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Autocatalysis in Lithium Diisopropylamide-Mediated Ortholithiations

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Abstract: Ortholithiation of 3-fluorophenyl-*N*,*N*-diisopropyl carbamate by lithium diisopropylamide (LDA) in THF at -78 °C affords unusual rate behavior including linear decays of the carbamate, delayed formation of LDA-aryllithium mixed dimers, and evidence of autocatalysis. A mechanistic model in conjunction with numeric integration methods accounts for the time-dependent changes in concentration. The two critical rate-limiting steps in the model entail (1) an LDA dimer-based metalation of arylcarbamate, and (2) a rate-limiting condensation of the resulting aryllithium with the LDA dimer to form two isomeric LDA-ArLi mixed dimers. One isomer elicits a highly efficient (post-rate-limiting) metalation of aryl carbamate, in turn, regenerating aryllithium. The prevalence and implications of such autocatalysis are discussed.

Introduction

Lithium diisopropylamide (LDA) is often the reagent of choice for reactions requiring a strong base.¹ The central importance of LDA motivated us to examine how solvation and aggregation influence its reactivity.² Studies spanning two decades have documented more than a dozen mechanisms that differ in solvation and aggregation at the rate-limiting transition structures. Recent studies of the LDA-mediated ortholithiation of aryl carbamates^{3,4} underscore the potential complexity by revealing both monomer and dimer-based pathways. A pronounced autoinhibition was traced to intervening mixed aggregates, which are often observed in reactions of LDA.⁵



LDA-mediated metalation of aryl carbamate 1, the most reactive of all carbamates we had studied to date, displayed two perplexing behaviors. First, treating carbamate 1 with 1.0



Figure 1. Reaction of 0.10 M carbamate 1 with 0.10 M LDA at -78 °C in neat THF as monitored by ¹⁹F NMR spectroscopy. The solid and dashed lines represent least-squares fits to first- and second-order decays, respectively.

equiv of LDA in neat THF at -78 °C—conditions used for many LDA-mediated reactions-affords aryllithium monomer 2 to the exclusion of mixed aggregates. The concentration of 1 also follows a first-order decay to a reasonable approximation (Figure 1); a second-order fit (dashed line) is clearly inferior. Thus, on first inspection, we appeared to have discovered the most outwardly simple LDA-mediated reaction reported to date. Something was wrong, however: This inherently bimolecular reaction should not be first order, regardless of whether the ratelimiting transition structure is based on an LDA monomer or dimer. The second strange observation appeared when a large excess of LDA was used. The large excess-so-called pseudofirst-order conditions-usually forces a higher-order reaction to behave as if it is first order (thus the name). When such pseudofirst-order conditions are used, we observe a linear decay of 1 (Figure 2). The decay is clearly *not* first order in carbamate 1, appearing instead to be *zeroth*-order.

We describe herein structural and rate studies showing that numerous deviant rate behaviors for the ortholithiation of 1 stem

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Figure 2. Reaction of 0.010 M carbamate 1 with 0.10 M LDA at -78 °C in neat THF as monitored by 19 F NMR spectroscopy.

Scheme 1



from mixed aggregates and autocatalysis^{6,7} as summarized generically in Scheme 1.^{5–11} *It is the most mechanistically complex reaction of LDA reported to date.* Autocatalysis results from intervening LDA–ArLi mixed dimers **4** and **5**. Although metalation of arene **1** by minor isomer **5** (Step III) is an important step in this unusual autocatalysis, the condensation of aryllithium **2** with LDA dimer **3** (Step IV) is rate limiting.

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Mounting evidence suggests that autocatalysis may be prevalent in LDA/THF-mediated reactions at -78 °C.

Results

General Methods. LDA, [⁶Li]LDA, and [⁶Li,¹⁵N]LDA were purified as white crystalline solids.¹² The structural assignments derive from routine ⁶Li, ¹³C, and ¹⁵N NMR spectroscopic methods.^{9,13}

Rate studies were carried out using in situ IR spectroscopy¹⁴ and ¹⁹F NMR spectroscopy¹⁵ following standard protocols. The results from the two forms of spectroscopy are fully consistent. Unless stated otherwise, pseudo-first-order conditions were established by maintaining the concentration of carbamate 1 at 0.004–0.008 M. LDA and THF were maintained at high yet adjustable concentrations using hexane as the cosolvent.¹⁶ Although pseudo-first-order conditions are used routinely to preclude the effects of mixed aggregation and other conversion-dependent effects,^{2,8b,11} mixed aggregation effects are pronounced under all conditions used to lithiate carbamate 1.^{17,18} IR spectroscopic analysis of the lithiation of carbamate 1 with ≥ 1.0 equiv of LDA in THF reveals that the reaction proceeds to full conversion. Under no conditions do we observe carbamate–LDA complexation. Time-dependent concentrations

- (13) Spectral data: ⁶Li NMR; **2** (δ 1.27, s), **3** (δ 1.97, t, J = 5.0 Hz), **4** (δ 1.79, d, J = 5.3 Hz); ¹³C NMR; **4** (δ 150.5, dq, J = 121.7, 5.6 Hz); ¹⁵N NMR; **3** (δ 74.7, q, J = 5.0), **4** (δ 76.4, q, J = 5.3); ¹⁹F NMR; **1** (δ -113.0, -113.2), **2** (δ -74.5), **4** (δ -79.4), **5** (δ -80.5). ⁶Li and ¹⁵N NMR spectra were recorded in 11.1 M THF/pentane at-90 °C; ¹³C NMR spectra were recorded in 3.1 M THF/pentane at-90 °C; ¹⁹F NMR spectra were are singlet.
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- (16) The concentration of LDA, although expressed in units of molarity, refers to the concentration of the monomer unit (normality).
- (17) Carbamate 1 and structurally related diisopropylamine-derived carbamates and carboxamides¹⁸ display distinct conformational properties that are not observed in less sterically congested analogues. Carbamate 1 affords normal NMR spectra at ambient temperature. On cooling the NMR probe below-70 °C, the single ¹⁹F resonance (-113.1 ppm) appears as a pair of resonances (-113.2 and-113.0 ppm) in equal proportions. Similarly in the ¹³C NMR spectrum, two methyne resonances of the diisopropyl moiety are observed at ambient temperatures, whereas four are observed at \leq -90 °C. One can explain these results by assuming a slow conformer exchange illustrated below and discussed previously.¹⁸ A crystal structure of 1 affords the following structure displaying a highly typical orientation of the diisopropyl moiety.



