Electron Solvation and the Unique Liquid Structure of a Mixed-Amine Expanded Metal: The Saturated Li–NH$_3$–MeNH$_2$ System

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Abstract: Metal–amine solutions provide a unique arena in which to study electrons in solution, and to tune the electron density from the extremes of electrolytic through to true metallic behavior. The existence and structure of a new class of concentrated metal-amine liquid, Li–NH$_3$–MeNH$_2$, is presented in which the mixed solvent produces a novel type of electron solvation and delocalization that is fundamentally different from either of the constituent systems. NMR, ESR, and neutron diffraction allow the environment of the solvated electron and liquid structure to be precisely interrogated. Unexpectedly it was found that the solution is truly homogeneous and metallic. Equally surprising was the observation of strong long-range order in this mixed solvent system. This is despite the heterogeneity of the cation solvation, and it is concluded that the solvated electron itself acts as a structural template. This is a quite remarkable observation, given that the liquid is metallic.

Alkali metals demonstrate an exceptional solubility in NH$_3$, yielding intensely colored conducting solutions that have fascinated chemists since the time of Sir Humphry Davy.$^{[1,2]}$ Varying the concentration of metal in these liquids dramatically alters the electronic, magnetic, and structural properties of the solutions, and enables us to experimentally determine the manner in which liquid systems accommodate excess electron density. At a low concentration of metal/electrons, the solutions are electrolytic, whereby the metal valence electrons have been ionized into solution and exist as solvated electrons propagating between solvent cavities.$^{[3]}$ Increasing the concentration results in metallization in the liquid phase, which for the Li–NH$_3$ system occurs at a mere 4 mol% metal (MPM).$^{[3]}$ Interestingly at lower temperatures, below $T_c = 210$ K, the point of the Mott-type metal–insulator transition (MIT) is obscured by a pronounced liquid–liquid phase separation.$^{[2]}$ This illustrates that the localized and delocalized electron states do not readily co-exist, which is dramatically manifested by the fact that the more concentrated metallic solution floats above the dilute electrolytic phase for $T < T_c$. Above 8 MPM the solutions do not exhibit phase separation, appearing golden up to the concentration limit of 20 MPM,$^{[3]}$ the expanded metal Li(NH$_3$)$_4$.$^{[6,7]}$ The concentration and temperature dependence of this liquid–liquid phase separation has been mapped through multi-element NMR spectroscopy.$^{[8]}$

Along with metal concentration, chemical tunability of the electronic properties of these systems can be achieved by varying the amine. Lithium will also dissolve in MeNH$_2$ to yield a system whereby solvated electrons transition to a metallic state.$^{[8,9–12]}$ The MIT in Li–MeNH$_2$ occurs at 15 MPM, a notably higher concentration than in Li–NH$_3$. No liquid–liquid phase separation has been detected across the full concentration range of Li–MeNH$_2$, and the solution remains a deep blue, albeit with a metallic luster. The conductivity of liquid Li(MeNH$_2$)$_3$ is 400 $\Omega^{-1}$ cm$^{-1}$ (compared to 15000 $\Omega^{-1}$ cm$^{-1}$ in Li(NH$_3$)$_4$), which is close to Mott’s minimum conductivity limit, demonstrating that the Li–MeNH$_2$ system lies just on the metallic side of the MIT.

