Interconversion of Contact and Separated Ion Pairs in Silyl- and Arylthio-Substituted Alkyllithium Reagents

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ABSTRACT

Ether-solvated contact and separated ion pairs (CIP and SIP) for two lithium reagents, tris(trimethylsilyl)methyllithium (1) and bis(3,5 bistrifluoromethylphenylthio)methyllithium (2), have been characterized and observed for the first time under conditions of slow exchange by NMR spectroscopy, and barriers to interconversion have been measured. A Saunders isotope perturbation experiment was used to support identification of the CIP and SIP species for 2.

Organolithium reagents in solution exist as a variety of structures, depending in a complex way on solvent donicity, the presence of Lewis basic additives, and the structure of the carbanion.^{1a,2,3,4a,5a} In nonpolar media, they tend to be

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aggregated, but most are converted progressively to lower aggregates as solvent donicity increases. This is a consequence of the fact that lithium is normally tetracoordinate: only one basic group can coordinate to each lithium in a cubic tetramer, two can coordinate to each lithium in a fourcenter dimer, and three can coordinate to the lithium in a monomer. If the carbanion has sufficient charge delocalization, ion separation can occur (conversion of a contact ion pair (CIP) to a separated ion pair (SIP)) driven by tetra (or even higher) coordination of the lithium by basic solvents or additives.4a,6a The conversion of CIP to SIP can lead to enormous increases, $6b,7,8$ as well as decreases, $4b,6b$ in reactivity. Selectivity changes have also been detected.^{6c,6d,9} We report on NMR spectroscopic studies of two systems which address the barrier to interconversion of CIP and SIP structures.

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The lithium reagent **1**, formed by deprotonation of tris(trimethylsilyl)methane,^{10a} Li/Hg exchange of $[(Me₃ Si$ ₃C]₂Hg,¹¹ or Li/Se exchange^{12a} of (Me₃Si)₃CSePh, has unusual structural properties: in the solid state, it forms a four-center dimer when no solvent is present¹¹ and a triple ion when solvated by THF (**1T**).10a An NMR study10b suggested that the triple ion is also present in solution, in addition to a second species, which was tentatively assigned the structure **1C**, the monomeric contact ion pair. We have reinvestigated the solution structure of **1** by multinuclear NMR spectroscopy of natural abundance as well as of ^{13}C and ⁶Li-enriched material in ether-THF solvent mixtures
and found that below -80 °C there are actually three species and found that below -80 °C there are actually three species present *in slow exchange on the NMR time scale*: the triple ion **1T**, the contact ion pair **1C**, and the separated ion pair **1S**.

Evidence for this assignment was provided by the ^{13}C , ${}^{7}Li$, ${}^{6}Li$, ${}^{29}Si$, and ${}^{1}H$ NMR spectra at various temperatures and in several solvents. Two of the species, **1C** and **1T**, can be readily identified from the NMR spectra of the 13Cenriched sample (Figure 1). For **1T**, the 13C NMR signal of the C-Li carbon at δ 1.8 with lithium at natural abundance is a broad doublet, a consequence of unusually fast quadrupolar relaxation of ⁷ Li which partially averages the peaks of

Figure 1. NMR spectra of **1** in 3:2 THF/ether. 13C NMR spectra of (a) 13C-enriched and (b) 13C-6Li doubly enriched **¹**. 6Li NMR spectra of (c) 6 Li-enriched and (d) 13 C $-{}^6$ Li doubly enriched 1.

the expected 1:1:1:1 quartet, superimposed on which is the natural abundance (7.4%) 1:1:1 triplet of the ⁶Li isotopomer (Figure 1a).¹³ The ⁶Li-enriched sample shows a 1:2:1 triplet at δ 2.8 in the ⁶Li (Figure 1d) and a 1:1:1 triplet in the ¹³C NMR spectra (Figure 1b), defining a carbon bonded to one lithium and a lithium bonded to two carbons. Compound **1C** shows the expected doublet at δ -0.2 in the ⁶Li and a 1:1:1 quartet (patural abundance) or a 1:1:1 triplet (⁶Li 1:1:1:1 quartet (natural abundance) or a 1:1:1 triplet (6Li enriched) at δ -2.2 in the ¹³C NMR spectra ($J_{\text{Li-C}}$ = 8.5 Hz).