We have no evidence that conformation influences the mathematical form of the rate law described herein. To the contrary, the conformers are shown to maintain the equilibrium proportions throughout the course of the reaction.

measured by ¹⁹F NMR spectroscopy are fit to a mechanistic model (see Discussion) expressed by a set of differential equations (see Experimental Section) using numeric integration, yielding functions describing concentration versus time for multiple species.

Density functional theory (DFT) computations were carried out at the B3LYP/6-31G(d) level.¹⁹ The isopropyl moieties of carbamate **1** were approximated with methyl groups, and THF was modeled using Me₂O. Free energies are reported in kcal/ mol at -78 °C. The computational results are described sparingly in appropriate contexts below; additional results are archived in Supporting Information, including some calculations using LDA rather than Me₂NLi.

Solution Structures. In this section, we focus only on structure; the time-dependent aggregate concentrations are examined extensively in subsequent sections. ⁶Li and ¹⁵N NMR spectroscopic studies using 2.0 equiv of [⁶Li,¹⁵N]LDA reveal aryllithium **2**, LDA dimer **3**, and mixed dimer **4** (Figure 3).^{13,20} An additional species (ca. 3%), putative mixed dimer **5**, was detected using ¹⁹F NMR spectroscopy (below) but was not observed in the ⁶Li spectra.

Aryllithium **2** shows no ¹⁵N coupling, consistent with the absence of an LDA fragment. Limited solubility prevented ¹³C NMR spectroscopic analysis of the ⁶Li⁻¹³C coupling, but analogous aryllithiums with ortho fluoro moieties²¹ or other potentially coordinating ortho substituents are monomeric.²² Computational studies show minima for mono- and disolvated aryllithium monomer and suggest that disolvated monomer is preferred. Mixed dimer **4** is easily assigned from ⁶Li⁻¹³C and ⁶Li⁻¹⁵N coupling patterns as described previously.^{5a}



Monitoring ortholithiations of **1** using ⁶Li and ¹⁹F NMR spectroscopies allowed us to assign ¹⁹F resonances to aryllithium monomer **2** and mixed dimer **4**; a representative spectrum is shown in Figure 4. We also observe an additional species (**5** in Figure 4) initially suspected to be a mixed trimer akin to those characterized previously.^{5,8,10} The ratio of **4** and **5**, however, is invariant with large changes in ArLi, LDA, and THF concentrations (see Supporting Information). Thus, **4** and **5** are *isomers*.

Computations were used to investigate the solvation of 4 and the more elusive gross structure and solvation of 5 (Scheme 2). Serial solvation shows minima for mono- and disolvated mixed dimers 6-9. Trisolvates failed to minimize, resulting in des-

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Figure 3. ⁶Li NMR spectrum of 0.10 M [6 Li, 15 N]LDA at -90 °C with (A) 0.050 M 1; (B) 0.10 M 1; (C) 0.10 M 1 at -90 °C after warming at -40 °C for 60 min.



Figure 4. 19 F NMR spectrum of 0.10 M LDA with 0.02 M 1 at -78 °C.

olvation in each attempt. Mono- and disolvated dimers 6 and 7 are isoenergetic, whereas 8 and 9 are significantly less stable. Dimer 7, bearing a C-Li-N-Li connectivity characteristic of open dimers^{23,24} but with an unusual Li-F interaction, is the only detectable structure bearing two solvents coordinated to



the lithium distal to the carbamoyl moiety. No minimum was found corresponding to the analogue of **7** with a single solvent on the distal lithium.

Experimental data show that **4** and **5** are isomeric, whereas computations suggest such closed and ring-expanded forms prefer different solvation numbers. We are convinced by the experimental data and, given the approximations used, are comfortable offering **9** and **7** as computational models of isomeric mixed dimers **4** and **5** (respectively). The unusual structural features in **7** reappear in calculated transition structures (vide infra) and in mechanistic discussions.

Linear Decays. In theory, the linear decay for the ortholithiation of **1** (illustrated in Figure 2) could arise if an aggregation event such as a deaggregation is rate limiting (eqs 2 and 3).²⁵ The rate of ortholithiation would be dictated by the rate at which monomer is formed²⁶ and independent of the initial substrate concentration (Figure 5).

$$(i-\Pr_2 NLi)_2 \hookrightarrow 2 \ i-\Pr_2 NLi \tag{2}$$

$$i - \Pr_2 NLi + ArH \xrightarrow{fast} [(i - \Pr_2 NLi)(ArH)]^{\ddagger}$$
 (3)

In practice, the scenario in eqs 2 and 3 did not withstand scrutiny. Figure 6 reveals reaction rates that depend markedly on the initial concentration of the carbamate. The nearly coincident *x*-intercepts in Figure 6 are strikingly similar to those in plots of initial rates arising from a first-order reaction.²⁶ A plot of the slopes from Figure 6 versus initial substrate concentration (Figure 7) illustrates the concentration-dependent rates. The highest substrate concentrations in Figure 6 far exceed those necessary to preserve pseudo-first-order conditions, yet the linearities persist.

