Large-Scale, Precision Xenon Doping of Liquid Argon

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Abstract

The detection of scintillation light from liquid argon is an experimental technique key to a number of current and future nuclear/particle physics experiments, such as neutrino physics, neutrinoless double beta decay and dark matter searches. Although the idea of adding small quantities of xenon (doping) to enhance the light yield has attracted considerable interest, this technique has never been demonstrated at the necessary scale or precision. Here we report on xenon doping in a 100 l cryogenic vessel. Xenon doping was performed in four concentrations of 1.00\(\pm\)0.06 ppm, 2.0\(\pm\)0.1 ppm, 5.0\(\pm\)0.3 ppm, and 10.0\(\pm\)0.5 ppm. These measurements represent the most precise xenon doping measurements as of publishing. We observed an increase in average light yield by a factor of 1.92\(\pm\)0.12(syst)\(\pm\)0.02(stat) at a dopant concentration of 10 ppm.

Keywords: neutrinoless double beta decay, xenon doping, liquid argon, Birk’s constant, Geant4

1. Introduction

Liquid noble-gas detectors (LNGD) are widely used or are planned to be used in many different particle physics applications, including dark matter (DARWIN \cite{11}), neutrino tracking (DUNE \cite{13}), and neutrinoless double beta decay (0\(\nu\beta\beta\)) (GERDA \cite{7,8}, LEGEND \cite{2}). Although He and Ne have been considered \cite{13}, Ar and Xe monolithic LNGDs have advantages due to the large scintillation yields, longer scintillation wavelengths, and fast scintillation time constants. For the liquid Ar (LAr) case, the boiling temperature is
near the preferred operating temperature of Ge detectors and Ar can provide substantial radiation shielding. These features led to the GERDA concept of operating Ge detectors bare within a LAr environment [7]. If the light detection of the LAr in GERDA could be significantly enhanced, it would result in improved background rejection. Doping the LAr with Xe is one technique to improve the light detection and is being considered for LEGEND-1000. However, a Xe-doped LAr (XeDLAr) volume of sufficient size to demonstrate its use for LEGEND has not yet been demonstrated. It is important to demonstrate that the doping can be done at a large scale in a controlled manner, and to determine if XeDLAr mixture is stable.

1.1. Xenon Doped Argon

Doping LAr with 0.1-1000 ppm of Xe as shown by [24] (10-1000ppm Xe as shown by [10, 27, 32, 26, 22]) shifts the wavelength of emitted scintillation light from 128 nm, where LAr scintillates, to 175 nm where Xe scintillates. This wavelength shift is advantageous for several reasons. Argon has a long attenuation length at 175 nm [26] but only a 50-60 cm attenuation length at 128 nm [25, 22]. Additionally, devices sensitive to detect both 128 nm and 175 nm are more sensitive at 175 nm. In order to detect the 128 nm light with conventional photo-multipliers (PMT), the light must first be wavelength shifted. The coupling of light detection devices to wavelength shifting materials (e.g. TPB [16]) typically occurs at the surface of the device, and therefore nearly half the light does not directly enter the PMT because the secondary re-emission process is isotropic. In contrast, the XeDLAr wavelength shifting occurs approximately at the interaction point.

Obviously there are many advantages to a XeDLAr LNGD but in order to accurately quantify them, XeDLAr needs to be demonstrated on a large volume. The largest active volume previously used was 13.6 l [27], the other active volumes used in previous works were 31 mL [10], 56 mL [25], 1.6 l [22], and 3.14 l [32]. One concern is the possibility of clumping: reference [22] (1.6 l active volume) claims to measure clumping at Xe concentrations of 3%, while a chemical analysis shows that Xe is 16% soluble in LAr at 87 K [12]. Another concern is the stability of the mixture, since Xe has a much higher freezing point than Ar (161.40 K versus 83.81 K).

Xenon doping was performed in previous smaller experiments using one of two methods. Either a room temperature argon-xenon mixture was injected into the liquid volume, or the mixture was prepared in the gaseous phase and then condensed into liquid. The warm gas injection technique will
not work on large scale experiments because xenon homogeneity will take a long time to be achieved, while a pre-mixed gas is not feasible due to the large gas storage volumes required. Additionally, these doping techniques result in large total-concentration uncertainties making reported scintillation time constants difficult to compare. The smallest doping concentration uncertainty reported to date is 23.5% \cite{27}. New injection techniques need to be considered if xenon doping is to be pursued on a large scale.

