

Preliminary communication

STABILIZATION OF A CARBANION BY α -Me₃Si GROUPS. THE KINETIC ACIDITY OF TRIS(TRIMETHYLSILYL)METHANE

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Summary

In Me₂SO/H₂O/KOH at 50°C, (Me₃Si)₃C³H undergoes detritiation about 5-7 times as rapidly as Ph₃C³H, indicating that α -Me₃Si groups substantially stabilize carbanions.

The metallation which occurs at C—H bonds α to Me₃Si groups [1,2] can reasonably be attributed to some stabilization of the carbanions by ($p \rightarrow d$) _{π} interaction with the silicon atom. Such stabilization was considered to contribute to the low reactivity of (Me₃Si)₃CLi [2], but steric effects probably dominate in this case. Steric effects also account for the fact that additional α -Me₃Si groups hinder the base cleavage of a C—SiMe₃ bond [e.g. PhCH(SiMe₃)₂ is less reactive than PhCH₂SiMe₃] in spite of the expected additional stabilization of the separating carbanion [3].

Recently *ab initio* calculations in this School indicated that the α -H₃Si group should substantially stabilize a carbanion, possibly more so than an α -Ph group [4], and there is recent indirect evidence from preparative studies that an α -Me₃Si does stabilize more than an α -Ph group [5]. To obtain direct information we have now compared the kinetic acidities of (Me₃Si)₃CH and Ph₃CH.

With Ph₃C³H, detritiation in a mixture of Me₂SO(6 vol) and 1.0 M aqueous KOH (1 vol) at 50°C gave a good first-order plot to more than 75% completion of reaction, with a residual count of <5% of the initial count; the first order constant is $3.6 \times 10^{-5} \text{ s}^{-1}$. With (Me₃Si)₃C³H, in a period equal to 25 times that for loss of the first 25% count the count fell by only 75%, i.e. there was an unexpected residual activity of ca. 25%, though this "residual" count continued to fall slowly. The residual count of ca. 25% could in principle be due to contamination of the (Me₃Si)₃C³H by material tritiated in a Me₃Si group, and effectively inert. A first order plot based on this extreme assumption is linear to about

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70% reaction and corresponds with a rate constant of $2.3 \times 10^{-4} \text{ s}^{-1}$. However, GLC analysis showed that $(\text{Me}_3\text{Si})_2\text{CH}_2$ was formed, and the residual count is probably mainly due to conversion of $(\text{Me}_3\text{Si})_3\text{C}^3\text{H}$ into $(\text{Me}_3\text{Si})_2\text{CH}^3\text{H}$ at a rate equal to about one-quarter of that of the exchange. On this assumption and the further simplifying assumption that the $(\text{Me}_3\text{Si})_2\text{CH}^3\text{H}$ is effectively inert to exchange, the first order plot is linear to about 65% of reaction and corresponds to a rate constant of $1.9 \times 10^{-4} \text{ s}^{-1}$. If the residual count is ignored, and the assumption made that the count is tending to zero, the first order plot is linear over only about 35% of reaction, and the corresponding rate constant is ca. $1.8 \times 10^{-4} \text{ s}^{-1}$. Probably the $(\text{Me}_3\text{Si})_3\text{C}^3\text{H}$ both contained inert tritiated material and gave $(\text{Me}_3\text{Si})_2\text{CH}^3\text{H}$, with the latter undergoing detritiation at a significant rate, but further analyses seems unprofitable, since the uncertainties do not cast doubt on the conclusion that $(\text{Me}_3\text{Si})_3\text{C}^3\text{H}$ undergoes exchange about 5-7 times as fast as $\text{Ph}_3\text{C}^3\text{H}$.

This evidence that the kinetic acidity of $(\text{Me}_3\text{Si})_3\text{CH}$ is greater than that of Ph_3CH does not necessarily mean that the equilibrium acidities would be in this order, though this is likely. Furthermore, since for steric reasons the 3 phenyl groups in the Ph_3C^- ion are thought to be unable to exert their maximum delocalizing potential, the acidity of Me_3SiCH_3 would not necessarily be greater than that of PhCH_3 . It is clear, however, that $\alpha\text{-Me}_3\text{Si}$ groups can substantially stabilize carbanions.

The fairly ready cleavage of $(\text{Me}_3\text{Si})_3\text{CH}$ to $(\text{Me}_3\text{Si})_2\text{CH}_2$ is itself a reflection of the influence of the Me_3Si groups in stabilizing the $(\text{Me}_3\text{Si})_2\text{HC}^-$ anion.

Experimental

Materials

The sample of $\text{Ph}_3\text{C}^3\text{H}$ was made by adding tritiated water to Ph_3CLi obtained by treating Ph_3CH with $n\text{-BuLi}$ in ether/THF. The product was recrystallized from methanol to constant activity (1580 c.p.s./mg).

The $(\text{Me}_3\text{Si})_3\text{C}^3\text{H}$ (activity 8680 c.p.s./mg) was made analogously from $(\text{Me}_3\text{Si})_3\text{CLi}$ obtained by metallation of $(\text{Me}_3\text{Si})_3\text{CH}$ with MeLi in THF/ Et_2O [2]. It had b.p. 216°C and GLC revealed no impurity.

Rate measurements

A solution of the tritiated material (ca. 5 mg) in Me_2SO (30 ml) at 50°C was mixed with 1.0 *M* aqueous KOH (5 ml) and the mixture was placed in a thermostat at 50.0°C . At appropriate intervals 1 ml aliquots were withdrawn and added to a mixture of water (20 ml) with toluene (10 ml) containing a scintillator [6]. The toluene layer was washed with water ($3 \times 20 \text{ ml}$) and dried, and a 5 ml sample taken for counting.

For $(\text{Me}_3\text{Si})_3\text{C}^3\text{H}$, a typical pattern of the variation with time (min. unless otherwise indicated) of the percentage count (in parentheses) was as follows: 0 (100); 10(90); 20(80); 30(72.5); 50(60); 60(55); 70(51); 80(48); 100(44); 130(41); 6 h(27); 24 h(25); 48 h(23). If the residual activity is due to formation of $(\text{Me}_3\text{Si})_2\text{CH}^3\text{H}$ at a rate 0.25 that of the exchange, when the count has fallen to $(100 - x)\%$, the proportion, P , of $(\text{Me}_3\text{Si})_3\text{C}^3\text{H}$ which has lost its tritium is given by $(100 - x)/(100 - 0.25x)$; a plot of $\log(1 - P)$ against time is linear

down to $P = 0.35$ (i.e. 65% exchange).

Products

GLC analyses (5% Carbowax M) of the mixture obtained from $(\text{Me}_3\text{Si})_3\text{CH}$ after about 15 h revealed that in addition to $(\text{Me}_3\text{Si})_3\text{CH}$ some $(\text{Me}_3\text{Si})_2\text{CH}_2$ was present, this was identified by its retention time and mass spectrum in a linked GLC-mass spectrometer system. [The peak of highest mass was at 145 ($M - 15$) (strong), just as that from $(\text{Me}_3\text{Si})_3\text{CH}$ was at 217 ($M - 15$)].

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