Optimum lithium loading of a liquid scintillator for neutron and neutrino detection

D. E. Bergeron*, H. P. Mumm*, M. A. Tyra, J. La Rosa, S. Nour
National Institute of Standards and Technology, Gaithersburg, Maryland 20899, USA
T. J. Langford
Wright Laboratory, Yale University, New Haven, CT 06511, USA

Abstract
Neutral particle detection in high-background environments is greatly aided by the ability to easily load $^6$Li into liquid scintillators. We describe a readily available and inexpensive liquid scintillation cocktail stably loaded with a Li mass fraction up to 1%. Compositions that give thermodynamically stable microemulsions (reverse-micellar systems) were explored, using a Compton spectrum quenching technique to distinguish these from unstable emulsions. Scintillation light yield and transmittance were characterized. Pulse shape discrimination (PSD) was measured using a $^{252}$Cf source, showing that electron-like and proton-like recoil events are well-resolved even for Li loading up to 1%, providing a means of background suppression in neutron/neutrino detectors. While samples in this work were prepared with $^{nat}$Li (7.59 % $^6$Li), the neutron capture peak was clearly visible in the PSD spectrum; this implies that while extremely high capture efficiency could be achieved with $^6$Li-enriched material, a very inexpensive neutron-sensitive detector can be prepared with $^{nat}$Li.

Keywords: Capture gating, pulse shape discrimination, microemulsion, micellar phase boundary, phase separation, light yield, inverse beta decay, Li-6

1. Introduction
A common problem in low count rate experiments is poor signal-to-background ratio. A widely used technique to isolate a signal of interest
in such a situation is to search for a unique coincidence between multiple components of the signal. Coincidence conditions can be imposed in the time domain, the space domain, or both. Capture gating, for instance, yields powerful background reduction in scintillation-based detectors [1, 2, 3, 4]. In such a scheme, separate gates are set for an initial particle interaction and a subsequent capture interaction. The capture can be identified by energy or, in an appropriate detector, via pulse shape discrimination (PSD) [5, 6, 7, 8]. Depending on circumstances, capture gating alone can yield background suppression of orders of magnitude. Capture gating with PSD is useful where the interaction of interest is easily confused with other signals. Particular examples include fast neutron spectroscopy and neutrino detection through Inverse Beta Decay (IBD) [9, 10, 11, 12]. In fast neutron spectrometry using typical hydrogenous scintillator-based methods, the spectrum is built by adding the energy deposition from multiple proton recoils. As the primary neutron thermalizes, recoils will decrease dramatically in energy and light output will be quenched. These low-energy events are easily confused with more frequent gamma interactions, and simple subtraction of backgrounds with the signal absent is not feasible in many applications (e.g., where the signal is always present or the production mechanism yields both neutrons and gammas). Neutrino detection through IBD suffers analogous problems; although the initial recoil positron carries the energy of the neutrino (which can be several MeV depending on application), unless the annihilation products can be isolated, neutrino events cannot easily be distinguished from Compton-scattered gammas.

In both of the previous examples, efficient detection of thermal neutrons enhances detector sensitivity. If the detector medium is doped with an appropriate isotope, neutron capture yields a unique signal. Capture times on the order of 10 µs are readily obtainable, nicely separating the capture from the prompt interaction while still effectively reducing uncorrelated backgrounds. Modern examples of both neutron spectrometers and neutrino detectors are driven to similar designs. Compact and segmented detectors are optimized for energy resolution and background rejection [10, 13, 14, 15]. In detectors with this design, neutron capture on $^6\text{Li}$, yielding short-range alpha and triton particles, provides the ideal, topologically compact, capture events that make discreet gating possible.
Liquid scintillators\(^1\) are particularly useful in radioactivity measurements, where a particle-emitting radionuclide can be added directly to the detection medium (‘internal’ detection). The short range of alpha and beta particles means that 4\(\pi\) geometry is complemented by high (100 % for alpha particles) detection efficiencies, achieving very high overall counting efficiencies.

Liquid scintillators also have properties that make them highly effective ‘external’ detectors. They are uniform without requiring the growth of a large single crystal or plastic bar, self-healing, generally less expensive than solid alternatives, efficient, and can be made to accommodate different neutron capture dopants [16, 17, 18].

Because \(^6\)Li is most typically available in a salt form (carbonate or chloride), approaches to suspending aqueous material in nonpolar organic liquid scintillators are necessary. By adding an appropriate non-ionic surfactant, or combination of surfactants, a thermodynamically-stable reverse-micellar solution, alternatively called a microemulsion, can be formed. In environmental analysis, nuclear power, and nuclear medicine settings, radionuclides are most commonly encountered as metal salts in aqueous solution. To accommodate these radioactive samples, commercially available liquid scintillators are variously optimized for aqueous loading capacity, high ionic strength, and/or PSD. Previous work on \(^6\)Li-loaded cocktails described good performance [8, 19, 20, 21], but the specific scintillators studied are no longer available. This drove the desire to find an inexpensive alternative and study its properties in the context of various expected end-uses. As a way to narrow the space of possibilities, we focused on liquid scintillators designed for radioactivity measurements with which we had previous experience.