These results present a paradox. The linearities suggest zerothorder substrate dependencies, but the substrate concentrationdependent slopes are akin to those for first-order substrate dependencies. Autocatalysis is the culprit.²⁷

Evidence of Autocatalysis. We routinely carry out rate studies under pseudo-first-order conditions in which the substrate is kept at low concentration. By allowing no more than a 5% accumulation of new lithium salts, one usually precludes autoin-



Figure 5. Theoretical depiction of substrate dependence for a zeroth-order reaction.



Figure 6. Plot showing IR absorbance of $1 (1721 \text{ cm}^{-1})$ versus time for different initial concentrations of carbamate 1.



Figure 7. Plot of Δ [1]/ Δt (from Figure 6) versus [1] for the ortholithiation of 1 with 0.10 M LDA in 12.2 M THF/hexane cosolvent at -78 °C.

hibition and autocatalysis that can arise from the intervention of mixed aggregates.² The successful exclusion of such mixed aggregation effects is demonstrated by a simple control experiment: on completion of a reaction, injection of a second small aliquot of substrate provides a decay and a second value for k_{obsd} that is unchanged (±10%). By inference, the products of the reaction do not influence the rate or mechanism.

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Figure 8. Absorbance versus time for the reaction of (0.004 M) **1**- d_1 with (0.10 M) LDA in 12.2 M THF/hexane cosolvent at -78 °C. (1- d_1 , 1721 cm⁻¹; **2** + **4**, 1660 cm⁻¹).

One might expect a second aliquot of 1 would afford an analogous linear decay that is superimposable on the first $(\pm 10\%)$. Experimentally, however, serial injection of two aliquots of carbamate 1 to a large excess of LDA in THF at -78 °C reveals a linear decay in the first and an almost instantaneous reaction in the second. Significant accelerations in the second aliquot are observable under many conditions studied, suggesting to us that the linear decays in Figure 6 derive, at least in part, from the superposition of autocatalysis on a normal (first-order) decay.⁶ Pronounced mechanistic changes likely derive from LDA–aryllithium mixed aggregates.²⁸

Autocatalysis emerged in a more conventional form^{6,7} with minor changes in the reaction conditions. By example, sigmoidal curvature (anomalous acceleration) is observed at low THF concentration using hexane as a cosolvent.²⁹ Similar sigmoidal decays are observed for the lithiation of $1-d_1$ (Figure 8). A second aliquot of $1-d_1$ is consumed nearly instantaneously. Comparing the initial rates (slopes at early conversion) of **1** and $1-d_1$ affords a small kinetic isotope effect ($k_H/k_D = 2-3$). Ortholithiations, including LDA-mediated ortholithiations of structurally related carbamates,⁵ display isotope effects covering a wide range ($k_H/k_D = 3$ to >20).^{30,31} The high reactivity of the LDA-ArLi mixed aggregates—an essential component of autocatalysis—correlates with the delayed onset of mixed aggregation as shown below.

Delayed Mixed Aggregation. Monitoring ortholithiations of **1** using ¹⁹F NMR spectroscopy reveals very odd time-dependent changes in concentrations illustrated emblematically in Figures 9–11. The origins of the functions drawn through the data are described in the discussion. Several qualitative observations and control experiments are immediately noteworthy.

The linear loss of carbamate **1** in Figure 9 correlates quantitatively with appearance of aryllithium monomer **2**. Oddly, mixed dimers **4** and **5** appear after the ortholithiation of **1** is

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- (29) Sigmoidal curvature for the loss of aryl carbamate is also observed using tetrahydropyran and 2-methyltetrahydrofuran.



Figure 9. Concentrations versus time for the reaction of 0.0074 M **1** with 0.10 M LDA in 12.2 M THF at -78 °C monitored by ¹⁹F NMR spectroscopy. The functions result from a least-squares fit to the model outline in eqs 9–12 and described mathematically by eqs 16–20.



Figure 10. Concentrations versus time for the reaction of 0.0074 M 1 with 0.10 M LDA in 4.9 M THF at $-78 \text{ }^{\circ}\text{C}$ monitored by ¹⁹F NMR spectroscopy. The functions result from a least-squares fit to the model outline in eqs 9-12 and described mathematically by eqs 16-20.



Figure 11. Concentrations versus time for the reaction of 0.01 M **1** with 0.40 M LDA in 12.2 M THF at -100 °C monitored by ¹⁹F NMR spectroscopy. The functions result from a least-squares fit to the model outline in eqs 9–12 and described mathematically by eqs 16–20.

complete and quite slowly: Aggregates are not completely equilibrated on the time scale of the ortholithiation.²⁵ At low THF concentrations (Figure 10), the mixed dimers begin to appear before the metalation of 1 is complete. The inflection point in the formation of mixed dimer 4—the point of maximum rate—correlates with the maximum concentration of aryllithium 2, suggesting that 2 is a key precursor to 4.³² Importantly, the ratio of 4 and 5 remains constant at the equilibrium value throughout the reaction coordinate, indicating that 4 and 5 exchange rapidly on the time scale of the metalation.

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Figure 12. Plot of initial rates versus [LDA] for the lithiation of carbamate 1 (0.0074 M) in 12.2 M THF at -78 °C. The curve depicts an unweighted least-squares fit to rate = $k[LDA]^n$ ($k = (4.2 \pm 0.2) \times 10^1$; $n = 1.02 \pm 0.03$).



Figure 13. Plot of initial rates versus [THF] for the lithiation of carbamate 1 (0.004 M) in 0.1 M LDA using hexanes as cosolvent at -78 °C. The curve depicts an unweighted least-squares fit to rate = k[THF] + k' (k = (1.0 ± 0.2) × 10⁻⁶; k' = (2.3 ± 0.1) × 10⁻⁵).

