Structure Effects of Ion Pair Separation: Planar and Pyramidal Sulfur- and Silicon-Substituted Carbanions¹

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The conversion of a lithium reagent to a free carbanion (contact to separated ion pairs) is expected to cause structural changes such as decreased pyramidalization, 22 decreased inversion barriers at carbon, and increased charge delocalization to substituent groups.2b Meaningful comparisons have not been possible since many well-studied lithium reagents, such as those substituted by strong electron-attracting groups (α -phosphoryl-, $^3\alpha$ -sulfinyl-, and α-sulfonylalkyllithiums⁴ and lithium enolates⁵) coordinate through oxygen and not carbon and others are so delocalized (benzhydryl, trityl, fluorenyl, etc.) that C-Li association is weak.^{2a,6} We know of only one system for which X-ray data are available for both a contact and solvent-separated ion pair: Ph₃CLi(TMEDA)^{7a} and Ph₃CLi(12-crown-4)₂.7b The contact ion is slightly pyramidalized, the separated ion planar. Although the effects of THF and more strongly coordinating media (crown ethers, HMPA) on reactivity8 and solution structure of many organolithium reagents have been studied, the extent of ion separation is usually unknown.4a,3b,8c,9

The discovery^{1a,b,10} that solvation and ion pair status of many lithium reagents in THF/HMPA can be reliably determined by NMR spectroscopy has made it possible to make a detailed correlation between lithium coordination and carbanion geometry.

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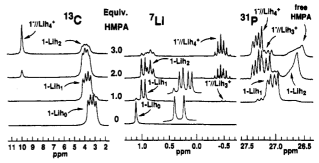


Figure 1. HMPA titration of 0.13 M 1-Li in 3:2 THF/ether at -132 °C. The 7Li and 31P spectra are at natural isotopic abundance. Insets in the ⁷Li spectra at 0 and 1 equiv and the ¹³C signals of C-2 are from ¹³Cenriched samples of 1. 7Li shifts were referenced to external 0.3 M LiCl

Scheme I

Table I. ¹³C δ and J_{CH} Values in THF/Ether at -100 °C

		$\delta_{ m ipso}$		$\delta_{ m para}$			
reagent	CX	S	Si	S	Si	δ_{CX}	¹ <i>J</i> _{CH}
S H	CH	139.2		126.2		35.9	156
	C–Li	150.2		122.9		27.8	123
S'2X	C-//Li+	152.0		120.8		27.8	123
PhS SPh	CH	136.0		126.5		35.9	154
Χ ,	C-Li	150.2		121.9		20.3	141
н x 4	C-//Li+	158.6		120.4		4.3	183
PhS SiPh ₃	CH	141.1	133.9	124.6	130.7	13.7	135
\times	C–Li	151.8	143.3	120.7	127.7	-1.8	131
$H \times 3$	C-//Li+	160.5	146.6	119.6	126.6	-1.4	151

We report the results of a low-temperature ¹³C, ⁷Li, ³¹P, and ²⁹Si NMR study of the sulfur- and silicon-substituted organolithium reagents 1-411 in which we show that ion separation may give either planar or pyramidal carbanions. We will abbreviate Li-(HMPA), as Lih, and use the symbol // to indicate separated ions (Scheme I).

The 7Li, 31P, and 13C NMR spectra obtained during an HMPA titration of 1-Li¹² in THF/ether (Figure 1) identify 1-Lih₀, 1-Lih₁, and 1-Lih₂ as contact ions (s, d, and t in the ⁷Li NMR due to ²J_{LiP}, observation of ${}^{1}J_{CLi}$ of 24.3, 21.4, and 18.8 Hz in the ${}^{7}Li$ and ${}^{13}C$ NMR spectra of ¹³C-enriched compound). Above 2 equiv of HMPA, separated ions 1-//Lih₃+ and 1-//Lih₄+ predominate, and the ¹J_{CLi} coupling disappears. ^{1b} Similar HMPA titrations were performed for 2-Li, the acyclic models 3 and 4 (Table I), and a series of related compounds.11 All are contact ion pairs in THF; addition of HMPA forms separated ion pairs.

Examination of the ¹³C NMR spectra of 1-Li (Figure 2) shows that 1-Lih₀ (Figure 2B) and 1-Lih₁ (Figure 2C) have diastereotopic Si(Ph)₂ signals. A coalescence temperature of -40 °C for 1-Lih₀ $(\Delta G^*$ for inversion 10.8 kcal/mol) and -100 °C for 1-Lih₁ $(\Delta G^*$ = 9.3 kcal/mol) was found. These data, together with the decrease

⁽¹¹⁾ For spectroscopic data on PhS(PhSe)CHLi, (PhSe)2CHLi, PhS-(PhMe2Si)CHLi, PhSe(PhMe2Si)CHLi, and Ph(PhSe)CHLi, see supplementary material.