A large Xe doping test stand is also critical for understanding the effects of the long attenuation length at 175 nm in XeDLAr. This long attenuation length can result in an increase in light yield as observed previously by Ref. \cite{27} (×1.4 increase at 300±80 ppm) and Ref. \cite{32} (×1.25 increase at 1093±547 ppm). The discrepancies in these increased light yield measurements can be attributed to different detectors sizes, with the larger increase being measured in the larger detector.

2. Argon and Xenon Scintillation

When ionizing radiation interacts with a noble gas, dimers form which, when relaxing back to the ground state, produce scintillation light. Scintillation light is emitted through the transition from one of the two lowest molecular excited states, $^3\Sigma_u^+$ (triplet) or $^1\Sigma_u^+$ (singlet), to the ground state $^1\Sigma_g^+$. These states cannot be easily distinguished spectroscopically but do have different relaxation times. The $^1\Sigma_u^+$ state for liquid argon has a half life of around 5 ns while the $^3\Sigma_u^+$ state half lives for pure liquid argon have been measured to be between 1.2-1.6 $\mu$s \cite{19,23,5}. This large difference in measured $^3\Sigma_u^+$ state lifetimes for “pure” argon is likely due to small concentrations (~1 ppm) of residual atmospheric xenon\footnote{Atmospheric argon concentrations are 0.934\% by volume in Earth’s atmosphere, while atmospheric xenon concentrations are 87±1 ppb \cite{20} by volume. If the argon distillation process does not remove any xenon, as much as 9.3 ppm of xenon can remain.}. The $^1\Sigma_u^+$ state for liquid xenon has a half life of 2-4 ns and the $^3\Sigma_u^+$ state 21-28 ns \cite{9}. The main feature of the argon and xenon scintillation spectra is the Gaussian feature centered at 128 nm and 175 nm for argon and xenon respectively. The summed probability distribution function for the $^1\Sigma_u^+$ and $^3\Sigma_u^+$ components can be written as:

$$I(t) = \frac{A_1}{\tau_1} e^{-t/\tau_1} + \frac{A_3}{\tau_3} e^{-t/\tau_3}$$ (1)
where $\tau_1$ and $\tau_3$ are the time constants for the $^1\Sigma_u^+$ and $^3\Sigma_u^+$ state respectively and $A_1$ and $A_3$ are the relative intensities of each components. The integral of $I(t)$ over all time is normalized to unity and therefore $A_1 + A_3 = 1$.

2.1. Xenon Doping

When Xe is inserted into LAr, it quenches the Ar dimers through collisional de-excitation. Unlike other contaminants (e.g. N), this collision allows for effective energy transfer between the Ar dimer and Xe, allowing the Xe atom to create a dimer of its own, as:

\[
\ce{Ar_2^* + Xe + collision \rightarrow (ArXe)^* + Ar} \tag{2a}
\]
\[
\ce{(ArXe)^* + Xe + collision \rightarrow Xe_2^* + Ar} \tag{2b}
\]

Here collision refers to a molecular collision between a Xe atom and a Ar$_2^*$ or ArXe$^*$ dimer. Based on estimates of the diffusion rate from [15], this WLS process occurs no more than 500 $\mu$m from the initial argon interaction for Xe concentrations considered in this work.

Xe and Ar have very different scintillation time profiles with the biggest difference being their $^3\Sigma_u^+$ state lifetimes ($\sim 1$ $\mu$s versus $\sim 30$ ns). When Xe is injected into Ar at low concentrations, the $^3\Sigma_u^+$ state of the Ar is quenched by collisions, thus the time distribution can be described [10]:

\[
I(t) = \frac{A_f}{\tau_f} e^{-t/\tau_f} + \frac{A_s}{\tau_s} e^{-t/\tau_s} - \frac{A_d}{\tau_d} e^{-t/\tau_d} \tag{3}
\]

where $\tau_f$ and $\tau_s$ are the decay times of the fast and slow components of the Ar-Xe mixture respectively; $\tau_d$ is a time characterizing the conversion of Ar$_2^*$ to Xe$_2^*$. $A_f$, $A_s$, and $A_d$ are the intensities of these three terms. In order to normalize Equation 3, the following relation is defined: $A_f + A_s - A_d = 1$. These time constants have been measured by several groups and the summary of these results can be found in Ref. [10]. These measurements have unresolved tension, which is likely due to the Xe doping techniques and/or residual Xe or contaminants in the Ar. These groups also consider higher Xe concentrations which are not considered by this work.