Typically, these scintillators are engineered with various combinations of ionic and nonionic surfactants and alcohols to accommodate aqueous solutions encountered in common applications. The optimization of the specific proprietary formulations in the various scintillators leads them to exhibit significant variation in loading capacity and performance characteristics. In particular, certain formulations tolerate higher concentrations of ionic species which interact strongly with polar or ionic surfactant head groups. In order

\(^1\)The terms “scintillator”, “scintillant”, and “cocktail” are often used ambiguously in the literature. To avoid confusion, in this paper, we use “fluor” to refer to the fluorescent molecule; we use “scintillator” to refer to the fluor with solvent, wavelength shifters, and surfactants but no added aqueous material; and “cocktail” to refer to the scintillator with added aqueous material.
to find a formulation that provided optimal loading with good scintillation ef-

ciency, we surveyed multiple scintillators, loading them with LiCl solutions
of different concentrations. These included, HiSafe2, HiSafe3, HionicFluor,
Ultima Gold, and Ultima Gold AB (all PerkinElmer, Waltham, MA)\(^2\). This
initial survey, using the quench indicating parameter (QIP) as a measure of
stability, indicated that Ultima Gold AB (UGAB), a scintillator designed
specifically for optimal PSD, exhibited the highest LiCl loading capacity.
Building on this survey, we describe here studies suggesting that LiCl-loaded
UGAB (Li-UGAB) holds promise as a neutron-sensitive liquid scintillation
detector.

2. Sample preparation

The maximum aqueous loading capacity for a given scintillator is expected
to vary with ionic strength. In order to assure that a decrease in aqueous
loading capacity caused by increased ionic strength did not limit our overall
LiCl loading in the cocktail, we explored loading with a range of aqueous LiCl
concentrations. Thus, we confirmed that the highest concentrations of Li in
the cocktails were achieved with the highest concentration LiCl solutions.

Most samples were prepared from 8 mol/L aqueous LiCl solutions because
we were able to purchase a commercial solution (Sigma Aldrich, St. Louis,
Missouri) at this concentration. A 10 mol/L solution was also prepared from
LiCl powder (Section 2.1) to look at the possibility of using higher concen-
trations; while LiCl is soluble in cold water up to approximately 14 mol/L
(20 mol/kg)\(^2\),\(^3\) we adopted the conservative limit of 10 mol/L in our
studies in order to avoid any solution instabilities that might arise from
small impurities.

2.1. Preparation of 10 mol/L LiCl

The 10 mol/L solution was prepared from a LiCl salt\(^3\), which was as-
sumed to have a natural Li isotopic composition with Li atomic weight of

\(^2\)Certain commercial equipment, instruments, or materials are identified in this paper to
foster understanding. Such identification does not imply recommendation by the National
Institute of Standards and Technology, nor does it imply that the materials or equipment
identified are necessarily the best available for the purpose

\(^3\)Sigma product no. L9650-500G, lot no. BCBM2697V. Assay LiCl \(\geq 99\%\) (mole fraction) with the following impurities listed: \(\text{SO}_4^{2-} \leq 0.01\%, \text{Ba} \leq 0.003\%, \text{Fe} \leq 0.001\%\),
\(\text{K} \leq 0.01\%, \text{Na} \leq 0.20\%, \) heavy metals (as Pb) \(\leq 0.002\%.\)
6.941 g/mol. The 10 mol/L solution was prepared by slow addition of distilled water to a large beaker containing a pre-weighed amount of LiCl (see online supplemental material for details). The presence of insoluble material required multiple filtration steps, but ultimately yielded a clear solution with a measured density of 1.21 g/mL at room temperature. This sample was used for all of the measurements described herein.

Another 10 mol/L LiCl solution was prepared from a Li$_2$CO$_3$ solution with natural Li isotopic composition. This solution was prepared as an exercise to test and refine the procedures intended for use with isotopically enriched $^6$Li$_2$CO$_3$ for the PROSPECT (Precision Reactor Oscillation and SPECTrum) experiment [24, 25]. In this case, the solution was prepared by reacting the carbonate with concentrated HCl (see online supplemental material for details), resulting in a yellow solution. The solution was passed over an anion exchange resin, removing the yellow coloration (presumably caused by Fe(III) impurities). The clear column-purified 10 mol/L LiCl solution had a measured density of 1.21 g/mL at room temperature.

2.2. Preparation of Li-loaded cocktails

To achieve a range of Li and aqueous loading fractions ($f_{Li}$ and $f_{aq}$, reported as mass fractions), aqueous solutions of LiCl were prepared at several different concentrations and added to liquid scintillator.

All additions were performed volumetrically using dispensettes (PerkinElmer, Waltham, Massachusetts; quoted accuracy of 0.5 %) and micropipettes (Eppendorf, Westbury, New York; research, adjustable volume series; quoted accuracy 0.6 % to 3.0 %). Loading fractions, $f_{Li}$ and $f_{aq}$, are calculated from the volumes using measured densities. The uncertainties on the $f_{Li}$ are on the order of 3 % with the uncertainty on the LiCl concentration (as quoted by the manufacturer) in the stock solutions being the main contributor, followed by the uncertainties on the volumetric additions.

All small volume samples for quenching and spectroscopy measurements were prepared and stored in 20 mL borosilicate scintillation vials (PerkinElmer, Waltham, MA, USA). Samples were prepared to have the same total volume (scintillator plus aqueous material) in order to eliminate possible volume effects in the quenching measurements [26]. Larger volume samples for capture time and quantitative light output measurements were prepared using similar volumetric techniques, but in larger volume glass bottles (Wheaton, Millville, NJ, USA).
3. Quenching measurements

In many organic/surfactant systems the microemulsion phase is bordered on the phase diagram by a premicellar phase at lower aqueous fractions and an emulsion phase at higher aqueous fractions. Whereas microemulsions feature nanoscale aqueous reverse micelles and exhibit thermodynamic stability, emulsions feature larger aqueous domains that tend to agglomerate over time, ultimately resulting in separation into organic-rich and aqueous-rich phases [27]. We expect that the microemulsion phase will be optimal for high Li-loading and since phase boundaries can be sensitive to environmental conditions, we must assure that formulations intended for use in detectors are prepared with compositions reasonably far from phase boundaries. Thus, it is important to know where those boundaries lie.

