Sodium Diisopropylamide: Aggregation, Solvation, and Stability

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**Supporting Information

**ABSTRACT:** The solution structures, stabilities, physical properties, and reactivities of sodium diisopropylamide (NaDA) in a variety of coordinating solvents are described. NaDA is stable for months as a solid or as a 1.0 M solution in N,N-dimethylethylamine (DMEA) at −20 °C. A combination of NMR spectroscopic and computational studies show that NaDA is a disolvated symmetric dimer in DMEA, N,N-dimethyl-n-butylamine, and N-methylpyrrolidine. Tetrahydrofuran (THF) readily displaces DMEA, affording a tetrasolvated cyclic dimer at all THF concentrations. Dimethylethylamine (DME) and N,N,N′,N′-tetramethylethylenediamine quantitatively displace DMEA, affording doubly chelated symmetric dimers. The trifunctional ligands N,N,N′,N″,N″-pentamethyldiethylenetriamine and diglyme bind the dimer as bidentate rather than tridentate ligands. Relative rates of solvent decompositions are reported, and rate studies for the decomposition of THF and DME are consistent with monomer-based mechanisms.

**INTRODUCTION**

Several groups, most recently that of Mulvey, have underscored the merits of sodium dialkylamides, but the response of the synthetic organic community remains muted compared with the enthusiastic embrace of lithium diisopropylamide (LDA) and related lithium dialkylamides. Sodium diisopropylamide (NaDA) is an excellent case in point. Since the initial preparation of NaDA by Levine in 1959 and subsequent improved preparations, NaDA has languished in relative obscurity, appearing in the literature only a dozen times over half a century. This scarcity is somewhat perplexing on first inspection. NaDA is easily prepared, stable as a solid if refrigerated, and a powerful Brønsted base. We surmised that potential users must be leery of the rapid destruction of standard ethereal solvents by NaDA and its insolubility in inert hydrocarbons. In short, NaDA is inconvenient.

In our first paper of this series, we reported that 1.0 M solutions of NaDA in neat N,N-dimethylethylamine (DMEA) or DMEA–hydrocarbon mixtures are stable for weeks at room temperature and for months under refrigeration. Moreover, NaDA–DMEA can be prepared in 15 min from technical-grade reagents without pre-purification or pre-drying. Metalations of a dozen substrates with a broad range of functionality have shown that NaDA–DMEA is often orders of magnitude more reactive than LDA–THF toward orthometalations, dehydrohalogenations, diene metalations, and epoxide eliminations. The steric and chemoselectivities of the two bases are often complementary. Even concerns that the sodiated intermediates and products in DMEA might be insoluble have proved unfounded. Overall, the results of our initial studies were highly encouraging, and, as a reviewer noted, “running reactions in trialkylamine solvent is not crazy.”

In this second paper, we examine the solution structure and stability of NaDA solvated by DMEA, other trialkylamines, and a number of synthetically important coordinating ligands including tetrahydrofuran (THF), N,N,N′,N″-tetramethyl-ethylenediamine (TMEDA), and 1,2-dimethoxyethane (DME). The cyclic dimer motif (1a–h) is the only detectable form. DMEA is substitutionally labile, which is a critical prerequisite for the addition of other ligands before metalation to modulate the reactivity of NaDA and after metalation to control reactivity of the resulting sodium salts. Mechanistic studies of solvent decomposition offer the first glimpse into NaDA reactivity. This paper is intended to detail the structural foundations underpinning NaDA structure–reactivity–selectivity principles of potential interest to synthetic organic chemists.

**RESULTS**

**Methods.** We modified the dissolving metal method first described by Wakefield and co-workers by using kinetically inert and solubilizing DMEA as the bulk solvent. Although the procedure affords 1.0 M stock solutions of NaDA adequate

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for synthetic applications, a precautionary crystallization from DMEA−hexane was included for spectroscopic and rate studies. We hasten to add that the typical user will detect little or no difference between the pre- and post-purified reagent.

