

Simultaneous Measurement of Oscillation Parameters in Beam and Atmospheric Neutrino Data from Tokai-to-Kamioka and Super-Kamiokande Experiments



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A thesis submitted for the degree of

Doctor of Philosophy

Michaelmas 2022

Abstract

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1

Introduction

2

Neutrino Oscillation Physics

When first proposed, neutrinos were expected to be massless fermions that only interact through weak and gravitational forces. This meant they were very difficult to detect as they can pass through significant amounts of matter without interacting. Despite this, experimental neutrino physics has developed with many different detection techniques and neutrino sources being used today. In direct tension with standard model physics, neutrinos have been determined to oscillate between different lepton flavours, requiring them to have mass.

The observation techniques which led to the discovery of the neutrino are documented in section 2.1. The theory underpinning neutrino oscillation is described in section 2.2 and includes the approximations which can be made to simplify the understanding of neutrino oscillation in the two-flavour approximation. Past, current, and future neutrino experiments are detailed in section 2.3, including the reactor, atmospheric, and long-baseline accelerator neutrino sources that have been used to successfully constrain oscillation parameters. Finally, the current state of oscillation parameter measurements are summarised in section 2.4.

2.1 Discovery of Neutrinos

At the start of the 20th century, the electrons emitted from the β -decay of the nucleus were found to have a continuous energy spectrum [1, 2]. This observation seemingly broke the energy conservation invoked within that period's nuclear models. Postulated in 1930 by Pauli as the solution to this problem, the neutrino (originally termed "neutron") was theorized to be an electrically neutral spin-1/2 fermion with a mass of the same order of magnitude as the electron [3]. This neutrino was to be emitted with the electron in β -decay to alleviate the apparent breaking of energy conservation. As a predecessor of today's weak interaction model, Fermi's theory of β -decay developed the understanding by coupling the four constituent particles: electron, proton, neutron, and neutrino, into a consistent model [4].

Whilst Pauli was not convinced of the ability to detect neutrinos, the first observations of the particle were made in the mid-1950s when neutrinos from a reactor were observed via the inverse β -decay (IBD) process, $\bar{\nu}_e + p \rightarrow n + e^+$ [5, 6]. The detector consisted of two parts: a neutrino interaction medium and a liquid scintillator. The interaction medium was built from two water tanks. These were loaded with cadmium chloride to allow increased efficiency of neutron capture. The positron emitted from IBD annihilates, $e^+ + e^- \rightarrow 2\gamma$, generating a prompt signal and the neutron is captured on the cadmium via $n + ^{108}Cd \rightarrow ^{109*}Cd \rightarrow ^{109}Cd + \gamma$, producing a delayed signal. An increase in the coincidence rate was observed when the reactor was operating which was interpreted as interactions from neutrinos generated in the reactor.

After the discovery of the ν_e , the natural question of how many flavours of neutrino exist was asked. In 1962, a measurement of the ν_μ was conducted at the Brookhaven National Laboratory [7]. A proton beam was directed at a beryllium target, generating a π -dominated beam which then decayed via $\pi^\pm \rightarrow \mu^\pm + (\nu_\mu, \bar{\nu}_\mu)$, and the subsequent interactions of the ν_μ were observed. As the subsequent interaction of the neutrino generates muons rather than

electrons, it was determined the ν_μ was fundamentally different from ν_e . The final observation to be made was that of the ν_τ from the DONUT experiment [8]. Three neutrinos seem the obvious solution as it mirrors the known number of charged lepton (as they form weak isospin doublets) but there could be evidence of more. Several neutrino experiments have found anomalous results [9, 10] which could be attributed to “sterile” neutrinos. These hypothesised particles are not affected by gauge interactions in the standard model so their presence can only be inferred through the observation of non-standard oscillation modes. However, cosmological observations indicate the number of neutrino species $N_{eff} = 2.99 \pm 0.17$ [11], as measured from the cosmic microwave background power spectrum. LEP also measured the number of active neutrino flavours to be $N_\nu = 2.9840 \pm 0.0082$ [12] from measurements of the Z-decay width, but this does not strongly constrain the number of sterile neutrinos.

2.2 Theory of Neutrino Oscillation

A neutrino generated with lepton flavour α can change into a different lepton flavour β after propagating some distance. This phenomenon is called neutrino oscillation and requires that neutrinos must have a non-zero mass. This behaviour has been characterised by the Pontecorvo-Maki-Nakagawa-Sakata (PMNS) [13–15] mixing matrix which describes how the flavour and mass of neutrinos are associated. This is analogous to the Cabibbo-Kobayashi-Maskawa (CKM) [16] matrix measured in quark physics.

2.2.1 Three Flavour Oscillations

The PMNS parameterisation defines three flavour eigenstates, ν_e , ν_μ and ν_τ (indexed ν_α), which are eigenstates of the weak interaction and three mass eigenstates, ν_1 , ν_2 and ν_3 (indexed ν_i). Each mass eigenstate is the superposition

of all three flavour states,

$$|\nu_i\rangle = \sum_{\alpha} U_{\alpha i} |\nu_{\alpha}\rangle. \quad (2.1)$$

Where U is the 3×3 PMNS matrix which is unitary and connects the mass and flavour eigenstates.

The weak interaction couples to flavour eigenstates so neutrinos interact with leptons of the same flavour. The propagation of a neutrino flavour eigenstate, in a vacuum, can be re-written as a plane-wave solution to the time-dependent Schrödinger equation,

$$|\nu_{\alpha}(t)\rangle = \sum_i U_{\alpha i}^* |\nu_i\rangle e^{-i\phi_i}. \quad (2.2)$$

The probability of observing a neutrino of flavour eigenstate β from one which originated as flavour α can be calculated as,

$$P(\nu_{\alpha} \rightarrow \nu_{\beta}) = |\langle \nu_{\beta} | \nu_{\alpha}(t) \rangle|^2 = \sum_{i,j} U_{\alpha i}^* U_{\beta i} U_{\alpha j} U_{\beta j}^* e^{-i(\phi_j - \phi_i)} \quad (2.3)$$

The ϕ_i term can be expressed in terms of the energy, E_i , and magnitude of the three momenta, p_i , of the neutrino, $\phi_i = E_i t - p_i x$ (t and x being time and position coordinates). Therefore,

$$\phi_j - \phi_i = E_j t - E_i t - p_j x + p_i x. \quad (2.4)$$

For a relativistic particle, $E_i \gg m_i$,

$$p_i = \sqrt{E_i^2 - m_i^2} \approx E_i - \frac{m_i^2}{2E_i}. \quad (2.5)$$

Making the approximations that neutrinos are relativistic, the mass eigenstates were created with the same energy and that $x = L$, where L is the distance traveled by the neutrino, Equation 2.4 then becomes

$$\phi_j - \phi_i = \frac{\Delta m_{ij}^2 L}{2E}, \quad (2.6)$$

where $\Delta m_{ij}^2 = m_i^2 - m_j^2$. This, combined with further use of unitarity relations results in Equation 2.3 becoming

$$\begin{aligned} P(\nu_\alpha \rightarrow \nu_\beta) &= \delta_{\alpha\beta} - 4 \sum_{i>j} \Re \left(U_{\alpha i}^* U_{\beta i} U_{\alpha j} U_{\beta j}^* \right) \sin^2 \left(\frac{\Delta m_{ij}^2 L}{4E} \right) \\ &\quad + (-) 2 \sum_{i>j} \Im \left(U_{\alpha i}^* U_{\beta i} U_{\alpha j} U_{\beta j}^* \right) \sin \left(\frac{\Delta m_{ij}^2 L}{2E} \right) \end{aligned} \quad (2.7)$$

Where $\delta_{\alpha\beta}$ is the Kronecker delta function and the negative sign on the last term is included for the oscillation probability of antineutrinos. As an important point to note, the observation of oscillation probability requires that a non-zero value of Δm_{ij}^2 , which in turn requires that neutrinos have differing masses.

Typically, the PMNS matrix is parameterised into three mixing angles, a charge parity (CP) violating phase δ_{CP} , and two Majorana phases $\alpha_{1,2}$,

$$U = \underbrace{\begin{pmatrix} 1 & 0 & 0 \\ 0 & c_{23} & s_{23} \\ 0 & -s_{23} & c_{23} \end{pmatrix}}_{\text{Atmospheric, Accelerator}} \underbrace{\begin{pmatrix} c_{13} & 0 & s_{13} e^{-i\delta_{CP}} \\ 0 & 1 & 0 \\ -s_{13} e^{-i\delta_{CP}} & 0 & c_{13} \end{pmatrix}}_{\text{Reactor, Accelerator}} \times \underbrace{\begin{pmatrix} c_{12} & s_{12} & 0 \\ -s_{12} & c_{12} & 0 \\ 0 & 0 & 1 \end{pmatrix}}_{\text{Reactor, Solar}} \underbrace{\begin{pmatrix} e^{i\alpha_1/2} & 0 & 0 \\ 0 & e^{i\alpha_2/2} & 0 \\ 0 & 0 & 1 \end{pmatrix}}_{\text{Majorana}}. \quad (2.8)$$

Where $s_{ij} = \sin(\theta_{ij})$ and $c_{ij} = \cos(\theta_{ij})$. The oscillation parameters are often grouped: (1,2) as “solar”, (2,3) as “atmospheric” and (1,3) as “reactor”. Many neutrino experiments aim to measure the PMNS parameters from a wide array of origins, as is the purpose of this thesis.

The Majorana phase, $\alpha_{1,2}$, included within the fourth matrix in Equation 2.8 is only included for completeness. For an oscillation analysis experiment, any terms containing this phase disappear due to taking the expectation value of the PMNS matrix. Measurements of these phases are typically performed by experiments searching for neutrino-less double β -decay [17].

A two flavour approximation can be obtained when one assumes the third mass eigenstate is degenerate with another. As discussed in section 2.3, it is found that $\Delta m_{21}^2 \ll |\Delta m_{31}^2|$. This results in the two flavour approximation being reasonable for understanding the features of the oscillation. In this two flavour case, the mixing matrix becomes,

$$U_{2\text{ Flav.}} = \begin{pmatrix} \cos(\theta) & \sin(\theta) \\ -\sin(\theta) & \cos(\theta) \end{pmatrix}. \quad (2.9)$$

This culminates in the oscillation probability,

$$\begin{aligned} P(\nu_\alpha \rightarrow \nu_\alpha) &= 1 - \sin^2(2\theta) \sin^2\left(\frac{\Delta m^2 L}{4E}\right), \\ P(\nu_\alpha \rightarrow \nu_\beta) &= \sin^2(2\theta) \sin^2\left(\frac{\Delta m^2 L}{4E}\right). \end{aligned} \quad (2.10)$$

Where $\alpha \neq \beta$. For a fixed neutrino energy, the oscillation probability is a sinusoidal function depending upon the distance over which the neutrino propagates. The frequency and amplitude of oscillation are dependent upon $\Delta m^2/4E$ and $\sin^2 2\theta$, respectively. The oscillation probabilities presented thus far assume $c = 1$, where c is the speed of light in vacuum. In more familiar units, the maximum oscillation probability for a fixed value of θ is given at $L[\text{km}] / E[\text{GeV}] \sim 1.27/\Delta m^2$. It is this calculation that determines the best L/E value for a given experiment to be designed around for measurements of a specific value of Δm^2 .

2.2.2 The MSW Effect

The theory of neutrino oscillation in a vacuum has been described in subsection 2.2.1. However, the beam neutrinos and atmospheric neutrinos originating from below the horizon propagate through matter in the Earth. The coherent scattering of neutrinos from a material target modifies the Hamiltonian of the system. This results in a change in the oscillation probability. Notably, charged current scattering ($\nu_e + e^- \rightarrow \nu_e + e^-$, propagated by a W boson) only affects electron neutrinos whereas the neutral current scattering ($\nu_l + l^- \rightarrow \nu_l + l^-$,

propagated by a Z^0 boson) interacts through all neutrino flavours equally. In the two-flavour approximation, the effective mixing parameter becomes

$$\sin^2(2\theta) \rightarrow \sin^2(2\theta_m) = \frac{\sin^2(2\theta)}{(A/\Delta m^2 - \cos(2\theta))^2 + \sin^2(2\theta)}, \quad (2.11)$$

where $A = 2\sqrt{2}G_F N_e E$, N_e is the electron density of the medium and G_F is Fermi's constant. It is clear to see that there exists a value of $A = \Delta m^2 \cos(2\theta)$ for $\Delta m^2 > 0$ which results in a divergent mixing parameter. This resonance is termed the Mikheyev-Smirnov-Wolfenstein (MSW) effect (or more colloquially, the matter resonance) which regenerates the electron neutrino component of the neutrino flux [18–20]. The density at which the resonance occurs is given by

$$N_e = \frac{\Delta m^2 \cos(2\theta)}{2\sqrt{2}G_F E}. \quad (2.12)$$

At densities lower than this critical value, the oscillation probability will be much closer to that of vacuum oscillation. For antineutrinos, $N_e \rightarrow -N_e$ [21]. The resonance occurring from the MSW effect depends on the sign of Δm^2 . Therefore, any neutrino oscillation experiment which observes neutrinos and antineutrinos which have propagated through matter can have some sensitivity to the ordering of the neutrino mass eigenstates.

2.3 Neutrino Oscillation Measurements

As evidence of beyond standard model physics, the 2015 Nobel Prize in Physics was awarded to the Super-Kamiokande (SK) [22] and Sudbury Neutrino Observatory (SNO) [23] collaborations for the first definitive observation of solar and atmospheric neutrino oscillation [24]. Since then, the field has seen a wide array of oscillation measurements from a variety of neutrino sources. As seen in subsection 2.2.1, the neutrino oscillation probability is dependent on the ratio of the propagation baseline, L , to the neutrino energy, E . It is this ratio that determines the type of neutrino oscillation a particular experiment is sensitive to.

As illustrated in Figure 2.1, there are many neutrino sources that span a wide range of energies. The least energetic neutrinos are from diffuse supernovae and terrestrial neutrinos at $O(1)$ MeV whereas the most energetic neutrinos originate from atmospheric and galactic neutrinos of $> O(1)$ TeV.

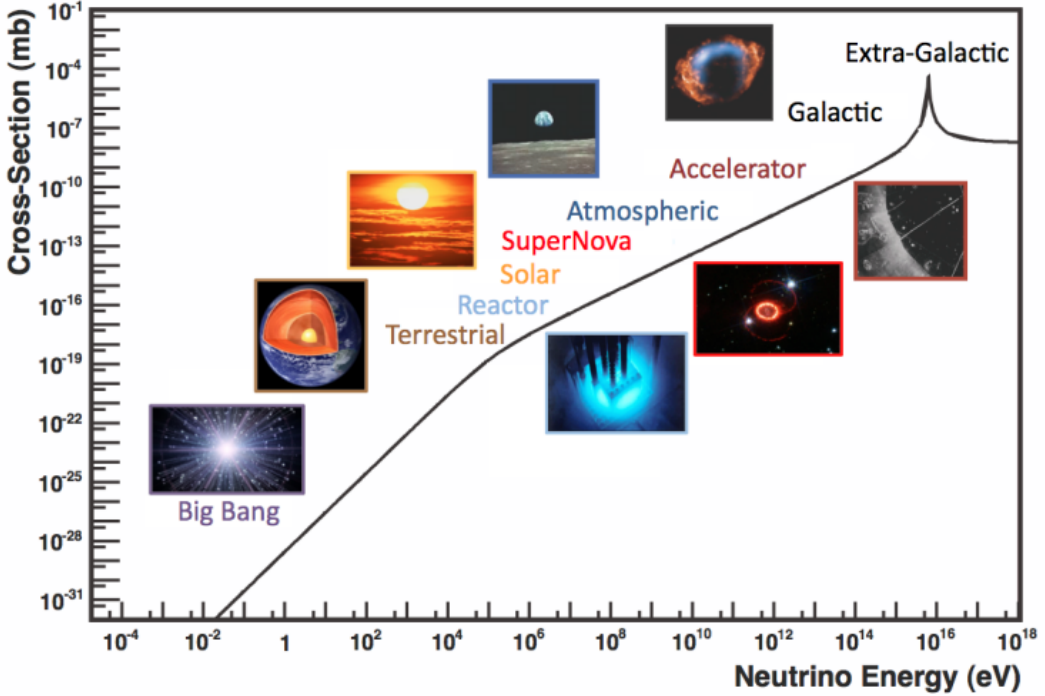


Figure 2.1: The cross-section of neutrinos from various natural and man-made sources as a function of neutrino energy. Taken from [25]

2.3.1 Solar Neutrinos

Solar neutrinos are emitted from fusion reaction chains at the center of the Sun. The solar neutrino flux, given as a function of neutrino energy for different fusion and decay chains, is illustrated in Figure 2.2. Whilst proton-proton fusion generates the largest flux of neutrinos, the neutrinos are of low energy and are difficult to reconstruct due to the IBD interaction threshold of 1.8MeV. Consequently, most experiments focus on the neutrinos from the decay of 8B (via ${}^8B \rightarrow {}^8Be^* + e^+ + \nu_e$), which are higher energy.

The first measurements of solar neutrinos observed a significant reduction in the event rate compared to predictions from the Standard Solar Model [27, 28]. A

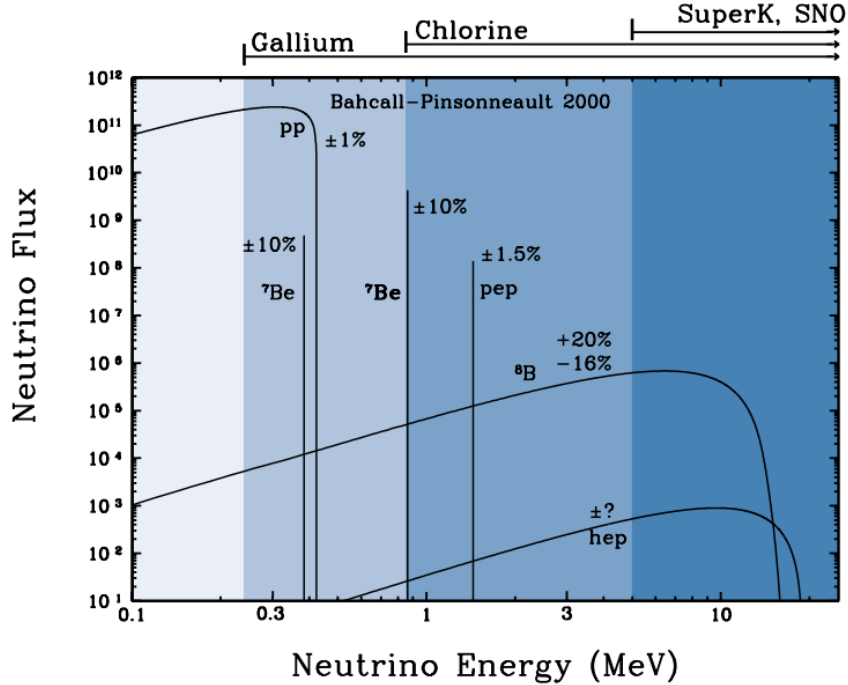
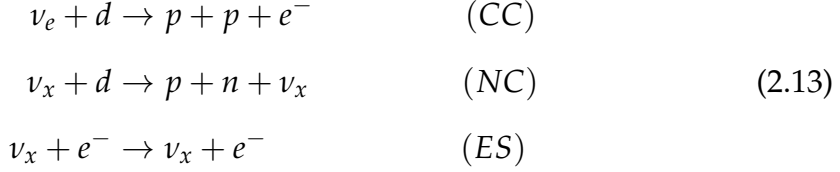


Figure 2.2: The solar neutrino flux as a function of neutrino energy for various fusion reactions and decay chains as predicted by the Standard Solar Model. Taken from [26].

proposed solution to this “solar neutrino problem” was $\nu_e \leftrightarrow \nu_\mu$ oscillations in a precursory version of the PMNS model [29]. The Kamiokande [30], Gallex [31] and Sage [32] experiments confirmed the ~ 0.5 factor deficit of solar neutrinos.

The conclusive solution to this problem was determined by the SNO collaboration [33]. Using a deuterium water target to observe 8B neutrinos, the event rate of charged current (CC), neutral current (NC), and elastic scattering (ES) interactions (Given in Equation 2.13) was simultaneously measured. CC events can only occur for electron neutrinos, whereas the NC channel is agnostic to neutrino flavour, and the ES reaction has a slight excess sensitivity to electron neutrino interactions. This meant that there were direct measurements of the ν_e and ν_x neutrino flux. It was concluded that the CC and ES interaction rates were consistent with the deficit previously observed. Most importantly, the NC reaction rate was only

consistent with the others under the hypothesis of flavour transformation.



Many experiments have since measured the neutrino flux of different interaction chains within the sun [34–36]. The most recent measurement was that of CNO neutrinos which were recently observed with 5σ significance by the Borexino collaboration. Future neutrino experiments aim to further these spectroscopic measurements of different fusion chains within the Sun [37–39].

2.3.2 Atmospheric Neutrinos

The interactions of primary cosmic ray protons in Earth's upper atmosphere generate showers of energetic hadrons. These are mostly pions and kaons which when they decay produce a natural source of neutrinos spanning energies of MeV to TeV [40]. The main decay is via

$$\begin{aligned}
 \pi^\pm &\rightarrow \mu^\pm + (\nu_\mu, \bar{\nu}_\mu) \\
 \mu^\pm &\rightarrow e^\pm + (\nu_\mu, \bar{\nu}_\mu) + (\nu_e, \bar{\nu}_e)
 \end{aligned} \tag{2.14}$$

such that for a single pion decay, three neutrinos are typically produced. The atmospheric neutrino flux energy spectra as predicted by the Bartol [41], Honda [42–44], and FLUKA [45] models are illustrated in Figure 2.3. The flux distribution peaks at an energy of $O(10)\text{GeV}$. The uncertainties associated with these models are dominated by the hadronic production of kaon and pions as well as the primary cosmic flux.

Unlike long-baseline experiments which have a fixed baseline, the distance atmospheric neutrinos propagate is dependent upon the zenith angle at which they interact. This is illustrated in Figure 2.4. Neutrinos that are generated directly above the detector ($\cos(\theta) = 1.0$) have a baseline equivalent to the

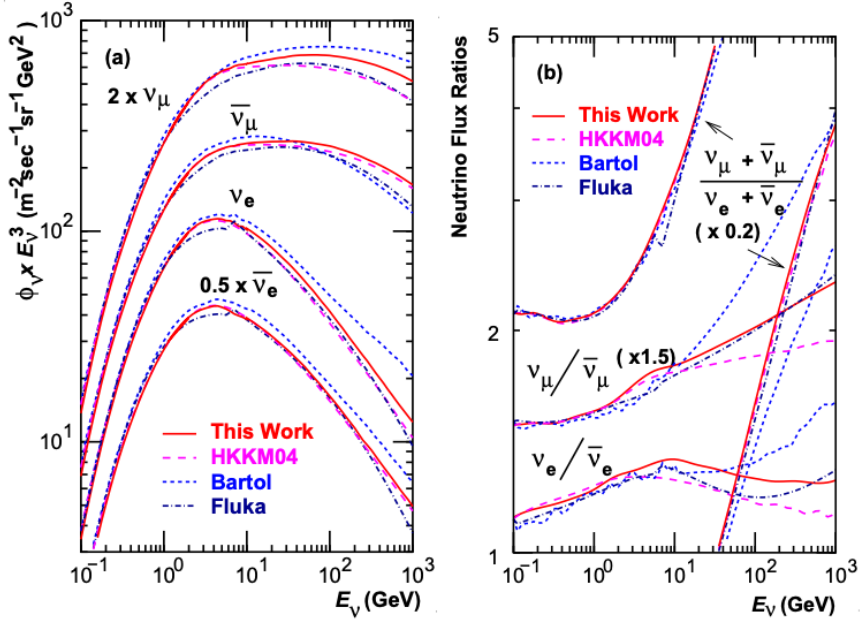


Figure 2.3: Left panel: The atmospheric neutrino flux for different neutrino flavours as a function of neutrino energy as predicted by the 2007 Honda model (“This work”) [42], the 2004 Honda model (“HKKM04”)[43], the Bartol model [41] and the FLUKA model [45]. Right panel: The ratio of the muon to electron neutrino flux as predicted by all the quoted models. Both figures taken from [42].

height of the atmosphere whereas neutrinos that interact directly below the detector ($\cos(\theta) = -1.0$) have to travel a length equal to the diameter of the Earth. This means atmospheric neutrinos have a baseline that varies from $O(20)$ km to $O(6 \times 10^3)$ km. Any neutrino generated at or below the horizon will be subject to matter effects as they propagate through the Earth.

Figure 2.5 highlights the neutrino flux as a function of the zenith angle for different slices of neutrino energy. For medium to high-energy neutrinos (and to a lesser degree for low-energy neutrinos), the flux is approximately symmetric around $\cos(\theta) = 0$. To the accuracy of this approximation, the systematic uncertainties associated with atmospheric flux for comparing upward-going and down-going neutrino cancels. This allows the down-going events, which are mostly insensitive to oscillation probabilities, to act as an unoscillated prediction (similar to a near detector in an accelerator neutrino experiment).

Precursory hints of atmospheric neutrinos were observed in the mid-1960s searching for $\nu_\mu^- + X \rightarrow X^* + \mu^\pm$ [47], although it was called an anomaly

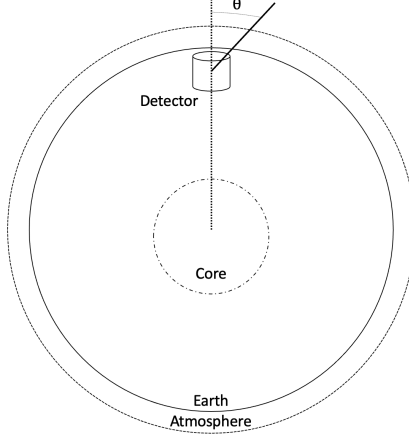


Figure 2.4: A diagram illustrating the definition of zenith angle as used in the Super Kamiokande experiment [46].

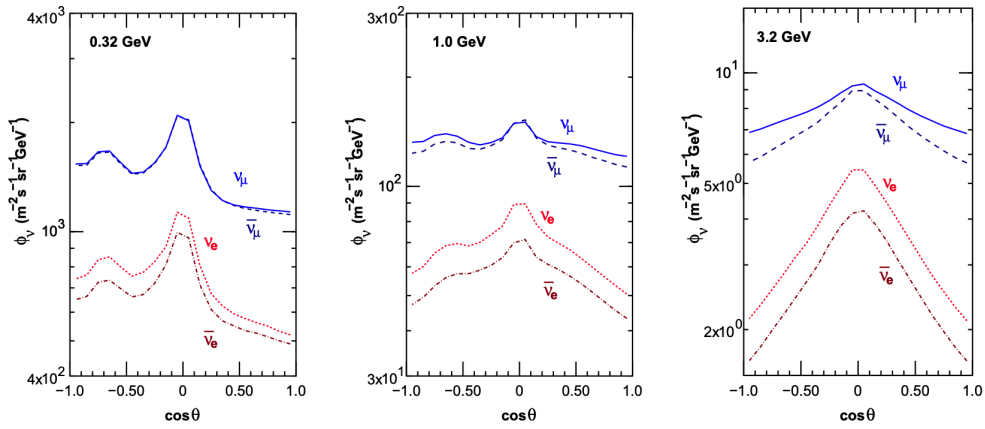


Figure 2.5: Prediction of $\nu_e, \bar{\nu}_e, \nu_\mu, \bar{\nu}_\mu$ fluxes as a function of zenith angle as calculated by the HKKM model [44]. The left, middle and right panels represent three values of neutrino energy, 0.32GeV, 1.0GeV and 3.2GeV respectively. Predictions for other models including Bartol [41], Honda [42] and FLUKA [45] are given in [46].

at the time of measurement. This was succeeded with the IMB-3 [48] and Kamiokande [49] experiments which measured the ratio of muon neutrinos compared to electron neutrinos $R(\nu_\mu/\nu_e)$. Both experiments were found to have a consistent deficit of muon neutrinos, with $R(\nu_\mu/\nu_e) = 0.67 \pm 0.17$ and $R(\nu_\mu/\nu_e) = 0.60^{+0.07}_{-0.06} \pm 0.05$. Super-Kamiokande (SK) [46] extended this analysis by fitting oscillation parameters in $P(\nu_\mu \rightarrow \nu_\tau)$ which found best fit parameters $\sin^2(2\theta) > 0.92$ and $1.5 \times 10^{-3} < \Delta m^2 < 3.4 \times 10^{-3}$ eV².

Since then, atmospheric neutrino experiments have been making precision

measurements of the $\sin^2(\theta_{23})$ and Δm_{32}^2 oscillation parameters. Atmospheric neutrino oscillation is dominated by $P(\nu_\mu \rightarrow \nu_\tau)$, where SK observed a 4.6σ discovery of ν_τ appearance [50]. Figure 2.6 illustrates the current estimates on the atmospheric mixing parameters from a wide range of atmospheric and accelerator neutrino observatories.

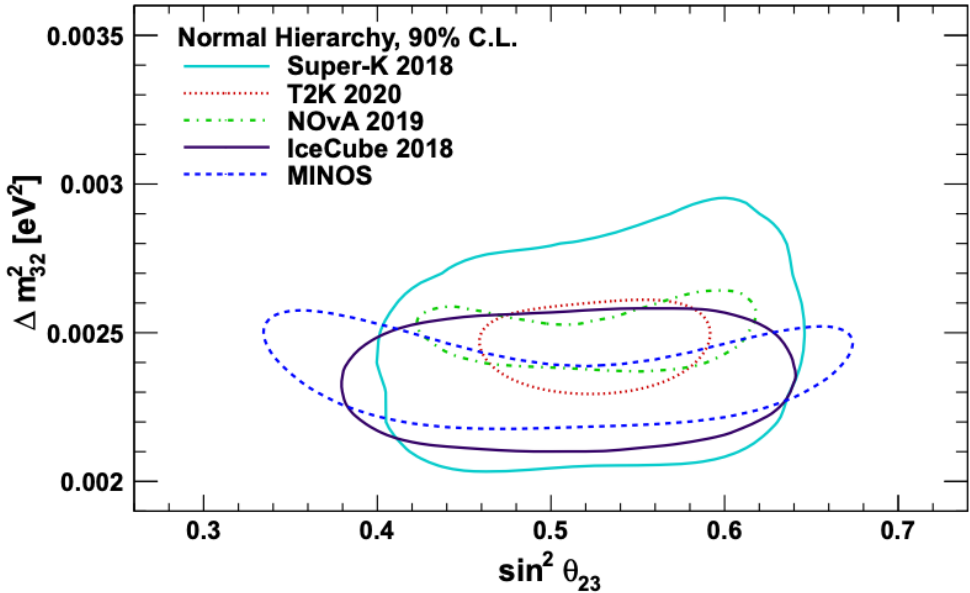


Figure 2.6: Constraints on the atmospheric oscillation parameters, $\sin^2(\theta_{23})$ and Δm_{32}^2 , from atmospheric and long baseline experiments: SK [51], T2K [52], NOvA [53], IceCube [54] and MINOS [55]. Figure taken from [56].