3. Experimental Setup

3.1. Vacuum, Cryogenics, and Recirculation Systems

Our stainless steel cryostat consists of two nested cylinders (inner (IV) and outer (OV) vessels) (Fig. 1). The IV can hold approximately 100 l
of liquid. The space between IV and OV is held near vacuum to prevent convective heating. An aluminized infrared-reflective wrapping (Fig. 1, right) surrounds the inner vessel to reduce radiative heating from the warm outer can. The inner vessel is attached to the lid of the outer vessel with four G-10 rods.

The vessel is drip cooled using an AL60 Cryomech cold head providing approximately 60 W of cooling power at 80 K. The cold head is mounted on top of the OV and is connected to the IV through a thin walled, 2.75-in ConFlat bellows. The cold head drips condensed Ar down a thin stainless steel rod directly to the bottom of the inner vessel. The cold head is surrounded by a Teflon funnel, which is used to separate the warm and cold Ar gas. As Ar evaporates, it flows up the funnel toward the cold head, recondenses, and drips back into the liquid. Warm Ar gas is inserted above the cold head and falls as it cools. The cold head is instrumented with a 50 W heater and a thermocouple, which maintains the cold head at LAr temperature.

The inner vessel was filled from the ullage of a research grade LAr cylin-
Both inner and outer vessels are leak checked prior to each fill. No thorough vessel pump and bake cycling was found necessary as a result of our re-circulation system described below. Gas was flowed through a SAES PS4-MT3/15-R getter, which can purify noble gasses to less than 1 ppb of all common contamitnates, but not noble gases \(^\text{[17]}\). The quantity of Ar in the inner vessel is measured using a shipping scale accurate to ±0.5 lbs. The mass of LAr in the inner vessel is measured as the vessel filled at a liquefaction rate of 0.58 kg/hr. This implies a net cooling power of about 44 W, assuming the Ar gas started at a temperature of 300 K and is liquified at 87.15 K.

In order to purify the liquid argon after the initial fill, gaseous argon in the ullage of the IV is pumped through the getter using a stainless steel bellows pump that uses PTFE and stainless steel for all wetted surfaces. The flow is controlled using a mass flow controller \(^\text{[21]}\) and set to 6 slpm to match the optimal flow rate of the getter. The flow rate fluctuates no more than 1% around the set point. The purified gas is then condensed again at the coldhead. The ullage pressure was initially set to 550 torr but then raised to 640 torr (described in Section 5). At both pressures, the weight of the LAr and pressure of the GAr in the ullage do not change while recirculating. This implies that our coldhead is able to supply enough power to match the heat load of recirculation system.

3.2. Gas Doping System

The gas doping system consists of a small measured cylindrical volume, 19±1 ml, connected to a supply gas bottle on one side and the recirculation system on the other (Fig. 2). The gas pressure is measured using a high pressure Baratron gauge with a precision of 1 Torr and a range from 0 to 2000 Torr. Prior to any doping run, the doping volume is pumped out using a turbo pump to ensure cleanliness. The doping gas is inserted in steps of \(~0.5\) ppm (\(~1500\) Torr) with the largest achievable step size of \(~0.9\) ppm. Larger doping steps are not considered due to the constraints from the maximum range of the high pressure gauge and the small size of the doping volume. The pressure in the doping volume is chosen to be larger than that of the recirculation system to ensure the gases mix adequately once the dop-

\(^2\)upper limit from vendor for research grade LAr states purity to be at least 99.999% pure.
ing volume is exposed to the recirculation system. The remaining gas in the
doping volume is expected to mix based on the diffusion rate of the binary
gas mixture \((33, 31)\), exposure time and small doping volume. About 30
minutes pass between each doping step to ensure that the doping volume has
had time to fully mix with the recirculated gas. The doping volume is evac-
uated before each doping step using a dry scroll pump down to a pressure
of 10 mTorr. We estimate that a negligible amount of gas is lost while evac-
uating the doping volume. During the last doping step, the doping volume
is over pressured and the dopant is slowly leaked into the vacuum system
using a \(\frac{1}{4}\)-in bellows valve until the desired pressure is reached. This last
step gives excellent control over the total amount of dopant inserted per run.

