Measurement Of 8b Solar Neutrinos In The Sno+ Water Phase And A Model Of Vacuum-Enhanced Neutrino Mixing

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Measurement Of $^{8}$B Solar Neutrinos In The SNO+ Water Phase And A Model Of Vacuum-Enhanced Neutrino Mixing

Abstract

This thesis describes two separate results in the study of solar neutrinos. The first is a measurement of the $^{8}$B solar neutrino flux using a 69.2 kt·day dataset acquired with the SNO+ detector during its water commissioning phase. Using an energy range of 5 MeV to 15 MeV, fits of the solar neutrino event direction yielded an observed flux of $2.53^{+0.31}_{-0.28}$(stat.)$^{+0.13}_{-0.10}$(syst.)$\times10^6$ cm$^{-2}$s$^{-1}$, assuming no neutrino oscillations. This flux is consistent with standard matter enhanced neutrino oscillations and measurements done by other experiments. At energies above 6 MeV the dataset is an extremely pure sample of solar neutrino elastic scattering events, allowing an accurate measurement with relatively little exposure. In that energy region the best fit background rate is $0.25^{+0.09}_{-0.07}$ events/kt·day, significantly lower than the measured solar neutrino event rate in that energy range, which is $1.03^{+0.13}_{-0.12}$ events/kt·day.

The second topic is the development of a novel model for neutrino mixing that allows for a potential that couples to neutrino flavor and is only significant in areas of near zero matter density. A neutrino mixing simulation of this model was developed and is described. Special consideration is given to a previously observed tension in the measured value for $\Delta m^{2}_{21}$ by the KamLAND reactor experiment and a combined solar neutrino experiment measurement. Performing a fit to solar neutrino and reactor neutrino data it’s determined that this new neutrino mixing model reduces the tension between solar neutrino measurements and the KamLAND measurement from $\Delta \chi^{2}=4.2$ to $\Delta \chi^{2}=0.85$ with two new degrees of freedom. Providing a modest preference for this new model over standard neutrino mixing. Finally, future improvements and generalizations to this result are discussed.

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MEASUREMENT OF $^8$B SOLAR NEUTRINOS IN THE SNO+ WATER PHASE AND A MODEL OF VACUUM-ENHANCED NEUTRINO MIXING

Eric Marzec

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MEASUREMENT OF $^8$B SOLAR NEUTRINOS IN THE SNO+ WATER PHASE AND A MODEL OF VACUUM-ENHANCED NEUTRINO MIXING

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Eric Marzec

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ABSTRACT

MEASUREMENT OF $^8$B SOLAR NEUTRINOS IN THE SNO+ WATER PHASE AND A MODEL OF VACUUM-ENHANCED NEUTRINO MIXING

Eric Marzec
Joshua R. Klein

This thesis describes two separate results in the study of solar neutrinos. The first is a measurement of the $^8$B solar neutrino flux using a 69.2 kt-day dataset acquired with the SNO+ detector during its water commissioning phase. At energies above 6 MeV the dataset is an extremely pure sample of solar neutrino elastic scattering events, owing primarily to the detector’s deep location, allowing an accurate measurement with relatively little exposure. In that energy region the best fit background rate is $0.25^{+0.09}_{-0.07}$ events/kt-day, significantly lower than the measured solar neutrino event rate in that energy range, which is $1.03^{+0.13}_{-0.12}$ events/kt-day. Also using data below this threshold, down to 5 MeV, fits of the solar neutrino event direction yielded an observed flux of $2.53^{+0.31(\text{stat.})+0.13(\text{syst.})}_{-0.28} \times 10^6$ cm$^{-2}$s$^{-1}$, assuming no neutrino oscillations. This rate is consistent with matter enhanced neutrino oscillations and measurements from other experiments.

The second topic is the development of a novel model for neutrino mixing that allows for a potential that couples to neutrino flavor and is only significant in areas of near zero matter density. A neutrino mixing simulation of this model was developed and is described. Special consideration is given to a previously observed tension in the measured value for $\Delta m^2_{21}$ by the KamLAND reactor experiment and a combined solar neutrino experiment measurement. Performing a fit to solar neutrino and reactor neutrino data it’s determined that this new neutrino mixing model reduces the tension between solar neutrino measurements and the KamLAND measurement from $\Delta \chi^2 = 4.2$ to $\Delta \chi^2 = .85$ with two new degrees of freedom. Providing a modest preference for this new model over standard neutrino mixing. Finally, future improvements and generalizations to this result are discussed.
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Chapter 1

Introduction

This thesis discusses two separate results on the topic of solar neutrinos. The first topic describes a measurement of the $^8$B solar neutrino flux with the initial SNO+ water phase dataset. The measured value for the flux is consistent with, though less precise, similar measurements done by other solar neutrino experiments. Exceptionally low backgrounds were observed in the higher energy region of the analysis, this provides the purest sample of solar neutrino elastic scatter events so far detected. The precision of this measurement is ultimately limited by the available statistics of the dataset, and so can be improved with further data taking.

The second topic is the description, development, and evaluation of a novel model for neutrino mixing that proposes neutrinos might mix differently in areas of near zero matter density. Part of the motivation for this model is the fact that solar neutrino experiments have not yet observed an expected rise in the solar neutrino survival probability at low energies. This thesis considers if a proposed model for a neutrino flavor coupled potential that is only significant in the vacuum of space could provide an explanation for the unexpectedly flat solar neutrino survival probability; also considered is how much preference existing measurements show for this model.
1.1 Neutrino History

This chapter provides an introduction and overview of the physics topics relevant to solar neutrinos, neutrino detection, and neutrino oscillation. Following that begins discussion of the SNO+ $^8\text{B}$ solar neutrino flux measurement. Chapter 2 provides an overview of topics relevant to the measurement, providing a basic introduction to the relevant topics that will be discussed in greater detail in the following sections. Chapter 3 provides description of the SNO+ detector and DAQ, highlighting some of my own contributions to trigger and DAQ system. In Chapter 4 the simulation methods used for the analysis are described. The techniques used for reconstructing event vertex information from detector observables is discussed in Chapter 5. The calibration sources used as for evaluating systematics are discussed in Chapter 6. Chapter 7 describes the cuts placed on detected and simulated data to remove backgrounds. The analysis and relevant systematics are described in Chapter 8. And finally the fit to data and results of the fit are given in Chapter 9 also discussed are possible future improvements to this measurement and future solar neutrino measurements for SNO+.

Following the description of the $^8\text{B}$ measurement is the description of the vacuum enhanced neutrino mixing model that I developed. First Chapter 10 provides a technical description of the model, along with motivations for the model. Next Chapter 11 describes existing measurements of $\Delta m^2_{21}$ and their relevance to solar neutrino measurements. Simulation of the vacuum-enhanced mixing model is described in Chapter 12. The data used to constrain this model is described in Ch. 13 and finally the results of fitting this model to the data is given in Chapter 14.

1.1 Neutrino History

Neutrinos were first hypothesized by Wolfgang Pauli in 1930. The motivation for the proposal was the apparent violation of energy conservation in $\beta$ decay. Several years after Pauli’s speculative proposal Enrico Fermi offered a thorough model of beta decay that
1.1 Neutrino History

conserved energy using the neutrino \(\nu\). Nearly two decades after its initial proposal, Frederick Reines & Clyde Cowan performed an experiment that involved bombarding a tank of cadmium-doped water with anti-neutrinos from nuclear reactor \(\nu\). Their detector was able to observe the rate and energy of inverse \(\beta\) decays that occurred. The experiment’s results were consistent with Fermi’s model of \(\beta\) decay and were considered experimental confirmation of the neutrino’s existence.

The first experimental evidence for neutrino flavor came in 1962 from an experiment \(\nu\) that studied the interactions of neutrinos from muon decay and from beta decay. It was observed that neutrinos that came from muon decay would only produce muons upon interacting in a detector, and neutrinos produced from \(\beta\) decay would only create electrons in the detector. This led to the conclusion that there are two different flavors of neutrino, the \(\nu_e\) and the \(\nu_\mu\), and the idea that lepton flavor is conserved in weak interactions. The third lepton generation, the \(\tau\), was discovered 13 years later in 1975 \(\nu\). The accompanying neutrino, the \(\nu_\tau\) was discovered 25 years later by the DONUT collaboration \(\nu\).

![Figure 1.1: The evolution in the predicted and detectable flux of neutrinos from the sun. Figure from Ref. (10).](image)

Modern modelling of the Sun and stars could be said to have begun in 1920 with Arthur
Eddington’s proposal that nuclear reactions fuel stellar burning (11). Bethe’s 1938 calculations of solar energy production (12, 13) followed Eddington’s proposal and produced the first model for stellar burning and neutrino production within stars. This model was developed over the years as knowledge of nuclear interaction improved, eventually resulting in Bahcall’s comprehensive summary of and prediction for the Sun’s rates of neutrino production (14). Simultaneous with improvements in solar and nuclear modeling, particle detection techniques had developed and the first solar neutrino detector, the Homestake experiment, began operations in 1967 (15). The Homestake experiment provided the first experimental measurement of solar neutrinos and measured a flux that was approximately one-third what was expected (16). Figure 1.1 from Bahcall and Davis (10), depicts the evolution of solar neutrino predictions and the limits of experimental observation.

The persistent deficit observed by the Homestake experiment eventually became known as the “Solar Neutrino Problem”. It was the first evidence that the flavor of neutrinos oscillates with time, and that therefore neutrinos must have mass. Conclusive evidence for neutrino oscillation came from atmospheric neutrino measurements by the Super-Kamiokande and Irvine-Michigan-Brookhaven (IMB) experiments (17, 18). And the Solar Neutrino Problem was conclusively resolved by flavor independent solar neutrino measurement done by the Sudbury Neutrino Observatory (SNO) (19, 20, 21). More will be said about the Super-Kamiokande and SNO experiments in Sec. 1.6.

1.2 Neutrino Interactions

The neutrino interacts almost exclusively via the weak interaction. In principle neutrinos also interact gravitationally, and they have a non-vanishing magnetic moment so can interact electromagnetically, but these interaction potentials are so small they can be neglected in all practical cases (22). The weak interaction has a number of aspects that limit the sort of neutrino interactions that can occur. One aspect is lepton flavor conservation; all weak interactions conserve both the total lepton number of a system and also the total lepton...
flavor of the system as well. This leads to nearly all interactions involving a neutrino also involving the same flavor charged lepton. The second aspect is that the weak interaction is chiral, only left-handed particles and right-handed anti-particles interact weakly. Since neutrinos only interact weakly, the only detectable varieties of neutrino is the left-handed neutrino $\nu^L$ and the right-handed anti-neutrino $\bar{\nu}^R$.

Discussed here are the neutrino interactions that occur in the energy regime relevant for solar neutrinos. At higher energies, $E_\nu \gtrsim 100$ MeV, neutrinos can interact with nuclear targets through quasi-elastic scattering, resonant pion-production, or deep inelastic scattering. A more complete overview of neutrino interactions is available in Ref. (24).

![Feynman Diagrams](image)

**Figure 1.2:** Charged-current (a) and neutral-current (b) neutrino nuclear interactions.

Significant at lower energies is the charged-current (CC) nuclear interaction $\nu_e + n \rightarrow p + e^-$. Figure 1.2a gives the Feynman diagram for this process. The charged-current nuclear interaction played a significant role in early radio-chemical solar neutrino experiments (16, 25, 26, 27). SNO made use of the charged-current nuclear interaction and the neutral current (NC) interaction (Fig 1.2b). SNO and its measurements will be discussed in greater detail in Sec 1.6.1.

Neutrino-electron elastic scattering (ES) is an important neutrino interaction channel for this work — it is the main interaction through which neutrinos are detected in the
1.2 Neutrino Interactions

(a) Charged-current elastic scattering interaction  
(b) Neutral-current elastic scattering interaction

Figure 1.3: The Feynman diagrams for the charged-current elastic scattering interaction (a) and the neutral current elastic scattering interaction (b).

measurement discussed in Part I. The ES process is $\nu_x + e^- \rightarrow \nu_x + e^-$ or $\bar{\nu}_x + e^- \rightarrow \bar{\nu}_x + e^-$. The tree level Feynman diagrams for the charged-current neutrino elastic scattering interaction is shown in Fig. 1.3(a) and the neutral-current elastic scattering interaction in Fig. 1.3(b). Neutral-current interactions involve $\nu_x$ where $x = e, \mu, \tau$, and the charged-current interaction involves only $\nu_e$.

In principle the charged-current interaction is available to neutrinos of all flavors, however, for solar neutrinos only the electron flavor version of the interaction is available. The initial electron on the left-hand side of the interaction is usually understood to be from an atom within detector target, and therefore at rest within the lab frame. So the total energy in the interaction is simply $E = E_\nu + m_e$. For even the highest energy solar neutrinos $E < 20 \text{ MeV}$, far less than muon rest mass of $m_\mu = 105.7 \text{ MeV}/c^2$ or tau mass $m_\tau = 1776.8 \text{ MeV}/c^2$. Meaning a muon or tau cannot be created; the electron with a rest mass of $m_e = 0.511 \text{ MeV}/c^2$ is the only charged lepton that can be created from the CC-ES interaction for solar neutrinos. And since lepton flavor is conserved in weak interactions, the electron neutrino is the only neutrino that can produce an electron, therefore the electron neutrino is the only neutrino flavor that can undergo the charged-current elastic scattering
interaction.

This flavor disparity in the interaction means that the cross-section for the elastic scattering process is larger for $\nu_e$ than it is for a $\nu_\mu$ or $\nu_\tau$, and the cross-section for a $\nu_\mu$ is the same as that for a target $\nu_\tau$. The differential cross section for the diagrams shown in Fig 1.3a and Fig. 1.3b can be calculated as

$$
\frac{d\sigma}{dT_e}(E_\nu, T_e) = \frac{\sigma_0}{m_e} \left[ g_1^2 + g_2^2 \left( 1 - \frac{T_e}{E_\nu} \right)^2 - g_1 g_2 \frac{m_e T_e}{E_\nu^2} \right],
$$

where,

$$
\sigma_0 = \frac{2G_F^2 m_e^2}{\pi}.
$$

For $\nu_x = \nu_e$

$$
g_1 = \frac{1}{2} + \sin^2 \theta_W,
$$

and

$$
g_2 = \sin^2 \theta_W.
$$

For $\nu_x = \nu_\mu$ or $\nu_x = \nu_\tau$ $g_1$ and $g_2$ are given by,

$$
g_1 = \sin^2 \theta_W - \frac{1}{2},
$$

$$
g_2 = \sin^2 \theta_W.
$$

Since the electron scattering is elastic, the kinematics of the interaction leave only one free parameter, the outgoing electron direction with respect to the incoming neutrino direction, $\theta$. For a recoil electron with a given value for $\theta$ the electron kinetic energy is given by

$$
T_e = \frac{2m_e E_\nu^2 \cos^2 \theta}{(m_e + E_\nu)^2 - E_\nu^2 \cos^2 \theta},
$$

7
This relationship can be used to produce the differential cross-section in $\cos \theta$,

$$
\frac{d\sigma}{d\cos \theta} = \sigma_0 \frac{4E_\nu^2(m_e + E_\nu^2 \cos^2 \theta)^2 \cos \theta}{[(m_e + E_\nu^2 - E_\nu^2 \cos^2 \theta)]^2}
$$

$$
\left[ g_1^2 + g_2^2 \left( 1 - \frac{2m_eE_\nu \cos^2 \theta}{(m_e + E_\nu)^2 - E_\nu^2 \cos^2 \theta} \right)^2 - g_1g_2 \frac{2m_e \cos^2 \theta}{(m_e + E_\nu)^2 - E_\nu^2 \cos^2 \theta} \right]. \quad (1.8)
$$

And finally the maximum energy a recoil electron will have can be found by setting $\cos \theta = 1$ in equation (1.7), this yields

$$
T_{e}^{\text{max}}(E_\nu) = \frac{2E_\nu^2}{m_e + 2E_\nu^2}. \quad (1.9)
$$

These results are derived from first principles in Ref. (29).

Figure 1.4 shows the differential cross-sections for a 10 MeV neutrino. The fact that the scattering cross-section is peaked near $\cos \theta = 1$ is useful for neutrino experiments; the observed electron direction will almost always be nearly co-linear with the incoming neutrino direction. Unfortunately, since the cross-section is nearly flat in $T_e$ the electron energy provides almost no information about the incoming neutrino energy. The kinematics and cross-section of the out-going neutrino is also well predicted with respect to $\cos \theta$, however the out-going neutrino is typically of little interest because it cannot be detected.

Experimentally, neutrino elastic scattering has been studied in a number of venues historically and recently (30, 31, 32). Famously the first experimental evidence for the $Z$ boson came from the observations of neutrino elastic scattering at the Gargamelle experiment (33). It’s also been a useful detection method for water-cherenkov solar neutrino experiments because the directionality of the interaction allows events that originate from the Sun to be identified (19, 34, 35).

1.3 Neutrino Oscillations

Neutrino oscillation is a phenomenon of massive neutrinos, and its observation is currently the only experimental evidence that neutrinos do indeed have mass. Neutrino oscillation
1.3 Neutrino Oscillations

Figure 1.4: The electron recoil (a) cross-section and electron angular (b) cross-section for neutrino-electron elastic scattering with a 10 MeV incoming neutrino energy.
results from the fact that neutrino flavors do not have well-defined masses, instead neutrino flavor states are a quantum superposition of neutrino mass states. And conversely, mass states can be described as a superposition of flavor states. Stated more precisely,

\[ |\nu_i\rangle = U_{i\ell} |\nu_\ell\rangle \quad (1.10) \]

Where \( |\nu_\ell\rangle \) represents the neutrino flavor states, \( |\nu_i\rangle \) represents the mass states, and \( U_{i\ell} \) describes the mixing of these states. \( U \) is known as the Pontecorvo-Maki-Nakagawa-Sakata (PMNS) matrix, and it is analogous to the Cabibbo-Kobayashi-Maskawa (CKM) matrix used to describe quark mixing. In the simplest case where the weak states and the mass states are the same \( U \) would just be the identity matrix. Under the assumption that there are three flavor states and three mass states \( U_{i\ell} \) must be unitary so that the probability of observing a neutrino in any state is never less than one. It is known from observations of Z boson decay products that there are only three “active” neutrino flavors \( (36) \). Active meaning that the neutrino participates in weak interactions. It is possible that there is a neutrino that does not interact weakly, this is referred to as a sterile neutrino and is the subject of a number of experimental searches \( (37, 38, 39, 40) \). If there were a sterile neutrino state, then the \( 3 \times 3 \) PMNS matrix would no longer be unitary.

It is typical to characterize \( U \) with three angles \( (\theta_{12}, \theta_{13}, \theta_{23}) \) and a CP violating phase \( \delta_{cp} \). Doing so allows for the general SU(3) mixing matrix to be decomposed into two SO(2) matrices and one SU(2) matrix,

\[
U_{12} = \begin{pmatrix} \cos \theta_{12} & \sin \theta_{12} & 0 \\ -\sin \theta_{12} & \cos \theta_{12} & 0 \\ 0 & 0 & 0 \end{pmatrix}, \\
U_{13} = \begin{pmatrix} \cos \theta_{13} & 0 & \sin \theta_{13} e^{-i\delta_{cp}} \\ 0 & 0 & 0 \\ -\sin \theta_{13} e^{-i\delta_{cp}} & 0 & \cos \theta_{13} \end{pmatrix}, \\
U_{23} = \begin{pmatrix} 0 & 0 & 0 \\ 0 & \cos \theta_{23} & \sin \theta_{23} \\ 0 & -\sin \theta_{23} & \cos \theta_{23} \end{pmatrix}.
\]
1.3 Neutrino Oscillations

The product of these matrices produces the full mixing matrix, $U = U_{23}U_{13}U_{12}$.

The mixed nature of neutrino flavor and mass states gives rise to oscillations in the flavor content of propagating neutrinos. By definition the neutrino mass states are eigenstates of the vacuum Hamiltonian

$$H |\nu_i\rangle = E_i |\nu_i\rangle,$$

(1.11)

where the energy is given by the standard relativistic energy equation,

$$E_i = \sqrt{p_i^2 + m_i^2}.$$  

(1.12)

It is typical to make the assumption that all neutrino mass states have the same momentum, $p_i = p$. The equal momentum assumption allows for a straightforward derivation of the correct description of neutrino oscillations, but it is not well motivated. A discussion of the equal momentum assumption and its validity is available in Ref. (41).

Applying Schrödinger’s equation,

$$i \frac{d}{dt} |\nu_i(t)\rangle = H |\nu_i(t)\rangle = E_i |\nu_i(t)\rangle,$$

(1.13)

results in,

$$|\nu_i(t)\rangle = e^{-iE_i t} |\nu_i(t = 0)\rangle.$$  

(1.14)

This time evolution of mass eigenstates can be used to then describe the state of a neutrino that is created in an electron flavor eigenstate.

$$|\nu(t = 0)\rangle = |\nu_e\rangle = U_{1e} |\nu_1\rangle + U_{2e} |\nu_2\rangle + U_{3e} |\nu_3\rangle = \sum_{i=1}^{3} U_{ie} |\nu_i\rangle$$

(1.15)

Each of the mass states’ time evolution can be immediately written down from Equation (1.14),

$$|\nu(t)\rangle = \sum_{i=1}^{3} U_{ie} e^{-iE_i t} |\nu_i\rangle.$$  

(1.16)
From Eqn. (1.16) the quantity that’s often of the most interest is the survival probability, defined as

\[ P_{ee}(t) = |\langle \nu_e | \nu(t) \rangle|^2. \]  

(1.17)

\( P_{ee}(t) \) can be understood as the probability that a neutrino, produced in an electron flavor state, will be detected as an electron flavor state a time \( t \) later. The survival probability for the state given in Eqn. (1.16) is

\[ P_{ee}(t) = \sum_{i,j} |U_{ei}|^2 |U_{ej}|^2 e^{-i(E_i - E_j)t}. \]  

(1.18)

It is useful to separate out the terms where \( i = j \),

\[ P_{ee}(t) = \sum_i |U_{ei}|^4 + \sum_{i,j, j \neq i} |U_{ei}|^2 |U_{ej}|^2 e^{-i(E_i - E_j)t}. \]  

(1.19)

Here we can see there are terms that vary with time, and terms that are constant.

A few more simplifications are commonly done, using the equal momentum assumption mentioned earlier and the fact the neutrino rest mass is very small, the energy differences can be simplified,

\[ E_i - E_j = \sqrt{p^2 + m_i^2} - \sqrt{p^2 + m_j^2} \approx p^2 - p^2 + \frac{m_i^2}{2p} - \frac{m_j^2}{2p}. \]  

(1.20)

To good approximation \( p^2 = E^2 \), so

\[ E_i - E_j = \frac{\Delta m_{ij}^2}{2E}, \]  

(1.21)

where the definition of the mass-squared splittings is used,

\[ \Delta m_{ij}^2 = m_i^2 - m_j^2. \]  

(1.22)

The final simplification is to assume that the neutrino is moving very close to the speed of light, therefore any flavor oscillations over time will also occur at a distance \( L \) from the
neutrino's creation. All this gives

$$P_{ee}(L,E) = \sum_{i} |U_{ei}|^4 + \sum_{i,j, j \neq i} |U_{ei}|^2 |U_{ej}|^2 e^{-i \left( \frac{\Delta m_{ij}^2}{2E} L \right)}.$$  \hspace{1cm} (1.23)$$

For the mixing parameters given in (42) the average survival probability for an electron neutrino is $P_{ee} = 55.8\%$.

It is common to represent neutrino mixing in the following form

$$i \frac{d}{dx} |\Psi_\alpha\rangle = H |\Psi\rangle,$$  \hspace{1cm} (1.24)$$

where

$$H = \frac{1}{2E} U M^2 U^\dagger |\Psi_\alpha\rangle.$$  \hspace{1cm} (1.25)$$

Where $U$ is the PMNS matrix, $|\Psi_\alpha\rangle$ is the neutrino state in the flavor basis, and

$$M^2 = \begin{bmatrix} 0 & 0 & 0 \\ 0 & \Delta m_{21}^2 & 0 \\ 0 & 0 & \Delta m_{31}^2 \end{bmatrix}.$$  \hspace{1cm} (1.26)$$

This form of the mixing equation is the same as Eqn. (1.13) written in the flavor basis.

Except for the CP violating phase $\delta_{cp}$, all mixing parameters and mass-squared splitting have been measured experimentally [1]. None of the parameters are predicted in the standard model, so they are theoretically unconstrained. The measurements of the mass-squared splittings and other parameters are discussed in Sec. 1.6. The absolute mass of each neutrino state is not yet known. There are upper limits placed on the sum of the neutrino masses from observations of tritium decay [43], and from cosmological observations and modelling [44].