Given the absence of resolved $^{23}$Na−$^{15}$N coupling owing to the quadrupolar $^{23}$Na nucleus and with an eye toward expanding the application of the method of continuous variations (MCV) to aggregated organometallic species lacking NMR-active metal nuclei, we turned to MCV to assign the solution structure. In short, an ensemble generated from constitutionally similar species of unknown aggregation number ($A_n$ and $B_n$, eq 1) is monitored with NMR spectroscopy as a function of mole fraction $X_A$ or $X_B$. The number of heteroaggregates attests to the aggregation state. Plotting the relative proportions versus mole fraction affords a Job plot confirming the assignment.

$$A_n + B_n \rightarrow A_{n-1}B_1 + A_{n-2}B_2 + \ldots + B_n$$ (1)

For this study, NaDA was paired with sodium dicyclohexylamide (NaDCA)$^{5,10}$ and sodium isopropylcyclohexylamide (NaICA).$^{9a}$ NaICA was previously suggested to be dimeric, as evidenced by two stereoisomers ($cis$-2 and $trans$-2, eq 2; Cy = cyclohexyl).$^{9a}$ We confirmed the dimer assignment for NaICA and showed that both NaDA and NaDCA are dimers 1 and 3, respectively, by examining NaDA−NaICA and NaDA−NaDCA mixtures using $^{13}$C and $^{15}$N NMR spectroscopies. NaDA−NaDCA pairings are superior and are emphasized below. Considerable data for the NaDA−NaICA pairings are archived in the Supporting Information. Although $^1$H NMR spectroscopy proved especially valuable in MCV-based studies of sodium enolates,$^{9a,11}$ the resolution was inadequate for the study of sodium dialkylamide aggregation.

The strategies and quantitative insights into solution solvation numbers of NaDA are notable in our opinions. Confirmations and experimentally elusive details are provided with density functional theory (DFT) computations at the B3LYP/6-31G(d) level,$^{12}$ with single-point calculations at the MP2 level of theory.$^{13}$

**Solution Structure: NaDA in DMEA and Related Amines.** $^{13}$C NMR spectra of NaDA−NaDCA mixtures in DMEA showed the dimer ensemble depicted in eq 3 with resolution of all $^{13}$C methine resonances of the isopropyl and cyclohexyl moieties (Figure 1a). A plot of the relative integrations of 1, 3, and 4 afforded the Job plot illustrated in Figure 2, which is characteristic of statistically distributed dimers.

A $^{15}$N NMR spectrum of a 1:1 mixture of $[^{15}$N]NaDA and $[^{15}$N]NaDCA shows signals corresponding to two homodimers and a heterodimer distributed statistically (Figure 1b). A Job plot based on $^{15}$N NMR data is analogous to that shown in Figure 2 (Supporting Information).

MCV analyses using the NaDA−NaICA mixtures (eq 4) were effective, but they have been largely relegated to Supporting Information. Several interesting observations are worth noting, however. The $^{13}$C NMR data resolved two methinyl resonances in each of the two NaICA homodimers as well as the four methinyl resonances in heterodimer 5 (eq 4), readily affording the corresponding Job plot. $^{15}$N NMR
spectroscopy using $^{15}$NNaDA and $^{15}$NNaICA showed that the $^{15}$N resonances of the homo- and heteroaggregated NaICA fragments are poorly resolved, whereas homo- and hetero-aggregated NaDA fragments are well-resolved. Poor resolution on one pair is not critical; the resulting Job plot is consistent with a statistical distribution of dimers.

$^{13}$C and $^{15}$N NMR spectroscopic investigations of the NaDA−NaDCA pair in N,N-dimethylbutylamine (DMBA) and N-methylpyrrolidine showed that NaDA is dimeric (Supporting Information). DFT computations showed that sequential solvation of dimeric NaDA by DMEA is highly exothermic and affords disolvate 1a, but no minima corresponding to tri- or tetrarsolvated dimers were found. The sterically less demanding and computationally simpler Me$_3$N gave similar results. This outcome contrasts sharply with ethereal and multidentate ligands (vide infra).

Table 1. Molecular Weight (MW) of Sodium Diisopropylamide in Dimethylethylamine and Tetrahydrofuran Determined with $^1$H Diffusion-Ordered Spectroscopy

<table>
<thead>
<tr>
<th>aggregate</th>
<th>MW</th>
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<tr>
<td>NaDA (1)</td>
<td>393</td>
</tr>
<tr>
<td>NaDCA (2)</td>
<td>534</td>
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</table>

Owing to our experience with DFT computations in lithium chemistry, we were comfortable with the assignment of 1a as disolvated. Nonetheless, we applied several experimental methods to determine the solution as follows. Proton diffusion-ordered NMR spectroscopy ($^1$H-DOSY), first applied to organolithium aggregates by Williard,$^{15}$ is now proliferating within the field$^{16}$ and has been applied to organosodliums.$^{15}$ Underlying assumptions about molecular shape and influence on the diffusion constant have always left us uneasy, however.