To identify phase boundaries and assure a balance between stability and LiCl loading, we used Compton spectrum quenching, relying on QIP determinations [28], and optical transmittance and fluorescence spectroscopy to find discontinuities associated with increased scattering due to micelle formation. Further, we used a custom-built apparatus to measure light yield and PSD from Compton spectra.

3.1. Quench indicating parameters

QIPs were measured on either a Packard Tri-Carb A2500TR (PerkinElmer, Waltham, MA) or a Beckman Coulter LS6500 (Beckman Coulter, Fullerton, CA, USA). These counters are equipped with internal $\gamma$-radiation sources ($^{133}$Ba and $^{137}$Cs, respectively) which produce Compton electrons in the sample. The Compton spectrum is used to derive a QIP in terms of the “special index of the transformed external standard spectrum” (tSIE; decreases with increased quenching) for the Packard counter or the Horrocks number (H#; increases with increased quenching) for the Beckman. The tSIE corresponds to the energy bin that is intersected by the extrapolation of a line drawn between the points corresponding to 20% and 10% of the total counts in the Compton spectrum, while the H# is the inflection point at the Compton edge [29].

In the experiments described herein, samples were measured over multiple cycles, with the QIPs providing a measure of sample stability. We observed a few unstable cocktails undergoing phase separation on the timescale of the experiment (Fig. 1): cocktails prone to separation (emulsions) exhibited
decreased quenching over time as the more dense aqueous phase sank below the organic region being probed by the Compton source.

As Figure 1 illustrates, the highest concentrations of LiCl afforded the highest overall Li loading. Figure 1(a) and 1(b) show QIP results for series of Li-UGAB samples prepared with 1 mol/L and 8 mol/L LiCl solutions. In the 1 mol/L series, samples with $f_{Li} > 0.0017$ ($f_{aq} > 0.25$) show decreased quenching over the first few measurement cycles. The QIP for these samples seems to converge at approximately $tSIE = 425$, a value similar to that for the $f_{Li} = 0.0017$ sample. This may be taken as an indication of equilibration to a common saturation point where the organic phase can accommodate no more aqueous LiCl. In the 8 mol/L series, the sample with $f_{Li} = 0.026$ ($f_{aq} = 0.47$) exhibits decreased quenching after the first measurement cycle. The samples with $f_{Li} \leq 0.021$ ($f_{aq} \leq 0.37$) appear to be stable over time. The phase separated samples with $f_{Li} > 0.0017$ in the 1 mol/L series or $f_{Li} = 0.026$ in the 8 mol/L series exhibit the visual characteristics of a Winsor 2 type system, where a nearly pure aqueous phase is in equilibrium with a microemulsion phase [30]. In Figure 1 a, the sample with $f_{Li} = 0.0037$ does not approach equilibrium monotonically, but exhibits an oscillation. This behavior was observed in several samples and appears to be a spontaneously occurring example of the curious phenomenon referred to as oscillating phase separation [31, 32, 33].

Figure 1c shows the same data for the 1 mol/L and 8 mol/L LiCl series and also includes data for 2 mol/L and 4 mol/L series. In this panel, bars indicating a wide spread in the QIP values at a given value for $f_{Li}$ reflect sample instability over a number of measurement cycles (as reflected in panels a and b). So, it is clear that the samples in the 2 mol/L series with $f_{Li} \approx 0.003$ and 0.005 ($f_{aq} \approx 0.25$ and 0.36) are unstable, as is the sample in the 4 mol/L series with $f_{Li} \approx 0.007$ ($f_{aq} \approx 0.25$). Figure 1c illustrates clearly that the highest overall Li loading is achieved with the highest concentration of LiCl.

QIP (H#) data were obtained for samples prepared with 8 mol/L and 10 mol/L LiCl solutions (Figure 2a). The data for the 8 mol/L curves are taken from two separate experiments and overlap nicely. These measurements were performed with the Beckman counter instead of the Packard,

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4In this work, $f_{Li}$ is reported as a mass fraction assuming a natural isotopic abundance for Li. Care should be taken when comparing compositions for scintillation cocktails prepared with solutions enriched with $^6$Li since using an aqueous solution of LiCl with the same concentration by mole (e.g., 10 mol/L) would give a different $f_{Li}$ for the same $f_{aq}$. 

Figure 1: Quench indicating parameters (QIP) for different concentrations of \( ^{nat}\text{LiCl} \) loaded in Ultima Gold AB. a) QIP results for cocktails prepared with 1 mol/L LiCl solution over repeated measurement cycles spanning several hours. Results are shown for different Li loading fractions, \( f_{Li} \). b) QIP results for 8 mol/L LiCl solution. c) QIP results vs Li loading fraction, \( f_{Li} \), for different concentrations of LiCl solutions. The symbols represent points from the first measurement cycle, while the vertical bars illustrate the range covered in repeat measurements. The large bars indicate unstable cocktails undergoing phase separation.
Figure 2: Quench indicating parameter (H#) as a function of natLiCl loading in Ultima Gold AB. Series prepared with 8 mol/L and 10 mol/L LiCl were matched for Li content, as can be clearly seen in (a). Plotting against the aqueous fraction (b) shows that the higher LiCl concentration at the same aqueous loading results in slightly higher quenching. Uncertainty bars correspond to the standard deviation on repeat QIP measurements.
which was unavailable at the time of these experiments.