The “hierarchy” of the neutrino masses, i.e. which mass states are heavier or lighter than the others is also unknown. Solar neutrino measurements have established that the $\nu_2$ state is more massive than the $\nu_1$ state [19, 45], but so far no experiment has been able to resolve if $\nu_3$ is more or less massive than $\nu_1$ and $\nu_2$ states [46]. The case where the $\nu_1$
state is the lightest is referred to as the “normal hierarchy” because then lightest neutrino mass state is also the most electron flavored, similar to the charged leptons. The case where $\nu_3$ is the least massive neutrino is known as the “inverted hierarchy”. Experimental measurements and cosmological considerations currently prefer the normal hierarchy over the inverted hierarchy at approximately $2.4\sigma$ statistical significance (47).

The model of neutrino mixing as described here assumes that neutrinos are Dirac, as opposed to Majorana. In the case that the neutrino is a Majorana particle the PMNS matrix gains two additional complex “Majorana” phases, similar to the CP violating phase $\delta_{CP}$. However, neutrino oscillation observables, such as the survival and transition probability, are in general not affected by the Majorana phase (48). Therefore, it is accurate to ignore the presence of Majorana phases in typical neutrino mixing calculations regardless of the true Majorana or Dirac nature of the neutrino.

### 1.3.1 Matter-Enhanced Oscillations

When neutrinos propagate through matter, as opposed to vacuum, their oscillation is altered by the potential introduced from available nuclear and leptonic interactions. The interaction comes primarily from a neutral-current interaction of the form shown in Fig. 1.2b, and a flavor-dependent charged-current interaction shown in Fig. 1.2a and the neutrino-electron elastic scattering interaction shown in Fig. 1.3a and Fig. 1.3b. The potential for the nuclear charged-current interaction is negligible because it modifies the state of the nucleon it interacts with, and is thus considered an incoherent nuclear interaction. It is shown in Ref. (49) that incoherent interactions will contribute negligibly to the interaction potential unless the neutrinos are extremely high energy or in regions of extremely high matter density.

The potential associated with the charged-current ES interaction is given by

$$V_{CC} = \sqrt{2}G_F N_e,$$

(1.27)
where $N_e$ is the local electron density. Similarly, the neutral-current interactions contribute a flavor independent shift to the potential given by

$$V_{NC} = -\frac{\sqrt{2}}{2} G_F N_n$$

(1.28)

where $N_n$ is the local neutron density, the contributions to this potential from electron and proton interactions cancel out. This derivation assumes the neutrino is traveling through matter composed primarily of protons, neutrons, and electrons. A safe assumption for almost all cases.

The mixing described by Eqn. (1.24) is modified to include the potential terms described above.

$$i \frac{d}{dt} |\Psi_\alpha\rangle = [H_{vac} + H_I] |\Psi\rangle$$

(1.29)

$$H_{vac} = \frac{1}{2E} U M^2 U^\dagger$$

(1.30)

$$H_I = A_{CC} + A_{NC}.$$ (1.31)

The mixing potentials for the charged and neutral-current interactions are given by

$$A_{CC} = \begin{bmatrix} V_{CC} & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix},$$ (1.32)

$$A_{NC} = \begin{bmatrix} V_{NC} & 0 & 0 \\ 0 & V_{NC} & 0 \\ 0 & 0 & V_{NC} \end{bmatrix}. $$ (1.33)

It can be show that since $A_{NC}$ is identity-like it can be neglected from consideration; identity-like matrices produce no observable effect on the overall neutrino mixing. The full mixing Hamiltonian is given by

$$H = \frac{1}{2E} U M^2 U^\dagger + A_{CC}.$$ (1.34)

These results are derived in detail in Ref. (49) and Ref. (29).
It’s clear that the electron density through which a neutrino propagates can modify the effective mass-splitting and mixing angle for the electron neutrino. That is to say, any matter-enhanced mixing can also be described as vacuum mixing with just a different set of mixing parameters. For a given neutrino energy, $E_{\nu}$, there exists an electron density for which the effective mixing angle is maximal, this is known as resonant density or a Mikheyev–Smirnov–Wolfenstein (MSW) resonance \([49, 50]\). The resonance density is given by

$$N_e = \frac{\Delta m^2 \cos 2\theta}{2\sqrt{2}E_G F}.$$  \hspace{1cm} (1.35)

This resonance is important for solar neutrinos and will be discussed further in the next section.

Matter effects also introduce the question of adiabaticity into neutrino oscillations. As a neutrino travels through changing matter potentials it’s possible for the effective potential to evolve much faster than the neutrino oscillates. A neutrino’s propagation is said to be fully adiabatic if its mass state composition does not change between production and detection. A specific example is if a neutrino state is a pure vacuum mass-1 state,

$$|\Psi_{\nu}\rangle = |\nu_1\rangle$$

suddenly enters a region of significant electron density where the mass states are now $|\nu'_k\rangle$. The neutrino state does not have time to evolve at all so the state does not change, but it is not longer a pure eigenstate of the mixing Hamiltonian,

$$|\Psi_{\nu}\rangle = |\nu_1\rangle = \sum_{k=1}^{3} \langle \nu'_k | \nu_1 \rangle |\nu'_k\rangle.$$

Since the neutrino is no longer in a pure eigenstate the state will now oscillate.

In contrast, if the neutrino enters the region of significant electron density more slowly, then the neutrino state will smoothly evolve with the eigenstate, $|\nu_1\rangle \rightarrow |\nu'_1\rangle$. And so the final state of the neutrino in the adiabatic case will be,

$$|\Psi_{\nu}\rangle = |\nu'_1\rangle.$$
1.4 Solar Neutrinos

Nuclear reactions in the core of the Sun provides the main fuel for stellar burning and prevents the Sun from collapsing due to gravitation pressure. There exists two separate chains of nuclear reactions that are present in typical stellar conditions, the pp-chain and the CNO-cycle \(^{14}\text{(14)}\). Figure \[1.5\] and \[1.6\] shows these two reaction chains. For the Sun the
1.4 Solar Neutrinos

\[ ^{12}\text{C} + p \rightarrow ^{13}\text{N} + \gamma \]
\[ ^{15}\text{N} + p \rightarrow ^{12}\text{C} + ^{4}\text{He} \]
\[ ^{13}\text{N} \rightarrow ^{13}\text{C} + e^+ + \nu_e \ (^{13}\text{N}) \]
\[ ^{15}\text{O} \rightarrow ^{15}\text{N} + e^+ + \nu_e \]
\[ ^{14}\text{N} + p \rightarrow ^{15}\text{O} + \gamma \]
\[ ^{17}\text{O} + p \rightarrow ^{14}\text{N} + ^{4}\text{He} \]
\[ ^{17}\text{O} \rightarrow ^{17}\text{F} + \gamma \]
\[ ^{16}\text{O} + p \rightarrow ^{17}\text{F} + \gamma \]
\[ ^{17}\text{F} \rightarrow ^{17}\text{O} + e^+ + \nu_e \ (^{17}\text{F}) \]

**Figure 1.6:** The CNO cycle reactions with branching ratios for the Sun, values from [51].
1.4 Solar Neutrinos

$pp$ chain provides 99% of the generated nuclear energy, and the CNO-cycle provides the remaining 1%. For stars significantly more massive than the Sun, the CNO-cycle is the main energy generating mechanism.

![Energy spectrum for neutrinos from the Sun with normalization uncertainties.](image)

Figure 1.7: Energy spectrum for neutrinos from the Sun with normalization uncertainties. Figure from [52].

Within the $pp$-chain there are five processes that produce neutrinos. Since the Q-value of the processes in the $pp$ chain are all below the rest mass of a muon or tau, the only charged lepton generated is electrons. And so from lepton flavor conservation only electron flavor neutrinos are generated. And because the neutrino producing processes are all fusion reactions or $\beta^+$ decay, only neutrinos are produced and not any anti-neutrinos. Neutrinos are produced with an energy spectrum shown in Fig. 1.7.

The $hep$ and $^8B$ reactions produce neutrinos with the highest energies. Since the $hep$ reaction branching ratio is so low the flux of $hep$ neutrinos is also very low compared to that of $^8B$ neutrinos; the $hep$ flux is expected to be approximately 0.15% of the $^8B$ flux. So for water-Cherenkov detectors that have a typical threshold of a few MeV, $^8B$ neutrinos are the primary source of detectable solar neutrinos. The uncertainty on the predicted $^8B$ flux
is relatively large, this comes mostly from the uncertainty on the cross-sections and how 
those cross-sections change with temperature, and uncertainties on the temperature profile 
within the core of the Sun. And because the $^8$B reaction has five preceding reactions the 
uncertainty on those reactions are part of the uncertainty on the $^8$B flux (52).

The uncertainty on the $pp$ and $pep$ neutrinos is much lower for two reasons. First, be-
because they are at early stage of the reaction chain, so their reaction rate is not dependent on 
any other preceding interaction. The $pp$ reaction is also the main energy generating mech-
anism for the Sun, so measurements of the total solar luminosity provide strict constraints 
on the $pp$ flux as well.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{solar_neutrinos}
\caption{The radial PDF for each solar neutrino production process. Values from BS05OP.}
\end{figure}

Predictions for solar neutrinos rely heavily on solar modelling, which in turn is con-
strained by helioseismological and photospheric measurements of the Sun. Detailed solar 
models are used to simulate the evolution of the Sun from a proto-star with a given mix-
ture of chemical abundances and produce predictions for what we would expect to observe
1.4 Solar Neutrinos

Figure 1.9: Electron density in units of Avogadro’s Number as function of radius in the Sun. Values from BS05OP.

today. Following this procedure a temperature and density profile for the Sun can be determined which in turn leads to predictions for solar neutrino production. For this work the BS05OP (52) solar model predictions are used. Figure 1.8 shows the solar neutrino production distribution within the Sun. Figure 1.9 shows the electron density profile within the Sun. The radial production distributions and electron density are necessary inputs for solar neutrino mixing calculations.

There are two standard estimates for the solar chemical abundances, GS98 (53) and AGS (54). The BS05 model uses both as inputs and produces separate predictions for each, in general this thesis uses the predictions from the GS98 abundances. The most significant discrepancy between the two models is in the value for the solar metallicity, the AGS prediction for the metallicity is lower by roughly a factor of two. AGS estimates make use of a 3-dimensional model of the solar photosphere, as opposed to the GS98 estimates which use a 1-dimensional model, however the AGS model also produces discrepancies
1.5 Solar Neutrino Mixing

Mixing for solar neutrinos is historically very important as it was the measurements of solar neutrinos that provided the first evidence that neutrinos mix at all \cite{16,21}. Those measurements began the Solar Neutrino Problem. The eventual resolution of which, by SNO and Super-Kamiokande, led to the initial measurements of the neutrino mixing parameters \cite{17,20}. Still today the Sun provides a unique source of neutrinos that travel through densities and distances that cannot be produced by any other source, thus making them a valuable object for study. Chapter 10 discusses the study of mixing with solar neutrinos further.

Neutrinos created in the solar core can experience significant matter enhanced mixing effects from local electron density, including an MSW resonance. The required electron density for the resonance is given in Eqn. \ref{eqn:msw}. This condition is met for a 10 MeV at an electron density of \( \frac{N_e}{N_A} \approx 20 \), \( N_A \) being Avagadro’s number. This condition occurs at a solar radius of \( 0.25R_\odot \), for the mixing parameters given in \ref{chap:1}. For neutrinos with energy below approximately 5 MeV this condition is not met at any point within the sun, and so those neutrinos do not experience an MSW resonance. They can still experience significant matter effects however. Once a neutrino created in the core of the sun has travelled past a solar radius of approximately \( 0.5R_\odot \) the solar electron density has dropped far enough that matter effects are no longer significant and neutrinos are effectively travelling through vacuum.

For solar neutrinos the neutrino states will generally evolve adiabatically from matter effect dominated states to vacuum states. However, when the neutrino transitions through
the MSW-resonance the flavor composition of the mass states changes rapidly, which can lead to a non-adiabatic transition between the mass-1 and mass-2 state. For solar neutrinos the effect this transition has on the neutrino survival probability is characterized with \( P_{\text{jump}} \),

\[
P_{\text{jump}} = \exp \left[ -\frac{\pi}{2} \sin^2 2\theta_{12} \frac{\Delta m_{21}^2}{2E} \frac{dN}{dx} |_{x_R} \right]
\]

(1.36)

Where \( x_R \) indicates the position at which the resonant density is crossed and \( N \) is electron density as a function of position. For experimentally determined values of \( \theta_{12} \) and \( \Delta m_{21}^2 \) in the so-called Large Mixing Angle (LMA) regime the effect of \( P_{\text{jump}} \) is negligible.

One final feature of solar neutrino mixing is the “day-night effect”. The mantle, crust, and core of the Earth have sufficient electron density to provide a non-negligible change to the effective neutrino mixing Hamiltonian. Earth’s atmosphere, however, is not dense enough to provide a significant matter enhancement. The relatively low density of the Earth’s atmosphere, coupled with the fact that at higher energies neutrino mixing lengths

\[\text{Figure 1.10: } ^8\text{B survival probability during the day and night.}\]
are \((O)(100\text{km})\), means that neutrinos that travel a short distance \((<\sim 100\text{km})\) through the Earth will be detected in a vacuum eigenstate. Neutrinos that travel a large distance in Earth’s crust, mantle, or core, will experience significant, potentially non-adiabatic matter effects, and will reach the detector in a matter enhanced state. For solar neutrinos the result is that neutrinos detected during the day, when the Sun is directly above the detector, will not travel significantly through the Earth; neutrinos that are detected at night, when the detector is “shadowed” by the Earth, will travel through large portions of the Earth. Thus, the effective survival probability is different during the day than the night. Figure 1.10 shows the survival probability during the day and night.

1.6 Neutrino Experiments

There’s a long and diverse list of neutrino experiments that have contributed to our current understanding of neutrinos, neutrino oscillations, and solar neutrinos. I won’t attempt to list them all here, but rather highlight the most immediately relevant to this work; These experiments play a significant role in the analysis described in Chapter 10. A more comprehensive review of neutrino experiments can be found in Ref. (29).

1.6.1 SNO

The Sudbury Neutrino Observatory (SNO) was a water-Cherenkov detector located 2 km underground near Sudbury Ontario in Canada, it ran from 1999 to 2006 and detected primarily \(^{8}\text{B}\) solar neutrinos. SNO had the unique feature of being able to detect neutrinos through three different interaction channels, each channel had its own sensitivity to different flavor neutrinos. Combining measurements from each interaction channel allowed for a measurement of the \(^{8}\text{B}\) solar neutrino flux that was not dependent on the flavor composition of the incoming neutrinos. This was accomplished by using a heavy-water \((^2\text{H}_2\text{O a.k.a. D}_2\text{O})\) target and detecting interactions through the electron-neutrino elastic scattering, as
well as charged and neutral-current nuclear interactions on the deuterium nuclei. These interactions are described generally in Sec. 1.2. An NC interaction on deuterium can break apart the neutron and proton that comprises the nucleus, $\nu_x + d \rightarrow p + n + \nu_x$. The free-neutron can then capture on another deuterium atom forming tritium ($^3\text{H}$) and emitting an 6.2 MeV gamma which can be detected; this process has no neutrino flavor dependence. The CC interaction $\nu_e + d \rightarrow p + p + e^-$ is detected by observing the Cherenkov cone of the electron and will only occur for electron flavor neutrinos. ES interactions are similarly detected from the scattered electron, but the electron direction has a strong correlation with solar direction, allowing the rate of ES interactions to be determined separately from the rate of CC interactions. The ES interaction occurs for neutrinos of all flavors, but with an increased cross-section for electron flavor neutrinos.

SNO ran in the three phases, each phase with a different target designed to increase the detector’s ability to observe neutrons from neutrino interactions. The first phase had a pure D$_2$O target. The second phase loaded salt (NaCl) into the heavy-water to increase the neutron capture efficiency through observations of neutron capture on chlorine instead of deuterium. In the third phase an array of $^3\text{He}$ proportional counters were added to the detector, to once again increase neutron detection efficiency (57).

A few main results of the SNO experiment are summarized in Figures 1.11a and 1.11b. Their combined 3-phase result for the $^8\text{B}$ flux is

$$\Phi_{^8\text{B}} = (5.25 \pm 0.16(\text{stat.})^{+0.11}_{-0.13}(\text{syst.})) \times 10^6 \text{cm}^{-2}\text{s}^{-1}.$$  

Their best fit mixing parameter estimates measurements are

$$\Delta m_{21}^2 = (5.13^{+1.49}_{-0.98}) \times 10^{-5} \text{eV}^2$$  \hspace{1cm} (1.37)$$

$$\tan^2 \theta_{12} = 0.436^{+0.048}_{-0.036}.$$  \hspace{1cm} (1.38)$$

Values from Ref. (45).
1.6 Neutrino Experiments

Figure 1.11: (a) The extracted ES, CC, and NC rates in the SNO experiment and how they relate to the effective $\nu_e$ flux and $\nu_{\mu,\tau}$ flux; figure from [20]. (b) The combined best fit survival probability from the combined three-phase SNO analysis (Ref. [45]), the red band is the best fit and uncertainty from a maximum likelihood fit, the blue from a Bayesian fit.

1.6.2 Super-Kamiokande

Super-Kamiokande (Super-K) is a 50 kton cylindrical water Cherenkov detector. It started running in 1996 and has since made the most precise measurements of atmospheric neutrinos and $^8$B solar neutrinos [58, 59]. It’s the successor to the Kamiokande experiment, which was a significantly smaller and had a higher energy threshold for event detection. Super-K can detect $^8$B solar neutrinos through only neutrino-electron elastic scattering, they do not use a D$_2$O target and so are not sensitive to the nuclear interactions similar to that of SNO. Their extremely large detector volume though provides them extraordinary statistics with which they’ve produced the most precise measurement of the $^8$B elastic scattering rate,

$$\Phi_{ES} = (2.308 \pm 0.020(stat.)^{+0.039}_{-0.040}(syst.)) \times 10^6 cm^{-2}s^{-1}$$
Constraining the full $^8\text{B}$ flux to the SNO result they’ve measured the solar mixing parameters as,

$$\Delta m_{21}^2 = 4.8^{+1.5}_{-0.8} \times 10^{-5} \text{eV}^2$$

$$\sin^2 \theta_{12} = 0.334^{+0.027}_{-0.023} \text{[58]}.\$$

### 1.6.3 Borexino

Borexino is 300 ton spherical liquid scintillator detector [60]. Their detector apparatus is similar to that of a water Cherenkov detector, the significant difference is that the water is replaced with pseudocumene, a liquid scintillator. A charged particle moving through scintillator generates roughly 50-100 times more light than is produced by Cherenkov radiation alone. Water-Cherenkov detector are typically limited in energy threshold and energy resolution by the number of photons produced and detected, a liquid scintillator detector solves this problem. Scintillation light, unlike Cherenkov light, is isotropic and provides no information about the direction the particle was moving in.

Water-Cherenkov detector are able to measure solar neutrinos by correlating the direction of detected events with the position of the sun. Since Borexino is not able to determine the direction of events within their detector, they instead perform a spectroscopic measurement. The measurement requires all sources of backgrounds to be accounted for and constrained from *ex-situ* measurements. Figure 1.12 shows the observed spectrum by Borexino and the spectra of the constituent solar fluxes and backgrounds.

Borexino took data from 2007 to 2016, with a pause in 2010 to remove source of radioactive backgrounds and improve the radio-purity of their detector. With that data they’ve measured $^7\text{Be}$, pep, pp, and $^8\text{B}$ neutrino fluxes; they’ve also placed upper limits on the flux of neutrinos from the CNO cycle and from the *hep* solar reaction. They’re currently the only experiment to have measured the pp and pep neutrino fluxes. Figure 1.13 shows these results as the best fit survival probability for each measured flux.
Figure 1.12: Borexino’s observed distribution of event energies with best fit signal and background model. $N_h$ is the number of prompt PMT hits. Figure from [61].
1.6 Neutrino Experiments

1.6.4 KamLAND

The Kamioka Liquid Scintillator Anti-neutrino Detector (KamLAND) is a liquid-scintillator detector. Its primary physics goals were the detection of reactor anti-neutrinos via inverse beta-decay. In addition to reactor anti-neutrinos KamLAND is also sensitive to solar neutrinos. Performing a fit to the observed energy spectrum they were able to measure the flux of $^7$Be and $^8$B solar neutrinos. The $^8$B flux is reported as the “elastic-scattering” flux $\Phi_{ES}$, the flux of pure electron flavor neutrinos that would produce the observed event rate. They measure

$$\Phi_{ES,^8B} = 2.77 \pm 0.26{\text{(stat.)}} \pm 0.32{\text{(syst.)}} \times 10^6 \text{cm}^{-2}\text{s}^{-1}.$$ 

KamLAND’s reported $^7$Be flux, accounting for oscillation effects, is

$$\Phi_{^7\text{Be}} = (5.82 \pm 1.02) \times 10^9 \text{cm}^{-2}\text{s}^{-1}.$$
Figure 1.14: The observed reconstructed event energy spectrum for the Kamland Be7 flux measurement (a) and the 8B flux measurement (b). Figures from [2] and [3].
Interestingly, KamLAND's reactor neutrino measurements [62] are more relevant to the study of solar neutrinos than their solar neutrino measurements. The long baseline (~180 km) and low energy (~3 MeV) of reactor neutrinos provides unique sensitivity to $\Delta m_{21}^2$. Other reactor neutrino experiments, such as Daya Bay, RENO, & Double Chooz, are primarily sensitive to neutrinos with too short a baseline to be strongly affected by $\Delta m_{21}^2$.

\[ \Delta m_{21}^2 = (7.58^{+0.14}_{-0.13}(\text{stat.})^{+0.15}_{-0.15}(\text{syst.})) \times 10^{-5}\text{eV}^2 \]

and,

\[ \tan^2 \theta_{12} = 0.56^{+0.10}_{-0.07}(\text{stat.})^{+0.10}_{-0.06}(\text{syst.}) \]

Figure 1.15: Reactor anti-neutrino energy spectrum observed by KamLAND. Figure from [62]. Figure 1.15 shows reactor anti-neutrino energy spectrum measured by the KamLAND experiment, with the best fit mixing parameters compared to the expected spectrum with no oscillations. From this measurement the best fit value of $\Delta m_{21}^2$ is $7.58^{+0.14}_{-0.13}(\text{stat.})^{+0.15}_{-0.15}(\text{syst.}) \times 10^{-5}\text{eV}^2$ and $\tan^2 \theta_{12} = 0.56^{+0.10}_{-0.07}(\text{stat.})^{+0.10}_{-0.06}(\text{syst.})$. 

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was determined.

The value for $\Delta m^2_{21}$ measured by KamLAND is in disagreement with the value determined by solar experiments, although it cannot be ruled out that the disagreement is a result of a statistical fluctuation. This discrepancy will be discussed further in Sec. 10.
Part I

$^{8}\text{B Neutrino Flux Measurement}$
Chapter 2

Analysis Overview

Described here is the analysis used for extracting the $^8$B solar neutrino flux and interaction rate in the SNO+ detector during its initial water-phase data taking run. This first section provides a broad overview of the data, simulation and analysis methods used for this result. A more detailed description of these topics is provided in the following sections. A report of these results is also published in Anderson et. al. (63).

Neutrinos interact in the SNO+ detector by elastic scattering off electrons in the detector volume. Depending on the energy transferred in the interaction the scattered electron will produce Cherenkov radiation, which is detected by the PMT array. The direction of the Cherenkov cone is expected to be pointing directly away from the Sun at the time of the event for solar neutrino interactions. The principle of this analysis is to search for the rate of events that exhibit this solar direction correlation, and interpret that rate as a solar neutrino flux. This analysis is similar to solar neutrino measurements done by SuperK (58) and, SNO (45).

Data for this analysis was taken from May 2017 to January 2018, with pauses for maintenance, calibration, and commissioning of detector components. The majority of data taken was blinded for an eventual search for invisible nucleon decay (64). A two week period of
time at the start of data taking was left un-blinded to allow for analysis methods to be evaluated, this period of time is referred to as the open dataset.

Simulation is used to determine expected event rates and distributions of detector observables for SNO+. The simulation software used is “RAT”, which performs Monte Carlo simulation of events within the SNO+ detector. Beyond simulation RAT also provides methods for reconstructing event observables, e.g. energy, direction etc. And so RAT is also used for processing detector data in addition to simulated data. There exists an open-source version of this software called RAT-PAC that is largely the same as the SNO+ version of RAT, the most significant difference is the implementations of a detector model and event reconstruction methods. RAT was originally developed largely by Stan Siebert at University of Texas as part of the Braidwood Collaboration, and later developed further for SNO+ here at Penn. RAT-PAC was adapted for more general use and open-sourced by Andy Mastbaum while at Penn.

The $^8$B solar neutrino signal is extracted from the detector dataset using a number of cuts to isolate likely solar neutrino events, and remove background events. Two broad categories of cuts exist, data cleaning and analysis cuts. Data cleaning aims to remove events that originate from instrumental effects using low-level criteria, such as the distribution of charge across the hit PMTS. Analysis cuts are designed to select the events originating from the physics of interest, in this case $^8$B solar neutrino events. The analysis cuts make use of reconstructed information, such as event energy or position.