We examined the structure of NaDA in DMEA using the double bipolar pulse pair stimulated echo sequence with convection compensation pulse sequence to measure the diffusion coefficients of NaDA dissolved in DMEA. An analogous experiment independently measured diffusion coefficients for DMEA, THF, tetramethylsilane, anisole, 1,3-dimethoxybenzene, and 18-crown-6 to obtain the molecular weights for NaDA shown in Table 1. Although the data recorded at ambient temperature seemed to confirm the disolvated dimer, the molecular weight at $-80$ °C was 25% low. Similar temperature dependencies have been noted by others.$^{16,17}$ We have no evidence, however, that the reduced value at $-80$ °C is based on a structural change: low temperature promotes rather than retards solvation owing to negative enthalpy. Also, the solubility of NaDA in DMEA is nearly temperature-independent. The results from THF solvate 1d (see Table 1) are discussed below.

A more traditional probe of solubility provided significant insight.$^{18}$ The concentration of DMEA-solubilized NaDA in a suspension of NaDA in toluene-$d_8$ was monitored as a function of added DMEA using $^1$H NMR spectroscopy and benzene as an internal standard; the two-phase equilibrium is described in eq 5.$^{19}$ If DMEA coordinates to and solubilizes NaDA quantitatively ($K_{\text{solv}} \gg 1$), the concentration of NaDA will be proportional to the concentration of DMEA until full solubility is achieved, with the solvation number being extractable from the slope and the end point (eq 8). We see this behavior for superior solvents (vide infra). In the limit of weak binding ($K_{\text{solv}} \ll 1$), the solubility will be low while manifesting curvature diagnostic of the coordination number (eq 9). The measured concentration of NaDA in solution versus added DMEA (Figure 3) showed non-limiting behavior: it is upwardly curving with a fit to eq 7 showing disolvated dimer ($n = 2$). Forcing the model to a tetrarsolvated dimer ($n = 4$) provided a markedly inferior fit to the data.

$$\begin{align*}
(A)_\text{solid} + nS & \rightleftharpoons (A_2S_n)_\text{solution} \\
K_{\text{solv}} &= [A_2S_n]/[S]^n \\
[A_2S_n] &= K_{\text{solv}}([S]_0 - n[A_2S_n])^n \\
[A_2S_n] &= [S]_0/n \\
[A_2S_n] &= K_{\text{solv}}[S]_0^n
\end{align*}$$

Thus, recalcitrant solubilization of NaDA by DMEA reflects weak binding. This notion is reinforced by studies of additional monodentate trialkylamines (Supporting Information). As steric demands increase, the binding constant, $K_{\text{solv}}$, decreases. The poor solubility of NaDA in triethylamine, for example, stems from weak coordination rather than a low solubility of

Figure 3. Plot of NaDA concentration versus total DMEA concentration in toluene at room temperature fit to eq 7 and presuming one DMEA per sodium (solid line, $n = 2$ [set]),$^{20}$ $K_{\text{solv}} = 0.072 \pm 0.004$ and two DMEA per sodium (dashed line, $n = 4$ [set], $K_{\text{solv}} = 0.007 \pm 0.002$).
the doubly solvated dimer. We return to this idea in the Discussion.

**Solution Structure: NaDA–THF.** MCV using $^{13}$C and $^{15}$N NMR spectroscopies for NaDA–NaICA and NaDA–NaDCA pairs demonstrated that THF-solvated NaDA is dimeric at both low and high THF concentrations (Supporting Information). As usual, determining the solvation state demanded several strategies. Incremental additions of THF to NaDA in 2.0 M DMEA (Figure 4) showed clear saturation of the changing chemical shifts of the methine proton and $^{15}$N resonances resulting from ligand substitution consistent with strongly preferential THF coordination. However, curvature with saturation occurring at $\sim$3.0 equiv of THF per sodium (as opposed to a linear dependence with a sharp end point) belied non-quantitative substitution, thereby obscuring the stoichiometry. Fortunately, titration of a suspension of NaDA in toluene, as described for trialkylamines (Figure 5 and eqs 5–7), revealed a linear dependence of the measured titer on added THF and a constant 2:1 THF/NaDA ratio in solution up to full solubilization at 2.0 equiv. This outcome is fully consistent with tetrasolvate 1d.

