Position Reconstruction in LUX

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Abstract: The \((x, y)\) position reconstruction method used in the analysis of the complete exposure of the Large Underground Xenon (LUX) experiment is presented. The algorithm is based on a statistical test that makes use of an iterative method to recover the photomultiplier tube (PMT) light response directly from the calibration data. The light response functions make use of a two-dimensional functional form to account for the photons reflected on the inner walls of the detector. To increase the resolution for small pulses, a photon counting technique was employed to describe the response of the PMTs. The reconstruction was assessed with calibration data including \(^{83m}\text{Kr}\) (releasing a total energy of 41.5 keV) and \(^3\text{H}\ (\beta^- \text{ with } Q = 18.6 \text{ keV})\) decays, and a deuterium-deuterium (D-D) neutron beam (2.45 MeV). In the horizontal plane, the reconstruction has achieved an \((x, y)\) position uncertainty of \(\sigma = 0.82\) cm for events of only 200 electroluminescence photons and \(\sigma = 0.17\) cm for 4,000 electroluminescence photons. Such signals are associated with electron recoils of energies \(~0.25\) keV and \(~10\) keV, respectively. The reconstructed position of the smallest events with a single electron emitted from the liquid surface has a horizontal \((x, y)\) uncertainty of 2.13 cm.

Keywords: dark matter, position reconstruction, xenon
1 Introduction

The LUX detector, a 370 kg liquid-gas dual-phase xenon time projection chamber (TPC) [1], has as its main scientific goal the observation of nuclear recoils resulting from hypothetical dark matter particle candidates, called Weakly Interacting Massive Particles. The excited xenon atoms and ionization electrons from the recoil in the liquid phase are observed through two different light signals: the prompt scintillation signal (called S1) and the charge signal (called S2). S1 arises from the direct production of xenon excited states and the recombination of some electrons with ions in the liquid. S2 is generated by the drifting of the ionization electrons that do not recombine towards the liquid surface, where they are extracted and accelerated into the gas phase producing electroluminescence. The light from both S1 and S2 is observed by two arrays of photomultiplier tubes (PMTs) placed on the top and bottom of the detector. The detection of both signals ensures a good reconstruction of the position of the interaction: the depth of the interaction is obtained from the time separation between S1 and S2, while the \((x, y)\) coordinates are obtained from the distribution of the S2 light among the PMTs. Assuming a parallel field geometry, the \((x, y)\) position of the interaction can be taken as that of the electroluminescence production (i.e. the S2 signal). The energy depositions of the recoils in the liquid xenon are considered point-like in the energy range
of interest. For example, a 10 keV electron has a range of about 2 µm while that of a 5 MeV alpha particle is less than 50 µm. These numbers are much smaller than the typical size of the electron cloud in the \((x, y)\) plane due to electron diffusion along the transverse direction. At the center of the LUX chamber (drift time = 160 µs), the standard deviation of the electron transverse diffusion in the \(x\) and \(y\) direction is between 1 mm (for a diffusion constant of 30 cm\(^2\)/s [2]) and 1.4 mm (for a diffusion constant of 60 cm\(^2\)/s [3]).

A good 3D position reconstruction of the events is paramount in the analysis of dual phase detectors to ensure a good description of the observed events. Most of the radiological background events are localized closer to the walls, the photomultipliers or the electrode grids, leaving the center of the detector with a very low background [4]. Some especially troublesome backgrounds originate from the radioactive decay of \(^{210}\)Pb, \(^{210}\)Bi and \(^{210}\)Po plated on the Polytetrafluoroethylene (PTFE) walls of the detector (pp. 127–133, [5]). These decays mimic low energy nuclear recoils and cannot be discriminated from a true recoil of a xenon nucleus using the ratio between S2 and S1. We reject those events by preferably searching for signals in the detector central volume — the fiducial volume. Consequently, a precise and accurate event position reconstruction, especially close to the walls of the detector and to small S2 sizes (S2 = 0–4,000 detected photons, phd hereafter), is essential to accurately model external backgrounds from high rates at the walls to progressively lower rates towards the center of the detector.

Position reconstruction is also essential to correct for variations of the number of S1 and S2 photons throughout the chamber. These variations are due to the light collection dependence on the interaction position, the finite electron lifetime due to the presence of impurities, and to local distortions of the electric field [6]. The detector calibration with respect to nuclear recoils also requires a precise reconstruction of the vertex of neutron interactions in the detector [7].

Several \((x, y)\) position reconstruction algorithms have been employed in scintillation detectors. The oldest is the center of gravity or centroid method (often used in the Anger camera [8]), in which the position of the interaction is obtained from a weighted average of the PMT responses. A simple weighted average is biased towards the center of the array due to the finite extent, but the position can be partially corrected using a lookup table with corrections that depend on the position or an alternative parameterization [9]. Another choice of position algorithm involves the use of artificial neural networks [10], which have been employed by the XENON collaboration [11].

Other methods make use of a statistical test, such as the \(\chi^2\) test or other maximum likelihood techniques [12]. They determine the position of interactions by matching expected PMT outputs with the observed values. These methods have the advantage of giving an estimator (e.g. \(\chi^2\) minimum) that can be used to assess the quality of the position reconstruction. For each PMT, the expected outputs as a function of the event position are usually stored in a Monte-Carlo generated lookup table (e.g. ref. [13]) or described by an empirical light response function (LRF), which characterizes the response of the PMT as a function of the position of the emission of the light \((x, y)\).

All the aforementioned algorithms require the use of some sort of calibration data for which the position of interaction is already known. In large detectors such as LUX, it is not feasible to use radioactive sources to produce energy depositions at desirable locations in the inner region of the detector, due to the large volume and self-shielding properties of liquid xenon. Thus, the solution usually adopted is to make use of simulated data, which has limitations since the simulations may
not describe accurately all the intricacies of the light collection such as the precise optical properties of the materials in these detectors.

The position reconstruction algorithm employed in the LUX experiment (named Mercury, originally developed for the ZEPLIN-III dark matter experiment [14]) is a statistical-based algorithm using a maximum likelihood test to find the best set of output parameters. Mercury uses S2 photons to reconstruct the \((x, y)\) position of the event. It employs LRFs to predict the response of each PMT for interactions at an arbitrary distance from that PMT. The innovative aspect and major advantage of this method is the way in which the LRFs are obtained: it employs a virtual scan method, using the detector’s own data with minimal or no reliance on simulations. In this virtual scan, the LRFs are obtained iteratively through a sequence of fits to the calibration data until the response functions converge simultaneously. This method is described in detail in section 2.

LUX was designed to achieve a very low energy threshold (down to 1 keV for a nuclear recoil) aiming to be very sensitive to low mass WIMPs. This means that the signals of interest can be of very small amplitude (a few detected photons per PMT) which imposes new challenges to the position reconstruction. To optimize the precision and accuracy of the algorithm for this energy region, a new method was implemented (section 3) to count individual photons in channels with very low signal. Another major difference from the original version of Mercury is that the LRFs in this work are 2-dimensional (axial and polar) instead of 1-dimensional to account for the reflection on the walls of the detector (section 2). The ZEPLIN-III detector had low-reflectance internal walls, and thus the axial symmetry was a good approximation.

