

Directionality in Kilo-ton Scale Scintillating Neutrino Detectors

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Liquid scintillator detectors are some of the most prevalent detectors due to their good energy resolution and scalability to large volumes. This has made them particularly attractive for neutrino measurements, and this technology is at the heart of many of the most important measurements in this field. Their weakness is that they only provide information on the energy of the particle. The extraction of a second signal like the particle direction would greatly enhance the scientific reach of these detectors especially for searches for neutrinoless double-beta decay. In this paper, we develop a technique for extracting particle direction and evaluate different detector advances that could be used to make direction reconstruction a reality in a kilo-ton scale detector.

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I. INTRODUCTION

Liquid scintillator based detectors are responsible for several of the critical measurements that have cemented our understanding of three-neutrino oscillations. These measurements include KamLAND's measurement reactor antineutrino oscillation at a distance of ~ 200 km, Borexino's measurement of ^7Be solar neutrino oscillation and most recently the short baseline reactor antineutrino experiments that measured oscillations due to θ_{13} at a distance of 1 km: Daya Bay, Double Chooz, and RENO. Scintillating neutrino detectors will continue to be important for the next set of neutrino measurements from the determination of the neutrino mass hierarchy to elastic scattering measurements and sterile neutrino searches and for non-proliferation applications. Their scalability to large volumes makes them particularly attractive for neutrinoless double-beta ($0\nu\beta\beta$) decay searches, and currently one of the best limits for the $0\nu\beta\beta$ half-life comes from KamLAND-Zen.

The advantage of liquid scintillators for measurements in the ~ 1 MeV range is their simple scalability from 1 ton to 1 kiloton while providing energy resolutions of $\sim 5\%$. This is roughly a factor of two better than water Cerenkov detectors, the other technology that is easily scaled to these large masses. While the energy resolution is good due to the abundant scintillation light, this light is isotropic and does not retain the directional information of the primary particle. In contrast, the direction of the particle can be reconstructed from the Cerenkov cones in water-based detectors, but the energy resolution rapidly degrades below ~ 5 MeV.

In a scintillating detector, Cerenkov light is produced although most is absorbed and reemitted as part of the scintillation processes; however, some fraction retains its directional information. If this directional Cerenkov light can be isolated from the copious isotropic scintillation light then it may be possible to reconstruct the direction of the primary particle or at least determine something about the topography of the event. The addition of directionality is a powerful tool for background rejection and could be used to search for new physics especially in the case that $0\nu\beta\beta$ is observed. In this paper, we develop a technique for separating the Cerenkov and

scintillation using the photon arrival times and evaluate different detector technologies that would allow the realization of direction reconstruction in kilo-ton scale scintillating neutrino detectors.

II. LIGHT PRODUCTION IN LIQUID SCINTILLATORS

Liquid scintillators are cocktails of aromatic hydrocarbons. When charged particles move through a scintillator, the molecules are excited, predominantly the non-localized electrons in the π -bonds of the phenyl groups. Vibrational and rotational modes of the molecules are turned into heat within picoseconds through collisions with other molecules. Within ~ 10 picoseconds, the π -electrons de-excite to the first excited state from higher levels through radiationless transitions. The first excited state de-excites through photon emission. There are two characteristic times for this de-excitation depending if the singlet state or the triplet state was excited. The singlet state will de-excite within nanoseconds while the triplet state de-excites on the order of 10's or 100's of nanoseconds. These two processes are fluorescence and phosphorescence respectively. The exact time constants for these processes are determined by the composition of the scintillator.

