c IR spectra on the distillation residues indicated the presence of Si-phenyl, Si-OH, and Si-O-Si groups. ^d Prepared from 15 mmoles of LiAlH₄ and 5 mmoles of AlCl₃, according to the method of Wiberg³⁸. ^c Octaphenylcyclotetrasilane. ^f Decaphenyl-cyclopentasilane; yield based on unrecovered (C_6H_5)₈Si₄. ^g Freshly sublimed; dissolved in THF at -30° to prevent interaction between solvent and AlCl₃. ^h The resulting light-pink solution gave a negative or very weakly positive Color Test I²⁶. ⁱ Prior to hydrolysis, the reddish-brown mixture gave a marginally positive Color Test I²⁶. Job Victor Test I²⁶ was essentially negative after 15 min. Aprox. one-fourth of the solution was withdrawn and hydrolyzed in dil. DCl; the ratio of deuterated to non-deuterated triphenylsilane was about 1:10, resp. Octaphenylcyclotetrasilane had been added to the rest of the reaction mixture, which was subsequently refluxed for 24 h; following hydrolysis in dil. DCl, octaphenylcyclotetrasilane was recovered nearly quantitatively.

Perphenylated cyclic polysilanes and other metallic hydrides

Various other metallic hydride systems were used in attempts to effect siliconsilicon bond reductions. The reagents, reaction conditions, and results are listed in Table 5. The reaction mixtures were all hydrolyzed in crushed ice containing excess dilute acid. Any solids were filtered off, washed with benzene, and dried. The soluble materials were worked up in the usual manner.

Organopolysilanes and various reducing agents (attempted)

The reagents, reaction conditions and results are given in Table 6.

Analyses of mixtures of triphenylsilane and triphenylsilane-d

IR analyses of mixtures of triphenylsilane and triphenylsilane-d were carried

J. Organometal. Chem., 15 (1968) 43-56

CLEAVAGE OF ORGANOPOLYSILANES BY LIAIH4

5	3
	-

Compound		Reducing	Reaction conditions	Starting	Products ^b		
(g)	(g)	agent		material (%)		(%)	
$(C_6H_5)_8Si_4$	10	LiAIH	Reflux 7 days in ether	98.7			
,	10	NaH	Reflux 20 h in THF	0	(C ₆ H ₅) ₁₀ Si ₅	77	
	10	LiH	Reflux 72 h in THF	88	(C6H5)10Si5	3	
	10	NaBH₄	Reflux 5 days in dioxane	64 [.]	Siloxane ^b	ca. 20	
	10	NaH/AlCl	Reflux 72 h in THF	97			
	10	LiAlH ₄ /2AlCl ₃	Reflux 6 days in THF	95			
(C ₆ H ₅) ₁₀ Si ₅	10.9	LiAlH	Reflux 6 days in ether	99			
(-0-5)[0-5	10.9	NaH	Reflux 18 h in THF	93.2			
	10.9	LiH	Reflux 72 h in THF	92.8			
	10.9	NaBH ₄	Reflux 5 days in dioxane	96			
	10.9	LiAIH /2AICI	Reflux 25 h in THF	95.8			
$(C_6H_5)_6Si_2$	5.2	LiAIH4	Reflux 54 h in pyridine	96			

TABLE 5

ATTEMPTED REDUCTION OF PERPHENYLATED POLYSILANES WITH METALLIC HYDRIDES⁴

^a The mole ratio of reactants was 1:1 in each case. ^b The IR spectrum of the remaining material, m.p. range 170–190°, showed bands characteristic of siloxane groups, but no silicon hydrogen groups. Material having a similar m.p. range and IR spectrum was also obtained during an attempted recrystallization of octaphenyl-cyclotetrasilane from boiling dioxane.