The differing metallic properties of Li–NH$_3$ and Li–MeNH$_2$ are also reflected in their distinct liquid structures.$^{[13–16]}$ All metallic metal amines share the trait of being highly structured liquids,$^{[13–16]}$ with distinct $M_{(sub)}$–$M_{(sub)}$ correlations. In both Li–NH$_3$ and Li–MeNH$_2$ solutions, Li is found to be four-coordinate, which has subsequently been found to be the case in the gas phase,$^{[17–19]}$ and through computational investigation.$^{[11,19–21]}$ The volumetric expansion of these liquid metals across their insulator–metal transition is also accompanied by the appearance of void spaces, which is presumably linked to the conduction electron density. In Li–NH$_3$ these voids take the form of channels between Li(NH$_3$)$_4$ units,$^{[18]}$ whereas in Li(MeNH$_2$)$_3$ the voids are spatially isolated.$^{[16]}$ This is concordant with magnetic measurements that suggest electronic conduction in Li(MeNH$_2$)$_3$ is via rapid migration of electrons between these polaronic voids, which begin to localize in the solid state.$^{[22–25]}$
The question then is how a mixture of Li(NH₃)₄ and Li(MeNH₂)₄ will behave. Surprisingly, perhaps, these mixed amine solutions have only previously been studied by optical absorption towards the limit of infinite dilution.[24] Herein we report the first studies of the liquid structure of Li-NH₃-MeNH₂ in the concentrated regime at 20 MPM, an empirical stoichiometry of Li(NH₃)₄·(MeNH₂)₄. We note that after equilibration these solutions appear to the eye as homogeneous red/bronze liquids (Figure 1), which upon dilution demonstrate a pronounced liquid–liquid phase separation whereby the lustrous phase floats above a deep-blue solution. We have observed this phase separation across a very large concentration range of Li, from 2–19 MPM. As confirmation of the homogeneity at 20 MPM, Figure 1 shows that only a single feature is witnessed in the corresponding ⁷Li NMR spectrum. This has proven to be an extremely sensitive indication of homogeneity in previous metal–amine studies.[8]

Interestingly, the ⁷Li peak appears at higher ppm than both Li(NH₃)₄ and Li(MeNH₂)₄,[8,12,25] suggesting a higher conduction electron spin-density at the Li nucleus (or indeed larger molar spin susceptibility) in the mixed amine than in either of the parent amines. The temperature dependence of the ⁷Li Knight shift[8] and the Dysonian lineshape of the ESR are also indicative of a homogeneous metallic system.[22–27]

Neutron diffraction is a uniquely powerful method for determining the liquid structure of metal amines, allowing for a number of isotopically distinct samples to be measured (defined in the Experimental Section below). Figure 2 presents an example fit to an experimental total structure factor, F(Q), from samples A and H in Table 1. Also shown is the difference spectrum [F₁₁(Q)−F₋₁₁(Q)] and its Fourier transform, the Li-centered total pair correlation function, ΔGₖ(r). The single principal peak at 1.75 Å⁻¹, and importantly the existence of a sharp pre-peak at 0.97 Å⁻¹, attest to the homogeneity of the 20 MPM solution. The pre-peak is a structural feature witnessed in solvated-electron systems as they transition to the metallic state, and arises owing to strong intermediate range ordering in the liquid.[13–15] Integration of the principal peak in the ΔGₖ(r) distribution corresponds to the coordination number of Li, and is consistent with approximately 4 solvent molecules, as reported previously for other Li–RNH₃ systems.[11,14]

**Figure 1.** Top left: ⁷Li NMR for 20 MPM Li(NH₃)₄·(MeNH₂)₄. Top right: Temperature dependence of the ⁷Li Knight shift. Bottom left: Homogenization of the 20 MPM system showing initial phase separation (left) and homogenous liquid (right). Bottom right: ESR line shape for 20 MPM Li(NH₃)₄·(MeNH₂)₄.

**Figure 2.** Top: Experimental total structure factors for 20 MPM ⁷Li–ND₃·CD₃ND₂ and ⁷⁷⁷Li–ND₃·CD₃ND₂ solutions. Middle: First-order Li difference structure factor. Bottom: Corresponding Li partial pair correlation.

**Table 1:** Isotopic substitutions in samples of 20 MPM Li–NH₃–MeNH₂.

| Sample | Li         | NH₃       | MeNH₂     |
|--------|------------|-----------|-----------|
| A      | ⁷⁷⁷Li      | ND₃       | CD₃ND₂    |
| B      | ⁷⁷⁷Li      | NH₃       | CD₃NH₂    |
| C      | ⁷⁷⁷Li      | [NH₃·ND₃] | [CD₃NH₂·CD₃ND₂] |
| D      | ⁷⁷⁷Li      | ND₃       | CH₃ND₂    |
| E      | ⁷⁷⁷Li      | ND₃       | [CD₃ND₂·CH₃ND₂] |
| F      | ⁷⁷⁷Li      | NH₃       | CH₃NH₂    |
| G      | ⁷⁷⁷Li      | [NH₃·ND₃] | [CD₃ND₂·CH₃NH₂·CD₃NH₂·CH₃ND₂] |
| H      | ⁷⁷⁷Li      | ND₃       | CD₃ND₂    |
To determine the spatial arrangement of lithium and amine species in the system, the empirical potential structure refinement (EPSR) technique was invoked to produce a structural model that was refined simultaneously to all the isotopically unique experimental $F(Q)$ functions (Table 1).[28]