2.3.3 Accelerator Neutrinos

The concept of using a man-made “neutrino beam” was first realised in 1962 [57]. Since then, many experiments have followed which all use the same fundamental concepts. Typically, a proton beam is aimed at a target producing charged mesons that decay to neutrinos. The mesons can be sign-selected by the use of magnetic focusing horns to generate a neutrino or antineutrino beam. Pions are the primary meson that decay and depending on the orientation of the magnetic field, a muon (anti-)neutrino beam is generated via $\pi^+ \rightarrow \mu^+ + \nu_\mu$ or $\pi^- \rightarrow \mu^- + \bar{\nu}_\mu$. The decay of muons and kaons does result in an irreducible intrinsic electron neutrino background. In T2K, this background contamination is $O(< 1\%)$ [58]. There is

also an approximately $\sim 5\%$ “wrong-sign” neutrino background of $\bar{\nu}_\mu$ generated via the same decays. As the beam is generated by proton interactions (rather than anti-proton interactions), the wrong-sign component in the antineutrino beam is larger when operating in neutrino mode.

Tuning the proton energy in the beam and using beam focusing techniques allows the neutrino energy to be set to a value that maximises the disappearance oscillation probability in the L/E term in Equation 2.10. This means that accelerator experiments are typically more sensitive to the mixing parameters as compared to a natural neutrino source. However, the disadvantage compared to atmospheric neutrino experiments is that the baseline has to be shorter due to the lower flux. Consequently, there is typically less sensitivity to matter effects and the ordering of the neutrino mass eigenstates.

A neutrino experiment measures

$$R(\vec{x}) = \Phi(E_\nu) \times \sigma(E_\nu) \times \epsilon(\vec{x}) \times P(\nu_\alpha \rightarrow \nu_\beta), \quad (2.15)$$

where $R(\vec{x})$ is the event rate of neutrinos at position \vec{x} , $\Phi(E_\nu)$ is the flux of neutrinos with energy E_ν , $\sigma(E_\nu)$ is the cross-section of the neutrino interaction and $\epsilon(\vec{x})$ is the efficiency and resolution of the detector. In order to leverage the most out of an accelerator neutrino experiment, the flux and cross-section systematics need to be constrained. This is typically done via the use of a “near detector”, situated at a baseline of $O(1)\text{km}$. This detector observes the unoscillated neutrino flux and constrains the parameters used within the flux and cross-section model.

The first accelerator experiments to precisely measure oscillation parameters were MINOS [59] and K2K [60]. These experiments confirmed the ν_μ disappearance seen in atmospheric neutrino experiments by finding consistent parameter values for $\sin^2(\theta_{23})$ and Δm_{23}^2 . The current generation of accelerator neutrino experiments, T2K and NO ν A extended this field by observing $\bar{\nu}_\mu \rightarrow \bar{\nu}_e$ and lead the sensitivity to atmospheric mixing parameters as seen in Figure 2.6 [61]. The two experiments differ in their peak neutrino energy, baseline, and detection technique. The NO ν A experiment is situated at a baseline of 810km

from the NuMI beamline which delivers 2GeV neutrinos. The T2K neutrino beam is peaked around 0.6GeV and propagates 295km. The NO ν A experiment also uses functionally identical detectors (near and far) which allow the approximate cancellation of detector systematics whereas T2K uses a plastic scintillator technique at the near detector and a water Cherenkov far detector. The future generation experiments DUNE [62] and Hyper-Kamiokande [63] will succeed these experiments as the high-precision era of neutrino oscillation parameter measurements develops.

Several anomalous results have been observed in the LSND [9] and Mini-BooNE [10] detectors which were designed with purposefully short baselines. Parts of the neutrino community attributed these results to oscillations induced by a fourth “sterile” neutrino [64] but several searches in other experiments, MicroBooNE [65] and KARMEN [66], found no hints of additional neutrino species. The solution to the anomalous results is still being determined.

2.3.4 Reactor Neutrinos

As illustrated in the first discovery of neutrinos (section 2.1), nuclear reactors are a very useful man-made source of electron antineutrinos. For reactors that use low-enriched uranium ^{235}U as fuel, the antineutrino flux is dominated by the β -decay fission of ^{235}U , ^{238}U , ^{239}Pu and ^{241}Pu [67] as illustrated in Figure 2.7.

Due to their low energy, reactor electron antineutrinos predominantly interact via the inverse β -decay (IBD) interaction. The typical signature contains two signals delayed by $O(200)\mu\text{s}$; firstly the prompt photons from positron annihilation, and secondly the photons emitted ($E_{tot}^\gamma = 2.2\text{MeV}$) from de-excitation after neutron capture on hydrogen. Searching for both signals improves the detector’s ability to distinguish between background and signal events [69].

There are many short baseline experiments ($L \sim O(1)\text{km}$) that have measured the $\sin^2(\theta_{13})$ and Δm_{23}^2 oscillation parameters. Daya Bay [70], RENO [71] and Double Chooz [72] have all provided precise measurements, with the first discovery of a non-zero θ_{13} made by Daya Bay and RENO (and complemented by T2K



Figure 2.7: Reactor electron antineutrino fluxes for ^{235}U (Black), ^{238}U (Green), ^{239}Pu (Purple), and ^{241}Pu (Orange) isotopes. The inverse β -decay cross-section (Blue) and corresponding measurable neutrino spectrum (Red) are also given. Top panel: Schematic of Inverse β -decay interaction including the eventual capture of the emitted neutron. This capture emits a γ -ray which provides a second signal of the event. Taken from [68].

[72]). The constraints on $\sin^2(\theta_{13})$ by the reactor experiments lead the field and are often used as external inputs to accelerator neutrino experiments to improve their sensitivity to δ_{CP} and mass hierarchy determination. JUNO-TAO [73], a small collaboration within the larger JUNO experiment, is a next-generation reactor experiment that aims to precisely measure the isotopic antineutrino yields from the different fission chains.

Kamland [74] is the only experiment to have observed reactor neutrinos using a long baseline (flux weighted averaged baseline of $L \sim 180\text{km}$) which allows it to have sensitivity to Δm_{12}^2 . Combined with the SK solar neutrino experiment, the combined analysis puts the most stringent constraint on Δm_{12}^2 [75].

2.4 Summary Of Oscillation Parameter Measurements

Since observing the first evidence of neutrino oscillations in the late 1990's, numerous measurements of the mixing parameters have been made. Many experiments use neutrinos as a tool for discovery of new physics (diffuse supernova background, neutrinoless double beta decay and others) so the PMNS parameters are summarised in the Particle Data Group (PDG) review tables. The analysis presented in this thesis focuses on the 2020 T2K oscillation analysis presented in [76] where the 2020 PDG constraints [77] were used. These constraints are outlined in Table 2.1.

Parameter	2020 Constraint
$\sin^2(\theta_{12})$	0.307 ± 0.013
Δm_{21}^2	$(7.53 \pm 0.18) \times 10^{-5} \text{ eV}^2$
$\sin^2(\theta_{13})$	$(2.18 \pm 0.07) \times 10^{-2}$
$\sin^2(\theta_{23})$ (I.H.)	0.547 ± 0.021
$\sin^2(\theta_{23})$ (N.H.)	0.545 ± 0.021
Δm_{32}^2 (I.H.)	$(-2.546^{+0.034}_{-0.040}) \times 10^{-3} \text{ eV}^2$
Δm_{32}^2 (N.H.)	$(2.453 \pm 0.034) \times 10^{-3} \text{ eV}^2$

Table 2.1: The 2020 Particle Data Group constraints of the oscillation parameters taken from [77]. The value of Δm_{23}^2 is given for both normal hierarchy (N.H.) and inverted hierarchy (I.H.) and $\sin^2(\theta_{23})$ is broken down by whether its value is below (Q1) or above (Q2) 0.5.

The $\sin^2(\theta_{13})$ measurement stems from the electron antineutrino disappearance, $P(\bar{\nu}_e \rightarrow \bar{\nu}_e)$, and is take as the average best-fit from the combination of Daya Bay, Reno and Double Chooz. It is often used as a prior uncertainty within other neutrino oscillation experiments, typically termed the reactor constraint. The $\sin^2(\theta_{12})$ parameter is predominantly measured through electron neutrino disappearance, $P(\nu_e \rightarrow \nu_{\mu,\tau})$, in solar neutrino experiments. The long-baseline reactor neutrino experiment Kamland also has sensitivity to this parameter and is used in a joint fit to solar data from SNO and SK, using the reactor constraint. Measurements of $\sin^2(\theta_{23})$ are made by long-baseline and atmospheric neutrino experiments. The PDG value is a joint fit of T2K, NO ν A , MINOS and IceCube DeepCore experiments. The latest T2K-only measurement, provided at

Neutrino2020 and is the basis of this thesis, is given as $\sin^2(\theta_{23}) = 0.546^{+0.024}_{-0.046}$ [76]. The PDG constraint on Δm_{12}^2 is provided by the KamLAND experiment using solar and geoneutrino data. This measurement utilised a $\sin^2(\theta_{13})$ constraint from accelerator (T2K, MINOS) and reactor neutrino (Daya Bay, RENO, Double Chooz) experiments. Accelerator measurements make some of the most stringent constraints on Δm_{23}^2 although atmospheric experiments have more sensitivity to the mass hierarchy determination. The PDG performs a joint fit of accelerator and atmospheric data, in both normal and inverted hierarchy separately. The latest T2K-only result is $\Delta m_{32}^2 = 2.49^{+0.058}_{-0.082} \times 10^{-3} \text{ eV}^2$ favouring normal hierarchy [76]. The value of δ_{CP} is largely undetermined. CP-conserving values of 0 and π were rejected with $\sim 2\sigma$ intervals, as published in Nature, although more recent analysis have reduced the rejection intervals to 90%. Since the 2020 PDG publication, there has been a new measurement of $\sin^2(\theta_{13}) = (2.20 \pm 0.07) \times 10^{-2}$ [78], alongside updated Δm_{23}^2 and $\sin^2(\theta_{23})$ measurements.

Throughout this thesis, several sample spectra predictions and contours are presented which require oscillation parameters to be assumed. Table 2.2 defines two sets of oscillation parameters, with “Asimov A” set being close to the preferred values from a previous T2K-only fit [79] and “Asimov B” being CP-conserving and further from maximal θ_{23} mixing.

Parameter	Asimov A	Asimov B
Δm_{12}^2	$7.53 \times 10^{-5} \text{ eV}^2$	
Δm_{32}^2	$2.509 \times 10^{-3} \text{ eV}^2$	
$\sin^2(\theta_{12})$	0.304	
$\sin^2(\theta_{13})$	0.0219	
$\sin^2(\theta_{23})$	0.528	0.45
δ_{CP}	-1.601	0.0

Table 2.2: Reference values of the neutrino oscillation parameters for two different oscillation parameter sets.

3

T2K and SK Experiment Overview

As the successor of the Kamiokande experiment, the Super-Kamiokande (SK) collaboration has been leading atmospheric neutrino oscillation analyses for over two decades. The detector has provided some of the strongest constraints on proton decay and the first precise measurements of the Δm_{23}^2 and $\sin^2(\theta_{23})$ neutrino oscillation parameters. The history, detection technique, and operation of the SK detector is described in section 3.1.

The Tokai-to-Kamioka (T2K) experiment was one of the first long-baseline experiments to use both neutrino and antineutrino beams to precisely measure the charge parity violation within the neutrino sector. The T2K experiment observed the first hints of a non-zero $\sin^2(\theta_{13})$ measurement and continues to lead the field with the constraints it provides on $\sin^2(\theta_{13})$, $\sin^2(\theta_{23})$, Δm_{23}^2 and δ_{CP} . In section 3.2, the techniques that T2K use to generate the neutrino beam and constrain systematic parameter through near detector constraints are described.

3.1 The Super-Kamiokande Experiment

The SK experiment began taking data in 1996 [80] and has had many modifications throughout its operation. There have been seven defined periods of data taking as noted in Table 3.1. Data taking began in SK-I which ran for five

years. Between the SK-I and SK-II periods, approximately 55% of the PMTs were damaged during maintenance [81]. Those that survived were equally distributed throughout the detector in the SK-II era, which resulted in a reduced 19% photo-coverage. From SK-III onwards, repairs to the detector meant the full suite of PMTs was operational recovering the 40% photocoverage. Before the start of SK-IV, the data acquisition and electronic systems were upgraded. Between SK-IV and SK-V, a significant effort was placed into tank open maintenance and repair/replacement of defective PMTs, a task for which the author of this thesis was required. Consequently, the detector conditions were significantly different between the two operational periods. SK-VI marked the start of the SK-Gd era, with the detector being doped with gadolinium at a concentration of 0.01%. SK-VII, which started during the writing of this thesis, has increased the gadolinium concentration to 0.03% for continued operation [82].

The oscillation analysis presented within this thesis focuses on the SK-IV period of running and the data taking within it. This follows from the recent SK analysis presented in [83]. Therefore, the information presented within this section focuses on that period.

Period	Start Date	End Date	Live-time (days)
I	April 1996	July 2001	1489.19
II	October 2002	October 2005	798.59
III	July 2006	September 2008	518.08
IV	September 2008	May 2018	3244.4
V	January 2019	July 2020	461.02
VI	July 2020	May 2022	583.3
VII	May 2022	Ongoing	N/A

Table 3.1: The various SK periods and respective live-time. The SK-VI live-time is calculated until 1st April 2022. SK-VII started during the writing of this thesis.

3.1.1 The SK Detector

The basic structure of the Super-Kamiokande (SK) detector is a cylindrical tank with a diameter 39.3m and height 41.1m filled with ultrapure water [81]. A diagram of the significant components of the SK detector is given in Figure 3.1.

The SK detector is situated in the Kamioka mine in Gifu, Japan. The mine is underground with roughly 1km rock overburden (2.7km water equivalent overburden) [84]. At this depth, the rate of cosmic ray muons is significantly decreased to a value of $\sim 2\text{Hz}$. The top of the tank is covered with stainless steel which is designed as a working platform for maintenance, calibration, and location for high voltage and data acquisition electronics.

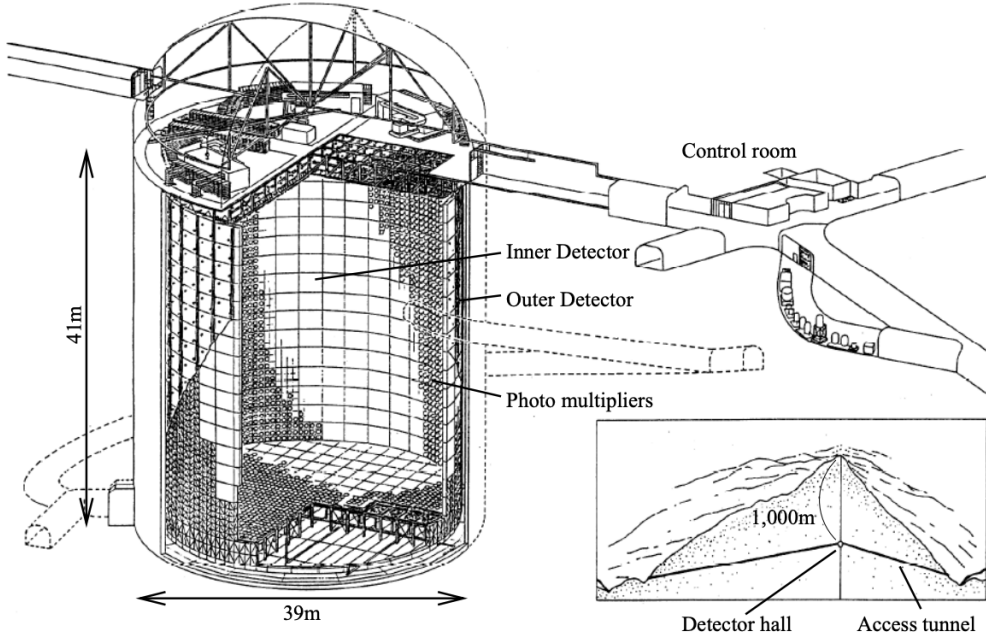


Figure 3.1: A schematic diagram of the Super-Kamiokande Detector. Taken from [85].

A smaller cylindrical structure (36.2m diameter, 33.8m height) is situated inside the tank, with an approximate 2m gap between this structure and the outer tank wall. The purpose of this structure is to support the photomultiplier tubes (PMTs). The volume inside and outside the support structure is referred to as the inner detector (ID) and outer detector (OD), respectively. In the SK-IV era, the ID and OD are instrumented by 11,129 50cm and 1,885 20cm PMTs respectively [81]. The ID contains a 32kton mass of water. Many analyses performed at SK use a “fiducial volume” defined by the volume of water inside the ID excluding some distance to the ID wall. This reduces the volume of the detector which is sensitive to neutrino events but reduces radioactive backgrounds and allows for

better reconstruction performance. The nominal fiducial volume is defined as the area contained inside 2m from the ID wall for a total of 22.5kton water [86].

The two regions of the detector (ID and OD) are optically separated with opaque black plastic hung from the support structure. The purpose of this is to determine whether an event entered or exited the ID. This allows cosmic ray muons and partially contained events to be tagged and separated from neutrino events entirely contained within the ID. This black plastic is also used to cover the area between the ID PMTs to reduce photon reflection from the ID walls. Opposite to this, the OD is lined with a reflective material to allow photons to reflect around inside the OD until collected by one of the PMTs. Furthermore, each OD PMT is optically coupled with $50 \times 50\text{cm}$ plates of wavelength shifting acrylic which increases the efficiency of light collection [84].

In the SK-IV data-taking period, the photocathode coverage of the detector, or the fraction of the ID wall instrumented with PMTs, is $\sim 40\%$ [84]. The PMTs have a quantum efficiency (the ratio of detected electrons to incident photons) of $\sim 21\%$ for photons with wavelengths of $360\text{nm} < \lambda < 390\text{nm}$ [87]. The proportion of photoelectrons that produce a signal in the dynode of a PMT, termed the collection efficiency, is $> 70\%$ [84]. The PMTs used within SK are most sensitive to photons with wavelength $300\text{nm} \leq \lambda \leq 600\text{nm}$ [84]. One disadvantage of using PMTs as the detection media is that the Earth's geomagnetic field can modify its response. Therefore, a set of compensation coils is built around the inner surface of the detector to mitigate this effect [88].

As mentioned, the SK detector is filled with ultrapure water, which in a perfect world would contain no impurities. However, bacteria and organic compounds can significantly degrade the water quality. This decreases the attenuation length, which reduces the total number of photons that hit a PMT. To combat this, a sophisticated water treatment system has been developed [84, 89]. UV lights, mechanical filters, and membrane degasifiers are used to reduce the bacteria, suspended particulates, and radioactive materials from the water. The flow of water within the tank is also critical as it can remove stagnant bacterial growth

or build-up of dust on the surfaces within the tank. Gravity drifts impurities in the water towards the bottom of the tank which, if left uncontrolled, can create asymmetric water conditions between the top and bottom of the tank. Typically, the water entering the tank is cooled below the ambient temperature of the tank to control convection and inhibit bacteria growth. Furthermore, the rate of dark noise hits within PMTs is sensitive to the PMT temperature [90] so controlling the temperature gradients within the tank is beneficial for stable measurements.

SK-VI is the first phase of the SK experiment to use gadolinium dopants within the ultrapure water [82]. As such, the SK water system had to be replaced to avoid removing the gadolinium concentrate from the ultrapure water [91]. For an inverse β -decay (IBD) interaction in a water target, the emitted neutron is thermally captured on hydrogen. This process releases 2.2MeV γ ray which are difficult to detect as the resulting Compton scattered electrons are very close to the Cherenkov threshold, limiting detection capability. Thermal capture of neutrons on gadolinium generates γ rays with higher energy (8MeV [69]) meaning they are more easily detected and reconstructed. SK-VI has 0.01% Gd loading (0.02% gadolinium sulphate by mass) which causes $\approx 50\%$ of neutrons emitted by IBD to be captured on gadolinium[92, 93]. Whilst predominantly useful for low energy analyses, Gd loading allows better $\nu/\bar{\nu}$ separation for atmospheric neutrino event selections [94]. Efforts are currently in place to increase the gadolinium concentrate to 0.03% for $\approx 75\%$ neutron capture efficiency on gadolinium [95]. The final stage of loading targets 0.1% concentrate targeting $\approx 90\%$ neutron capture efficiency on gadolinium.

3.1.2 Calibration

The calibration of the SK detector is documented in [81] and summarised below. The analysis presented within this thesis is dependent upon ‘high energy events’ (Charged particles with $O(> 100)$ MeV momenta). These are events that are expected to generate a larger number of photons such that each PMT will be hit with multiple photons. The reconstruction of these events depends

upon the charge deposited within each PMT and the timing response of each individual PMT. Therefore, the most relevant calibration techniques to this thesis are outlined.

Before installation, 420 PMTs were calibrated to have identical charge responses and then distributed throughout the tank in a cross-shape pattern (As illustrated by Figure 3.2). These are used as a standardised measure for the rest of the PMTs installed at similar geometric positions within SK to be calibrated against. To perform this calibration, a xenon lamp is located at the centre of the SK tank which flashes uniform light at 1Hz. This allows for geometrical effects, water quality variation, and timing effects to be measured in-situ throughout normal data-taking periods.

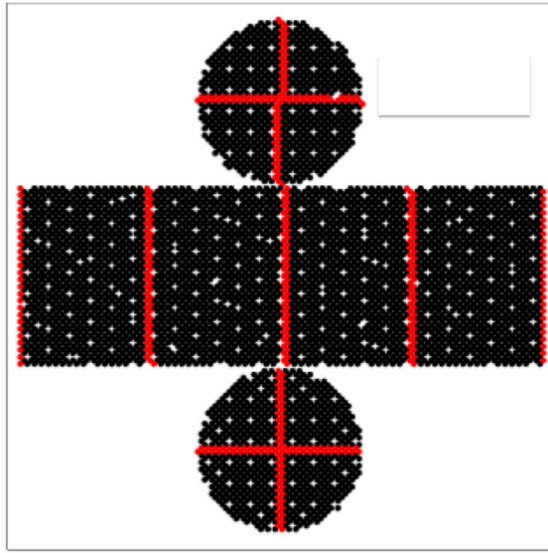


Figure 3.2: The location of “standard PMTs” (red) inside the SK detector. Taken from [81].

When specifically performing calibration of the detector (in out-of-data taking mode), the water in the tank was circulated to avoid top/bottom asymmetric water quality. Any non-uniformity within the tank significantly affects the PMT hit probability through scattering or absorption. This becomes a dominant effect for the very low-intensity light sources discussed later which are designed such that only one photon is incident upon a given PMT.

The “gain” of a PMT is defined as the ratio of the total charge of the signal produced compared to the charge of photoelectrons emitted by the photocathodes within the PMT. To calibrate the signal of each PMT, the “relative” and “absolute” gain values are measured. The relative gain is the variation of gain among each of the PMTs whereas the absolute gain is the average gain of all PMTs.

The relative gain is calibrated as follows. A laser is used to generate two measurements: a high-intensity flash that illuminates every PMT with a sufficient number of photons, and a low-intensity flash in which only a small number of PMTs collect light. The first measurement creates an average charge, $Q_{obs}(i)$ on PMT i , whereas the second measurement ensures that each hit PMT only generates a single photoelectron. For the low-intensity measurement, the number of times each PMT records a charge larger than 1/4 photoelectrons, $N_{obs}(i)$, is counted. The values measured can be expressed as

$$\begin{aligned} Q_{obs}(i) &\propto I_H \times f(i) \times \epsilon(i) \times G(i), \\ N_{obs}(i) &\propto I_L \times f(i) \times \epsilon(i), \end{aligned} \tag{3.1}$$

Where I_H and I_L is the intensity of the high and low flashes, $f(i)$ is the acceptance efficiency of the i^{th} PMT, $\epsilon(i)$ is the product of the quantum and collection efficiency of the i^{th} PMT and $G(i)$ is the gain of the i^{th} PMT. The relative gain for each PMT can determined by taking the ratio of these quantities.

The absolute gain calibration is performed by observing fixed energy γ -rays of $E_\gamma \sim 9\text{MeV}$ emitted isotropically from neutron capture on a NiCf source situated at the centre of the detector. This generates a photon yield of about 0.004 photoelectrons/PMT/event, meaning that $> 99\%$ of PMT signals are generated from single photoelectrons. A charge distribution is generated by performing this calibration over all PMTs, and the average value of this distribution is taken to be the absolute gain value.

As mentioned in subsection 3.1.1, the average quantum and collection efficiency for the SK detector PMTs is $\sim 21\%$ and $> 70\%$ respectively. However, these values do differ between each PMT and need to be calibrated accordingly.

Consequently, the NiCf source is also used to calibrate the “quantum \times collection” efficiency (denoted “QE”) value of each PMT. The NiCf low-intensity source is used as the PMT hit probability is proportional to the QE ($N_{obs}(i) \propto \epsilon(i)$ in Equation 3.1). A Monte Carlo prediction which includes photon absorption, scattering, and reflection is made to estimate the number of photons incident on each PMT and the ratio of the number of predicted to observed hits is calculated. The difference is attributed to the QE efficiency of that PMT. This technique is extended to calculate the relative QE efficiency by normalizing the average of all PMTs which removes the dependence on the light intensity.

Due to differing cable lengths and readout electronics, the timing response between a photon hitting the PMT and the signal being captured by the data acquisition can be different between each PMT. Due to threshold triggers (Described in subsection 3.1.3), the time at which a pulse reaches a threshold is dependent upon the size of the pulse. This is known as the ‘time-walk’ effect and also needs to be accounted for in each PMT. To calibrate the timing response, a pulse of light with width 0.2ns is emitted into the detector through a diffuser. Two-dimensional distributions of time and pulse height (or charge) are made for each PMT and are used to calibrate the timing response. This is performed in-situ during data taking with the light source pulsing at 0.03Hz.

The top/bottom water quality asymmetry is measured using the NiCf calibration data and cross-referencing these results to the “standard PMTs”. The water attenuation length is continuously measured by the rate of vertically-downgoing cosmic-ray muons which enter via the top of the tank.

Dark noise is where a PMT registers a pulse that is consistent with a single photoelectron emitted from photon detection despite the PMT being in complete darkness. This is predominately caused by two processes. Firstly there is intrinsic dark noise which is where photoelectrons gain enough thermal energy to be emitted from the photocathode, and secondly, the radioactive decay of contaminants inside the structure of the PMT. Typical dark noise rate for PMTs used within SK are $O(3)$ kHz [84]. This is lower than the expected number of

photons generated for a ‘high energy event’ (As described in subsection 3.1.4) but instability in this value can cause biases in reconstruction. Dark noise is related to the gain of a PMT and is calibrated using hits inside a time window recorded before an event trigger [96].

3.1.3 Data Acquisition and Triggering

As the analysis presented in this thesis only uses the SK-IV period of the SK experiment so this subsection focuses on the relevant points of the data acquisition and triggering systems to that SK period. The earlier data acquisition and triggering systems are documented in [97, 98].

Before the SK-IV period started, the existing front-end electronics were replaced with “QTC-Based Electrons with Ethernet, QBEE” systems [99]. When the QBEE observes a signal above a 1/4 photoelectron threshold, the charge-to-time (QTC) converter generates a rectangular pulse. The start of the rectangular pulse indicates the time at which the analog photoelectron signal was received and the width of the pulse indicates the total charge integrated throughout the signal. This is then digitized by time-to-digital converters and sent to the “front-end” PCs. The digitized signal from every QBEE is then chronologically ordered and sent to the “merger” PCs. It is the merger PCs that apply the software trigger. Any triggered events are passed to the “organizer” PC. This sorts the data stream of multiple merger PCs into chronologically ordered events which are then saved to disk. The schematic of data flow from PMTs to disk is illustrated in Figure 3.3.

The software trigger (described in [101]) operates by determining the number of PMT hits within a 200ns sliding window, N_{200} . This window coincides with the maximum time that a Cherenkov photon would take to traverse the length of the SK tank [98]. For lower energy events that generate fewer photons, this technique is useful for eliminating background processes like dark noise and radioactive decay which would be expected to separate in time. When the value of N_{200} exceeds some threshold, a software trigger is issued. There are several trigger thresholds used within the SK-IV period which are detailed in Table 3.2.

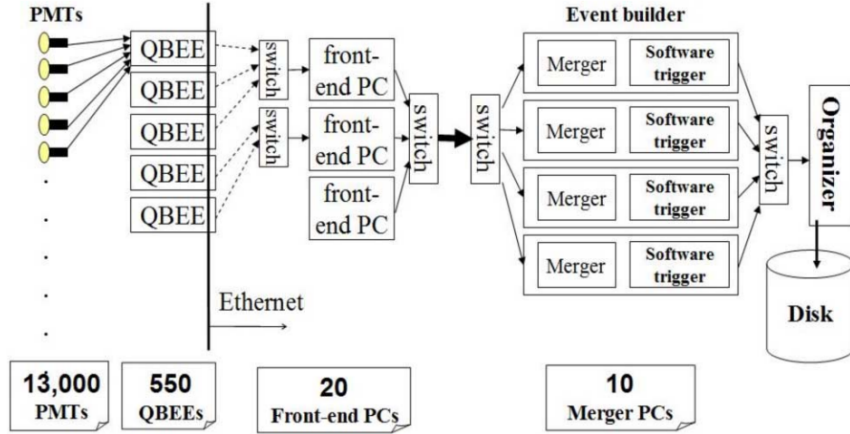


Figure 3.3: Schematic view of the data flow through the data acquisition and online system. Taken from [100].

If one of these thresholds is met, the PMT hits within an extended time window are also read out and saved to disk. In the special case of an event that exceeds the SHE trigger but does not exceed the OD trigger, the AFT trigger looks for delayed coincidences of 2.2MeV gamma rays emitted from neutron capture in a $535\mu\text{s}$ window after the SHE trigger. A similar but more complex “Wideband Intelligent Trigger (WIT)” has been deployed and is described in [102].

Trigger	Acronym	Condition	Extended time window (μs)
Super Low Energy	SLE	>34/31 hits	1.3
Low Energy	LE	>47 hits	40
High Energy	HE	>50 hits	40
Super High Energy	SHE	>70/58 hits	40
Outer Detector	OD	>22 hits in OD	N/A

Table 3.2: The trigger thresholds and extended time windows saved around an event which were utilised throughout the SK-IV period. The exact thresholds can change and the values listed here represent the thresholds at the start and end of the SK-IV period.

3.1.4 Cherenkov Radiation

Cherenkov light is emitted from any highly energetic charged particle traveling with relativistic velocity, β , greater than the local speed of light in a medium [103].

Cherenkov light is formed at the surface of a cone with characteristic pitch angle,

$$\cos(\theta) = \frac{1}{\beta n}. \quad (3.2)$$

where n is the refractive index of the medium. Consequently, the Cherenkov momentum threshold, P_{thres} , is dependent upon the mass, m , of the charged particle moving through the medium,

$$P_{thres} = \frac{m}{\sqrt{n^2 - 1}} \quad (3.3)$$

For water, where $n = 1.33$, the Cherenkov threshold momentum and energy for various particles are given in Table 3.3. In contrast, γ -rays are detected indirectly via the combination of photons generated by Compton scattering and pair production. The threshold for detection in the SK detector is typically higher than the threshold for photon production. This is due to the fact that the attenuation of photons in the water means that typically $\sim 75\%$ of Cherenkov photons reach the ID PMTs. Then the collection and quantum efficiencies described in subsection 3.1.1 result in the number of detected photons being lower than the number of photons which reach the PMTs.