The molecules in liquid scintillators are not isolated. Radiationless processes transfer energy between the molecules. The probability of energy transfer between molecules increases as the overlap between the molecule's absorption and emission spectra increase. The absorption and emission spectra overlap at some level for all molecules. Therefore, if there is only one type of molecule in the scintillator cocktail, the light output would be reduced due to inefficiencies in the energy transfer through multiple absorption and reemission processes. Aromatic solutes or fluors are added to the primary solvent to shift the wavelength of the photons to longer wavelengths where the scintillator is more transparent. This wavelength-shifting is also used to match the quantum efficiency as a function of wavelength for the photodetectors being used. One typical scintillator mixture uses pseudo-cumene as the solvent with 1-5 g/L of PPO as the fluor. This mixture has a peak emission at about 400 nm where photomultiplier tubes (PMTs) are most sensitive and the pseudo-cumene is relatively transparent.

A good liquid scintillator will produce $\sim 10,000$ photons isotropically per MeV of deposited energy. Although less abundant, if a particle is moving faster than the speed of light in the medium then Cerenkov light will be produced as well. This light is emitted in a cone pointed in the direction of the particles trajectory and with a continuous spectrum wavelengths weighted toward shorter wavelengths but extending well into the red,

$$\frac{dN}{d\lambda dx} = \frac{2\pi\alpha Z^2}{\lambda^2} \left(1 - \frac{1}{\beta^2 n(\lambda)^2}\right) \quad (1)$$

where $n(\lambda)$ is the wavelength-dependent index of refraction and β is the velocity of the incoming particle. The Cerenkov light produced at wavelengths shorter than the absorption cutoff of the scintillator will be absorbed and re-emitted as isotropic light, but wavelengths longer than this cutoff will propagate across the detector retaining their directional information, roughly 60 photons per MeV assuming a 400 nm absorption cutoff. These undisturbed Cerenkov photons will have timing determined by the group velocity in the liquid,

$$v_g(\lambda) = \frac{c_{vacuum}}{n(\lambda) + dn(\lambda)/d\log(\lambda)}. \quad (2)$$

These longer wavelengths Cerenkov photons arrive in principle before the scintillation light which is slowed by both the scintillation processes and the shorter wavelengths involved. Thus, with sufficient timing resolution and sensitivity to longer wavelength, it should be possible to separate the directional Cerenkov light and the isotropic scintillation light in time and then reconstruct the direction of the initial particle.

III. GEANT4 SIMULATION

In order to study the effects relevant to directionality reconstruction in liquid scintillators, a Geant4 [?] simulation has been set up. The presented simulation application has been developed using standard Geant4 functionality for the optical model of the liquid scintillator. Version Geant4.9.6 is used and the simulation application has been built from a customized official application examples. The simulation is kept simple to provide generally applicable information about the main factors in directionality reconstruction.

The implemented detector geometry is a sphere of 6.5 m (or 0.65 m) diameter filled with liquid scintillator. The scintillator properties have been chosen to match the KamLAND scintillator as the default simulation settings [?]: 80 % n-Dodecane, 20 % Pseudocumene (1,2,4-Trimethylbenzene) and 1.52 g/l PPO (2,5-Diphenyloxazole). The implemented scintillator properties include the atomic composition and density ($\rho = 0.7752$ g/ml), the wavelength-dependent attenuation length and refractive index, the scintillation emission spectrum, emission rise time ($\tau_r = 1.0$ ns) and emission decay time constants ($\tau_{d1} = 6.9$ ns and $\tau_{d2} = 8.8$ ns with relative weights of 0.87 and 0.13), scintillator light yield (LY, 9030.5 photons/MeV) and the Birks constant ($kB = 0.106$ mm/MeV)

[?]. Variations from the baseline KamLAND case are discussed below when applicable. The sphere itself is the photodetector and the following detector properties have been varied: Transit time spread (TTS, default $\sigma = 0.1$ ns) and wavelength-dependent quantum efficiency (QE) for photoelectron production. The default is the bialkali photocathode of the Double Chooz photomultiplier tubes (PMTs) [?] which has been verified to be the same type as the KamLAND photocathode. We used the Double Chooz QE because the data was available with finer spacing in wavelength. The sphere surface is treated as fully absorbing (no reflections) with a coverage of 100 %. Reemission of absorbed photons in the scintillator bulk volume and scattering has not been included. Reflections, reemission and scattering are expected to be corrections to the main effects studied in this paper and are subject of future work.