The position reconstruction method presented here was used in the analysis of the LUX data from 2013 (WS2013), including both the original analysis (85.3 live days of data, [15]) and the reanalysis (95.0 live days, [16–18]), as well as in the analysis of the data collected from 2014 until 2016 (WS2014–16, 332.0 live days, [19, 20]) and the calibration data used to monitor and calibrate the detector [7, 21, 22].

Mercury has also been used with some modifications in the DarkSide-50 experiment [23], the Panda-X experiment [24], and proven to work well for a gamma camera for medical imaging [25].

This article is organized as follows: the method of obtaining the LRFs (virtual scan) is described in section 2; in section 3, we introduce the method implemented to find the position of emission of S2 light; in section 4, we discuss the use of the position reconstruction algorithm on calibration data, the associated uncertainties, and the position resolution.

## 2 The Light response functions

The determination of the shape of the light response functions (LRFs) is an essential feature in Mercury as they are used to estimate the expected response of the detector. The light response function of a PMT \(i\), \(\mathcal{H}_i\), is proportional to the probability that a S2 photon emitted at \((x, y)\) and detected by any of the PMTs is detected in channel \(i\). These functions may change from PMT to PMT because the light collection depends on the relative position of the PMT in the array. Besides the \((x, y)\) position, the minimization method described in section 3 also estimates the associated uncertainties and the value of the log-likelihood ratio, \(q_{\text{min}}\), associated to the reconstructed position.
**Figure 1.** Top array of the LUX detector with the variables used in the definition of the functional form of the LRFs. The yellow star corresponds to the point of emission of the light from an electro-luminescence signal (S2). $\rho$ is the distance between the PMT and the light source; $(R, \Phi)$ and $(r, \phi)$ are coordinates of the PMT and of the emission of the light position, respectively. The PMTs are grouped according to the distance to the center of the chamber. The numbering scheme reflects the group which the PMT belongs to.

In this section, we start with the description of the LUX detector geometry followed by the discussion of the influence of reflections from the PTFE walls on the S2 light collection. Finally we describe the fitting procedure used to derive the LRFs.

### 2.1 The LUX detector and calibrations

The detector light collection, which determines the $(x, y)$ dependence of the LRFs, depends on many factors such as the geometry of the detector, the optical properties of the internal surfaces, and the PMT characteristics. In LUX, the active liquid xenon region has a dodecagonal shape with a height of 52.4 cm (with a 5.5 cm gas phase layer on top of the liquid), and a distance between the center of the side faces and the center of the detector of $23.65 \pm 0.05$ cm [1]. Both S1 and S2 signal result from far ultraviolet photons with a wavelength of $\sim 175$ nm [26, 27]. This light is detected by 122 Hamamatsu R8778 PMTs organized in two hexagonal arrays of 61 PMTs each, one placed immediately above and the other at the bottom of the sensitive volume [28]. The geometry of the top array is shown in figure 1. Each PMT has a diameter of 57 mm and an active photocathode diameter of 45 mm, except for the PMTs on the periphery which are partially covered by the detector walls.

Each of the 12 side walls of the detector and the space between the photomultipliers are covered by PTFE. The high reflectance of PTFE for the xenon scintillation light (>97% in the liquid phase [29]) ensures a good light collection for both the S1 and S2 signals.

During the WIMP search run, the detector was calibrated using a variety of radioactive sources, both internal ($^{83m}$Kr and $^3$H, injected in the gas system of the detector) and external ($2.45$ MeV deuterium-deuterium (D-D) neutron source, $^{252}$Cf, Am-Be, and $^{137}$Cs) [30]. These calibrations...
were not only used to study the response of the detector for both electronic and nuclear recoils as a function of the deposited energy, but also to develop and study the position reconstruction method. The $^{83m}\text{Kr}$ calibration is particularly important for the latter. It provided the data to i) obtain the LRFs of the PMTs (section 2.3), ii) monitor the quality of the position reconstruction, and iii) measure the position resolution (section 4.1).

The $^{83m}\text{Kr}$ isotope decays with a half-life of 1.83 h occurring in two transitions of 32.1 keV and 9.4 keV respectively, the half-life of the intermediate state being 154 ns [21]. Given the short time between these two decays, the two pulses overlap producing a single S2 signal that has between 4,000 and 20,000 phd distributed among several PMTs, depending on the depth of the event, which gives a sizable signal on the top PMT array but far from the saturation point (~10,000 phd per PMT). After being injected in the circulation system, $^{83m}\text{Kr}$ distributes itself uniformly throughout the sensitive volume of the detector [31]. The uniformity of the reconstructed $^{83m}\text{Kr}$ events gives us insight about the quality of the position reconstruction. Furthermore, the large number of $^{83m}\text{Kr}$-decay events collected during the science runs (10 million events in WS2013) is sufficient to characterize with great detail the S2 light collection dependence on the event position. Single electrons are also a possible source of calibration. They are the lowest possible S2 signal ($\langle S_2 \rangle \sim 22 \text{ phd}$), and thus can be used to study the systematics and statistical errors affecting the reconstruction of the lowest pulses which are of great interest in the LUX analysis.

The $(x, y)$ position reconstruction makes use of the signals from the PMTs of the top array except for two malfunctioning units (marked by a cross in figure 1). Information from the bottom array PMTs is only used for the determination of the total S2 size, as the S2 light collection efficiency in each individual bottom PMT is almost independent on the $(x, y)$ position of emission in this high-reflectance chamber design. As for the S1 signal, its amplitude is relatively small on both PMT arrays, being inadequate to reconstruct the position. From the S2 signal of each PMT $i$ on the top array, two quantities are extracted for the $(x, y)$ position reconstruction: the pulse area, $A_i$, and the photon counts, $N_i$. The pulse area, $A_i$, is obtained by the software integration of the PMT signal along the duration of the pulse, while the number of detected photons, $N_i$, is estimated by identifying and counting the individual discrete single detected photon pulses observed during the emission of the pulse (typically some microseconds for an S2 signal). The PMT signals are recorded in a dedicated Struck board with a sampling period of 10 ns [32], making it possible to implement photon counting software. In the simple method employed, a photon is counted each time the photomultiplier waveform crosses a certain threshold (>1.4 mV, $5\sigma$ above the baseline noise). Both $A_i$ and $N_i$ are used in the statistical method described in the section 3.

2.2 Influence of the wall reflection in the S2 light collection

In LUX, the LRFs are described using two-dimensional analytic functions, dependent on the radial position of the S2 light emission, $r$, and the distance between the PMT and the and the position of the S2 light emission, $\rho$. This is unlike the simpler approach in ZEPLIN-III described in [14], in which one dimensional LRFs were used instead as lateral reflections could be ignored to good approximation due to the low reflectivity of copper (the bi-hemispherical reflectance [33] of polished copper in xenon gas and for $\lambda \sim 175$ nm is less than 13% [34]). In LUX the active region is defined by highly reflective PTFE panels extending up until the PMTs in the top array in order to increase the light collection of the S1 signal and thus increase the detection efficiency for low
energy nuclear recoils. The bi-hemispherical reflectance of these panels in contact with gas was found to be 70% as measured by [35] and >75% as determined in LUX from comparison with Monte Carlo simulations [30]. The bi-hemispherical reflectance of the panels immersed in liquid xenon is much higher (>95% [29]). While the light reflected on the PTFE walls in the gas gap has a significant impact in the light distribution pattern, the reflection of the scintillation light on the walls in contact with the liquid xenon does not influence the shape of the LRFs.