A fit is performed to the dataset after cuts, searching for the rate above background of events that are directed away from the Sun as the time of the event. The source of these events is assumed to be $^8$B solar neutrinos, and so the flux $^8$B events is estimated from the observed event rate. Simulation is used to produce the PDFs used in the fit and provide a map observed event rate to neutrino flux. Since simulation plays a significant role in this analysis, data taken using a deployed $^{16}$N source is used to evaluate the accuracy of the
simulation, assign systematic uncertainties to distributions of simulated observables and similarly provide empirical corrections to those observables.
Chapter 3

SNO+ Detector

The SNO+ detector inherits much of its infrastructure from the SNO experiment (66). The detector is mostly simply described as a large volume of a target material that is deep underground and is observed by a large array of photomultiplier tubes (PMTs). The target material dictates the physical processes observable with the detector. SNO was originally designed for a heavy-water target, the benefits of a heavy-water target are discussed in Sec 1.6.1. For SNO+ the detector will operate in three separate phases, each with a different target medium. For all results in this work the target medium is light-water (H$_2$O); this phase is referred to as the “water phase”. In the next phase the water is to be replaced by a liquid-scintillator, LAB-PPO (67), that will be the “pure scintillator phase” or just the “scintillator phase”. After the scintillator phase the LAB-PPO will be doped with tellurium for a neutrinoless double-beta decay search, this is referred to as the “tellurium phase”. The motivations for each phase is discussed in Ref. (68).

The detector’s target volume is encapsulated within a 6-meter radius spherical acrylic vessel (AV), which is held suspended in a large cavity filled with ultra-pure water (UPW). The acrylic sphere has a 6.8 m acrylic chimney, called the “neck”, at its top to allow access to the detector volume. Surrounding the acrylic vessel is an array of inward pointing PMTs.
Figure 3.1: (a) Drawing of the SNO+ showing the AV, PSUP, cavity, neck, ropes, and upper deck. (b) Schematic depiction of the AV and hold-up and hold down ropes.
The structure holding these PMTs is referred to as the PMT Support Structure (PSUP). Figure 3.1a depicts the SNO+ detector.

There are roughly 90 PMTs on the PSUP that point outward, towards the surrounding cavity walls. These outward looking tubes are called OWLs and are for the purpose of tagging particle interactions that occur outside the PSUP. There are an additional three tubes mounted at the top of the neck of the AV, these are referred to as NECK tubes.

SNO+ has 9385 inward looking PMTs all mounted in the PSUP at an average radius of 8.4 m from the center of the detector. All PMTs are Hamamatsu R1408 8-inch PMTs. The inward looking PMTs are all housed within a plastic cassette, and each PMT is also surrounded by an array of reflective petals arranged in a Winston cone (69). The reflective petals

Figure 3.2: Schematic diagram of the SNO/SNO+ PMT and PMT housing. Dimensions in cm. Figure from (66).
petals serve to increase the effective photo-sensitive area of the PMT and are typically referred to as the PMT “concentrator”. With the concentrators the approximate geometric coverage of the PMT array is 55%, accounting for the angular acceptance and reflectivity of the concentrators gives an effective coverage of approximately 50%. Figure 3.2 shows a schematic diagram of the PMT and its housing.

For the scintillator and tellurium-loaded phases the AV volume will have a lower density than the surrounding water volume, resulting in significant buoyant forces on the AV. To counteract the buoyancy a hold-down rope-net was installed across the top of the AV, and anchored to the cavity floor. A number of PMTs were removed from the PSUP to allow the rope to pass through. Figure 3.1b depicts the rope-net.

Above the cavity volume is an optically isolated deck on which all the detector readout and trigger electronics are kept. The PMT housing provides a BNC-like connection that allows the 2kV power supply to be provided to each PMT and allows for the PMT signals to be read out.

### 3.1 Detection Mechanism

The goal of the SNO+ detector is to detect and record information about as many of the photons produced within the AV as possible. By observing photons information about the physical processes that produced the light can be inferred. Cherenkov radiation is the primary photon generating process of interest in SNO+’s water-phase. Cherenkov radiation is produced by any charged particle moving with velocity ($\vec{v}$) such that $|\vec{v}| > \frac{c}{n}$, where $c$ is the speed of light and $n$ is the index of refraction in the target medium (70).

Cherenkov light has a few properties that make it a desirable method for particle detection. The first property is that the number of photons produced scales nearly linearly with the energy of the particle propagating, therefore the energy of the particle can be estimated by simply counting the number of detected photons. The second desirable property is that
3.1 Detection Mechanism

the produced photons will be emitted in a cone, with an opening angle of approximately $42^\circ$ with respect to the particle’s direction of travel. Meaning by detecting Cherenkov light the particle’s direction of travel can be inferred. And finally, if the time of each detected photon is well-known then the requirement that all photons travel from the same point at the same speed allows the position of the particle to be deduced. SNO+ takes advantage of each of these aspects of Cherenkov radiation to reconstruct the position, time, energy, and direction of all interacting particles in the detector. The methods used for deducing each value is detailed in Sec 5.

For the scintillator and tellurium-loaded phases scintillation processes will be the dominant photon production mechanism. The Cherenkov light production is approximately the same in scintillator as it is in water, however the ratio of light produced by scintillation processes to Cherenkov light is approximately 100:1. With SNO+ it’s not expected to be possible to infer which detected photons are from scintillation processes and which are Cherenkov radiation, so it can be treated as though all light is from scintillation. Many people are developing methods to separate Cherenkov light from scintillation light [71, 72, 73], and so this may be possible for SNO+ in the future.

The primary neutrino interaction that SNO+ is sensitive to is elastic scattering (ES) off electrons. Charged and neutral-current nuclear interactions occur as well with the oxygen in the water, however these are rare and cannot be uniquely identified against radioactive backgrounds, and so are ignored. The cross-section for the ES interaction is discussed in 1.2. The forward peaked angular cross-section means that information about the neutrino direction is maintained in the interaction. But the differential cross-section for recoil energy is nearly flat below the end point. Meaning relatively little information about the incoming neutrino energy is preserved by the interaction.
3.1 Detection Mechanism

3.1.1 Electronics And DAQ

The hardware the connects to and reads out the PMT array forms the SNO+ data acquisition (DAQ) system. The SNO+ data acquisition (DAQ) inherits much of its design and components from SNO. There are a few notable upgrades that were made for the purpose of handling the higher light yield and event rate that SNO+ has compared to SNO. The DAQ hardware can be described as a few separate systems, the trigger system, the readout system, and the PMT interface system. The PMT interface provides an approximately 2 kV (HV) supply to each PMT and provides the signal from the PMT to the rest of the DAQ electronics. The trigger system’s purpose is to decide when an interesting interaction within the detector has occurred, and to start the readout process when such an interaction has occurred. The readout process is responsible for ensuring enough information about each PMT signal is recorded such that offline analysis is possible.

The first step of the PMT interface system is the PMT base. The base is responsible for fanning-out the supplied HV to the PMT dynode pins and connecting the PMT output to a PMT cable. The PMT base is housed within a water-tight cassette. The PMT cables pass through penetrations in the cavity ceiling where they then connect to the rest of the electronics. The PMT cable connects first to a PMT interface board (PMTIC). The physical connection occurs on a daughter card, called a “paddle card”, that accommodates up to eight PMT cables; Each PMTIC hosts four paddle cards. The PMTIC is responsible for fanning out the PMT high voltage to each PMT and providing channel level adjustment to the voltage each PMT receives; the voltage adjustment is done with a series of swappable resistors. The PMTIC is also responsible for separating the PMT signal from the supplied HV, this is achieved with a capacitative decoupling circuit. Once the two signals are separated the PMTIC sends the PMT signal to a front end card (FEC) via a board-to-board connector, where it enters the readout and trigger system.

That signal is compared to a threshold, if the signal is over threshold a “hit” has occurred — this threshold is often called the “channel threshold”. At the time of the channel
3.1 Detection Mechanism

threshold crossing the following processes occur, the name for each is given in parentheses: a 100 ns long fixed-height square pulse is created (N100), a 20 ns fixed height square pulse is created (N20), a high gain copy of the signal is created (ESUMH), a low gain copy of the signal is created (ESUML), a linear voltage ramp begins (TAC ramp), the signal is integrated for 50 ns with high gain (QHS), the signal is integrated for up to 400 ns with high gain (QHL), and the signal is integrated for 50 ns with a low gain (QLX). These signals and values are created on a few different custom ASICs on the daughter boards. The trigger system uses the first of those 4 signals (N100, N20, ESUMH, and ESUML). The readout system uses the latter four values (TAC, QHS, QHL, QLX).

The trigger signals are all combined with their counterparts from other PMT channels across the detector, i.e. the N100 signals from all channels will be combined and separately all the N20 signals will be combined, etc. The signals are combined through analog summation, summing is done on a few different circuit boards within the detector. The FEC sums its top and bottom sixteen channels separately, the crate trigger card (CTC) sums the signals from the sixteen FECs that are in each electronics crate. The signals from each of the nineteen CTCs are all summed on the Master Trigger Card - Analog (MTCA+). The SNO+ MTCA+ is an upgraded version of the SNO MTCA; more information about the MTCA+ is available in Sec. 3.2.2.

Separate, but identical, MTCA+s are used for each of the four trigger signals. Each MTCA+ performs the analog summation with three different gains, resulting in a total of twelve signals spread across four different boards. Each of the twelve signals are separately compared to a threshold; each of the twelve thresholds are independent of each other. These thresholds are called “trigger thresholds”.

The different gains are in place due to the practical difficulty of maintaining a good signal-to-noise ratio (SNR) without limiting the range of the system. For example, if there exists 10 mV of noise in the system a 20 mV pulse would give a 2:1 SNR, however this would mean if 5000 PMT hits occurred simultaneously the signal would be 100 V in size.
3.1 Detection Mechanism

It is not practical to have a system with 100 V range and 20 mV resolution, so the three different gain paths allow for three different trade-offs between SNR/resolution and range. The highest gain signal has the best SNR, but the smallest range, and so usually the highest gain signal has the lowest effective threshold. The reason being that it’s more important to have single hit resolution at a threshold of 8-hits than it is at a threshold 25 hits. The different gains on each signal are therefore labelled by their threshold (not their gain), e.g. the high, medium and low gain paths for the N100 signal are respectively called N100 Low (N100L), N100 Medium (N100M), and N100 High (N100H).

Although there are twelve signal-gain combinations available only seven are used: N100-Low, N100-Med., N100-High, N20-Low, N20-Med (also called just N20), ESUMH-Low, and ESUML-Low. Since the ESUMH and ESUML each only use one gain path, they’re usually referred to simply as ESUMH and ESUML with their gain path understood to be the high gain path.

When a trigger signal goes over its threshold a 20 ns digital pulse is emitted for that signal. This pulse is called a “raw-trigger” and there is one for each of the seven used trigger signals. The raw-trigger signals are sent from the MTCA+s to the Master Trigger Card-Digital (MTCD). Finally, each of these seven raw-trigger signals can be masked in or masked out on the MTCD; if a raw-trigger is masked out, nothing happens when it fires, if it is masked in, then the raw trigger creates a “global-trigger” (GT) signal. That global trigger signal is fanned out to all the data crates which in turn sends the GT to all front end cards and daughter boards. When the GT signal is created the MTCD also generates a signal called Lockout (LO). Lockout is typically a 420 ns long pulse and while the signal is high the MTCD will not create any more global triggers.

Once the global trigger is created the trigger cycle is complete and the readout process begins. The raw-trigger signal that caused the global trigger, as well as any other raw-trigger signals that were high within a 20 ns window of the global trigger, are recorded and
3.1 Detection Mechanism

readout, this is known as the “trigger word”. When the GT is created a counter, called the

The four values that are created by the PMT signal crossing the channel threshold
(TAC, QHS, QHL, QLX) are stored in analog memory cells on the daughter boards. They
are stored for a length of time known as “GT_VALID”, if a GT does not arrive before
GT_VALID expires the TAC, QHS, QHL, & QLX values are discarded. A typical value for
GT_VALID is 400\,ns, although there exists some channel-to-channel variation. If a GT
signal does arrive at the channel before GT_VALID expires the values in the memory cells
are digitized and readout to a memory buffer on the FEC. The TAC ramp starts when the
PMT signal crosses channel threshold and stops when the GT signal arrives at the channel.
Since the TAC voltage increases linearly with time, the value of the TAC indicates when
the hit occurred relative to the GT signal.

The FEC stores those values and adds information to identify which channel’s data
is stored, it also records the value of its own GTID. Each FEC keeps a counter that is
incremented every time it receives a global trigger signal, in principle the value of this
counter will always be the same as the MTCD GTID, and the same as the counter in every
other FEC in the detector. The GTID counter is our only way of associating recorded hit
data with each other and with the trigger word.

In practice, it is possible for a channel’s GTID to become out of sync with the GTIDs of
all other channels. This can result in the hits on a particular channel being associated with
the wrong event. To mitigate this problem every \(2^{16}\)th and \(2^{24}\)th GT respectively creates a
SYNC and SYNC24 signal, those signals are sent by the MTCD to each FEC & DB. If a
FEC or DB receives either of these synchronization pulses but its own GTID counter is not
at an increment of \(2^{16}\) or \(2^{24}\) then the channel is identified as out of sync. If this happens,
the GTID counter is adjusted to the correct value and the next hit to read out from the
out of sync channel/channels is accompanied by a flag to indicate that it was out of sync.
This system ensures a channel is never out of sync for more than 65536 events.
A short while after the data and the associated identifying information and status flags are buffered in FEC memory, the data is read out by a crate level readout card, the “XL3”. The XL3 is new to SNO+; it replaces the XL1 and XL2 from SNO, more will be said about the XL3 in Sec. 3.2.1.

Each crate has its own XL3, all XL3s read out and serve data asynchronously. The data-server process receives data from each XL3 and relays that data to any clients that have subscribed to the PMT data feed. A similar process is done for the trigger word data. The MTCD sends trigger data to the data-server, the data-server relays that data to any clients that have subscribed clients.

The primary client to the data server is what’s known as the “Event Builder”, sometimes called the EB or just the “Builder”. The Builder receives data from the data-server and uses GTID information to associate trigger words and hits with each other. Once all the hits for an event have been associated with their trigger word the event has been “built” it is written to disk and the read out process for that event is complete. Data is typically taken in hour long chunks referred to as a “run”; every run has a unique number associated with it and a “run type” number that gives basic context to the detector circumstances and settings in which the data was taken. The Builder, in addition to building events, is responsible for associating events with their run number and run type.

There are a few ancillary systems within the DAQ electronics, all of which are new to SNO+. The first is the CAEN v1720, commonly referred to as just “CAEN”, which is a 12-bit digitizer board. It’s role follows from the Analog Measurement Board (AMB) used in SNO. The CAEN is used to digitize and readout the trigger signals. It has eight available input channels that it can digitize, however, typically only three signals are actually used, those channels digitize ESUMH, N20L, and N100L. The CAEN’s digitization window and sampling rate can be varied, most commonly the digitization window is 420 ns and the sample width is 4 ns. The CAEN receives a copy of the global trigger allowing and it keeps its own GTID counter so its data can later be associated with the appropriate hit and trigger
data. It also receives a copy of the SYNC and SYNC24 signal so its synchronization can be ensured.

The input voltage range for the CAEN is an adjustable 2 V window. The voltage range for the trigger signals is 10 V. The difference in ranges necessitates some way of reducing the range of the trigger signals before they're sent to the CAEN. The simplest way of reducing the voltage range is to use a voltage divider to attenuate the signal by a factor of five. Attenuation has a few undesirable effects though. The full range of the trigger signal is 10 V, but the vast majority of events will only use a small fraction of that range. So for events that use a small amount of the available 10 V a factor of five attenuation will make the signal much smaller than it needs to be, resulting in loss of information because the signal will be smaller than the analog noise, or from the noise digitization process itself. And for the purpose of most analyses that use the data from the CAEN it's more important to be able to resolve a single hit than to resolve the height of the full pulse if the pulse is very large.

A scheme for fitting the trigger signal into the CAEN’s available range that optionally allows for either single hit resolution or preserving the full signal range was adopted. The trigger signal is clipped within the first 2 V, thereby retaining full resolution for small signals, but losing resolution for signals that go over 2 V. A board was designed to optionally clip or attenuate the trigger signals before digitization, for the vast majority of the data taking the signals were clipped. I designed the board that performs this analog signal processing, it’s discussed in further in 3.2.3.

### 3.2 Electronics Upgrades

As previously mentioned, in SNO+’s scintillator and tellurium-loaded phases the amount of light produced by any interaction is expected to be roughly a factor of 100 greater than the light that would be produced by a water target. This increase in photon production
translates to an equivalent increase in the current in the trigger system and the necessary data read out rate. Between the time when the original SNO DAQ electronics was designed and now, the availability and sophistication of commercial computing and DAQ hardware has increased dramatically. To accommodate the increased current and data volume and to take advantage of modern hardware a few key pieces of the SNO DAQ was upgraded as part of the change from SNO to SNO+.

Beyond the need to improve the DAQ for the higher expected light yield there were a few aspects of the SNO electronics that motivated several features in the upgraded electronics. Most notable of these is the unstable analog trigger baseline. The voltage of any trigger signal’s baseline can vary. The baseline of each trigger signal is the voltage observed when there are zero hits in the analog sum, As the baseline moves closer to or further from the trigger threshold the number of PMT hits required to trigger the detector changes. The result is a difficult to understand trigger efficiency and bursts in the trigger rate that can overwhelm the DAQ.

The baselines are sensitive to a number of known factors such as the ambient temperature, the PMT noise environment, and settings on the front-end. There are also a number of factors that effect that are more difficult to identify, such as transistors on the CTC performing malfunctioning due to age or other unknown factors. These factors lead to the baseline for any trigger signal varying by up to a few hits over the course of a few hours. A common source of baseline variation is known as “dropout”; dropout is single hit shifts in the baseline that persist over an indefinite time period with no clear signature in the readout. Dropout will be discussed in detail in 3.3.1.

3.2.1 XL3

The SNO DAQ used a centralized serial readout system, where each crate of electronics was read out in one after the other. As part of the electronics upgrade from SNO to SNO+ this system was changed to an asynchronous, parallel readout system, allowing for a significant
3.2 Electronics Upgrades

increase in the data readout speed. The board responsible for this is the XL3 which hosts a Xilinx ML403 Evaluation platform. The ML403 uses a Xilinx Vertex-4 FPGA as its primary logic chip and has 64-MB of supporting SDRAM and persistent memory provided by a CompactFlash card. The XL3 & ML403 interface with the FECs in a crate through VME-like communication across the “SNOBUS” backplane.

![Diagram of the SNO+ DAQ](image)

**Figure 3.3:** Diagram of the SNO+ DAQ. Figure from [74].

The XL3 reads data from the FECs across the backplane, and buffers that data in on-board memory before reading it out to a central DAQ computer over Ethernet. The maximum data rate each XL3 is capable of is approximately 14 MB/s, yielding a detector-wide rate of 250 MB/s. This translates to being able to read out roughly one-hundred thousand hits per crate per second. Figure 3.3 shows schematically how the XL3 is incorporated into the SNO+ DAQ.
Figure 3.4: Schematic diagram of analog summation in the SNO+ trigger system. Figure courtesy of Andy Mastbaum.
3.2.2 MTCA+

The SNO MTCA was not expected to be able to operate at the expected hit rate and occupancy of SNO+ in its scintillator or tellurium-loaded phases. So the primary goal for the MTCA+ is to simply replace the analog summing and discriminating capabilities of the SNO MTCA, and perform stably at the higher rates expected in SNO+. For this reason the MTCA+ performs the analog multiplicity sum using a series of high-current voltage feedback operational amplifiers.

One of the most transformative changes that the MTCA+ introduces into the SNO+ trigger system is its baseline restoration circuitry. In SNO typical baseline variations could be tolerated because the threshold was far from the baseline, so variations of a few hits did not typically have a very large effect. In SNO+, due to many of the upgrades, a significantly lower threshold was achieved, so a variation a few hits causes a much larger change in the trigger rate, leading to a higher likelihood of triggering faster than the maximum possible readout rate.

The MTCA+ provides two ways to mitigate baseline variations. The first is that the MTCA+ provides a relay to dynamically enable or disable each crate’s participation in the trigger sum. This is useful if a CTC fails, its trigger sums can be disabled to prevent it from pulling up/down the entire trigger sum to the point that stable triggering is no longer possible. This situation is rare, but the ability to disable individual crates in those situations is useful for diagnosing and isolating issues.

The second is baseline restoration circuit on the MTCA+. At the final stage of each analog sum the output sum is fed through a long-pass filter to extract the average voltage over an \( \approx 1 \text{s} \) period. That voltage is buffered and fed back into the non-inverting input of the operational amplifier used for the final stage of the analog sum. The effect of this feedback loop is to subtract any long term voltage offsets from the trigger sum.

Using the long term average of the trigger sum is a good way of determining the sum baseline in the limit that variations from PMT hits have a small effect on the average.
Since the average is performed over a period of \( \approx 1\, \text{s} \) and the trigger signals are \( \approx 100\, \text{ns} \) wide, then a hit rate of \( \approx 10^7 \, \text{hits/second} \) is required for a significant effect on the average trigger signal. Since there are \( \approx 10^4 \) PMTs participating in the trigger sum at any time, and each has a typical dark rate of \( \approx 1\, \text{kHz} \), the criteria of \( 10^7 \, \text{hits/second} \) is met. This means the baseline will be adjusted to account for the dark-rate hits. The PMT dark rate is the dominant source of hits for the detector when it has a water target, it’s suspected, but not known, that this will still be true for a scintillator target as well. Since typical variations in the baseline from thermal and other environmental effects occur on the \( \approx 1\, \text{hour} \) timescale the \( 1\, \text{s} \) time scale for the baseline restoration provides adequate correction for those sorts of effects.

### 3.2.3 TUBii

I designed an auxiliary trigger board for SNO+ called the Trigger Utility Board Mk.II (TUBii) the spiritual successor to Trigger Utility Board from SNO. TUBii has a number of different roles in the SNO+ DAQ, as already mentioned it provides the analog shaping for trigger signals sent to the CAEN, but it also provides digital signaling to allow different DAQ & calibration subsystems to communicate, and it has its own triggering functionality as well. TUBii has approximately 90 inputs and outputs, most of which are transmitted through BNCs on its front or back panel, Fig 3.5a shows a picture of TUBii’s front panel. Figure 3.5b shows the board in its box, with connections between the board and box.

The CAEN analog buffer operates by taking the MTCA+ trigger signals and splitting the signal along two different paths. The first path simply removes the \(-5\, \text{V} \) baseline from the signal through a unity-gain op-amp. That signal is meant to go to a monitoring oscilloscope for detector operators. The second signal also goes to a similar op-amp that removes the \(-5\, \text{V} \) baseline, but the series of input resistors to that op-amp is controlled by a reed relay. For one set of resistors the op-amp has unity gain, for the other set of resistors the signal is attenuated by a factor of five. That signal, attenuated or not, is clipped by
3.2 Electronics Upgrades

Figure 3.5: The front panel (a) for TUBii show its main inputs and outputs and (b) the box interior showing the board itself and the connections between the board and box.

a series of diodes to fit within an approximate 4 V range. Without this analog buffer the CAEN digitizer would not have been useful for digitizing trigger signals. Data from the digitizer was used in a detector monitoring and modeling tool that I developed, described in Sec 3.3.1 and in a cut to remove instrumental backgrounds that I developed, described in Appendix A.

TUBii also has its own triggering functionality that supports the main trigger system. It can receive raw-trigger signals from the MTCA+s or any other source and apply custom trigger logic to them and emit its own raw-trigger pulses which are sent to the MTCD. TUBii also receives the global trigger signal and produces its own trigger word based upon which raw trigger pulses it had received. The TUBii trigger word is synchronized with the rest of the data for each event through its own global trigger counter and through the SYNC/SYNC24 signal. Beyond triggering on input digital signals TUBii has two inputs for analog signals. TUBii has an analog discriminator that can produce its own raw-trigger signal and that signal is sent through re-trigger logic that mimics the re-trigger logic on
3.2 Electronics Upgrades

the MTCA+ CPLD. This analog discrimination and re-trigger logic is referred to as the “MTCA Mimic” functionality because it provides a “plug-and-play” way to trigger the detector off an arbitrary analog signal.

TUBii is used as an interface board for some detector calibration systems. These systems emit light into the detector and usually need to be synchronized with the trigger system. This synchronization requires a variety of pulses and delays tuned to account for the time it takes for signals and light propagate throughout the detector and DAQ system; TUBii provides those pulses and delays.

TUBii’s customizable complex trigger logic allows it to create trigger pulses from its inputs. The input trigger signals are fed into a Xilinx MicroZed, which is an FPGA and micro-controller. The MicroZed allows for nearly any logical combination of trigger signals including using recent trigger signals to inform the current trigger logic.