Particle	Threshold Momentum (MeV)	Threshold Energy (MeV)
Electron	0.5828	0.7751
Muon	120.5	160.3
Pion	159.2	211.7
Proton	1070.0	1423.1

Table 3.3: The threshold momentum and energy for a particle to generate Cherenkov light in ultrapure water, as calculated in Equation 3.2 in ultrapure water which has refractive index $n = 1.33$.

The Frank-Tamm equation [104] describes the relationship between the number of Cherenkov photons generated per unit length, dN/dx , the wavelength of the photons generated, λ , and the relativistic velocity of the charged particle,

$$\frac{d^2N}{dxd\lambda} = 2\pi\alpha \left(1 - \frac{1}{n^2\beta^2}\right) \frac{1}{\lambda^2}. \quad (3.4)$$

where α is the fine structure constant. For a 100MeV momentum electron, approximately 330 photons will be produced per centimeter in the $300\text{nm} \leq \lambda \leq 700\text{nm}$ region which the ID PMTs are most sensitive to [84].

3.2 The Tokai to Kamioka Experiment

The Tokai to Kamioka (T2K) experiment is a long-baseline neutrino oscillation experiment located in Japan. Proposed in the early 2000s [105, 106] to replace K2K [107], T2K was designed to observe electron neutrino appearance whilst precisely measuring the oscillation parameters associated with muon neutrino disappearance [108]. The experiment consists of a neutrino beam generated at the Japan Proton Accelerator Research Complex (J-PARC), a suite of near detectors situated 280m from the beam target, and the Super Kamiokande far detector positioned at a 295km baseline. The cross-section view of the T2K experiment is drawn in Figure 3.4.

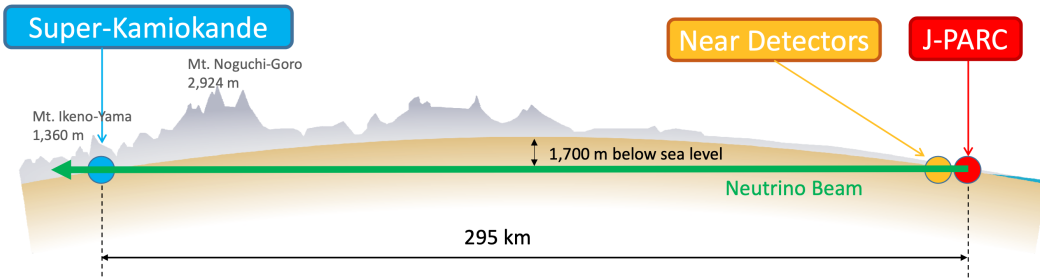


Figure 3.4: The cross-section view of the Tokai to Kamioka experiment illustrating the beam generation facility at J-PARC, the near detector situated at a baseline of 280m and the Super Kamiokande far detector situated 295km from the beam target.

The T2K collaboration makes world-leading measurements of the $\sin^2(\theta_{23})$, Δm_{23}^2 , and δ_{CP} oscillation parameters. Improvements in the precision and accuracy of parameter estimates are still being made by including new data samples and developing the models which describe the neutrino interactions and detector responses [109]. Electron neutrino appearance was first observed at T2K in 2014 [110] with 7.3σ significance.

The near detectors provide constraints on the beam flux and cross-section model parameters used within the oscillation analysis by observing the unoscillated neutrino beam. There are a host of detectors situated in the near detector hall (As illustrated in Figure 3.5): ND280 (subsection 3.2.2), INGRID (subsection 3.2.3), NINJA [111], WAGASCI [112], and Baby-MIND [113]. The latter three are not currently used within the oscillation analysis presented within this thesis.

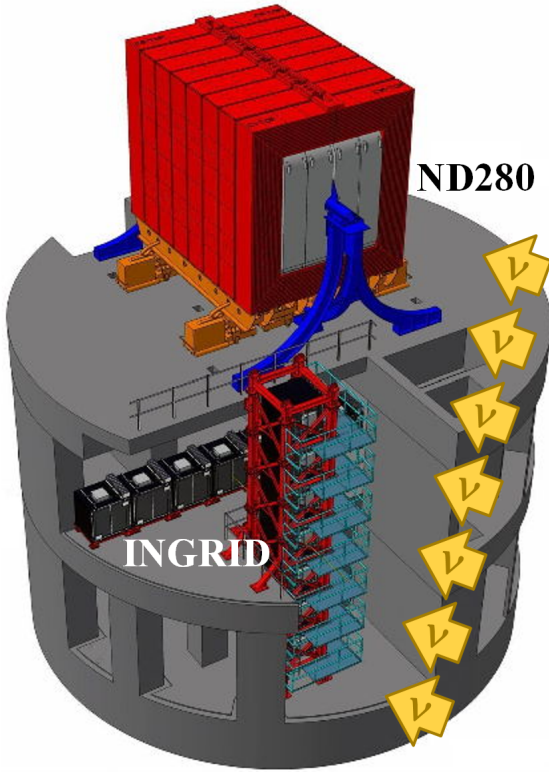


Figure 3.5: The near detector suite for the T2K experiment showing the ND280 and INGRID detectors. The distance between the detectors and the beam target is 280m.

Whilst this thesis presents the ND280 in terms of its purpose for the oscillation analysis, the detector can also make many cross-section measurements at neutrino energies of $O(1)$ GeV for the different targets within the detector [114, 115]. These measurements are of equal importance as they can lead the way in determining the model parameters used in the interaction models for the future high-precision era of neutrino physics.

There are two independent fitters, MaCh3 and BANFF, which perform the near detector fit. MaCh3 is the basis of this analysis and uses a bayesian Markov

Chain Monte Carlo fitting technique, whereas BANFF uses a frequentist gradient descent technique. The output of each fitter is converted into a covariance matrix to describe the error and correlations between all the flux and cross-section parameters. This is then propagated to the far-detector oscillation analysis group for use in the P-Theta and VALOR fitting framework. As MaCh3 can handle both near and far detector samples, it does not use this covariance matrix and instead opts for a simultaneous fit of the two detector measurements. This is an analysis choice which removes the assumption of Gaussian posterior distributions required when building the post-fit covariance matrix.

DB: MaCh3 vs PTheta and Valor

There are three particular tunes of the T2K flux and low energy cross section model typically considered. Firstly, the “generated” tune which is the set of dial values with which the Monte Carlo was generated. Secondly, the set of dial values which are taken from external data measurements and used as inputs. These are the “pre-fit” dial values. The reason these two sets of dial values are different is that the external data measurements are continually updated but it is very computationally intensive to regenerate a Monte Carlo prediction after each update. The final tune is the “post-fit”, “post-ND fit” or “post-BANFF” dial values. These are the values taken from the fit to the beam near detector data.

3.2.1 The Neutrino Beam

The neutrino beam used within the T2K experiment is described in [58, 116] and summarised below. The accelerating facility at J-PARC is composed of two sections; the primary and secondary beamlines. Figure 3.6 illustrates a schematic of the beamline, focusing mostly on the components of the secondary beamline. The primary beamline has three accelerators that progressively accelerate protons; a linear accelerator, a rapid-cycling synchrotron, and the main-ring (MR) synchrotron. Once fully accelerated by the MR, the protons have a kinetic energy of 30GeV. Eight bunches of these protons, separated by 500ns, are extracted per “spill” from the MR and directed towards a graphite target (a rod of length

91.4cm and diameter 2.6cm). Spills are extracted at 0.5Hz with $\sim 3 \times 10^{14}$ protons contained per spill.

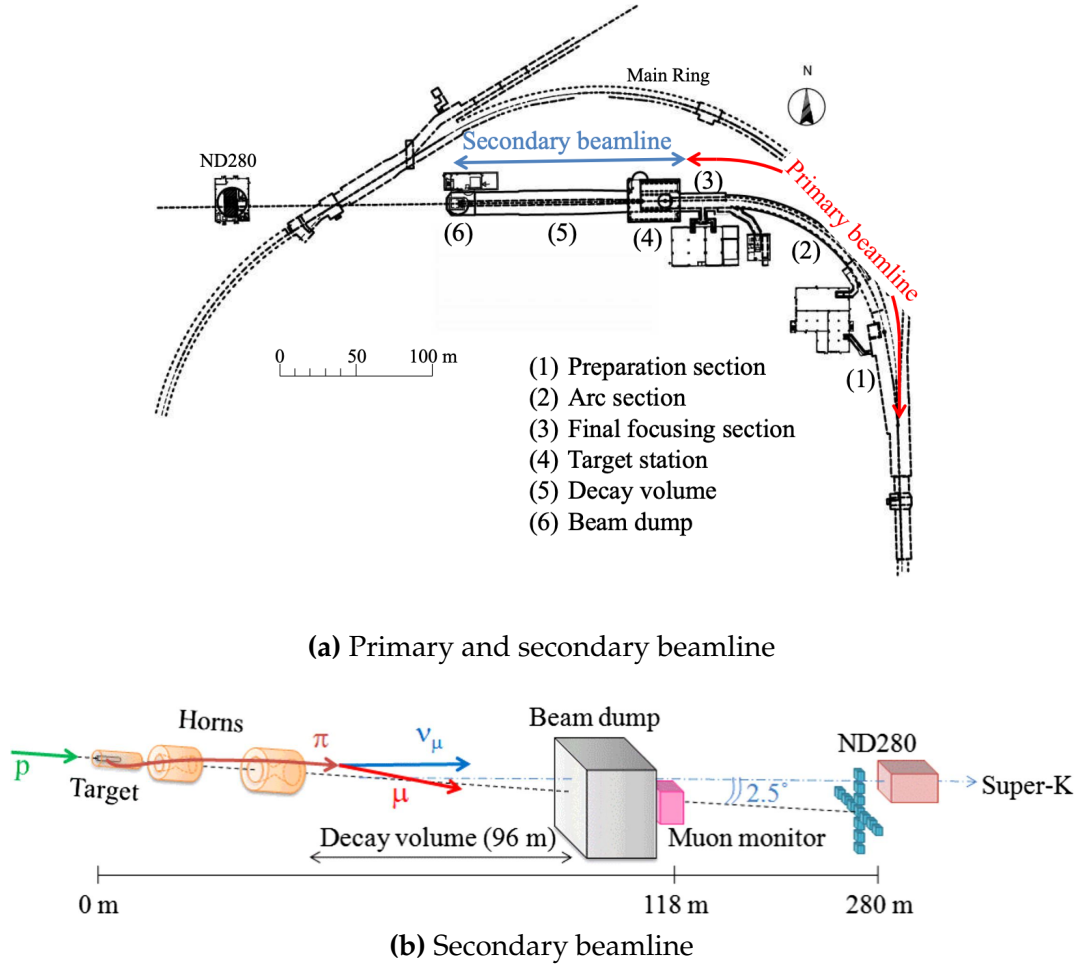


Figure 3.6: Top panel: Bird's eye view of the most relevant part of primary and secondary beamline used within the T2K experiment. The primary beamline is the main-ring proton synchrotron, kicker magnet, and graphite target. The secondary beamline consists of the three focusing horns, decay volume, and beam dump. Figure taken from [116]. Bottom panel: The side-view of the secondary beamline including the focusing horns, beam dump and neutrino detectors. Figure taken from [117].

The secondary beamline consists of three main components: the target station, the decay volume, and the beam dump. The target station is comprised of the target, beam monitors, and three magnetic focusing horns. The proton beam interacts with the graphite target to form a secondary beam of mostly pions and kaons. The secondary beam travels through a 96m long decay volume, generating neutrinos through the following decays [58],

$$\begin{array}{ll}
\pi^+ \rightarrow \mu^+ + \nu_\mu & \pi^- \rightarrow \mu^- + \bar{\nu}_\mu \\
K^+ \rightarrow \mu^+ + \nu_\mu & K^- \rightarrow \mu^- + \bar{\nu}_\mu \\
\rightarrow \pi^0 + e^+ + \nu_e & \rightarrow \pi^0 + e^- + \bar{\nu}_e \\
\rightarrow \pi^0 + \mu^+ + \nu_\mu & \rightarrow \pi^0 + \mu^- + \bar{\nu}_\mu \\
K_L^0 \rightarrow \pi^- + e^+ + \nu_e & K_L^0 \rightarrow \pi^+ + e^- + \bar{\nu}_e \\
\rightarrow \pi^- + \mu^+ + \nu_\mu & \rightarrow \pi^+ + \mu^- + \bar{\nu}_\mu \\
\mu^+ \rightarrow e^+ + \bar{\nu}_\mu + \nu_e & \mu^- \rightarrow e^- + \nu_\mu + \bar{\nu}_e
\end{array}$$

The electrically charged component of the secondary beam is focused towards the far detector by the three magnetic horns. These horns direct charged particles of a particular polarity towards SK whilst defocusing the oppositely charged particles. This allows a mostly neutrino or mostly antineutrino beam to be used within the experiment, denoted as “forward horn current (FHC)” or “reverse horn current (RHC)” respectively.

Figure 3.7 illustrates the different contributions to the FHC and RHC neutrino flux. The low energy flux is dominated by the decay of pions whereas kaon decay becomes the dominant source of neutrinos for $E_\nu > 3\text{GeV}$. The “wrong-sign” component, which is the $\bar{\nu}_\mu$ background in a ν_μ beam, and the intrinsic irreducible ν_e background, are predominantly due to muon decay for $E_\nu < 2\text{GeV}$. As the antineutrino production cross-section is smaller than the neutrino cross-section, the wrong-sign component is more dominant in the RHC beam as compared to that in the FHC beam.

The beam dump, situated at the end of the decay volume, stops all charged particles other than highly energetic muons ($p_\mu > 5\text{GeV}$). The MuMon detector monitors the penetrating muons to determine the beam direction and intensity which is used to constrain some of the beam flux systematics within the analysis [117, 119].

The T2K experiment uses an off-axis beam to narrow the neutrino energy distribution. This was the first implementation of this technique in a long-baseline neutrino oscillation experiment after its original proposal [120]. Pion decay, $\pi \rightarrow \mu + \nu_\mu$, is a two-body decay. Consequently, the neutrino energy,

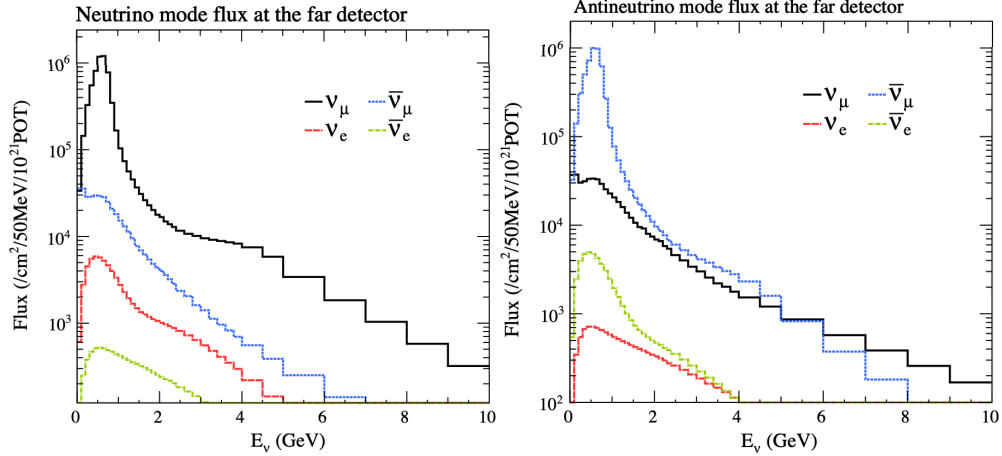


Figure 3.7: The Monte Carlo prediction of the energy spectrum for each flavour of neutrino (ν_e , $\bar{\nu}_e$, ν_μ and $\bar{\nu}_\mu$) in the neutrino dominated beam FHC mode (Left) and antineutrino dominated beam RHC mode (Right) expected at SK. Taken from [118].

E_ν , can be determined based on the pion energy, E_π , and the angle at which the neutrino is emitted, θ ,

$$E_\nu = \frac{m_\pi^2 - m_\mu^2}{2(E_\pi - p_\pi \cos(\theta))}, \quad (3.5)$$

where m_π and m_μ are the mass of the pion and muon respectively. For a fixed energy pion, the neutrino energy distribution is dependent upon the angle at which the neutrinos are observed from the initial pion beam direction. For the 295km baseline at T2K, $E_\nu = 0.6\text{GeV}$ maximises the electron neutrino appearance probability, $P(\nu_\mu \rightarrow \nu_e)$, whilst minimising the muon disappearance probability, $P(\nu_\mu \rightarrow \nu_\mu)$. Figure 3.8 illustrates the neutrino energy distribution for a range of off-axis angles, as well as the oscillation probabilities most relevant to T2K.

3.2.2 The Near Detector at 280m

Whilst all the near detectors are situated in the same “pit” located at 280m from the beamline, the “ND280” detector is the off-axis detector which is situated at the same off-axis angle as the Super-Kamiokande far detector. It has two primary functions; firstly it measures the neutrino flux and secondly it counts the event rates of different types of neutrino interactions. Both of these constrain the flux



Figure 3.8: Top panel: T2K muon neutrino disappearance probability as a function of neutrino energy. Middle panel: T2K electron neutrino appearance probability as a function of neutrino energy. Bottom panel: The neutrino flux distribution for three different off-axis angles (Arbitrary units) as a function of neutrino energy.

and cross-section systematics invoked within the model for a more accurate prediction of the expected event rate at the far detector.

As illustrated in Figure 3.9, the ND280 detector consists of several sub-detectors. The most important part of the detector for this analysis is the tracker region. This is comprised of two time projection chambers (TPCs) sandwiched between three fine grain detectors (FGDs). The FGDs contain both hydrocarbon plastics and water targets for neutrino interactions and provide track reconstruction near the interaction vertex. The emitted charged particles can then propagate into the TPCs which provide particle identification and momentum reconstruction. The FGDs and TPCs are further described in subsubsection 3.2.2.1 and subsubsection 3.2.2.2 respectively. The electromagnetic calorimeter (ECAL)

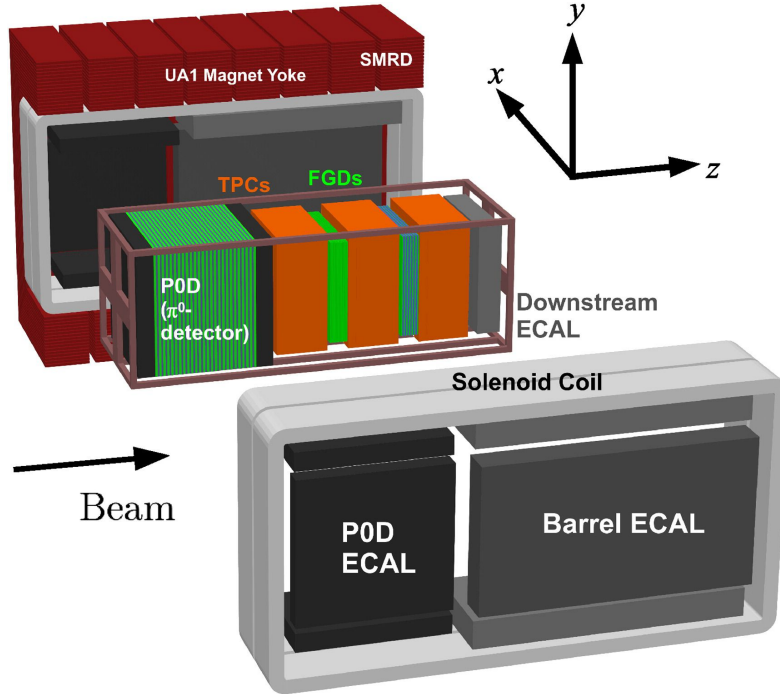


Figure 3.9: The components of the ND280 detector. The neutrino beam travels from left to right. Taken from [116].

encapsulates the tracker region alongside the π^0 detector (P0D). The ECAL measures the deposited energy from photons emitted from interactions within the FGD. The P0D constrains the cross-section of neutral current interactions which generate neutral pions, which is one of the largest backgrounds in the electron neutrino appearance oscillation channel. The P0D and ECAL detectors are detailed in subsubsection 3.2.2.3 and subsubsection 3.2.2.4 respectively. The entire detector is located within a large yoke magnet which produces a 0.2T magnetic field. This design of the magnet also includes a scintillating detector called the side muon range detector (SMRD) which is used to track high-angle muons as well as acting as a cosmic veto. The SMRD is described in subsubsection 3.2.2.5.

3.2.2.1 Fine Grained Detectors

The T2K tracker region is comprised of two fine grained detectors (FGD) and three Time Projection Chambers (TPC). A detailed description of the FGD design, construction, and assembly is found in [121] and summarised below. The FGDs are the primary target for neutrino interactions with a mass of 1.1 tonnes per FGD.

Alongside this, the FGDs are designed to be able to track short-range particles which do not exit the FGD. Typically, short-range particles are low momentum and are observed as tracks that deposit a large amount of energy per unit length. This means the FGD needs good granularity to resolve these particles. The FGDs have the best timing resolution ($\sim 3\text{ns}$) of any of the sub-detectors of the ND280 detector. As such, the FGDs are used for time of flight measurements to distinguish forward going positively charged particles from backward going negatively charged particles. Finally, any tracks which pass through multiple sub-detectors are required to be track matched to the FGD.

Both FGDs are made from square scintillator planes of side length 186cm and width 2.02cm. Each plane consists of two layers of 192 scintillator bars in an X or Y orientation. A wavelength shifting fiber is threaded through the centre of each bar and is read out by a multi-pixel photon counter (MPPC). FGD1 is the most upstream of the two FGDs and contains 15 planes of carbon plastic scintillator which is a common target in external neutrino scattering data. As the far detector is a pure water target, 7 of the 15 scintillator planes in FGD2 have been replaced with a hybrid water-scintillator target. Due to the complexity of the nucleus, nuclear effects can not be extrapolated between different nuclei. Therefore having the ability to take data on one target which is the same as external data and another target which is the same as the far detector target is beneficial for reliable model parameter estimates.

The integrated deposited energy is used for particle identification. The FGD can distinguish protons from other charged particles by comparing the integrated deposited energy from data to Monte Carlo prediction as seen in Figure 3.10.

3.2.2.2 Time Projection Chambers

The majority of particle identification and momentum measurements within ND280 are provided by three Time Projection Chambers (TPCs) [122]. The TPCs are located on either side of the FGDs. They are located inside of the

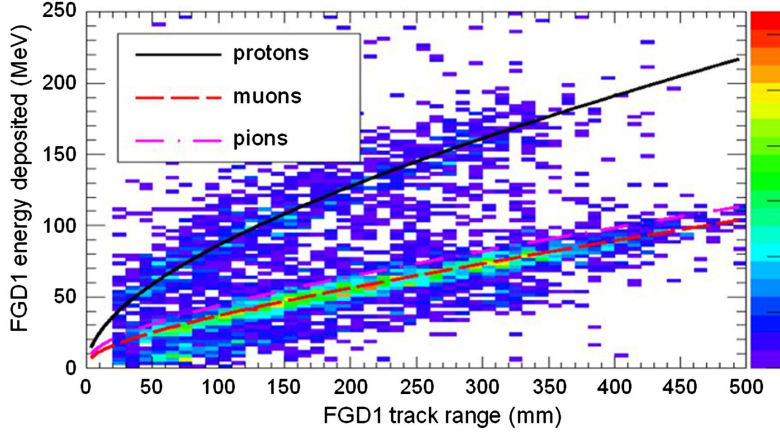


Figure 3.10: Comparison of data to Monte Carlo prediction of integrated deposited energy as a function of track length for particles that stopped in FGD1. Taken from [121].

magnetic field meaning the momentum of a charged particle can be determined from the bending of the track.

Each TPC module consists of two gas-tight boxes, as shown in Figure 3.11, which are made of non-magnetic material. The outer box is filled with CO₂ which acts as an electrical insulator between the inner box and the ground. The inner box forms the field cage which produces a uniform electric drift field of $\sim 275\text{V/cm}$ and is filled with an argon gas mixture. Charged particles moving through this gas mixture ionize the gas and the ionised charge is drifted towards micromegas detectors which measure the ionization charge. The time and position information in the readout allows a three-dimensional image of the neutrino interaction.

The particle identification of tracks that pass through the TPCs is performed using dE/dx measurements. Figure 3.12 illustrates the data to Monte Carlo distributions of the energy lost by a charged particle passing through the TPC as a function of the reconstructed particle momentum. The resolution is $7.8 \pm 0.2\%$ meaning that electrons and muons can be distinguished. This allows reliable measurements of the intrinsic ν_e component of the beam.

3.2.2.3 π^0 Detector

If one of the γ -rays from a $\pi^0 \rightarrow 2\gamma$ decay is missed at the far detector, the reconstruction will determine that event to be a charge current ν_e -like event.

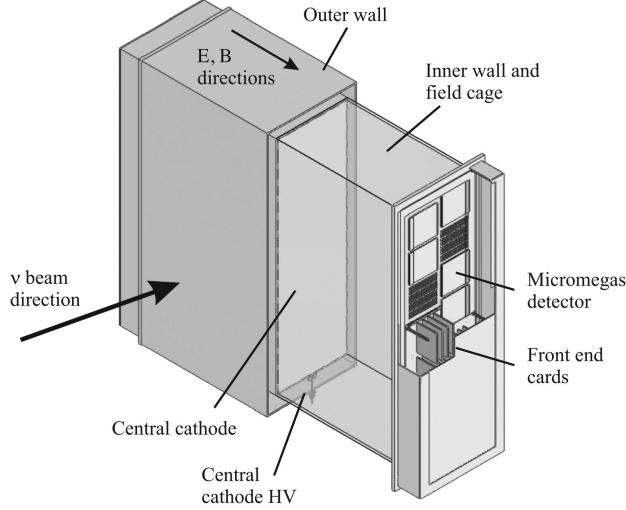


Figure 3.11: Schematic design of a Time Projection Chamber detector. Taken from [122].

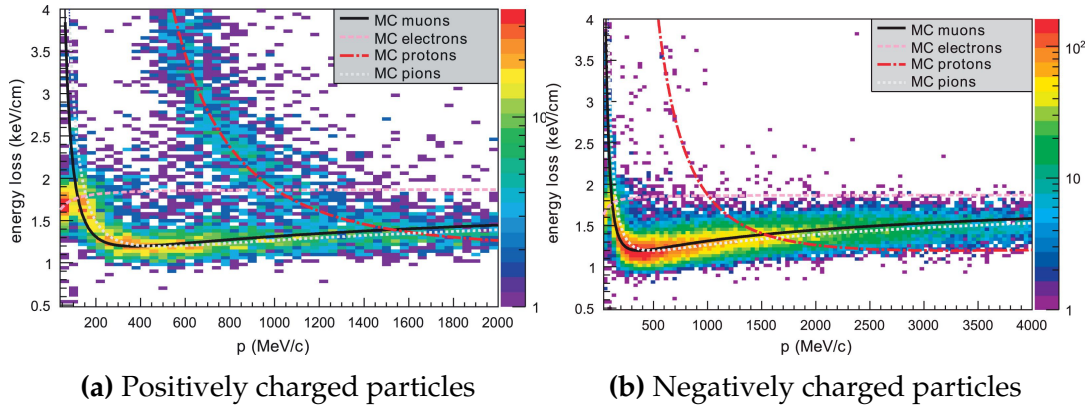


Figure 3.12: The distribution of energy loss as a function of reconstructed momentum for charged particles passing through the TPC, comparing data to Monte Carlo prediction. Taken from [122].

This is one of the main backgrounds hindering the electron neutrino appearance searches. The π^0 detector (P0D) measures the cross-section of the neutral current induced neutral pion production on a water target to constrain this background.

The P0D is a cube of approximately 2.5m length consisting of layers of scintillating bars, brass and lead sheets, and water bags as illustrated in Figure 3.13. Two electromagnetic calorimeters are positioned at the most upstream and most downstream position in the sub-detector and the water target is situated in between them. The scintillator layers are built from two triangular bars orientated in opposite directions to form a rectangular layer. Each triangular scintillator bar is threaded with optical fiber which is read out by MPPCs. The high-Z brass and

lead regions produce electron showers from the photons emitted in π^0 decay.

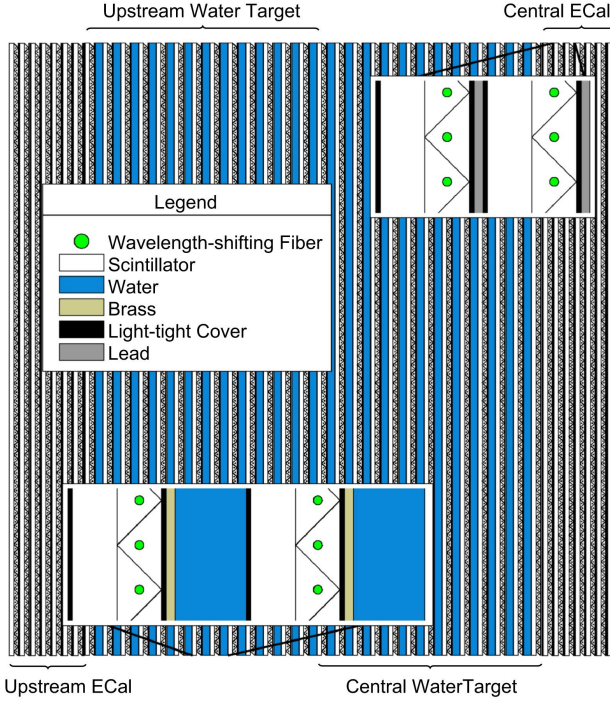


Figure 3.13: A schematic of the P0D side-view. Taken from [123].

The sub-detector can generate measurements of NC1 π^0 cross-sections on a water target by measuring the event rate both with and without the water target, with the cross-section on a water target being determined as the difference. The total active mass is 16.1 tonnes when filled with water and 13.3 tonnes when empty.

3.2.2.4 Electromagnetic Calorimeter

The electromagnetic calorimeter [124] (ECal) encapsulates the P0D and tracking sub-detectors. Its primary purpose is to aid π^0 reconstruction from any interaction in the tracker. To do this, it measures the energy and direction of photon showers from $\pi^0 \rightarrow 2\gamma$ decay. It can also distinguish pion and muon tracks depending on the shape of the photon shower deposited.

The ECal is comprised of three sections; the P0D ECal which surrounds the P0D, the barrel ECal which encompasses the tracking region, and the downstream ECal which is situated downstream of the tracker region. The barrel and downstream ECals are tracking calorimeters that focus on electromagnetic showers

from high-angle particles emitted from the tracking sub-detectors. Particularly in the TPC, high-angle tracks (those which travel perpendicularly to the beam-axis) can travel along a single scintillator bar resulting in very few hits. The width of the barrel and downstream ECal corresponds to ~ 11 electron radiation lengths to ensure a significant amount of the π^0 energy is contained. As the P0D has its own calorimetry which reconstructs showers, the P0D ECal determines the energy which escapes the P0D.

Each ECal is constructed of multiple layers of scintillating bars sandwiched between lead sheets. The scintillating bars are threaded with optical fiber and read out by MPPCs. Each sequential layer of the scintillator is orientated perpendicular to the previous which allows a three dimensional event reconstruction. The target mass of the P0D ECal, barrel ECal, and downstream ECal are 1.50, 4.80 and 6.62 tonnes respectively.

3.2.2.5 Side Muon Range Detector

As illustrated in Figure 3.9, the ECal, FGDs, P0D, and TPCs are enclosed within the UA1 magnet. Originally designed for the NOMAD [125] experiment and reconditioned for use in the T2K experiment [126], the UA1 magnet provides a uniform horizontal magnetic field of 0.2T with an uncertainty of 2×10^{-4} T.