In sections ?? to ??, we effectively include vertex reconstruction uncertainties. To this end, we use the KamLAND vertex resolution of 12.0 cm [?] (σ in one dimension) to randomly draw a vertex around the known true vertex for each event. This vertex is subsequently used as a reconstructed vertex and consequently, a time of flight (TOF) correction relative to the center of the sphere is applied for each photon hit depending on the distance between the reconstructed vertex and the hit position on the sphere. For the TOF correction a single effective speed of light is used. This effective speed was extracted at the mean wavelength of the photons which create PEs (408 nm). The 12.0 cm vertex resolution corresponds to a time resolution broadening of about 0.6 ns (σ). Note that the vertex resolution itself is dependent on scintillator and detector properties. For example, the PMTs used in KamLAND have a TTS of 1.28 ns (σ) and significant improvement of the vertex resolution could be possible with faster photodetectors. In section ??, we discuss first results on more realistic reconstruction.

Before we discuss the simulation results for different simulation settings in the following sections, we highlight the effects which contribute to the timing of the scintillator detector system. First, the simulated travel time of the initial 5 MeV electron is between 0.10 and 0.15 ns, while the travel distance is about 3 cm. Second, the scintillation light emission follows a scintillator-specific distribution characterized by rise and decay time(s). Past neutrino experiments were not as interested in the effect of scintillation rise time which is the reason why there is a lack of accurate numbers. Before the solutes in liquid scintillator can emit optical photons, the energy has to be transferred from the solvent to the solute. The time constant of this energy transfer accounts for a rise time in scintillation. We assume a rise time of 1 ns, more detailed studies are needed in the future. The two time constants used to describe the falling edge of the scintillator emission time distribution (quoted above) are values specific to the KamLAND scintillator. Third, chromatic dispersion turns out to be an important effect in a 6.5 m diameter detector. Due to the wavelength-dependence of the refractive index the speed of light in the scintillator (the group velocity is used in Geant4 for normal dispersion) is also wavelength dependent. In order to study the size of this effect, we extracted results from a simplified sim-

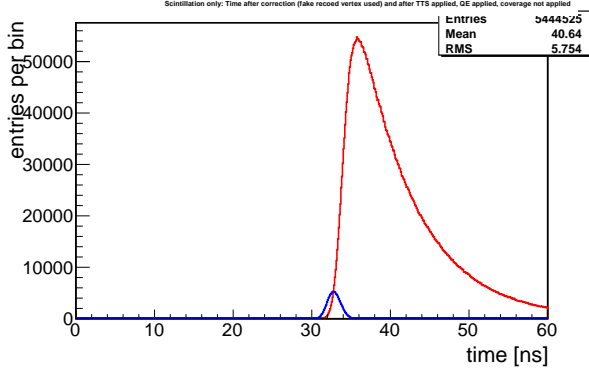


FIG. 1: Effectively reconstructed PE times for the simulation of 1000 electrons (5 MeV) with default settings: Detector diameter = 6.5 m, bialkali photocathode, KamLAND scintillator emission spectrum, TTS = 0.1 ns (σ). PEs from Cerenkov light only (blue) and scintillation light only (red) are compared. The number of PEs per event after a 33.0 ns time cut is 37 from scintillation and 64 from Cerenkov.