Something like this is desirable for identifying and ensuring the detector will be sensitive to time-correlated events. An example of this would be that the decay chain of $^{214}$Bi → $^{214}$Po → $^{210}$Pb, this decay chain is referred to as BiPo214. The signature of this decay is an electron from β decay, followed, with a half-life of 4 µs, by an α decay. It’s very important that the α decay is detected so that the β-α decays can be identified as likely from a BiPo. If the α is not observed the β can be mis-identified and potentially leak into a signal region. TUBii is able to mitigate this risk by having a trigger that is particularly sensitive to the initial β decay and can trigger off a lower threshold input for a short time after the β trigger; ensuring that the α is detected.

TUBii also provides general purpose and “glue” functionality, facilitating different circuits from different boards in the DAQ to communicate. An example of this is that the CAEN requires the global trigger and other synchronization pulses be sent to it using Low-Voltage Differential Signaling (LVDS), but the global trigger is created using Emitter Coupled Logic (ECL). And so TUBii provides translation between these two digital signaling protocols, allowing the CAEN to remain synchronized.
3.3 Electronics Calibration

There are three primary calibrations performed for electronics to ensure that the detector behaves predictably and that readout values can be interpreted for a physics analysis. The first calibration is the ECAL (Electronics Calibration), the next is the ECA (also Electronics Calibration), and the final one is the PCA (PMT Calibration).

Both the ECA and ECAL use the PEDESTAL and PULSE_GT signals. Both signals are produced on the MTCD by a pulser. The PULSE_GT simply produces GT signals at a fixed rate. The PEDESTAL signal is sent to the FECs, and they fake a PMT hit occurring, i.e. a hit occurs in the electronics regardless of if the PMT has produced a signal or not. The channels that do or do not receive the PEDESTAL can be arbitrarily chosen. Since the PEDESTAL signal does not change the PMT signal that is measured, the QHS, QHL, and QLX will always read out with the same value. The same is true for the TAC, the PEDESTAL is always emitted a fixed time before the PULSE_GT signal, meaning the time between the PEDESTAL hit and the GT readout will always be the same. The time delay between the PEDESTAL and PULSE_GT can be adjusted from 24 ns to 2574 ns.

The goal of an ECAL is to provide settings for each channel that will result in a uniform detector response. Put differently, the ECAL attempts to minimize channel-to-channel variation across the detector. A number of factors need to be accounted for to produce a uniform detector response for example, the slope at which the time it takes for the TAC ramp to complete, the value for the channel threshold, the length of the GT_VALID signal, etc. The ECAL does this through a suite of separate tests and calibrations. ECALs are only ran as needed and typically an ECAL is only need after a board within the detector is replaced or repaired.

The ECA is generally used for determining how values from the detector map to absolute physical values. There are two varieties of ECA, PDST and TSLP. The PDST ECA consists of sending many PEDESTAL signals to each channel in the detector and measuring the distribution of charge values (QHS, QHL, and QLX) from each channel. This provides a
3.3 Electronics Calibration

determination of which values of each charge correspond to zero PMT signal and how much
those values can vary. This zero-point measurement is where the PEDESTAL signal derives
its name; it measures the charge pedestal upon which the PMT signal sits, so to speak.

The TSLP calibration follows a similar procedure, but varies the delay between the
PEDESTAL and PULSE_GT. The result is a precise determination of the mapping between
time (in ns) and TAC value. Beyond providing a mapping between physical values and
recorded values the ECA also provides information about which electronics channels are
working reliably and which are not capable of producing useful data. Channels that cannot
produce useful data are removed in later analysis but are typically not modified within the
electronics, except in the case where they can be repaired or replaced. Both varieties of
ECA are run on an approximately weekly basis to account for variations that may occur
with time in the read out values and to quickly identify when a channel becomes unreliable.

The final electronics calibration, the PCA, is the only one to make use of the PMTs.
The PCA is used for identifying the charge associated with the detection of a single photon
by each PMT. There exists some variation in that value from differences in the electronics
and the PMTs themselves, the PCA attempts to measure those variations. For SNO+ there
exists two ways of performing a PCA, the first is with a deployed light source called the
“laserball”. More information about the laserball can be found in Ref. [75]. The laserball is
typically placed within the center of the detector and emits light isotropically. For a typical
laserball PCA the amount of light emitted is very small, such that only a few PMTs detect
anything in a single event; this ensures that no PMT is likely to observe more than a single
photon. Data are taken this way for a long time so that every PMT is hit many times over
many events. The data is later analyzed to extract how much charge corresponds to a single
photon for each channel.

For SNO+ a similar procedure can be done using a newly installed laser/LED system
mounted on the PSUP called ELLIE (Embedded Laser/LED Light Injection Entity). The
ELLIE system consists of a number of fibres that project light from one side of the PSUP,
across the detector, to the PMTs on the other side. The fibres are placed at a number of different positions around the PSUP. ELLIE can be used for a number of calibration purposes, including playing a similar role to the laserball for a PCA.

3.3.1 Dropout

Dropout comes from an error in the design of one of the ASICs on the DB; the error results in the N100 and N20 trigger pulses from a channel being much longer than they should be, e.g. 1 ms wide instead of 100 ns. The nominal effect of dropout is to lower the effective threshold for the period that the channel is dropped out. Since the typical length of a single dropout pulse is short compared to the baseline correction time constant, individual channels being dropped out are not corrected for.

Dropout does have a somewhat indirect effect on the baseline correction though. The baseline correction will move the “zero” of each trigger signal to its most common value. So if there exists, on average, two channels dropped out in the detector at any time, then that average will be shifted by approximately two nhit. If the threshold for the detector is set to 10 nhit, the effect of dropout and baseline correction will leave the effective threshold at 10 nhit, but for short periods when there is fewer or greater than two dropped out channels there will be an effectively higher or lower threshold, respectively. Because of baseline correction, dropout can cause the effective threshold to move in both directions, higher or lower. In SNO, without baseline correction, dropout would only ever lower the effective threshold.

For physics analyses with a signal near threshold, it’s important that the threshold for the detector be as well modeled as possible. So a method for accurately modelling dropout is desirable to avoid large systematic uncertainties associated with the trigger system. Since dropout is the result of a design error in the front-end system, the readout system is not sensitive to it, so there is no straightforward way of measuring how many or which channels are dropped out at any time. Using the data recorded by the CAEN I was able to develop
3.3 Electronics Calibration

a method for determining how many channels are dropped out during certain triggered events. Using that measurement I was able to estimate the rate of channels dropping out in the detector as a whole. This information is included in our simulation of the detector DAQ system to improve our model of the detector response.

I developed a method for extracting the dropout from the CAEN data recorded by the detector. The method is to measure the baseline of the CAEN recorded N100-Lo and N20-Lo trigger signals. The measured baseline is histogrammed and then the function

$$\text{Pr}(x) = \sum_{k=0}^{\infty} \text{Pr}(x|k) \text{Pr}(k) = \sum_{k=0}^{\infty} \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\frac{(x - (S \ast k + C))^2}{2\sigma^2}} e^{-\lambda \frac{\lambda^k}{k!}}$$

(3.1)

is fit to the histogram. Eqn. 3.1 describes a series of Gaussian distributions each with width $\sigma$ and separation $S$, and an overall shift $C$. The normalization of the $k^{th}$ Gaussian is given by the value of the Poisson distribution for an average rate $\lambda$. The parameter $S$ gives the separation in ADC counts between the baseline shifts from each dropped out channel, $C$ gives the baseline value when there are zero dropped out channels. The width of each Gaussian $\sigma$ corresponds to the noise on the baseline measurement, i.e how much the baseline varies at a fixed amount of dropout. The parameter $\lambda$ corresponds the average dropout for the detector. All parameters are determined from a fit.

Figure 3.6 shows the measured dropout for run 101357. The best fit extracts an average dropout of $\lambda = 2.7$ channels. For each run a similar dropout measurement is performed, the resulting dropout rate is later used in simulation as part of the DAQ simulation.

3.3.2 Nhit Monitor

During standard data taking a calibration, called the nhit monitor, is performed to measure the effective threshold of the trigger N100 and N20 triggers. The goal is to estimate the function $P_{\text{trig}}(N)$, called the trigger efficiency curve, it’s the probability of triggering at any given number of hits $N$. The process for estimating that curve is to steadily increase
3.3 Electronics Calibration

Figure 3.6: [black solid line] The measured baselines for run 101357 and [red] the best fit dropout model with the individual Gaussian distributions [black dashed line] that make up the best fit.
the number of PEDESTAL hits occurring at a single time and observe at which number of hits the detector triggers, and how frequently it triggers. For the entirety of the data taking only channels in crate 4 of the detector were pedestalled for the nhit monitor. This introduces the possibility of bias into the nhit monitor estimate for the trigger efficiency curve. Although all channels in the detector are designed to function identically, this is not something that is closely monitored or tested, so it could be the case that the channels on crate 4 are not representative of the detector as a whole. For this reason the estimates of the trigger efficiency from the nhit monitor are usually compared to other estimates from laserball data.

A value related to the number of hits in an event, called the “in-time hits” or $\tilde{n}$, is typically used for estimating the trigger efficiency. The in-times hits is an estimate of the maximum height of the trigger pulse in units of nhit, it includes effects from the pulse width of each trigger and the average pulse rise-time. Each event has two in-time nhit estimates one for the N100, $\tilde{n}_{100}$ and one for the N20, $\tilde{n}_{20}$. Figure 3.7 shows the trigger efficiency from nhit monitor data compared to data from simulation for the N100 trigger.
3.3 Electronics Calibration

Figure 3.7: Comparison of the measured trigger efficiency and simulated trigger efficiency for run 107640. Plot from (76).
Chapter 4

Simulation

4.1 RAT

Figure 4.1: The $^8$B neutrino flux normalized to the solar reaction rate predicted by BS05-0P \cite{52}
4.1 RAT

RAT, a Monte Carlo simulation of particle interactions in the detector is used for predicting detector observables for solar neutrino and background events. RAT is a Geant4-based (77) simulation that contains a detector and DAQ simulation in addition to simulation of particle interactions and photon propagation.

The first step in the Monte Carlo event simulation process is to generate event vertex information, including the interaction location, and the neutrino and recoil electron energy and direction. The distribution for the $^8$B neutrino energy from Winters et al. (78) was used, normalized to a nominal flux of $\Phi_{\nu_e} = 9.67 \times 10^9\text{cm}^{-2}\text{s}^{-1}$ and $\Phi_{\nu_\mu} = 5.46 \times 10^{10}\text{cm}^{-2}\text{s}^{-1}$. These values are the full BS05OP (52) $^8$B flux scaled by a factor of 1700 for the $\nu_e$ and a factor of 9600 for the $\nu_\mu$. The overall flux normalization is arbitrary in this analysis, but the enhanced rate ensures that the statistical fluctuations observed in the MC datasets are negligible compared to the detected dataset.

![Figure 4.2: The differential cross section for $\nu_e$ electron elastic scattering (a) and $\nu_\mu,\tau$ (b) as used by RAT. Units for Z-axis is $10^{-42}\text{cm}^{-2}\text{MeV}^{-2}$.](image)

The rate of solar neutrino interactions for a given flux follows from the cross-section for interaction. The only interaction relevant for this analysis is the neutrino-electron elastic scattering interaction, the cross-section of which is discussed in Sec. 1.2. The model used for
event generation in RAT includes radiative corrections as described in Bahcall et. al (79). The differential cross-section as a function of $E_\nu$ is shown in Fig. 4.2.

Convolving the differential cross section with $^8\text{B}$ neutrino flux spectrum provides the expected event rate as a function of $T_e$. RAT performs this convolution through Monte Carlo sampling of the neutrino and cross-section PDFs. The results of the MC sampling are show in Figure 4.3.

Simulated recoil electrons are distributed uniformly throughout the SNO+ AV volume,
with the $\theta_{\text{sun}}$ given by,

$$\cos \theta_{\text{sun}} = \sqrt{\frac{T_e (m_e + E_\nu)^2}{2m_e E_\nu^2 + E_\nu^2 T_e}}.$$  \hfill (4.1)

This equation assumes that the direction of the neutrino is directly from the center of sun. Averaged over many events that is a safe assumption; the additional angular uncertainty introduced by considering the radial neutrino production distribution is negligible.

Once the simulated position, direction, and energy of the recoil electron are determined the electron interactions and photon production and propagation in the detector are simulated by Geant4 \footnote{\cite{77}}. The goal of this simulation is to transform the kinematics of the event to a collection of PMT hits in the simulated detector. In the simulation, once a photon interacts with a PMT photo-cathode, a check is performed to simulate the detection efficiency for that photon, which includes the photo-cathode quantum efficiency and PMT collection efficiency. These efficiencies are determined from both \textit{ex-situ} PMT measurements and \textit{in-situ} calibration measurement. The collection efficiencies are taken to be uniform across the detector. If the photon passes that check, a PMT signal is created in the DAQ simulation. Once all the photons in an event have been simulated and have either exited the detector, been detected, or been absorbed, the DAQ simulation is run on the detected photon signals.

The goal of the DAQ simulation is to transform simulated PMT hits to detector observables, \textit{e.g.} QHS, hit times, trigger information etc. The DAQ simulation accomplishes this by replicating the trigger system of the real SNO+ detector. It starts by creating waveforms for every simulated PMT signal, the size of the PMT signal is drawn from the charge spectrum for each PMT as determined by the PCA calibration. Electronics noise is added to each simulated waveform, then each waveform is compared to a discriminator threshold, the value for which is matched to detector settings. For signals that pass the discriminator threshold a PMT hit is created and N100, N20, ESUMH and ESUML trigger signals are produced. The waveforms for the trigger signals are simulated with electronic noise and dropout, both of which match detector measurements. The trigger signals are
then summed detector wide and compared to “MTCA+” thresholds. If any of the threshold crossing signals are included in the simulated trigger mask each hit’s QHS, QHL, QLX, and threshold crossing time, are calculated from the original PMT waveform. The trigger waveforms are also digitized in a simulated 12-bit digitizer with sampling rate matched to that of the CAEN v1720 digitizer.

![Probability Density vs. Reconstructed Energy](image)

**Figure 4.4:** The distributions of recoil electron truth energy (black) and reconstructed energy (red) for simulated $^8\text{B}$ $\nu_e$ elastic-scatter events. Both histograms are normalized to 1.0.

The result of the DAQ simulation is a dataset that can be used in exactly the same way the detector data can be used. Reconstruction is performed on the simulated detector observables and the reconstructed quantities can be compared to the corresponding MC truth values to assess expected detector performance. The comparison between true energy and reconstructed energy is shown in Fig. 4.4 for the simulated $^8\text{B}$ $\nu_e$ dataset. The probability of an event having a reconstructed energy below approximately 3 MeV is small because events with low energy will often not trigger the detector, or fail to reconstruct at all. Additionally, to prevent the reconstruction process from being too computationally...
expensive only 10% of events with fewer than 15 PMT hits are reconstructed because those events are generally not useful for analysis.

RAT is used to generate two MC datasets, a solar $\nu_e$ dataset and a solar $\nu_\mu$ dataset. No $\nu_\tau$ dataset is generated because the SNO+ detector has no way to discriminate between $\nu_\mu$ and $\nu_\tau$ elastic scatters. So the $\nu_\mu$ dataset can be considered to represent both the $\nu_\mu$ and the $\nu_\tau$ components of the solar neutrino flux. The simulated solar neutrino datasets restrict interactions to the AV volume, further datasets were generated that simulate interactions exclusively in the volume between the AV and PSUP. These external datasets contribute negligibly to the final dataset after all analysis cuts are performed, and so they are in general not considered.

Also, generated are a number of datasets for the expected backgrounds from radioactive isotopes in various detector components. These datasets are not directly used in this analysis, they were used however for assessing the effectiveness of the analysis cuts that are used.

This analysis uses what is referred to as “run-by-run” simulation. This means every detector data run used in the analysis has a simulated run that uses the detector and electronic calibration settings from the run as input to the DAQ simulation, and simulates a length of time and time of day equal to the detector run. The purpose of this is to remove the possibility that changes in detector settings or circumstances will bias any result. This method is not perfect, there exists features of the real detector that are not simulated. For example, it’s known that human activity around the DAQ electronics can generate noise on the front end, this sort of noise source can be difficult to directly observe in the detector data and cannot be simulated. For this reason detector runs that have difficult to simulate conditions are typically not included in the analysis, this is discussed further in Sec. 7.
4.2 Survival Probability Simulation

Figure 4.5: Survival probability for $^8$B solar neutrinos for mixing parameters given in table 1 of Capozzi et. al. [1].
4.2 Survival Probability Simulation

The neutrino survival probability is simulated outside of RAT. The MC $\nu_e$ and $\nu_\mu$ datasets are combined as one of the last stages of this analysis to ensure that MC data does not need to be regenerated for any change in the simulated survival probability.

The survival probability is calculated using a three-flavor adiabatic approximation. The implementation used is the Physics interpretation Sun-Earth Large Mixing Angle Adiabatic Approximation (PSelmaa) used in SNO (45, 80). The survival probability calculation use the GS98 (53) solar abundances and the BS05OP radial production distributions and solar density profile (52). Figure 4.5 shows an example survival probability, using the best fit mixing parameters from a global fit to neutrino oscillation data (1). The uncertainties on those values are also considered as will be discussed in Sec 8.2.
Chapter 5

Reconstruction

A series of reconstruction algorithms are run over all events that pass data cleaning. These algorithms estimate the position, time, direction, and energy of the event. All events are reconstructed under the hypothesis that the PMT hits are from Cherenkov radiation produced by a single electron. Additionally, the reconstruction algorithms use only the hits in a prompt time window to ensure only light that travelled directly from the event origin is used, though each algorithm defines differently window is used. Only hits that are from well calibrated, online PMTs and detector channels are used. The same reconstruction algorithms are used on both simulated and detected events.

The direction ($\vec{d}$), time ($t_0$), and position ($\vec{p}$) are determined by performing a likelihood fit to the time and position of PMT hits. Only hits within a 100 ns window centered around the mode of the raw hit time distribution are used. The algorithm evaluates the likelihood of a hypothesized event position and time by calculating the time residual for each hit PMT,

$$t_{res} = t_{PMT} - t_{transit} - t_0. \quad (5.1)$$

The transit time is calculated by considering the distance the photon would traverse in inner volume water, the external water, and inside the AV acrylic, as it travels from the
hypothesized position to the PMT, accounting for refractions at boundaries. From those
distances the transit time is simply calculated as
\[ t_{\text{transit}} = \frac{d_{\text{AV}}}{v_{\text{AV}}} + \left( \frac{d_{\text{inner}} + d_{\text{external}}}{v_{\text{water}}} \right), \] (5.2)
the group-velocity for 400 nm light is used as the velocity.

![Figure 5.1:](image)

Figure 5.1: (black) Distribution of time residuals for simulated 6 MeV electrons. (red) Simplified timing distribution used for reconstruction. Figure from [74].

The PDF for \( t_{\text{res}} \), \( P(t_{\text{res}}) \), is determined from prior simulation, Fig. 5.1 shows this PDF compared to a full timing simulation. The position and time that minimize the quantity
\[ \sum_{i=0}^{N_{\text{PMT}}} P(t_{\text{res}}) \] (5.3)
is used as the event position and time.
Event direction is determined by evaluating $\theta_{\text{PMT}}$ for each hit where $\theta_{\text{PMT}}$ is defined by,

$$\cos \theta_{\text{PMT}} = \vec{d} \cdot (\vec{p}_{\text{PMT}} - \vec{p}).$$  \hspace{1cm} (5.4)

The likelihood, $P(\theta_{\text{PMT}})$, is determined from a prior simulation of 6 MeV electrons at the center of the detector, Fig 5.2 shows this PDF. The direction that minimizes

$$\sum_{i=0}^{N_{\text{PMT}}} P(\theta_{\text{PMT}})$$  \hspace{1cm} (5.5)

is used as the reconstructed event direction.

**Figure 5.2:** The PDF used by RAT when evaluating the likelihood of a hypothesized direction. The peak at approximately $40^\circ$ corresponds to the Cherenkov cone opening angle.

The kinetic energy of the event is determined after the event position, time and direction are determined. The position and time are used to determine the number of PMT hits that occurred in a prompt 18 ns window. Then the number of photons that would most likely produce that number of PMT hits is estimated using a combination of analytic calculation
and PDFs from prior simulation. A look up table is used to estimate the most likely electron kinetic energy that would produce the determined number of photons. This method of energy reconstruction is called “EnergyRSP”, which stands simply for Energy Response.

![Fractional error of fitted energy for solar $\nu_e$ events with simulated energy above 5 MeV.](image)

**Figure 5.3:** Fractional error of fitted energy for solar $\nu_e$ events with simulated energy above 5 MeV.

The uncertainty in the energy fit is dominated by the uncertainty introduced by the photon-statistics associated with Cherenkov radiation. For electrons the number of detected photons per MeV kinetic energy is approximately 7. Meaning a 5 MeV electron event will on average produce 35 PMT hits with one-sigma statistical fluctuations of roughly $\sqrt{35} \approx 5.9$, or about 17%. Figure 5.3 shows the residuals for fit results on MC simulated solar $\nu_e$ events.
Chapter 6

Calibration

6.1 $^{16}$N

The accuracy of simulated events is evaluated with data taken while a radioactive source was deployed within the detector volume. The data taken with the source deployed is used to assign systematics to reconstructed values and, in the case of the energy reconstruction, determine an empirical correction to improve the agreement between detected and MC simulated events. For this analysis an $^{16}$N source was used. The SNO+ $^{16}$N source was developed for SNO, as their primary energy calibration source (81). The methods and results of SNO+’s $^{16}$N calibration are summarized here but are described in greater detail by (82).

The $^{16}$N source uses a commercial deuterium and tritium generator (DT-generator) to produce gaseous $^{16}$N. The gas is pumped into the deployed source where it can undergo $\beta$-decay to an excited state of $^{16}$O, the $^{16}$O will then de-excite and typically emit a 6.1 MeV gamma particle. Higher or lower energy gammas are emitted at a lower rate, the primary branching ratios for the de-excitation gammas are shown in Fig 6.1.

A small block of plastic scintillator, observed by a PMT, is embedded within the source cannister. The PMT detects the $\beta$ from the initial $^{16}$N → $^{16}$O* decay. That PMT signal is
6.2 Trigger Efficiency

Figure 6.1: The most significant branching ratios for $^{16}\text{N}$ decaying to an excited state of $^{16}\text{O}$. Figure from [81].

used as a tag in the detector DAQ to identify events from the deployed source.

The source position within the AV was varied in a 3-dimensional scan. A 1-dimensional scan was done along the z-axis between the AV and PSUP. Scanning many positions allowed for a position dependent evaluation of systematics and a position dependent correction to the reconstructed energy. The analyses that were done using $^{16}\text{N}$ data to constrain systematics relevant to the solar neutrino analysis are discussed in Sec 8.2.

6.2 Trigger Efficiency

The trigger efficiency for this analysis is defined to be the probability that the detector will trigger on an event as a function of the number of “in-time” hits produced by that event. Here, in-time hits is the effective maximum number of hits as seen by the analog trigger system for an event. For each event the in-time nhit, $\tilde{n}_{100}$, is well estimated by the maximum number of hits in a 100 ns window within the event. Effects from the rise-time of
6.2 Trigger Efficiency

Figure 6.2: (a) The trigger efficiency during the high threshold period for the N100 trigger as measured by nhit monitor, laserball, and TELLIE data, for N100L (left three curves) N100M (center three curves) and N100L (right three curves). (b) The trigger efficiency for the lower threshold data taking period from nhit monitor data.
6.2 Trigger Efficiency

trigger pulses and the limited band-width of the trigger system are applied as corrections to that simple estimate.

The trigger efficiency is estimated in two different ways, using laserball data, and using nhit monitor data. Nhít monitor measurements of the trigger efficiency is discussed in §3.3.2. The laserball can measure the trigger efficiency because the same signal that generates the light for the laserball also produces a trigger for the DAQ. So even if the light produced does not trigger the detector, the laserball signal will ensure that the data is still readout. So data was taken with a very low average light output from the laserball and with that the trigger efficiency curve was measured near threshold. The trigger efficiency curve produced by the nhit monitor and the laserball data disagree by a small, but non-negligible amount, the reason for the disagreement is not well-known, but the differences are taken as a systematic uncertainty. Figure 6.2 shows the trigger efficiency curves for nhit monitor, laserball, and TELLIE data. Here, trigger efficiency is defined to be the probability that a raw-trigger is emitted for a given event.