Built into the UA1 magnet, the side muon range detector (SMRD)[127] monitors high-energy muons which leave the tracking region and permeate through the ECal. It additionally acts as a cosmic muon veto and trigger.

3.2.3 The Interactive Neutrino GRID

The Interactive Neutrino GRID (INGRID) detector is situated within the same “pit” as the other near detectors. It is aligned with the beam in the “on-axis” position and measures the beam direction, spread, and intensity. The detector was originally designed with 16 identical modules [116] (two modules have since been decommissioned) and a “proton” module. The design of the detector is cross-shaped with length and height 10m \times 10m as illustrated in Figure 3.14.

Each module is composed of iron sheets interlaced with eleven tracking scintillator planes for a total target mass of 7.1 tonnes per module. The scintillator design is an X-Y pattern of 24 bars in both orientations, where each bar contains wave-length shifting fibers which are connected to multi-pixel photon counters (MPPCs). Each module is encapsulated inside veto planes to aid the rejection of charged particles entering the module.

The proton module is different from the other modules in that it consists of entirely scintillator planes with no iron target. The scintillator bars are also smaller than those used in the other modules to increase the granularity of the detector and improve tracking capabilities. The module sits in the centre of the beamline and is designed to give precise measurements of quasi-elastic charged current interactions to evaluate the performance of the Monte Carlo simulation of the beamline.

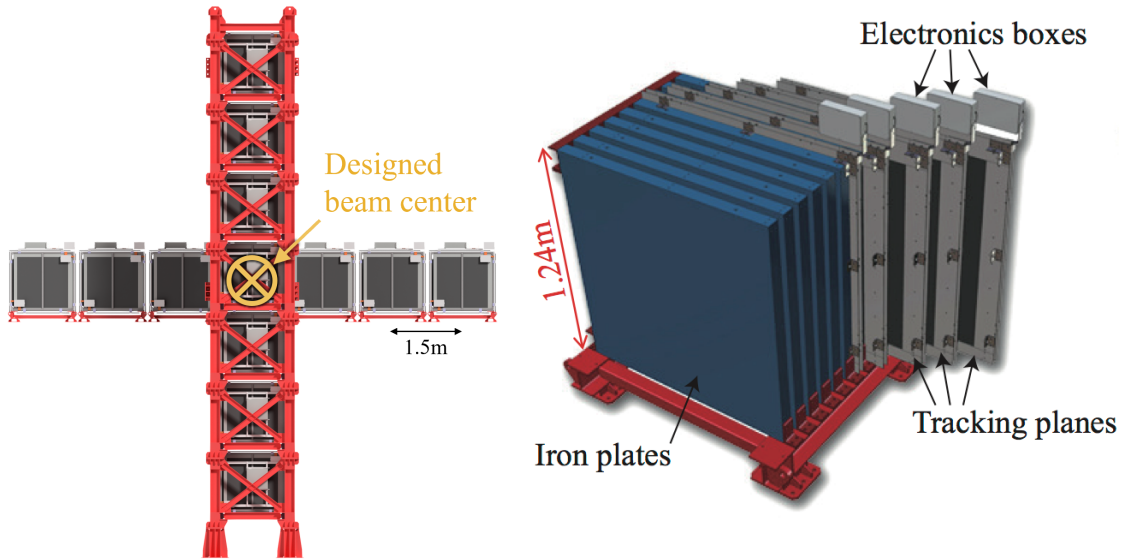


Figure 3.14: Left panel: The Interactive Neutrino GRID on-axis Detector. 14 modules are arranged in a cross-shape configuration, with the centre modules being directly aligned with the on-axis beam. Right panel: The layout of a single module of the INGRID detector. Both figures are recreated from [116].

The INGRID detector can measure the beam direction to an uncertainty of 0.4mrad and the beam centre within a resolution of 10cm [116]. The beam direction in both the vertical and horizontal directions is discussed in [128]

and it is found to be in good agreement with the MUMON monitor described in subsection 3.2.1.

4

Bayesian Statistics and Markov Chain Monte Carlo Techniques

This thesis presents a Bayesian oscillation analysis. To extract the oscillation parameters, a Markov Chain Monte Carlo (MCMC) method is used. This chapter explains the theory of how parameter estimates can be determined using this technique and condenses the material found in the literature [129–132].

The oscillation parameter determination presented within this thesis is built upon a simultaneous fit to neutrino beam data in the near detector, beam data at SK and atmospheric data at SK. In total, there are four oscillation parameters of interest ($\sin^2(\theta_{23})$, $\sin^2(\theta_{13})$, Δm_{23}^2 , and δ_{CP}), two oscillation parameters to which this study will not be sensitive ($\sin^2(\theta_{12})$, Δm_{12}^2) and many nuisance parameters that control the systematic uncertainty models invoked within this study.

The MCMC technique generates a multi-dimensional probability distribution across all of the model parameters used in the fit. To determine the parameter estimate of a single parameter, this multi-dimensional object is integrated over all other parameters. This process is called Marginalisation and is further described in subsection 4.3.1. Monte Carlo techniques approximate the probability distribution of each parameter within the limit of generating infinite samples. As ever, generating a large number of samples is time and resource-dependent. Therefore,

an MCMC technique is utilised within this analysis to reduce the required number of steps to sufficiently sample the parameter space. This technique is described in further detail in subsection 4.2.1.

4.1 Bayesian Statistics

Bayesian inference treats observable data, D , and model parameters, $\vec{\theta}$, on equal footing such that a probability model of both data and parameters is required. This is the joint probability distribution $P(D, \vec{\theta})$ and can be described by the prior distribution for model parameters $P(\vec{\theta})$ and the likelihood of the data given the model parameters $P(D|\vec{\theta})$,

$$P(D, \vec{\theta}) = P(D|\vec{\theta})P(\vec{\theta}). \quad (4.1)$$

The prior distribution, $P(\vec{\theta})$, describes all previous knowledge about the parameters within the model. For example, if the risk of developing health problems is known to increase with age, the prior distribution would describe the increase. For the purpose of this analysis, the prior distribution is typically the best-fit values taken from external data measurements with a Gaussian uncertainty. The prior distribution can also contain correlations between model parameters. In an analysis using Monte Carlo techniques, the likelihood of measuring some data assuming some set of model parameters is calculated by comparing the Monte Carlo prediction generated at that particular set of model parameters to the data.

It is parameter estimation that is important for this analysis and as such, we apply Bayes' theorem [133] to calculate the probability for each parameter to have a certain value given the observed data, $P(\vec{\theta}|D)$, which is known as the posterior distribution (often termed the posterior). This can be expressed as

$$P(\vec{\theta}|D) = \frac{P(D|\vec{\theta})P(\vec{\theta})}{\int P(D|\vec{\theta})P(\vec{\theta})d\vec{\theta}}. \quad (4.2)$$

The denominator in Equation 4.2 is the integral of the joint probability distribution over all values of all parameters used within the fit. For brevity, we say that the posterior distribution is

$$P(\vec{\theta}|D) \propto P(D|\vec{\theta})P(\vec{\theta}). \quad (4.3)$$

In subsection 4.3.1, we see that for the cases used within this analysis, it is reasonable to know the posterior to some normalisation constant.

4.1.1 Application of Prior Knowledge

The posterior distribution is proportional to the prior uncertainty applied on each parameter, as illustrated by Equation 4.3. This means that it is possible to change the prior after the posterior distribution has been determined. The prior uncertainty of a particular parameter can be ‘divided’ out of the posterior distribution and the resulting distribution can be reweighted using the new prior uncertainty that is to be applied. The methodology and implementation of changing the prior follows that described in [134].

An example implementation that is useful for this analysis is the application of the “reactor constraint”. As discussed in section 2.4, an external constraint on $\sin^2(\theta_{13})$ is determined from measurements taken from reactor experiments. However, the sensitivities from just using the T2K and SK samples is equally as important. Without this technique, two fits would have to be run, doubling the required resources. Therefore, the key benefit for this analysis is the fact that only a single ‘fit’ has to be performed and can be used to build the two posterior distributions of the with and without reactor constraint applied.

4.2 Monte Carlo Simulation

Monte Carlo techniques are used to numerically solve a complex problem that does not necessarily have an analytical solution. These techniques rely on

building a large ensemble of samples from an unknown distribution and then using the ensemble to approximate the properties of the distribution.

An example that uses Monte Carlo techniques is to calculate the area underneath a curve. For example, take the problem of calculating the area under a straight line with gradient $M = 0.4$ and intercept $C = 1.0$. Analytically, one can calculate the area under the line is equal to 30 units for $0 \leq x \leq 10$. Using Monte Carlo techniques, one can calculate the area under this line by throwing many random values for the x and y components of each sample and then calculating whether that point falls below the line. The area can then be calculated by the ratio of points below the line to the total number of samples thrown multiplied by the total area in which samples were scattered. The study is shown in Figure 4.1 highlights this technique and finds the area under the curve to be 29.9 compared to an analytical solution of 30.0. The deviation of the numerical to analytical solution can be attributed to the number of samples used in the study. The accuracy of the approximation in which the properties of the Monte Carlo samples replicate those of the desired distribution is dependent on the number of samples used. Replicating this study with a differing number of Monte Carlo samples used in each study (As shown in Figure 4.2) highlights how the Monte Carlo techniques are only accurate within the limit of a high number of samples.

Whilst the above example has an analytical solution, these techniques are just as applicable to complex solutions. Clearly, any numerical solution is only as useful as its efficiency. As discussed, the accuracy of the Monte Carlo technique is dependent upon the number of samples generated to approximate the properties of the distribution. Furthermore, if the positions at which the samples are evaluated are not ‘cleverly’ picked, the efficiency of the Monte Carlo technique significantly drops. Given the example in Figure 4.1, if the region in which the samples are scattered significantly extends passed the region of interest, many calculations will be calculated but do not add to the ability of the Monte Carlo technique to achieve the correct result. For instance, any sample evaluated at a $y \geq 5$ could be removed without affecting the final result. This does bring in

an aspect of the ‘chicken and egg’ problem in that to achieve efficient sampling, one needs to know the distribution beforehand.

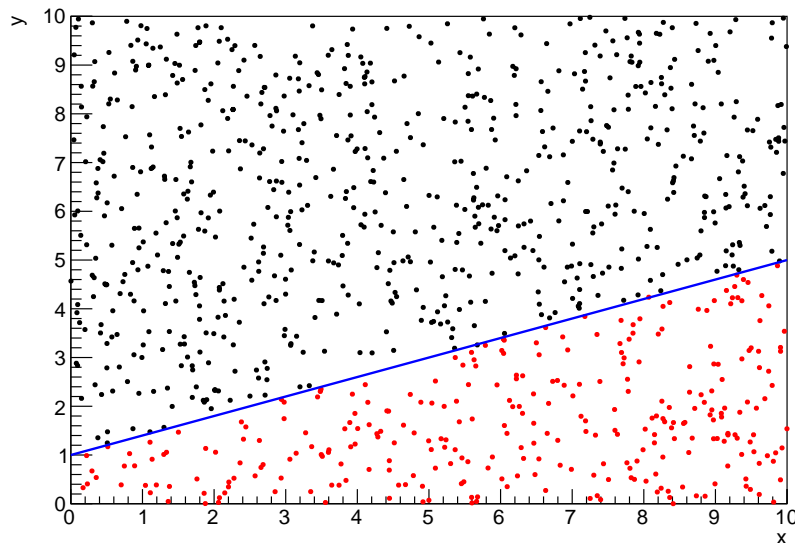


Figure 4.1: Example of using Monte Carlo techniques to find the area under the blue line. The gradient and intercept of the line are 0.4 and 1.0 respectively. The area found to be under the curve using one thousand samples is 29.9 units.

4.2.1 Markov Chain Monte Carlo

This analysis utilises a multi-dimensional probability distribution, with some dimensions being significantly more constrained than others. This could be from prior knowledge of parameter distributions from external data or unphysical regions in which parameters can not exist. Consequently, the Monte Carlo techniques used need to be as efficient as possible. For this analysis, the Markov Chain Monte Carlo (MCMC) technique is chosen. An MCMC technique is a Monte Carlo technique that uses a Markov chain to select which points at which to sample the parameter distribution. This technique performs a semi-random stochastic walk through the allowable parameter space. This builds a posterior distribution which has the property that the density of sampled points is proportional to the probability density of that parameter. This does mean that

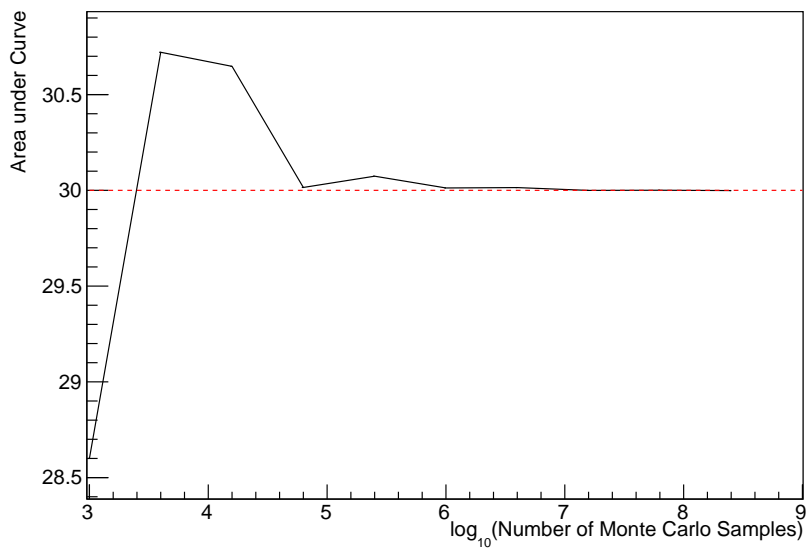


Figure 4.2: The area under a line of gradient 0.4 and intercept 1.0 for the range $0 \leq x \leq 10$ as calculated using Monte Carlo techniques as a function of the number of samples used in each repetition. The analytical solution to the area is 30 units as given by the red line.

the samples produced by this technique are not statistically independent but they will cover the space of the distribution.

A Markov chain functions by selecting the position of step \vec{x}_{i+1} based on the position of \vec{x}_i . The space in which the Markov chain selects samples is dependent upon the total number of parameters utilised within the fit, where a discrete point in this space is described by the N-dimensional space \vec{x} . In a perfectly operating Markov chain, the position of the next step depends solely on the previous step and not on the further history of the chain (\vec{x}_0 , \vec{x}_1 , etc.). However, in solving the multi-dimensionality of the fit used within this analysis, each step becomes correlated with several of the steps preceding itself. This behaviour is further explained in subsection 4.2.3. Providing the MCMC chain is well optimised, it will begin to converge towards a unique stationary distribution. The period between the chain's initial starting point and the convergence to the unique stationary distribution is colloquially known as the burn-in period. This is discussed further in subsection 4.2.3. Once the chain reaches the stationary distribution, all points sampled after that point will look like samples from that distribution.

Further details of the theories underpinning MCMC techniques are discussed in [130] but can be summarised by the requirement that the chain satisfies the three ‘regularity conditions’:

- Irreducibility: From every position in the parameter space \vec{x} , there must exist a non-zero probability for every other position in the parameter space to be reached.
- Recurrence: Once the chain arrives at the stationary distribution, every step following from that position must be samples from the same stationary distribution.
- Aperiodicity: The chain must not repeat the same sequence of steps at any point throughout the sampling period.

The output of the chain after burn-in (ie. the sampled points after the chain has reached the stationary distribution) can be used to approximate the posterior distribution and model parameters $\vec{\theta}$. To achieve the requirement that the unique stationary distribution found by the chain be the posterior distribution, one can use the Metropolis-Hastings algorithm. This guides the stochastic process depending on the likelihood of the current proposed step compared to that of the previous step. Implementation and other details of this technique are discussed in subsection 4.2.2.

4.2.2 Metropolis-Hastings Algorithm

As a requirement for MCMCs, the Markov chain implemented in this technique must have a unique stationary distribution that is equivalent to the posterior distribution. To ensure this requirement and that the regularity conditions are met, this analysis utilises the Metropolis-Hastings (MH) algorithm [135, 136]. For the i^{th} step in the chain, the MH algorithm determines the position in the parameter space to which the chain moves to based on the current step, \vec{x}_i , and the proposed step, \vec{y}_{i+1} . The proposed step is randomly selected from some

proposal function $f(\vec{x}_{i+1}|\vec{x}_i)$, which depends solely on the current step (ie. not the further history of the chain). The next step in the chain \vec{x}_{i+1} can be either the current step or the proposed step determined by whether the proposed step is accepted or rejected. To decide if the proposed step is selected, the acceptance probability, $\alpha(\vec{x}_i, \vec{y}_i)$, is calculated as

$$\alpha(\vec{x}_i, \vec{y}_{i+1}) = \min\left(1, \frac{P(\vec{y}_{i+1}|D)f(\vec{x}_i|\vec{y}_{i+1})}{P(\vec{x}_i|D)f(\vec{y}_{i+1}|\vec{x}_i)}\right). \quad (4.4)$$

Where $P(\vec{y}_{i+1}|D)$ is the posterior distribution as introduced in section 4.1. To simplify this calculation, the proposal function is required to be symmetric such that $f(\vec{x}_i|\vec{y}_{i+1}) = f(\vec{y}_{i+1}|\vec{x}_i)$. In practice, a multi-variate Gaussian distribution is used to throw parameter proposals from. This reduces Equation 4.4 to

$$\alpha(\vec{x}_i, \vec{y}_{i+1}) = \min\left(1, \frac{P(\vec{y}_{i+1}|D)}{P(\vec{x}_i|D)}\right). \quad (4.5)$$

After calculating this quantity, a random number, β , is generated uniformly between 0 and 1. If $\beta \leq \alpha(\vec{x}_i, \vec{y}_{i+1})$, the proposed step is accepted. Otherwise, the chain sets the next step equal to the current step and this procedure is repeated. This can be interpreted as if the posterior probability of the proposed step is greater than that of the current step, ($P(\vec{y}_{i+1}|D) \geq P(\vec{x}_i|D)$), the proposed step will always be accepted. If the opposite is true, ($P(\vec{y}_{i+1}|D) \leq P(\vec{x}_i|D)$), the proposed step will be accepted with probability $P(\vec{x}_i|D)/P(\vec{y}_{i+1}|D)$. This ensures that the Markov chain does not get trapped in any local minima in the potentially non-Gaussian posterior distribution. The outcome of this technique is that the density of steps taken in a discrete region is directly proportional to the probability density in that region.

4.2.3 MCMC Optimisation

As discussed in subsection 4.2.2, the proposal function invoked within the MH algorithm can take any form and the chain will still converge to the stationary distribution. At each set of proposed parameter values, a prediction of the same spectra has to be generated which requires significant computational resources.

Therefore, the number of steps taken before the unique stationary distribution is found should be minimised as only steps after convergence add information to the oscillation analysis. Furthermore, the chain should entirely cover the allowable parameter space to ensure that all values have been considered. Tuning the distance that the proposal function jumps between steps on a parameter-by-parameter basis can both minimise the length of the burn-in period and ensure that the correlation between step \vec{x}_i and \vec{x}_j is sufficiently small.

The effect of changing the width of the proposal function is highlighted in Figure 4.3. Three scenarios, each with the same underlying stationary distribution (A Gaussian of width 1.0 and mean 0.), are presented. The only difference between the three scenarios is the width of the proposal function, colloquially known as the ‘step size σ ’. Each scenario starts at an initial parameter value of 10.0 which would be considered an extreme variation. For the case where $\sigma = 0.1$, it is clear to see that the chain takes a long time to reach the expected region of the parameter. This indicates that this chain would have a large burn-in period and does not converge to the stationary distribution until step ~ 500 . Furthermore, whilst the chain does move towards the expected region, each step is significantly correlated with the previous. Considering the case where $\sigma = 5.0$, the chain approaches the expected parameter region almost instantly meaning that the burn-in period is not significant. However, there are clearly large regions of steps where the chain does not move. This is likely due to the chain proposing steps in the tails of the distribution which have a low probability of being accepted. Consequently, this chain would take a significant number of steps to fully span the allowable parameter region. For the final scenario, where $\sigma = 0.5$, you can see a relatively small burn-in period of approximately 100 steps. Once the chain reaches the stationary distribution, it moves throughout the expected region of parameter values many times, sufficiently sampling the full parameter region. This example is a single parameter varying across a continuous distribution and does not fully reflect the difficulties in the many-hundred multi-variate parameter

distribution used within this analysis. However, it does give a conceptual idea of the importance of selecting the proposal function and associated step size.

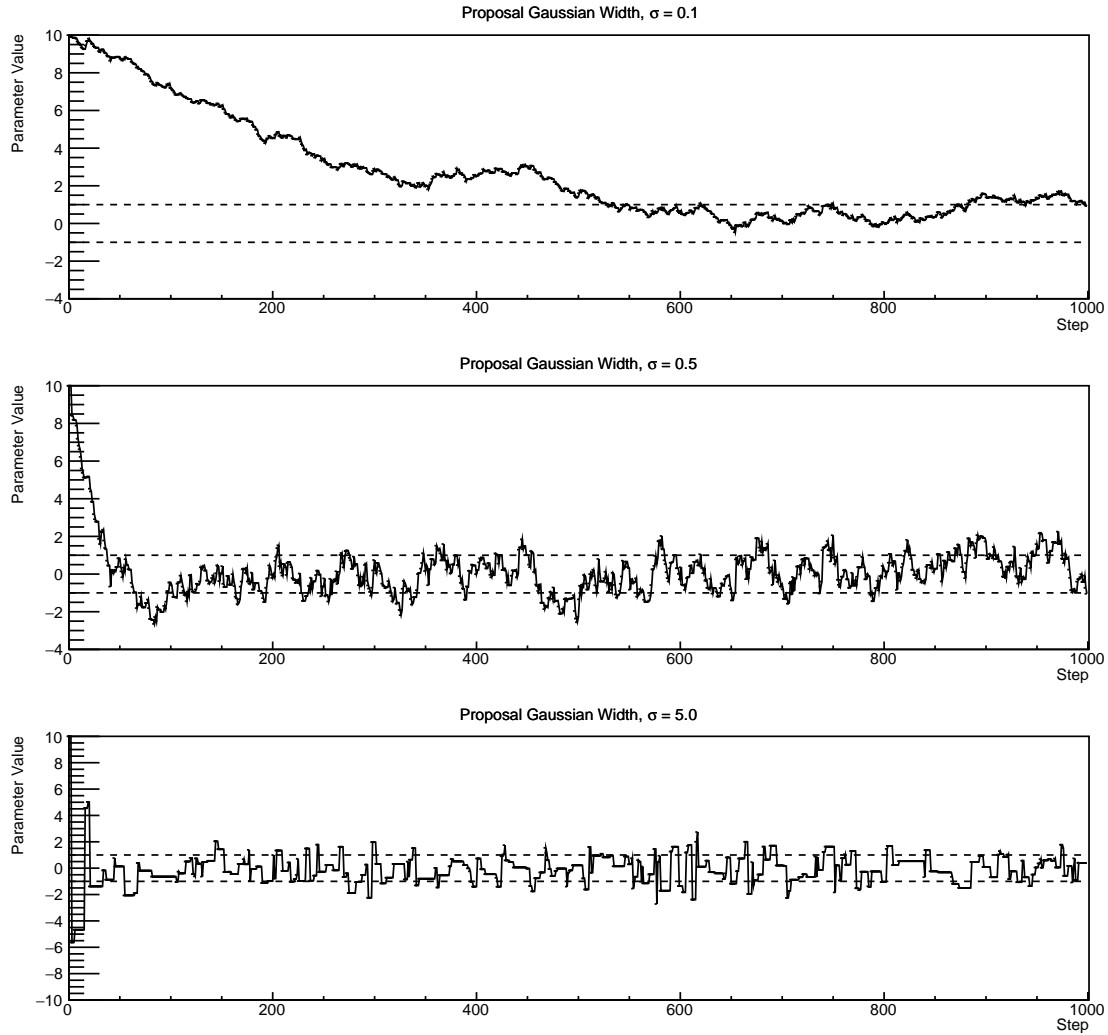


Figure 4.3: Three MCMC chains, each with a stationary distribution equal to a Gaussian centered at 0 and width 1 (As indicated by the black dotted lines). All of the chains use a Gaussian proposal function but have different widths (or ‘step size σ ’). The top panel has $\sigma = 0.1$, middle panel has $\sigma = 0.5$ and the bottom panel has $\sigma = 5.0$.

As discussed, step size tuning directly correlates to the average step acceptance rate. If the step size is too small, many steps will be accepted but the chain moves slowly. If the opposite is true, many steps will be rejected as the chain proposes steps in the tails of the distribution. Discussion in [137] suggests that the ‘ideal’ acceptance rate of a high dimension MCMC chain should be

approximately $\sim 25\%$. An “ideal” step size [137] of

$$\sigma = \frac{2.4}{N_p}, \quad (4.6)$$

where N_p is the number of parameters included in the MCMC fit. However, the complex correlations between systematics mean that some parameters have to be hand tuned and many efforts have been taken to select a set of parameter-by-parameter step sizes to approximately reach the ideal acceptance rate.

Figure 4.3 highlights the likelihood as calculated by the fit in DB: [Link to AsimovA Sensitivity Section](#) as a function of the number of steps in each chain. In practice, many independent MCMC chains are run simultaneously to parallelise the task of performing the fit. This figure overlays the distribution found in each chain. As seen, the likelihood decreases from its initial value and converges towards a stationary distribution after $\sim 1 \times 10^5$ steps.

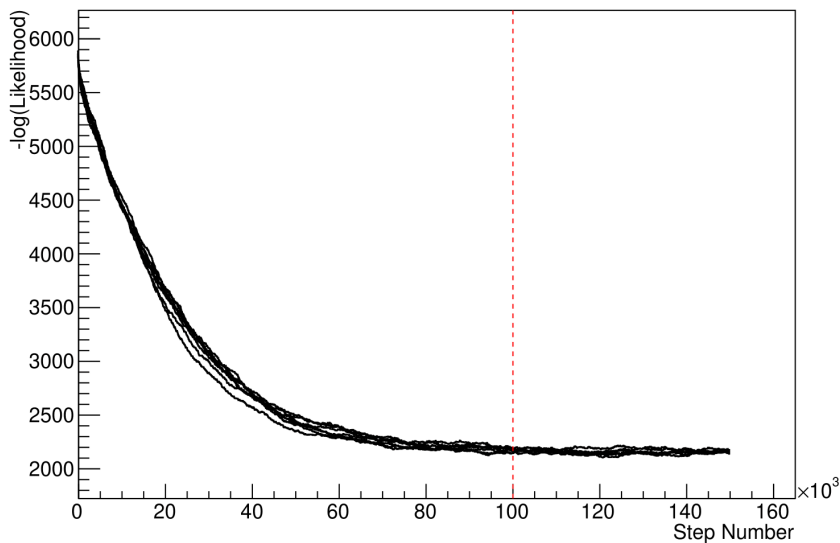


Figure 4.4: The log-likelihood from the fit detailed in DB: [Link to AsimovA Sensitivity Section](#) as a function of the number of steps accumulated in each fit. Many independent MCMC chains were run in parallel and overlaid on this plot. The red line indicates the 1×10^5 step burn-in period after which the log-likelihood becomes stable.

Multiple configurations of this analysis have been performed throughout this thesis where different samples or systematics have been used. For all of these configurations, it was found that a burnin period of 1×10^5 was sufficient in all cases.

4.3 Understanding the MCMC Results

The previous sections have described how to generate the posterior probability distribution using Bayesian MCMC techniques. However, this analysis focuses on oscillation parameter determination. The posterior distribution output from the chain is a high dimension object, with as many dimensions as there are parameters included in the oscillation analysis. However, this multi-dimensional object is difficult to conceptualize so parameter estimations are often presented in one or two-dimensional projections of this probability distribution. To do this, we invoke the marginalisation technique highlighted in subsection 4.3.1.

4.3.1 Marginalisation

The output of the MCMC chain is a highly dimensional probability distribution which is very difficult to interpret. From the standpoint of an oscillation analysis experiment, the one or two-dimensional ‘projections’ of the oscillation parameters of interest are most relevant. Despite this, the best fit values and uncertainties on the oscillation parameters of interest should correctly encapsulate the correlations to the other systematic uncertainties (colloquially called ‘nuisance’ parameters). For this joint beam and atmospheric analysis, the oscillation parameters of interest are $\sin^2(\theta_{23})$, $\sin^2(\theta_{13})$, Δm_{23}^2 , and δ_{CP} . All other parameters (Including the oscillation parameter this fit is insensitive to) are deemed nuisance parameters. To generate these projections, we rely upon integrating the posterior distribution over all nuisance parameters. This is called marginalisation. A simple example of this technique is to imagine the scenario where two coins are flipped. To determine the probability that the first coin returned a ‘head’, the exact result of the second coin flip is disregarded and simply integrated over. For the parameters of interest, $\vec{\theta}_i$, we can calculate the marginalised posterior by integrating over the nuisance parameters, $\vec{\theta}_n$. In this case, Equation 4.2 becomes

$$P(\vec{\theta}_i|D) = \frac{\int P(D|\vec{\theta}_i, \vec{\theta}_n)P(\vec{\theta}_i, \vec{\theta}_n)d\vec{\theta}_n}{\int P(D|\vec{\theta})P(\vec{\theta})d\vec{\theta}} \quad (4.7)$$

Where $P(\vec{\theta}_i, \vec{\theta}_n)$ encodes the prior knowledge about the uncertainty and correlations between the parameters of interest and the nuisance parameters. In practice, this is simply taking the one or two-dimensional projection of the multi-dimensional probability distribution.

Whilst in principle an easy solution to a complex problem, correlations between the interesting and nuisance parameters can bias the marginalised results. A similar effect is found when the parameters being marginalised over have non-Gaussian probability distributions. For example, Figure 4.5 highlights the marginalisation bias in the probability distribution found for a parameter when requiring a correlated parameter to have a positive parameter value. Due to the complex nature of this oscillation parameter fit presented in this thesis, there are correlations occurring between the oscillation parameters of interest and the other nuisance parameters included in the fit.

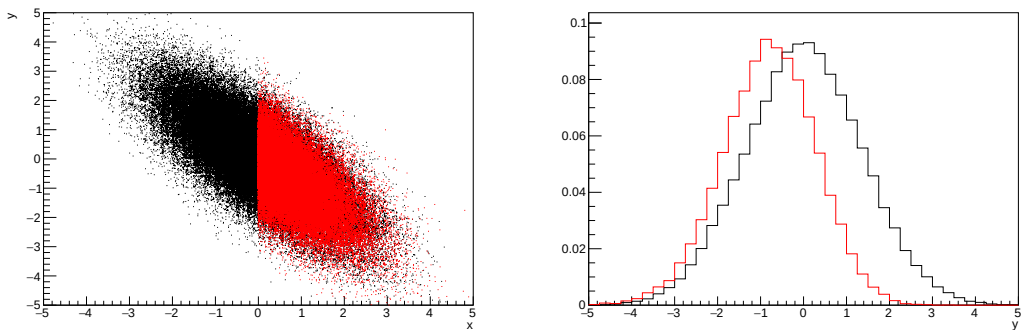


Figure 4.5: Left: The two dimensional probability distribution for two correlated parameters x and y . The red distribution shows the two dimensional probability distribution when $0 \leq x \leq 5$. Right: The marginalised probability distribution for the y parameter found when requiring the x to be bound between $-5 \leq x \leq 5$ and $0 \leq x \leq 5$ for the black and red distribution, respectively.