ulation of 5 MeV electrons at the center of the sphere where we used instantaneous scintillation and did apply neither TTS nor TOF correction. The true time distribution of photoelectrons were analyzed for scintillation light and Cerenkov light separately. Photoelectrons coming from Cerenkov light are created about 0.5 ns earlier than PEs from scintillation light on average. The RMS values for both the Cerenkov and scintillation light true PE time distributions are both about 0.5 ns. Note that these numbers include the effect of the finite electron travel time. On the detector side, the time information of single photoelectrons is affected by the TTS of the photodetectors, a number which can be different by orders of magnitude depending on the detector type. The default TTS of 0.1 ns (σ) is a value which can be achieved with LAPPDs [?]. In fact, even significantly lower TTS numbers are realistic with this novel technology. Finally, the accuracy of the vertex reconstruction is reflected in broadening and distortions of the time spectrum (after the TOF correction has been applied). Another effect which is to be addressed in future work is the granularity of the photodetectors which adds to the uncertainty in the light path and thus the TOF correction.

IV. DETECTOR TIMING

The settings of the default simulation are described in the previous section. Figure ?? shows the TOF corrected photoelectron detection time for 1000 simulated electrons with 5 MeV energy in the center of the detector with initial momentum direction coinciding with the x axis. The photoelectrons induced by Cerenkov light arrive earlier, as expected due to the instantaneous emission and the differences in the photon speed. There is however overlap with the PE time distribution coming from scintillation light. In order to compare simulations with different parameters to each other a fixed time cut

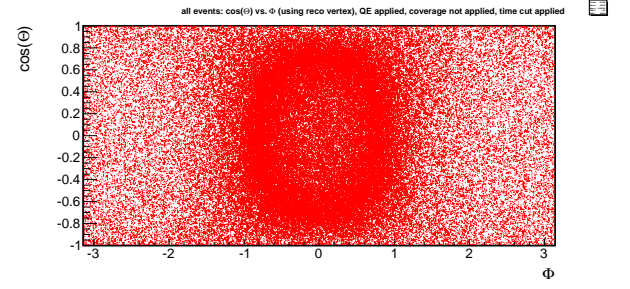


FIG. 2: Photoelectron hit positions for 1000 events (5 MeV electrons in x direction) in the default simulation after the effective position reconstruction effect was included and the 33.0 ns time cut has been applied. The azimuthal angle Φ is zero in x direction and $\pi/2$ in y direction, while $\cos(\Theta)$ ranges from -1 (-z direction) to 1 (z direction).

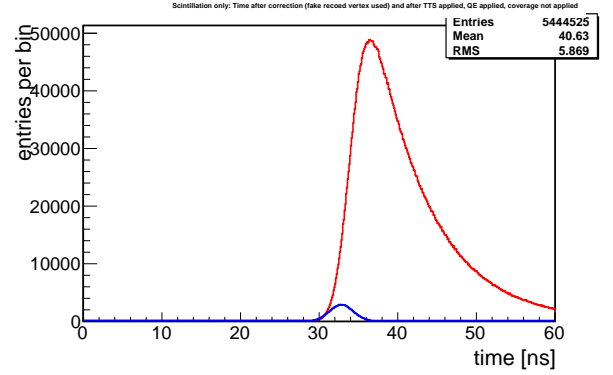


FIG. 3: Effectively reconstructed PE times for the simulation of 1000 electrons (5 MeV) with TTS = 1.28 ns: Detector diameter = 6.5 m, bialkali photocathode, KamLAND scintillator emission spectrum, TTS = 1.28 ns (σ). PEs from Cerenkov light only (blue) and scintillation light only (red) are compared. The number of PEs per event after a 33.0 ns time cut is 153 from scintillation and 60 from Cerenkov.

$t \leq 33.0$ ns is done (using the truth information of the electron starting time) instead of a more realistic event-by-event time reconstruction. For the default simulation case, the number of PEs coming from Cerenkov light after the time cut (64) is about 58 % of the total number of PEs from Cerenkov light (110). For scintillation the number of PEs after the time cut (37) is only 0.68 % (5445).