We studied the influence of the reflectance of the inner surfaces on the light collection distribution at each PMT of the top array using the S2s from a LUXSim $^{83m}\text{Kr}$ simulation with full light propagation [36]. Two scenarios were considered: i) all the PTFE inner surfaces are diffusely reflective with a diffuse albedo of 100%, and ii) all the PTFE inner surfaces are non-reflective. In both cases, specular reflection was not considered. For each case, 20,000 $^{83m}\text{Kr}$ events uniformly distributed in the active liquid volume were generated, and the light was propagated until it was detected or absorbed. Figure 2 shows the S2 light collected at a PMT close to a detector wall. As shown, for the case with non-reflective inner surfaces, the PMTs collect almost only direct S2 light, which is emitted isotropically, with exception of two small components: light that is reflected on the liquid/gas interface and light that crosses that interface and returns to the gas via Rayleigh scattering in the liquid. On the contrary, for reflective surfaces, the light collection is no longer axially symmetric, showing a saddle extending from the center of the photomultiplier towards the wall. It is clear from this study that in LUX the light collection depends not only on the distance between the PMT and the S2 position but also on the distance of the S2 position to the walls, requiring two variables to describe the LRFs.
2.3 LRF determination

We implemented a new model for the LRFs that takes into account the light reflected on the walls and takes advantage of the symmetries of the detector by adding an axial ($\eta$) and a polar ($\varepsilon$) component calculated for each PMT $i$ on the top array. This function can be written as

$$H_i(r,\rho) = C_i [\eta(\rho) + \varepsilon_i(r,\rho)],$$

(2.1)

where $\rho$ corresponds to the distance between PMT $i$ centered at ($R_i, \Phi_i$) and the light emission at ($r, \phi$) (figure 1). $C_i$ are normalization constants to equalize the response of the PMTs for the same number of incident photons per unit of PMT window area under the same geometrical conditions. Differences in the values of $C_i$ are due to different quantum efficiencies of the photomultipliers for xenon scintillation and different exposed photocathode areas as some PMTs are partially covered with PTFE tiles reducing the light collection.

The functional form of the axial component was determined using LUXSim [36], being described by the following empirical function

$$\eta(\rho) = \frac{A}{(1 + \gamma^2 \rho^2)^\frac{3}{2}} + m\rho + b,$$

(2.2)

where $A$, $\gamma$, $m$ and $b$ are the fitting parameters. The first term, a bivariate-Cauchy function, describes the light that goes directly to the PMT or is reflected on the gas/liquid interface, while the polynomial term is necessary to describe the light that is reflected inside the liquid bulk or in the PTFE trefoils placed between the PMTs.

The polar component, $\varepsilon$, cannot be described using the same function for all PMTs, as in the case of $\eta$, given that PMTs closer to the walls detect more reflected light. However, the symmetry of the LUX detector allows us to use the same LRF for PMTs with the same radial position $R_i$. Therefore, the function $\varepsilon$ has to be found for each of the 8 different groups of PMTs in the top array (identified in figure 1 with the numbers 2–9). This approach significantly reduces the number of parameters and is more robust as the symmetry of the chamber is directly incorporated in the LRF model. In the initial iterations, only the PMT groups closer to the walls ($R_i > 18$ cm), where the polar component is more significant, are used. After at least 5 iterations, all the PMT groups are incorporated in this fit.

The shape of the curve used to fit the polar component to the experimental data was also inferred from simulations of the light collection in the detector. These simulations showed that this component is described in first approximation by the following empirical function:

$$\varepsilon_g(\rho, r) = \kappa_g \exp\left(\frac{r}{\xi_g}\right) \exp\left(-\frac{\rho}{\zeta_g}\right),$$

(2.3)

where the index $g$ indicates the PMT group ($g = 2$–9) and $\kappa_g$, $\xi_g$ and $\zeta_g$ are the fit parameters. This function ensures that the polar component increases with the radial coordinate of the light emission location, $r$, as expected, since the effect of the wall reflection is larger for events at larger radii. However, it was clear that this simple function was not sufficient to describe the intricacies of the light collection in the detector. Hence, in the final iterations of the process of the LRF determination...
Figure 3. Flowchart of the LRF fitting process.

that will be described next, an alternate format of eq. (2.3) that included more fitting parameters was used:

\[
\varepsilon_g (\rho, r) = \left[ \kappa_g \exp \left( \frac{r}{\xi_g} \right) + \alpha_g (r) \right] \exp \left( -\frac{\rho}{\zeta_g + \beta_g (r)} \right) \\
\alpha_g (r) = k_{g,1} \exp \left[ -\frac{(r - k_{g,2})^2}{2 (k_{g,3})^2} \right] + k_{g,4} r + k_{g,5} \\
\beta_g (r) = k_{g,6} \exp \left[ -\frac{(W_c - r)^2}{2 (k_{g,7})^2} \right],
\]

(2.4)

where \((\kappa_g, \xi_g, \zeta_g, k_{g,1}, k_{g,2}, \ldots, k_{g,7})\) correspond to the fitting parameters for each group of PMTs and \(W_c = 24.5\) cm is the position of the corners of the wall in the LUX dodecagon. Although this function is more complex than eq. (2.3), the new terms only introduce small corrections to the LRFs. \(\kappa_g\) is larger than \(\alpha_g\), and the constant term \(\zeta_g\) dominates with an average value \(\sim 0.19\) cm, while \(k_{g,6}\) is only, on average, \(\sim 0.04\) cm.

Eq. (2.1) is used for all the PMTs with the exception of the central channel \((R = 0)\). As this PMT has axial symmetry relative to the active volume of the chamber, its LRF can be described using the axial component, \(\eta(\rho)\), only (given by eq. (2.2) but in an independent fit).

The experimental data used to obtain the LRFs consisted of a sample of \(^{83m}\)Kr calibration S2 pulses with unknown positions. For this, we selected 100,000 \(^{83m}\)Kr events close to the top of the detector (at depths ranging from \(\sim 8\) cm to \(\sim 12\) cm). This number of events ensures the fitting is done in a reasonable time and is sufficient to describe the light collection in the detector. For each \(^{83m}\)Kr event and for each PMT, the pulse fractions, \(F_i\), are defined as

\[
F_i = \mathcal{A}_i / \mathcal{A}_T \quad \text{or} \quad F_i = N_i / \mathcal{A}_T,
\]

(2.5)
\( \mathcal{F}_i \) being a sample of the function \( \mathcal{H}_i (r, \rho) \). \( \mathcal{A}_F \) is the total pulse area of the S2 pulse summed over all the photomultipliers in the chamber.