Figure 2: Piping diagram for the gas insertions.

It is assumed that given enough time, the dopant inserted in the gas will
adequately mix with the LAr through diffusion at the liquid/gas interface.
This hypothesis is validated from the results N doping (see Section 5). It is
known to the authors that dopants will have different equilibrium concen-
trations in GAr versus LAr. For example, it is known that the vapor pressure
of N in GAr is about three times larger than in LAr \([5]\) while vapor pressure
of Xe in GAr is about 4000 times less than in LAr \([15]\). In either case, as
long as the ratio of the dopant’s vapor pressure in GAr compared to LAr is much smaller than the ratio of the GAr mass to LAr mass (~4000 for our case), it is assumed that almost all of the dopant will dissolve in the liquid. For these measurements, only dopants that are soluble in LAr are chosen.

The use of high purity gases ensures that the number of mols of the dopant could be calculated using the ideal gas law (PV = nRT). Molar concentration in ppm can then be calculated by using the total weight of the liquid argon volume measured using a shipping scale (119±0.2 kg). The measured temperature of the lab room (~295 K), after the doping system has reached equilibrium, is taken as the doping gas temperature. This temperature typically fluctuates about 2 K around the set point of the room thermostat. As mentioned above, the measured pressure fluctuates by about 1 Torr for each doping step. Each doping step has approximately the same measurement uncertainty (~5%) due to insertion of gas at the same pressure, volume, and temperature. Hence, as the total number of doping cycles increased, the relative measurement error on the total concentration decreases, causing the total error to be dominated by the systematic error of the doping volume.

3.3. Electronic Systems

The inner vessel is instrumented with two 12-stage Hamamatsu, 3-in PMTs (R11065) \[28\] which are box&linear-focused, with synthetic silica 3-in windows and bi-alkali photo-cathodes designed to operate at LAr temperatures. The PMT cathode ground scheme is chosen to allow signals and high voltage to share a cable, reducing the heat load from the wires. Each PMT is instrumented with a fiber optic cable and a blue LED which are used to check the stability of the PMTs. One PMT was lost while the cryostat was being filled with LAr though the other PMT remained stable and had constant gain through out data taking. The PMT signals are amplified using a Phillips Scientific Octal Variable gain amplifier Nim Model 777 \[30\]. The gain of our single PMT was estimated using dark count to be \((4.64 \pm 1.45) \times 10^6\) after accounting for the \(\times 2.86\) amplification of the pre-amp circuit. The conflat feed-throughs are mounted on the warm side of the cryostat to ensure thermal cycling did not damage the ceramic/steel interface. Signals are digitized using a Tetronix TDS 3052B digital oscilloscope and read out to a laptop using custom Python software. The scope has a slow read-out rate of ~1 Hz though it can digitize at 1 GS/s allowing for 1 ns bins over 10 µs.
3.4. Pulse Finding

A pulse finding method was adapted from Ref. [34], which uses a smoothed derivative method to identify signal pulses as large (3.5σ) fluctuations in the noise. Four threshold crossings of the derivative define a pulse. An example of this pulse finding can be seen in Fig. 3. Local minima are identified using the derivative allowing individual pulses to be distinguished in the event of pulse pileup. This feature proved essential for our measurement of the \( ^3\Sigma_u^+ \) lifetime of Ar.

3.5. TPB Coating

The PMT windows have a mounted tetraphenyl butadiene (TPB) coated acrylic disk to wavelength shift the Ar scintillation light into the sensitive region of the PMTs. To prevent the PMTs from floating in the LAr, the TPB disks are tied down to the Al holders with stainless steel wire. The average quantum efficiency over the TPB emission spectrum is \( \sim 29\% \), making it well suited for measuring VUV scintillation [4]. TPB is vapor deposited to a thickness of about 2 µm. The QE of the TPB coating was never measured.
because we planned to do a relative light yield measurement.