Figure ?? displays the PE hit pattern after the 33.0 ns time cut has been applied (1000 events plotted together). Although the time cut is an oversimplification of actual time reconstruction effects, we can use it to indicate the spatial distribution of hits after timing information has been used to separate Cerenkov and scintillation light. The Cerenkov ring structure can be clearly seen (opening angle between 40° and 50°) which demonstrates that the directional signal conveyed by the Cerenkov photons is not erased by scattering of the initial 5 MeV electrons.

If the 17 inch KamLAND PMTs [?] (TTS = 1.28 ns) are

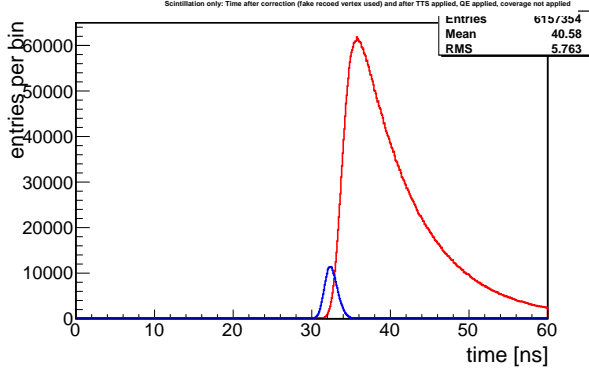


FIG. 4: Effectively reconstructed PE times for the simulation of 1000 electrons (5 MeV) with red-sensitive photocathode: Detector diameter = 6.5 m, red-sensitive GaAsP photocathode, KamLAND scintillator emission spectrum, TTS = 0.1 ns (σ). PEs from Cerenkov light only (blue) and scintillation light only (red) are compared. The number of PEs per event after a 33.0 ns time cut is 50.8 from scintillation and 171 from Cerenkov.

used in the simulation, the broadening of the time distributions leads to a strongly decreased ratio of Cerenkov over Scintillation light after the time cut (see FIG. ??). This demonstrates that the TTS of the photodetectors is critical for directionality reconstruction. Note that the same KamLAND vertex resolution of 12.0 cm has been used in both cases. For detectors with TTS = 0.1 ns (σ) this number is pessimistic since faster detectors can improve the vertex resolution accuracy.

V. DETECTOR WAVELENGTH RESPONSE

In the previous section we saw that low photodetector TTS is highly beneficial for discrimination between Cerenkov and scintillation light. The second important photodetector parameter is the wavelength-dependent QE. Since Cerenkov photons which passed 6.5 m of scintillator have higher average wavelengths than scintillation photons, a photodetector which is more sensitive at high wavelengths increases not only the absolute number of PEs but also the ratio between Cerenkov- and scintillation-induced PEs. Therefore, a simulation has been run with the QE of an extended red-sensitive GaAsP photocathode (the data has been digitized using Hamamatsu R3809U-63 QE data).

Figure ?? shows the results for the modified simulation with high QE in the red spectral region. The higher absolute number of PE coming from Cerenkov light and the increased Cerenkov/scintillation ratio after the time cut (almost doubled compared to the default case) would significantly enhance the directionality reconstruction capabilities.

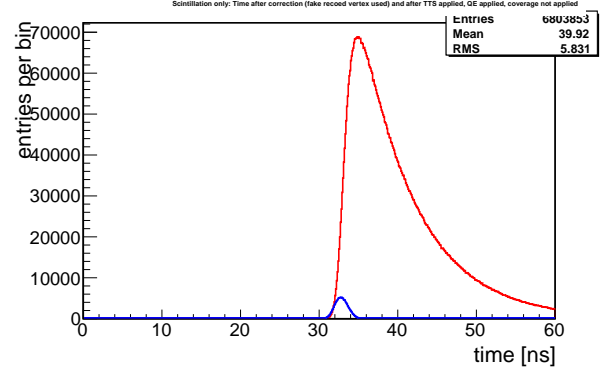


FIG. 5: Effectively reconstructed PE times for the simulation of 1000 electrons (5 MeV) with QD emission spectrum at 461 nm: Detector diameter = 6.5 m, bialkali photocathode, QD emission spectrum at 461 nm, TTS = 0.1 ns (σ). PEs from Cerenkov light only (blue) and scintillation light only (red) are compared. The number of PEs per event after a 33.0 ns time cut is 180 from scintillation and 64 from Cerenkov.