The functions \( \mathcal{H}_i (r, \rho) \) are obtained by an iterative process. The flowchart of the fitting process that follows is shown in figure 3. In each iteration \( m \), \( C_i^{(m)} \), \( \eta_i^{(m)} (\rho) \), and \( \varepsilon_i^{(m)} (r, \rho) \) are fitted in this order. The functional forms of \( \eta (\rho) \) and \( \varepsilon_i (r, \rho) \) are fitted to their values calculated with eq. (2.1) using the experimental values of \( \mathcal{H}_i \) given by \( \mathcal{F}_i \) and the event position estimates from the previous iteration. The constants \( C_i^{(m)} \) and the functions \( \eta_i^{(m)} \) and \( \varepsilon_i^{(m)} \) are then employed to get better estimates of the positions of the experimental events using the statistical method described in section 3. These positions are used in the next iteration. The event positions used in the first iteration are obtained with a corrected centroid method [14].

The constants \( C_i^{(m)} \) are obtained by equalizing the radial response of the PMTs from the top array. The method used to determine \( C_i^{(m)} \) is as follows: the observed axial component, \( \eta_{\text{obs}} \), is computed as \( \eta_{\text{obs}} = \mathcal{F}_i / C_i^{(m-1)} - \varepsilon_i^{(m-1)} \) (in the first iteration \( C_i^{(m-1)} \) and \( \varepsilon_i^{(m-1)} \) are set equal to 1 and 0, respectively). The values of \( \eta_{\text{obs}} \) with \( \rho \in [0, 15] \) cm are grouped in sections 1 cm wide in \( \rho \). For each section, we calculate the ratio between the average over that section of the axial function for PMT \( i \) and that for all the PMTs of the top array. The factor \( C_i^{(m)} \) is given by the average over all the sections of these ratios multiplied by \( C_i^{(m-1)} \).

To obtain \( \eta_i^{(m)} (\rho) \), the function given by eq. (2.2) is fitted to its values at the experimental values given by \( \mathcal{F}_i / C_i^{(m)} - \varepsilon_i^{(m-1)} \) (in the first iteration \( \varepsilon_i^{(m-1)} \) is set equal to zero).

For the determination of both \( C_i^{(m)} \) and \( \eta_i^{(m)} \), only events within a strip with thickness \( d \) along the radial line between the current PMT center and the farthest wall are used, selected by the conditions

\[
| r \sin (\phi - \Phi_i) | < d \quad \text{and} \quad r \cos (\phi - \Phi_i) < R_i.
\]

In this fit, we set \( d = 0.25 \) cm.

To determine \( \varepsilon_i^{(m)} \), the function given by eq. (2.3) (or by eq. (2.4) for the final iterations) is fitted to its values at the experimental event positions given by \( \mathcal{F}_i / C_i^{(m)} - \eta_i^{(m)} \).

The \( C_i^{(m)} \), \( \eta_i^{(m)} \), and \( \varepsilon_i^{(m)} \) are then used to update the estimated coordinates of the positions of the experimental events (using the statistical method described in section 3) that will be used in the next iteration. In the first 5 iterations, a uniformization of the event positions (see section IV in ref. [21]) obtained in the previous iteration was performed in the radial direction, in order to speed-up the process.

The iteration process finishes once the average value of the \( q_{\min} \) (minimum of the likelihood ratio defined next, eq. (3.3)) does not change significantly between consecutive iterations (\( |\Delta \langle q_{\min} \rangle | < 0.1 \) typically). For LUX, the number of iterations needed to find a good set of functions \( \{ \mathcal{H}_i \} \) was around 20, clearly larger than than the 5 iterations needed in ref. [14] when the simpler axial approximation was used.

The good agreement between the data and the LRFs is illustrated in figures 4 and 5. The former shows the final iteration of the fit of eq. (2.2) to the experimental data; in figure 5, the colors represent the observed fractions \( \mathcal{F}_i \) (panels a and b) or the corresponding LRFs (panels c and d) as a function of the reconstructed interaction position \( (x, y) \) for a PMT in the central region \( (R_i = 10.4 \) cm) and a PMT close to the walls of the detector \( (R_i = 18 \) cm). The LRFs are mostly axial symmetric for PMTs in the central region, approaching an oval shape for PMTs closer to the
walls due a larger polar component. In the latter case, the position of the maximum is not coincident with the center of the PMT being pushed towards the wall of the detector as already expected from the simulation results shown in figure 2.

3 The statistical method

A maximum likelihood algorithm [37, 38] was implemented to search for the \((x, y)\) position of the interaction that would produce in an array of \(N\) PMTs the expected outputs \(S = \{S_i(x, y)\}\) \((i = 1, ..., N)\) which are the closest to the observed outputs \(\mathcal{D} = \{\mathcal{D}_i\}\). This consists in finding the position \((x, y)\) that maximizes the likelihood function,

\[
\mathcal{L}(\mathcal{D}|S) = \prod_{i=1}^{N} p_i(\mathcal{D}_i|S_i),
\]  

Figure 4. The axial component of the LUX PMTs. \(\{\mathcal{F}_i/C_i - e_i\}\) is shown as function of \(\rho\) — the distance between the center of the PMTs and the position of the event. The polar component as predicted by the function \(e\) is being subtracted. The curve is the fit with eq. (2.2).

Figure 5. 2D histogram of the fractions \(\mathcal{F}_i\) from \(^{83}\text{Kr}\) for \(R_i = 10.4\) cm (panel a) and \(R_i = 18\) cm (panel b) events and the respective LRFs (panel c and d). The contours of the photomultipliers and the border of the chamber are shown in gray, while the center of the PMT under analysis is marked by a black cross.
where $p_i$ is the probability for the PMT $i$ to have the observed output $D_i$ given the expected output $S_i$. Provided that the response of the PMTs and the subsequent electronics is linear, the expected outputs can be written as

$$S_i = D_T H_i (x, y), \quad \text{with} \quad D_T = \sum_{i=1}^{\text{all PMTs}} D_i, \quad (3.2)$$

where $D_T$ is the total number of detected photons summed over all the photomultipliers in the chamber.

Instead of maximizing $L(D|S)$, it is more convenient to minimize the log-likelihood ratio [39] given by

$$q = -2 \left[ \ln L - \ln L_0 \right] \quad \text{with} \quad L_0 = L(D|D), \quad (3.3)$$

where $L_0$ corresponds to the likelihood maximized in an unconstrained way (in this analysis, we assumed $S = D$ in the computation of $L_0$).

The probability $p_i$ can be written as

$$p_i (D_i|S_i) = \sum_{n=0}^{+\infty} P_i(n; S_i) \cdot u_i(n; D_i), \quad (3.4)$$

where $P_i(n; S_i)$ describes the fluctuations of the number of detected photons, $n$, for an expected output $S_i$; and $u_i(n; D_i)$ characterizes the response of the PMT $i$ and the respective signal processing electronics, being equal to the probability that the response is $D_i$ for $n$ detected photons. In this work we assume $P_i(n; S_i)$ to be the same for all the PMTs and given by a Poisson distribution:

$$P_i(n; S_i) = \left[ S_i^n \exp(-S_i)/n! \right]. \quad (3.5)$$

As previously mentioned, the PMT outputs $D_i$ can be assessed by pulse areas or by photon counting. The formulation of the maximum likelihood method in each of these cases is described next, along with an innovative mixed method, which uses both as proxies for $D_i$ depending on the pulse area recorded for each PMT.