For data taken after the detector threshold change discussed in Sec 7.1 all methods agree that the trigger is 100% efficient for $\tilde{n}_{100} > 10$; only events with energy significantly below the analysis threshold will have a $\tilde{n}_{100} \leq 10$, and so the discrepant estimates of the trigger efficiency do not have any effect on the solar analysis for the second trigger period. For the first trigger period the trigger was not 100% efficient until $\tilde{n}_{100} \approx 23$, which is much closer to the analysis energy threshold, and therefore uncertainties cannot be neglected. Section 8.2 discusses how the observed trigger efficiency uncertainties were handled in the solar analysis.
Chapter 7

Data Selection

Events are included or removed from the dataset across three stages of selection. First entire runs are either included or removed based upon whether they meet certain criteria for data quality. The events within selected runs are then rejected or approved by a set of low-level cuts that attempt to remove events caused by instrumental backgrounds and other sources of unwanted events. Within events that pass the low-level cuts, hits can be rejected from consideration if they’re deemed unlikely to have originated from light within the detector. Following that, analysis level cuts are applied to the reconstructed quantities for each event. The analysis cuts are designed to maximize the signal efficiency for dataset and minimize the contamination from background sources. Each of these steps are detailed below.

7.1 Dataset

Data for this analysis was taken in SNO+’s initial water-phase data-taking run. Data taking started May 05, 2017 (exactly 4 years after I started my PhD) and ended December 25 the same year. The data taking started with run 100000 and concluded with run 108416, with each run being typically 1 hour long, though not all runs are suitable for analysis. In those
7.2 Run Selection

runs approximately 185 days worth of useable data was taken, with significant pauses for commissioning of ancillary detector components, such as the water circulation systems, and for detector maintenance. Data taking was restarted several months after the end of this dataset, after the AV circulation pipes had been replaced to help reduce backgrounds in the detector.

In the dataset there are a three different periods of data taking that are given special treatment in this analysis to be considered. The first is the hot-spot period of runs; an external water circulation pipe was observed to have introduced backgrounds into the upper portion of the detector, between the PSUP and AV. Analysis cuts were adjusted for the data taken during this period, the cuts will be discussed in Sec 7.5. The elevated background levels lasted from run 100400 to 102048.

The other period that requires special treatment within the analysis is due to significant adjustments to the detector trigger settings done approximately half-way through the dataset. So the dataset is split into a high and low trigger period, the high trigger period being first. The high trigger period started at run 100000 the low trigger period started at run 104613. The adjustment made was to raise the threshold on all channel threshold by a single DAC count. The result was a significant reduction in the observed dropout and noise rate in the detector, which allowed for the trigger threshold to be pushed further down, and an overall increase the detector’s efficiency.

7.2 Run Selection

Runs are rejected from the dataset if they fail to meet certain criteria for data quality and detector stability. It’s first required that all meta-info about a run be created and stored within the RAT Database successfully. Those tables have information about the state of the hardware and DAQ, as well as information about the run including the run type and length. This information is necessary for assessing if the run is capable of being used for
Figure 7.1: N20 dropout in the high (a) and low (b) trigger period, and the average dropout measured across all runs in the dataset (c); the various “jumps” in the dropout are generally due to bad fits.
7.2 Run Selection

a physics analysis. It’s further necessary that all meta-info be available so that the run can be simulated. It’s very rare for the meta-info for a run not to be generated and stored properly, so this check has almost no effect on the dataset.

It’s required that the run type of the run be “PHYSICS”, this indicates that the DAQ settings were not changed at any point in the run and no external sources of events were present, and that the thresholds for triggering were set at a level deemed sufficient for most physics analyses. Further checks are done that require all electronics crates have high voltage on their PMTs throughout the run. It’s also required that all channels that are at high voltage are capable of reading out data, and that all crates are participating in the trigger sums. A number of separate checks are performed to ensure that all necessary DAQ processes were running well throughout the run, to ensure that the data taken during the run was not interrupted by a lapse in the DAQ.

Beyond checks on the detector state and stability, a number of checks are placed on the data taken within the run. Most of these checks are placed on the rate of certain events within the detector. It’s required that the average ESUMH and N100L trigger rate be greater than 5Hz and that the total trigger rate be less than 7kHz. These checks ensure that the data taken during the run triggered the detector at a rate consistent with standard running, during which the typical trigger rate is near 1kHz. Similarly, it’s required that fewer than 15% of all events be retrigger events (events that fall within 420 ns of the previous event). At a nominal rate of 1kHz it’s very unlikely for two events to be within 420 ns by chance, so a high rate of retrigger events might indicate an abnormally high level of noise in the trigger system or light in the detector. There are however events that can occur in the detector, such as followers after a cosmic muon, that can produce re-trigger events. The cut threshold is designed to allow for retriggers from natural source but still flag detector abnormalities. These checks all ensure that the data in a run is likely to be useful for a physics analysis, but are loose enough not to bias the dataset in a way that might influence results.
7.3 Livetime

While most event selection removes individual events based upon whether they pass or fail certain criteria, some cuts remove all events that occur for a period before or after some criteria is met. These cuts are said to introduce a deadtime into the dataset. The most significant example of this is the muon follower data cleaning cut, which cuts all events for 30 seconds after every muon interaction in the detector. The livetime for each run is then defined as

\[ t_{\text{live}} = t_{\text{run}} - t_{\text{dead}}, \]

where \( t_{\text{run}} \) is the time between the first and last valid event within a run and \( t_{\text{dead}} \) is the sum of all deadtimes introduced into that run by cuts. The livetime is used to calculate the total exposure represented by the dataset. For simulated events many of the effects that necessitate deadtime are not simulated, so no deadtimes are added into the simulated runs. The livetime represented by the dataset is 120.2 days, after subtracting the calculated deadtimes the effective livetime 114.7 days.

7.4 Data Cleaning

There are a number of instrumental effects that can cause an event to be recorded by the detector, these events typically have some sort of distinguishing feature or features that set them apart from events that originate from particle interactions within the detector. A number of algorithms and cuts have been designed to identify and remove these events from the dataset. These algorithms are said to “clean” the data by removing events of instrumental origin. This definition is extended to include removing hits within an event that are likely of instrumental origin as well.

The primary type of instrumental event that must be removed is “flashers” and “shark-fin” events. Both of these result from charge build-up on the PMT-base causing a spark. For a flasher event the light from the spark escapes through the PMT face and illuminates
the PMTs on the other side of the detector. Flashers occur at a rate of a few per minute. A shark-fin is similar but the spark is either small enough or located in a position such that the light does not escape the PMT. In both types of events the PMT in which the spark occurs will have a very high-charge hit, and the channels next to it on the FEC will have low-charge hits from electronic pickup. For shark-fin event no other channels will be hit, except possibly by an accidental coincidence; for a flasher hit a number of hits will occur from the light that escaped the PMT. Since the number of PMT hits that occur in a flasher event can vary significantly, anywhere between tens of hits and hundreds, they can reconstruct to a wide range of energies and possibly contaminate a signal region. Many data cleaning cuts are designed to ensure that all flasher events are identified and removed from the dataset.

Within an event hits that are deemed unlikely to have come from a photon interacting with a PMT are removed from the analysis through hit cleaning. For this analysis the only sort of hits that were removed were those identified as coming from cross-talk between adjacent channels in the detector. Hits from cross-talk arise from stray capacitve coupling between adjacent, or nearby, channels on a single daughter board. Typically, the noise from cross-talk will only be large enough to cause a hit on adjacent channels if the original signal is relatively large. The cross-talk hits will usually be especially low in charge because they’re the result of bi-polar noise, rather than a PMT signal, and the cross-talk hit will always show up after the original hit. These criteria are codified as a cut on any hits that show up within six channels from a hit that has a pedestal subtracted QHS greater than 50 ADC counts. Of those hits if it has a pedestal subtracted QHS between 10 and −30 and are between 9 and 25 ns after the high charge hit, then the hit is flagged a cross-talk hit and removed from the analysis.

Events of instrumental origin are not well modeled within our simulation, so simulation is not used for evaluating the efficiencies and or false-positive rates of data cleaning cuts. Instead, a data-driven approach is used that relies primarily on calibration data from the
\( ^{16}N \) source, that analysis is detailed in (83). The basic approach is to use tagged \( ^{16}N \) events as a source of known non-instrumental events, and evaluate what fraction of the time those events are identified as instrumentals, this provides an estimate of the data-cleaning false-positive rate. The results of this analysis estimated a 1.2\% of signal events would be cut by data-cleaning.

An estimate of the signal contamination was performed using a method developed by SNO (84) and applied to the dataset (83); the number of instrumental events leaked into the signal region was estimated to be roughly 0.5 events over the entire dataset. However, a contamination estimate is not an input to the solar analysis, so that value is not used beyond a check that the instrumental background is reduced to an acceptable level.

The data cleaning cuts that were used in this analysis are given in Appendix A. I also detail there the cuts used that I developed.

### 7.5 Analysis Cuts

The dataset of events passing all data cleaning cuts is further reduced by requiring all events pass cuts on reconstructed quantities. The cuts are designed to minimize the number of events in the dataset from non-solar interactions.

Necessarily, the first of these cuts is the requirement that the reconstruction fits produce to a valid position, time and energy. The reconstruction algorithms can fail to converge if an event occurs in an optically complicated region of the detector, \textit{e.g.} near the detector neck. The fitting algorithms rely on the assumption that the majority of the produced light will travel directly from the event vertex to PMT array. For events in optically complicated regions this assumption is not a good one. These regions modelled in simulation, so the Monte Carlo simulation estimate of the efficiency of reconstruction to produce valid fits for solar neutrino events is used.

A cut, called the “trigger efficiency” cut, is placed on the number of in-time nhits in each event. This cut ensures the dataset occupies a region where the detector trigger efficiency
is well understood and near 100%. This cut ensures the analysis is minimally effected by uncertainties associated with the detector’s trigger.

As mentioned in Sec. 7.1, the detector trigger threshold were adjusted part way through data-taking. The in-time nhit cut was adjusted to account for this. For the first trigger period all events were required to have an in-time Nhit greater than or equal to 23; for the second trigger period this threshold was reduced to 10. This cut is similar to an energy cut because nhit is the best energy estimator for an event. But, as discussed below, events passing the analysis energy cut are very unlikely to fail the trigger efficiency cut.

The next analysis cut is a fiducial volume (FV) cut that requires all events be within 5.3 m of the center of the detector. This cut is designed to reduce the background from radioactive decays within the AV or from the water outside the AV. During the hot-spot time period the FV cut was modified to require any events in the top half of the detector (z > 0) fall within a radius of 4.2 m, events in the bottom half of the detector were still subject to the standard 5.3 m FV cut. The more restrictive cut was in place for 13% of the dataset livetime.

The time residual, defined in equation 5.1 for a PMT hit is an extremely useful quantity because in general light that travels directly from an interaction will have a very small time residual. Light that is produced by another source, or reflects off of a detector component between production and detection will have a larger time residual.

The fraction of hits that satisfy

\[-2.5 < t_{res} < 5\]

is known as the “in-time ratio” (ITR).

The quantity \(\beta_{14}\) is used to quantify how isotropic the hits in an event is. It is defined as

\[\beta_{14} = \beta_1 + 4\beta_4\]
where

\[ \beta_\ell = \frac{2}{N(N-1)} \sum_{i=1}^{N-1} \sum_{j=i+1}^{N} P_\ell(\cos \theta_{ij}). \]  

(7.4)

\( P_\ell \) is the \( \ell \)th Legendre Polynomial and \( \theta_{ij} \) is the angle subtended by the vectors pointing from the reconstructed position of the event to the \( i \)th and \( j \)th hit PMT.

Cuts are placed on the ITR and \( \beta_{14} \) value for each event. These cuts aim to remove events that have an ITR or \( \beta_{14} \) that is inconsistent with the event originating from Cherenkov light produced by a solar neutrino event. The ITR of events is required to be greater than 55% and the \( \beta_{14} \) value must be between –0.12 and 0.95. These cuts are similar in purpose to the data cleaning cuts, as they attempt to remove events that are produced by sources other than Cherenkov light within the detector.

The final analysis cut is on the reconstructed kinetic energy of each event. The energy region for this analysis is \( 5.0 < T_e < 15.0 \) MeV. This region is chosen to minimize contamination from atmospheric neutrino interactions, and radioactive decays within the detector. Additionally, the only solar neutrino flux that is significant across this energy range are the neutrinos from the \( ^8B \) solar reaction. Neutrinos from the \( hep \) interaction also fall within the same energy range, however their flux is expected to be much lower than that of the \( ^8B \) neutrino flux that their presence can be largely neglected.

In principle the lower energy threshold could be lowered or removed to increase the efficiency for solar neutrino events, however, the rate of backgrounds from radioactive decays increases rapidly at lower energies. So no additional sensitivity to the solar neutrino interaction rate would be gained with a lower energy threshold. The 5 MeV threshold was chosen as the lowest energy from which a solar signal could still be resolved.
Chapter 8

Analysis

8.1 Signal Extraction

Once an MC simulated dataset and a detector dataset are selected, the analysis of those events is performed. The first steps of the analysis is to bin all events in a two-dimensional histogram of reconstructed energy and $\cos \theta_{\text{sun}}$. Events are distributed across $N_\theta$ equal width bins in $\cos \theta_{\text{sun}}$ from $-1$ to 1 and $N_E$ bins in energy from 5.0 to 15.0 MeV. For this analysis $N_\theta$ is 40 and $N_E$ is 6. From 5.0 to 10.0 MeV 5 bins of width 1 MeV are used and a single bin from 10.0 to 15.0 MeV is used. Simulated and detected events are placed into separate histograms.

Simulated $\nu_e$ and $\nu_\mu$ events are histogrammed separately, but given different weights in their respective histogram according to the expected survival probability for each event. The weight for a $\nu_e$ event with neutrino energy $E_\nu$ is given by,

$$w_e = P_{ee}(E_\nu),$$  \hspace{1cm} (8.1)

and the weight for a $\nu_\mu$ event is given by,

$$w_\mu = P_{e\mu} = 1 - P_{ee}(E_\nu).$$  \hspace{1cm} (8.2)
8.1 Signal Extraction

As mentioned in Sec 4, the effective flux of the solar simulations are much larger than the expected flux of the detected dataset; this scaled flux is the same as if the simulated dataset had a much higher livetime but a standard flux value. So the simulated histograms are scaled to by \( \frac{t_{\text{live}}}{t_{\text{sim}}} \) to make the effective simulated livetime match the livetime of the detector dataset. An additional scaling is done to the simulated histograms to account for the data cleaning, this is needed because data cleaning is not applied to simulated events. The data cleaning rejection rate was determined to be 1.2%, so the simulated histograms are scaled by a factor of 0.988.

Once these scales are applied the \( \nu_e \) and \( \nu_\mu \) histograms respectively represent the expected distribution and event rate for the charged current and neutral current interactions in the dataset for the nominal \( ^8 \text{B} \) solar neutrino flux used in simulation. The \( \nu_e \) and \( \nu_\mu \) histograms are then combined by simply adding their bin contents together to get the expected mixed flavor elastic scattering interaction rate as a function of \( T_e \) and \( \cos \theta_{\text{sun}} \). Applying additional scaling to this combined histogram is done to represent a hypothesized scaling to the overall solar neutrino flux. So the rate of solar neutrino events is given by

\[
R_\nu(\cos \theta_{\text{sun}}, T_e) = S \phi(R) \tag{8.3}
\]

Equation (8.3) is modified slightly to include a parameter related to the detector angular resolution, \( \delta_\theta \),

\[
R_\nu(\cos \theta_{\text{sun}}, T_e) \rightarrow R_\nu(\cos \theta_{\text{sun}}, T_e, \delta_\theta). \tag{8.4}
\]

This modification is discussed more in Sec 8.2.3.

No simulation or measurements were done for the expected backgrounds for this analysis, so a simple background model is adopted. It is assumed that the direction of any background event will be uncorrelated with the position of the sun, this is what makes \( \cos \theta_{\text{sun}} \) such a useful variable in this analysis. The distribution of background events is given by

\[
R_B(\cos \theta_{\text{sun}}, T_e) = R_B(T_e) = \frac{1}{N_\theta} n_B(T_e) \tag{8.5}
\]

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Where \( n_b(T_e) \) is number of background events in the histogram energy bins corresponding to the energy \( T_e \). The number of background events in each energy bin are not known \textit{a priori} and are treated as a nuisance parameter in the remainder of the analysis.

The total expected events in each bin can be expressed by

\[
R(\cos \theta_{\text{sun}}, T_e) = R_B(T_e) + R_\nu(\cos \theta_{\text{sun}}, T_e, \delta_\theta) .
\] (8.6)

The unknown parameters of this rate are the 6 background rates and the solar rate and \( \delta_\theta \). A fit to data is performed for equation (8.6) to extract those parameters. Goodness-of-fit is evaluated using a likelihood given by

\[
\mathcal{L}(S, B, \delta_\theta | n, \mu_\theta, \sigma_\theta) = N(\delta_\theta, \mu_\theta, \sigma_\theta) \prod_{j=0}^{N_e} \prod_{i=0}^{N_\theta} \text{Pois} \left( n_{ij}, B_j + S p_{ij}(\delta_\theta) \right) .
\] (8.7)

The parameters \( \mu_\theta \) and \( \sigma_\theta \) are respectively the best fit and the constraint on \( \delta_\theta \) from the \(^{16}\text{N} \) source analysis. \text{Pois} \( (k, \lambda) \) is the value of the Poisson distribution at the value \( k \) for a rate parameter \( \lambda \).

Equation (8.7) can be modified to fit for the solar rate in individual energy bins, rather than constraining the solar rate to be the same across all energy bins. This is done by replacing the product over energy bins, \( \prod_{j=0}^{N_e} \), with the selection of \( j = 0, 1, \text{ etc.} \). Fitting for a solar rate in each bin allows for a spectral measurement of the solar neutrino flux, as opposed to an integrated flux measurement.

### 8.2 Systematics

Systematics associated with event reconstruction, the mixing parameters, and trigger efficiency are considered for this analysis. The event reconstruction systematics are uncertainties on the energy reconstruction scale and resolution, position reconstruction resolution.
and scale, and the resolution of the direction reconstruction. The position and energy systematics must be known to assess uncertainties on the efficiencies for the fiducial volume cut and for what fraction of solar neutrino interactions will fall into each energy bin. Since the solar event rate is extracted from a fit to the angular distribution of events with respect to the solar direction, it’s important that the detector’s angular resolution be well constrained. The uncertainties for these reconstructed quantities are determined from analysis of $^{16}$N data. All systematics are treated in the same, or a similar, way, to propagate their effect to the flux result. First the systematic uncertainty on a reconstructed quantity (e.g. energy, position etc.) is determined from an analysis of the $^{16}$N data. Then, the relevant reconstructed Monte Carlo quantity is modified according to the one-sigma systematic uncertainty. From this a 2D histogram in energy and $\cos \theta_{sun}$ is produced using the modified event quantities, applying the standard cuts. Finally, the flux result is found by performing a fit to data using the systematically adjusted histogram as input. The difference between the best fit value found using unmodified event quantities and using the modified quantities is taken to be the one-sigma systematic uncertainty. How each variable is modified, and any deviations from this process of propagating systematics is detailed below. All systematics are treated as uncorrelated; variables are modified according to only one systematic at a time.

### 8.2.1 Energy Systematics

The detector resolution $\sigma_E$ and relative energy scale $\delta_E$ are determined from the $^{16}$N detected energy spectrum. The detected energy spectrum is modeled by $P(T_e)$ in electron equivalent energy, and is given by $P_{source}(T_e)$ convolved with a normalized Gaussian distribution.

$$P(T_e) = N \int P_{source}(T_e') \frac{1}{\sqrt{2\pi}\sigma_E} e^{-\frac{((1 + \delta_E)T_e - T_e')^2}{2\sigma_E^2}} dT_e'. \quad (8.8)$$

$P_{source}(T_e)$ represents the distribution of deposited energy in the detector from the $^{16}$N source. Since the $^{16}$N emits gammas into the detector the mapping between gamma energy
and electron equivalent energy is done by finding the electron energy that can produce the same number of Cherenkov photons as each gamma; this is not a one-to-one mapping because the same electron or gamma energy will not always produce the same number of photons. The mapping is determined from simulation and is shown in Fig. 8.1. The gamma to electron energy mapping is then applied to the simulated $^{16}\text{N}$ gamma energy spectrum to determine $P_{\text{source}}(T_e)$.

![Figure 8.1: The map between electron energy and expected Cherenkov photons production. Used for the energy calibration to map between gamma energy and equivalent electron energy. This is the same PDF used by EnergyRSP to estimate energy from the observed PMT hits.](image)

The values for $\sigma_E$ and $\delta_E$ are extracted from (8.8) by performing a fit to the reconstructed $^{16}\text{N}$ energy spectrum. The fit is done to both simulated $^{16}\text{N}$ data and to detector data, each determining their own values for $\sigma_E$ and $\delta_E$. As is show in Fig 8.2, this fit is only performed over the energy range 3.25 to 9.6 MeV, at energies outside this range difficult to model source-container effects dominate the energy spectrum. It’s worth noting that $\sigma_E$ represents only the resolution provided by detector effects, resolution from effects such as photon statistics are accounted for in $P_{\text{source}}(T_e)$.

Values for $\sigma_E$ and $\delta_E$ are extracted for data taken, or simulated, with the $^{16}\text{N}$ source.
8.2 Systematics

Figure 8.2: (a) The comparison between reconstructed energy for \(^{16}\text{N}\) data and Monte Carlo. (b) Fit of Eqn. (8.8) to distribution of reconstructed energies for detected central \(^{16}\text{N}\) events.

at many position, allowing for a position dependent determination of the energy scale and resolution. Fitting to both simulated and to detected data allows for a correction to be created that can make the two datasets match better, however the data used to create the correction cannot then be used to determine systematics. So the \(^{16}\text{N}\) data was split into two datasets, one for determining what correction should be applied to simulation, the other for evaluating the remaining uncertainties after the correction is applied.

The data for the correction is further divided into position bins in \(z\) and \(\rho\), where \(\rho = \sqrt{x^2 + y^2}\). The choice of binning is motivated by the symmetry of the detector, the detector is very symmetric for an interchange of \(x\) and \(y\) or \(x, y \rightarrow -x, -y\). There exists, however, significant asymmetries along the \(z\) axis from the detector neck and from the rope-net along the top of the AV. The data is divided into 4-bins along the \(\rho\) direction each 200 cm long and bins of 57 cm height along the \(z\) axis. The number of bins along the \(z\)-axis varies for each slice in \(\rho\) because data was primarily taken within the AV. Figure 8.3 shows position dependence of the fits for \(\delta_E\) and \(\sigma_E\) in each bin for both simulated and detected events.
8.2 Systematics

Figure 8.3: The best fit energy scale before for detected (a) and Monte Carlo simulated (b) $^{16}$N events as a function of position. And the best fit energy resolution parameter for detected (c) and simulated (d) $^{16}$N events
8.2 Systematics

Figure 8.4: The energy scale non-uniformity correction for data (a) and MC simulation (b). The measured energy scale as a function of position after the application of the non-uniformity correction for data (c) and MC simulation (d).
8.2 Systematics

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data</td>
<td>2.53e-2</td>
<td>1.48e-9</td>
<td>-5.44e-6</td>
<td>2.14e-9</td>
<td>6.49e-13</td>
</tr>
<tr>
<td>Simulation</td>
<td>3.33e-2</td>
<td>9.48e-10</td>
<td>3.77e-6</td>
<td>4.46e-10</td>
<td>1.43e-13</td>
</tr>
</tbody>
</table>

Table 8.1: Best fit values for (8.9) for simulated and detected data, determined using units of mm for $z$ and $\rho$.

Variations in $\delta_E$ along $z$ and $\rho$ were modeled by a polynomial given by,

$$
\delta_E(\rho^2, z) = A + [(1 + B\rho^2)(1 + Cz + Dz^2 + Ez^3) - 1].
$$

(8.9)

Values for $A$, $B$, $C$, $D$, and $E$ are extracted from a fit to the observed spatial variation of $\delta_E$ for simulation and data and are given in table 8.1. The reconstructed energies of simulated and detected events are then corrected according to (8.9) by their respective best fit values. Figure 8.4 shows the correction as a function of position in the detector and the energy scale measured across the detector after the application of the correction. The energy resolution is evaluated as a function of position but no correction is determined from it.

After the correction is applied to remaining half of the calibration dataset $\sigma_E$ and $\delta_E$ are determined once more as a function of position. The bin-by-bin differences in $\sigma_E$ and $\delta_E$ between simulated and detected data are taken as the systematic uncertainty for those parameters, with additional fit uncertainties added in quadrature. Averaging the bin-by-bin systematic uncertainty over the detector volume relevant for the solar analysis yields a remaining uncertainty on the energy scale of $\delta_E = 1 \pm 2.0\%$ and $\delta_{\sigma} = +0.018$, $-0.016$ to be the uncertainty on the energy resolution.