VI. SCINTILLATOR EMISSION SPECTRUM

One important contributing factor to the separation in time between Cerenkov and scintillation photon hits is the higher average light speed for the photons coming from the Cerenkov effect due to their higher average wavelength. In this section we present two more sets of simulations where the scintillator emission spectrum was changed.

Recently, the use of quantum dots (QDs) in liquid scintillators has been studied as a possibility to improve future large scale neutrino experiments [?]. One important motivation is the control of the emission spectra by tuning the size or composition of the quantum dots. The emission spectrum of alloyed core/shell $\text{CdS}_x\text{Se}_{1-x}/\text{ZnS}$ Trilite450 [?] quantum dots was measured. This spectrum is a symmetric peak centered around 461 nm with FWHM = 29 nm.

In order to isolate the effect of a different emission spectrum, the other simulation settings, including the KamLAND absorption spectrum are kept unchanged. In FIG. ?? the result for this simulation is presented. Compared to the default case shown in FIG. ?? the separation is worse because the scintillation light wavelengths are higher than in the KamLAND emission spectrum. However, advances in the production of commercial quantum dot samples could yield quantum dots which have similar, single peak emission shapes at lower wavelengths. This case has been simulated using the same spectral shape of the measured Trilite450 emission but shifted to lower wavelengths such that the emission peak is centered at 384 nm. This peak emission value has been measured for other types of QDs, however with a much more pronounced tail [?]. The resulting PE time distribution shows (as expected) a clearer separation of Cerenkov and Scintillation light. After the simple 33.0 ns time cut we obtain slightly better numbers compared to the default simulation: The number of Cerenkov-induced PE after the time cut is unchanged

(64) while the number of PEs coming from scintillation light is decreased to 25 (compared to 37 for the default simulation).

VII. ENERGY DEPENDENCE AND DETECTOR SIZE

include this?

The number of Cerenkov photons decreases disproportionately with decreasing electron energy. The number of PEs after the 33 ns time cut for 1 MeV electrons is 8.8 PE from scintillation and 7.7 PE from Cerenkov light. This illustrates that directionality reconstruction benefits from higher initial energies but even at 1 MeV there might be accessible directionality information.

An additional set of simulations of a 0.65 m diameter ($\approx 1 \text{ m}^3$) detector with otherwise default parameters was performed. In the smaller detector the chromatic dispersion is not as important and absorption in the scintillator bulk is reduced. The latter effect increases the absolute number of photoelectrons. The former effect, on one hand, reduces the separation in time between Cerenkov and Scintillation light. On the other hand, it reduces the broadening of both time distributions which helps more sophisticated vertex and track reconstruction with the help of fast photodetectors like for example the LAPPDs. However, with the simplified effective vertex reconstruction and a time cut of 3.265 ns we get 87 PEs from scintillation and 87 PEs from Cerenkov. The time cut was adjusted such that the fraction of selected PEs from Cerenkov light matches the fraction in the large detector ($\approx 58\%$). The

QD spectrum discussed in section ?? (both shifted and unshifted) did not change the PE time distribution significantly. We also simulated the red-sensitive photocathode discussed in section ?? in the small detector. The amount of PE increases again significantly and with the 3.265 ns time cut the number of PE coming from Cerenkov light is 182 while the number of PE from scintillation light is 96.

In summary, the photodetector properties (TTS and QE) become more important relative to the scintillator emission and absorption spectra when the detector size is decreased from 6.5 m to 0.65 m.

VIII. RECONSTRUCTION

Hopefully results

IX. CONCLUSION

This is going to work.

X. ACKNOWLEDGEMENTS

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