After completing an initial test run, it was found that TPB from the acrylic disks had dislodged and redeposited in a ring on the wall of the cryostat at the approximate height of the PMT windows (Figure 4). Additionally there were pieces scattered on the bottom of the IV that were likely floating on top of the LAr. Though TPB has been shown to be stable in liquid argon [6], it is speculated that the volatile boundary between the gas and liquid phases of the Ar was abrasive to the vapor deposited TPB. As the liquid level rose above the PMTs, no further abrasion was observed. If this abrasion happened in for the data presented in this paper, it is not known what effect this has on the light yield of the TPB. Light from the ring of TPB should not significantly contribute to the light seen in the PMT because the TPB ring is at the horizon of the PMT’s view. WLS light from the suspected floating TPB particulate likely could be seen. We estimate that this WLS light contributes negligibly to the total light seen because of the small relative area of the TPB flakes when compared to the total surface area of the liquid.

4. Triplet Lifetime Measurement

4.1. After-pulsing

After-pulsing was noticed in the data between 1 $\mu$s and 3 $\mu$s. In order to understand the effects of after-pulsing, a large LED pulse was used to generate after-pulses. The charge weighted pulse arrival time distribution of
the after pulses can be seen in Figure 5. The first after-pulse has an area that is 20.5% of the initial peak, the second pulse has an area that is 5.2% of the initial peak and the third after-pulse has an area of about 1.5% of the initial peak. The exact area of third after-pulse is harder to determine because the dark count has a large contribution to its total area. The effects of after-pulsing cannot be reduced without decreasing the single photon resolution and thus the nominal operation voltage of 1600 V was chosen.

Based on the literature describing after-pulsing (111), it is clear that after-pulsing will most likely be triggered from the singlet component of the scintillation light due to the large size and short duration of the initial signal. In contrast, the probability of triggering an after-pulse from a single photo electron is estimated to be small (<1%) based on the number of large events seen in dark rate calibration data. The relative contribution of after-pulses should not change with dopant concentration because the singlet intensity does not change with these concentrations. Thus when doing a light yield comparison, the effects of after-pulsing will not drop out when looking at the relative ratio of photo-electrons before and after doping but rather be a constant effect regardless of doping concentration.

Figure 5: A large LED pulse can be seen at ∼1 µs and after pulses can be seen peaking at 1.6 µs, 2.1 µs, and 2.6 µs

To estimate the contribution of after-pulsing to the total charge of a distribution, an exponential is fit in two side-bands windows around the first after-pulse peak. The windows are chosen to be 1.30 to 1.45 µs and 1.75 to
1.90 $\mu$s. This exponential is assumed to be the underlying distribution around the first after-pulse peak. The fit is then subtracted from the waveform leaving behind an estimate of the after-pulse. An example of this procedure performed on pure LAr data can be seen in Figure 6. Through this technique, the contribution of after-pulses to the total charge of a waveform can be made and later added in as a systemic effect when reporting on light yield. Finally, it is worth mentioning that an attempt to subtract out the after-pulsing peaks using the shapes found in the after-pulsing calibration data was performed. This method ultimately failed because although the shapes appear similar by eye, they do not match well enough to adequately remove the after-pulsing without distorting the distributions.

Figure 6: An example of how after-pulsing is estimated using an exponential fit. The blue markers are data, while the solid red line is the exponential fit to the side bands of the first after-pulse peak, and the solid green histogram is the waveform subtracted estimate of the after-pulses. A zoom in of the after pulse region is shown in the insert.

4.2. Fitting the Triplet Lifetime

We investigated mitigating the after-pulsing effect on the LAr $^3\Sigma_u^+$ lifetime using two fitting methods. In both methods the singlet was not fit because the time resolution of the initial trigger was too large. Data was taken using a random trigger in which the trigger level was set to $\sim 15$ PE.
Figure 7: Two fits to the $^3\Sigma^+_u$ scintillation distribution in pure LAr. The left plot is the fit with a single exponential and a constant from 2.5 $\mu$s to 10 $\mu$s. The right plot shows the fit with a Landau shape for the first after-pulse and a Gaussian shape for the second after-pulse with an exponential and a constant fit the tail. The fit to the after-pulsing region is shown in the insert.