To create the systematically modified energy resolution histograms for the solar analysis the reconstructed energy of each MC simulated event is mapped to a normalized Gaussian distribution with a mean value of the event’s energy and a variance given by

$$
\sigma^2 = \sigma_E^2 \left( (1 + \delta_{\sigma})^2 - 1 \right).
$$

(8.10)
This process of mapping a single energy value to a Gaussian distribution is referred to as “smearing”. Here $\sigma_E$ is given by $\sqrt{E}$ to match the functional form used in Eqn 8.8. The idea behind this smearing is to compensate for the possibility that our Monte Carlo simulation could have a systematically smaller energy resolution than occurs in real data. So by applying a smearing the Monte Carlo energy resolution is artificially deteriorated, and the uncertainty on the resolution is accounted for. A similar process does not exist to account for the possibility that the MC simulation has a poorer energy resolution than data taken from the real detector; there’s no way to “un-smear” the reconstructed MC event energy. To account the effect of an over-estimated energy resolution the error on the result is assumed to be symmetric. As a penalty for this assumption the larger uncertainty between the positive and negative uncertainty on the energy resolution is used. For each event, and for each energy bin of the systematically modified histogram is filled by an amount equal to the integral of the event’s Gaussian across that energy bin. The Gaussian generated from each event is integrated across each energy bin.

Systematically varied histograms for the energy scale is generated by simply modifying the reconstructed kinetic energy of each event according to

$$T'_{e} = \delta_{E} T_{e}. \quad (8.11)$$

This leads to a straightforward shift of $\pm 2.0\%$ for all energies. At all points in the analysis afterwards $T'_{e}$ is used instead of $T_{e}$ and the systematic histogram is generated the same way the standard analysis histograms are.

### 8.2.2 Fiducial Volume

Uncertainty on the fiducial volume comes from systematics associated with the position reconstruction. If the position reconstruction is more likely to pull an event towards the middle of the detector in MC simulation than in data, it will result in an over prediction of the number of events that will pass the FV cut. This possibility is accounted for by shifting
The reconstructed position of simulated events according to the uncertainty, the fiducial volume cut is applied to those shifted positions. Shifting the events results in modified PDFs, those PDFs are used in the fit for the solar event rate, the difference between the best fit value extracted with the modified PDFs and the best fit value from the standard PDFs is taken to be the systematic uncertainty.

The systematic uncertainty on position reconstruction is evaluated using $^{16}\text{N}$ data and simulation. The difference between the source position and the reconstructed position of each event is determined and histogrammed. A fit to that distribution is performed using a model of a Gaussian distribution with exponential tails convolved with a distribution for the first gamma interaction distance. The equation for this is given by,

$$ P(x) = A \cdot \left[ \left( \frac{1 - \alpha}{\sqrt{2\pi}\sigma} e^{-\frac{(x-\mu)^2}{2\sigma^2}} + \frac{\alpha}{2\tau} e^{-\frac{|x-\mu|}{\tau}} \right) \ast P_{\gamma}(x) \right]. \quad (8.12) $$

Where $\mu$ and $\sigma$ are respectively the center and width of the Gaussian, $\tau$ represents the decay rate for the exponential tails, and $\alpha$ represents the relative strength of the exponential vs. the Gaussian; $P_{\gamma}(x)$ is distribution of distance travelled by an $^{16}\text{N}$ gamma before it’s first interaction, it is determined from a separate MC simulation. Finally, $A$ is an overall normalization to account for the number of events included in the distribution. The Gaussian and exponential portion of (8.12) represents the spread introduced by the detector and position reconstruction, the $P_{\gamma}$ term represents the intrinsic spread in interaction positions from the source itself. The distribution used for $P_{\gamma}$ in show in Fig 8.5. Figure 8.6 shows a comparison between data and Monte Carlo simulation for reconstructed position and an example fit to the data for a central $^{16}\text{N}$ dataset.

With this model three types of position uncertainties are considered, a shift uncertainty, a resolution uncertainty, and a scale uncertainty. Here a position shift is the value for $\mu$ in equation (8.12) averaged over the entire detector volume, $\langle \mu \rangle$; the position shift systematic then is the difference in $\langle \mu \rangle$ from MC simulation and as determined by detector data. Rather than averaging over all source positions $\langle \mu \rangle$ is determined averaging over scans along the
8.2 Systematics

Figure 8.5: The distribution of first interaction distances for gamma ray produced in the $^{16}$N source. Produced from MC simulation and used in Eqn. (8.12).

Figure 8.6: (a) Reconstructed position of $^{16}$N detected events and MC simulated events for a central $^{16}$N run. (b) An example fit to detected events, also for a central $^{16}$N run.
8.2 Systematics

| $\langle \mu \rangle$ Systematic Uncertainty (mm) |
|---------------------|-----------------|
| x                   | +16.4, -18.2    |
| y                   | +22.3, -19.2    |
| z                   | +38.4, -16.7    |

Table 8.2: Position shift systematic uncertainties

| $\langle \sigma \rangle$ Systematic Uncertainty (mm) |
|---------------------|----------------|
| x                   | 104.0           |
| y                   | 98.2            |
| z                   | 106.2           |

Table 8.3: Position resolution systematic uncertainties

$x$, $y$ and $z$ axis and so a position shift for each axial direction is determined. Only source positions along each axis are used to avoid possible correlations in each direction’s position shift. The resulting systematic uncertainties along each axial direction are given in table 8.2.

The position resolution systematics is evaluated in a similar way as the position shift systematic, comparing values for $\sigma^2$ in equation (8.12) instead of $\mu$, but otherwise following the same procedure. Table 8.3 gives the extracted position resolution systematics uncertainties in mm. The uncertainties are given as one-sided because a resolution uncertainty, unlike the shift uncertainty, can only be applied to MC simulation by applying addition smearing.

The final position systematic considered is the position scale uncertainty, which represents any position dependent shift in $\mu$ between simulation and data. Unlike the previous two uncertainties this systematic can effect the number of events that would be predicted to fall within a volume if the events are distributed uniformly throughout space. For this reason the position scale systematic is sometimes called the fiducial volume systematic.
### 8.2 Systematics

<table>
<thead>
<tr>
<th></th>
<th>Position Scale Systematic Uncertainty (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>x</td>
<td>+0.91, -1.01</td>
</tr>
<tr>
<td>y</td>
<td>+0.92, -1.02</td>
</tr>
<tr>
<td>z</td>
<td>+0.91, -0.99</td>
</tr>
</tbody>
</table>

**Table 8.4:** Position scale systematic uncertainties

The position scale for simulation and data is determined by fitting the values of $\mu$ as a function of position, along each axis, with a linear function. The best fit slope for that line gives the position dependence of the position shift. The value for that shift is defined to be zero at the center of the detector. The position scale systematic can be thought of as the positional divergence introduced by the MC simulation compared to the detector data. **Table 8.4** gives the position scale systematic uncertainty along each axis.

These uncertainties, the position shift, scale, and resolution are applied by modifying the positions of simulated events. For the shift uncertainty the shift uncertainties are simply added to the reconstructed position. In general the shift uncertainty has no effect at all on the final result because just as many events will be shifted into the fiducial volume as are shifted out.

For the resolution a three-dimensional Gaussian centered at the reconstructed position is created, with uncorrelated widths given by the position resolution uncertainty along each axis. From that Gaussian 1000 samples are drawn, and the fraction of those samples that fall within the fiducial volume is taken to be the weight that event adds to the histogram for this systematic. Similar to the position shift uncertainty, the position resolution uncertainty tends to bring as many events into the fiducial volume as it brings out. So the position resolution systematic has a negligible effect on the final result.

The position scale systematic shifts the simulated event positions by

$$\vec{p}' = (1 + \text{scale})\vec{p},$$

(8.13)
where the vector \( \vec{\text{scale}} \) is the position scale systematic uncertainties. The fiducial volume cut is applied to the scaled position to generate a systematically modified histogram.

### 8.2.3 Direction

![Cartoon depicting “true” angle reconstruction with \(^{16}\text{N}\) source. The blue direction is the formed by reconstructed position of the interaction and the known position of the source. That direction is compared to the reconstructed direction (black arrow).](image)

**Figure 8.7:** Cartoon depicting “true” angle reconstruction with \(^{16}\text{N}\) source. The blue direction is formed by reconstructed position of the interaction and the known position of the source. That direction is compared to the reconstructed direction (black arrow).

Like position and energy, the direction reconstruction systematics are determined using data from the \(^{16}\text{N}\) source. For each \(^{16}\text{N}\) event the direction of the gamma is estimated as co-linear with the vector from the source position to the reconstructed event position. The dot product of that vector with the reconstructed event direction is taken, this gives the
value $\cos \theta$ for that event,

$$
\cos \theta = \frac{\vec{p}_{\text{fit}} - \vec{p}_{\text{source}}}{|\vec{p}_{\text{fit}} - \vec{p}_{\text{source}}|} \cdot \vec{d}_{\text{fit}}.
$$

(8.14)

Figure 8.7 schematically shows how $\cos \theta$ is determined.

A fit is then performed to the distribution of events in $\cos \theta$ using the model of a double exponential,

$$
P(\cos \theta) = \alpha \beta_s e^{\beta_s (\cos \theta - 1)} + (1 - \alpha) \beta_l e^{\beta_l (\cos \theta - 1)}.
$$

(8.15)

Where $\beta_s$ and $\beta_l$ represent the “short” and “long” decay constants for the two exponentials, and $\alpha$ represents the relative strength of the short exponential vs the long one. This model was developed by the SNO experiment (85), the two decay constants represent the single and multiple-scattering contributions. Here Eqn. (8.15) is used here simply as an empirical method to parameterize the distribution of events in $\cos \theta$. Figure 8.8 shows a comparison between data and Monte Carlo for a central $^{16}$N run and an example fit to the data.

![Figure 8.8: (a) The comparison between data and Monte Carlo simulation for a central $^{16}$N run. (b) The fit of Eqn. (8.15) to data for a central $^{16}$N run.](image)

shown in (86) that the systematic uncertainties on the parameters derived from (8.15) can
Table 8.5: The relative difference in best fit values of Eqn (8.15) from $^{16}$N data and simulation. Here $\mu$ represents the best fit value and $\sigma$ is the uncertainty on that value.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>$\mu$ [%]</th>
<th>$\sigma$ [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta(\alpha)<em>{rel} \equiv (\alpha</em>{data} - \alpha_{MC})/\alpha_{MC}$</td>
<td>7.1</td>
<td>10.97</td>
</tr>
<tr>
<td>$\Delta(\beta_s)<em>{rel} \equiv (\beta</em>{data} - \beta_{s,MC})/\alpha_{MC}$</td>
<td>-6.9</td>
<td>11.6</td>
</tr>
<tr>
<td>$\Delta(\beta_l)<em>{rel} \equiv (\beta</em>{l, data} - \beta_{l, MC})/\beta_{s,MC}$</td>
<td>-2.9</td>
<td>10.4</td>
</tr>
</tbody>
</table>

be transformed to a shift in $\cos \theta$ given by,

$$\cos \theta' = 1 + (\cos \theta - 1)(1 + \delta_\theta), \quad (8.16)$$

where $\delta_\theta$ is the relative systematic uncertainty of $\beta_s$ and $\beta_l$. These relative uncertainties are given in Tbl. 8.5. Transformed this way the systematic uncertainty for the direction reconstruction is given by

$$\delta_\theta = +0.08, -0.13,$$

the full derivation of these uncertainties can be found in Ref (82).

The angular resolution uncertainty is propagated to the final result differently from the other uncertainties because the distribution of events in $\cos \theta_{sun}$ is directly related to the direction resolution. To minimize the impact the angular resolution has on the result it is used as one of the parameters in the fit to the $\cos \theta_{sun}$ solar neutrino distribution, and constrained by the results of the $^{16}$N analysis.

The angular resolution systematic is applied using Eqn. (8.16). Where $\theta$ is the angle between the true event direction and the reconstructed event direction, and $\delta$ is the angular resolution systematic uncertainty. Using Eqn. (8.16) has the unfortunate downside producing unphysical values for $\cos \theta'$ for values of $\cos \theta \approx -1$. For values of $\cos \theta'$ below $-1$ the value is instead replaced with a random value drawn from a uniform distribution $[-1, 1]$. The logic behind this choice is that when an event reconstructs with a direction that’s nearly 180° from the correct value, then the reconstruction has likely failed to such a
degree that the reconstructed values are uncorrelated with the true values, and so drawing from a uniform random distribution preserves that uncorrelated nature without adding any additional bias.

Once the systematically varied value for $\cos \theta$ is determined, the new angle needs to be transformed into a corresponding direction vector for the particle. To do this first a vector that is normal to the plane spanned by the reconstructed and true direction vector is found by taking the cross-product between those vectors,

$$\vec{d}_{\text{norm}} = \vec{d}_{\text{true}} \times \vec{d}_{\text{recon}}.$$  \hfill (8.17)

Then $\vec{d}_{\text{recon}}$ is rotated around $\vec{d}_{\text{norm}}$ such that the rotated vector $\vec{d}'_{\text{recon}}$ now has an angle of $\theta'$. The direction $\vec{d}'_{\text{recon}}$ is then used in the analysis to generate event distributions in $\cos \theta_{\text{sun}}$.

Following this procedure PDFs for $\cos \theta_{\text{sun}}$ are generated for many values of $\delta \theta$, producing $P(\cos \theta_{\text{sun}}, \delta \theta)$. The constraint on $\delta \theta$ produced by the $^{16}\text{N}$ analysis are included in this two-dimensional PDF. $P(\cos \theta_{\text{sun}}, \delta \theta)$ is used in the remainder of the analysis treating $\delta \theta$ as a nuisance parameter.

### 8.2.4 Mixing Parameters

The central values and uncertainties of the neutrino mixing parameters, $\Delta m^2_{21}$, $\theta_{12}$ and $\theta_{13}$ is taken from Ref. (1). The values and uncertainties for the mixing parameters are summarized in Table 8.6. A survival probability curve is generated for each of the mixing parameters shifted by their positive and negative one-sigma uncertainty. These systematically adjusted PDFs are used in the analysis replacing the standard survival probability curve to propagate the uncertainties to the flux result.
Table 8.6: A summary of the mixing parameters and their uncertainties as used in propagation of systematic uncertainties. Values from Ref (1).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Uncertainty</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta m_{21}^2$</td>
<td>$7.37 \times 10^{-5}$ MeV/c$^2$</td>
<td>+0.17, -0.16</td>
</tr>
<tr>
<td>$\theta_{12}$</td>
<td>33.02°</td>
<td>0.537, -0.455</td>
</tr>
<tr>
<td>$\theta_{13}$</td>
<td>8.41°</td>
<td>–</td>
</tr>
<tr>
<td>$\Delta m_{31}^2$</td>
<td>$2.5 \times 10^{-3}$ MeV/c$^2$</td>
<td>–</td>
</tr>
</tbody>
</table>

Table 8.7: Weight given to events with values for $n_{100}$ above cut value for the high threshold period, but where the nhit-monitor and laserball disagree.

8.2.5 Trigger Efficiency

The uncertainty on the trigger efficiency is described in Sec 6.2. PDFs for $\cos \theta_{\text{sun}}$ are generated using the more pessimistic trigger efficiency curves measured by the laserball and TELLIE. Simulated events that have an in-time nhit that is predicted by the nhit-monitor to be 100% efficient are de-weighted to match the laserball efficiency measurement. The histogram that results from the de-weighted events are used as the systematically adjusted histogram to account propagate the trigger efficiency uncertainty to the flux result. The weight used for each value of $n_{100}$ is given in Tbl. 8.7.
Chapter 9

Results

9.1 Fit

The analysis outlined in the previous chapter was performed in two steps. First the analysis was done with a single energy bin from 5 to 15, this provided an idea of the signal and background rate, but was not used in the final flux result. The principal purpose of the single bin analysis was to constrain the value of $\delta_\theta$, the angular resolution systematic. A fit was done to this distribution in three dimensions, signal normalization, background normalization and $\delta_\theta$. In this fit $\delta_\theta$ was constrained by the separate $^{16}$N results, a bifurcated Gaussian was used to model the likelihood of the $^{16}$N results. The fit determined the systematic uncertainty on $\delta_\theta$ was determined to be $+0.07, -0.12$ a small improvement compared to the values from the $^{16}$N analysis, $+0.08, -0.013$.

The analysis was then performed again in six energy bins, five equal width bins from 5 to 10 MeV and a single bin from 10 to 15 MeV. In each energy bin a two-dimensional fit to the distribution of events in $\cos \theta_{\text{sun}}$ was done to determine the signal and background normalization. Then the likelihood space from each fit was combined, marginalizing over each energy bin’s background rate, to find the most likely solar neutrino flux normalization. The best fit solar rates in each energy bin also serve to provide a measurement of the spectral
Figure 9.1: Distribution of event direction with respect to solar direction. The systematic error bar includes angular correlated and uncorrelated errors.
shape of the solar neutrino flux. The spectral fit was also done as a three-dimensional fit in each bin, allowing \( \delta \theta \) to vary. This provided negligible difference in the final result and was significantly more computationally expensive.

For each of these steps the fitting routine was a simple grid scan over the parameter space. The solar and background normalizations were both scanned in 2000 steps from 0 to an value equal to 1.5 times the number of events in the energy bin. For fits involving \( \delta \theta \) 500 steps were used scanning values across double the \( ^{16}\text{N} \) constraint. The uncertainty introduced by the sampling step size was, in all cases, negligible compared to the systematic and statistical uncertainty.

Figure 9.1 shows the distribution of events in \( \cos \theta_{\text{sun}} \) for events over the entire energy range of 5 to 15 MeV and the fit to that distribution. The value \( \delta \theta \) for the best fit line is fixed to zero, but the systematic uncertainties are shown in the blue band. The fit gives a solar event rate of 1.30 \( \pm \) 0.18 events/kt-day and background rate of 10.23 \( \pm \) 0.38 events/kt-day.

From the energy bin-by-bin fit yielded the best fit solar event rate as a function of energy shown in Fig 9.2 and an overall flux

\[
\Phi_{ES} = 2.53^{+0.31}_{-0.28} \text{(stat.)}^{+0.13}_{-0.10} \text{(syst.)} \times 10^6 \text{cm}^{-2}\text{s}^{-1}.
\]

This value assumes the neutrino flux consists purely of electron flavor neutrinos. The result agrees with the elastic scattering flux published by Super-K, \( \Phi_{ES} = (2.345 \pm 0.039) \times 10^6 \text{cm}^{-2}\text{s}^{-1} \) \cite{58}, combining statistical and systematic errors.

Including the effects of solar neutrino oscillations, using the neutrino mixing parameters given in Ref. \cite{42} and the solar production and electron density distributions given in Ref. \cite{52} gave a best fit solar flux of

\[
\Phi_{^{8}\text{B}} = 5.95^{+0.75}_{-0.71} \text{(stat.)}^{+0.28}_{-0.30} \text{(syst.)} \times 10^6 \text{cm}^{-2}\text{s}^{-1}.
\]

This result is consistent with the \(^{8}\text{B} \) flux as measured by the SNO experiment, \( \Phi_{^{8}\text{B}} = (5.25 \pm 0.20) \times 10^6 \text{cm}^{-2}\text{s}^{-1} \) \cite{43}, combining statistical and systematic uncertainties. Figure 9.2 shows the best fit solar neutrino \(^{8}\text{B} \) event rate in each energy bin along with the
Figure 9.2: (Top) The extracted solar neutrino elastic scattering event rate as a function of reconstructed electron kinetic energy $T_e$. (Bottom) The same, as a fraction of the expected rate. The red and blue lines show the MC simulation predicted spectrum normalized to the best fit flux and the SNO flux measurement [45], respectively. The uncertainty on the SNO result includes reported uncertainty combined with mixing parameter uncertainties. The black points are the results of the fits to the $\cos \theta_{\text{sun}}$ distribution in each energy bin, with error bars indicating the combined statistical and systematic uncertainty, including energy-correlated uncertainty. A horizontal dash is placed on each error bar indicating the statistics only uncertainty; for all points the statistical error is dominant and the systematic error bar is not visible above the dash.
9.1 Fit

<table>
<thead>
<tr>
<th>Systematic</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy Scale</td>
<td>3.9%</td>
</tr>
<tr>
<td>Fiducial Volume</td>
<td>2.8%</td>
</tr>
<tr>
<td>Angular Resolution</td>
<td>1.7%</td>
</tr>
<tr>
<td>Mixing Parameters</td>
<td>1.4%</td>
</tr>
<tr>
<td>Energy Resolution</td>
<td>0.4%</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>5.0%</strong></td>
</tr>
</tbody>
</table>

Table 9.1: Effect of each systematic uncertainty on the extracted solar neutrino flux. Systematic uncertainties with negligible effects are not shown. For asymmetric uncertainties, the larger is shown.

predicted energy spectrum scaled to the best fit flux, and scaled to the flux measured by SNO. Each statistical error bar on the measured rate is affected by both the solar neutrino and background rates in that energy bin. Table 9.1 details how each systematic uncertainty affects this result.

The upper five energy bins, 6.0–15.0 MeV, were an extremely low background region for this analysis. There was very little background contamination from cosmogenically produced isotopes due primarily to depth of the detector. The comparatively high rate of backgrounds in the 5.0–6.0 MeV bin comes primarily from decays of radioactive isotopes, such as radon, within the detector. Figure 9.3 shows the distribution in $\cos\theta_{\text{sun}}$ of events at energies above 6 MeV, illustrating the low background rate. In that energy region the best fit background rate was $0.25^{+0.09}_{-0.07}$ events/kt-day, much lower than the measured solar rate in that energy range: $1.03^{+0.13}_{-0.12}$ events/kt-day. For the region above 6 MeV, this is the lowest background elastic scattering measurement of solar neutrinos in a water Cherenkov detector. Figure 9.4 shows the distribution of events in $\cos\theta_{\text{sun}}$ with best fit MC prediction in each energy bin of the analysis.
Figure 9.3: Distribution of event directions with respect to solar direction for events with energy in 6.0–15.0 MeV.
Figure 9.4: Distribution of events in $\cos \theta_{\text{sun}}$ in all six energy bins with best fit signal+background distribution.
9.2 Outlook For Water and Scintillator Phase

The primary result of this analysis is a measurement of the $^8$B flux with the SNO+ detector. But, by using the $^8$B flux measured by the SNO experiment, the data for this analysis can be used to measure the solar survival probability instead of the flux. This analysis is statistically limited, so a measurement of the survival probability done this way would have uncertainties much larger than the similar measurements done by the Super Kamiokande collaboration. But, the low backgrounds observed at higher energies could allow for a good measurement of the day-night effect once more data are acquired. A measurement of the day-night effect could allow for relatively competitive constraints on the standard neutrino mixing parameters.

Since the completion of this measurement SNO+ has acquired approximately double the livetime, with lower backgrounds. The decreased background rates may allow for an expanded fiducial volume at higher energies, which would increase the available statistics for this measurement or a possible measurement of the day-night effect. A likelihood fit in both $\cos \theta_{\text{sun}}$ and radius might allow for an expanded fiducial volume as well.

SNO+ has begun its partial fill period, in which the AV water is replaced with liquid scintillator. Once complete SNO+ will be able to take data for low-energy solar neutrino analysis. The expected increase in radio-purity and energy resolution will allow for solar neutrinos to be identified above background at much lower energies than was possible in the water phase. Figure 9.5 shows the expected solar neutrino and background spectrum at low energies for SNO+, scintillator purity related backgrounds are assumed to be at Borexino levels.

One of SNO+'s advantages for making solar neutrino measurements is the very large overburden, which provides a low rate of cosmogenic muons passing through the detector, which results in a low rate of cosmogenic isotope production within the detector. One of the most significant cosmogenic isotopes for low energy solar is the $^{11}$C, which has a half-life of 20 minutes and produces a $\beta^+$ with an endpoint energy approximately 1 MeV. The long
Figure 9.5: Expected interaction rates for solar neutrinos and backgrounds in SNO+’s scintillator phase
9.2 Outlook For Water and Scintillator Phase

half-life means the $\beta^+$ cannot easily be associated with the muon that produced it, and the energy spectrum for the decay is at the end point for the pep and CNO solar neutrino fluxes. So a low rate of cosmogenic muons will mean the $^{11}$C background does not preclude a significant measurement of the pep and CNO neutrino flux. However, the low rate of CNO neutrinos compared to pep neutrinos and other backgrounds will still make a strong measurement of the CNO flux very difficult. With 6 months of scintillator data it’s expected that SNO+ will be able to measure the pep flux to 13% uncertainty, almost a factor of two improvement on existing measurements. With the same data a measurement of $^7$Be flux to 5% and the $^8$B flux to 10% is expected.
Part II

Vacuum-Enhanced Mixing
Chapter 10

Model Description

Considered here is a novel phenomenological theory that modifies the solar neutrino survival probability. Whereas many theories of modified neutrino mixing alter the way neutrinos interact with matter, similar to standard matter enhanced-mixing, here we consider a modification to neutrino mixing in areas of very low matter density. Motivations for a potential with strength inversely related to local matter density is considered in Sec. 10.1. The effect a potential of this form is might have on neutrino mixing is parameterized as follows

\[ H = U M^2 U^\dagger + A_{\text{vac}}, \]  

(10.1)

where \( A_{\text{vac}} \) represents a potential only significant in regions of near-zero matter density. \( A_{\text{vac}} \) is given by

\[ A_{\text{vac}} = \begin{pmatrix} A_e & 0 & 0 \\ 0 & A_\mu & 0 \\ 0 & 0 & A_\tau \end{pmatrix}. \]  

(10.2)

Each diagonal entry represents a potential felt in the vacuum by the neutrino of the given flavor state. This parameterization can be simplified somewhat by subtracting off an identity like term.

\[ A'_{\text{vac}} = \begin{pmatrix} A_e & 0 & 0 \\ 0 & A_\mu & 0 \\ 0 & 0 & A_\tau \end{pmatrix} - \begin{pmatrix} A_e & 0 & 0 \\ 0 & A_\mu & 0 \\ 0 & 0 & A_\tau \end{pmatrix} = \begin{pmatrix} A'_e & 0 & 0 \\ 0 & A'_\mu & 0 \\ 0 & 0 & 0 \end{pmatrix}. \]  

(10.3)
10.1 Motivation

This makes no difference because neutrino mixing is only affected by differences between the neutrino states, so an identity-like matrix can always be subtracted from the mixing Hamiltonian without affecting the result. Given this the matrix $A'$ and the parameters $A'_e$ and $A'_\mu$ will be used for all discussion after this and the prime will be removed.