### 3.1 Pulse areas

In the pulse area method, the PMT response for a single detected photon is $u(1)$ with $\langle u(1) \rangle = 1 \text{phd}$. Therefore, the response for $n$ detected photons ($n > 1$) is obtained by the following recurrent convolution

$$u(n) = u(n-1) * u(1). \quad (3.6)$$

The probability density $p_i$ is given by the sum of the response for $n$ detected photons with weights given by Poisson distributions and taking the pulse areas, $\mathcal{A} = \{A_i\}$, as estimators for the measured outputs, $\{D_i\}$, leading to

$$p_i (\mathcal{A}_i|S_i) = e^{-S_i} \left[ \delta (\mathcal{A}_i) + \sum_{n=1}^{+\infty} \frac{(S_i)^n}{n!} u_i (n; \mathcal{A}_i) \right], \quad (3.7)$$
where the Dirac delta function, $\delta(A_i)$, corresponds to the case $n = 0$, and the expected outputs, $S_i$, are obtained from

$$S_i = \mathcal{A}_T \mathcal{H}_i(x, y), \quad \text{with} \quad \mathcal{A}_T = \sum_{i=1}^{\text{All PMTs}} A_i.$$  

(3.8)

To obtain the likelihood ratio $q$, we can use the probability distribution from eq. (3.7) in eq. (3.3).

When the number of photons detected in each PMT is large enough, Gaussian statistics can be used for $P_i$ instead of the Poisson distribution. In the asymptotic Gaussian limit (Wilks’ theorem [40]), $q$ is distributed as the following simple $\chi^2$ minimization

$$q \equiv \chi^2 = \sum_{i=1}^{N} \frac{(S_i - \mathcal{A}_i)^2}{S_i (1 + \sigma_i^2)},$$

(3.9)

where the term $(1 + \sigma_i^2)$ represents the excess noise factor for each PMT [41]. This factor measures the degradation of the signal compared with Poissonian statistics. It can be shown that $\sigma_i$ is the standard deviation of the response to single photons.

The pure pulse area method was used in the initial analysis of the WS2013 data [15]. In this analysis, the single photon response, $u(1)$, was obtained from LED calibrations ($\lambda = 420$ nm). It was found that $u(1)$ was well described by a Gaussian for all PMTs with the value of $\sigma_i$ ranging from 0.4 to 0.6 phd [42]. The simpler $\chi^2$ minimization was used for $\mathcal{A}_T > 2,000$ phd.

### 3.2 Photon counting

In the photon counting technique, the response of a PMT for $n$ detected photons is $u(n) = \delta(n)$; thus, $p_i$ follows a Poisson law, $q$ being given by (equation 39.16 in ref. [43])

$$q = 2 \sum_{i=1}^{N} \left[ S_i - N_i + N_i \ln \frac{N_i}{S_i} \right].$$

(3.10)

For the PMTs with no photon detection ($N_i = 0$), we have $N_i \cdot \ln(N_i / S_i) = 0$.

The main advantages of using the counting over the pulse area method are that it is much less computationally expensive, and the excess noise factor is 1, which improves the energy resolution. Moreover, the pulse areas method has a poor performance for very low signals given that the fluctuations in the baseline noise are integrated in the signal, while in the photon counting method most of the baseline noise is well rejected by the threshold discriminator. The latter fact can be directly assessed from figure 6 which shows a comparison of the resolution between these two metrics using pulses from a single extracted electron.

Another main advantage of photon counting is the manner in which the double photoelectron emission from a single xenon scintillation photon is treated [44]. For xenon scintillation light, 20% of the photons produce two photoelectrons per photon on the photocathode of the PMT. In this case, the single photon response function, $u(1)$, can no longer be approximated by the Gaussian single photon response of the PMTs obtained with blue LED light, in which case the photons do not have enough energy for double photoelectron production. This greatly complicates the use of the pulse area integration method that requires the knowledge of $u(1)$, but has no effect on the photon counting method, as the two photoelectrons are emitted simultaneously and thus counted as corresponding to a single photon.
**Figure 6.** Distribution of the pulse areas (PA) and photon counts (PC) for pulses with a single extracted electron from the liquid. The single electrons were emitted between the S1 and the S2 of the $^{83m}$Kr. Both histograms are fitted with Gaussian distributions for pulse size between 10 and 27 phd (fit-range indicated by the dashed vertical lines), the upper bound defined to avoid contamination from double electron pulses. The fit to the pulse areas is centered at 22.05±0.09 phd with a resolution ($\sigma/\mu$) of 23.1±0.6%, while the fit to the photon counting histogram has a mean of 22.04±0.06 phd and a resolution of 20.5±0.4%. This improvement of 2.6±0.7% matches well the expected 2.7% estimated from the reduction of the noise factor. Only events with a reconstructed radius smaller than 16 cm were considered in this analysis.

Photon counting is limited by the pile-up that occurs when two or more photons are detected almost simultaneously, producing a signal indistinguishable from that of single photon. The probability of pile-up depends on the time resolution, $\tau$, assumed to be the same for all the photomultipliers, and on the photon flux. The time resolution, $\tau$, is the minimum time difference between two detected photons to ensure they are not merged into a single count. It was estimated from the analysis of S1 signals to be 30 ns.

To estimate the effect of pile-up, we approximated the light flux during the emission of the S2 signal to a Gaussian distribution defined by a standard deviation $\sigma_{S2}$, estimated for each S2 pulse. $\sigma_{S2}$ increases with the drift time of the event due to longitudinal diffusion of the electrons while drifting to the liquid surface [45]. From the analysis of the S2 signals, it was found that the average value of $\sigma_{S2}$ ranges from 360 ns to 830 ns for events from the top and bottom of the liquid layer, respectively. The probability of pile-up was computed in a dedicated Monte-Carlo simulation for different values of $\sigma_{S2}$ and pulse areas of the PMTs. These results were then used in the $q$ of the mixed method described in the following subsection.

### 3.3 Mixed method

In the mixed method, the response of the PMTs is described with the photon counting method when the pulse area, $A_i$, observed for a PMT $i$, is smaller than a certain threshold, $T$, and the pulse areas method above that threshold. The threshold depends on the pulse width of the event, and the log-likelihood ratio is given by the value of $q$ for the PMTs described by pulse areas plus the value
of $q$ for the PMTs described by photon counting. By combining eqs. (3.9) and (3.10) we have:

$$q = \sum_{i=1}^{N} 2 \left[ S_i - N_i \frac{N_i}{S_i} \right] + \sum_{i=1}^{N} \frac{(S_i - A_i)^2}{S_i \left( 1 + \sigma_i^2 \right)}. \quad (3.11)$$

The threshold, $T$, is set such that the contamination of the pile-up in the photon counting is smaller than 5%, determined from the Monte-Carlo simulations described in the previous sections. It was found to be given by

$$T = 0.39 \cdot \frac{\sigma_{S2}}{\tau} \text{ (phd)}, \quad (3.12)$$

where $\tau = 30$ ns.