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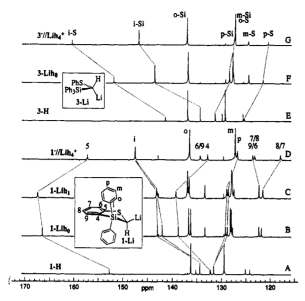


Figure 2. ¹³C NMR spectra at -100 °C of 1 and 3 in 3:2 THF/ether. A, 1-H; B, 1-Li(THF); C, 1-Lih₁; D, 1-//Lih₄ '(6 equiv HMPA); E, 3-H; F, 3-Li(THF); G, 3-//Lih₄ (6 equiv HMPA).

in ${}^{1}J_{\text{CH}}^{13}$ and ${}^{1}J_{\text{CSi}}^{13a}$ on going from 1-H (135 and 51 Hz) to 1-Li (114 and 49 Hz), indicate that the contact ions are pyramidal. The solvent-separated ion $1^{-}/\text{Lih}_4^+$ (Figure 2D) shows a loss of Si(Ph)₂ diastereotopicity and large ${}^{1}J_{\text{CH}}$ (154 Hz) and ${}^{1}J_{\text{CSi}}$ (95 Hz) values, which define the carbanion as having a planar or near planar structure. The acyclic models 3, 4, and others 11 also show a decrease in J_{CH} on going from C-H to C-Li, signaling a small *increase* in pyramidality and an increase on ion separation, indicating that substantial *flattening* occurs when the C-Li contact is lost. In sharp contrast, ${}^{1}J_{\text{CH}}$ for the bis-thio lithium reagent 2-Li does not change on ion separation (123 Hz for 2-//Lih₄+ vs 183 Hz for the model 4-//Lih₄+), and the free carbanion 2- is, therefore, pyramidal.

A rationalization for the different behavior of 1-Li and 2-Li on ion separation is suggested by an analysis of the ¹³C chemical shifts of the aromatic carbons (Table I, Figure 2). In compound 2-Li and the model systems 3-Li and 4-Li, the ipso carbons of the S-Ph and Si-Ph groups move downfield on going from C-H to C-Li and the separated ion C-//Li+ (Figures 2E-G). The para carbons move upfield. Other S-, Se-, and Si-substituted carbanions show similar behavior.¹¹ Carbon chemical shift effects near anionic centers are complex. In benzyl anions, charge polarization effects are thought to account for the nearly equal

downfield shifts for the ipso and upfield shifts for the para carbons between C-H and C-Li.^{2b,14} In compounds 1-Li to 4-Li, the charge is not directly conjugated with the benzene ring, and the small upfield para shifts reflect a correspondingly small degree of charge polarization. We feel that the large ipso shifts cannot be solely due to charge polarization and propose that they are a consequence of $n \to \sigma^*$ hyperconjugative interactions (R₂C-S-Ph \leftrightarrow R₂C=S-Ph).¹⁵ Such interactions represent an important carbanion-stabilizing mechanism.¹⁶

The ring quaternary carbons C-4 and C-5 of 1 shift downfield, just as do related carbons in 3 on going from C-H to contact C-Li. However, an upfield shift of these carbons was observed on going to the solvent-separated species 1-/Lih₄+ (Figure 2D). We interpret this upfield shift in terms of a reduction of hyperconjugative interaction into the ring. The Si-Ph groups show significantly higher than normal ipso carbon downfield and para carbon upfield shifts on ion separation, and this may signal an increase in the interaction of the carbanion center within the exocyclic silyl phenyls.

The divergent behavior of 1 and 2 can be rationalized by considering two limiting structures. The first has a pyramidal carbon and a puckered ring; carbanion-lithium interaction and $n_C \rightarrow \sigma^*$ overlap with endocyclic C-S and C-Si (for 1-Li) bonds is maximized and $n_C - n_S$ repulsion minimized. Both 1-Li and 2-Li adopt this geometry in the contact ion. For 1-Li, removal of lithium tips the balance to a second structure with a planar carbon and perhaps a less strained flat ring in which the major $n_C \rightarrow \sigma^*$ interactions are with the exocyclic Si-Ph groups, as shown in Scheme I. In contrast, 2-Li remains pyramidal in the separated ion and the ring nonplanar, since a planar ring and planar carbanion places the lone pair orthogonal to the C-S bonds and in a position to interact in a destabilizing fashion with sulfur lone pairs.

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Supplementary Material Available: ¹³C chemical shifts and coupling constants for PhS(PhSe)CHLi, (PhSe)₂CHLi, PhS-(PhMe₂Si)CHLi, PhSe(PhMe₂Si)CHLi, and Ph(PhSe)CHLi; HMPA titration of 2 (2 pages). Ordering information is given on any current masthead page.

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