The tetrasolvated form is also strongly supported computationally (eq 10). Given the exothermicity and our experience that DFT computations tend to falter with congested systems, we are confident in the tetrasolvate assignment.

**Solution Structure: NaDA–TMEDA.** MCV showed that the solution aggregation state of NaDA–TMEDA is akin to the reported doubly chelated crystal structure (Supporting Information). We hoped to observe free and bound TMEDA in the slow-exchange limit to directly measure the number of ligands on the bound form, but the exchange was rapid at $-100^\circ$C. Two titration methods showed strong coordination by TMEDA and supported a 1:1 proportion of NaDA/TMEDA:

1. Substitution of DMEA by TMEDA was evidenced by a downfield shift of the methinyl $^1$H resonance and an upfield shift of the $^{15}$N resonance after substitution of TMEDA by DMEA. As found for THF, however, the stoichiometry of the substitution was obscured by non-quantitative binding (Supporting Information).

2. Titration of a suspension of NaDA with TMEDA (analogous to that in Figure 5) showed a linear dependence of the measured titer on TMEDA concentration, a constant 1:1 proportion of TMEDA to NaDA in solution up to complete dissolution, and a hard solubility end point corresponding to 1:1 stoichiometry of TMEDA/NaDA.

DFT computations showed that the displacement of DMEA by chelated TMEDA (eq 11) is exothermic. This result contrasts with that observed with LDA, which forms a weakly ligated $\eta^1$-TMEDA-solvated dimer in solution.

**Solution Structure: NaDA–DME.** DME acts as an ethereal analogue of TMEDA in every respect: (1) $^{13}$C NMR spectroscopy in conjunction with MCV showed dimers, (2) incremental addition (akin to those in Figure 5) to NaDA–DMEA showed semi-quantitative binding consistent with one DME per sodium, (3) titration of a NaDA suspension showed a linear dependence of the titer on DME with a hard solubility end point at 1:1 stoichiometry, and (4) DFT computations supported a highly exothermic substitution of DMEA by DME without MP2 correction. We encountered a unique situation in which uncorrected energies and geometries corroborated experiment, yet MP2 correction provided highly questionable 1.6 kcal/mol/Na solvation energies (eq 11) for reasons we do not yet understand.
failed to identify. We explored a variety of larger basis sets with no obvious improvements. All evidence, including the results of experimental competition studies, suggested that the MP2 correction is indeed spurious.

Doubly chelated dimer 1f contrasts with LDA, in which two DMEs coordinated to the dimer are unchelated.\textsuperscript{2,24,25} We also competed TMEDA and DME by swapping one for the other at a fixed total ligand concentration corresponding to 2.0 equiv per sodium and monitored the chemical shifts of the time-averaged free and bound ligand. TMEDA is the stronger ligand by approximately 10-fold.

**Solution Structure: NaDA with Other Ligands.** The results of \textsuperscript{15}N NMR spectroscopy and MCV showed that trifunctional ligands diglyme and \textit{N,N,N,N',N\texttextendash}pentamethyldiethylenetriamine (PMDTA) afford dimers rather than monomers that could have arisen from tridentate ligation. Their binding affinities are slightly greater than those of the bidentate analogues DME and TMEDA. Thus, these potentially tridentate ligands function as bidentate ligands, as demonstrated by 1g and 1h.\textsuperscript{26,27} Computations suggest that the third ligand, although not coordinated to sodium as evidenced by long \begin{math} \text{Na}^-\text{OMe} \end{math} and \begin{math} \text{Na}^-\text{NMe}_2 \end{math} bonds, display distinct preferences for orienting the lone pairs toward the sodium nuclei. We discuss the potential synthetic importance of bidentate rather than tridentate coordination below.