The samples prepared with the 8 mol/L solution are slightly more quenched than the 10 mol/L samples with the same $f_{Li}$ (Figure 2a). Figure 2b plots the same data as a function of the aqueous mass fraction, $f_{aq}$, showing that the higher concentration of LiCl results in slightly more quenching at the same overall $f_{aq}$.

Finally, the data in Figure 2 are consistent with earlier measurements made with UGAB that indicated a micellar phase boundary occurring at $f_{aq} \approx 0.03$ to 0.05 [34, 35, 28]. Using linear extrapolation intersection methods described previously [28], phase boundaries were identified at $f_{aq} = 0.048(2)$ for the 8 mol/L series and at $f_{aq} = 0.042(1)$ for the 10 mol/L series. The standard uncertainties are calculated from the combined fit uncertainties and the estimated uncertainty on sample $f_{aq}$. Uncertainty due to the curvature of the traces near the phase boundary (previously accounted via data assignment sensitivity (DAS)) is neglected here due to sparse sampling of the phase space near the discontinuities, so the stated uncertainties are underestimates. It is possible that higher ion concentrations promote more ordered solvent structure, pushing the phase boundary to lower $f_{aq}$ by reducing the entropic cost of micelle formation.

### 3.2. Phase stability

The QIP results described in Section 3.1 indicated that some formulations are prone to phase separation. Since some emulsions can initially appear indistinguishable from microemulsions, we conducted additional experiments to probe the susceptibility of our samples to phase separation. As described in detail in a previous publication [36], we followed two matched sets of samples (prepared with 8 mol/L LiCl) over a period of 3 weeks, subjecting one set to centrifugation twice. QIPs were measured periodically to monitor for phase separation. In addition to the samples undergoing spontaneous phase separation (see Section 3.1 and Figure 1), the sample with $f_{Li} = 0.017$ ($f_{aq} = 0.31$) separated visibly upon centrifugation. Samples with $f_{Li} \leq 0.011$ ($f_{aq} \leq 0.21$) showed no phase separation and are considered stable. Stable samples stored for more than 9 months at room temperature show no visible signs of deterioration or phase separation. Additionally, samples that were sparged with inert gas and stored for more than two years showed no signs of deterioration or phase separation.
Figure 3: $^{137}$Cs Compton spectra of UGAB samples loaded with 8 mol/L $^{nat}$LiCl compared to EJ-309 which has a known light yield in terms of photons per electron equivalent pulse energy (MeVee).

Figure 4: Absolute light yield in terms of photons per MeV as a function of $^6$Li concentration. The values and error bars are determined through a $\chi^2$ fit of spectra normalized to the EJ-309 expected spectrum.
3.3. Light Yield

Light yield was determined using a small purpose-built setup with a 7.62 cm Hamamatsu R6091 Photomultiplier Tube (PMT) coupled to an acrylic vial holder with RTV-615 potting compound. The holder is a 7.62 cm right cylinder with a 2.5 cm diameter borehole along its axis to receive the vial. Vials are placed in the borehole without any additional coupling compound to ensure consistent optical properties between samples. The acrylic holder is wrapped on all surfaces with polytetrafluoroethylene (PTFE) reflectors to increase light collection efficiency. The PMT+acrylic holder is mounted upright inside a thin-walled aluminum tube with light-tight KF flanges. The system allows a convenient way to quickly swap samples contained in standard 20 mL scintillation vials in a reproducible configuration. Samples were prepared gravimetrically (15.0(1) g each) to assure consistency.

Prior to measurement, each sample was stored in a reduced-oxygen environment to reduce the effects of oxygen quenching, which has been shown to be problematic in Li-loaded scintillators in the past [24]. For the measurements reported here, Li-UGAB samples were gravimetrically prepared with 8 mol/L \(^{nat}\)LiCl with \(f_{\text{Li}} = 0, 0.001, 0.004, \) and \(0.010 (f_{\text{aq}} = 0, 0.02, 0.08, \) and 0.21).

The PMT was powered to -1700 V, which was shown to provide an acceptable dynamic range. To ensure a stable PMT performance for each measurement, the system was allowed to warm up for approximately 5 min after initial biasing. Each sample was irradiated with a \(^{137}\)Cs source placed directly above the vial on the top KF flange. The same system was used for the PSD measurements (Section 5) by placing the \(^{252}\)Cf source to the side of the vial behind a 5 cm lead shield to reduce the gamma flux. Digitized PMT signals are integrated to determine the deposited energy in each scattering event. The spectra of the samples’ response to a \(^{137}\)Cs gamma source is shown in Figure 3. The relative light yield of each sample was determined by a \(\chi^2\) fit of the normalized energy spectrum with a multiplicative scale factor free parameter. Light yields of the four samples were determined (see Figure 4) from the known light yield of the reference sample (11 500 ph/MeV) [24]. Total combined uncertainties in light yields are estimated to be 2 %. The data for the sample with \(f_{\text{Li}} = 0\) is consistent with an earlier measurement reported in [37]. These data can be combined with the QIP measurements discussed in Section 3.1 to provide a normalization and allow the interpretation of light yield as a function of loading. The light yield of the Li-UGAB sample with \(f_{\text{Li}} = 0.001\) is approximately 20 % lower than the similarly loaded cocktail used in the PROSPECT experiment [37]. However, Li-UGAB is simpler to
prepare, due to commercial availability of both LiCl and UGAB.