In the first fitting method, the $^3\Sigma^+_u$ lifetime was found using a single exponential plus a constant fit. A constant is added to account for GAr scintillation light coming from the ullage. A start time of 2.5 $\mu$s was chosen in order to avoid the first two the after-pulses. A $^3\Sigma^+_u$ lifetime of 1456 $\pm$ 38 ns was measured with this method. Alternatively, we fit a Landau shape to the first after-pulse and a Gaussian shape to the second after-pulse. These functions are added to the first function and the sum was fit starting from 1.75 $\mu$s. The average $\chi^2/\text{NDF}$ for this second fitting method was 1.44 $\pm$ 0.12 indicating that this chose of functions is reasonable. This method yields a systematically larger value of the $^3\Sigma^+_u$ lifetime of 1502 $\pm$ 30 ns. An example of these two different fitting methods can be seen in Fig. 7. It is likely that this second fitting method yields a larger value because the two after pulse function account for some of the light at early times thus flattening the exponential component of the fit. An average was taken as the true $^3\Sigma^+_u$ lifetime, measured to be 1479 $\pm$ 38 (stat) $\pm$ 23 (sys) ns where the statistical uncertainty was taken to be the larger of the two different fitting method results and the systematic uncertainty was taken to be difference in the two fitted $^3\Sigma^+_u$ lifetimes. It is estimated, using the technique mentioned above, that after-pulsing contributes about 6.4% to the total charge.

5. Nitrogen Doping

5.1. Measuring Birk’s Constant

In order to validate the gas doping and recirculation system, a small measured quantity of N$_2$ was injected using the methods described in Section
Following the injection, the N\textsubscript{2} was filtered out using the getter. The effects of N on Ar scintillation light are well known and have been characterized in Refs. [5] and [18]. The relationship between the measured triplet lifetime (τ\textsuperscript{3′}) and the true triplet lifetime (τ\textsuperscript{3}) follows Birk’s law:

\[ τ_3' = \frac{τ_3}{1 + τ_3 \cdot k \cdot C} \]  

(4)

where \( k \) is Birk’s constant and \( C \) is the molar concentration of impurities measured in parts per million. Birk’s constant has been measured to be 0.11 ppm\(^{-1}\)µs\(^{-1}\) and 0.13 ppm\(^{-1}\)µs\(^{-1}\) for N in LAr by the two previously mentioned authors. They obtained \( τ_3 = 1.26 \) µs compared to our value of 1.48 µs. Reference [5] was aware that their \( τ_3 \) was smaller than that typically measured (23, 1.46 µs) and speculated that if they used the fitting method of Ref. [23] and extrapolated their \( τ_3 \) using Eqn. 4 to pure LAr, their value becomes \( τ_3 = 1.45 \) µs, which is in good agreement with our measurement once errors are included.

Before doping, the recirculation and purification system was run and \( τ_3 \) measured frequently. Once the \( τ_3 \) measurement was stable, it was assumed that the Ar was pure. The nitrogen contaminant was then inserted and \( τ_3 \) decreased according Eqn. 4. As the Ar was purified, it is assumed that the contamination concentration decreased as:

\[ C(t) = C_0 \cdot e^{-t/τ_{filter}} \]  

(5)

where \( C_0 \) is the initial concentration measured in molar parts per million and \( τ_{filter} \) is a fitted time constant characterizing the filtration process.

We performed one \( N_2 \) doping run injecting 1.06±0.07 ppm of \( N_2 \). The nitrogen was injected in two steps using the techniques outlined in Section [3]. The triplet lifetime dropped from 1479 ± 38(stat) ± 23(sys) ns to 1238 ± 15(stat) ± 20(sys) ns as seen in Fig. 8. The quenched triplet lifetime and associated errors are calculated in the same way as in Section [4]. It is found that after-pulsing contributes about 7.7% to the total charge seen in these quenched waveforms. From this reduced lifetime a Birk’s constant of 0.12±0.03 ppm\(^{-1}\)µs\(^{-1}\) was found, consistent with the previous measurements. Using this measured value of Birk’s constant, Eqn. 4 was fit to the triplet purification curve in order to evaluate the effectiveness of the recirculation system. Initial measurements of the filtration time constant found it to be 9.5±0.6 d (left fitted curve in Fig. 9). To increase the filtration rate, the
Figure 8: Triplet lifetime (µs) versus run time (days) for the nitrogen doping run. The nitrogen was injected at run time -1.1 days. It can be seen that the nitrogen thoroughly mixed after about three hours.