TABLE 6

ATTEMPTED REDUCTIVE CLEAVAGES OF PERPHENYLATED ORGANOPOLYSILANES BY VARIOUS REDUCING AGENTS

Compound ^e	Reducing agent	Mole ratio compound red. ag.	Reactions conditions	Recovery (%)
(C ₆ H ₅) ₈ Si₄	Zn(Hg)/HCl	7:100	Reflux until H_2 evol. ceased (16 h) in a suspension of conc. HCl, toluene and dioxane	90
(C ₆ H ₅) ₁₀ Si ₅	Zn(Hg)/HCl	7:100	Reflux until H_2 evol. ceased (16 h) in a suspension of conc. HCl, toluene, and dioxane	98
(C ₆ H ₅) ₈ Si ₄	3% Na(Hg)/HC ₂ H ₃ O ₂	1:10	Room temp. until H_2 evol. ceased (8 h)	98
(C ₆ H ₅) ₁₀ Si ₅	3% Na(Hg)HC ₂ H ₃ O ₂	1:10	Room temp. until H_2 evol. ceased (8 h)	90
(C ₆ H ₅) ₈ Si₄	H1/P(red)	1:2	Reflux in xylene (64 h)	0°
(C6H5)10Si5	HI/P(red)	1:2	Reflux in xylene (64 h)	71 ⁶
(C ₆ H ₅) ₈ Si ₄	$Al(iso-OC_3H_7)_3$	1:1	Reflux (48 h) in benzene	96
(C ₆ H ₅) ₈ Si ₄	Mg ^c	1:10	Reflux (5 days) in THF	96
(C ₆ H ₅) ₈ Si ₄	Mg/MgI ₂ ^d	1 :10	Reflux (30 h) in THF	96
$(C_6H_5)_6Si_2$	Mg/Mgl ₂ ^d	1 :10	Stirred with a few ml of THF (20 h), reflux with 60 ml THF (24 h)	99
(C ₆ H ₅) ₆ Si ₂	Al	1:4	Reflux (5 days) in THF	.99

^a Five grams of compound was used in each case. ^b There was a considerable amount of material admixed with the red phosphorus, which was insoluble in organic solvents. An IR spectrum as a KBr pellet showed no absorption bands for organic groups. The material was apparently a mixture of silica and silicic acid. Refluxing octaphenylcyclotetrasilane with conc. HI in xylene for 64 h produced a similar material which was analyzed for silicon. (Found : Si, 42.39, 41.11. SiO₂ calcd.: Si, 46.72%).^c Powdered Mg, which had been washed successively with dil. HCl, ethanol, and ether, followed by drying at 130° for 3 h. ^d Prepared by the reaction of iodine with magnesium turnings in ether, using a 1:2 g-atom ratio, respectively. The ether was removed by distillation and replaced with THF.

J. Organometal. Chem., 15 (1968) 43-56

out on a Perkin Elmer "Model 21" double-beam spectrophotometer using a 0.5 mm, sodium chloride cell. The samples were analyzed for triphenylsilane-*d* using the absorption band at 6.45 microns, which is characteristic of silicon-deuterium bonds³³. The solvent used was Eastman spectral grade bromoform. The determinations were carried out at 25° . The results were an average of at least three runs. Conformity with Beer's law was established by the standardization plot, which was a good straight line nearly passing through the origin. The analysis of a prepared mixture containing 12.2 mg/ml of triphenylsilane-*d* and 40.0 mg/ml of triphenylsilane gave a result of 12.8 mg/ml, a value 5% higher than the theoretical. The amount of triphenylsilane in the mixtures was calculated by subtracting the quantity of triphenylsilane-*d* from the total weight of the sample. The results are given with each individual case.

Triphenylsilane and LiAlH₄

A mixture of 10.0 g (0.04 mole) of triphenylsilane (free of di- and tetraphenylsilane) and 1.5 g (0.04 mole) of LiAlH₄ was refluxed in *ca.* 100 ml of THF for 46 h. Subsequent to hydrolysis by slowly pouring upon a mixture of crushed ice and dilute acid, and the usual work-up, the products were distilled under reduced pressure to give 0.2 g (2.8%) of impure diphenylsilane, b.p. 60–70° (0.15 mm), n_D^{20} 1.5828 (lit. value³⁴, n_D^{20} 1.5975), identified by its IR spectrum and characteristic odor. Further distillation afforded 8.5 g (85%) of recovered starting material, b.p. 125–130° (0.17 mm), m.p. 44–46° (mixed m.p.). The distillation residue partially solidified on cooling. The solid material was washed with petroleum ether (b.p. 60–70°) and recrystallized from ethyl acetate to give 0.2 g (1.5%) of tetraphenylsilane, m.p. 232–235° (mixed m.p.).

Similar results were obtained in a second run under the same conditions.

Triphenylsilane and LiAlD₄

A mixture of 5.2 g (0.02 mole) of triphenylsilane and 0.84 g (0.02 mole) of LiAlD₄ was refluxed in 65 ml of THF for 24 h. Subsequent to hydrolysis and work-up as described in the preceding experiment, there was obtained 0.1 g (2.8%) of impure diphenylsilane, b.p. 40–50° (0.17 mm) and 4.1 g (78%) of triphenylsilane, b.p. 125–130° (0.17 mm), m.p. 42–45° (mixed m.p.). The IR spectra of these materials contained strong absorption bands for both silicon-hydrogen and silicon-deuterium bonds. From the distillation residue, there was obtained 0.1 g (1.5%) of tetraphenylsilane, m.p. 232–235° (mixed m.p.).