Mixed dimers appeared to be the source of autocatalysis. Indeed, injecting carbamate 1 directly into solutions containing LDA dimer 3 and mixed dimers 4 and 5 within the NMR probe²⁵ showed that both 4 and 5 disappear in seconds with concomitant appearance of aryllithium monomer 2 (Step III, Scheme 1). The much slower condensation of aryllithium 2 with LDA to regenerate the mixed dimers (Step IV, Scheme 1) occurs subsequently.

Temperature Effects. Ortholithiation at -100 °C (Figure 11) causes mixed dimer 4 to build to an appreciable and relatively constant concentration, whereas mixed dimer 5 does not. The ratio of 4 and 5 is also *not* constant and is markedly perturbed from its equilibrium value until the carbamate is consumed. *The isomerization of* 4 *and* 5 *is slow on the time scale of the metalation at* -100 °C. This temperature effect proves to be important in showing that 5 is the key reactive form (vide infra).

Reaction of LDA Dimer with ArH. The first stage in teasing apart the overall mechanism is to investigate key steps individually. The method of initial rates offers insight into the reaction of LDA and carbamate 1 before the onset of autocatalysis (Step I, Scheme 1). (By virtue of the linearities illustrated in Figure 6, the slopes and initial rates are equivalent.) Using ¹⁹F NMR spectroscopic data, we showed a first order in LDA (Figure 12) and zeroth order in THF (Figure 13). The idealized rate law³³



Figure 14. Plot of k_{obsd} versus [LDA] for the condensation of LDA with aryllithium **2** (0.0074 M) in 12.2 M THF at -78 °C. The curve depicts an unweighted least-squares fit to $k_{obsd} = k[LDA]^n + c$ ($k = (60 \pm 10)$; $n = 1.1 \pm 0.1$).

in eq 4, in conjunction with the assignment of LDA dimer **3** as a disolvate,^{2,9a} is consistent with the generic mechanism in eq 5. Open-dimer-based transition structure **10** is shown to be plausible by computation.⁸ Analogous open-dimer-based transition structures have been implicated on many occasions.²³

$$-d[\text{ArH}]/dt = k'[\text{LDA}][\text{THF}]^{0}[\text{ArH}]$$
(4)

$$(i-\Pr_2 \operatorname{NLi}_3)_2(\operatorname{THF})_2 + \operatorname{ArH}_1 \xrightarrow{\kappa} \left[(i-\Pr_2 \operatorname{NLi})_2(\operatorname{ArH})(\operatorname{THF})_2\right]^{\ddagger} (5)$$



Reaction of LDA Dimer with ArLi. After substrate 1 is consumed, the loss of aryllithium 2 via condensation with LDA dimer 3 to form mixed dimer 4 (Step IV, Scheme 1) follows clean pseudo-first-order behavior. Analogous reactivity was observed by adding a solution of LDA dimer 3 to aryllithium monomer 2. Monitoring the pseudo-first-order rate constants as a function of LDA and THF concentrations (Figures 14 and 15) shows that the condensation is first order in LDA and an inverse first order in THF.³⁴ The idealized rate law (eq 6) is consistent with a rate-limiting transition structure of stoichiometry $[(i-Pr_2NLi)_2(ArLi)]^{\dagger}$ (eq 7).³⁵ (The THF ligands are not included because the solvation number of ArLi 2 was studied only computationally, but disolvation seems likely.) This slow aggregation event is important in the formulation of an overall mechanistic hypothesis. Although we cannot even hazard a guess as to what transition structure 11 might look like, the relationship of transition structures 10 and 11 may be important (vide infra).

$$-d[\operatorname{ArLi}]/dt = k[\operatorname{LDA}][\operatorname{ArLi}][\operatorname{THF}]^{-1}$$
(6)

$$(i-\Pr_2 \operatorname{NLi})_2 + \operatorname{ArLi}_2 \xrightarrow{k} [(i-\Pr_2 \operatorname{NLi})_2 \operatorname{ArLi}]^{\dagger}$$
(7)

Reaction of Mixed Dimers with ArH. The reaction of arene **1** with the mixed dimers (Step III, Scheme 1) was too fast to obtain useful rate data as evidenced by the nearly instantaneous reaction when a carbamate **1** is added to a solution of **4** and **5**. DFT computations, however, provide insights that we find quite provocative. Given the circumstantial evidence supporting disolvation of **4** and **5**, we focused our computations on transition structures bearing the carbamate carbonyl moiety and

^{(31) (}a) Dyke, A. M.; Gill, D. M.; Harvey, J. N.; Hester, A. J.; Lloyd-Jones, G. C.; Munoz, M. P.; Shepperson, I. R. *Angew. Chem., Int. Ed.* **2008**, *47*, 5067. (b) Metalation of carbamate **1** in *t*-BuOMe proceeds from an LDA-carbamate complex via a normal (first-order) disappearance and displays $k_{\rm H}/k_{\rm D} = 3-4$.

Scheme 3



Figure 15. Plot of k_{obsd} versus [THF] for the condensation of LDA with aryllithium **2** (0.0074 M) and 0.1 M LDA at -78 °C. The curve depicts an unweighted least-squares fit to $k_{obsd} = k$ [THF]^{*n*} ($k = (39 \pm 7)$; $n = -1.1 \pm 0.2$).

one or two solvents. The transition structures bearing only one solvent proved to be much more stable computationally. (We have no experimental evidence to support either solvation number.) The results focusing on monosolvated transition structures **12** and **13** are illustrated in Scheme 3.

We believe the most logical scenario to explain the data at -100 °C is that **5** reacts directly with carbamate **1** (modeled by the conversion of **7** to **13**) and that **4** reacts by a rate-limiting conversion to **5** (modeled by the conversion of **9** to **7**). We note that the lack of Li–C–Li contacts in **7** and **13** is intriguing.