4.3.2 Parameter Estimation and Credible Intervals

The purpose of this analysis is to determine the best fit values for the oscillation parameters that the beam and atmospheric samples are sensitive to: $\sin^2(\theta_{23})$, $\sin^2(\theta_{13})$, Δm_{23}^2 , and δ_{CP} . Typically, the results presented take the form of one or two-dimension marginalised probability distributions for the appearance

($\sin^2(\theta_{13})$ and δ_{CP}) and disappearance ($\sin^2(\theta_{23})$ and Δm_{23}^2) parameters. The posterior probability density taken from the output MCMC chain is binned in these parameters. The parameter best-fit point is then taken to be the value that has the highest posterior probability. This is performed in both one and two-dimensional projections.

However, the single best-fit point in a given parameter is not of much use on its own. We would also like to determine the uncertainty, or credible interval, on that best-fit point. The definition of the 1σ credible interval is that we have 68% belief that the parameter is within those bounds. For a more generalised definition, the credible interval is the region, R , of the posterior distribution that contains a specific fraction of the total probability, such that

$$\int_R P(\theta|D)d\theta = \alpha \quad (4.8)$$

Where θ is the parameter on which we calculate the credible interval. This technique then calculates the $\alpha \times 100\%$ credible interval.

In practice, this analysis uses the highest posterior density (HPD) credible intervals which are calculated through the following method. First, the probability distribution is area-normalised such that it has an integrated area equal to 1.0. The bins of probability are then summed from the highest to lowest until the sum exceeds the 1σ level (0.68 in this example). This process is repeated for a range of credible intervals, notably the 1σ , 2σ and 3σ along with other levels where the critical values for each level can be found in [77]. This process can be repeated for the two-dimensional probability distributions by creating two-dimensional contours of credible intervals rather than a one-dimensional result.

4.3.3 Bayesian Model Comparisons

Due to the matter resonance, this analysis has some sensitivity to the mass hierarchy of neutrino states (whether Δm_{23}^2 is positive or negative) and the octant of $\sin^2(\theta_{23})$. The Bayesian approach utilised within this analysis gives an intuitive method of model comparison by determining which hypothesis is most

favourable. Taking the ratio of Equation 4.3 for the two hypotheses of normal hierarchy, NH , and inverted hierarchy, IH , gives

$$\frac{P(\vec{\theta}_{NH}|D)}{P(\vec{\theta}_{IH}|D)} = \frac{P(D|\vec{\theta}_{NH})}{P(D|\vec{\theta}_{IH})} \times \frac{P(\vec{\theta}_{NH})}{P(\vec{\theta}_{IH})}. \quad (4.9)$$

The middle term defines the Bayes factor which is a data-driven interpretation of how strong the data prefers one hierarchy to the other. For this analysis, equal priors on both mass hierarchy hypotheses are chosen ($P(\vec{\theta}_{NH}) = P(\vec{\theta}_{IH}) = 0.5$). In practice, the MCMC chain proposes a value of $|\Delta m_{23}^2|$ and then applies a 50% probability that the value is sign flipped. Consequently, the Bayes factor can be calculated from the ratio of the probability density in either hypothesis. This equates to counting the number of steps taken in the normal and inverted hierarchies and taking the ratio. The same approach can be taken to compare the upper octant (UO) compared to the lower octant (LO) hypothesis of $\sin^2(\theta_{23})$.

Whilst the value of the Bayes factor should always be shown, the Jeffreys scale [138] (highlighted in Table 4.1) gives an indication of the strength of preference for one model compared to the other. Other interpretations of the strength of preference of a model exist, e.g. the Kass and Raferty Scale [139].

$\log_{10}(B_{AB})$	B_{AB}	Strength of Preference
< 0.0	< 1	No preference for hypothesis A (Supports hypothesis B)
0.0 – 0.5	1.0 – 3.16	Preference for hypothesis A is weak
0.5 – 1.0	3.16 – 10.0	Preference for hypothesis A is substantial
1.0 – 1.5	10.0 – 31.6	Preference for hypothesis A is strong
1.5 – 2.0	31.6 – 100.0	Preference for hypothesis A is very strong
> 2.0	> 100.0	Decisive preference for hypothesis A

Table 4.1: Jeffreys scale for strength of preference for two models A and B as a function of the calculated Bayes factor ($B_{AB} = B(A/B)$) between the two models [138]. The original scale is given in terms of $\log_{10}(B(A/B))$ but converted to linear scale for easy comparison throughout this thesis.

4.3.4 Comparison of MCMC Output to Expectation

To ensure the fit is performing well, a best-fit spectrum is produced using the posterior probability distribution and compared with the data, allowing easy

by-eye comparisons to be made. A simple method of doing this is to perform a comparison in the fitting parameters (For instance, the reconstructed neutrino energy and lepton direction for T2K far detector beam samples) of the spectra generated by the MCMC chain to ‘data’. This ‘data’ could be true data or some variation of Monte Carlo prediction. This allows easy comparison of the MCMC probability distribution to the data. To perform this, N steps from the post burn-in MCMC chain are randomly selected (Where for all plots of this style in this thesis, $N = 3000$). From these, the Monte Carlo prediction at each step is generated by reweighting the model parameters to the values specified at that step. Due to the probability density being directly correlated with the density of steps in a certain region, parameter values close to the best fit value are most likely to be selected.

In practice, for each bin of the fitting parameters has a probability distribution of event rates, with one entry per sampled MCMC step. This distribution is binned where the bin with the highest probability is selected as the mean and an error on the width of this probability distribution is calculated using the approach highlighted in subsection 4.3.2. Consequently, the best fit distribution in the fit parameter is not necessarily that which would be attained by reweighting the Monte Carlo prediction to the most probable parameter values.

A similar study can be performed to illustrate the freedom of the model parameter space prior to the fit. This can be done by throwing parameter values from the prior uncertainty of each parameter. This becomes troublesome for parameters with no prior uncertainty as the range is technically infinite. Where applicable solutions to remove these have been addressed.

5

Simulation, Reconstruction, and Event Reduction

As a crucial part of the oscillation analysis, an accurate prediction of the expected neutrino spectrum at the far detector is required. This includes modeling the flux generation, neutrino interactions, and detector effects. All of the simulation packages required to do this are briefly described in section 5.1. The reconstruction of neutrino events inside the far detector, including the `fitQun` algorithm, is documented in section 5.2. This also includes data quality checks of the SK-V data which the author performed for the T2K oscillation analysis presented at Neutrino 2020 [76]. Finally, section 5.3 describes the steps taken in the SK detector to trigger on events of interest whilst removing the comparatively large rate of cosmic ray muon events.

5.1 Simulation

In order to generate a Monte Carlo prediction of the expected event rate at the far detector, all the processes in the beam and atmospheric flux, neutrino interaction, and detector need to be modeled. Each of these parts is individually modeled and each of them is detailed below.

The beamline simulation consists of three distinct parts: the initial hadron interaction modeled by FLUKA [140], the target station geometry and particle tracking performed by JNUBEAM, [141, 142] and any hadronic re-interactions simulated by GCALOR [143]. The primary hadronic interactions are $O(10)\text{GeV}$, where FLUKA matches external cross-section data better than GCALOR [144]. However, FLUKA is not very adaptable so a small simulation is built to model the interactions in the target and the output is then passed to JNUBEAM and GCALOR for propagation. The hadronic interactions are tuned to data from the NA61/SHINE [145–147] and HARP [148] experiments. The tuning is done by reweighting the FLUKA and GCALOR predictions to match the external data multiplicity and cross-section measurements, based on final state particle kinematics [144]. The culmination of this simulation package generates the predicted flux for neutrino and antineutrino beam modes which are illustrated in Figure 3.7.

The atmospheric neutrino flux is simulated by the HKKM model [42, 44]. The primary cosmic ray flux is tuned to AMS [149] and BESS [150] data assuming the US-standard atmosphere '76 [151] density profile and includes geomagnetic field effects. The primary cosmic rays interact to generate pions and muons. The interaction of these secondary particles to generate neutrinos is handled by DPMJET-III [152] for energies above 32GeV and JAM [44, 153] for energies below that value **DB: Question for Giles: Why different generators for above/below 32GeV?**. These hadronic interactions are tuned to BESS and L3 data [154, 155] using the same methodology as the tuning of the beamline simulation. The energy and cosine zenith predictions of $\nu_e, \bar{\nu}_e, \nu_\mu, \bar{\nu}_\mu$ flux are given in Figure 2.3 and Figure 2.5, respectively. The flux is approximately symmetrical and peaked around the horizon ($\cos(\theta_Z) = 0.0$). This is because horizontally-going pions and kaons can travel further than their vertically-going counterparts resulting in a larger probability of decaying to neutrinos. The symmetry is broken in lower-energy neutrinos due to geomagnetic effects, which modify the track of the primary cosmic rays. Updates to the HKKM model are currently ongoing [156].

Once a flux prediction has been made for all three detectors, NEUT 5.4.0 [157, 158] models the interactions of the neutrinos in the detectors. For the purposes of this analysis, quasi-elastic (QE), meson exchange (MEC), single meson production (PROD), coherent pion production (COH), and deep inelastic scattering (DIS) interactions are simulated. These interaction categories can be further broken down by whether they were propagated via a W^\pm boson in Charged Current (CC) interactions or via a Z^0 boson in Neutral Current (NC) interactions. CC interactions have a charged lepton in the final state, which can be flavour-tagged in reconstruction to determine the flavour of the neutrino. In contrast, NC interactions have a neutrino in the final state so no flavour information can be determined from the observables left in the detector after an interaction. This is the reason why neutrinos which interact through NC modes are assumed to not oscillate within this analysis. Both CC and NC interactions are modeled for all the above interaction categories, other than MEC interactions which are only modeled for CC events.

As illustrated in Figure 5.1, CCQE interactions dominate the cross-section of neutrino interactions around $E_\nu \sim 0.5\text{GeV}$. The NEUT implementation adopts the Llewellyn Smith [159] model for neutrino-nucleus interactions, where the nuclear ground state of any bound nucleons (neutrino-oxygen interactions) is approximated by a spectral-function [160] model that simulates the effects of Fermi momentum and Pauli blocking. The cross-section of QE interactions is controlled by vector and axial-vector form factors parameterised by the BBBA05 [161] model and a dipole form factor with $M_A^{QE} = 1.21\text{GeV}$ fit to external data [162], respectively. NEUT implements the Valencia [163] model to simulate MEC events, where two nucleons and two holes in the nuclear target are produced (Often called 2p2h interactions).

For neutrinos of energy $O(1)\text{GeV}$, PROD interactions become dominant. These predominantly produce charged and neutral pions although γ , kaon, and η production is also considered. To simulate these interactions, the Berger-Sehgal [164] model is implemented within NEUT. It simulates the excitation of a

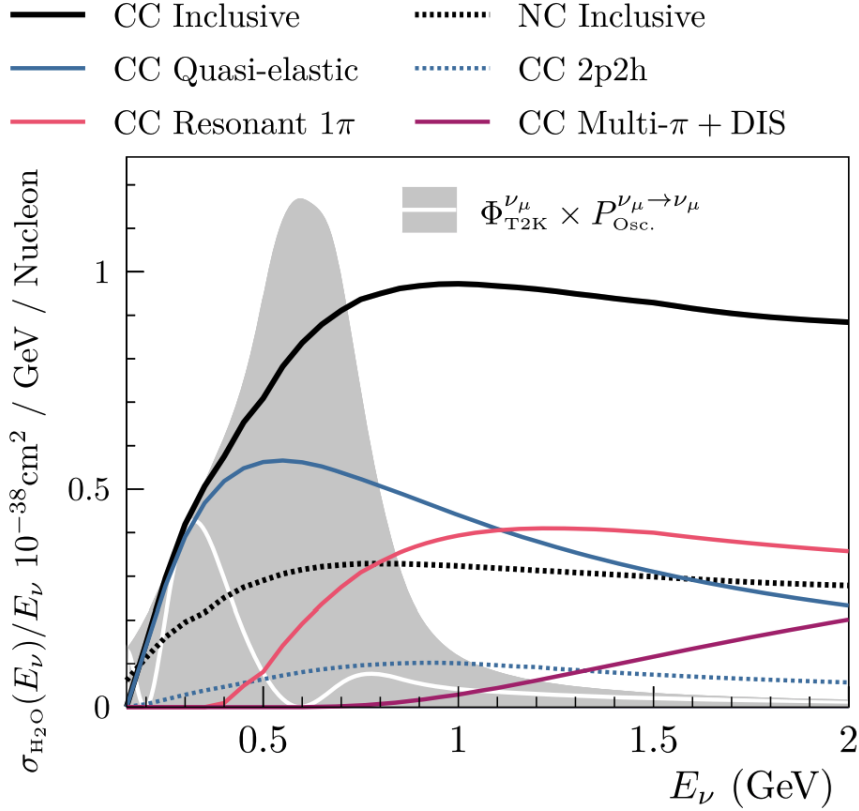


Figure 5.1: The NEUT prediction of the ν_μ -H₂O cross-section overlaid on the T2K ν_μ flux. The charged current (black, solid) and neutral current (black, dashed) inclusive, charged current quasi-elastic (blue, solid), charged current 2p2h (blue, dashed), charged current single pion production (pink), and charged current multi- π and DIS (Purple) cross-sections are illustrated. Figure taken from [157].

nucleon from a neutrino interaction, production of an intermediate baryon, and the subsequent decay to a single meson or γ . Pions can also be produced through COH interactions, which occur when the incoming neutrino interacts with the entire oxygen nucleus leaving a single pion outside of the nucleus. NEUT utilises the Berger-Sehgal [165] model to simulate these COH interactions.

DIS and multi- π producing interactions become the most dominant for energies $> O(5)\text{GeV}$. PYTHIA [166] is used to simulate any interaction with invariant mass $W > 2\text{GeV}/c^2$, which produces at least one meson. For any interaction which produces at least two mesons but has $W < 2\text{GeV}/c^2$, the Bronner model is used [167]. Both of these models use parton distribution functions based on the Bodek-Yang model [168–170].

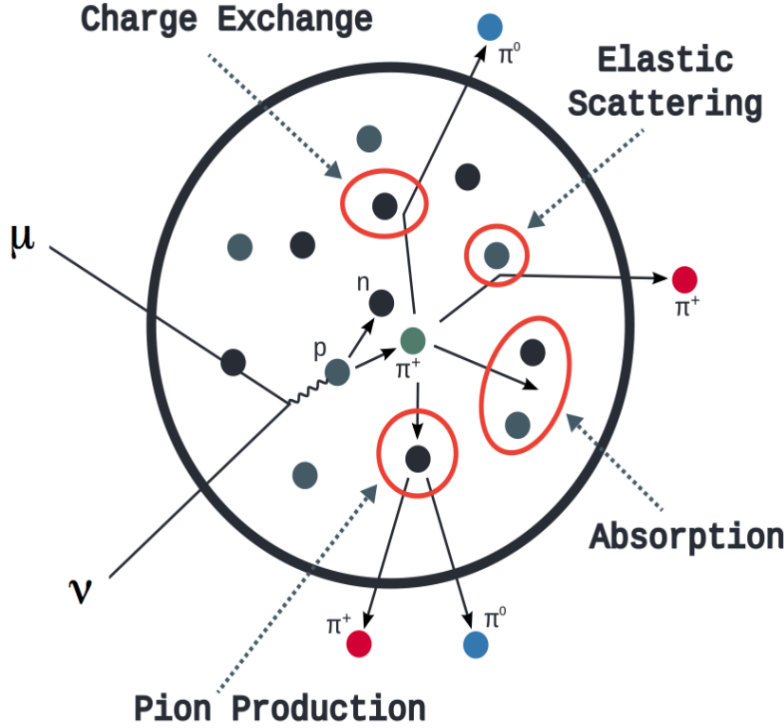


Figure 5.2: Illustration of the various processes which a pion can undergo before exiting the nucleus. Taken from [171].

Any pion which is produced within the nucleus can re-interact through final state interactions before it exits, as illustrated by the scattering, absorption, production, and exchange interactions in Figure 5.2. These re-interactions alter the observable particles within the detector. For instance, if the charged pion from a CC PROD interaction is absorbed, the observables would mimic a CC QE interaction. To simulate these effects, NEUT uses a semi-classical intranuclear cascade model [157]. This cascade functions by stepping the pion through the nucleus in fixed-length steps equivalent to $dx = R_N/100$, where R_N is the radius of the nucleus. At each step, the simulation allows the pion to interact through scattering, charged exchange, absorption, or production with an interaction-dependent probability calculated from a fit to external data [172]. This cascade continues until the pion is absorbed or exits the nucleus.

Once the final state particle kinematics have been determined by NEUT, they are passed into the detector simulation. The near detectors, ND280 and

INGRID, are simulated using a GEANT4 package [116, 173] to simulate the detector geometry, particle tracking, and energy deposition. The response of the detectors is simulated using the elecSim package [116]. The far detector simulation is based upon the original Kamiokande experiment software which uses the GEANT3-based SKDETSIM [116, 174] package. This simulates the interactions of particles in the water as well as Cherenkov light production. The water quality and PMT calibration measurements detailed in subsection 3.1.2 are also used within this simulation to make accurate predictions of the detector response.

5.2 Event Reconstruction at SK

Any event which generates optical photons that occurs in SK will be observed by the PMT array, where each PMT records the time and accumulated charge. This recorded information is shown in event displays similar to those illustrated in Figure 5.3. To be useful for physics analyses, this series of PMT hit information needs to be reconstructed to determine the particle’s identity and kinematics (or track parameters): four-vertex, direction, and momenta. The reconstruction uses the fact that the charge and timing distribution of photons generated by a particular particle in an event is dependent upon its initial kinematics. Electron and muon rings are distinguished by their “fuzziness”. Muons are heavier and less affected by scattering or showering meaning they typically produce “crisp” rings. Electrons are more likely to interact via electromagnetic showering or scattering which results in larger variations of their direction from the initial direction. Consequently, electrons typically produce “fuzzier” rings compared to muons.

For the purposes of this analysis, the `fiTQun` reconstruction algorithm is utilised. Its core function is to compare a prediction of the accumulated charged and timing distribution from each PMT, generated for a particular particle identity and track parameters, to that observed in the neutrino event. It determines the preferred values by minimising a likelihood function which includes information from PMTs which were hit and those that were not hit. `fiTQun` performs a

simultaneous fit of particle kinematics and identity, improving both the accuracy of the fit parameters and the rejection of neutral current π^0 events [175, 176]. The `fitQun` algorithm is based on the key concepts of the MiniBooNE reconstruction algorithm [177] and is described in [178] which is summarised below. The `fitQun` algorithm improves upon the `APFit` reconstruction algorithm which has been used for many previous SK analyses. `APFit` fits the vertex from timing information and then fits the momentum and direction of the particle from PMT hits within a 43 deg Cherenkov cone (which assumes an ultra-relativistic particle). It then fits the particle identity once the track parameters have been fit.

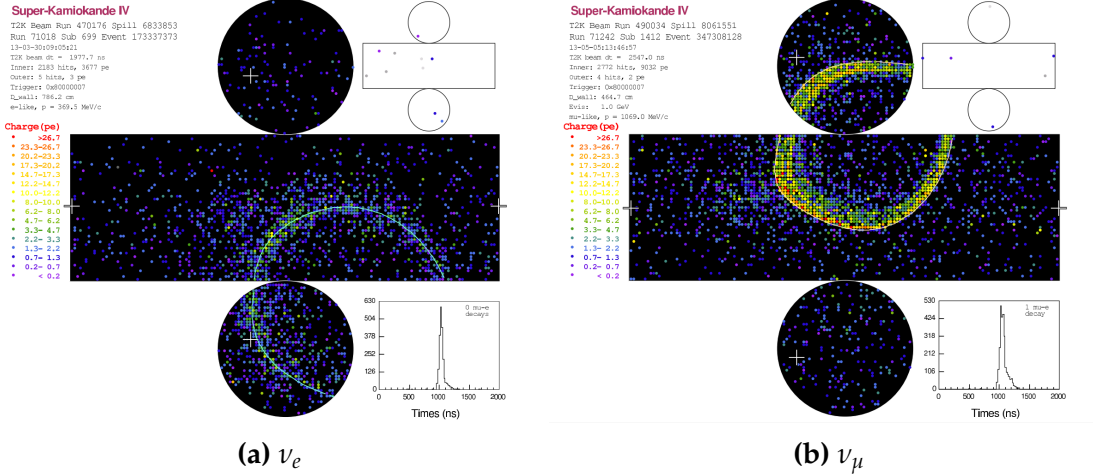


Figure 5.3: Event displays from Super Kamiokande illustrating the “crisp” ring from a muon and the typically “fuzzy” electron ring. Each pixel represents a PMT and the color scheme denotes the accumulated charge deposited on that PMT. Figures taken from [179].

The `fitQun` reconstruction algorithm proceeds by:

- **Vertex pre-fitting:** An estimate of the vertex is made using a goodness-of-fit metric
- **Peak finding:** The initial time of the event is determined by clustering events by time residuals
- **Single-ring fits:** Given the pre-fit vertex and estimated time of interaction, a maximum likelihood technique searches for a single particle generating light

- **Multi-ring fits:** Seeded from the single-ring fits, hypotheses with multiple light-producing particles are considered using the same maximum likelihood technique

An event in SK can consist of a primary and decay particles. For example, a charged current muon neutrino interaction can generate two particles that have the potential of generating Cherenkov photons: the primary muon, and the secondary decay-electron from the muon. To ensure the particles are reconstructed separately, each event is divided into time clusters which are called “subevents”. Subevents after the primary subevent are considered to be decay electrons. To find all the subevents in an event, a vertex goodness metric is calculated for some vertex position \vec{x} and time t ,

$$G(\vec{x}, t) = \sum_i^{\text{hit PMTs}} \exp \left(-\frac{1}{2} \left(\frac{T_{Res}^i(\vec{x}, t)}{\sigma} \right)^2 \right) \quad (5.1)$$

where

$$T_{Res}^i(\vec{x}, t) = t^i - t - |R_{PMT}^i - \vec{x}| / c_n \quad (5.2)$$

is the residual hit time. It is the difference in time between the PMT hit time, t^i , of the i^{th} PMT and the expected time of the PMT hit if the photon was emitted at the start of the vertex. R_{PMT}^i is the position of the i^{th} PMT, c_n is the speed of light in water and $\sigma = 4\text{ns}$ which is comparable to the time resolution of the PMT. When the proposed fit values of time and vertex are close to the true values, $T_{Res}^i(\vec{x}, t)$ tends to zero resulting in subevents appearing as spikes in the goodness metric. The proposed fit vertex and time are grid-scanned, and the values which maximise the goodness metric are selected as the “pre-fit vertex”. Whilst this predicts a vertex for use in the clustering algorithm, the final vertex is fit using the higher-precision maximum likelihood method described below.

Once the pre-fit vertex has been determined, the goodness metric is scanned as a function of t to determine the number of subevents. A peak-finding algorithm is then used on the goodness metric, requiring the goodness metric to exceed

some threshold and drop below a reduced threshold before any subsequent additional peaks are considered. The thresholds are set such that the rate of false peak finding is minimised while still attaining good data to Monte Carlo agreement. To improve performance, the pre-fit vertex for each delayed subevent is re-calculated after PMT hits from the previous subevent are masked. This improves the decay-electron tagging performance. Once all subevents have been determined, the time window around each subevent is then defined by the earliest and latest time which satisfies $-180 < T_{Res}^i < 800\text{ns}$. The subevents and associated time windows are then used as seeds for further reconstruction.

For a given subevent, the `fitQun` algorithm constructs a likelihood based on the accumulated charge q_i and time information t_i from the i^{th} PMT,

$$L(\Gamma, \vec{\theta}) = \prod_j^{\text{unhit}} P_j(\text{unhit}|\Gamma, \vec{\theta}) \prod_i^{\text{hit}} \{1 - P_i(\text{unhit}|\Gamma, \vec{\theta})\} f_q(q_i|\Gamma, \vec{\theta}) f_t(t_i|\Gamma, \vec{\theta}), \quad (5.3)$$

where $\vec{\theta}$ defines the track parameters; vertex position, direction vector and momenta, and Γ represents the particle hypothesis. $P_i(\text{unhit}|\Gamma, \vec{\theta})$ is the probability of the i^{th} tube to not register a hit given the track parameters and particle hypothesis. The charge likelihood, $f_q(q_i|\Gamma, \vec{\theta})$, and time likelihood, $f_t(t_i|\Gamma, \vec{\theta})$, represents the probability density function of observing charge q_i and time t_i on the i^{th} PMT given the specified track parameters and particle hypothesis.

The predicted charge is calculated based on contributions from both the direct light and the scattered light. The direct light contribution is determined based on the integration of the Cherenkov photon profile along the track. PMT angular acceptance, water quality, and calibration measurements discussed in subsection 3.1.2 are included to accurately predict the charge probability density at each PMT. The scattered light is calculated in a similar way, although it includes a scattering function that depends on the vertex of the particle and the position of the PMT. The charge likelihood is calculated by comparing the prediction to the observed charge in the PMT.

The time likelihood is approximated to depend on the vertex \vec{x} , direction \vec{d} , and time t of the track as well as the particle hypothesis. The expected time for PMT hits is calculated by assuming unscattered photons being emitted from the midpoint of the track, S_{mid} ,

$$t_{exp}^i = t + S_{mid}/c + |R_{PMT}^i - \vec{x} - S_{mid}\vec{d}|/c_n, \quad (5.4)$$

where c is the speed of light in a vacuum. The time likelihood is then expressed in terms of the residual difference between the PMT hit time and the expected hit time, $t_{Res}^i = t^i - t_{exp}^i$. The particle hypothesis and momentum also affect the Cherenkov photon distribution. These parameters modify the shape of the time likelihood density since in reality not all photons are emitted at the midpoint of the track. As with the charge likelihood, the contributions from both the direct and scattered light to the time likelihood density are calculated separately, which are both calculated from particle gun studies.

The track parameters and particle identity which maximise $L(\Gamma, \vec{\theta})$ are defined as the best-fit parameters. In practice MINUIT [180] is used to minimise the value of $-\ln L(\Gamma, \vec{\theta})$. The `f1TQun` algorithm considers an electron-like, muon-like, and charged pion-like hypothesis for events with a single final state particle, denoted “single-ring events”. The particle’s identity is determined by taking the ratio of the likelihood of each of the hypotheses. For instance, electrons and muons are distinguished by considering the value of $\ln(L(e, \vec{\theta}_e)/L(\mu, \vec{\theta}_\mu))$ in comparison to the reconstructed momentum of the electron hypothesis, as illustrated by Figure 5.4. The coefficients of the discriminator between electron-like and muon-like events are determined from Monte Carlo studies [178]. Similar distributions exist for distinguishing electron-like events from π^0 -like events, and muon-like events from pion-like events. The cuts are defined as,

$$\begin{aligned} \text{Electron/Muon} : & \ln(L_e/L_\mu) > 0.2 \times p_e^{rec} [\text{MeV}], \\ \text{Electron}/\pi^0 : & \ln(L_e/L_{\pi^0}) < 175 - 0.875 \times m_{\gamma\gamma} [\text{MeV}], \\ \text{Muon/Pion} : & \ln(L_\mu/L_{\pi^\pm}) < 0.15 \times p_\mu^{rec} [\text{MeV}], \end{aligned} \quad (5.5)$$

as taken from [181], where p_e^{rec} and p_μ^{rec} are the reconstructed momentum of the single-ring electron and muon fits, respectively. $m_{\gamma\gamma}$ represents the reconstructed invariant mass of the two photons emitted from π^0 decay. Typically, the distance between a particular entry in these two-dimensional distributions and the cut-line is termed the PID parameter and is illustrated in Figure 5.5.

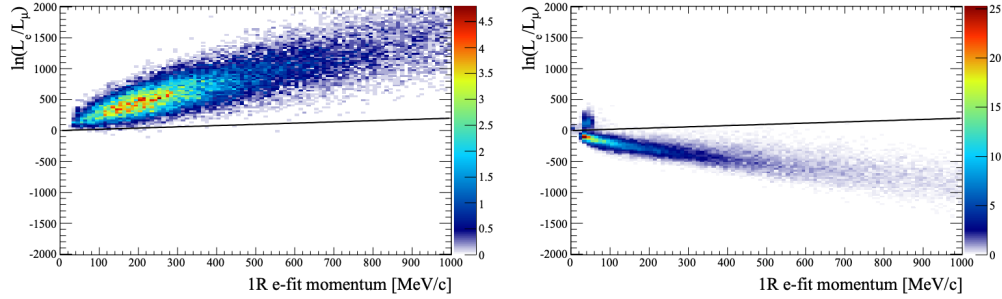


Figure 5.4: The difference of the electron-like and muon-like log-likelihood compared to the reconstructed single-ring fit momentum for atmospheric ν_e (left) and ν_μ (right) samples. The black line represents the cut used to discriminate electron-like and muon-like events, which coefficients obtained from Monte Carlo studies. Figures taken from [178].

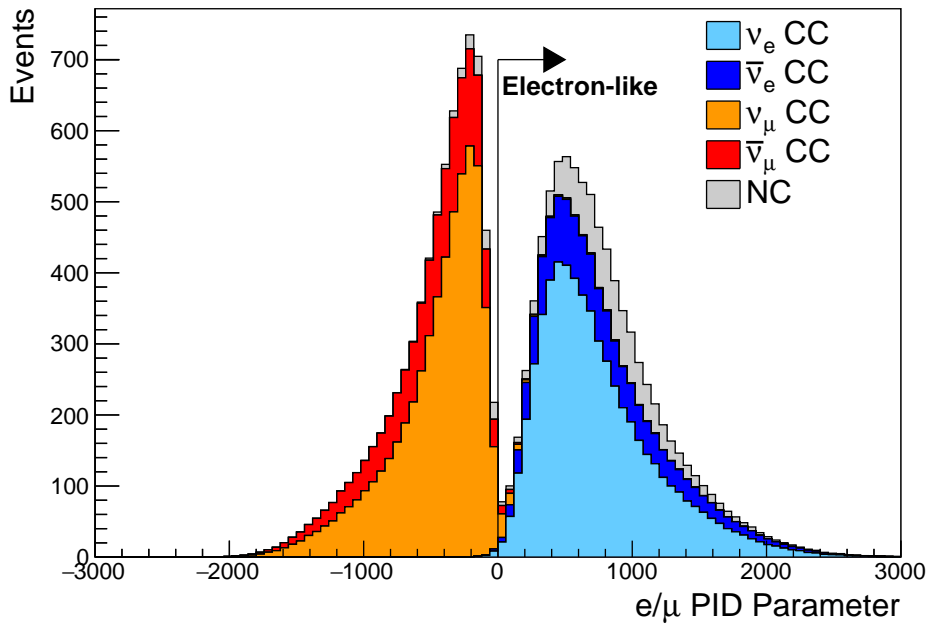


Figure 5.5: The electron/muon PID separation parameter for all sub-GeV single-ring events in SK-IV. The charged current (CC) component is broken down in four flavours of neutrino (ν_μ , $\bar{\nu}_\mu$, ν_e and $\bar{\nu}_e$). Events with positive values of the parameter are determined to be electron-like.