Tetraphenylsilane and LiAlH₄

A solution of 10.0 g (0.03 mole) of tetraphenylsilane and 1.5 g (0.4 mole) of LiAlH₄ in 100 ml of THF was refluxed for 89 h. The reaction mixture was poured onto a slurry of Dry Ice and ether. After allowing the carbonation mixture to warm to room temperature, dilute acid was added, and the solids filtered. The solids were washed with ether and dried to give 9.3 g (93%) of tetraphenylsilane, m.p. 234-236° (mixed m.p.). The ether washings and the filtrate were combined and extracted with 5% sodium hydroxide solution. The aqueous layer was acidified with conc. hydrochloric acid, and extracted with ether. Evaporation of the ether gave only a trace of solid, m.p. 222-228°, identified as impure tetraphenylsilane by its IR spectrum. An additional 0.3 g (3%) of tetraphenylsilane was obtained from the original organic layer. The small

amount of residue obtained from evaporation of the mother liquor did not show absorption bands for SiH, SiOH or SiOSi groups.

Chlorotriphenylsilane and aluminum (attempted)

Run I. A mixture of 6.0 g (0.02 mole) of chlorotriphenylsilane and 1.0 g (0.04 g-atom) of freshly turned aluminum metal chips was refluxed in 25 ml of THF for 5 days. The reaction mixture was hydrolyzed in excess dilute acid. After hydrogen evolution ceased, there remained no insoluble material, *i.e.*, hexaphenyldisilane. Subsequent to the usual work-up, the solvents were removed by distillation and petroleum ether (b.p. 60-70°) added. The solids were removed by filtration and the filtrate chromatographed on an alumina column. Evaporation of the solvent from the petroleum ether eluate left only a trace of material, the IR spectrum of which did not show the presence of triphenylsilane. The petroleum ether insoluble material was identified as triphenylsilanol, resulting from hydrolyzed starting material.

Run 2. The same quantities of starting materials and solvent which were used in Run 1, were heated at $140-150^{\circ}$ for 18 h and at $220-250^{\circ}$ for 3 h in a steel bomb, agitated by a rocking assembly, under nitrogen pressure (initially at 500 p.s.i.g. at room temp.). The work-up and results were identical to those described in Run 1.

ACKNOWLEDGEMENT

This research was supported in part by the United States Air Force under Contract AF 33(616)-6463 monitored by Materials Laboratory, Directorate of Laboratories, Wright Air Development Center, Wright-Patterson AFB, Ohio. Samples of lithium aluminum deuteride were obtained through the generosity of Dr. CHARLES W. HEITSCH, who also gave of his time in discussions on experimental details. The infrared spectral analyses were performed by Mrs. IKUE OGAWA.

REFERENCES

- 1 H. GILMAN AND H. J. S. WINKLER, in H. ZEISS (Ed.), Organometallic Chemistry, Reinhold, New York, 1960, pp. 270–345; D. WITTENBERG AND H. GILMAN, Quart. Rev. (London), 13 (1959) 116.
- 2 H. GILMAN, R. K. INGHAM AND A. G. SMITH, J. Org. Chem., 18 (1953) 1743.
- 3 H. GILMAN AND G. D. LICHTENWALTER, unpublished studies, reported in ref. 1.
- 4 D. WITTENBERG AND H. GILMAN, unpublished studies, reported in ref. 1.
- 5 C. A. KRAUS AND W. K. NELSON, J. Amer. Chem. Soc., 56 (1934) 195.
- 6 M. P. BROWN AND G. W. A. FOWLES, J. Chem. Soc., (1958) 2111.
- 7 H. GILMAN AND M. B. HUGHES, unpublished studies, reported in ref. 1.
- 8 H. GILMAN AND A. G. SMITH, unpublished studies, reported in ref. 1.
- 9 H. GILMAN AND T. C. WU, unpublished studies, reported in ref. 1.
- 10 U. G. STOLBERG, Z. Naturforsch., B, 18 (1963) 765; Angew. Chem. 75 (1963) 206.
- 11 H. GILMAN AND R. A. TOMASI, J. Organometal. Chem., 9 (1967) 223.
- 12 T. C. WU AND H. GILMAN, J. Org. Chem., 23 (1958) 913.
- 13 H. G. GUTOWSKY AND E. O. STEJSKAL, J. Chem. Phys., 22 (1954) 939.
- 14 A. E. FINHOLT, A. G. BOND, JR., K. E. WILZBACH AND H. I. SCHLESINGER, J. Amer. Chem. Soc., 69 (1947) 2692.
- 15 M. A. RING AND D. M. RITTER, J. Amer. Chem. Soc., 83 (1961) 802; J. A. MORRISON AND M. A. RING, Inorg. Chem., 6 (1967) 100.
- 16 W. A. INGLE, E. A. GROSCHWITZ AND M. A. RING, Inorg. Chem., 6 (1967) 1429.
- 17 H. GILMAN, D. J. PETERSON, A. W. P. JARVIE AND H. J. S. WINKLER, J. Amer. Chem. Soc., 82 (1960) 2076.