The mixed method was used both in the reanalysis of the WS2013 results [16] and in the WS2014–16 results [19]. In both analyses, the pulse areas are calibrated using $^{83m}$Kr calibration data in order to ensure compatibility between photon counting and pulse areas [30]. The mean values of the two histograms on figure 6 agree within uncertainties, showing good agreement between the photon counting and the pulse areas methods.

3.4 Position uncertainties

The analysis of the shape of $q$ around the minimization point ($q_{\text{min}}$) is used to estimate the uncertainties associated with the position reconstruction. Figure 7 shows the contour lines of the $q$ minimization for three different small S2 events ($A_T < 4,000$ phd). As shown, the $q$ surface is smooth, well behaved and has an unambiguous global minimum. The $q$ is axially symmetric for events in the central region of the detector ($r < 20$ cm), while for events near the walls the surface is elongated along the radial direction, reflecting the larger uncertainty on that coordinate.

Considering the symmetry of the $q$ contours, the uncertainties in the reconstructed positions are best expressed in cylindrical coordinates ($\sigma_r, \sigma_\phi$). To obtain them, we select the curve in the $(x, y)$ space where $q = q_{\text{min}} + 1$ [43]. The azimuthal, $\sigma_\phi$, and the radial, $\sigma_r$, uncertainties are given
by the distance between this curve and the position of the minimum along the azimuthal and the radial directions, respectively. Even though this method is only exact for a Gaussian distribution, the results presented in section 4 show that these values are a good estimate of the position uncertainty for the smaller S2 pulses.

The uncertainties in the reconstructed positions decrease with increasing number of top array PMTs involved in the reconstruction. However, that improvement is only relevant up to 20 PMTs (including all 59 working channels instead of 20 PMTs improves the position uncertainty by only 3%). For this reason, we considered in the position reconstruction procedure only the 20 PMTs closest to the initial position estimate (obtained by the corrected centroid algorithm [14]). This is done in order to minimize the interference from noise sources (such as the emission of single electrons or afterpulsing in a PMT), which typically lead to events located near the wall of the detector to be reconstructed towards the center. It also improves the speed of the reconstruction by 10%.

The position reconstruction fails or is affected by a large systematic error when the reconstructed signal is not from a single scatter S2 or the S2 is affected by some source of noise such as afterpulsing in a PMT or the presence of an unstable PMT etc. Those events have, on average, a larger minimum value of $q_{\text{min}}$, and thus a quality cut based on the value of $q_{\text{min}}$ can be used to remove those events. In the WS2014–2016, we considered only events with $q_{\text{min}} < 40 + \mathcal{A}_{\text{tot}}/42$. The efficiency of this cut calculated using calibration data (namely $^3\text{H}$) was above 95% [19] for $\mathcal{A}_{\text{tot}}$ between 200 and 4,000 phd.

4 Results and discussion

The position of each interaction is an essential quantity in the LUX analysis, since both the radius and the azimuthal angle are two of the observables ($r$, drift time, $S_1$, $S_2$, and $\phi$ in WS2014–16) of the profile likelihood analysis used for the determination of the exclusion limits [19]. It is thus crucial to assess the quality of the position reconstruction and determine the uncertainties of the reconstructed positions. As mentioned before, due to the detector design and size, it is not practical to obtain experimental data with known positions of interaction allowing the direct comparison of the reconstructed positions with the original positions. However, calibration data from dispersed sources have a known distribution of events in the detector. The most useful calibrations for the position reconstruction assessment are the $^{83m}\text{Kr}$ (as previously mentioned, used for the construction of the LRFs), the $^3\text{H}$ ($\beta^-$ with $Q = 18.6$ keV), and the 2.45 MeV D-D neutron calibrations.

In the analyses that follows, WS2013 data are used unless stated otherwise. However, both runs use the same set of LRFs and the same position reconstruction method with no observable changes in the light collection. Therefore, the main conclusions presented here are still valid for the WS2014–16 data. The calibration data used here was processed in the same manner as in the analysis of recently published LUX science results [16, 19].

4.1 Krypton-83m data

The reconstructed ($x, y$) distribution of $^{83m}\text{Kr}$ decays in the detector is shown in figure 8 for events occurring at the top (drift time between 4–10 $\mu$s) and bottom of the detector (drift time between 290–320 $\mu$s). From figure 8, it can be seen that the reconstructed coordinates do not extend to the
edge of the top PMT array. This effect is stronger for longer drift times. Detailed simulations of the electric field in the LUX chamber (described in ref. [6]) explain this observation by the existence of a radial component of the drift field, pushing electrons towards the center of the detector as they drift upwards and hence shifting radially inwards the position of S2 relative to that of the interaction. In the WS2013, the main origin of the radial field component is the electrical transparency of the cathode grid and the lateral field rings of the TPC, which allow some field leakage. A nearly identical effect is observed in the XENON100 detector [46]. This radial field also depends on the azimuthal angle, an effect which is also visible in figure 8. This may be caused by a combination of azimuthally-varying field leakage across the grid and possibly a non-uniform distribution of accumulated charge on insulating surfaces of the detector. In the WS2014–16 run, the radial field was much stronger compared with that in the WS2013 run (see ref. [6] for details), but with no direct impact on the LRFs.

As background rejection by fiducialization requires the knowledge of true interaction positions, i.e. the vertex location, the reconstructed positions have to be corrected from the radial field effect. For that, the $^{83}$Kr events, assumed to be uniformly distributed in the volume of the detector, were used to create a map $M$ that converts the reconstructed S2 position into the real vertex positions $x_{\text{ver}}$ and $y_{\text{ver}}$

$$(x_{\text{ver}}, y_{\text{ver}}) = M(x, y, \text{drift time}).$$  

A thorough discussion of the creation of those maps is presented in ref. [21]. In the following analyses, S2 reconstructed positions are used and not the real vertex positions unless stated otherwise.
Figure 9. Electron trajectories (in white) and electric potential (color-map) near gate grid wires obtained by simulation. The gate grid is shown at $z = 0$ cm by the white dots and the surface of the liquid is represented by the black line. Note that the line density is not intended to illustrate the field strength.

In figure 8, one can observe a striped pattern in the event density, parallel to the wires forming the gate grid (both the gate grid and the observed pattern are at an angle of 15° to the $x$ axis). The grid wires are 0.1 mm in diameter and 5 mm apart. The grid plane is ~4 mm below the surface of the liquid and 1 cm below the anode grid. This grid separates the drift field (defined by the cathode and gate grids, the former being 48 cm below the gate grid) and the extraction/electroluminescence fields (defined by the gate and anode grids, the latter being 4.8 cm below the top PMTs) in the detector, allowing these two fields to be set independently. Given the large difference between the drift field (180±20 V/cm in the WS2013) and the extraction field (2.84±0.16 kV/cm in liquid in WS2013), the drift field lines are compressed as they pass through the gate plane; any electrons leaving the drift volume appear only in narrow strips between each pair of gate wires. This effect is shown in figure 9, with the path of the electrons along the field lines represented by the white lines and the color-map representing the electric potential. We estimated, using a Monte Carlo simulation, that this effect contributes 1.1 mm to the position resolution of the vertex true event $(x_{\text{ver}}, y_{\text{ver}})$, independent of the event depth.