Figure 3 shows the EPSR fits to the experimental data, with the EPSR model fitting well the experimentally obtained partial structure factors. The corresponding real-space distribution of solvent molecules around the lithium ions is given in

![Image](52x499 to 291x661)

**Figure 3.** Experimental (black) and modeled (red) partial structure factors for 20 MPM Li–NH$_3$–MeNH$_2$.

Figure 4. The Li–N distance for both NH$_3$ and MeNH$_2$ is about 2.0 Å, in close agreement with that observed for both Li(NH$_3$)$_4$ and Li(MeNH$_2$)$_4$. Integration of this first solvation shell gives an average co-ordination number of 2.01 and 1.78 for Li–MeNH$_2$ and Li–NH$_3$, respectively. The orientational distribution of the coordinated solvent about Li can be extracted from the EPSR model (Figure 4a–d). Only a small distortion from tetrahedral geometry for a given Li$_{\text{('solv)}}$ unit is found, with the MeNH$_2$ ligands acting to compress the tetrahedra slightly along the tetrahedral $C_3$ axis. The dipole moment of NH$_3$ is oriented directly towards the cation, whereas the Li–MeNH$_2$ angle is about 109°. The relative orientation of neighboring Li$_{\text{('solv)}}$ units is not isotropic, with the vertex (that is, the NH$_3$/MeNH$_2$ molecules) of one tetrahedral unit approaching the face and edges of the adjacent tetrahedral unit.

![Image](52x153 to 291x247)

**Figure 4.** Left: Li-centered partial distribution functions for 20 MPM Li–NH$_3$–MeNH$_2$. Right: Li-centered three-dimensional configurations for radial range $r=1.5–2.5$ Å, 80% probabilities: a) Li–N (NH$_3$), b) Li–N (MeNH$_2$), c) Li–N (MeNH$_2$), d) Li–N $r=2.5–6.0$ Å.

Importantly, the EPSR model also reveals the statistical distribution of ammonia and methylamine molecules in the [Li(NH$_3$)$_4$(MeNH$_2$)$_2$]$^+$ tetrahedral complexes (Supporting Information, Figure S2). This distribution shows that for around 40% of cations ($n,m$) is (2,2), for around 50% it is (1,3) or (3,1), and for the remaining 10% it is (0,4) or (4,0). The cation solvation is therefore significantly heterogeneous, and it is all the more surprising that we observe a sharp diffraction pre-peak in our data, indicative of longer-range order in the liquid. In the absence of a single cationic motif, we must conclude that the solvated electron itself acts as a structural template that drives homogenization. This hypothesis is consistent our observation of a single 7 Li Knight shift (Figure 1), and is truly remarkable given that the system is metallic.

The manner in which excess electrons are accommodated can be determined by examining the liquid structures for void regions: spherical volumes with circa 2.5 Å radii containing no atoms. Previous studies have demonstrated the existence of these regions in each of the metallic metal amines, with stark differences in their size and orientation, which correlate with their differing metallic properties. In the highly conducting Li–NH$_3$ liquid these voids are centered above the protons of the NH$_3$, forming channels between solvated Li(NH$_3$)$_4$ units (hence the moniker expanded metal). In the poor metal, Li–MeNH$_2$, the cavities are spatially isolated from one another, removed from the primary solvation environment of Li and bounded by the methyl groups of coordinated MeNH$_2$. The void–void distribution function in our mixed amine solution is presented in the Supporting Information, Figure S3, and shows weak inter-void correlations beyond around 4 Å.

Figure 5 shows the orientation of void regions relative to an NH$_3$ and MeNH$_2$ molecule in the 20 MPM Li–NH$_3$–MeNH$_2$ system. It is seen that the voids (and conduction electron density) lie around the circumference of NH$_3$ and above the amine group of MeNH$_2$. This structure is distinctly different to either the Li–NH$_3$ or Li–MeNH$_2$ system, and indicates that the electron density is drawn closer to Li, around the circumference of the solvating NH$_3$/MeNH$_2$ molecules.