Discussion

We have described rate studies of the LDA-mediated metalation of carbamate 1 (eq 1). A host of observations that are unusual when placed in the context of previous mechanistic studies point to the importance of autocatalysis in which the resulting ArLi forms highly reactive mixed aggregates (Scheme 1). A summary of the key observations is followed by a detailed discussion of a mechanistic model depicted generically in Scheme 1.

Strange Rate Behavior. Ortholithiation of carbamate 1 under standard conditions—equimolar LDA in THF at -78 °C—shows a notable simplicity exemplified by an apparent first-order decay (Figure 1) and an absence of mixed aggregates. Surprisingly,

metalations using >2.0 equiv of LDA display unusual linear decays of carbamate 1 to full conversion (Figure 2). Linear decays of substrate versus time typically stem from zeroth-order dependencies on substrate that occur because the substrate is *not* involved in the rate-limiting step.^{7c} The carbamate concentration dependence of the decays in Figure 6, however, suggests a first-order dependence on carbamate 1. The two conclusions seem paradoxical. Indeed, the downwardly curving (sigmoidal) decays observed when using deuterated carbamate 1 (Figure 8) or low THF concentrations suggest autocatalysis is the culprit,²⁹ and the high reactivities of LDA–ArLi mixed dimers are supportive.

¹⁹F NMR spectroscopy offers a vivid picture of the odd timedependent aggregate concentrations (Figures 9–11). The linear disappearance of carbamate 1 in Figure 9 correlates quantitatively with an analogous linear formation of aryllithium monomer 2. Mixed dimer 4 and traces of isomeric mixed dimer 5 eventually appear, but only after the ortholithiation is complete. (Depiction of 5 as a fluorine-chelated open dimer^{23,24} derives from computational rather than spectroscopic data.) This delayed mixed aggregation was difficult to understand at the outset.

Reducing the THF concentration (Figure 10) causes aryllithium 2 to reach a lower maximum concentration and makes aggregates 4 and 5 visible before the carbamate has been fully consumed. Lowering the reaction temperature to -100 °C produces several notable changes including a measurable and surprisingly constant concentration of mixed dimer 4 but not 5 (Figure 11). The deviation of isomeric mixed dimers 4 and 5 from their equilibrium ratio is significant.

Mechanistic Model. It is an understatement to say that the conventional mechanisms for LDA-mediated reactions² fail to account for the results summarized above. Many unconventional models were considered and also found wanting. We offer a single model described by eqs 9-12 (referred to as simply "the model") based on the four steps summarized in Scheme 1.

LDA Dimer-Based Metalation (Step I)

$$(i-\Pr_2 \underset{\mathbf{3}}{\operatorname{NLi}})_2 + \underset{\mathbf{1}}{\operatorname{ArH}} \xrightarrow{k_1} (i-\Pr_2 \underset{\mathbf{4}}{\operatorname{NLi}})_2 (\operatorname{ArLi}) + i-\Pr_2 \operatorname{NH}$$
(9)

Mixed Dimer Isomerization (Step II)

$$(i-\Pr_2 \operatorname{NLi}_4)(\operatorname{ArLi}) \stackrel{k_2}{\underset{k_{-2}}{\rightleftharpoons}} (i-\Pr_2 \operatorname{NLi}_5)(\operatorname{ArLi})^*$$
(10)

Mixed Dimer-Based Metalation (Step III)

$$(i-\Pr_2 \operatorname{NLi}_5)(\operatorname{ArLi})^* + \operatorname{ArH}_1 \xrightarrow{\kappa_3} 2\operatorname{ArLi}_2 + i-\Pr_2 \operatorname{NH}$$
 (11)

LDA-ArLi Condensation (Step IV)

$$(i-\Pr_2 NLi)_2 + \operatorname{ArLi}_2 \stackrel{\star_4}{\underset{k_{-4}}{\longrightarrow}} (i-\Pr_2 NLi)(\operatorname{ArLi}) + \frac{1}{2}(i-\Pr_2 NLi)_2 \quad (12)$$

A discussion of the model is prefaced by a few essential points. First, we used standard kinetic methods to measure the full rate laws for the key rate-limiting steps (eqs 9 and 12). The THF dependencies were also measured, but they add more complexity than clarity. Thus, we exclude them from the discussion and simply note that the model is consistent with data acquired at all THF concentrations. Similarly, the differential equations used in the numeric integrations add little to the discussion; they are presented and discussed in the Experimental Section.

⁽³²⁾ Hughes, E. J. Chem. Educ. 1989, 66, 46.

⁽³³⁾ We define the idealized rate law as that obtained by rounding the observed reaction orders to the nearest rational order.

⁽³⁴⁾ The rates rose with decreasing THF concentrations. A seemingly analogous inverse THF dependence for the reaction of an aryllithium has been reported.²¹

Second, models containing large numbers of differential equations and adjustable parameters are said to be "sloppy" if they are highly under determined.³⁶ Cannot *any* model with such a plethora of adjustable parameters fit the data? Can the model "fit an elephant and make it wag its tail"?³⁷ The answer is no (for the first question, at least). The model is *not* sloppy. The adjustable parameters corresponding to the six rate constants are well defined. Moreover, many other models that were poked and prodded included provisions for autocatalysis, linear (or sigmoidal) decays, or delayed mixed aggregate formation but failed to account for *all* behaviors. The model has one compelling strength when compared with all other models that we considered and dismissed: *It fits the data in every instance*. The numeric integrations illustrated in Figures 9–11 are emblematic of the quality of the fits.

To facilitate the discussion, the model is described as four discrete steps, and the origins of each as well as the portions of the model that are negotiable are discussed. To minimize confusion, we also note at the outset that, although facile mixed dimer-based lithiation of carbamate **1** is necessary, the condensation of ArLi with LDA dimer in eq 12—an aggregation event, not a metalation²⁵—is the key rate-limiting step in the autocatalysis.