4. Spectroscopic measurements

Transmittance ($T$) measurements were performed on an Hitachi U3900 UV-Vis spectrophotometer (Hitachi, Northridge, CA, USA). Samples were measured in quartz cuvettes with an air blank. The air blank is not ideal since reflection losses at the interfaces will be different for the reference and sample cells; in fact, we often saw $T$ in excess of 100% because the difference in refractive indices between the diisopropyl naphthalene (DIN)-based cocktails and quartz is smaller than that than between air and quartz. For the purposes of comparing different cocktail compositions, however, our experimental approach was satisfactory. The sample cuvette was washed thoroughly with methyl alcohol between measurements. The sample and blank cuvettes were not exchanged so that variability from cuvette placement was minimized (i.e., only the sample cuvette was removed between each measurement). Scans were taken over the range of 350 nm to 600 nm with 1 nm resolution at a scan rate of 10 nm/s. Three repeat measurements were performed for each sample with relative standard deviations on the transmittance values $< 0.2\%$.

Fluorescence measurements were performed with an Hitachi F7000 fluorescence spectrophotometer (Hitachi, Northridge, CA, USA). Samples were again measured in quartz cuvettes which were cleaned thoroughly with methanol between each measurement. Excitation and emission spectra were collected for each sample.

Excitation spectra were collected over a range of excitation wavelengths, $\lambda_{EX} = 250$ nm to 450 nm with an open fluorescence detection window. The excitation slit width was 1 nm and the scan rate was 4 nm/s. The excitation peak appeared to blueshift very slightly with increased Li loading, but the shift was within the instrument resolution ($< 1$ nm).

Emission spectra were collected with $\lambda_{EX} = 407$ nm, consistent with the peak in the excitation spectrum. The detected fluorescence wavelength was measured over a range of $\lambda_{EM} = 350$ nm to 500 nm with 1 nm slit widths for both excitation and emission and a scan rate of 4 nm/s. The maximum transmittance for Li-UGAB is reached at approximately 440 nm (Figure 5); below 400 nm, the transmittance is near zero (i.e. there is almost 100% absorbance). For our purposes, the most important region for transmittance is where the scintillator fluoresces. As Figure 5 illustrates, the peak in the
fluorescence emission spectrum occurs at 425 nm; at this wavelength, all measured transmittances were $> 94.5\%$. Transmittance values at 500 nm, on the spectral plateau, were also always $> 94.5\%$.

For microemulsions, the fluorescence emission spectra included significant contributions from scattering of the 407 nm excitation beam. This was confirmed by varying the excitation wavelength and observing the corresponding change in the position of the sharp feature in the fluorescence emission spectrum. To facilitate analysis, we attempted a deconvolution of the fluorescence and scattering contributions. Figure 6 shows the fluorescence emission spectrum for a UGAB sample with $f_{Li} = 0.014$ ($f_{aq} = 0.21$). The major spectral features were fit to Gaussians using a least squares approach in order to deconvolute scattering and true fluorescence contributions. Fit residuals consistently showed that the largest mismatch occurred around the scattering peak. Since the scattering peak should have a wavelength defined by the excitation slit width, a departure from Gaussian peak shape is not surprising. The amplitudes from the fits provided a much better approximation of scattering contributions than integrating under the scatter peak and so they were used in subsequent analyses (Figure 7).

Figure 7 shows the dependence of transmittance, fluorescence, and scattering on $f_{aq}$. Figure 7a shows the measured transmittance for samples prepared with 8 mol/L and 10 mol/L LiCl solutions. The curves show two distinct regions. At low $f_{aq}$, transmittance increases with increasing $f_{aq}$. Near the phase boundary discussed above, there is a transition to a region
Figure 6: Fluorescence emission spectrum for a UGAB sample with $f_L = 0.014$ ($f_{aq} = 0.21$). The presence of reverse micelles in the samples leads to significant scattering of the excitation light ($\lambda_{EX} = 407$ nm). A least squares fit to the measurement data was used to deconvolute scattering and fluorescence contributions.

where transmittance decreases with increasing $f_{aq}$.

For a sample containing a mixture of multiple absorbing substances, Beer’s law can be written in terms of transmittance ($T$) as

$$T = e^{-\sum_i (\epsilon_{a,i} + \epsilon_{s,i}) b c_i}, \quad (1)$$

where $\epsilon_{a,i}$ and $\epsilon_{s,i}$ are, respectively, the absorption and scattering extinction coefficients for the $i^{th}$ component of the cocktail. The path length, $b$, comes from the 1 cm cuvette and is constant in all of our experiments. The concentration of the $i^{th}$ element, $c_i$, is in practice a function of the aqueous fraction of the sample, $f_{aq}$. As $f_{aq}$ increases, the concentration of the organic components of the scintillator are diluted in the cocktail. At the same time, as $f_{aq}$ increases, the concentration of micelles and premicellar aggregates increases.

The two regions in Figure 7 a, then, can be explained simply as the dilution region, where the fluor and other organic absorbers (with relatively large $\epsilon_{a,i}$ components) are diluted, and the scattering region, where the organization of the surfactants about the added aqueous material leads to the formation of large micelles (with relatively large $\epsilon_{s,i}$ components).