turbo pump was valved off from the guard vacuum while leaving the roughing pump on thereby decreasing the insulating vacuum from $5.0 \times 10^{-7}$ Torr to $1.4 \times 10^{-3}$ Torr. Additionally the cold-head set point was raised from -190°C to -187°C thus increasing the internal pressure of the cryostat from 550 Torr to 640 Torr. A modification to Eqn. 4 was made to account the for non-zero start time to obtain:

$$\tau_3' = \frac{\tau_3'}{1 + \tau_3' \cdot k \cdot C_0 \cdot e^{-\frac{t - t_0}{\tau_{filter}}}}$$

(6)

where $t_0$ was the non zero start time of the fit. The filtration time constant measured by fitting Eqn. 6 to the triplet purification curve after these changes was measured to be 4.5±0.3 d (right fitted curve in Fig. 9).

By monitoring the cosmic ray muon peak before and after doping (see Section 6), it was found that the most probable value (MPV) shifted from $334\pm21\pm7$ (stat) PE to $283\pm22\pm7$ (stat) PE, which corresponds to a $15.3\pm1.5\%$ drop. This drop is compatible with Ref. 5 who measured the total quenching of LAr waveforms doped with 1 ppm N$_2$ using gammas from a $^{60}$Co source.
6. Xenon Doping

Having validated our doping procedure with nitrogen, we then used xenon. Xe doping was done in four steps of 1.00±0.06 ppm, 2.0±0.1 ppm, 5.0±0.3 ppm, and 10.0±0.5 ppm. Xenon was injected in 0.5 ppm increments until the desired molar concentration was reached. High concentrations of Xe (≈100 ppm) were not considered\(^3\). Once the concentration was stable and fully mixed, 50,000 random background events were collected at each concentration to create the scintillation distributions seen in Fig. 10. Scintillation distributions are created by binning pulse arrival times weighted by the pulse charge. The waveforms were fit from 1.05 \(\mu\)s to 10 \(\mu\)s to use as much of the waveform as possible while avoiding the singlet contribution. The start time of the fit was varied by \(\pm 25 \text{ ns}\) in order to estimate the systematic uncertainty associated with the fit window. After-pulsing contributions to the total charge for 1, 2, 5, and 10 ppm were estimated to be 2.7%, 2.5%, 2.0%, and > 1% respectively.

\(^3\)The largest possible insertion step is \(\sim 0.9 \text{ ppm}\) and each 0.5 ppm injection took about 30 min. Additionally, the next Xe concentration of interest is at \(\sim 100 \text{ ppm}\) where previous studies have been performed. Thus achieving this next concentration would take in total 2 days of continuous doping.
A modified version of Eqn. 3 was used to measure the time constants of the mixture in which the fast component was removed:

\[ I(t) = \frac{A_s}{\tau_s} e^{-t/\tau_s} - \frac{A_d}{\tau_d} e^{-t/\tau_d} + \frac{A_{\text{long}}}{\tau_{\text{long}}} e^{-t/\tau_{\text{long}}} \]  

(7)

where \( \tau_s \) is the slow component, \( \tau_d \) is the energy transfer time between the Xe and triplet state of the Ar, \( A_1, A_2 \) and \( A_3 \) are the intensities, and \( \tau_{\text{long}} \) is an additional time constant to account for the exponential tail at late times. This artifact is consistent with light coming from the cold gas in the ullage [15].

The slow component, \( \tau_s \), represents the time constant for the combination of unquenched Ar dimer light and Xe dimer light created through the mechanism outline in Section 2.1. Though Xe decays rather quickly compared to Ar, the population of Xe dimers is pumped by the Ar-Xe mixed state, thus is able to have a much longer decay time compared to pure liquid Xe. The total concentration of Xe dimers and Ar dimers in the triplet state is \( A_1 e^{-t/\tau_s} - A_2 e^{-t/\tau_d} \). The fitted waveforms are seen in Fig. 11 and the results from these fits are summarized in Table 1. As the Xe concentration increases,
the energy transfer process becomes more effective. This is reflected in the fact that $\tau_d$ is anti-correlated with Xe concentration. Additionally, $\tau_s$ decreases because the relative population of Xe dimers to Ar dimers in the triplet state increases with increased Xe concentration.