LDA Dimer-Based Metalation. The first stage of the metalation of carbamate 1 (Scheme 1, Step I)—the reaction before the appearance of lithium salt products—was shown using standard methods to be LDA dimer-based as described by eq 9 and represented by transition structure 10. Temperature dependencies (discussed below) are consistent with mixed dimer 4 as the first-formed product.

Mixed Dimer Isomerization. Equation 10 represents the exchange between cyclic dimer **4** and open dimer **5** (Scheme 1, Step II). Several experiments showed that at -78 °C **4** and **5** interconvert rapidly on the time scales that they are formed or consumed. By contrast, at -100 °C, the rate of exchange becomes slow relative to the rate of mixed dimer-based metalations, which allows us—actually forces us—to distinguish which dimer is the reactive form (vide infra).

Mixed Dimer-Based Metalation. It almost goes without saying that an autocatalytic organolithium reaction involves highly reactive mixed aggregates as described by eq 11 (Scheme 1, Step III). Spectroscopic studies using rapid-injection methods^{25a,b} confirm that the resulting mixed dimers react almost instantaneously at -78 °C. In some sense, this mixed dimer-based metalation is of lesser consequence to the model than one might suspect because it is a post-rate-limiting step. Their high reactivities, however, explain two empirical observations: (1) the delayed appearance of the mixed dimers **4** and **5**, and (2) the highly temperature sensitive concentration dependencies. Both are discussed below.

LDA-ArLi Condensation. The rate-limiting step in the autocatalysis is the condensation of aryllithium 2 with LDA dimer 3 (Scheme 1, Step IV), which proceeds via an LDA dimer-based mechanism.³⁵ The reaction coordinate is likely to be more complex than described by eq 12, however, in that the

first-formed product is logically some form of the $(i-Pr_2NLi)_2(ArLi)$ trimer. Whether the putative trimer is akin to previously characterized cyclic trimers is unknown,^{5,9} but it is too fleeting to detect spectroscopically and is not germane to the numeric integrations.

Origins of Delayed Mixed Aggregation. Following an organolithium reaction using NMR spectroscopy reveals that the starting organolithium aggregates are replaced by new product-derived mixed and homoaggregates.⁵ The aggregate distributions change as the reaction proceeds, but equilibrium populations are maintained³⁸ *provided that aggregate exchange is rapid on the time scales of the reaction.* This is not the case in the metalation of carbamate **1**. Mixed aggregates **4** and **5** appear only after carbamate **1** is consumed. The rapid metalation in eq 11 accounts for the delayed appearance of the mixed dimers; the mixed dimers do indeed form but are consumed rapidly by carbamate **1** as they form.

Relative Mixed Dimer Reactivities. Unraveling the relative importance of isomeric mixed dimers 4 and 5 was particularly challenging. It was *very* tempting at the outset to ignore dimer 5 altogether given that 5 forms in low concentrations ($\approx 3\%$). Although 5 was confirmed to be an isomer of 4, its three-dimensional structure was assigned using exclusively DFT computations. Moreover, using data collected at -78 °C, conditions under which 4 and 5 equilibrate rapidly on the time scales of the metalation, it is not constructive to consider which is the so-called reactive form; 4 and 5 simply represent two resting states. At -100 °C, however, 4 and 5 interconvert slowly on the time scale of the metalation. Under these conditions, it is valid to consider the relative reactivities of 4 and 5 toward arene 1.

The model described in eqs 9-12 that is based on metalations by mixed dimer **5** fits the data, whereas alternative models based on the reaction of **4** do not. The alternative models fail to include provisions for the elevated and constant concentration of **4** during the consumption of carbamate at -100 °C (Figure 11). When the model presented is expanded to include provisions for reaction of both **4** and **5**, the fit suggests <1% contribution from a **4**-based metalation.

The scenario that is fully consistent with the data is that 4 isomerizes to 5 and dimer 5 subsequently reacts with carbamate 1 (see Scheme 1). The isomerization is fast relative to the metalation at -78 °C, whereas the isomerization becomes rate limiting at -100 °C. It was in this context that we wondered whether the transition structure for the metalation may show a ring-expanded motif similar to that of 5 rather than a more conventional closed form with a Li–C–Li contact as in 4.

Computational studies of the mixed dimer-based metalation are summarized in Scheme 3. Dimers 4 and 5 are represented by simplified reactants 9 and 7, respectively. Transition structures 12 and 13 are analogous to 9 and 7 in that conventional Li-C-Li contacts in 9 and 12 are absent in 7 and 13. Curiously, the calculated activation energies are also qualitatively consistent with experiment. Most significant, the computations suggest that the ring-expanded motif is relatively unfavorable in reactant 7 yet stabilizing in transition structure 13.

Origins of Linear Decays. The model reproduces the linear decays by superimposing a downward curvature from autocatalysis on the normal upward curvature resulting from loss of

⁽³⁵⁾ The rate law provides the stoichiometry of the transition structure relative to that of the reactants: Edwards, J. O.; Greene, E. F.; Ross, J. J. Chem. Educ. 1968, 45, 381.

⁽³⁶⁾ Brown, K. S.; Sethna, J. P. Phys. Rev. E 2003, 68, 021904.

⁽³⁷⁾ A discussion and attribution of this famous aphorism is found in ref 36. Physicists, apparently with considerable free time, are reputed to have successfully drawn elephants using complex mathematics and numerous adjustable parameters.

^{(38) (}a) Casado, J.; Lopez-Quintela, M. A.; Lorenzo-Barral, F. M. J. Chem. Educ. 1986, 63, 450. (b) Alberty, R. A. J. Chem. Educ. 2004, 81, 1206. (c) Hammes, G. G. Principles of Chemical Kinetics; Academic Press: New York, 1978; pp 14–15.