The `fitQun` algorithm also considers a π^0 hypothesis. To do this, it performs a fit looking for two standard electron-hypothesis tracks which point to the same four-vertex. This assumes the electron tracks are generated from photon-conversion so the electron tracks actually appear offset from the proposed π^0 vertex. For these fits, the conversion length, direction, and momentum of each photon are also considered as track parameters which are then fit in the same methodology as the standard single-ring hypotheses.

Whilst lower energy events are predominantly single-ring events, higher energy neutrino events can generate final states with multiple particles which generate Cherenkov photons. These “multi-ring” hypotheses are also considered in the `fitQun` algorithm. When calculating the charge likelihood density, the predicted charge associated with each ring is calculated separately and then summed to calculate the total accumulated charge on each PMT. Similarly, the time likelihood for the multi-ring hypothesis is calculated assuming each ring is independent. Each track is time-ordered based on the time of flight from the center of the track to the PMT and the direct light from any ring incident on the PMT is assumed to arrive before any scattered light. To reduce computational resource usage, the multi-ring fits only consider electron-like and charged pion-like rings as the pion fit can be used as a proxy for a muon fit due to their similar mass.

Multi-ring fits proceed by proposing another ring to the previous fit and then fitting the parameters in the method described above. Typically, multi-ring fits have the largest likelihood because of the additional degrees of freedom introduced. A likelihood value is calculated for the n -ring and $(n + 1)$ -ring hypotheses, where the additional ring is only included if the likelihood value is above 9.35, based on Monte Carlo studies in [182].

5.2.1 Validation of Reconstruction in SK-V

As an example of how the reconstruction depends on the detector conditions, the author of this thesis assessed the quality of event reconstruction for SK-V data.

The detector systematics used in the T2K-only oscillation analysis are determined using data-to-Monte Carlo comparisons of the SK-IV data [183]. Due to tank-open maintenance occurring between SK-IV and SK-V, the dark rate of each PMT was observed to increase in SK-V due to light exposure for a significant time during the repairs. This increase can be seen in Figure 5.6. Run-10 of the T2K experiment was conducted in the SK-V period, so the consistency of SK-IV and SK-V data needs to be studied to determine whether the SK-IV-defined systematics can be applied to the run-10 data. This comparison study was performed using the stopping muon data set for both the SK-IV and SK-V periods. This data sample is used due to the high rate of interactions ($O(200)$ events per hour) as well as having similar energies to muons from CCQE ν_μ interactions from beam interactions. The rate of cosmic muons does depend on the solar activity cycle [184] but has been neglected in this comparison study. This is because the shape of the distributions is most important for the purposes of being compared to the detector systematics. The SK-IV and SK-V data samples consist of 2398.42 and 626.719 hours of data which equates to 686k and 192k events respectively.

The predicted charge calculated in the `fitQun` algorithm includes a contribution from the photoelectron emission due to dark noise. Therefore, the increase in the SK-V dark rate needs to be accounted for. In practice, the average dark rate in each SK period is calculated and used as an input in the reconstruction. This is calculated by averaging the dark rate per run for each period separately, using the calibration measurements detailed in subsection 3.1.2. The average dark rate from SK-IV and SK-V were found to be 4.57kHz and 6.30kHz, respectively. The charges associated with the muon and decay electron subevents are illustrated in Figure 5.7. The photoelectron emission from dark noise is more significant for events that have lower energy. This is because this contribution becomes more comparable to the number of photoelectrons emitted from incident photons in lower-energy events. This behaviour is observed in the data, where the charge deposited by the muon subevent is mostly unaffected by the increase in dark rate, whilst the charge associated with the decay-electron is clearly affected.

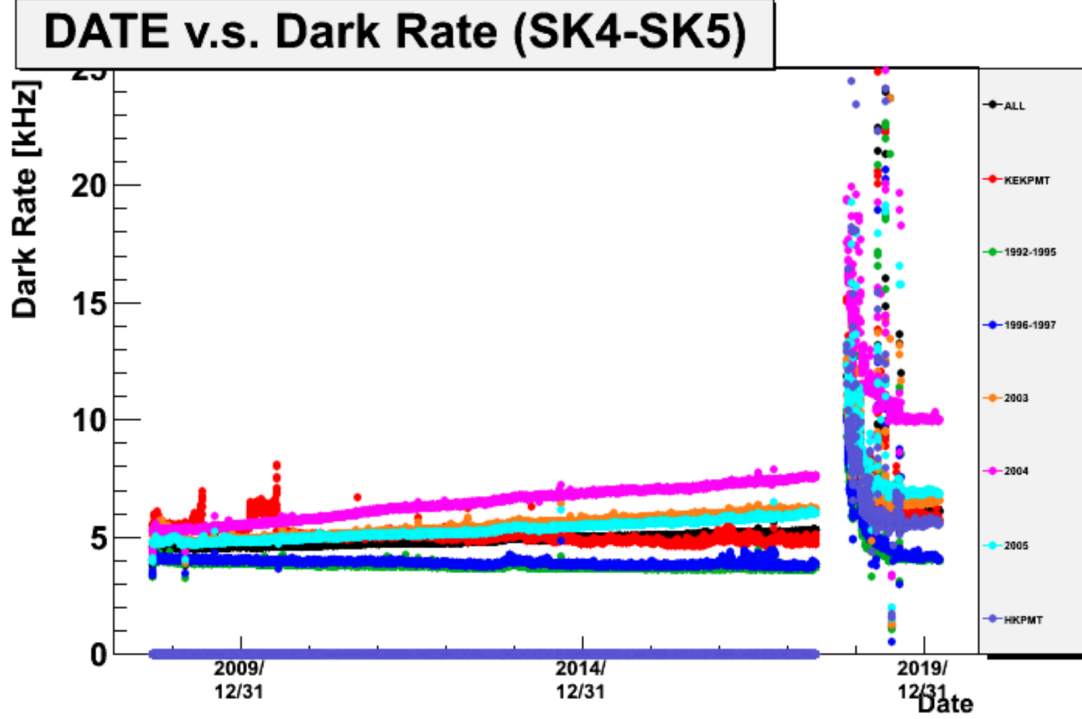


Figure 5.6: The variation of the measured dark rate as a function of date, broken down by PMT type. The SK-IV and SK-V periods span September 2008 to May 2018 and January 2019 to July 2020, respectively. The break in measurement in 2018 corresponds to the period of tank repair and refurbishment. Figure adapted from [183].

The energy scale systematic is estimated from data-to-Monte Carlo differences in the stopping muon sample in [185] and found to be 2.1%. To determine the consistency of SK-IV and SK-V with respect to the energy scale systematic, the muon momentum distribution is compared between the two SK periods. As the total number of Cherenkov photons is integrated across the track length, the reconstructed momentum divided by track length (or range) is compared between SK-IV and SK-V as illustrated in Figure 5.8.

The consistency between these distributions has been computed in two ways. Firstly, a Gaussian is fit to the peak of each distribution separately, whose mean is found to be $(2.272 \pm 0.003)\text{MeV/cm}$ and $(2.267 \pm 0.006)\text{MeV/cm}$ for SK-IV and SK-V respectively. The ratio of these is equal to 1.002 ± 0.003 . The means of the Gaussian fits are consistent with the expected stopping power of a minimum ionising muon for a target material (water) with $Z/A \sim 0.5$ [186]. The second

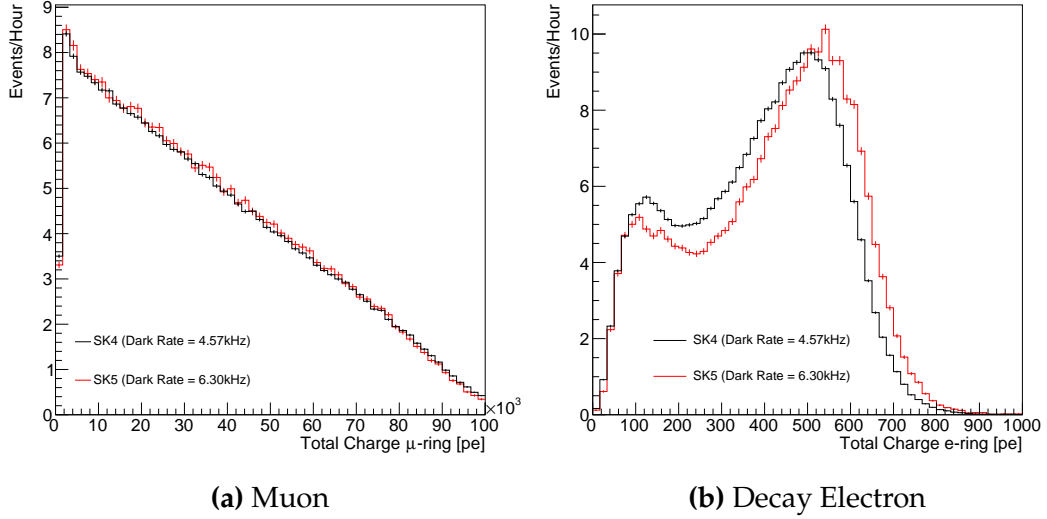


Figure 5.7: Comparison of the measured raw charge deposited per event from the stopping muon data samples between SK-IV (Blue) and SK-V (Red), split by the primary muon subevent and the associated decay electron subevent.

consistency check is performed by introducing a nuisance parameter, α , which modifies the SK-V distribution. The value of α which minimises the χ^2 value between the SK-IV and SK-V is determined by scanning across a range of values. This is repeated by applying the nuisance parameter as both a multiplicative factor and an additive shift. The χ^2 distributions for different values of α is illustrated in Figure 5.9. The values which minimise the χ^2 are found to be 0.0052 and 1.0024 for the additive and multiplicative implementations, respectively. No evidence of shifts larger than the 2.1% uncertainty on the energy scale systematic has been found in the reconstructed momentum distribution of SK-IV and SK-V.

5.3 Event Reduction at SK

Atmospheric neutrino events observed in the SK detector are categorised into three different types of samples: fully contained (FC), partially contained (PC) and up-going muon (Up- μ), using PMT hit signatures in the inner and outer detector (ID and OD, respectively). To identify FC neutrino events, it is required that the neutrino interacts inside the fiducial volume of the ID and that no significant OD activity is observed. For this analysis, an event is defined

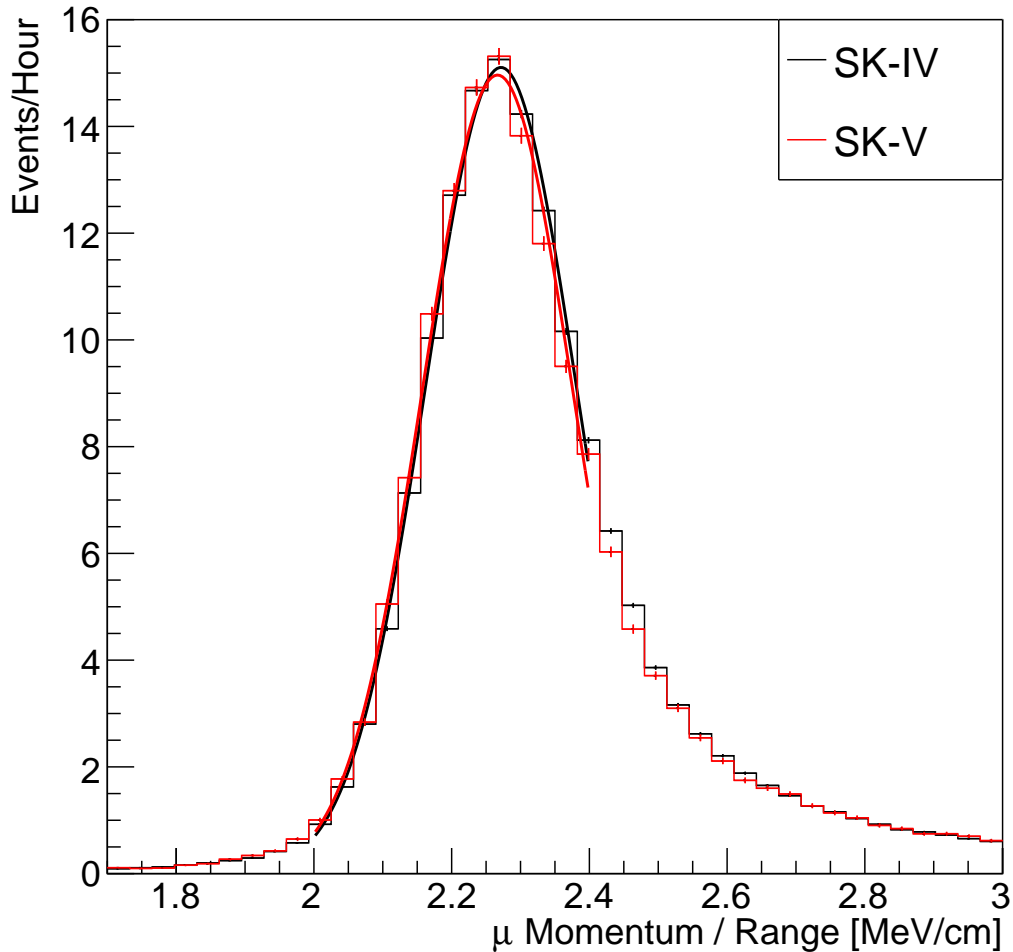


Figure 5.8: The distribution of the reconstructed momentum from the muon ring divided by the distance between the reconstructed muon and decay electron vertices as found in the stopping muon data sets of SK-IV (Black) and SK-IV (Red). Only events with one tagged decay electron are considered. A Gaussian fit is considered in the range [2.0, 2.4] MeV/cm and illustrated as the solid curve.

to be in the fiducial volume provided the event vertex is at least 0.5m away from the ID walls. PC events have the same ID requirements but can have a larger signal present inside the OD. Typically, only high energy muons from ν_μ interactions can penetrate the ID wall. The Up- μ sample contains events where muons are created from neutrino interactions in the OD water or rock below the tank. They then propagate upwards through the detector. Downward-going muons generated from neutrino interactions above the tank are neglected.

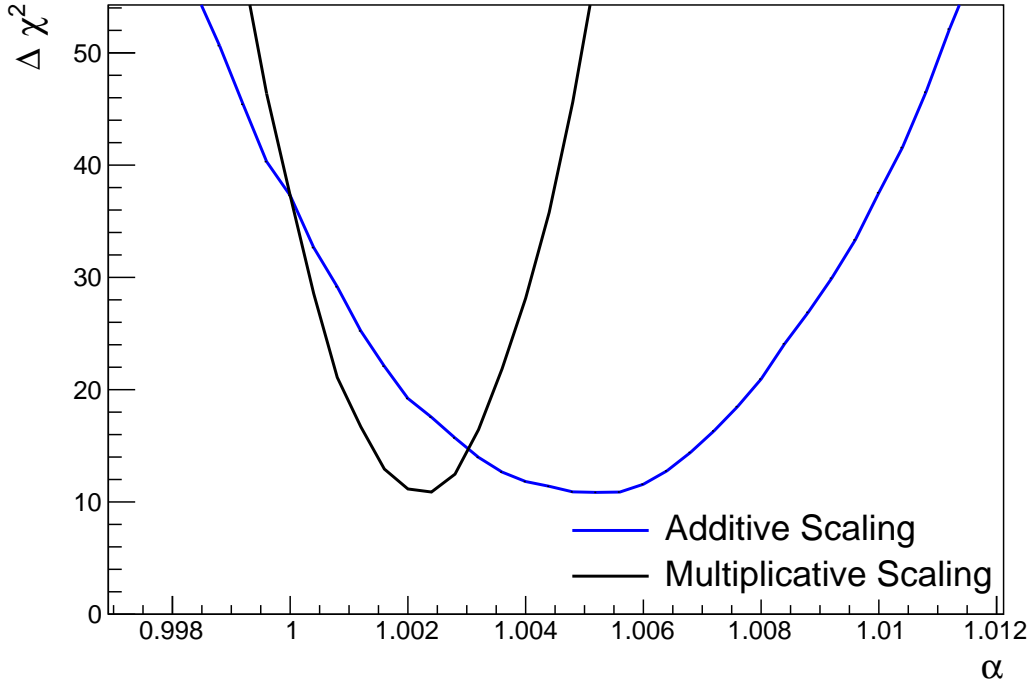


Figure 5.9: The χ^2 difference between the SK-IV and SK-V reconstructed muon momentum divided by range when the SK-V is modified by the scaling parameter α . Both additive (Blue) and multiplicative (Black) scaling factors have been considered. In practice, the additive scaling factor actually uses the value of $(\alpha - 1.0)$ but is illustrated like this so the results can be shown on the same axis range.

because of the difficulty in separating their signature from the cosmic muon shower background. The sample categories are visually depicted in Figure 5.10.

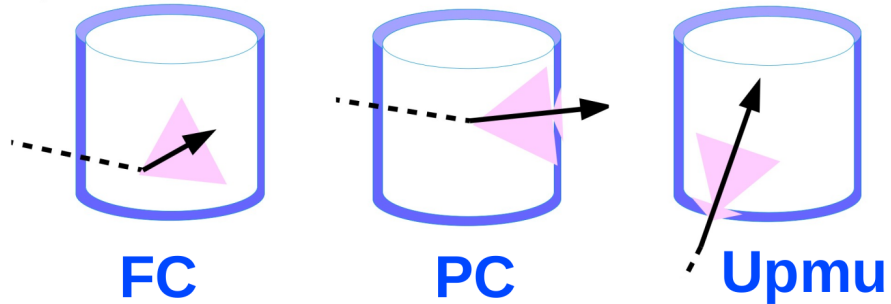


Figure 5.10: A depiction of the topology patterns for fully-contained (FC), partially-contained (PC) and up-going muon ($Up\mu$) samples included in this analysis.

Based on the event characteristics, as defined by the `fitQun` event reconstruction software, the FC events are categorised by

- **Visible Energy:** equal to the sum of the reconstructed kinetic energy of particles above the Cerenkov threshold for all rings present in the event. The purpose is to separate events into sub-GeV and multi-GeV categories.
- **Number of observed Cerenkov rings.** The purpose is to separate single-ring and multi-ring events, where single-ring events predominantly consist of quasi-elastic interactions and multi-ring events are typically resonant pion production or deep inelastic scattering events.
- **Particle identification parameter of the most energetic ring:** A value determined from the maximum likelihood value based on `fitQun`'s electron, muon, or pion hypothesis. The purpose is to separate electron-like and muon-like events.
- **Number of decay electrons:** The purpose is to separate quasi-elastic events (which have one decay electron emitted from the muon decay) and resonant pion production events (which have two decay electrons emitted from the muon and pion).

The PC and Up- μ categories are broken down into “through-going” and “stopping” samples depending on whether the muon leaves the detector. This is because the PC stopping events deposit the entire energy of the interaction into the detector, resulting in better reconstruction. The energy of events that exit the detector has to be estimated, with typically worse resolution, which introduces much larger systematic uncertainties. Through-going Up- μ samples are further broken down by whether any hadronic showering was observed in the event which typically indicates DIS interactions. The expected neutrino energy for the different categories is given in Figure 5.11. FC sub-GeV and multi-GeV events peak around 0.7GeV and 3GeV respectively, with slightly different peak energies for ν_e and $n\nu_\mu$ oscillation channels. PC and Up- μ are almost entirely comprised of ν_μ events and peak around 7GeV and 100GeV, respectively.

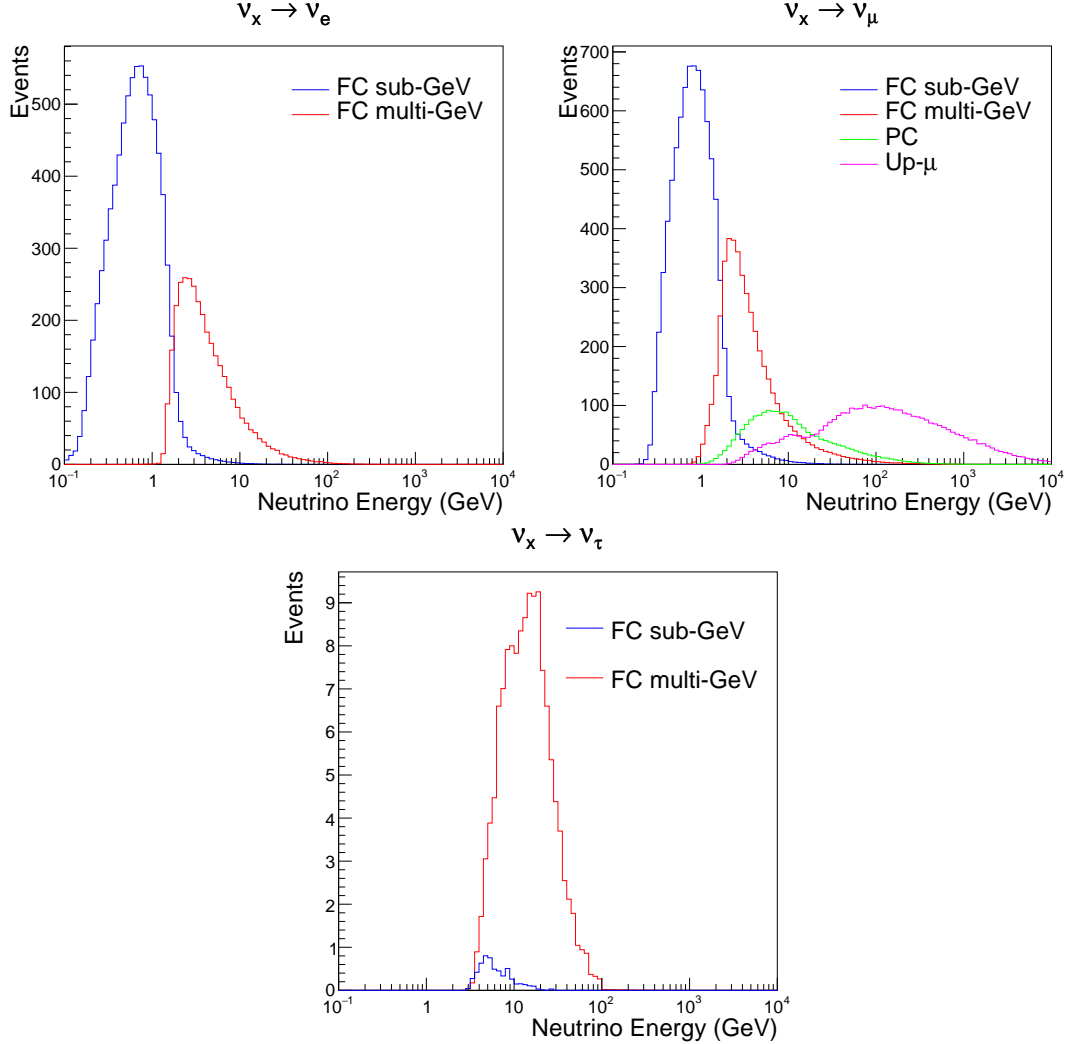


Figure 5.11: The predicted neutrino flux of the fully contained (FC) sub-GeV and multi-GeV, partially contained (PC), and upward-going muon (Up- μ) events. The prediction is broken down by the $\nu_x \rightarrow \nu_e$ prediction (top left), $\nu_x \rightarrow \nu_\mu$ prediction (top right) and $\nu_x \rightarrow \nu_\tau$ prediction (bottom). Asimov A oscillation parameters are assumed (given in Table 2.2).

In normal data-taking operations, the SK detector observes many background events alongside the beam and atmospheric neutrino signal events of physics interest for this thesis. Cosmic ray muons and flasher events, which are the spontaneous discharge of a given PMT, contribute the largest amount of background events in the energy range relevant to this thesis. Lower energy analyses like DSNB searches are also subject to radioactive backgrounds [187]. Therefore the data recorded is reduced with the aim of removing these background events. The reduction process is detailed in [46, 86] and briefly summarised below.

The first two steps in the FC reconstruction remove the majority of cosmic ray muons by requiring a significant amount of ID activity compared to that measured in the OD. Events that pass this cut are typically very high momentum muons or events that leave very little activity in the OD. Consequently, a third reduction step is then applied to select cosmic-ray muons that pass the initial reduction step. A purpose-built cosmic muon fitter is used to determine the entrance (or exit) position of the muon and a cut is applied to OD activity contained within 8m of this position. Flasher events are removed in the fourth reduction step which is based on the close proximity of PMT hits surrounding the PMT producing the flash. Events that pass all these reduction steps are reconstructed with the APFit algorithm. The fifth step of the reduction uses information from the more precise fitter to repeat the previous two steps with tighter cuts. Muons below the Cherenkov threshold can not generate optical photons in the ID but the associated decay electron can due to its lower mass. These are the types of events targeted in the fifth reduction step. The final cuts require the event vertex to be within the fiducial volume (0.5m from the wall although the nominal distance is 2.0m), visible energy $E_{vis} > 30\text{MeV}$ and fewer than 16 hits within the higher energy OD cluster. The culmination of the fully contained reduction results in 8.09 events/day in the nominal fiducial volume [83]. The uncertainty in the reconstruction is calculated by comparing Monte Carlo prediction to data. The largest discrepancy is found to be 1.3% in the fourth reduction step.

The PC and Up- μ events are processed through their own reduction processes detailed in [46]. Both of these samples are reconstructed with the APFit algorithm rather than `f1TQun`. This is because the efficiency of reconstructing events that leave the detector has not been sufficiently studied for reliable systematic uncertainties wth `f1TQun`. The PC and Up- μ samples acquire events at approximately 0.66 and 1.44 events/day.

Events due to beam neutrinos undergo the same reduction steps as FC events and are then subject to further cuts [188]. The GPS system which links the timing

between the beam facility and SK needs to be operating correctly and there should be no activity within the detector in the previous $100\mu\text{s}$ before the trigger. The events then need to triggered between $-2\mu\text{s}$ and $10\mu\text{s}$ of the expected spill timing.

The beam neutrino samples are not split by visible energy since their energy range is smaller than the atmospheric neutrino events. Following the T2K analysis in [76], only single-ring beam neutrino events are considered. Similar to atmospheric event selection, the number of decay electrons is used as a proxy for distinguishing CCQE and CCRES events. The expected neutrino energy, broken down by number of decay electrons, is given in Figure 5.12.

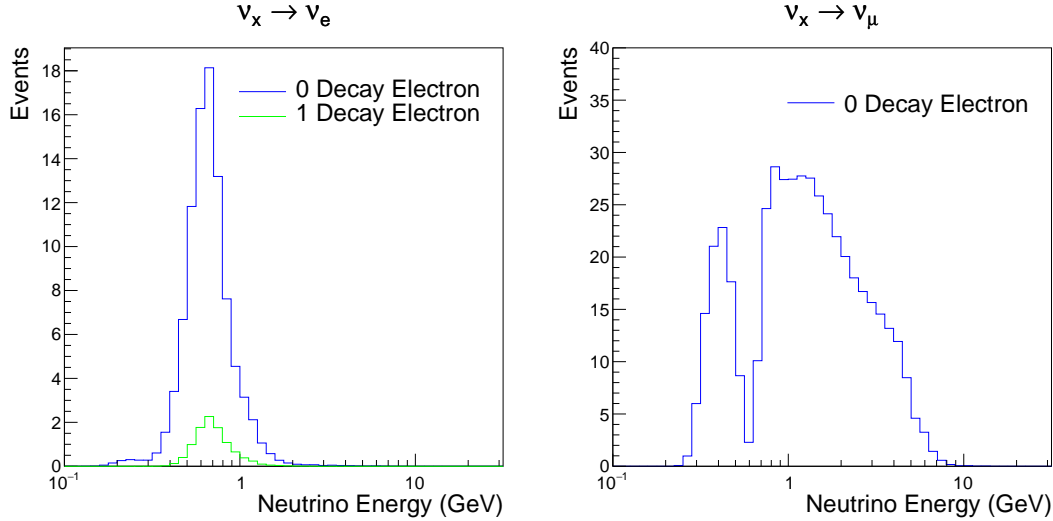


Figure 5.12: The predicted flux of beam neutrinos, as a function of neutrino energy. The predictions are broken down by the number of decay electrons associated with the particular events. Asimov A oscillation parameters are assumed (given in Table 2.2).

6

Sample Selections and Systematics

The oscillation analysis presented within this thesis is built upon a simultaneous fit to atmospheric data at SK, neutrino beam data in the near detector, and beam data measured at SK. The definitions of these samples are documented in section 6.1, section 6.2, and section 6.3, respectively. The data collected and used within this analysis is detailed in Table 6.1. The near and far detector data corresponds to T2K runs 2-9 and runs 1-10, respectively. The accumulated POT and beam power for runs 1 – 10 are illustrated in Figure 6.1.

Data Type	Total
Near Detector FHC	1.15×10^{21} POT
Near Detector RHC	8.34×10^{20} POT
Far Detector FHC	1.97×10^{21} POT
Far Detector RHC	1.63×10^{21} POT
Atmospheric SK-IV	3244.4 days

Table 6.1: The amount of data collected in each detector used within this analysis. The data collected at the near and far detector, for both neutrino beam (FHC) and antineutrino beam (RHC), is measured as the number of protons on target (POT).

The difference in POT recorded at the near and far detector is due to the difference in downtime. The SK detector is very stable with almost 100% of data recorded during beam operation. Due to various technical and operational

issues, the downtime of the near detector is significantly higher due to its more complex design and operating requirements.

The systematic parameters invoked within the flux, detector, and interaction models used within this analysis are documented in section 6.4. The standard configuration of the joint beam and atmospheric data fit utilises far detector systematics provided in the official inputs from the two experiments. Additionally, a correlated detector model which fits the parameters used in sample selections to data has been developed and documented in subsection 6.4.5.

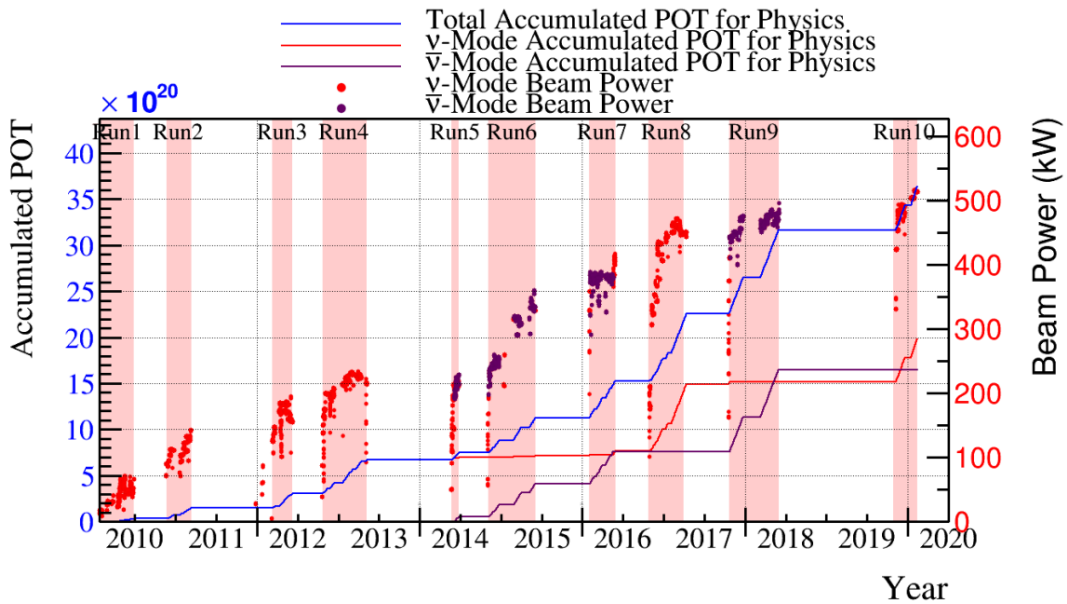


Figure 6.1: The accumulated beam data, measured as the number of protons on target (POT). The total data (blue) is given which comprises of the neutrino beam (red) and antineutrino (purple) components. The beam power for neutrino and antineutrino beams is given as the markers using the same colour scheme. The timescale runs from Run 1 which started in January 2010 until Run 10 which ended in February 2020. The ratio of accumulated data in neutrino and antineutrino beam is 54.7% : 45.3%.