In this model mixing is described by standard MSW-LMA mixing in areas of non-negligible matter density, and by the modified vacuum mixing potential in areas of near zero matter density. In areas where the matter effect on neutrinos is negligible but the local matter density is non-zero, the mixing is best described a standard vacuum mixing potential, \( H = U M^2 U^\dagger \). A model like this is desirable because it allows for the mixing of solar neutrinos, which travel for a significant distance in the vacuum of space between the Sun and Earth, to be modified, while not effecting neutrino mixing observed by terrestrial neutrino experiments.

For nearly all other detected neutrino sources, neutrinos are produced, propagate through, and are detected in areas of relatively high matter density. The exception to this is supernova neutrinos observed by from SN1987A (87, 88, 89), and neutrinos detected by the IceCube experiment from a blazar (90). Neither of these datasets are sufficient to make statements about neutrino mixing though. Ultra-high energy neutrinos from the GZK production process (91) would be a useful source for constraining this model, as GZK neutrinos not only travel through significant distances in vacuum, they’re also produced in vacuum. However, neutrinos from the GZK process have not yet been detected by any experiments. And so, solar neutrinos are the only currently available source of neutrinos that are sensitive to this model.

10.1 Motivation

There are broadly two motivating factors behind a vacuum-enhanced mixing model. The first is the tension in between solar neutrino measurements and the KamLAND reactor neutrino measurement of $\Delta m_{21}^2$. This tension is discussed in detail in Sec. 11. While the tension
10.1 Motivation

could be the result of statistical fluctuations or un-modelled systematic effects, the flatness of observed solar survival probability and unexpectedly large day-night asymmetry both independently prefer lower values for $\Delta m^2_{21}$. This motivates considerations of what physics might affect the solar survival probability and not the KamLAND result. Unexpected effects from the high density long baseline solar environment provide a neat answer. Similarly, unexpected effects from a near zero matter density environment provides an explanation for the observed tension between the KamLAND and solar results.

The second motivating factor is that there exists theories that suggest a potential in empty space across large distance scales may be the source of the observed expansion of the universe (92). This new potential is referred to as a “chameleon” force because it is screened by the local matter around it, and so it “blends in” in areas of relatively high matter density. But in empty space the potential is much stronger and thus able to provide a source of energy for the observed expansion of the universe. The chameleon theory for dark energy could also provide a neutrino flavor dependent potential that affects the oscillation of neutrinos in areas of very low matter density. The chameleon theory for dark energy here only provides a phenomenological basis for considering vacuum-enhanced neutrino mixing. The model proposed here is not necessarily dependent on the chameleon dark energy theory, or vice versa. This relationship is similar to how Mass Varying Neutrino (MaVaN) models for neutrino oscillation are motivated by acceleron theories for dark energy, but not specifically required (93).
Chapter 11

Measurements of $\Delta m_{21}^2$

The current global best fit value for $\Delta m_{21}^2$ and $\theta_{12}$ comes primarily from a combined fit to solar neutrino data from SNO and Super-K and Borexino, and reactor neutrino data from KamLAND. These fits have produced the values

$$\Delta m_{21}^2 = 7.39^{+0.21}_{-0.21} \times 10^5 \text{eV}^2$$ \hspace{1cm} (11.1)

$$\sin^2 \theta_{12} = 0.310^{+0.013}_{-0.012}. \hspace{1cm} (11.2)$$

However, KamLAND has much better sensitivity to $\Delta m_{21}^2$ than the combined SNO and Super-K result, so the best fit value for $\Delta m_{21}^2$ comes almost exclusively from the KamLAND reactor neutrino measurements. The KamLAND measurement is discussed briefly in Sec 1.6.4. The precision of their measurement comes from the fact that $\Delta m_{21}^2$ adjusts the “wavelength” in energy of oscillations in their observed reactor neutrino spectrum. Figure 11.1 shows KamLAND’s observed oscillation spectrum and best fit.

Figure 11.2 shows the solar survival probability, with labelled features. The transition region is where the survival probability changes from being very close to the vacuum survival probability at low energies, to being dominated by matter effects at the higher energies. The width of the transition region comes from the radial production distribution.
Figure 11.1: Reactor neutrino survival probability measured by the KamLAND experiment. Figure from [62].
Figure 11.2: Day (gold) and night (black) solar survival probability with approximate value for vacuum and matter dominated regions.
of solar neutrinos in the sun; Neutrinos at energies high enough to experience significant matter effects in the sun, may or may not be in an area of the Sun with sufficient electron density to provide a strong matter effect. The reason the solar neutrino experiments are relatively insensitive to $\Delta m_{21}^2$ is because the only effect it has on the solar neutrino survival probability is to shift the energy of the MSW-transition region, a smaller value for $\Delta m_{21}^2$ results a lower energy transition region. At energies in the middle of transition region, you’ll observe approximately as many neutrinos that never experience significant matter effects as neutrinos that did. For experiments, this means sensitivity to $\Delta m_{21}^2$ comes primarily from their ability to measure the survival probability in the transition region compared to the vacuum or matter dominated regions.

In principle this means for solar neutrino experiments to be sensitive to $\Delta m_{21}^2$, a good measurement of neutrino oscillation probability in the transition region is all that is required. However, the Sun does not provide an optimal source of neutrinos at energies in the transition region. The $^{8}\text{B}$ and $^{9}$hep reactions provide neutrinos at energies across the entirety of the transition region, but the flux of each of those sources is very small, especially at energies between $1 - 3 \text{ MeV}$; At energies below $15 \text{ MeV}$ the $^{8}\text{B}$ flux is dominant over the $^{9}$hep flux.

The challenge of detecting relatively low energy, low flux $^{8}\text{B}$ neutrinos is made more difficult by the fact that at those energies radioactive backgrounds are dominant in water Cherenkov detectors. Most notably backgrounds from $^{214}\text{Bi}$ and $^{208}\text{Tl}$ contamination which each provide a $\beta$ with a spectrum that respectively have a decay Q-value of $3.2 \text{ MeV}$ and $5.0 \text{ MeV}$. These background render $^{8}\text{B}$ neutrino measurements in the transition region nearly impossible for water Cherenkov detectors. For a liquid scintillator detector, target volume can be purified to a greater degree than a water detector, which mitigates background contamination somewhat. However, scintillator detectors cannot in general reconstruct the direction of the neutrino, and so solar neutrinos cannot be identified against backgrounds. And so even with higher radio-purity, at energies below approximately $3 \text{ MeV}$, the $^{8}\text{B}$ solar
neutrino flux is extremely difficult to measure due to the low flux and cross-section at those energies.

Beyond the difficulty of measuring solar neutrinos at energies in the transition region, is the fact that the most common way to measure solar neutrinos for water Cherenkov or scintillator detectors is through the electron elastic scattering interaction. And as discussed in Sec 1.2 the energy of the observable scattered electron is weakly correlated with the energy of the incoming neutrino. And so even if a measurement of $^8$B solar neutrino interaction rate at low energies was made, that interaction rate would include contributions from $^8$B neutrinos from the observed energy up to the end-point energy. Meaning any measurement performed in the transition region will contain many more interactions from neutrinos with energy above the transition region than within the transition region. This means any measurement will require a very high statistics dataset to make a good measurement of the solar neutrino survival probability in the transition region.

Despite these difficulties the $^8$B survival probability has been measured down to $3.5 \, MeV$, though uncertainties at the lower energies are significant. These measurements come from the Super-K elastic scattering measurements (58), combined with the three-phase SNO elastic scattering, charged-current and neutral-current measurements (45). The SNO measurement provides the strongest constraints on the full, all-flavor, $^8$B flux, and the Super-K measurement provides the strongest constraints on the flavor sensitive “elastic-scattering” flux. So far these experiments have not seen strong evidence for any upward transition in the survival probability. Figure 11.3 shows the SNO measurement and the combined SNO + Super-K measurement.

Because the “up-turn” in the survival probability has not yet been observed, solar neutrino measurements of $\Delta m_{21}^2$ prefer relatively low values that push the transition region to lower energies. Using only data from SNO and Super-K the best fit value for $\Delta m_{21}^2$ is

$$\Delta m_{21}^2 = (4.8^{+1.5}_{-0.8}) \times 10^{-5} \, eV^2 (58).$$

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Figure 11.3: Quadratic fit (a top) to solar survival probability and (b) linear fit to day-night asymmetry as measured by SNO. (b) (red) Measured ratio of oscillated solar neutrino spectrum to unoscillated and best fit. The green and blue lines are respectively the best fit survival probability for the SK only and the solar+KamLAND values of $\Delta m_{21}^2$. The orange and black lines are respectively the best fit quadratic and cubic functions. Figures from Ref (45) (a) and (58) (b).
In principle, the transition region could be probed as well by a precise measurement of the survival probability from pep or 7Be neutrinos. pep neutrinos are mono-energetic at $E_\nu = 1.4$ MeV, this means they should have a survival probability of 53% instead of the vacuum survival probability of about 56%. 7Be neutrinos are also mono-energetic at 0.861 MeV, providing them with an expected survival probability of 54%. A measured deviation of these survival probabilities from the expected vacuum mixing value would provide a measurement of $\Delta m_{21}^2$. But so far measuring these fluxes to that level of precision has not been possible. The current best measurement of these neutrino fluxes comes from the Borexino collaboration (61), their results are shown in Tbl. 11.1.

The day-night effect also provides sensitivity to $\Delta m_{21}^2$. The smaller $\Delta m_{21}^2$ is, the lower the energy threshold is for neutrinos to experience significant matter effects as they travel through the Earth. In practice the day-night effect has been difficult to measure due to the statistics required to measure a few percent change on the survival probability. And so the majority of sensitivity to $\Delta m_{21}^2$ for solar neutrino experiments comes from their ability to measure the transition region. The day-night asymmetry in the survival probability is often parameterized by the value $A_{ee}$, a value defined as,

$$A_{ee}(E_\nu) = \frac{P_{\text{ee, day}}(E_\nu) - P_{\text{ee, night}}(E_\nu)}{P_{\text{ee, day}}(E_\nu) + P_{\text{ee, night}}(E_\nu)}.$$  \hfill (11.3)

The SNO measurement of this asymmetry is shown in the bottom panel of 11.3a, their result is consistent with no asymmetry. The Super-K measured value for the asymmetry in

<table>
<thead>
<tr>
<th>Measured Survival Probability [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>$pp$</td>
</tr>
<tr>
<td>$56.6 \pm 9.2$</td>
</tr>
<tr>
<td>$7Be$</td>
</tr>
<tr>
<td>$53.2 \pm 5.4$</td>
</tr>
<tr>
<td>$pep$</td>
</tr>
<tr>
<td>$42.8 \pm 11.4$</td>
</tr>
</tbody>
</table>

Table 11.1: Measured survival probabilities for low energy solar neutrinos from the Borexino experiment (61).
the observed interaction rate (as opposed to the survival probability) is

\[ A_{SK} = (-3.3 \pm 1.0\,(\text{stat.}) \pm 0.5\,(\text{syst.}))\%. \]

Figure 11.4: Zenith angle dependence of elastic scatter interaction rate observed with Super-K. Figure from (58)

This value is inconsistent with the no asymmetry at the roughly two-sigma level. The predicted value for this asymmetry for the best fit KamLAND \( \Delta m_{21}^2 \) is \(-1.7\%\) the predicted value for the Super-K+SNO value of \( \Delta m_{21}^2 \) is \(-3.3\%\). So with relatively small significance, the solar measurement of \( \Delta m_{21}^2 \) from the day-night effect prefers a lower value than the KamLAND measurement. Figure 11.4 shows the Super-K asymmetry measurement in the observed interaction rate.

Figure 11.5 shows a summary of solar neutrino measurements with the solar survival probabilities as predicted by the KamLAND+solar and the solar only values for \( \Delta m_{21}^2 \).
Figure 11.5: Compilation of measurements to solar survival probability measurements overlaid onto the solar neutrino spectrum. The red band is a combined SNO+SuperK measurement, the data points come from Borexino [94], the CNO limit combines Borexino and radio-chemical measurements. The blue, teal and green lines are the MSW predicted survival probability for respectively the solar+Kamland, combined solar, and Super-K only mixing parameters. Figure from [58]. Figure from [58]
Figure 11.6: Likelihood contours for solar and KamLAND fit to mixing parameters. Contours are one-sigma, 90%, two-sigma, 99%, and three-sigma for two degrees of freedom. Figure from [95].
A global fit for solar neutrino parameters done by Esteban et. al. (95), shown in Fig. 11.6, the overall significance disagreement between the KamLAND+solar and the solar only measurements for $\Delta m^2_{21}$. They perform a fit to the solar data using solar model input from the GS98 solar abundances as well as the AGS abundances, and show that uncertainty introduced by the disagreement in those models is not sufficient to explain the consistently low value for $\Delta m^2_{21}$ observed by SNO and Super-K. They determine the disagreement between the solar and KamLAND values for $\Delta m^2_{21}$ to have a significance of $\Delta \chi^2 = 4.7$, or roughly a 3% chance of being the result of a statistical fluctuation.
Chapter 12

Simulation

Two different simulations of neutrino oscillations were developed to explore the effects of the vacuum-enhanced mixing potential. The first does a full simulation of the neutrino state as it propagates from the Sun, through the vacuum of space, to a detector at Earth. The second calculates the evolution of the mixing eigenstates for solar neutrinos, and does not simulate individual neutrinos. These simulations will respectively be referred to as the “neutrino” simulation and as the “mass-state” simulation. The goal of both simulations is to produce a solar neutrino survival probability for any given set of standard model mixing parameters and vacuum-enhanced mixing parameters. The reason for the two different methods is to allow for different trade-offs between simulation precision and computational speed.

12.0.1 Neutrino Simulation

For this method of simulation the neutrino state is propagated through the Sun with its mixing given by

$$i\frac{d}{dx} |\Psi\rangle = \frac{1}{2E} \left[ UM^2 U^\dagger + A_{CC} \right] |\Psi\rangle .$$

(12.1)
Where $|\Psi\rangle$ is 3-component vector, each value of which represents the fraction of the neutrino quantum state in the electron, muon, or tau flavor states,

$$|\Psi\rangle = \begin{bmatrix} \Psi_e \\ \Psi_\mu \\ \Psi_\tau \end{bmatrix}. \quad (12.2)$$

Since solar neutrinos are all produced as an electron neutrino the initial state is always

$$|\Psi(x = 0)\rangle = \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix}. \quad (12.3)$$

Given an energy and initial position within the Sun Eqn. (12.1) is evaluated numerically using the Runge-Kutta method of numerical integration. At each step of the integration the neutrinos position is advanced radially outward a small amount, and using the electron density at the new radius $A_{CC}$ is re-evaluated for the next step of integration. A linearly interpolated solar electron density curve from the BS05OP solar model is used. This process is repeated until the neutrino reaches a radius where the solar electron density is near zero and the mixing Hamiltonian is just standard vacuum oscillations. The reason the simulation stops there rather than going all the way to the edge of the sun is because once the matter effect has diminished to a negligible level, the state of the neutrino no longer evolves – the state oscillates around a central value, that value is static though. So, upon reaching the end of the simulation the final 5000 neutrino states of the integration are recorded to provide a full sampling of the possible neutrino states. Figure 12.1 shows the simulated survival probability of a single 10 MeV neutrino produced in the center of the sun.

Simulation was done for 200 neutrino energies spaced logarithmically from 0.1 MeV to 20 MeV and for each energy neutrinos at 8192 different starting radii spaced linearly from 0 solar radii to 0.6 solar radii were simulated. The result of the simulation is approximately 10 billion neutrino states to provide a representation of all possible neutrino states the
Figure 12.1: (a) The simulated survival probability for a 10 MeV solar neutrino produced in the center of the sun. The bottom panel (b) is the same as the top (a), but zoomed in around an arbitrary distance. X-axis units are meters.
sun can produce. This “library” of possible solar neutrino states is used as input to the vacuum-enhanced potential simulation.

The simulation of neutrinos this way is computationally expensive. A few methods were explored for ensuring this simulation could be performed in a reasonable amount of time. The method that was used for nearly all the results here was to perform the Runge-Kutta integration on a GPU, which each thread corresponding to a single sample in energy and production radius. Still, the production of all the simulated solar states requires approximately 2 GPU-months of processing. By construction though, the solar simulation is not affected by modified vacuum potential; the main inputs to the solar simulation are the standard model mixing parameters and the solar density profile. So, standard model mixing parameters taken from KamLAND and other non-solar neutrino experiments can be used for the solar simulation.

The modified vacuum portion of the simulation is done by first Monte Carlo sampling neutrino energies and production radii, and looking up the simulated neutrino state that corresponds to the sampled energy and production radius. The radial production PDFs for each neutrino type from the BS050P standard solar model are used for sampling production radii. The neutrino energy PDFs are also sampled however, only half of all sample energies are drawn from the relevant production spectra, the other half are drawn from a uniform distribution across the relevant energy region. The motivation for this sampling method is discussed further at the end of this section.

Since MC sampling produces energy and production radius values that fall between bins used when producing the solar state library, the states from the closest available radius bin is used. For energy a random choice is made between the energy bins immediately above and below the desired energy is performed, the probability of choosing either bin is dependent on how close the desired energy is to that bin. This process is repeated for 1000000 simulated neutrinos for each solar neutrino type, e.g. \(^{8}\)B, pep, etc.
Once Monte Carlo samples of solar neutrino states are calculated, the states are used as inputs to a simulation of the vacuum-enhanced potential. This simulation is in principle the same as the solar simulation, it simply involve evaluating Eqn. (12.1), where $A_{CC}$ is replaced by $A_{vac}$. Unlike the solar simulation where the electron density changes, the value for $A_{vac}$ is modeled as a constant as the neutrino propagates between the Sun and Earth. The constant value for the Hamiltonian means Eqn. (12.1) can be evaluated using a combination of analytic and numerical techniques instead of numerical integration.

The vacuum state evolution done by first evaluating the mixing Hamiltonian and diagonalizing the resulting matrix. Diagonalization is done using the GNU Scientific Library (96) linear algebra routines. The resulting eigenvectors are used to transform to a basis where the Hamiltonian is diagonal and the state can be evolved by evaluating

$$|\Psi_k(x)\rangle = e^{-iE_kx} |\Psi_k(t = 0)\rangle.$$ (12.4)

The energy $E_k$ is the $k^{th}$ energy eigenvalue determined by the diagonalization routine, $|\Psi_k(t = 0)\rangle$ is neutrino state, in the eigen-basis, at the time that it enters the modified vacuum potential; the states from the solar simulation are transformed to the eigen-basis and used for $|\Psi_k(t = 0)\rangle$. Equation (12.4) is evaluated for $x = 1\text{AU}$ and the new state is transformed back into the flavor basis. Performing this series of steps for each Monte Carlo sampled neutrino state provides the expected distribution of states as they arrive at Earth having travelled through a modified vacuum mixing potential.

The final step of the calculation is to evolve the sampled neutrino states through the Earth, to the detector. This is done similarly to the simulation of neutrino propagation through the Sun. The calculation for this is done for only a “day” path through the earth and a “night” path. The “day” path simulates the neutrino only travelling through the crust of the Earth. The “night” path simulates the neutrino travelling through the Earth, including the high density core. The one-dimensional earth density profile is from Ref. (97) is used.
The result of this chain of simulation steps is Monte Carlo samples of neutrino flavor states. The survival probability is calculated as,

\[ P_{ee}(|\Psi\rangle) = |\langle \nu_e | \Psi_{\nu} \rangle |^2 \]  

(12.5)

for each neutrino state. Performing an average of survival probabilities binned in neutrino energy gives the survival probability as a function of energy \( P_{ee}(E_\nu) \).

Since neutrino states are Monte Carlo sampled to calculate \( P_{ee}(E_\nu) \), each value has statistical uncertainty from the number of samples used. This problem was somewhat exacerbated by the distributions of some solar neutrino energy PDFs having small values in areas that are important for comparing to solar neutrino data. For example the low energy portion of the \(^8\)B solar neutrino flux is very important for solar neutrino experiments, but makes up a relatively small portion of the full \(^8\)B neutrino flux. To mitigate the problem of large sampling uncertainty for important regions in solar neutrino energy, energies were sampled according to a flat distribution and according to the PDFs for each solar neutrino flux. These two methods of sampling were performed in equal proportions for each flux type. Figure 12.2 shows an example survival probability from the vacuum-enhanced neutrino state simulation.

This method for calculating a vacuum-enhanced survival probability has the distinct drawback of being computationally expensive, to the point where it cannot be used in a fit to data. It does, however, have the benefit of simulating effects from a non-adiabatic transition from between different potentials that effect neutrino mixing. Non-adiabatic effects are important because the neutrino is neither produced nor detected in a vacuum, so for modified vacuum potential to have an effect that can be detected with a terrestrial detector there must be some non-adiabatic transition.
Figure 12.2: Survival probability with vacuum-enhanced mixing parameters $A_e = 6e - 13eV$, $A_\mu = A_\tau = 0$ from non-adiabatic simulation. Also shown are the survival probabilities for the best fit solar only and KamLAND only values for $\Delta m_{21}^2$. 
12.1 Mass State Simulation

For the mass state simulation instead of simulating individual neutrino states, only the eigen-states are simulated. For each production radius and production energy within the sun the mixing Hamiltonian is diagonalized,

$$H = U M U^\dagger + A_{CC} = P D P^\dagger.$$  \hfill (12.6)

Where $D$ is a diagonal matrix that gives the effective mass-squared difference between the mass-states. $P$ gives the flavor composition of the effective mass states, $|\nu^m_1\rangle$, $|\nu^m_2\rangle$, $|\nu^m_3\rangle$.

A neutrino produced in an electron flavor state, is given by

$$|\nu\rangle = \sum_{k=1}^{3} \langle \nu^m_k | \nu_e \rangle |\nu^m_k\rangle.$$  \hfill (12.7)

The neutrino state is then evolved adiabatically into a modified vacuum potential given by

$$H = U M U^\dagger + A_{\text{vac}}.$$  \hfill (12.8)

The eigenstates from the modified vacuum Hamiltonian gives the neutrino mass states $|\nu^{\text{vac}}_1\rangle$, $|\nu^{\text{vac}}_2\rangle$, $|\nu^{\text{vac}}_3\rangle$. So the neutrino state is now given by

$$|\nu\rangle = \sum_{k=1}^{3} \langle \nu^m_k | \nu_e \rangle |\nu^{\text{vac}}_k\rangle.$$  \hfill (12.9)

In general this equation could be used to evaluate the survival probability, as is shown in Section 1.3. However, equation 1.23 shows that this process produces terms that oscillate as the neutrino state evolves as well as constant terms. Solar neutrino experiments are not sensitive to oscillations in the survival probability because the production of neutrinos is distributed throughout the core of the Sun, which is many neutrino oscillation lengths across.
12.1 Mass State Simulation

Because solar neutrino experiments detect neutrinos incoherently, averaging over differences in oscillation lengths for neutrinos produced at different radii in the Sun, the survival probability can be expressed as,

$$P_{ee} = \sum_{k=1}^{3} \left| \langle \nu_{e} \mid \nu_{m}^{k} \rangle \right|^{2} \left| \langle \nu_{e} \mid \nu_{k}^{\text{vac}} \rangle \right|^{2}.$$  \hspace{1cm} (12.10)

Where terms that include oscillations from $|\nu_{m}^{k}\rangle$ to $|\nu_{j}^{\text{vac}}\rangle$ ($i \neq j$) average to zero.

This method for calculating the survival probability relies on the neutrino entering the modified vacuum potential adiabatically. Since the mechanism by which the neutrino is sensitive to the local matter density is not considered here, it’s plausible that the modified vacuum potential “turns-on” slowly enough that there’s no non-adiabatic transition. However, as mentioned earlier, since the solar neutrinos are neither detected nor created in a vacuum, a non-adiabatic transition is required for terrestrial neutrino detectors to be sensitive to a modified vacuum neutrino potential. For this model that non-adiabatic transition is assumed to be that the neutrino does not fully transition from being best described by a modified vacuum state by the time it’s detected. This is plausible because the oscillation length for higher energy neutrinos is approximately 200 km, the earth’s atmosphere is approximately the same size. So it could be the case that solar neutrinos, especially at higher energies, would not oscillate quickly enough to have an average survival probability best described by the standard oscillation probability. For any set of mixing parameters this assumption can be checked using the more computationally expensive neutrino state evolution simulation.