Of course, this simplified picture does not suffice to quantitatively explain the data. Interactions between the components are neglected. These are important since the relative concentrations of absorbers and scatterers may at times draw from the same molecular reservoir. In addition, interactions
Figure 7: (a) Transmittance at 425 nm for a series of cocktails prepared with 8 mol/L LiCl (blue triangles) and 10 mol/L LiCl (red diamonds). Uncertainty bars show the standard deviation on three repeat measurements. (b) Total fluorescence intensity for the 8 mol/L (blue circles) and 10 mol/L (red diamonds) series and scatter intensity for the 8 mol/L (teal triangles) and 10 mol/L (yellow squares) series. Lines between points are intended only to guide the eye. Uncertainty bars represent the uncertainty of the least squares fit used to determine peak intensities (see Figure 6 and text). The left axis applies to scatter intensities while the right axis applies to fluorescence intensities. Note that samples were prepared with the same $f_{Li}$ in each series, so the last point in the 8 mol/L series (with higher $f_{aq}$) corresponds to the same $f_{Li}$ as the last point in the 10 mol/L series.
between the components may contribute to shifts in the absorption maxima of the individual components through, e.g., exciplex formation near micellar interfaces. Finally, we have neglected terms for sample fluorescence. Still, Equation 1 reproduces the basic shape of the $T$ v. $f_{aq}$ curves observed in our experiments, demonstrating that dilution and scattering are the predominant (and competitive) mechanisms at play here.

The curvature of the transmittance data is due to the competition between dilution and scattering over the region between $f_{aq} \approx 0.02$ to 0.06. Because of the curvature, we have not attempted the type of analysis of intersects we performed for the QIP data. The transmittance data certainly indicate a phase boundary and in a $f_{aq}$ region consistent with the other determinations. At the highest $f_{aq}$, there is slight divergence of the 8 mol/L and 10 mol/L LiCl series, with the higher LiCl concentration samples exhibiting slightly higher transmittance.

Both fluorescence emission and scattering increase with $f_{aq}$ and both show some curvature or slope discontinuity near the phase boundary at $f_{aq} \approx 0.04$ (Figure 7 b). Below this threshold, there is effectively no scattering. Increased scattering above this threshold is consistent with the formation of strongly scattering reverse micelles. The absolute scatter intensities for the highest concentration samples in the 8 mol/L series are twice those with the same $f_{Li}$ in the 10 mol/L series. The formulations with higher LiCl concentration generally appear to scatter less 407 nm light. This is consistent with the series divergence noted in the transmittance data.

The emission intensity is a measure of the fluorescence output for excitation at 407 nm (Figure 7 b, right axis). Fluorescence emission intensity appears to increase very slightly with increased loading up to about $f_{aq} = 0.1$. This effect is much smaller in magnitude than the effect on scattering shown in Figure 7 b. The magnitude of change in fluorescence emission intensity is similar to the magnitude of change in transmittance (Figure 7 a). The increase in the observed fluorescence intensity may be attributed to increased transmittance of the fluorescence light. For $f_{aq} > 0.1$, increased scattering could be expected to produce more fluorescence light due to increased path length, but it appears that this effect is offset by reduced transmittance. Finally, the data are consistent with better transmittance of fluorescence light—presumably due to reduced scattering—for the 10 mol/L LiCl series.

The uncertainty bars in Figure 7 b represent only the uncertainty on the Gaussian fits used to estimate peak intensities. For the total fluorescence emission intensity, fit uncertainties were estimated using a Cholsky decom-
Figure 8: PSD \( (Q_{\text{tail}}/Q_{\text{tot}}) \) as function of pulse area for UGAB loaded to \( f_{\text{Li}} = 0.01 \) nat\( \text{Li} \) by mass. Excellent separation of the proton-like (upper) and electron-like (lower) recoil bands as well as a well defined capture peak at approximately 500 keVee.

position method to account for correlations in the Gaussian terms. For the fits of the scatter peaks, uncertainties of several percent appear small due to large changes over the series. Further, we expect that the total uncertainty on these intensities will be on the order of several percent and dominated by measurement systematics, especially due to sample preparation.

UGAB samples prepared with water (not aqueous LiCl) to achieve matched \( f_{\text{aq}} \) give similar results. Fluorescence quenching by LiCl is slight but observable, consistent with QIP results (see [36]). Despite this fluorescence quenching by LiCl, we find that at a given \( f_{\text{aq}} \), the more concentrated LiCl solution produces a cocktail with better transmittance and less scattering.

5. Pulse Shape Discrimination

A \(^{252}\text{Cf}\) source was used to characterize the Pulse Shape Discrimination (PSD) performance of the Li-UGAB with various \( f_{\text{Li}} \). The source, with an approximate activity of \( \sim 10^4 \) Bq, was positioned approximately 20 cm from the measurement system described above (section 3.3). For each digitized waveform recorded from the PMT a \( Q_{\text{full}} \) was defined by integration over a window from 12 ns before to 120 ns after the half-height of the waveform’s leading edge and a \( Q_{\text{tail}} \) as the charge integrated 40 ns to 120 ns after the leading edge half-height. A conventional tail-fraction PSD metric was then defined by \( Q_{\text{tail}}/Q_{\text{full}} \). Figure 8 shows this PSD metric plotted against electron equivalent pulse energy (MeVee). Two distinct bands are evident, the upper the result of proton-recoils from neutron scatters and the lower
Figure 9: PSD \((Q_{\text{tail}}/Q_{\text{tot}})\) in the approximate region of the neutron capture peak compared between various loading levels.

Figure 10: FOM (peak separation divided by the sum of the peak FWHM) in the approximate region of the neutron capture peak as a function of loading. As expected the trend closely follows the absolute light yield shown in Figure 4. The values and error bars are determined through a double-Gaussian fit of the distributions.
the result of electromagnetic interactions. Even though natural lithium was used for these studies, the peak from neutron capture on $^6\text{Li}$ (7.5 % natural abundance) is evident. Figure 9 shows PSD in the approximate region of the neutron capture peak compared between various $f_{\text{Li}}$. By fitting each peak to a Gaussian distribution and extracting the peak positions and the Full Width Half Max (FWHM) a PSD figure of merit (FOM) can be defined as peak separation divided by the sum of the peak widths. PSD performance remains good at all loading, but does fall off as expected with decreasing light yield as shown in Figure 10.