In conclusion, we have discovered that the mixed amine system Li(NH$_3$)$_4$(MeNH$_2$)$_2$ forms a highly structured homogenous liquid metal. Each cation is tetrahedrally coordinated by approximately four solvent molecules, but there is significant heterogeneity in the NH$_3$:MeNH$_2$ ratio around
individual Li\(^{+}\) ions. This lack of a single cationic motif makes our observation of strong intermediate-range order in the solutions particularly surprising. Even though the system is metallic, we therefore turn to the solvated electron as a potential structural template. Indeed, we find that the system appears unique amongst current expanded metal–amine solutions for the manner in which the excess electron density is accommodated. This is drawn closer to the lithium cations than in either the constituent lithium–ammonia or lithium–methylamine solutions, and we observe only a single Knight shift on the lithium nuclei. Mixed-solvent metal–amine solutions therefore present a new set of challenges for our understanding of electron solvation, and the nature of the concomitant phase separation of the localized and delocalized electronic states.

**Experimental Section**

H/D and \(^{7}\)Li/\(^{6}\)Li isotopic substitution experiments (where nat denotes natural abundance) were performed using the SANDALS diffractometer at the ISIS Spallation Neutron and Muon Source, UK.[7] A clean sample of Li metal was loaded under argon into a sealed flat-plate null-coherent scattering Ti/Zr container, with sample and wall thicknesses of 1 mm. The container and sample were attached to a stainless steel gas-rig and evacuated to \(10^{-9}\) mbar. To prepare a sample of 20 MPM Li–NH\(_2\)–MeNH\(_2\), an equimolar ratio of NH\(_2\)-MeNH\(_2\) were premixed to the exact volume required, and cryo-pumped onto the Li sample at 60 K. This avoided any problem in the different boiling temperatures of the amines. The sample was then isolated and warmed to 240 K. The samples were monitored for Bragg reflections in the warming process to ensure that the solutions were fully homogenized. Data were collected for a period of 8 h, and spectra were corrected for background, multiple scattering and absorption following standard protocols for SANDALS data.

The measured total structure factor for a given system comprising \(n\) chemical species can be expressed as in Eq. (1):

\[
F(Q) = \sum_{i=1}^{n} \sum_{j=1}^{n} c_i c_j b_i b_j S_{ij}(Q) - 1
\]

where \(c_i\) denotes the atomic fraction of species \(i\) and \(b_i\) is the bound coherent neutron scattering length for that species. \(S_{ij}(Q)\) is the Faber–Ziman partial structure factor, and \(Q\) the scattering vector. The detailed method for extraction the partial structure factor from isotopically varying \(F(Q)\) values is described elsewhere. \(F(Q)\) is the Fourier transform of the total pair correlation function \(G(r)\). The specific isotopic substitutions for prepared samples are given in Table 1.

The empirical potential structure refinement (EPSR) method was used to model the measured neutron total scattering data.[8] A structural model comprising 2000 species (Li and molecular NH\(_2\)-MeNH\(_2\)) is simultaneously refined to each of the isotopically unique experimental \(F(Q)\) spectra (samples A–H in Table 1), enabling a rigorous constraint to the resultant structural model, and extraction of atom specific pair-correlation functions, \(G_{ij}(r)\). The inter-atom potentials in EPSR are comprised of a Lennard-Jones potential and Coulombic term, and is iterated throughout the simulation until a suitable convergence to experimental \(F(Q)\) spectra is reached. The seed potentials used in this work were the same as used previously for metal–amine liquid structure studies.[12–15]

NMR spectra were recorded on a Bruker AVIIIHD 400 MHz spectrometer, referenced to LiCl in D\(_2\)O. ESR was performed on a Bruker EMXmicro spectrometer. Samples for each were prepared in the manner outlined above, within spectrosil quartz tubes that were subsequently sealed. All handling and transfer of samples was conducted under cryogenic conditions.

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**Conflict of interest**

The authors declare no conflict of interest.

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