- 18 H. GILMAN AND G. L. SCHWEBKE, J. Amer. Chem. Soc., 86 (1964) 2693.
- 19 A. W. P. JARVIE AND H. GILMAN, J. Org. Chem., 26 (1961) 1999.
- 20 H. GILMAN, A. W. P. JARVIE AND H. J. S. WINKLER, unpublished studies.
- 21 H. GILMAN, G. L. SCHWEBKE AND K. Y. CHANG, unpublished studies.
- 22 A. G. BROOK AND H. GILMAN, J. Amer. Chem. Soc., 77 (1955) 2322.
- 23 G. D. BARBARAS, C. DILLARD, A. E. FINHOLT, T. WARTIK, K. E. WILZBACH AND H. I. SCHLESINGER, J. Amer. Chem. Soc., 73 (1951) 4585; T. WARTIK AND H. I. SCHLESINGER, J. Amer. Chem. Soc., 75 (1953) 835.
- 24 G. WITTIG AND G. KEICHER, Naturwissenschaften, 34 (1947) 216; G. WITTIG AND O. BUB, Justus Liebigs Ann. Chem., 566 (1950) 113.
- 25 M. V. GEORGE, G. D. LICHTENWALTER AND H. GILMAN, J. Amer. Chem. Soc., 81 (1959) 978.
- 26 H. GILMAN AND F. SCHULTZ, J. Amer. Chem. Soc., 47 (1925) 2002.
- 27 C. A. EABORN, R. A. JACKSON AND R. W. WALSINGHAM, Chem. Commun., 14 (1965) 300; A. G. BEAUMONT, C. EABORN, R. A. JACKSON AND R. W. WALSINGHAM, J. Organometal. Chem., 5 (1966) 297; R. FIELDS, R. N. HASZELDINE AND R. E. HATTON, J. Chem: Soc., (1967) 2559.
- 28 N. S. VYAZANKIN, O. A. KRUGLAYA, G. A. RAZUVAEV AND G. S. SEMCHIKOVA, Dokl. Akad. Nauk SSR, 166 (1966) 99; Chem. Abstr., 64 (1966) 11245; J. Organometal. Chem., 6 (1966) 474.
- 29 D. SEYFERTH, G. RAAB AND S. O. GRIM, J. Org. Chem., 26 (1961) 3034.
- 30 K. RÜHLMANN AND H. HEINE, Z. Chem., 6 (1966) 421.
- 31 G. D. F. JACKSON, W. H. F. SASSE AND K. O. WADE, J. Organometal. Chem., 3 (1965) 177; W. H. F. SASSE AND K. O. WADE, Aust. J. Chem., 19 (1966) 1892.
- 32 H. GILMAN AND G. D. LICHTENWALTER, J. Amer. Chem. Soc., 80 (1958) 608.
- 33 R. N. KNISELY, V. A. FASSEL AND E. CONRAD, Spectrochim. Acta, 13 (1959) 651.
- 34 R. A. BENKESER, H. LANDESMAN AND D. J. FOSTER, J. Amer. Chem. Soc., 74 (1952) 648.
- 35 E. A. ZUECH, unpublished studies.
- 36 V. F. MIRONOV AND A. D. PETROV, Izv. Akad. Nauk SSSR, Otd. Khim. Nauk. (1957) 383; Chem. Abstr., 51 (1957) 15457.
- 37 G. A. RUSSELL, J. Org. Chem., 21 (1956) 1190.
- 38 E. WIBERG, Angew. Chem., 65 (1953) 16.
- J. Organometal. Chem., 15 (1968) 43-56