Figure 16. Influence of autocatalysis on the LDA-mediated metalation of carbamate 1 as described in eqs 9-12.

substrate.²⁷ That is not to say, however, that it is obvious *why* the model is successful or why the linearities persist over a wide range of initial LDA concentrations. This coincidence requires elaboration as follows.

Using some abstraction, we can distill the model to its simplest components. In the absence of autocatalysis, the metalation proceeds via an LDA dimer-based mechanism (eq 9). Let us recast this as eq 13 and the affiliated rate constant as k_{init} . Similarly, the autocatalytic pathway occurs via an LDA dimer-based rate-limiting condensation with ArLi. The autocatalysis simplifies to eq 14 with a rate constant denoted as k_{cat} .

$$(i-\Pr_2 NLi)_2 + ArH \xrightarrow{k_{init}} product$$
 (13)

$$(i-\Pr_2 \text{NLi})_2 + \text{ArLi} \xrightarrow{k_{\text{cat}}} \text{product}$$
 (14)

If only the uncatalyzed ortholithiation in eq 13 is operative, the reaction will follow a normal exponential decay (Figure 16, curve A). At 50% conversion of ArH, the reaction rate (the tangent to the curve A) would be 50% of the initial rate. For autocatalysis to straighten curve A to perfect linearity (curve B), the deceleration caused by the depletion of ArH must be precisely offset by the acceleration caused by the buildup of ArLi (eq 14). *Linearity is observed when* $k_{init} = k_{cat}$. Indeed, inspection of Figure 9 shows that the instantaneous slopes for the loss of carbamate 1 at early conversion and the loss of aryllithium monomer 2 (the initial slope to the right of its apex) are equal.

Is it a coincidence that the rates of the uncatalyzed and the catalyzed ortholithiation are equal? On one level, yes. Decreasing the THF concentration or introducing orthodeuteration results in sigmoidal behavior consistent with a relatively higher rate for the catalyzed process (Figure 16, curve C). (k_{init} is isotopically sensitive, whereas k_{cat} is not.) Nonetheless, through the reductionist approach described by eqs 13 and 14, one can see haunting similarities. Is there a common reactive intermediate such as an LDA open dimer²³ that reacts with ArH or ArLi (eq 15) with nearly equal efficacy? Although it is unclear why the ArH and ArLi react so similarly (and certainly further speculation is getting excessive), we wonder whether the Lewis basic³⁹ fluoro moieties are important.

$$(i-\Pr_2 NLi)_2 \rightleftharpoons (i-\Pr_2 NLi)_2^* \xrightarrow{\text{ArH or ArLi}} \text{products}$$
 (15)

Conclusion

LDA continues to harbor mysteries and offer insights into how aggregation and solvation influence organolithium reactivity. The LDA-mediated ortholithiation delineated in eq 1 afforded linear and sigmoidal decays—evidence of autocatalysis—and exceptional reactivities of mixed aggregates that are unexpected in the context of previously studied LDA-mediated metalations.² The model presented, although somewhat flexible in detail and vulnerable to revision, nicely accounts for the odd observations.

Taking a wider view, these results are not merely curiosities. As we peer into the past, we are reminded of LDA-mediated enolizations in THF at $-78 \,^{\circ}C^{10c,40,41}$ in which LDA–lithium enolate mixed aggregates markedly altered *E/Z* selectivities.⁴² We can also offer a peek into the future. Ongoing studies are uncovering many ortholithiations that are autocatalytic. Both the similarities and substrate-dependent differences are striking. We have also discovered that Michael additions to unsaturated esters, imine metalations, and ester enolizations display such anomalous behavior.⁴³ All of these reactions share a common denominator: They are effected using LDA in THF at $-78 \,^{\circ}C$. This is exciting from a mechanistic perspective. The prevalence of the autocatalysis, however, is also disquieting given the importance of this particular base–solvent–temperature combination in organic chemistry.

Experimental Section

Reagents and Solvents. THF, hexane, and pentane were distilled from blue or purple solutions containing sodium benzophenone ketyl. The hydrocarbon stills contained 1% tetraglyme to dissolve the ketyl. LDA was prepared from *n*-BuLi and subsequently rescrystallized.¹² The purified LDA was shown by both potentiometry⁴⁴ and ion chromatography⁴⁵ to contain $\leq 0.02\%$ LiCl. Airand moisture-sensitive materials were manipulated under argon or nitrogen using standard glovebox, vacuum line, and syringe techniques. Solutions of *n*-BuLi and LDA were titrated for active base using a literature method.⁴⁶

NMR Spectroscopic Analyses. All NMR samples were prepared using stock solutions and sealed under partial vacuum. Standard ⁶Li, ¹³C, ¹⁵N, and ¹⁹F NMR spectra were recorded on a 500 MHz spectrometer at 73.57, 125.79, 50.66, and 470.35 MHz (respectively). The ⁶Li, ¹³C, ¹⁵N, and ¹⁹F resonances are referenced to 0.30 M [⁶Li]LiCl/MeOH at -90 °C (0.0 ppm), the CH₂O resonance of THF at -90 °C (67.57 ppm), neat Me₂NEt at -90 °C (25.7 ppm), and C₆H₅F in neat THF at -78 °C (-113.15 ppm), respectively.