The third species was eventually assigned the separated ion pair structure **1S**, although not without some trepidation because the observation of distinct NMR signals for CIP/ SIP pairs in ethereal solvents is unprecedented.¹⁴ The evidence for this assignment can be summarized as follows. The signals for **1S** were present in all samples prepared by several different methods (metalation, Li/Se exchange, Li/I exchange) provided that the solvent contained a significant fraction of THF. The appropriate signals of all of the nuclei (⁶Li, ¹H, ¹³C, ²⁹Si) of **1S** and **1C** coalesced at around -70
^oC, showing that the signals assigned to **1S** were not a °C, showing that the signals assigned to **1S** were not a structurally different impurity. The species shows no C-Li coupling or signal broadening, even at -135 °C, and only one signal for the separated lithium cation of **1T** and **1S** was

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⁽¹⁴⁾ Separate NMR signals for SIP and CIP have been observed for crypt¹⁵ and HMPA^{6a} solvated species, where the SIP and CIP have different levels of crypt/HMPA solvation. Such strong ligands for lithium have relatively slow ligand exchange rates, so distinct solvates can be readily observed on the NMR time scale.

observed in the lithium NMR spectra. In dimethyl and diethyl ether, the signals for **1S** disappear due to poorer solvation, whereas the addition of HMPA led to complete conversion to $1S$, as is seen for many related reagents.^{6c,6e} The C-Si coupling for **1S** $(^1J_{Si-C} = 64.9 \text{ Hz})$ was significantly larger
than that for **1C** ($(^1L_{Co} = 42.0 \text{ Hz})$ which was close to the than that for **1C** $(^1J_{Si-C} = 42.0$ Hz, which was close to the value of 37.5 Hz in tris(trimethylsily) methane^{6h}) and **1T** value of 37.5 Hz in tris(trimethylsilyl)methane^{6h}) and 1T $(^1J_{Si-C} = 42.6 \text{ Hz})$, suggesting increased s character in the $C-Si$ bond on ion senaration 6e,16 Removal of the lithium $C-Si$ bond on ion separation.^{6e,16} Removal of the lithium cation should favor a more planar carbon to relieve strain and allow better π delocalization of charge into the C-Si *σ** bonds. Finally, dilution studies at room temperature showed that the concentrations of **1T** and the averaged **1C**/ **1S** signal changed in a way consistent with a dimeric structure for **1T** (a plot of log[**1C**-**S**] vs log[**1T**] had a slope of 1.91). However, at -105 °C, where **1S** and **1C** can be measured independently, comparison of **1T** with **1C** gave a molecularity of 2.2, whereas comparison with **1S** gave a value of 1.6. We do not understand the origin of these deviations from the expected value of 2, but they suggest a small degree of aggregation for **1S**.

DNMR studies of the ${}^{13}C$ and ¹H NMR spectra (Me₃Si signals) provided barriers to the interconversion of the three species.6h The barrier between **1S** and **1C** was over 9 kcal/ mol $(\Delta G^{\pm}{}_{-78} = 9.6 \text{ kcal/mol}, \Delta H^{\pm} = 8.1 \pm 0.2 \text{ kcal/mol},$
 $\Delta S^{\pm} = -8.0 \pm 0.8 \text{ AU}$. The ratio of **1S** and **1C** is essentially $\Delta S^{\dagger} = -8.0 \pm 0.8$ eu). The ratio of **1S** and **1C** is essentially temperature independent. Coalescence of **1T** with the averaged signals of **1C** and **1S** could barely be achieved in the 3:2 THF/ether medium because it occurred near the boiling point of the solution $(\Delta G^{\ddagger}{}_{41.3} = 16.4 \text{ kcal/mol}, \Delta H^{\ddagger} = 12.9 \pm 0.4 \text{ kcal/mol}, \Delta S^{\ddagger} = -11.1 \pm 1.1 \text{ eq. for conversion of }$ \pm 0.4 kcal/mol, $\Delta S^{\dagger} = -11.1 \pm 1.1$ eu for conversion of **1T** to **1C** and **1S**).

The extraordinary high barriers to interconversion of **1S** and **1C** encouraged us to look for other systems where this barrier could be measured. The requirements are stringentthe lithium reagent must be evenly balanced between CIP and SIP at low temperatures in THF-ether mixed solvent. We have previously reported on bis(3,5-bistrifluoromethylphenylthio)methyllithium (**2**) as an easily separable ion pair in connection with studies of the competition between 1,2 and $1,4$ -addition to enones.^{6c}

A study of the effect of solvent donicity revealed that in pure diethyl or dimethyl ether only **2C** was present, but it was converted completely to **2S** if several equivalents of HMPA were added. This can be seen from the 13C chemical shifts of the aryl C-S carbon, which, as observed for other lithium reagents of this type,^{6e} moved over 10 ppm downfield on ion separation. Figure 2 shows 13C NMR spectra of **2** in ether, where only **2C** is present, and various ether/THF and HMPA mixtures where the downfield progression of the ^C-S carbon signals indicates increasing fractions of **2S** (the spectra of Figure 2 were taken on a 60:40 mixture of D/H

Figure 2. Saunders isotope perturbation experiment: 13C NMR spectra of a THF titration of a 0.23 M solution of **2** (55% D-labeled) in 7:1 ether/pentane at -94 °C. Spectrum (e) contains 4 equiv of HMPA.

isotopomers, and the signals in Figure $2b-d$ are doubled as a consequence of isotope perturbation; see below).