The benefit of this method for calculating modified vacuum solar survival probabilities is that it’s computationally much faster than the full neutrino state evolution; A much wider search of the parameter space for solar neutrino mixing to be explored.

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Chapter 13

Data Sets

To determine solar neutrino and neutrino mixing parameters that is most consistent with experimental observations, a fit is done to published experimental results. Solar neutrino results from SNO [45], Super Kamiokande [35, 58, 98, 99], Borexino [61, 94], GNO [26, 27], SAGE [25], and Homestake [16] are used. Solar neutrino fluxes are constrained by the GS98 [53] solar abundance calculations. Reactor neutrino results from Daya Bay [100], KamLAND [62, 101], RENO [102] are also used to constrain neutrino mixing parameters. The KamLAND experiment has also published solar neutrino measurements [2, 3], only their reactor neutrino results are used however.

The experimental results for each experiment are modeled using methods developed by Richard Bonventre, Anthony LaTorre and Olivia Wasalski [74]. For all solar experiments, except for SNO, the expected event rate in each energy bin over the experiments sensitive energy region is calculated; for Super-K and Borexino the event rates are calculated in each energy bin. Event rates are calculated by performing a convolution of the neutrino flux spectrum with the interaction cross-section relevant for each experiment and with the hypothesized neutrino survival probability spectrum producing a spectrum of observable elastic-scatter electron energies, or an event rate for nuclear interactions. For Super-K and
Borexino the electron energy spectrum is convolved with a detector response function to produce expected event rates. For the Homestake experiment interaction cross-sections from Bahcall (103) are used. For the gallium experiments cross-sections for $^{71}\text{Ge}$ is given by the SAGE collaboration in Ref. (104) and Bahcall (105).

Each experiment provides its own measurement of detector related systematic uncertainties, these uncertainties are all treated as independent, except for the four Super-K datasets. Each Super-K dataset has its own associated systematic uncertainty, but each uncertainty is treated as fully correlated across datasets. Systematic uncertainties are treated as Gaussian, or a bifurcated Gaussian in the case of asymmetry uncertainties. Systematics are marginalized over when evaluating the likelihood of any hypothesized event rate.

For comparison to SNO data, the likelihood of a survival probability can be evaluated directly without calculating expected event rates. SNO reports their measurement of the $^8\text{B}$ survival probability as a quadratic parameterization,

$$P_{ee}(E_\nu) = c_0 + c_1(E_\nu - E_0) + c_2(E_\nu - E_0)^2$$

(13.1)

where $E_0 = 10 \text{ MeV}$, and a linear day-night asymmetry

$$A_{ee}(E_\nu) = a_0 + a_1(E_\nu - E_0).$$

(13.2)

A procedure recommended in Refs (106) and (45) is followed. A fit to of Eqn. (13.1) and (13.2) is done to the hypothesized the survival probability, exacting values for $c_0, c_1, c_2, a_0$ and $a_1$. The best fit values are compared to the best fit values reported by SNO to determine a likelihood.

For evaluating the likelihood of the KamLAND result for a set of mixing parameters and vacuum-enhanced parameters a simple lookup is done to their published $\Delta \chi^2$ map (101). A similar approach is used to evaluate the likelihood the Daya Bay and RENO reactor neutrino results. The hypothesized value for $\theta_{13}$ is compared to each experiment’s measured value with uncertainty to extract a likelihood.
Chapter 14

Fit And Results

Taken together the solar neutrino experimental results can be used to constrain to the vacuum-enhanced solar survival probability allowing up to 11 free parameters: $\Delta m_{21}^2$, $\theta_{12}$, $\theta_{13}$, each of the 5 pp-chain solar neutrino fluxes, the combined flux of the CNO neutrinos, and the two modified vacuum mixing potentials $A_e$ and $A_\mu$. A gradient descent fit was done for these parameters to evaluate the likelihood of the vacuum enhanced model, and to what degree the tension between solar measurements and KamLAND measurements for $\Delta m_{21}^2$ can be reduced. The fit allowed all values to vary except $\theta_{13}$, which does not strongly affect the solar survival probability; $\theta_{13}$ was fixed to 8.5$^{\text{deg}}$, a value preferred by other global fits to neutrino data (95).

The fit was done in four different manner, first using a solar only dataset and allowing the vacuum-enhanced mixing parameters to vary, then using solar only data and fixing vacuum-enhanced mixing parameters to zero. Then the fits are redone using the KamLAND constraints, allow the vacuum-enhanced mixing parameters to vary and fixing them to zero. Fitting is done using the ROOT Minuit algorithm (107).

It was noted that depending on the starting parameters the gradient descent would often converge at significantly different values. This suggests that gradient descent cannot
provide an accurate picture of the likelihood space. A full scan of the likelihood space is necessary to get a complete picture of the most likely values for each parameter. However, a thorough scan is too computation expensive for this result and so it is discussed more in the following section.

The results of each fit is summarized in Tbl. 14.1 and 14.2, not shown are the best fit neutrino flux values. The reported error on at least some fit results is unbelievably small. I believe this is a result of the non-Gaussian nature of the likelihood space causing the Minuit algorithm to poorly estimate errors and not an accurate estimate of the true errors. A more thorough fit that does not rely so heavily on assumptions about the likelihood space will be

$\begin{align*}
\Delta \chi^2 & \\
\Delta m^2_{21}[10^{-5}\text{eV}^2] & \\
\sin^2 \theta_{12} & \\
A_e[\text{eV}^2] & \\
A_\mu[\text{eV}^2] & 
\end{align*}$

$\begin{align*}
\Delta \chi^2 & \\
\Delta m^2_{21}[10^{-5}\text{eV}^2] & \\
\theta_{12} & \\
A_e[\text{eV}^2] & \\
A_\mu[\text{eV}^2] & 
\end{align*}$

Table 14.1: Fit results from an 11-parameter gradient descent fit to solar neutrino experimental measurements. Errors provided by the Minuit fitting algorithm and commented on in text.

<table>
<thead>
<tr>
<th>$\Delta \chi^2$</th>
<th>$\Delta m^2_{21}[10^{-5}\text{eV}^2]$</th>
<th>$\sin^2 \theta_{12}$</th>
<th>$A_e[\text{eV}^2]$</th>
<th>$A_\mu[\text{eV}^2]$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solar Only, Vacuum-Enhanced</td>
<td>$0$</td>
<td>$6.49 \pm 2.2$</td>
<td>$0.30 \pm 0.17$</td>
<td>$(-1.19 \pm 0.77) \times 10^{-11}$</td>
</tr>
<tr>
<td>Solar Only, MSW-LMA</td>
<td>$0.35$</td>
<td>$4.30 \pm 0.13$</td>
<td>$0.31 \pm 0.13$</td>
<td>$(-2.06 \pm 1.18) \times 10^{-11}$</td>
</tr>
</tbody>
</table>

Table 14.2: Fit results from an 11-parameter gradient descent fit to solar neutrino experimental measurements and the KamLAND reactor neutrino measurement. Errors provided by the Minuit fitting algorithm and commented on in text.
done in the future to estimate the errors. I believe the errors reported from the fit should be taken as a likely lower bound on the true 1-sigma error. However, these errors are not necessary for evaluating the consistency of the model with data.

These results show that the vacuum-enhanced mixing model can significantly reduce the tension between the KamLAND and solar measurements values for $\Delta m^2_{21}$. For the solar only fit a non-zero value for the vacuum-enhanced mixing parameters is preferred at small significance. For the solar+KamLAND fit the $\Delta \chi^2$ between the best fit vacuum-enhanced model and the standard MSW-LMA model is 3.75. The solar+KamLAND best fit has a $\Delta \chi^2 = 0.85$ to the solar only best fit value, evaluated using only solar neutrino measurements. This can be compared to the likelihood of the solar+KamLAND best fit parameters with only standard MSW-LMA mixing compared to the solar only best fit, this gives $\Delta \chi^2 = 4.18$. So the tension between the solar and KamLAND measurements is reduced from just over two-sigma to 0.92-sigma by allowing for vacuum enhanced mixing.

Figure 14.1 shows a comparison between the best fit survival probability from the solar only fit for the vacuum-enhanced mixing model and the standard MSW-LMA model. Similarly, Fig. 14.2 and Fig. 14.3 respectively show the comparison of the same survival probabilities to SNO and Super-K data. Using instead the best fit values from the solar and KamLAND constraints gives the similar comparisons shown in Figures 14.4, 14.5, and 14.6.

14.1 Conclusions and Future Work

The vacuum-enhanced mixing model is able to improve the fit to data when considering the KamLAND and solar neutrino measurements of $\Delta m^2_{21}$; the tension is less than one-sigma when allowing for vacuum-enhanced mixing compared to approximately two-sigma for standard mixing. This reduction is relatively weak considering the model adds two new degrees of freedom into the mixing calculations. I therefore do not consider this to be strong evidence in favor of the vacuum-enhanced mixing model that I developed here.
14.1 Conclusions and Future Work

**Figure 14.1:** Best fit survival probability for the vacuum-enhanced model and the standard MSW-LMA model using only solar neutrino data.

**Figure 14.2:** Comparison of SNO parabolic fit to the solar only best fit vacuum-enhanced survival probability and the solar only best fit MSW-LMA survival probability.
Figure 14.3: Comparison of Super-K data to the predicted elastic scattering rate from the solar only best fit vacuum-enhanced survival probability and the solar only best fit MSW-LMA survival probability. Top-left is the Super-K-I dataset, top-right is Super-K-II, bottom-left is Super-K-III, bottom-right is Super-K-IV.
14.1 Conclusions and Future Work

**Figure 14.4:** Best fit survival probability for the vacuum-enhanced model and the standard MSW-LMA model using solar and KamLAND neutrino data.

**Figure 14.5:** Comparison of SNO parabolic fit to the solar + KamLAND best fit vacuum-enhanced survival probability and the solar + KamLAND best fit MSW-LMA survival probability.
Figure 14.6: Comparison of Super-K data to the predicted elastic scattering rate from the solar + KamLAND best fit vacuum-enhanced survival probability and the solar + KamLAND best fit MSW-LMA survival probability. Top-left is the Super-K-I dataset, top-right is Super-K-II, bottom-left is Super-K-III, bottom-right is Super-K-IV.
14.1 Conclusions and Future Work

As mentioned before the fitting routine used here was observed to produce different results depending on the starting point for the gradient descent. In the future a more thorough search of the parameter space will be done, allowing for a full profile of the likelihood for different values of $\Delta m^2_{21}$ and a more exhaustive search of the best fit vacuum-enhanced mixing parameters and better error estimation. This sort of scan for the likelihood space was done, fixing all parameters except $\Delta m^2_{21}$, $\theta_{12}$ and the vacuum-enhanced mixing parameters. The results did not prove significantly useful for evaluating the vacuum-enhanced mixing model though, but are described in appendix B. There are a number of ways the simulation of vacuum-enhanced survival probabilities and the evaluation of experimental likelihood can be made more computationally tractable, including by offloading more of the computation to a GPU. For future works this will allow the likelihood space to be searched more thoroughly.

For the solar only best fit survival probability an up-turn exists at higher energies. The up-turn would be a relatively unique observable signature for this model as opposed to standard MSW-LMA mixing which does not allow for that sort of behavior. So future experiments that can measure the survival probability with extreme sensitivity at higher energies could allow for this model to be separated unambiguously from the standard mixing hypothesis. However, for the best fit survival probability when considering the KamLAND constraints on $\Delta m^2_{21}$ this high energy up-turn nearly disappears, or is much more subtle. So even with good sensitivity at higher energies this model may not be able to be uniquely identified using standard solar neutrino measurements.

The results found here were produced entirely using adiabatic neutrino mass state simulation, this choice was necessary to provide results in a reasonable amount of time. For future works it may be possible to use the non-adiabatic neutrino simulation to verify the results of the adiabatic simulation. This will allow for the portion of time the neutrino spends in Earth’s atmosphere to be simulated more accurately. Also, not considered here is any sort of annual variation that might be present in a vacuum-enhanced mixing scenario.
As the Earth moves closer or further from the Sun throughout the year neutrinos will move through more or less vacuum. If there exists significant non-adiabatic effects, it might be possible to observe some annual modulation to the neutrino signal.

Finally, the parameterization used here only considered a potential that couples to neutrino flavor, this was done as an arbitrary choice to limit the parameter space. For future work this choice should be reconsidered, allowing for an arbitrary coupling potential to see what provides the best fit to data, and what sort of experimental signature a model like that might have. The motivation for this would be that there’s no reason to expect a new vacuum potential to couple to Weak flavor states, and so you might expect them to couple to some other, arbitrarily mixed, combination of mass or flavor states. Adopting a model like this would require a significant amount of computation power to fully evaluate. So it’s hoped that a simulation of vacuum-enhanced mixing that relies more heavily on analytic calculation can be developed.
Appendices
Appendix A

Data Cleaning

Each data cleaning cut is associated with a bit in a 64-bit binary value called the data cleaning word or data cleaning mask. The cuts that each event passes or fails is tracked by its data cleaning mask. For the solar analyses all events are required to pass all cuts given by the data cleaning mask 0xFB0000017FFE. This corresponds to the following data cleaning cuts: Zero Zero Cut, Crate Isotropy Cut, FTS Cut, Flahser Geo Cut, ITC Time Spread Cut, Junk Cut, Muon Tag, Neck Cut, Owl Cut, QCluster Cut, QvNhit Cut, QvT Cut, Ring Of Fire Cut, Two-Pass Muon Follower, Short, Polling Cut, Retrigger Cut, Two-Pass Burst Cut, Missed Muon Follower Cut, Missing CAEN Data Cut, Ped Cut, Atmospheric Cut. Of those cuts I’ll detail here the motivation behind and evaluation of the cuts that I developed, the cuts that I did not directly develop are described in (83).

A.1 Ped Cut

During normal detector operations there are a few trigger calibration tests that are periodically ran. These tests use the PEDESTAL signal to inject a certain amount of fake hits into the detector, and events with those hits are inspected to evaluate the efficiency and
quality of the trigger response. It’s very important that these events are clearly identified and removed from the dataset so that the fake PEDESTAL hits are not confused for a real signal. Additionally, the trigger calibration processes usually include changing settings related to the PEDESTAL signal on the FEC, there’s reason to believe these sorts of changes can introduce noise to the front-end. So an aggressive approach of cutting all events that are within one second of a pedestal event is used. This not only cuts events but introduces a dead time into the dataset, this deadtime is subtracted from the overall livetime.

A.2 Missed Muon Follower Cut

The missed muon follower cut was a data cleaning cut used in SNO, but I adapted it for SNO+. In SNO it was very important to identify and cut neutrons that follow after a cosmic muon event, those neutrons could fake a neutral current solar neutrino event. In SNO+ this is not as much a problem because neutron captures in water will primarily produce a 2.2 MeV gamma, which is below the analysis threshold for solar neutrinos. However, there does exist events in the dataset which are observed to follow after high-nhit, events. The origin of these events is not well understood, they could likely be instrumental, or from spallation products within the detector. Since solar neutrino events are not expected to have any time correlations with other events in the detector, a cut can be placed on the time between events with relatively little sacrifice.

The cut considers three values, an initial and a follower event nhit, $N_1$ and $N_2$, and a time difference between the two events $\Delta t$; a threshold is placed on each of these three values, and any pairs of events that cross each threshold are cut and so is all other events that fall within the threshold time window. To evaluate the cut optimal cut criteria for $N_1$, $N_2$ and $\Delta t$ two different values are considered. The first is the probability that an event of nhit greater than $N_2$ will occur within a time window $\Delta t$ after an event of nhit greater than $N_1$. This value is called the follower probability, $P_F(N_1, N_2, \Delta t)$. The second values is
the probability that an event of nhit $N$ will occur within a time window $\Delta t$ that is chosen randomly, this value is called accidental probability or random coincidence probability, $P_R(N, \Delta t)$. Under the assumption that there is no time correlation with non-background events the probability of this cut sacrificing a signal event is

$$\text{Sacrifice} = P_R(N_1, \Delta t)P_S(N_2) + P_R(N_2, \Delta t)P_S(N_1),$$  \hspace{1cm} (A.1)$$

where $P_S(N)$ is the probability for a signal event to have an nhit above $N$.

![Figure A.1: $P_R(N_2 = 20, \Delta t)$ (red) vs $P_F(N_1 = 60, N_2 = 20, \Delta t)$ (blue).](image)

Figure A.1 shows the measured distributions for $P_F$ and $P_R$ evaluated during the unblinded portion of the dataset; a number of features exist in the observed distribution for $P_F$, the peak at very short $\Delta t$ comes from re-triggers within the detector following a high-nhit event. Figure A.2 shows this comparison in two dimensions, each bin along the Y-axis is normalized to account for the logarithmic change in effective bin size.

Since there exists a number of selections from the threshold values of $N_1$, $N_2$ and $\delta T$, the threshold value for $N_1$ was fixed at the SNO value of 60 (84). This value is motivated
A.3 CAEN Cut

by the fact that time correlated events are more likely to follow highly energetic events or events that can produce cosmogenic isotopes, and the nhit threshold for these processes is not effected significantly by the changes from SNO to SNO+. The threshold value for $N_2$ was selected based upon what nhit of event could serve as a background to the solar or nucleon decay analysis, a conservative value of 20 was selected, which corresponds to an electron kinetic energy of approximately 3 MeV. With the two nhit constraints selected the $\delta t$ window was selected to be as large as possible without incurring significant signal sacrifice, the threshold value of 1 ms was selected. From this a sacrifice of 0.01% percent was determined (83).

A.3 CAEN Cut

I developed a new data cleaning cut, called the “CAEN Cut”, that follows from the AMB Cut from SNO. The AMB Cut attempted to remove events from flashers the dataset by requiring that the integral and peak height of the ESUMH trigger sum (as measured by
A.3 CAEN Cut

the AMB) fall below some threshold value. The CAEN Cut performs a similar function, it calculates the baseline subtracted integral and peak height of the digitized ESUMH trigger signal and places a cut on those values.

The baseline value of each trace is calculated as the average value of the first 20 samples and the 65th to 85th samples. I chose to use two windows, one before the trigger pulse, one after the trigger pulse, to correct for any overall slope across the digitized window. The CAEN window is 104 samples long, the final 19 samples are not used because they often include a large noise pulse. The noise pulse comes from the GT pulse arriving at the front-end and generating electrical noise, it’s typically called “GT pickup”. The noise makes the last \( \approx 20 \) samples of the CAEN trace nearly useless.

The determined baseline is subtracted from the CAEN trace and the integral and maximum peak height are calculated from the samples between the two baseline windows. To pass the CAEN Cut the peak and integral must fall between an upper and lower, nhit dependent, cut value. The cut values are given by

\[
f(n) = C (1 - \sigma(n)) + \sigma(n) (mn + b). \tag{A.2}
\]

Here \( \sigma(x) \) indicates a sigmoid function,

\[
\sigma(x) = \frac{1}{1 + e^{-(x-x_0)/w}} \tag{A.3}
\]

The cut values are meant to be constant value at lower nhit, and then linear with nhit above \( \approx 15 \) nhit, the sigmoid allows for a smooth transition between those two functions; for both the upper and lower threshold the sigmoid position \( (x_0) \) and width \( (w) \) are 15 nhit and 5 nhit respectively. The constant value at lower nhit is \( C \) the slope of the line at higher nhit is given by \( m \) and the value \( b \) is required to be

\[
b = \frac{C}{mx_0} \tag{A.4}
\]

so that there is not discontinuity between the two cut regions. The values for these param-
Table A.1: The upper and lower bounds for the CAEN Cut as described in Eqn. (A.2). Constant values are in ADC Counts, slope values are ADC Counts per nhit.

<table>
<thead>
<tr>
<th></th>
<th>Constant</th>
<th>Slope</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upper Bound</td>
<td>658.8</td>
<td>-6.61</td>
</tr>
<tr>
<td>Lower Bound</td>
<td>-707.0</td>
<td>-15.9</td>
</tr>
</tbody>
</table>

The parameters are given in Table A.1.

The reason for the two cut regions is that at lower nhit the signal peak is smaller than the noise one the ESUMH signal, so the only requirement is that the peak and integral be consistent with a noise only trigger sum. At higher nhit the ESUMH signal scales linearly with nhit, each new hit adds approximately the same amount of height to the trigger pulse.

The cut parameters were determined from two calibration datasets, the first was tagged $^{16}$N events. The second was a sample of $PULSE\_GT$ triggers taken during normal running. The two datasets are used to determine the cut parameters for the two different cut regions. The $^{16}$N data was used to determine cut values for the higher nhit region, the $PULSE\_GT$ data was used for the lower nhit cut values.

For both regions the value of the integral or peak height that include 99% of the events at each nhit is found. Then the parameters of Eqn. (A.2) that best fit those points is determined. Then Eqn. (A.2) with the best fit upper and lower parameters to include 99% of the calibration data become the threshold values for rejecting flasher events. The 99% criteria was chosen arbitrarily to ensure that the fraction of “good” events rejected by this cut was similar to that of other data cleaning cuts. Figure A.3 shows how the ESUMH CAEN trace integral is distributed in the two calibration datasets and for standard physics data taking.
Figure A.3: Distribution of CAEN integral for $PULSE_{GT}$ (a), $^{16}N$ (b), and standard physics triggered (c) events. Integral values are in ADC counts and Z-axis values are counts. The thin red lines indicate the 99% inclusion values and the thick red lines indicate the best fit functions.
Appendix B

Initial Vacuum-Enhanced Mixing Fit

This appendix describes the initial attempt to profile the likelihood spaced for $\Delta m^2_{21}$ using vacuum-enhanced mixing model. A full scan of the 11 free parameters of the vacuum-enhanced neutrino mixing model is too computationally expensive given the speed of mixing simulation, and of the time required to evaluate the likelihood at each point. Instead, only the solar mixing parameters, $\Delta m^2_{21}$ and $\theta_{12}$, and the two enhanced vacuum mixing parameters, $A_e$ and $A_\mu$, are used in the fit. For this the adiabatic neutrino mass state simulation is used, using the non-adiabatic neutrino simulation would be too slow to be completed in reasonable amount of time.

First a profile likelihood search is done over the two standard model parameters, finding the values for $A_e$ and $A_\mu$ that minimize the likelihood function at each sampled value for $\Delta m^2_{12}$ and $\theta_{12}$. This method of search is used in part because the simulation can generate a survival probability for new values of $A_e$ and $A_\mu$ much faster than it can if $\Delta m^2_{21}$ and $\theta_{12}$ are modified, so it’s best to fix the two standard mixing parameters for as many samples as possible. The reason it is faster to generate a survival probability this way is because
the modified vacuum parameters do not change the neutrino’s propagation in the Sun, so that portion of the simulation does not need to be redone for new enhanced vacuum mixing parameters. Even with this scheme for search a single standard model single for $\Delta m_{21}^2$ and $\theta_{12}$ requires approximately 200 iterations, roughly 10 minutes, of search in $A_e$ and $A_\mu$ to find a local minimum.

This search uses only data from Solar experiments for the likelihood. The idea is that if the solar best fit value for $\Delta m_{21}^2$ was much closer to the KamLAND value while allowing for an arbitrary modified vacuum potential, that would be a compelling reason to believe a modified vacuum model is a good description for solar neutrino mixing. Performing the fit this way yielded the best fit value of $\Delta m_{21}^2 = 4.20^{+1.124}_{-0.907}$, profiling over values for $\theta_{12}$. Figure B.1 shows the likelihood space and the profile in $\Delta m_{21}^2$. The bumps in the one-dimensional profile comes from the relative paucity of samples in $\theta_{12}$.

It should also be noted that the KamLAND best fit value is excluded at nearly $3\sigma$; a
significantly larger disagreement than is determined by other global fits \cite{47}. This to be due to the fact that the fit is restricted in so many dimensions, if the solar fluxes were allowed to vary then the disagreement would not be so large. It’s therefore useful to compare the likelihood determined this way to the likelihood constraining modified vacuum parameters to be zero. This shows that in general the likelihood is worse when the modified vacuum parameters are constrained to zero; the KamLAND value is excluded by more than 3$\sigma$.

This scan of the likelihood space shows that even in a vacuum-enhanced mixing model, lower values for $\Delta m_{21}^2$ are preferred by solar neutrino measurements. The vacuum-enhanced parameters do however allow for a better fit to data, and do relieve the tension between the KamLAND and solar measurements for $\Delta m_{21}^2$ significantly, but not entirely. The exact level of the disagreement between the KamLAND and solar can’t be well estimated using this method, because many of the parameters that are correlated with $\Delta m_{21}^2$ are artificially fixed and not allowed to vary.
References

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