We carried out a brief survey of a number of ligands that lacked rigor but provided useful data nonetheless.\textsuperscript{28} Ethereal ligands such as \textit{Et}_2\textit{O} and 2,5-dimethyltetrahydrofuran substitute for DMEA but much more reluctantly than does THF as expected from binding measurements on lithium amides.\textsuperscript{23,26,27} Titration of NaDA/toluene suspensions with anisole displays chemical shift perturbations consistent with binding but no appreciable solubilization, suggesting that anisole is a poor ligand for sodium. The highly dipolar ligand \textit{N,N,N\texttextendash}dimethylpropyleneurea (DMPU) is quickly metalated by NaDA at \(-80\ °\text{C},\) as evidenced by the rapid appearance of extraneous resonances and disopropylamine observable with \textit{1H} NMR spectroscopy. The products of these decompositions have not been pursued.\textsuperscript{29}

**Solvent Decomposition: Products, Rates, and Mechanisms.**\textsuperscript{30,31} Whether in DMEA or in its solid form, NaDA at room temperature has a half-life of approximately 2 months, consistent with the thermal sensitivity noted by Wakefield.\textsuperscript{6b} This decomposition is mitigated by storage at \(-20\ °\text{C}\) in a standard laboratory freezer. However, facile decompositions of DME and DMPU underscore the possible limitations of NaDA when used in conjunction with standard ethereal solvents. The stability of NaDA in selected solvents at room temperature is illustrated in Table 2.

THF decomposition offered our first view of the mechanism of NaDA-mediated metalations. NaDA decomposition in THF–hexane mixtures at 25 °C forms partially soluble \textit{cis}- and \textit{trans}-alkoxides \textit{7} observable with \textit{1H} NMR spectroscopy (Scheme 1). Quenching afforded known alcohols \textit{8} and \textit{9} in the proportions shown in Scheme 1. The presumed intermediate salt \textit{6} is observed at low (equilibrium) levels owing to facile NaDA-mediated isomerization to \textit{7}; the isomerization of \textit{6} under the reaction conditions was confirmed by adding \textit{8} to NaDA–THF–\textit{d}_8.

Monitoring the rate of THF decomposition by tracking the loss of \textit{1d} (\(\delta\) 3.11 ppm in \textit{1H} NMR) and the formation of disopropylamine showed that the reaction does not follow a first-order decay, which indicated that the decomposition does not occur from the observable dimer. Fitting the traces to the nonlinear Noyes equation\textsuperscript{32} afforded an average order of 0.68 (Figure 6), approximating a half order. Plotting initial rates versus THF concentration revealed a second-order THF dependence (Figure 7). The half-order rate constants are nearly independent of the NaDA concentration, albeit with a slight upward drift (Figure 8). The idealized rate law\textsuperscript{33} (eq 12) is consistent with the mechanism described in eq 13.\textsuperscript{34} Rate-limiting proton transfer was evidenced by a substantial kinetic isotope effect determined by comparing THF to THF–\textit{d}_8 (\(k_{1a}/k_{1d} = 6.9\).

\[
-d[A_2(\text{THF})_4]/dt = k[A_2(\text{THF})_4]^{1/2}[\text{THF}]^2
\]

\[
1/2 A_2(\text{THF})_4 + 2\text{THF} \rightarrow k_{1a} [A(\text{THF})_4]^{\ddagger}
\]

This decomposition is mitigated by storage at \(-20\ °\text{C}\) in a standard laboratory freezer. However, facile decompositions of DME and DMPU underscore the possible limitations of NaDA when used in conjunction with standard ethereal solvents. The stability of NaDA in selected solvents at room temperature is illustrated in Table 2. THF decomposition offered our first view of the mechanism of NaDA-mediated metalations. NaDA decomposition in

<table>
<thead>
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<th>ligand\textsuperscript{a}</th>
<th>half-life</th>
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<tr>
<td>DMEA</td>
<td>2 months</td>
</tr>
<tr>
<td>TMEDA</td>
<td>1 month</td>
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<tr>
<td>THF</td>
<td>1 h</td>
</tr>
<tr>
<td>DME</td>
<td>10 s</td>
</tr>
<tr>
<td>DMPU\textsuperscript{b}</td>
<td>&lt;1 s</td>
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\textsuperscript{a}Neat. \textsuperscript{b}0.30 M DMPU in DMEA.