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IR Spectroscopic Analyses. Spectra were recorded using an in situ IR spectrometer fitted with a 30-bounce, silicon-tipped probe. The spectra were acquired in 16 scans at a gain of 1 and a resolution of 8 cm⁻¹. A representative reaction was carried out as follows: The IR probe was inserted through a nylon adapter and FETFE O-ring seal into an oven-dried, cylindrical flask fitted with a magnetic stir bar and a T-joint. The T-joint was capped by a septum for injections. Following evacuation under full vacuum and flushing with N_2 , the flask was charged with a solution of LDA (108.2 mg, 1.01 mmol) in THF (10.0 mL) and cooled in a freshly prepared dry ice/acetone cooling bath. (Condensation of moisture adversely affects temperature consistency.) After recording a background spectrum, carbamate 1 was added to the LDA/THF mixture at -78°C from a dilute stock solution (100 μ L, 0.404 M) with stirring. IR spectra were recorded over the course of the reaction. To account for mixing and temperature equilibration, spectra recorded in the first 1.5 min were discarded. All reactions were monitored to >5 half-lives.

Rate Studies Using ¹⁹**F NMR Spectroscopy.** The rates of metalation and mixed aggregation were monitored using ¹⁹F NMR spectroscopy. Spectroscopic samples were prepared using either standard methods described previously or by injecting carbamate **1** via 25 μ m i.d. flexible capillary tubing by gas-tight syringe directly into the solution of LDA at -78 °C in the NMR probe.^{25a,b} The methyl ¹H resonances on the magnetically inequivalent *i*-propyl groups on carbamate **1** coalesce on warming above -70 °C as described previously. Monitoring the decoalescence to give two distinct resonances showed that the temperature equilibration occurs in <30 s. T_1 relaxation times were determined for all species, and the delay between scans was set to $>5 \times T_1$ to ensure accurate integrations.

The condensation of **2** with LDA to form mixed aggregates **4** and **5** was easily monitored in situ because the condensation occurs only after **1** is consumed. Loss of **2** followed clean exponential decays, and formation of **4** correlated quantitatively. Results obtained from fitting the decay of **2** $[A = A_0 \exp(-k_{obsd}t) + C]$ or formation of **4**{ $A = A_0[1 - \exp(-k_{obsd}t)]$ } afforded equivalent first-order rate constants (±10%).

Numeric Integrations. The proposed mechanism (eqs 9-12) can be expressed as a series of differential equations as follows:

$$d[\text{ArH}]/dt = -k_1[\text{ArH}][\text{A}_2] - k_3[\text{A} \cdot \text{ArLi}^*][\text{ArH}]$$
(16)
$$d[\text{ArLi}]/dt = -k_4[\text{ArLi}][\text{A}_2] + k_4[\text{A} \cdot \text{ArLi}][\text{A}_2]^{0.5} + 2k_3[\text{A} \cdot \text{ArLi}^*][\text{ArH}]$$
(17)

 $d[\mathbf{A} \cdot \mathbf{ArLi}]/dt = k_1[\mathbf{ArH}][\mathbf{A}_2] - k_2[\mathbf{A} \cdot \mathbf{ArLi}] +$

$$k_{-2}[\mathbf{A} \cdot \mathbf{ArLi}^*] + k_4[\mathbf{ArLi}][\mathbf{A}_2] - k_{-4}[\mathbf{A} \cdot \mathbf{ArLi}][\mathbf{A}_2]^{0.3} \quad (18)$$

$$d[\mathbf{A} \cdot \mathbf{ArLi}^*]/dt = k_2[\mathbf{A} \cdot \mathbf{ArLi}] - k_{-2}[\mathbf{A} \cdot \mathbf{ArLi}^*] - k_{-2}[\mathbf{A}$$

$$\frac{k_{2}[A \cdot ArLi] - k_{2}[A \cdot ArLi]}{k_{3}[A \cdot ArLi^{*}][ArH]}$$
(19)

$$[A_2] = [A_2(init)] - 0.5[ArLi] - [A \cdot ArLi] - [A \cdot ArLi^*]$$
(20)

where ArH is arene 1, ArLi is monomer 2, A_2 is LDA dimer 3, A•ArLi is mixed dimer 4, A•ArLi* is mixed dimer 5, and [A₂(init)] is the initial concentration of LDA.

The time-dependent concentrations measured by ¹⁹F NMR spectroscopy are fit to a mechanistic model expressed by the differential equations. The curve fitting operation minimizes χ^2 in searching for the coefficient values (rate constants). The Levenberg–Marquardt algorithm⁴⁷ is used for the χ^2 minimization and is a form of nonlinear, least-squares fitting. The fitting procedure implements numeric integration based on the backward differentiation formula (BDF)⁴⁸ to solve the differential equations, yielding functions describing concentration versus time.

The results from the fits are as follows. From a single run, the rate constant for the LDA dimer-based metalation (k_1 in eq 9) can be uniquely determined. The equilibrium constant for the exchange of mixed dimers **4** and **5** (k_2/k_{-2}) is also well defined, yet only a lower limit can be established for the individual values of k_2 and k_{-2} (k_2 , $k_{-2} > 10 \cdot k_1$). The rate constant corresponding to metalation of carbamate **1** by mixed dimer **5** (k_3 in eq 11) has a lower boundary (>10⁵ \cdot k_1). The k_4/k_{-4} ratio corresponding to the monomer-mixed dimer equilibrium (eq 12) is well defined as are the separate values of k_4 and k_{-4} . By carrying out a global fit of multiple data sets, the quality of the individual fits are decreased somewhat as expected for a reaction that is sensitive to initial conditions. We find, however, that best-fit values of the rate constants from the global fit do not differ much from the individual fits.

The data recorded at -100 °C offer insights into the relative rate constants unavailable at -78 °C (vide infra). The individual values of k_2 and k_{-2} become well defined, which ultimately leads to the conclusion that LDA dimer-mediated metalation affords **4** and the mixed dimer-mediated metalation proceeds via **5** as described in the discussion.

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Supporting Information Available: NMR spectra, rate and computational data, experimental protocols, details on the numeric integrations, and a complete list of authors for ref 19. This material is available free of charge via the Internet at http:// pubs.acs.org.

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