Previous measurements [10, 32] of $\tau_s$ for 1 ppm Xe reported values between 1 µs and 2.5 µs, though the uncertainty on the measured concentration is also 1 ppm. At 10 ppm Xe, $\tau_s$ has been measured as 280 ns and 750 ns, with the concentration uncertainty at least 3 ppm. $\tau_d$ has not been measured for 1 ppm Xe, but for 10 ppm the value ranges from about 250 ns to about 700 ns. Past measurements of these time constants have a much larger Xe-concentration uncertainty making them difficult to compare with our measurements. Disagreements between the previous measurements is likely due the concentration uncertainty and the unknown initial xenon concentration.

Figure 11: Fit of Eqn. 7 for Xe doped waveforms with Xe concentrations of 1, 2, 5, and 10 ppm. The solid red line is the total fit, the dashed blue line is the long exponential component associated with $\tau_l$, and the magenta dashed line is the Xe dimer and Ar triplet component ($A_1 e^{-t/\tau_s} - A_2 e^{-t/\tau_d}$). Deviations from the fit at short times (1.5-2.5 µs) are due after-pulsing.
Table 1: Summary of Xe doping time constants, measured in ns, for various molar concentrations from 1 to 10 ppm Xe. The first uncertainty is a systematic uncertainty estimated by varying the fit start time from 1.025 µs to 1.075 µs and the second uncertainty is statistical.

7. Light Yield Comparison

The total increase in light yield resulting from Xe doping is evaluated with two methods. First, a direct comparison is made of the position of the muon peak seen in the pure Ar random trigger data to the muon peak in 10 ppm XeDLAr random trigger data. Second, the relative shift of the mean value of the random trigger distributions is measured. Cosmic muons can be seen in the random trigger data at large photo-electrons (PE) because the typical energies deposited are of order 100 MeV while nuclear decay energies are of order 1 MeV. The shape of the expected distributions were verified with an initial Geant4 simulation and from this simulation it is estimated that events with less than ~100 PE are likely due to nuclear decays and not cosmic muons. The muon peak in both data sets is well modeled by a Landau function. Both data sets were fit in two different ranges in order to estimate this uncertainty. For the Ar data, the lower boundary of the fit range is fixed at 100 PE while the upper boundary is varied from 450 to 600 PE. A MPV of 334±21(syst)±7(stat) PE is found. The fit for XeDLAr data had the lower bound fixed at 100 PE while the upper boundary is varied from 2000 to 10,000 PE. A MPV of 1004±47(syst)±4(stat) PE is found. The position of this peak increased by a factor of 3.00±0.24(syst)±0.06(stat). If the mean of each distribution is calculated, the pure LAr data set has a mean value of 1067±68(syst)±4(stat) PE while the 10 ppm XeDLAr data set has a mean value of 2050±20(syst)±14(stat) PE which leads to a relative average light yield increase of 1.92±0.12(syst)±0.02(stat). This second method yields a smaller increase because a large portion of the photo-electron spectrum in
the pure LAr spectrum falls above the Landau peak and as such the spectrum as a whole is not well modeled by a single function. On the other hand, the distribution in the XeDLAr spectrum is fit well by a Landau. This light yield increase is larger than past results ([32, 27]) due to the longer attenuation length in XeDLAr at 175 nm in a larger active volume.

8. Conclusion

It has been demonstrated that a small concentration of gas can be reliably and precisely injected into a cryostat containing \( \sim 120 \) kg of LAr. Nitrogen at a concentration of 1.06\( \pm 0.07 \) ppm was inserted and the quenching factor (Birk’s constant) was measured to be 0.12\( \pm 0.03 \) ppm\(^{-1}\)\( \mu s^{-1} \), consistent with past literature. This demonstrates that small quantities of gas can be precisely injected. The \( N_2 \) contaminant was then removed using a ullage recirculation and purification system with a filtration time constant as low as 3.6 d.

We investigated XeDLAr concentrations of 1.00\( \pm 0.06 \) ppm, 2.0\( \pm 0.1 \) ppm, 5.0\( \pm 0.3 \) ppm, and 10.0\( \pm 0.5 \) ppm with the largest uncertainty being 6% compared to past experiments which achieved 23.5%. We have measured the xenon doping time constants as outlined in past experiments and give confidence to these measurements due to the small uncertainty in our doping concentrations. A relative light yield increase of 1.92\( \pm 0.12 \) (syst)\( \pm 0.02 \) (stat) is found for 10 ppm Xe compared to pure Ar.

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