![Figure 6](image_url) **Figure 6.** Plot of NaDA concentration versus time for the decomposition of THF (Scheme 1) of NaDA at 25 °C. The red curve depicts an unweighted least-squares fit to the function \(f(t) = [a^{1-n} - k(t - (1 - n))^{1/1-n}]\), where \(a = 0.3484 \pm 0.0002\), \(k = 1.061 \times 10^{-5} \pm 9 \times 10^{-8}\), and \(n = 0.698 \pm 0.002\).
The [AS₄]⁺ stoichiometry is consistent with an α deprotonation via transition structure 11-α to generate oxacarbenoid 12 as a precursor to carbene 13 (Scheme 2). Alternatively, a β metalation whether via a concerted E2-like elimination or 11-β discrete carbanion 14 with post-rate-limiting elimination to 6a is plausible. Although the isomerization and consequent scrambling dissuaded us from sophisticated isotopic labeling studies, we collected evidence supporting the carbenoid pathway. Comparing THF to THF-d₄ (15) afforded kᵢ/kₒ ≈ 6, suggesting a rate limiting C–H(D) cleavage at the 3 position, but it does not distinguish the two possible mechanisms. Monitoring a mixture of NaDA, THF, and i-Pr₂ND by ¹H and ³H NMR spectroscopies shows isotopic exchange at the α protons of THF, consistent with α deprotonation. The comparable rates of exchange and decomposition suggest that neither metatation nor insertion are dominantly rate limiting. The existence of discrete carbene 13, however, is not even assured.

NaDA in DME–toluene undergoes decomposition to give methyl vinyl ether and sodium methoxide (eq 14). Following the proton resonance of 1f at 61.10 ppm shows that the decay approximates a half-order rather than a first-order decay (akin to Figure 6). Fitting multiple decays to the nonlinear Noyes equation to ascertain the order by best fit affords an average order of 0.52. Plotting the half-order rate constants versus NaDA concentration showed some upward drift (akin to that in Figure 7). A plot of kₜₜₜₜ versus DME concentration approximated first order with a slight upward curvature (Figure 9), possibly hinting at either low contributions from a highly solvent-dependent pathway or generalized medium effects. The structure of 1f in conjunction with the idealized rate law (eq 15) afforded the generic mechanism in eq 16. The doubly solvated monomer-based transition state, [A(DME)_2]⁺, is isostructural to THF-based 11 when ligand hapticity is considered. Although DFT computations showed 16 and 17 to be computationally viable, 16 was preferred by ~21 kcal/mol. The low coordination number—the absence of a "Na-
(DME)₃ fragment—argues against a free-ion-based mechanism.

\[
-d[A_2(DME)_2]/dt = k[A_2(DME)_2]^{1/2}[DME]^1
\]  
(15)

\[
1/2 A_2(DME)_2 + DME \rightarrow [A(DME)_2]^e
\]  
(16)

## DISCUSSION

The first paper in this series showed that NaDA is a highly efficacious Brønsted base when compared with LDA. Our seemingly trivial but potentially consequential contribution at the outset was to show that NaDA is soluble and stable in DMEA and related minimally hindered trialkylamines. We previously asserted that simple trialkylamines have been largely overlooked as ligands for lithium salts, an oversight we attribute to the unfortunate urge to use them in concert with strongly coordinating ethereal ligands.

The structural studies described in this second paper lay foundations for subsequent studies that will dovetail synthetic organic applications with mechanistic investigations. We must suppress the almost irresistible urge to rely on analogy to lithium amides. Parallel behaviors exist, but such analogies are imperfect and demand, at a minimum, experimental support.

**Structures.** A survey of NaDA reveals cyclic dimers in solutions containing a number of standard coordinating ligands, analogous to the dominance of LDA dimers (Scheme 3). Even the smallest trialkylamines are so sterically demanding as to afford disolvated dimers of NaDA (1a−1c) akin to those of LDA. Compared with lithium, however, the larger sodium can accommodate more ligands. LDA dimer, for example, never exceeds one solvent per lithium in the solid or solution state (see 19) and is trisolvated endothermically by THF in silico.

Even bifunctional ligands such as DME and TMEDA remain uncchelated on the LDA dimer (20). By contrast, NaDA in THF forms tetratsolvated dimer 1d, and both TMEDA and DME readily chelate dimeric NaDA (1e and 1f). All substitute DMEA exothermically (see eqs 6 and 7).

**Comments on Methods.** We use a combination of tactics to understand structure-reactivity relationships and develop new strategies whenever possible. MCV as a method to study aggregation, for example, has its origins in early work from several laboratories and has been of enormous importance to us for studying aggregation in systems in which M−X coupling is not observable. To this end, our expansion of the method to include ¹³C and ¹⁵N as observable nuclei is new and noteworthy.

We also explored DOSY as a means to ascertain the structure of NaDA using diffusion molecular weights as a proxy. DOSY is increasingly popular, and our results could be considered supportive. That said, however, we remain cautious owing to significant (up to 25%) temperature-dependent changes in measured molecular weights that do not appear to derive from changes in structure.