6.1 Atmospheric Samples

The atmospheric event selection follows the official SK-IV analysis presented in [86] and is documented below. The Monte Carlo prediction used within this analysis corresponds to 500 years worth of neutrino events, which is scaled down to match the SK-IV livetime of 3244.4 days.

The fully contained (FC), partially contained (PC), and upward going muon events ($\text{up-}\mu$) which pass the reduction cuts discussed in section 5.3 are further broken down into different samples based on reconstruction information. This section details the samples used within this oscillation analysis, alongside the chosen binning.

FC events are first separated by the visible energy deposited within the detector. This is calculated as the sum of the reconstructed kinetic energy above the Cherenkov threshold for all rings present in the event. Events are separated by whether they were above or below $E_{\text{vis}} = 1.33\text{GeV}$. This separates “subGeV” and “multiGeV” events. Typically, lower energy events consist of charged current quasi-elastic (CCQE) interactions which are better understood and simpler to reconstruct resulting in smaller systematic uncertainties. Events are further separated by the number of rings associated with the event due to similar reasoning. As the oscillation probability is dependent upon the flavour of neutrino, electron and muon events are separated using a similar likelihood method to that discussed in section 5.2. To reduce computational resources required for the reconstruction, only electron and pion hypotheses are considered so this separation cut depends on the ratio of the electron to pion likelihoods, $\log(L_e/L_\pi)$. Finally, the number of decay electrons is used to classify events. Charged current resonant pion production (CCRES) interactions generate a final-state pion. This can decay, mostly likely through a muon, into a decay electron. Therefore any electron-like event with one decay electron or muon-like event with two decay electrons was most likely produced by a CCRES interaction. Consequently, the number of decay electrons can be used to distinguish CCQE

and CCRES interaction modes. Ultimately, FC subGeV events are separated into the samples listed in Table 6.2.

Sample Name	Description
SubGeV- <i>e</i> like-0dcy	Single ring <i>e</i> -like events with zero decay electrons
SubGeV- <i>e</i> like-1dcy	Single ring <i>e</i> -like events with one or more decay electrons
SubGeV- <i>μ</i> like-0dcy	Single ring <i>μ</i> -like events with zero decay electrons
SubGeV- <i>μ</i> like-1dcy	Single ring <i>μ</i> -like events with one decay electrons
SubGeV- <i>μ</i> like-2dcy	Single ring <i>μ</i> -like events with two or more decay electrons
SubGeV- <i>π</i> 0like	Two <i>e</i> -like ring events with zero decay electrons and reconstructed π^0 mass $85 \leq m_{\pi^0} < 215$ MeV

Table 6.2: The fully contained subGeV samples, defined as events with visible energy $E_{vis} < 1.33$ GeV, used within this oscillation analysis.

In addition to the cuts discussed above, multiGeV samples also have additional cuts to separate samples which target neutrino and antineutrino events. As discussed in section 7.1, the matter resonance only occurs for neutrinos in normal hierarchy and antineutrinos in an inverted mass hierarchy. Therefore, having flavour-enriched samples aids in the determination of the mass hierarchy. For a CCRES interaction,

$$\begin{aligned}
 \bar{\nu}_e + N &\rightarrow e^+ + N' + \pi^-, \\
 \nu_e + N &\rightarrow e^- + N' + \pi^+ \\
 &\quad \downarrow \mu^+ + \nu_\mu \\
 &\quad \downarrow e^+ + \nu_e + \bar{\nu}_\mu.
 \end{aligned} \tag{6.1}$$

The π^- emitted from a $\bar{\nu}_e$ interaction is more likely to be absorbed within the oxygen nucleus than the π^+ from ν_e interactions [189]. These pions then decay, mostly through muons, to electrons. Therefore the number of tagged decay electrons associated with an event gives an indication of whether the interaction was due to a neutrino or antineutrino: zero for $\bar{\nu}_e$ events, and one for ν_e events. The ability to separate neutrino from antineutrino events is illustrated in Table 6.4, where the MultiGeV-*e*like-nue has 78% purity of CC neutrino interactions with only 7% antineutrino background, the rest consisting of NC backgrounds.

The number of decay electrons discriminator works reasonably well for single-ring events. However, this is not the case for multi-ring events. A multiGeV multiring electron-like (MME) likelihood cut was introduced in [190, 191]. This is a two-stage likelihood selection cut. Four observables are used in the first likelihood cut to distinguish $CC\nu_e$ and $CC\bar{\nu}_e$ events from background:

- The number of decay electrons
- The maximum distance between the vertex of the neutrino and the decay electrons
- The energy deposited by the highest energy ring
- The particle identification of that highest energy ring

Background events consist of $CC\nu_\mu$ and NC interactions. Typically, the majority of the energy in these background events is carried by the hadronic system. Additionally, muons tend to travel further than the pions from $CC\nu_e$ before decaying. Thus, the parameters used within the likelihood cut target these typical background interaction kinematics.

Sample Name	Description
MultiGeV-elike-nue	Single ring e -like events with zero decay electrons
MultiGeV-elike-nuebar	Single ring e -like events with one or more decay electrons
MultiGeV-mulike	Single ring μ -like events
MultiRing-elike-nue	Two or more ring events with leading energy e -like ring and passed both MME and $\nu/\bar{\nu}$ separation cuts
MultiRing-elike-nuebar	Two or more ring events with leading energy e -like ring and passed MME and failed $\nu/\bar{\nu}$ separation cuts
MultiRing-mulike	Two or more ring events with leading energy μ -like ring and only requires $E_{vis} > 0.6\text{GeV}$ DB: Why is this not }1.33\text{GeV}
MultiRing-Other1	Two or more ring events with leading energy e -like ring and failed the MME likelihood cut

Table 6.3: The fully contained multiGeV samples used within this oscillation analysis. Both the sample name and description are given.

Neutrino and antineutrino events are then separated by a second likelihood method ($\nu/\bar{\nu}$ separation) detailed in [51]. This uses the number of decay electrons,

the number of reconstructed rings, and the event’s transverse momentum. The last two parameters are used because higher-energy samples tend to have more pions produced above the Cherenkov threshold which results in more rings compared to an antineutrino interaction. Furthermore, the angular distribution also tends to be more forward peaked in antineutrino interactions as compared to neutrino interactions [86]. These FC multiGeV sample definitions are detailed in Table 6.3.

The PC and up- μ samples are split by the amount of energy deposited within the outer detector, into “stopping” and “through-going” samples. If an event leaves the detector, the energy it takes with it has to be estimated which increases the systematic uncertainty compared to events entirely contained within the inner detector. This estimation is particularly poor at high energies, thus the up- μ through-going events are not binned in reconstructed momentum. The through-going up- μ are further separated by the presence of any electromagnetic showering in the event, as the assumption of non-showering muon does not give reliable reconstruction for these types of events [46]. In total, 13 FC, 2 PC, and 3 up- μ atmospheric samples are included within this analysis.

The atmospheric samples are binned in direct observables: reconstructed lepton momentum and direction, as given by Table A.1. The distribution of the reconstructed lepton momentum (for samples that only have one bin in reconstructed zenith angle) and reconstructed direction for each atmospheric sample used within this analysis is illustrated in Figure 6.2. The by-mode breakdown of each of the atmospheric samples is given in Appendix A.

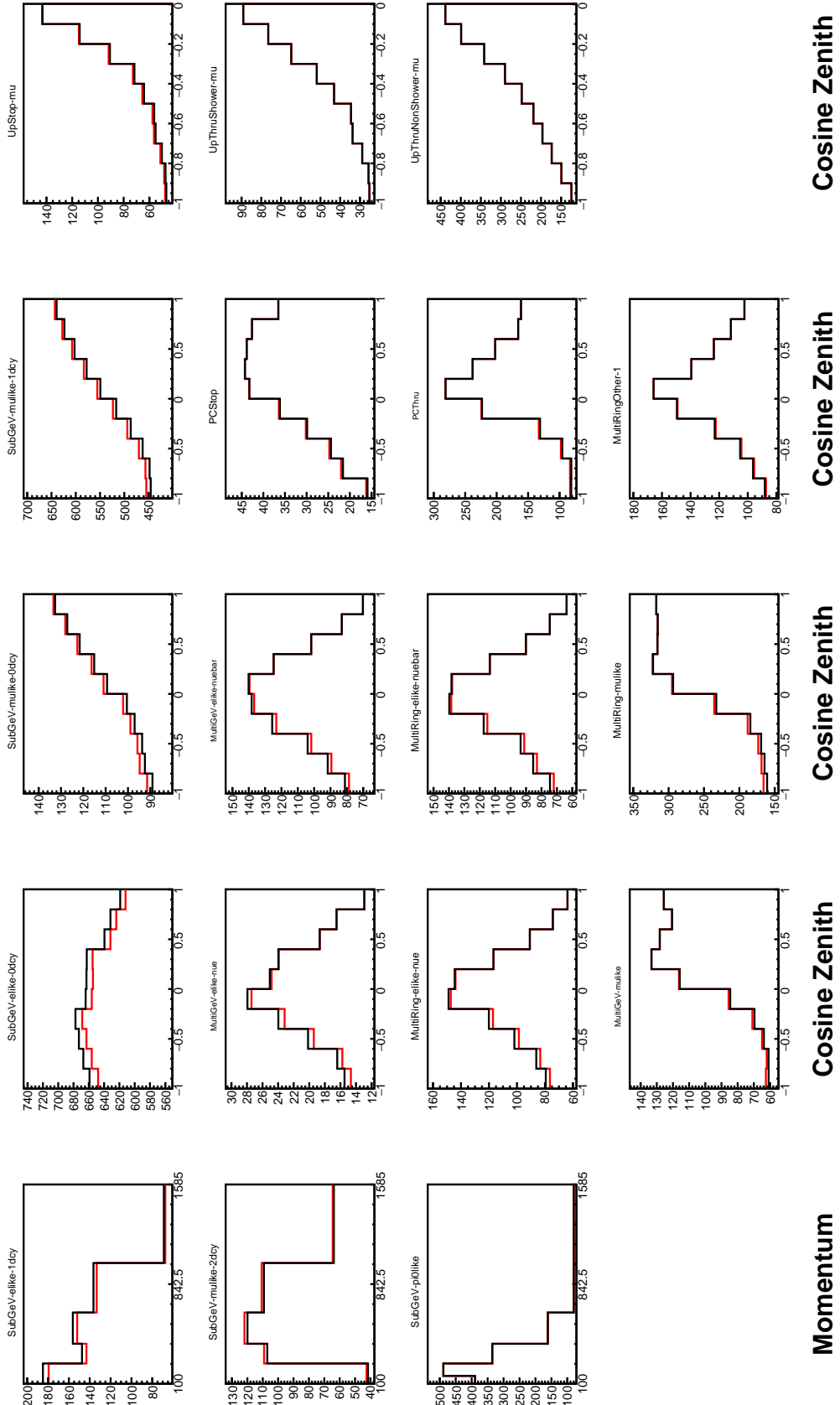


Figure 6.2: Comparison of the SK-IV atmospheric samples between predictions made with the CP-violating Asimov A (Black) and CP-conserving Asimov B (Red) oscillation parameter sets (given in Table 2.2). The subGeV samples CCRES and π^0 -like samples are given in their reconstructed lepton momentum. All other samples are presented in their reconstructed zenith angle projection.

Sample	$CC\nu_e$	$CC\bar{\nu}_e$	$CC(\nu_\mu + \bar{\nu}_\mu)$	$CC(\nu_\tau + \bar{\nu}_\tau)$	NC
SubGeV- <i>elike</i> -0dcy	72.17	23.3	0.724	0.033	3.77
SubGeV- <i>elike</i> -1dcy	86.81	1.773	7.002	0.062	4.351
SubGeV- <i>mulike</i> -0dcy	1.003	0.380	90.07	0.036	8.511
SubGeV- <i>mulike</i> -1dcy	0.023	0.	98.46	0.029	1.484
SubGeV- <i>mulike</i> -2dcy	0.012	0.	99.25	0.030	0.711
SubGeV- <i>pi0like</i>	6.923	2.368	0.928	0.011	89.77
MultiGeV- <i>elike</i> -nue	78.18	7.041	3.439	1.886	9.451
MultiGeV- <i>elike</i> -nuebar	56.68	37.81	0.174	0.614	4.718
MultiGeV- <i>mulike</i>	0.024	0.005	99.67	0.245	0.058
MultiRing- <i>elike</i> -nue	59.32	12.39	4.906	3.385	20
MultiRing- <i>elike</i> -nuebar	52.39	31.03	1.854	1.585	13.14
MultiRing- <i>mulike</i>	0.673	0.080	97.33	0.342	1.578
MultiRingOther-1	27.98	2.366	34.93	4.946	29.78
PCStop	8.216	3.118	84.45	0.	4.214
PCThrus	0.564	0.207	98.65	0.	0.576
UpStop-mu	0.829	0.370	98.51	0.	0.289
UpThruNonShower-mu	0.206	0.073	99.62	0.	0.103
UpThruShower-mu	0.128	0.054	99.69	0.	0.132

Table 6.4: The purity of each atmospheric sample used within this analysis, broken down by charged current (CC) and neutral current (NC) interactions and which neutrino flavour interacted within the detector. Asimov A oscillation parameter sets are assumed (given in Table 2.2). Electron neutrino and antineutrino events are separated to illustrate the ability of the separation likelihood cuts used within the multiGeV and multiring sample selections.

6.2 Near Detector Beam Samples

The near detector sample selections are documented in detail within [192] and summarised below. Samples are selected based upon the which Fine Grained Detector (FGD) that the vertex is reconstructed in as well as the operating mode of the beam: FHC or RHC. Wrong-sign neutrino samples are considered in the RHC mode in order to add additional constraints on model parameters. Samples from the wrong-sign component of the FHC beam mode are not included as they are statistically insignificant compared to those samples already listed.

For additional constraints on model parameters, wrong-sign neutrino samples are also considered when the beam is operating in RHC mode.

The reconstruction algorithm uses a clustering algorithm to group hits within the TPC. It then adds information from the upstream FGD to form a track which passes through both sub-detectors. In FHC(RHC), the highest momentum negative(positive) curvature track is defined as the muon candidate. Before being assigned a sample, these candidate muon events must pass CC-inclusive cuts, as defined in [193]:

- Event Timing: The DAQ must be operational and the event must occur within the expected beam time window consistent with the beam spill
- TPC Requirement: The muon-candidate track path must intercept one or more TPCs
- Fiducial volume: The event must originate from within the fiducial volume defined in [194].
- Upstream Background: Remove events that have muon tracks that originate upstream of the FGDs by requiring no high-momentum tracks within 150mm upstream of the candidate vertex. Additionally, events that occur within the downstream FGD are vetoed if a secondary track starts within the upstream FGD

- Broken track removal: All candidates where the muon candidate is broken in two are removed
- Muon PID: Measurements of dE/dx in a TPC are used to distinguish muon-like events, from electron-like or proton-like, using a likelihood cut

In addition to these cuts, RHC neutrino events also have to undergo the following cuts to aid in the separation of neutrino and antineutrino [195]:

- TPC Requirement: The track path must intercept TPC2
- Positive Track: The highest momentum track must have positive reconstructed charge
- TPC1 Veto: Remove any events originating upstream of TPC1

Once all CC-inclusive events have been determined, they are further split by pion multiplicity: CC0 π , CC1 π , and CCOther. These target specific interaction modes CCQE, CCRES, and other CC background interactions, respectively. Pions in the TPCs and FGDs are selected by requiring a second track to be observed, which is separate from the muon track and is in the same beam spill window and sub-detector. If the pion originated within a FGD, it must also pass through the sequential downstream TPC (TPC2 for FGD1, TPC3 for FGD2).

CC0 π , CC1 π , and CCOther samples are defined with the following cuts:

DB: Understand pion cuts at ND

- ν_μ CC0 π Selection: No electrons in TPC and no charged pions or decay electrons within the TPC or FGD
- ν_μ CC1 π Selection: Exactly one charged pion in either the TPC or FGD, where the number of charged pions in the FGD is equal to the number of decay electrons
- ν_μ CCOther Selection: All events which are not classified into the above two selections.

Counting the three selections for each FGD in FHC and RHC running, including the wrong-sign background in RHC, 18 near detector samples are used within this analysis. These samples are binned in reconstructed lepton momentum (illustrated in Figure 6.3) and direction with respect to the beam. The binning is chosen such that each event has at least 20 Monte Carlo events in each bin [194]. This is to ensure that the bins are coarse enough to ensure the reduction of statistical errors, whilst also being fine enough to sample the high-resolution peak regions. The exact binning is detailed in [194].

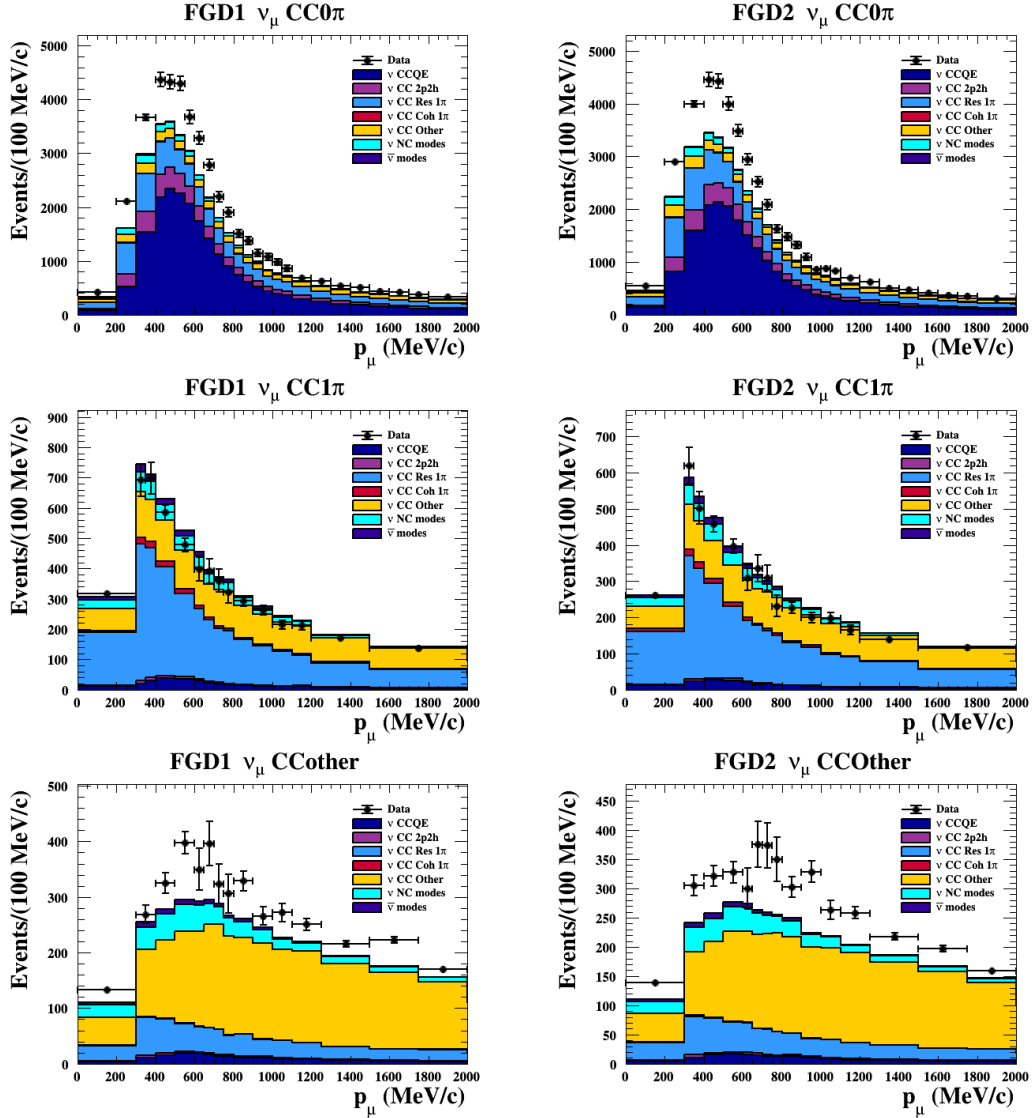


Figure 6.3: The nominal Monte Carlo predictions compared to data for the FGD1 and FGD2 samples in neutrino beam mode, broken down into the $CC\nu_\mu 0\pi$, $CC\nu_\mu 1\pi$ and $CC\nu_\mu$ Other categories. Figures taken from [192].

6.3 Far Detector Beam Samples

The beam neutrino events which occur at the SK detector, which pass the reduction cuts detailed in section 5.3, are separated based on whether the beam was operating in FHC or RHC mode. The events are then separated into three samples: electron-like ($1Re$), muon-like ($1R\mu$), and $CC1\pi^+$ -like ($1Re1de$) which are observed as electron-like events with an associated decay electron [183]. As discussed in section 6.1, positively charged pions emitted from neutrino interactions are more likely to produce decay electrons than negatively charged pions. Consequently, the $CC1\pi^+$ -like sample is only selected when the beam is operating in FHC mode. Therefore, five beam samples measured at SK are used in this analysis.

The fiducial volume definition for beam samples is slightly different from that used for the atmospheric samples. It uses both the distance to the closest wall ($dWall$) and the distance to the wall along the trajectory of the particle ($toWall$). This allows events that originate close to the wall but are facing into the tank to be included within the analysis, which would have otherwise been removed. These additional events are beneficial for a statistics-limited experiment. The exact cut values for both $dWall$ and $toWall$ are different for each of the three types of sample and are optimised based on T2K sensitivity to δ_{CP} [181, 196]. They are:

1Re event selection For an event to be classified as a $1Re$ -like, the event must satisfy:

- Fully-contained and have $dWall > 80\text{cm}$ and $toWall > 170\text{cm}$
- Total of one ring which is reconstructed as electron-like with reconstructed momentum $P_e > 100\text{MeV}$
- Zero decay electrons are associated with the event
- Passes π^0 rejection cut discussed in section 5.2

The zero decay electron cut removes non-CCQE interactions and the π^0 rejection cut is designed to remove neutral current π^0 background events which can be easily reconstructed as 1Re-like events.

The zero decay electron cut removes non-CCQE interactions and the π^0 rejection cut is designed to remove neutral current π^0 background events which can be easily reconstructed as 1Re-like events.

CC1 π^+ event selection This event selection is very similar to that of the 1Re sample. The only differences are that the `dWall` and `toWall` criteria are changed to $> 50\text{cm}$ and $> 270\text{cm}$, respectively, and exactly one decay electron is required from the π^+ decay.

1R μ event selection A 1R μ -like event is determined by the following cuts:

- Fully-contained and have `dWall` $> 50\text{cm}$ and `toWall` $> 250\text{cm}$
- Total of one ring which is reconstructed as muon-like with reconstructed momentum $P_\mu > 200\text{MeV}$
- Fewer than two decay electrons are associated with the event
- Passes π^+ rejection cut discussed in section 5.2

All of these samples are binned in reconstructed neutrino energy. This is possible under a particular interaction mode assumption, as the direction from the source is known extremely well. For the 1Re-like and 1R μ -like samples,

$$E_\nu^{rec} = \frac{(M_N - V_{nuc})E_l - m_l^2/2 + M_N V_{nuc} - V_{nuc}^2/2 + (M_P^2 + M_N^2)/2}{M_N - V_{nuc} - E_l + P_l \cos(\theta_{beam})} \quad (6.2)$$

Where M_N , M_P and m_l are the masses of the neutron, proton and outgoing lepton, respectively. $V_{nuc} = 27\text{MeV}$ is the binding energy of the oxygen nucleus [183], θ_{beam} is the angle between the beam and the direction of the outgoing lepton, and E_l and P_l are the energy and momentum of that outgoing lepton.

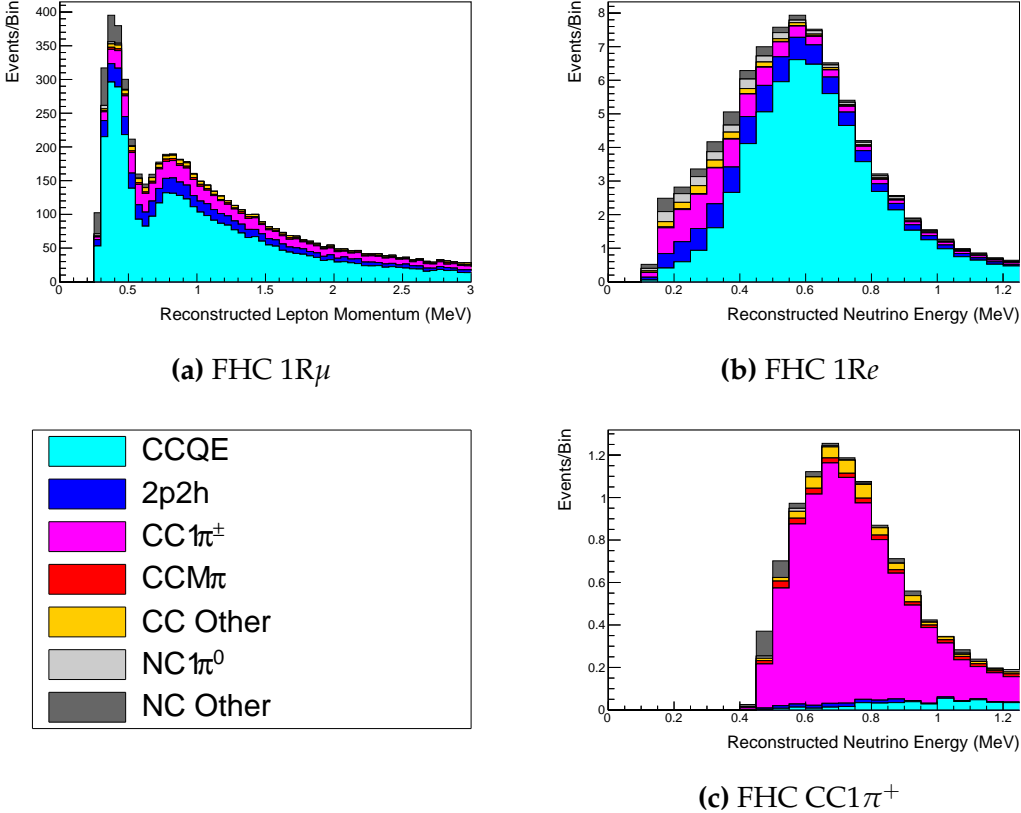


Figure 6.4: The reconstructed neutrino energy, as defined by Equation 6.2 and Equation 6.3, for the 1R μ -like, 1Re-like and CC1 π^+ -like samples. Asimov A oscillation parameter sets are assumed (given in Table 2.2). These samples are the FHC mode samples. For ease of viewing, the 1R μ sample only shows the $0 \leq E_\nu^{rec} < 3.0\text{GeV}$ but the binning extends to 30.0GeV.

The reconstructed neutrino energy of the CC1 π^+ -like events also accounts for the delta resonance produced within the interaction,

$$E_\nu^{rec} = \frac{2M_N E_l + M_{\Delta^{++}}^2 - M_N^2 - m_l^2}{2(M_N - E_l + P_l \cos(\theta_{beam}))} \quad (6.3)$$

Where $M_{\Delta^{++}}$ is the mass of the delta baryon. Binding energy effects are not considered as a two-body process with the delta baryon is assumed. This follows the T2K oscillation analysis presented in [76], although recent developments of the interaction model in the latest T2K oscillation analysis do include effects from binding energy in this calculation [197].

The reconstructed neutrino energy for the FHC samples is illustrated in Figure 6.4. As expected, the 1R μ -like and 1Re-like samples are heavily dominated

by CCQE interactions, with smaller contributions from 2p2h meson exchange and resonant pion production interactions. The CC1 π^+ -like sample predominantly consists of charged current resonant pion production interactions. The 1Re-like and CC1 π^+ -like samples are also binned by the angle between the neutrino beam and the reconstructed lepton momentum. This is to aid in charged current and neutral current separation, as indicated in Figure 6.5. This is because the neutral current backgrounds are predominantly due to π^0 -decays, where the opening angle of the two gammas alongside the different final state kinematics produces a slightly broader angular distribution compared to the final state particles originating from charged current ν_e interactions.

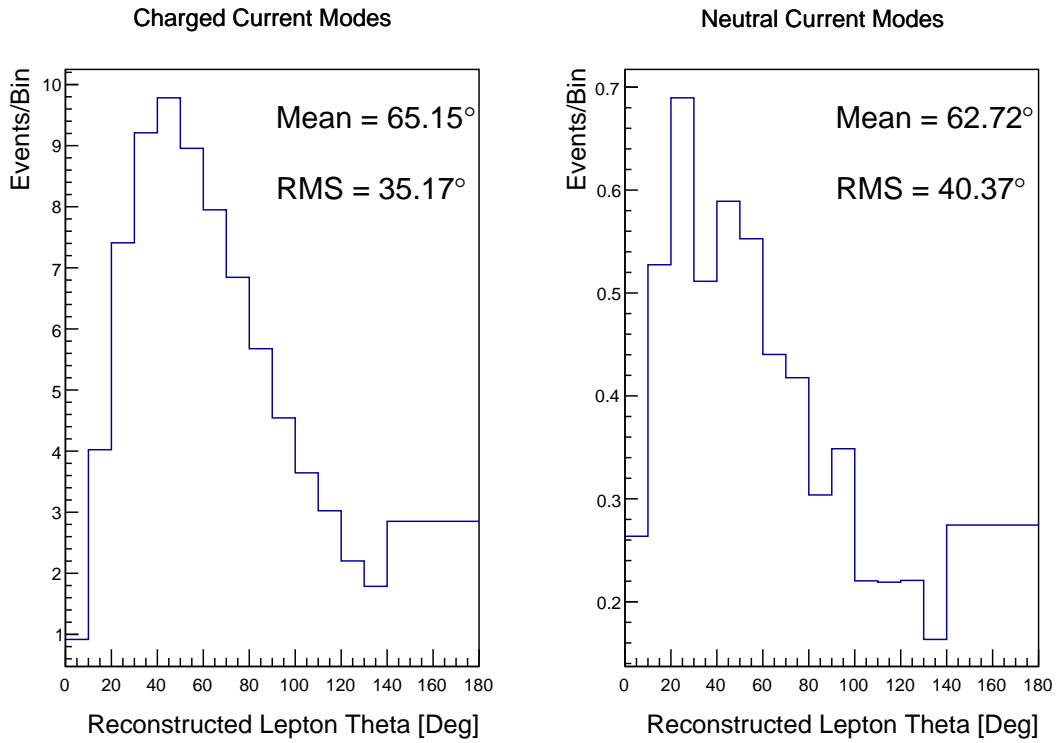


Figure 6.5: The distribution of the angle between the neutrino beam direction and the reconstructed final state lepton, for the FHC 1Re-like sample. The distribution is broken down by neutrino interaction mode into charged current (left) and neutral current (right) components. Asimov A oscillation parameter sets are assumed (given in Table 2.2). The RMS of the charged and neutral current plots are 35.17° and 40.37° , respectively.