The effect of quenching on PSD has been discussed by Pates et al. [38] and it appears that quenching induced by the addition of aqueous material and the formation of reverse micelles is impacting the PSDs measured here. Quenching shortens both the prompt and delayed components of scintillation pulses, but the delayed component is typically quenched more. The initial decrease in PSD going from $f_{\text{Li}} = 0$ to 0.001 (Figure 9) arises from the overall shortening of anode pulses with increased quenching. Figure 10 shows how increasing quenching preferentially impacts the proton-like pulses, reducing the FOM. Still, the separation between electronic-like and proton-like recoil events remains good (Figures 9 & 10). Quenching by micelles and effects of scattering on PSD are worthy of further study.

6. Conclusion

We have explored optimal loading of a liquid scintillator. We found that Ultima Gold AB (UGAB) accommodates up to 1.0 % nat Li in a microemulsion phase while preserving good pulse shape discrimination characteristics. Consistent with previous work with UGAB [36], we found that the presence of LiCl has a minimal impact on the quenching expected for a given aqueous fraction. We achieved the highest Li-loading fractions with the highest concentration LiCl solutions, but did not explore concentrations higher than 10 mol/L. Optical spectrophotometry measurements hint that incremental gains in light yield due to improved transmittance might be made with higher concentrations.

The presence of LiCl does affect micellar dynamics in UGAB. The phase boundary separating the premicellar regime from the reverse micellar (microemulsion) phase appears to occur at higher $f_{\text{aq}}$ with increasing LiCl concentration. For detector applications, compositions in the reverse micellar phase provide stable loading with high $f_{\text{Li}}$. Compositions near the phase
boundary are to be avoided since large changes in the optical properties (scattering, transmittance) could result from small changes in temperature or pressure, or from ionization events that would provide nucleation centers for metastable surfactant-solute clusters. For applications where lower $f_{Li}$ is acceptable, compositions below the micellar phase boundary may be worthy of further investigation, promising reduced scattering and thus substantially improved optical transmittance.

Pulse shape discrimination in the $^{nat}Li$-loaded UGAB cocktails showed clear signs of the neutron capture peak on $^6Li$. While $^{nat}Li$ is only 7.59 % $^6Li$ [39], the extremely high Li-loading we have achieved provides enough of the neutron-sensitive isotope for good capture gating. On one hand, this implies that a very inexpensive neutron-sensitive cocktail can be prepared. On the other hand, with an investment in $^6Li$-enriched material, a cocktail with extremely high capture-efficiency could be prepared.

The formulation described herein is easily prepared, exhibits excellent stability, has a higher $f_{Li}$ than alternatives, and preserves good PSD and optical properties. Studies are needed to further establish its promise for neutron and neutrino detection.

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8. Online supplemental material

8.1. Preparation of 10 mol/L LiCl from LiCl salt

On a top-loading balance (Mettler-Toledo PB3002), (212.19 ± 0.03) g of LiCl powder (equivalent to 5.0 mol of LiCl) were weighed out directly from the bottle into a tall-form, 1000 mL borosilicate glass beaker. This mass of powder had a volume of about 150 mL. The slow addition of 250 mL of distilled water (DW) was accompanied by considerable heat generation due to the exothermic hydration reaction. The goal was to dissolve all of the salt while carefully bringing the volume of the solution to 500 mL in order to
produce a nominally 10 mol/L LiCl solution. The salt formed a large, solid cake which dissolved slowly. The application of additional heat by means of a hotplate with occasional stirring dissolved the LiCl within 40 min, producing a volume of about 375 mL of hot solution. Further DW addition of 80 mL brought the volume to ~ 460 mL. The resulting solution, which was visibly turbid, was allowed to cool to room temperature over a period of 3 h. No apparent crystallization of LiCl occurred upon cooling. The solution was filtered under maximum suction but passed very slowly through a 47 mm, 0.45 µm porosity membrane filter (Pall GN-6 Metricel, product no. 63069). The filtrate was transferred to a 500 mL glass volumetric flask, diluted to the mark and mixed. However, this filtered 10 mol/L LiCl solution soon appeared cloudy. It was eventually decided to allow the 10 mol/L LiCl solution to stand for several days to permit complete precipitation of any insoluble matter. It has been observed that such behavior may occur when very concentrated solutions are prepared, presumably because the high ionic strength can affect the precipitation rate of even very insoluble compounds5. After 5 d of standing, the suspension was again filtered under suction through a GN-6 (0.45 µm porosity) membrane filter; the filtration rate was now relatively fast and the filtrate appeared to be clear and colorless. The 10 mol/L LiCl solution, which had a pH between 3 and 4 (measured by wide range indicator paper) and a density of 1.205 g/mL (±1 %, k=1) at room temperature, was placed in a clear borosilicate glass bottle for storage. No further turbidity was observed.

A suggested procedure for future preparation of 10 mol/L LiCl solutions from the salt would be to add the salt (212 g) in small portions to the water (~330 mL to 350 mL), dissolving the added material before the next addition. Then, the solution should be allowed to stand ~ 1 week to permit precipitation of insoluble matter before filtering to give the final solution.