**Solvent Decomposition.** We investigated the decomposition of THF and DME to understand the limitations of standard ethereal solvents as ligands for NaDA (Table 2) and get a first peek at the mechanism of NaDA-mediated metalations. THF decomposition occurred via tetrasolvated-monomer-based α-elimination (11). Although isotope effects show rate-limiting cleavage of the C−H(D) bonds at the β position (Scheme 2), unambiguous differentiation of a carbene-derived α-elimination (11-α) rather than E2-like β-metalation (11-β) was elusive owing to isotopic scrambling. From a synthetic organic (applications) perspective, the high solvent order shows that decomposition can be suppressed by using low ethereal ligand concentrations. Analogous rate suppression allows LDA/THF/hydrocarbon solutions to be sold commercially.

Facile decomposition of DME (eq 12) proceeded via a disolvated-monomer-based transition structure. Both 16 and 17 are computationally viable, with 16 energetically preferred. The

Scheme 3. Comparison of NaDA and LDA Structures
suppression of decomposition at low DME concentrations is especially crucial if DME is to find a niche given its lability and relatively high cost (see Table 2).

Thoughts on Solubilities. One might argue that our interest in the solubilities of NaDA in trialkylamines is excessive, but applications of NaDA–R₃N mixtures are predicated on the fact that high-molarity stock solutions are easily prepared, handled, and stored. Sodiated intermediates must also be soluble to be of value to synthetic chemists. There is, however, a previously undisclosed and subtle motivation for confirming NaDA solubility: ongoing rate studies of NaDA-mediated metalations in trialkylamines display odd rate behaviors that we cannot reconcile easily. Such discussions are beyond the scope of this paper but add to our obsession.

The solubility studies underscored some basic principles of alkali metal salt solubilities that, although not unprecedented, warrant further discussion. NaDA is highly soluble in DMEA and DMEA–hexane yet poorly soluble in triethylamine and TMEDA–hexane at low temperatures: why? This question is nuanced and is summarized in eq 18.

\[
\begin{align*}
K_{\text{sol}}(1) &= \frac{[A_2S_n]_{\text{solution}}}{[A_2S_n]_{\text{solid}}} \\
K_{\text{sol}}(2) &= \frac{[A_2S_n]_{\text{solid}}}{[A_2S_n]_{\text{solution}}} \\
&= \frac{1}{K_{\text{sol}}(1)}
\end{align*}
\]

Dissolving unsolvated solid NaDA, denoted as \((A_{n})_{\text{solid}}\), to form solubilized dissolved dimer stems largely from the enthalpy of primary shell solvation.\(^{14,42}\) the ligand must bind strongly enough to overcome the enthalpy of lattice deaggregation. The fact that the solubility of NaDA in trialkylamines is nearly temperature-independent shows that the two large enthalpic contributions cancel. The overall high solubility shows that \((A_2S_n)_{\text{solution}}\) is favorable relative to \((A_2S_n)_{\text{solid}}\) and \((A_{n})_{\text{solid}}\). Triethylamine, on the other hand, does not bind well; the poor solubility of NaDA correlates with a small binding constant, \(K_{\text{sol}}(1)\), rather than an inherent insolubility of \(A_2S_n\), reflected in \(K_{\text{sol}}(2)\). The failure of primary shell solvation causes the insolubility of NaDA in triethylamine.

By contrast, TMEDA binds strongly: \(K_{\text{sol}}(1)\) is large. However, doubly chelated dimer \(1e\) \((A_3S_1\) in eq 18) has limited solubility at low temperature. Therefore, to the extent that \(A_3S_1\) (and almost every organic molecule) is more soluble at high temperature, the solubilization of \((A_3S_1)_{\text{solid}}\) dimer is entropy-dominated.\(^{43}\) The failure of secondary shell solvation causes the insolubility of NaDA in TMEDA–hexane.

Synthetic Implications. In our first contribution,\(^{7}\) we emphasized the merits of NaDA–DMEA for metalations owing to its ease of handling and resistance to base-mediated solvent decomposition.\(^{30,31}\) However, the substutional lability of trialkylamines is also synthetically important. In forthcoming papers, we will explore the effects of ethereal ligands on reactivities of NaDA–DMEA mixtures. Ongoing studies show that THF elicits large accelerations relative to that of NaDA–DMEA while the metalation rate far exceeds the THF decomposition rate. The short half-life of NaDA in ethereal stock solutions by no means precludes their use as additives. Mono- and difunctional ethers readily displace coordinated DMEA in NaDA–DMEA stock solutions. Moreover, when robust trialkylamines are required to form a recalcitrant metalation, the resulting salts—arylsodims, for example—can be treated with ethereal ligands after the metalation to modulate their reactivities. Again, the ligand substitution will be highly favorable.