The visualization of this focusing effect of the field lines between the grid wires can be used to assess the quality of the position reconstruction. The histogram on figure 10 represents the density of the events occurring right below the gate plane (0 to 1 cm below the gate) and along the direction perpendicular to the gate wires. Each peak was fitted with a Gaussian distribution to obtain the central position and adjacent strips width. The results show that for the selected region the average distance between two wires is $5.13\pm0.07$ mm, matching the grid pitch (5 mm) within $2\sigma$. Additionally, the resolution of the wires determined from the Gaussian width is on average $\sigma = 0.965\pm0.028$ mm for the selected region, similar to the average uncertainty from the position reconstruction method ($\sigma = 0.86$ mm, see section 4.4). While the strips in the reconstructed position distribution are mostly uniform right below the gate grid (figure 8 left), large variations in the number of events in each strip are observed in the remaining volume (figure 8 right). These variations are mostly likely to be caused by small variations in the pitch of the wires producing...
Figure 10. Histogram of the density of $^{83m}$Kr events just below the gate grid represented on an axis perpendicular to the orientation of the gate wires. The original $(x, y)$ position of the events were rotated by $15^\circ$ in such a way that the direction of the new $x$-axis ($x'$) is parallel to the gate wires, and the direction of the new $y$-axis ($y'$) is perpendicular to the gate wires. Only $^{83m}$Kr events at a depth less than 1 cm below the gate are considered.

Figure 11. Histogram of the S2 event density for $^3$H $\beta$-decay events for the top of the detector (black curve) and for $^{83m}$Kr events for the top (blue curve), middle (green curve), and bottom (red curve) of the detector. For clarity we added a value of $+100$ events/cm$^2$ to the green curve, $+200$ events/cm$^2$ to the blue curve and $+300$ events/cm$^2$ to the black curve.

fluctuations of the bulk field which are not observed for the S2 right below the gate wires.

4.2 Tritium data

The detector was also calibrated using a tritiated methane source [22, 47], where the $Q$-value of the $^3$H $\beta$-decay is 18.59 keV. We collected more than 300,000 tritium events in a relatively high rate (10 Bq) calibration [22], which is sufficient to compare with $^{83m}$Kr calibrations. As in the case
of $^{83m}$Kr, the $^3$H events are expected to be uniformly distributed in the liquid, but the size of the S2 is smaller, up to 6,000 phd. The range of S2 for $^3$H is comparable to what is expected from a WIMP event in the search region. For this reason, most of the PMT pulses are described using the photon counting technique instead of pulse area integration as is the case with $^{83m}$Kr. Therefore, $^3$H is an excellent source to assess the quality of the position reconstruction for smaller S2 pulses and to confirm the consistency of the mixed model described in section 3.

Similarly to $^{83m}$Kr, the distribution of reconstructed tritium events exhibits the same striped pattern in the $(x, y)$ density of events and the presence of the same radial field. Figure 11 shows the event density profiles along the direction perpendicular to the wires for these two sources and for different drift times. As shown, the event density profiles are aligned as expected, showing no significant systematic error affecting the position reconstruction as function of the drift time and between the two sources.

Another test performed using tritium data was the comparison between the radial density of $^3$H and $^{83m}$Kr events (figure 12). The two histograms match well except for a small systematic difference for the events with $r^2 > 400 \text{ cm}^2$. This disagreement is caused by the larger position uncertainties of the tritium events due to their smaller size [30].

4.3 Neutron data

The response of the detector to nuclear recoils was calibrated using a D-D neutron generator that emits monoenergetic neutrons with an energy of 2.45 MeV [7, 48, 49]. The D-D neutrons are collimated by a 4.9 cm internal diameter air-filled tube which extends between the walls of the shielding water tank and the detector cryostat. Since the position of the first neutron scatter is located in the direction of the beam, the data from these calibrations were used to check for any systematic error affecting the reconstruction.

Figure 13 shows the reconstructed positions of the first neutron scatter. The color scale represents the event density within a circle with a radius of 1 cm normalized to the number of events.
along the neutron beam. No correction from the radial field effect was applied to data as, at the height of the neutron beam, this correction is small (only 4%).

To check the straightness of the reconstructed beam, we divided the beam along the $y$-axis in 2 cm thick slices, and then for each slice we fitted the histogram of the $x$ position to find the center of the beam. The fit is composed of a semi-circle function describing the elastic neutron events inside the beam and a skew-normal distribution describing the background from multiple scatter events and other non-nuclear recoil events. The results are represented in figure 13 by the black dots with the error bars corresponding to the uncertainty obtained in the fit plus a systematic contribution from the size and position of the bins. These data are well described by a linear fit represented by the gray line. The root-mean square of the difference between the center of the beam set by the center of the data slices and the fit is only 1.4 mm with the maximum absolute deviation being 4 mm.

### 4.4 Position uncertainties and resolution

The LUX position reconstruction code calculates two statistical position uncertainties ($\sigma_r, \sigma_\phi$) in the reconstructed positions (section 3.4). These uncertainties are used in: i) the neutron calibrations, to estimate the uncertainty in the scattering angle of double scatters [7]; and ii) calculating the wall-event contribution to the background model, to estimate the contamination of wall events in the fiducial volume [19].

Figure 14 shows both uncertainties as a function of the S2 area in the top array, $A_{\text{top}}$ ($\langle A_{\text{top}}/A_T \rangle = 0.566\pm0.019$ for $r < 20$ cm). This figure includes calibration data from both tritium (blue dots) and D-D nuclear recoil events (red crosses), for a reconstructed radius smaller
than 17 cm (dark red/blue markers) and larger than 17 cm (light red/blue markers). As shown, the uncertainties depend mostly on $\sigma^2 \propto 1/A_{\text{top}}$ reflecting the Poisson distribution of the photon statistics.

We can normalize the uncertainties to the number of photons, defining $\Upsilon_r$ and $\Upsilon_\phi$ as

$$\Upsilon_{(r,\phi)} = \sigma_{(r,\phi)} \sqrt{A_{\text{top}}}. \quad (4.2)$$

The average values of $\Upsilon_r$ and $\Upsilon_\phi$ as a function of the radius $r$ are represented on figure 15. As the figure confirms, $\Upsilon_r$ has a significant radial dependence, being about 40% larger for peripheral events than for central ones. Conversely, $\Upsilon_\phi$ is mostly constant. Additionally, the values of $\Upsilon_r$ and $\Upsilon_\phi$ depend on the distance of the event to the center of the nearest PMT. We observed that both uncertainties are on average 8% larger for S2 pulses generated right below the center of a PMT than for those near the border of a PMT. We can explain this by looking to the first derivative of the radial component, $|\eta'|$, (see figure 4) which is maximized for $\rho$ between 3 and 5 cm. The number of PMTs that fall into that region is larger when the light is emitted near the border of a PMT.