Although we have assigned the chemical shift changes of Figure 2 and the decoalescence of two signals described below to a SIP/CIP interconversion, there are a number of other structural changes which can cause chemical shift changes and signal doubling in NMR spectra. These include conformational equilibrations, aggregate exchanges, or variable solvation. Only the SIP/CIP interconversion is likely to involve a significant change in hybridization, so we performed the Saunders isotope perturbation experiment¹⁷ shown in Figure 2. In samples where significant fractions of both **2S** and **2C** were present, the signal of the deuterated compound was 0.30-0.34 ppm downfield of the protio isotopomer. These isotope shifts are far too large, and of the wrong sign, to be normal ³ Δ shifts, where $\Delta \delta$ of <0.05 normal up steed ¹⁸ In fact, no isotone shift ppm upfield would be expected.18 In fact, no isotope shift could be detected when only **2S** (in THF/ether/HMPA) or **2C** (in ether) was present. The observation of a large isotope shift signals a hybridization change of the $C-H(D)$ bond, as expected for a SIP/CIP equilibration.^{1b} The downfield shift of the D-isotopomer means that there is a larger fraction of **2S** in the deutero compound. This is as expected from the much larger ${}^{1}J_{CH}$ in **2S** vs **2C** (183 vs 141 Hz), indicating a change in hybridization of the carbanion toward $sp²$ in **2S**, hence a stronger C-H(D) bond and an energetic preference for $C-D$ in the SIP.^{6e} We believe that this is the first time that a SIP/CIP dichotomy has been established using the Saunders isotope perturbation method, although carbanion properties, such as the localization of allyllithiums^{5c,19,20} and

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propargyl/allenyllithiums, ^{6f} the cyclopropylcarbinyl-allyl interconversion, 21a,22 and the symmetry of carbanion-CH bonds,21b have been probed.

Samples of **2** (H only, no D) in mixtures of dimethyl ether and THF behaved similarly to those in ether and THF, with a large predominance of **2C** in pure dimethyl ether and an increasing fraction of **2S** as THF was added. Both the ⁷ Li and 13C spectra of such samples developed significant broadening at temperatures below -130 °C. This effect could be assigned to slow interconversion between **2C** and **2S**. In a sample with an appropriate mixture of THF and $Me₂O$ such that the averaged 13C NMR signal indicated nearly equal amounts of **2C** and **2S**, it was possible to decoalesce the two signals at -151 °C (Figure 3), giving peaks with

Figure 3. 13C and 7Li NMR spectra of a 0.16 M solution of **2** in Me₂O and a Me₂O/THF mixture. The upper lines are simulations (two-spin exchange). $6g,6h$

approximately the shifts of **2C** in ether (Figure 2a) and **2S** in THF/ether/HMPA (Figure 2e). Because of the smaller

chemical shift difference (714 vs 1007 Hz), the 7 Li signals were not quite decoalesced even at -155 °C, the lowest temperature that could be achieved.

Line shape simulation of the spectra gave a free energy of activation of 5.3 kcal/mol for interconversion of the contact and separated ion pairs of **2**. Previous measurements of the barrier to interconversion of lithium reagent SIP and CIP species have come from ESR studies of aromatic radical anions, with values of $2-4$ kcal/mol.²³ The slightly higher barrier in **2** seems reasonable, given the greater localization of charge and hence the stronger C-Li coordination expected for this system compared to aromatic radical anions. However, the much higher barrier for the SIP/CIP interconversion of tris(trimethylsilyl)methyllithium (**1**) must have its origin in the extraordinary steric hindrance at the carbanion center. It seems likely that solvent-induced ion pair separation occurs by an associative mechanism.24 In **1**, such a process may be sterically prohibited, requiring an S_E1 -type dissociation of a $Li(THF)₃⁺$ or perhaps even a $Li(THF)₂⁺$ cation with little or no solvent assistance. Similarly, the ion association process would then require predissociation of one or two solvent molecules from the $Li(THF)₄⁺$ cation before the C-Li
contact could be established contact could be established.

The even higher barrier for dissociation/association of **1T** must have similar origins: if the triple ion simply fragments to a molecule of **1C** and **1S**, then the departing **1C** fragment may be unable to coordinate adequately with solvent to assist cleavage of the $C-Li$ bond.²⁸ Conversely, if the triple ion fragments to two molecules of **1C** by electrophilic attack of $Li⁺$ on **1T**, then the steric inaccessibility of the C-Li bond of **1T** will require a substantially desolvated cation electrophile, as well as a poorly solvated or unsolvated departing lithium. A zinc analogue of **1T**, bis[tris(trimethylsilyl)methyl] zinc, shows legendary stability toward electrophiles^{10c} attributed to this kind of steric shielding of the metal and carbanion reactive centers.

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