5One potential component of the insoluble precipitate is BaSO4. If the Ba impurity was equal to 0.003% by weight, then 212 g of LiCl could produce as much as 0.010 g BaSO4. This would exceed the solubility in a water solution of 500 mL (0.00123 g BaSO4). In the present situation, the 500 mL of 10 mol/L LiCl certainly contains less than 500 mL of water, which may further reduce the BaSO4 solubility, although the high ionic strength of the solution may counteract this effect.
We prepared 100 mL of a 10 mol/L LiCl solution starting with Li$_2$CO$_3$ of presumably natural Li isotopic composition as an exercise to test and refine the procedures intended for use with isotopically enriched $^6$Li$_2$CO$_3$ for the PROSPECT experiment [25]. Almost exactly 37 g of lithium carbonate (from an old Baker and Adamson reagent-grade product$^6$), approximately equal to 1.0 mol of Li, were weighed (directly from bottle) into a 250 mL glass beaker. A stoichiometric amount$^7$ of concentrated HCl (37.5 % by mass), 85 mL, was introduced into a 500 mL Erlenmeyer flask. The Li$_2$CO$_3$ powder was added in small amounts using a plastic spoon to the Erlenmeyer flask over the course of 25 min with occasional swirling. The effervescence was easily under control; i.e., the spray and foaming were virtually completely contained within the flask, and the heating was moderated so that the solution only became warm. The residue of Li$_2$CO$_3$ in the 250 mL beaker was dissolved in a small amount of water containing a few drops of concentrated HCl and combined with the main liquid. However, the main solution in the Erlenmeyer flask was not clear. Concentrated HCl was added drop-wise with swirling until the effervescence ceased (approximately 4 mL). At this point, the solution was clear and distinctly light yellow-colored but with no visible insoluble matter. It was heated, boiled very briefly and allowed to cool over several hours to room temperature. The pH was 2 to 4 (using wide range indicator test paper) with a solution volume < 100 mL. The yellow color did not disappear upon cooling; it was suspected that the color derived from iron(III) impurity in the original salt since the previous solution prepared from LiCl salt was colorless. This yellow color was undesirable from the standpoint of producing color-quenching when mixed with liquid scintillator. Assuming it was due to Fe(III), the iron would most likely be present as the anionic chloro-complex FeCl$_4^-$ in this concentrated LiCl solution. As such it might be removed by using a strongly basic anion exchange resin such as AG 1 (quaternary amine functional group) just as Fe(III) is tightly retained on this resin from $\geq$ 6 mol/L HCl solution. Subsequently it was verified that when this LiCl solution was passed slowly through a 2 mL bed volume

$^6$Baker and Adamson (General Chemical Co.) reagent no. 1202, code 1872. Impurities (maximum, in %): Cl 0.005, SO$_4$ 0.25, Al$_2$O$_3$ 0.03, CaO 0.10, Fe 0.004, heavy metals (as Pb) 0.002.

$^7$Reaction: Li$_2$CO$_3$ + 2HCl = 2LiCl + H$_2$O + CO$_2$(g); the carbon dioxide gas is evolved from the solution.
column of AG 1-X4, 100 to 200 mesh resin in chloride form, the yellow color was absent from the column eluate. If needed, the small amount of LiCl remaining in the resin after passing the solution could be washed out with 9 mol/L HCl, leaving the Fe(III) in the resin. This 9 mol/L HCl – LiCl eluate could be evaporated, treated with 30 % H$_2$O$_2$ to destroy organic traces (from the resin), re-dissolved, and combined with the main body of purified LiCl solution$^8$. The column-purified LiCl solution (clear, colorless) was made up to 100 mL in a volumetric flask to give 10 mol/L LiCl with a measured density of 1.21 g/mL at room temperature.$^8$

The final preparation of 10 mol/L $^6$LiCl from $^6$Li$_2$CO$_3$ salt was carried out with the same technique. Out of 100 g of enriched-$^6$Li lithium carbonate (95 % enrichment, origin Oak Ridge National Laboratory), 36 g were taken to prepare 100 mL of 10 mol/L $^6$LiCl solution. The dissolution procedure followed closely that used for the natural Li$_2$CO$_3$, including a 500 mL Erlenmeyer flask. The resulting solution in HCl was also yellow-colored and contained visible, undissolved black particles. Based on our experience with the natural-isotopic Li$_2$CO$_3$, a 2 mL bed volume column of AG 1-X4, 100 to 200 mesh, chloride form anion exchange resin was prepared. The $^6$LiCl solution was first filtered through fast filter paper (Whatman no. 41) by gravity to remove the black particles before passing through the AG 1 column. The column successfully eliminated the yellow color from the eluate. $^6$LiCl remaining on the column was stripped with 9 mol/L HCl, evaporated to dryness and treated with 30 % H$_2$O$_2$ to destroy organic matter from the AG 1 resin$^9$. The $^6$LiCl recovered was about 1.5 g (out of 41.5 g $^6$LiCl expected). All of the $^6$LiCl was combined into a single 100 mL solution (nominally 10 mol/L $^6$LiCl) with a measured density of 1.20 g/mL at room temperature.

All $^6$LiCl solutions were stored in polyethylene bottles to allay concerns of

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$^8$Passage of 5 mL of 0.1 mol/L HCl into the column after the 9 mol/L HCl immediately eluted the yellow substance, which gave a positive (cherry-red) spot test with 5 mol/L NH$_4$SCN solution, a positive indicator for Fe(III). The solution was partially decolorized by the addition of HF or KF, converting the colored FeSCN$^{2+}$ to colorless FeF$_3^{-}$. While the residual yellow-colored impurity is not conclusively identified, it is removed by anion exchange chromatography.

$^9$Caution! 30 % H$_2$O$_2$ added to LiCl produces a very exothermic reaction resulting in immediate foaming/effervescence; therefore, cold H$_2$O$_2$ must be added slowly with provisions for intercepting the spray generated by the gas liberation.
possible boron (a strong neutron absorber) leaching from borosilicate glass.

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