An S2 due to a single electron that escapes from the liquid into the gas is the smallest electroluminescence signal that can be observed ($A_T \approx 22$ phd, figure 6); thus, the study of the single electron events is paramount to assess the reconstruction of very small signals and to determine the lower limit of the position resolution. In a detector such as LUX, the single electrons are typically caused by [50, 51]:

1. delayed single emission from a previous S2 event: electrons can accumulate under the surface barrier at the liquid/gas interface and escape into the gas later;
Figure 15. Average position uncertainties $\mathcal{U}_r$ ($\sigma_r \sqrt{A_{\text{top}}}$, blue curves) and $\mathcal{U}_\phi$ ($\sigma_\phi \sqrt{A_{\text{top}}}$, red curves) as a function of the reconstructed radius $r$. The colored regions correspond to the 1-$\sigma$ range on the distribution of the position uncertainties.

Figure 16. Radial statistical uncertainty, $\sigma_r$, as a function of the pulse area of the top PMT array. The data are from the neutron calibrations, tritium calibrations, single electrons from $^{83m}$Kr, and simulations of D-D neutron calibrations, covering a wide range of pulse areas. For the simulations, the uncertainty corresponds to the root mean square of the difference between the reconstructed radius and the true radial position of the event. Only the events with a reconstructed radius smaller than 20 cm and a drift time between 50 and 300 $\mu$s are shown. The uncertainty for single electron events is $\sigma_r = 2.13$ cm (average top pulse area of 13.8 phd). The dashed line corresponds to the S2 threshold used in the WS2013 analysis [16, 17].
Figure 17. Difference between the reconstructed $x$ position of a S2 $^{83m}$Kr signal, $x(S2)$, and the reconstructed $x$ position of the single electron observed after the $^{83m}$Kr signal, $x(SE)$. Only single electrons signals with a total pulse area from 10 to 35 phd, $r < 20$ cm, and $\Delta \tau < 20$ $\mu$s are analyzed. The dashed black curve corresponds to the fit with the sum of two Gaussians centered at zero, and the dashed gray curve corresponds to the contribution from the broader Gaussian distribution.

(2) photoionization of impurities in the liquid by the S2 or S1 photons.

The delayed electrons (1) are expected to have the same reconstructed $(x, y)$ position of the parent S2 signal, so they can be used to directly measure the position resolution for single electrons. On the contrary, single electrons from photoionization (2) are created anywhere along the path of the S2 (or S1) light in the liquid being almost uncorrelated with the position of the parent S2.

In this study, we selected single electrons emitted after the S2 pulse of $^{83m}$Kr events. The uncertainty in the $(x, y)$ position of the $^{83m}$Kr S2 pulse is negligible compared with that of the single electron since it is much larger (between 4,000 and 20,000 phd). This selection of single electrons includes both single electrons from delayed emission and photoionization. To reduce the background from the latter, only single electrons observed within a time window, $\Delta \tau$, less than $\Delta \tau < 20$ $\mu$s after the end of the $^{83m}$Kr S2 signal were accepted.

The distribution of the difference between the reconstructed $x$ position of the $^{83m}$Kr S2 signal and the reconstructed $x$ position of the associated single electron, $\Delta x$, is shown in figure 17. This histogram is fitted with the sum of two Gaussians with different widths, both centered at zero. The narrower distribution corresponds to the single electrons from the delayed emission while the wider Gaussian describes the contribution from photoionization single electrons. The resolution of the single electrons given by the standard deviation of the narrower Gaussian distribution was determined to be 2.24$\pm$0.04 cm, comparable to the average uncertainty in the $x$ position (2.13 cm) obtained for these single electrons in the position reconstruction. In contrast, the standard deviation of the wider Gaussian is about 11 cm. The results for the $\Delta y$ match exactly with the results for $\Delta x$.

The contamination of the single electrons from photoionization can be estimated by measuring the relative area of the wider Gaussian distribution. For our selection of $\Delta \tau < 20$ $\mu$s, the photoionization accounts for 46.8% of all the single electrons. For larger values of $\Delta \tau$, the number of single
electrons from delayed emission as function of $\Delta \tau$ follows an exponential decay law with a decay time constant of $\sim 40 \mu s$.

We checked the existence of any possible drift of single electrons from delayed emission along the liquid surface of the detector as this would deteriorate the position resolution. The existence of such drift was tested by measuring the average value of $\Delta x$ and $\Delta y$ as function of $\Delta \tau$. For this, the data were sliced in 20 $\mu s$ sections in $\Delta \tau$ up to a maximum value of 200 $\mu s$ and for each of those slices the $\Delta x$ and $\Delta y$ histograms were fitted using a double Gaussian with the mean of both Gaussian distributions not fixed. We observed that for $\Delta \tau < 200$ $\mu s$ the absolute value of the fitted mean is always smaller than 2 mm for both $\Delta x$ and $\Delta y$, and no significant increase in the standard deviation of the distribution was observed. This is a good indication that there is no significant movement of the charge along the surface of the liquid.

We performed this same analysis using only the pulse areas to reconstruct the position of single electrons instead of the mixed method adopted through this paper (both described in section 3.1). The resolution obtained in the double Gaussian fit was $2.47 \pm 0.04$ cm which corresponds to a degradation on the resolution of single electrons of $10.27 \pm 0.25\%$. This shows that the mixed method employed here improves the reconstruction of low energy events.

5 Summary

A statistical method was developed to obtain the $(x, y)$ position of an interaction in the LUX detector from the observed S2 signal distribution at the top PMT array. This method employs in situ calibrations to obtain the LRF for each photomultiplier. The presence of PTFE reflectors around the sensitive volume increases the complexity of the LRFs. These were written as the sum of two terms: an axial component $\eta(\rho)$ describing the light that goes directly to the PMTs or is reflected in the liquid surface and a polar component $\epsilon(\rho)$ characterizing the light reflected from the PTFE walls.

We assessed the quality of the position reconstruction using calibration data obtained during LUX detector operations. Although in these data the position of each individual interaction is not known, the resolution can be measured using both the effect of the focusing effect of the electrons going through the gate grid and the single electrons emitted right after a $^{83m}$Kr S2 event. This resulted in an observed resolution of $0.0965 \pm 0.0028$ cm for the $^{83m}$Kr S2 pulses (with average areas of 22,000 phd) and $2.24 \pm 0.04$ cm for single electrons (with average areas of 22 phd), which agree well with the calculated uncertainties obtained in the position reconstruction algorithm ($0.086$ cm for $^{83m}$Kr S2 pulses and $2.13$ cm for single electrons).

The analysis of systematic errors was done by looking to the uniformity in $(x, y)$ of both $^3$H and $^{83m}$Kr reconstructed events, the linearity of the neutron beam produced in the D-D neutron calibrations, and by measuring the average distance between two wires of the gate grid. All these tests verified the absence of significant systematic uncertainties affecting the position reconstruction.

For the position reconstruction of very small S2s ($A_T < 4,000$ phd), we employed a method where the response of a PMT is described using photon counting or pulse areas according to the pulse size observed in the PMT. This method was faster when compared to a pure maximum likelihood method based only on pulse areas. Moreover, we observed no significant systematic error on the low S2 pulse data ($^3$H and single electrons) when compared with the large S2 pulse
($^{83m}$Kr), and the analysis of the position resolution of single electrons obtained with the mixed method revealed an improvement of $10.27 \pm 0.25\%$ when compared with a pure pulse areas method.

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