Hemilabile Trifunctional Ligands. Trifunctional PMDTA and diglyme afford dimeric NaDA bound only as bidentate ligands (1g and 1h). This outcome segues to a topic of longstanding interest: hemilabile ligands.\(^{44}\) In contrast to transition metal chemists attempting to build weakly chelating di- and polyfunctional ligands that readily liberate coordination sites,\(^{45}\) we approach hemilability by identifying ligands that chelate selectively in the transition state to optimize accelerations.\(^{44}\) Imagine, for example, transformations in which fleeting intermediates bearing a trifunctional ligand (eq 19) are markedly accelerated by transition-state stabilization that is not offset by ground-state stabilization. The reader might also realize that we have evidence to support this notion.

CONCLUSIONS

We are enthusiastic about the promise of NaDA–DMEA mixtures to solve metalation problems that plague synthetic chemists. NaDA is both convenient and demonstrably effective. It can be prepared in minutes using standard glassware and stored for months with refrigeration as 1.0 M DMEA solutions. Metalation rates are typically orders of magnitude higher than those for LDA–THF. The obvious limitations include the higher costs and unpleasant odors of trialkylamines, but neither precludes application to difficult metalations. Another point that should not be overlooked is that DMEA is substitutionally labile. Stronger ligands, even potentially expensive ligands, can be added before the metalation to accelerate it—they do—or after the metalation to modulate the reactivity of the resulting sodiated intermediate. Furthermore, quite unlike LDA, NaDA shows little tendency to form mixed aggregates.\(^{44}\) We are encouraged by the well-defined aggregation and solvation states described herein that are required to unravel mechanistic details of NaDA-mediated metalations and correlate them with selectivities.

EXPERIMENTAL SECTION

Reagents and solvents. THF, DME, diethyl ether, TMEDA, PMDTA, diglyme, hexane, and all trialkylamines were distilled from blue or purple solutions containing sodium benzophenone ketyl. \([^{13}N]\)diisopropylamine, \([^{13}N]\)dicyclohexylamine,\(^{46}\) and \([^{13}N]\)-isopropylcyclohexylamine have been described previously. NaDA was prepared from diisopropylamine, isoprene, DMEA, and sodium dispersion using a modified procedure first reported by Wakefield.\(^{47}\) A precautionary crystallization was used for the work described herein. Solutions of NaDA were titrated using a literature method.\(^{49}\) NaDCA was prepared with an optimized dissolving-metal-based preparation analogous to that used to prepare NaDA.\(^{26,27}\)

NaDCA. NaDCA is more conveniently prepared using n-butyldisodium rather than sodium dispersion because of its lower solubility. To a dry 15 mL pear flask charged with n-butyldisodium (33.0 mg, 0.413 mmol) was added DMEA (3.3 mL) at room temperature. On complete dissolution of the n-butyldisodium, neat cyclohexylamine (66.0 µL, 0.33 mmol) was added to provide a stock solution.\(^{13}\)
NMR spectrum (125.72 MHz, DMEA) δ 61.9, 41.6, 27.3, 26.7; 13N NMR spectrum (50.66 MHz, DMEA) 86.9.

**NMR Spectroscopic Analyses.** NMR samples for reaction monitoring were routinely prepared using stock solutions of NaDA and sealed under partial vacuum. DMEA-free solutions of NaDA with added ligands used DMEA-free crystallized NaDA. Samples were routinely flame-sealed except when used in experiments involving serial titration with a coordinating ligand. Standard 1H, 13C, and 15N spectra were recorded on a 500 MHz spectrometer at 500, 125.79, and 50.66 MHz, respectively. The 13C and 15N resonances were referenced to the CH3O resonance of THF at −90 °C (67.57 ppm) and neat Me2Ne at −90 °C (25.7 ppm), respectively.

**ASSOCIATED CONTENT**

* Supporting Information
   The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b03061.

* Structural, kinetic, and computational data; complete ref 12 (PDF)

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**Notes**

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