6.4 Systematic Uncertainties

The systematic model parameters for this analysis are split into groups, or blocks, depending on their purpose. They consist of flux uncertainties, neutrino-matter interaction systematics, and detector efficiencies. There are also uncertainties on the oscillation parameters which this analysis is not sensitive to, namely Δm_{12}^2 and $\sin^2(\theta_{12})$. These oscillation parameters uncertainties are taken from the 2020 PDG measurements [77]. As described in chapter 4, each model parameter used within this analysis requires a prior uncertainty. This is provided via separate covariance matrices for each block. The covariance matrices can include prior correlations between parameters within a single block, but the separate treatment means prior correlations can not be included for parameters in different groups. Some parameters in these models have no reasonably motivated uncertainties and are assigned flat priors which do not modify the likelihood penalty. In practice, these flat prior parameters are actually assigned a Gaussian with a very large width to ensure the covariance matrix is positive definite. They are then checked at run time to determine if they contribute to the likelihood. The flux, neutrino interaction, and detector modeling simulations have already been discussed in section 5.1 and section 5.2. The uncertainties invoked within each of these models are described below.

6.4.1 Beam Flux

The neutrino beam flux systematics are based upon the uncertainty in the modeling of the components of the beam simulation. This includes the model of hadron productions and reinteractions, the shape, intensity, and alignment of the beam with respect to the target, and the uniformity of the magnetic field produced by the horn, alongside other effects. The uncertainty, as a function of neutrino energy, is illustrated in Figure 6.6 which includes a depiction of the total uncertainty as well as the contribution from individual components. The uncertainty around the peak of the energy distribution ($E_\nu \sim 0.6\text{GeV}$) is

dominated by uncertainties in the beam profile and alignment. Outside of this region, uncertainties on hadron production dominate the error.

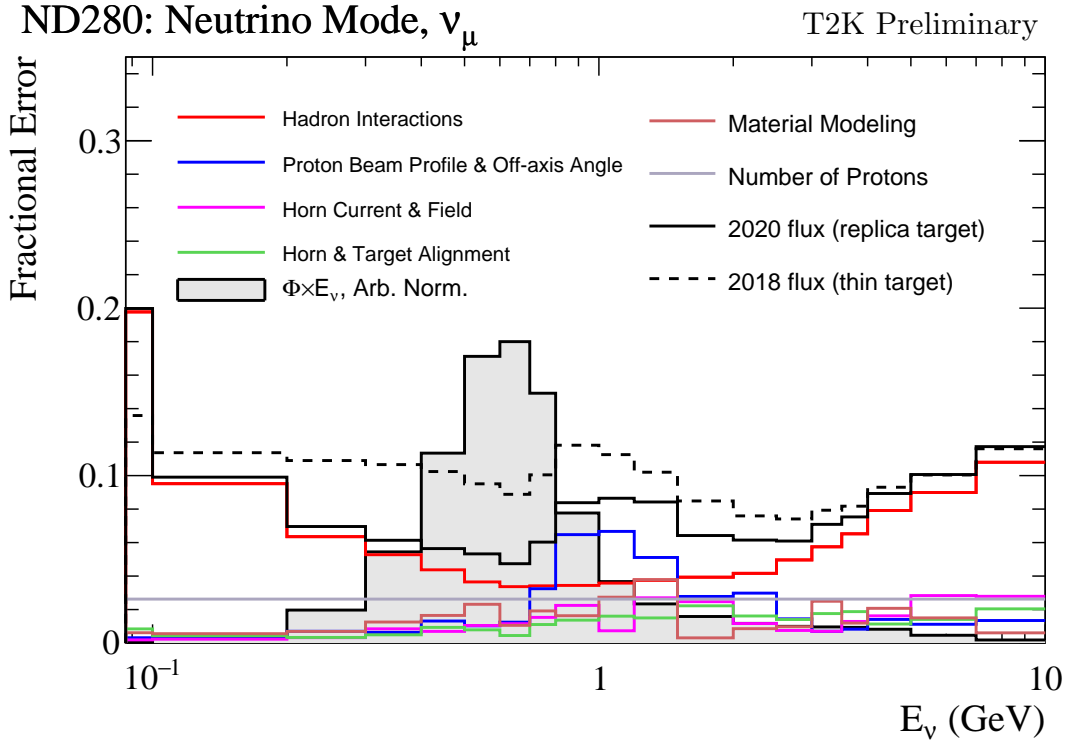


Figure 6.6: The total uncertainty evaluated on the near detector ν_μ flux prediction constrained by the replica-target data, illustrated as a function of neutrino energy. The solid(dashed) line indicates the uncertainty used within this analysis(the T2K 2018 analysis [198]). The solid histogram indicates the neutrino flux as a function of energy. Figure taken from [199].

The beam flux uncertainties are described by one hundred parameters. They are split between the ND280 and SK detectors and binned by neutrino flavour: ν_μ , $\bar{\nu}_\mu$, ν_e and $\bar{\nu}_e$. The response is then broken down as a function of neutrino energy. The bin density in the neutrino energy is the same for the ν_μ in FHC and $\bar{\nu}_\mu$ in RHC beams, and narrows for neutrino energies close to the oscillation maximum of $E_\nu = 0.6\text{GeV}$. This binning is specified in Table 6.5. All of these systematic uncertainties are applied as normalisation parameters with Gaussian priors centered at 1.0 and error specified from a covariance matrix provided by the T2K beam group [199].

Neutrino Flavour	Sign	Neutrino Energy Bin Edges (GeV)
μ	Right	0., 0.4, 0.5, 0.6, 0.7, 1., 1.5, 2.5, 3.5, 5., 7., 30.
μ	Wrong	0., 0.7, 1., 1.5, 2.5, 30.
e	Right	0., 0.5, 0.7, 0.8, 1.5, 2.5, 4., 30.
e	Wrong	0., 2.5, 30.

Table 6.5: The neutrino energy binning for the different neutrino flavours. “Right” sign indicates neutrinos in the FHC beam and antineutrinos in the RHC beam. “Wrong” sign indicates antineutrinos in the FHC beam and neutrinos in the RHC beam. The binning of the detector response is identical for the FHC and RHC modes as well as at ND280 and SK.

6.4.2 Atmospheric Flux

The atmospheric neutrino flux is modeled by the HKKM model [42]. 16 systematic uncertainties are applied to control the normalisation of each neutrino flavour, energy, and direction. They are summarised below:

- **Absolute Normalisation:** The overall normalisation of each neutrino flavour is controlled by two independent systematic uncertainties, for $E_\nu < 1\text{GeV}$ and $E_\nu > 1\text{GeV}$, respectively. This is driven mostly by hadronic interaction uncertainties for the production of pions and kaons [42]. The strength of the response is dependent upon the neutrino energy. The uncertainty is parameterized following Figure 11 in [42].
- **Relative Normalisation:** Uncertainties on the ratio of $(\nu_\mu + \bar{\nu}_\mu) / (\nu_e + \bar{\nu}_e)$ are controlled by the difference between the HKKM model [42], FLUKA [45] and Bartol models [41]. Three independent parameters are applied in the energy ranges: $E_\nu < 1\text{GeV}$, $1\text{GeV} < E_\nu < 10\text{GeV}$, and $E_\nu > 10\text{GeV}$.
- **$\nu/\bar{\nu}$ Normalisation:** The uncertainties in the π^+/π^- (and kaon equivalent) production uncertainties in the flux of $\nu/\bar{\nu}$. The response is applied using the same methodology as the relative normalisation parameters.
- **Up/Down and Vertical/Horizontal Ratio:** Similar to the above two systematics, the difference between the HKKM, FLUKA, and Bartol model

predictions, as a function of $\cos(\theta_Z)$, is used to control the normalisation of events as a function of zenith angle.

- **K/π Ratio:** Higher energy neutrinos ($E_\nu > 10\text{GeV}$) mostly originate in kaon decay. Measurements of the ratio of K/π production [200] are used to control the systematic uncertainty of the expected ratio of pion and kaon production.
- **Solar Activity:** As the 11-year solar cycle can affect the Earth's magnetic field, the flux of primary cosmic rays varies across the same period. The uncertainty is calculated by taking a ± 1 year variation, equating to a 10% uncertainty for the SK-IV period.
- **Atmospheric Density:** The height of the interaction of the primary cosmic rays is dependent upon the atmospheric density. The HKKM assumes the US standard 1976 [151] profile. This systematic controls the uncertainty in that model.

The total uncertainty is dominated by the absolute and relative normalisation parameters. The effect of which is illustrated in Figure 6.7. Generally, the uncertainty is large at low energy, reducing to $O(10\%)$ around the peak of the flux distribution and then increasing once the neutrino energy exceeds 10GeV.

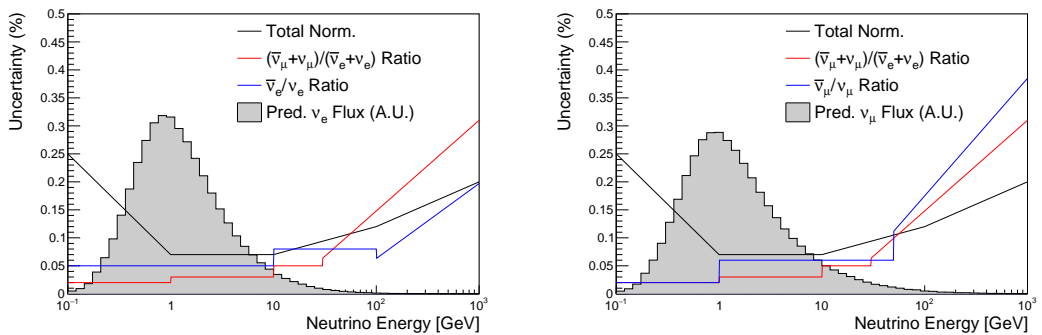


Figure 6.7: The uncertainty evaluated on the atmospheric ν_e (left) and ν_μ (right) flux predictions. The absolute normalisation and flavour ratio uncertainties are given. The solid histogram indicates the neutrino flux as a function of energy.

Updates to the HKKM and Bartol models are underway [156] to use a similar tuning technique to that used in the beam flux predictions. After those updates, it may be possible to include correlations in the hadron production uncertainty systematics for beam and atmospheric flux predictions.

6.4.3 Neutrino Interaction

Neutrino interactions in the detectors are modeled by NEUT. The two independent oscillation analyses, T2K-only [201] and the SK-only [51], have developed separate interaction models. To maximise sensitivity out of this simultaneous beam and atmospheric analysis, a correlated interaction model has been defined in [202]. Where applicable, correlations allow the systematic uncertainties applied to the atmospheric samples to be constrained by near detector neutrino beam measurements. This can lead to stronger sensitivity to oscillation parameters as compared to an uncorrelated model.

The low-energy T2K systematic model has a more sophisticated treatment of CCQE, 2p2h, and CCRES uncertainties, where extensive comparisons of this model have been performed to external data [201]. However, the model is not designed for high-energy atmospheric events, like those illustrated in Figure 5.11. Therefore the high energy systematic model from the SK-only analysis is implemented for the relevant multiGeV, PC, and up- μ samples. The T2K CCQE model is more sophisticated so it has been implemented for all samples within this analysis, where separate low-energy and high-energy dials have been implemented. The low-energy dials are constrained by the near detector measurements and are uncorrelated to their high-energy counterparts. The author of this thesis was responsible for implementing and validating the combined cross section model as documented in [202, 203].

The high energy systematic model includes parameters developed from comparisons of Nieves and Rein-Seghal models which affect resonant pion producing interactions, comparisons of the GRV98 and CKMT models which control DIS interactions, and hadron multiplicity measurements which modulate

the normalisation of multi-pion producing events. The uncertainty on the ν_τ cross-section is particularly large and is controlled by a 25% normalisation uncertainty. These uncertainties are applied via normalisation or shape parameters. The former linearly scales the weight of all affected Monte-Carlo events, whereas the latter can increase or decrease a particular event's weight depending on its neutrino energy and mode of interaction. The response of the shape parameters is defined by third-order polynomial splines which return a weight for a particular neutrino energy. To reduce computational resources for the far detector fit, the response is binned by neutrino energy and sample binning: lepton momentum and cosine zenith binning for atmospheric splined responses and reconstructed neutrino energy and direction binning for beam samples. In total, 17 normalisation and 15 shape parameters are included in the high-energy model within this analysis.

Figure 6.8 indicates the predicted neutrino energy distribution for both beam and subGeV atmospheric samples. There is clearly significant overlap in neutrino energy between the subGeV atmospheric and beam samples, allowing similar kinematics in the final state particles. Figure 6.9 illustrates the fractional contribution of the different interaction modes per sample.

Comparing beam and atmospheric samples which target CCQE interactions (S.G. e-like 0de, S.G. μ -like [0,1]de, [FHC,RHC] 1R μ -like and [FHC,RHC] 1R e-like samples), there is a very similar contribution of CCQE, CC 2p2h, and CC1 π^\pm interactions. The samples which target CC1 π^\pm interactions, (S.G. e-like 0de, S.G. μ -like 2de and FHC 1R+1d.e e-like) also consist of very similar mode interactions.

As a consequence of the similarity in energy and mode contributions, correlating the systematic model between the beam and subGeV atmospheric samples ensures that this analysis attains the largest sensitivity to oscillation parameters while still ensuring neutrino interaction systematics are correctly accounted for. Due to its more sophisticated CCQE and 2p2h model, the T2K systematic model was chosen as the basis of the correlated model.

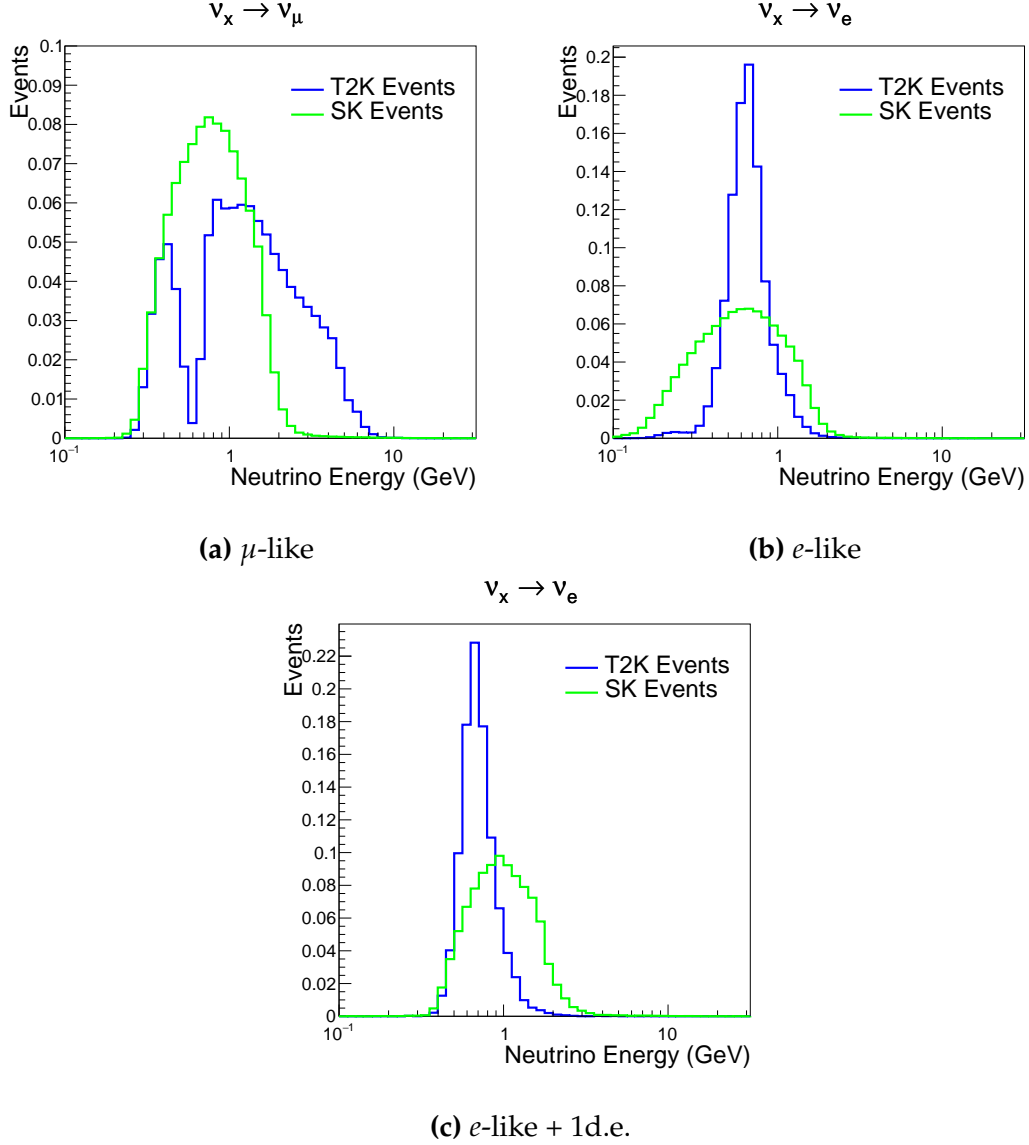


Figure 6.8: The predicted neutrino energy distribution for subGeV atmospheric and beam samples. FHC and RHC beam samples are summed together Asimov A oscillation parameters are assumed (given in Table 2.2). Beam and atmospheric samples with similar cuts are compared against one another.

The T2K systematic model [201] is applied in a similar methodology to the SK model parameters. It consists of 19 shape parameters and 24 normalisation parameters. Four additional parameters, which model the uncertainty in the binding energy, are applied in a way to shift the momentum of the lepton emitted from a nucleus. This controls the uncertainty specified on the 27MeV binding energy assumed within Equation 6.2. The majority of these parameters are

	CC QE	CC 2p2h	CC $1\pi^\pm$	CC $M\pi$	CC Other	NC $1\pi^0$	NC $1\pi^\pm$	NC $M\pi$	NC Coh.	NC Other
FHC 1R+1d.e. e-like	0.04	0.02	0.83	0.03	0.04	0.01	0.01	0.01	0.00	0.01
RHC 1R e-like	0.62	0.12	0.11	0.01	0.02	0.06	0.01	0.01	0.01	0.04
FHC 1R e-like	0.68	0.12	0.10	0.00	0.02	0.04	0.01	0.00	0.00	0.02
RHC 1R μ -like	0.62	0.13	0.17	0.02	0.03	0.00	0.02	0.00	0.00	0.00
FHC 1R μ -like	0.62	0.12	0.16	0.02	0.03	0.00	0.03	0.00	0.00	0.00
S.G. π^0 -like	0.05	0.01	0.02	0.00	0.01	0.68	0.06	0.07	0.06	0.04
S.G. μ -like 2de	0.04	0.01	0.80	0.10	0.04	0.00	0.00	0.00	0.00	0.00
S.G. μ -like 1de	0.72	0.11	0.12	0.01	0.02	0.00	0.01	0.00	0.00	0.00
S.G. μ -like 0de	0.68	0.11	0.10	0.01	0.02	0.01	0.05	0.01	0.00	0.02
S.G. e-like 1de	0.05	0.01	0.75	0.10	0.05	0.00	0.01	0.02	0.00	0.01
S.G. e-like 0de	0.73	0.11	0.10	0.01	0.02	0.02	0.00	0.00	0.00	0.00

Figure 6.9: The interaction mode contribution of each sample given as a fraction of the total event rate in that sample. Asimov A oscillation parameters are assumed (given in Table 2.2). The Charged Current (CC) modes are broken into quasi-elastic (QE), 2p2h, resonant charged pion production ($1\pi^\pm$), multi-pion production ($M\pi$), and other interaction categories. Neutral Current (NC) interaction modes are given in interaction mode categories: π^0 production, resonant charged pion production, multi-pion production, and others.

assigned a Gaussian prior uncertainty. Those that have no reasonably motivated uncertainty, or those which have not been fit to external data, are assigned a flat prior which does not affect the penalty term.

On top of the combination of the SK and T2K interaction models, several other parameters have been specifically developed for the joint oscillation analysis. The majority of the atmospheric samples' δ_{CP} sensitivity comes from the normalisation of subGeV electron-like events. These are modeled using a spectral function to approximate the nuclear ground state. However, the near detector is not able to constrain the model so an additional systematic is introduced which models applies an alternative Continuous Random Phase Approximation (CRPA) nuclear ground state. This dial approximates the event weights if a CRPA model

had been assumed rather than a spectral function. This dial only applies to ν_e and $\bar{\nu}_e$ as the near detector does not constraint ν_e cross-section measurements. It is applied as a shape parameter.

Further additions to the model have been introduced due to the inclusion of the subGeV π^0 atmospheric sample. This particularly targets charged current and neutral current π^0 producing interactions to help constrain the systematic uncertainties. Therefore, an uncertainty that affects neutral current resonant π^0 production is incorporated into this analysis. Comparisons of NEUT's NC resonant pion production predictions have been made to MiniBooNE [204] data and a consistent 16% to 21% underprediction is observed [202]. Consequently, a conservative 30% normalisation parameter is invoked.

Down-going events are mostly insensitive to oscillation parameters and can act similar to the near detector within an accelerator experiment (Details will be discussed in chapter 7). This region of phase space can act as a sideband and allows the cross-section model and near detector constraint to be studied. The distribution of events in this region is calculated using the technique outlined in subsection 4.3.4. The results are illustrated in Figure 6.10. For CCQE-targeting samples, the application of the near detector constraint is well within the statistical fluctuation of the down-going data such that no significant tension is observed between the data and the Monte Carlo prediction after the near detector constraint is applied. This is not the case for samples with target CCRES interactions. The electron-like data is consistent with the constrained prediction at high reconstructed momenta but diverges at lower momentum, whereas the muon-like sample is under-predicted throughout the range of momenta. To combat this disagreement, an additional cross-section systematic dial, specifically designed to inflate the low pion momentum systematics was developed in [202]. This is a shape parameter implemented through a splined response.

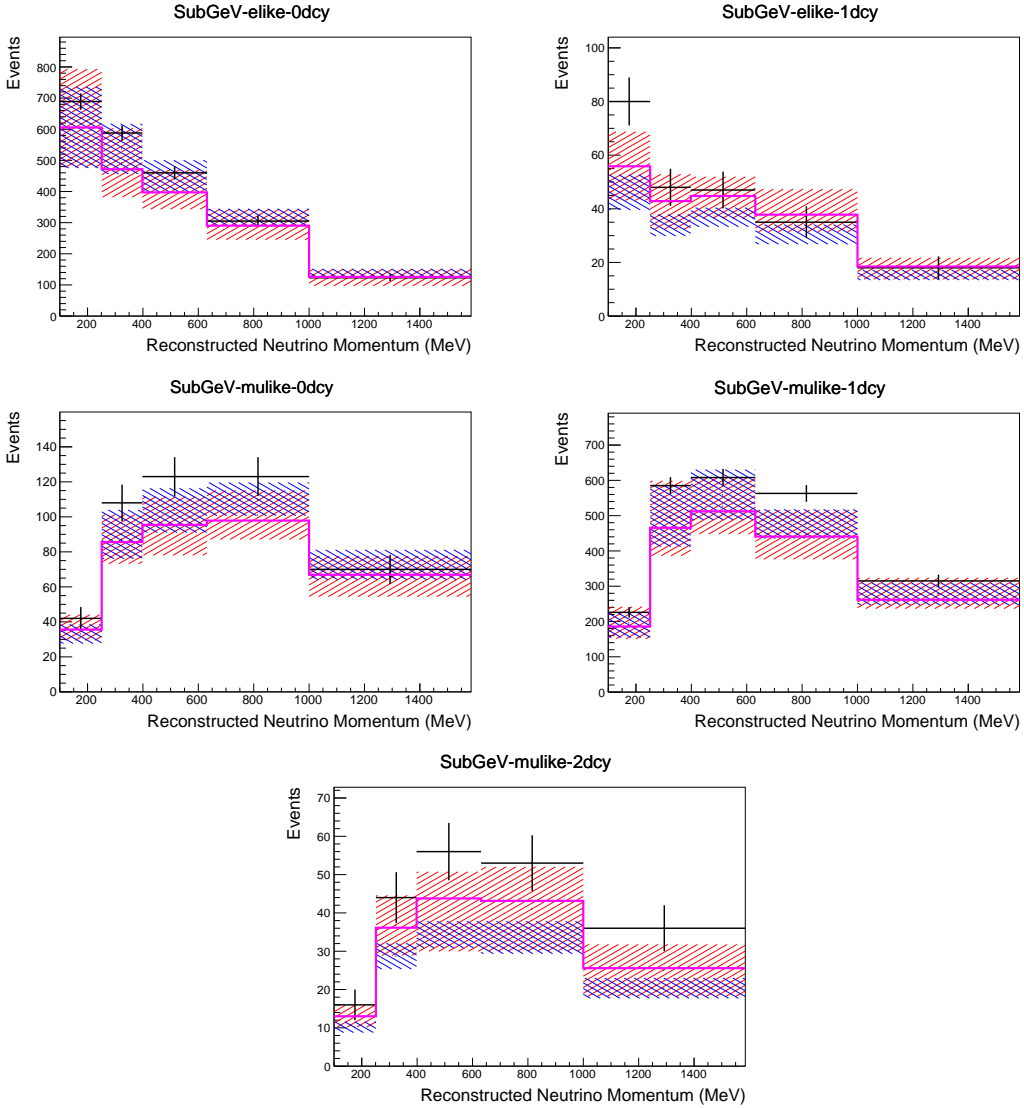


Figure 6.10: Down-going atmospheric subGeV single-ring samples comparing the mean and error of the pre-fit and post-fit Monte Carlo predictions in red and blue, respectively. The magenta histogram illustrates the Monte Carlo prediction using the generated dial values. The black points illustrate the down-going data with statistical errors given. The mean and errors of the Monte Carlo predictions are calculated by the techniques documented in subsection 4.3.4. The pre-fit spectrum is calculated by throwing the cross-section and atmospheric flux dial values from the pre-fit covariance matrix. The post-fit spectrum is calculated by sampling the cross-section dial values from an ND fit MCMC chain, whilst still throwing the atmospheric flux dials from the pre-fit covariance.

6.4.4 Near Detector

The systematics applied due to uncertainties arising from the response of the near detector is documented in [131]. The response is described by 574 normalisation parameters binned in the selected sample as well as momentum and angle,

P_μ and $\cos(\theta_\mu)$, of the final-state muon. These are applied via a covariance matrix with each parameter being assigned a Gaussian prior from that covariance matrix. These normalisation parameters are built from underlying systematics, e.g. pion secondary interaction systematics, which are randomly thrown and the variation in each $P_\mu \times \cos(\theta_\mu)$ bin is determined. Two thousand throws are evaluated and a covariance matrix response is created. This allows significant correlations between FGD1 and FGD2 samples, as well as adjacent $P_\mu \times \cos(\theta_\mu)$ bins. Statistical uncertainties are accounted for by including fluctuations of each event's weight from a Poisson distribution.

Similar to the cross-section systematics, MaCh3 and BANFF are used to constrain the uncertainty of these systematics through independent validations. Each fitter generates a post-fit covariance matrix which is compared and passed to the far-detector oscillation analysis working group. As the analysis presented within this thesis uses the MaCh3 framework, a joint oscillation analysis fit of all three sets of samples and their respective systematics is performed.

6.4.5 Far Detector

Two configurations of the far detector systematic model implementation have been considered. Firstly, the far detector systematic uncertainties for beam and atmospheric samples are taken from their respective analysis inputs, denoted “official inputs” analysis, with no correlations assumed between the beam and atmospheric samples. The beam- and atmospheric-specific inputs are documented in subsubsection 6.4.5.1 and subsubsection 6.4.5.2. Secondly, an alternative detector model has been developed which correlates the response of the SK detector systematics between the beam and atmospheric samples. Here, the distribution of parameters used for applying event cuts (e.g. electron-muon PID separation) is modified within the fit. It follows a similar methodology to the beam far detector systematics implementation but performs a joint fit of the beam and atmospheric data. This alternative implementation is detailed in subsubsection 6.4.5.3.

6.4.5.1 Beam Samples

There are 45 systematics which describe the response of the far detector to beam events [183], split into 44 normalisation parameters and one energy scale systematic. The energy scale systematic is applied as a multiplicative scaling of the reconstructed neutrino energy. It is estimated from data-to-Monte Carlo differences in the stopping muon sample in [185] and found to be 2.1%. The normalisation parameters are assigned a Gaussian error centred at one with width taken from a covariance matrix. A detailed breakdown of the generation of the covariance matrix is found in [196]. To build the covariance matrix, a fit is performed on atmospheric data which has been selected using beam sample selection cuts. These cuts use the variables, L^i , where the index i is detailed in Table 6.6. Each L^i is a smear, α , and shift, β parameter such that,

$$L_j^i \rightarrow \bar{L}_j^i = \alpha_j^i L + \beta_j^i \quad (6.4)$$

Where L_j^i (\bar{L}_j^i) correspond to nominal(varied) PID cut parameters given in Table 6.6. The shift and smear parameters are nuisance parameters with no prior constraints. They are binned by final-state topology, j , where the binning is given in Table 6.7. The final-state topology binning is because the detector will respond differently to events that have one or multiple rings. For example, the detector will be able to distinguish single-ring events better than two overlapping ring events, resulting in different systematic uncertainty for one-ring events compared to two-ring events. This approach is used to allow the cut parameter distributions to be modified within the fit, allowing for better data to Monte Carlo agreement.

Cut Variable	Parameter Name
0	<code>fitQun e/mu PID</code>
1	<code>fitQun e/pi0 PID</code>
2	<code>fitQun mu/pi PID</code>
3	<code>fitQun Ring-Counting Parameter</code>

Table 6.6: List of cut variables that are included within the shift/smear fit documented in [196].

Category	Description
$1e$	Only one electron above Cherenkov threshold in the final state
1μ	Only one muon above Cherenkov threshold in the final state
$1e+other$	One electron and one or more other charged particles above Cherenkov threshold in the final state
$1\mu+other$	One muon and one or more other charged particles above Cherenkov threshold in the final state
$1\pi^0$	Only one π^0 in the final state
$1\pi^\pm$ or $1p$	Only one hadron (typically charged pion or proton) in the final state
Other	Any other final state

Table 6.7: Reconstructed event topology categories on which the SK detector systematics [196] are based.

The mis-modeling of π^0 events is also considered. If one of the two rings from a π^0 event is missed, this will be reconstructed as a CC ν_e -like event. This is one of the largest systematics hindering the electron neutrino appearance analyses. Consequently, additional systematics have been introduced to constrain the mis-modeling of π^0 events in SK, binned by reconstructed neutrino energy. To evaluate this systematic uncertainty, a set of “hybrid- π^0 ” samples is constructed. These events are built by overlaying one electron-like ring from the SK atmospheric neutrino samples or decay electron ring from a stopping cosmic ray muon with one simulated photon ring. Both rings are chosen so that momenta and opening angle follow the decay kinematics of NC π^0 events from the T2K-MC. Hybrid- π^0 Monte Carlo samples with both rings from the SK Monte Carlo are produced to compare with the hybrid- π^0 data samples and the difference in the fraction of events that pass the ν_e selection criteria is used to assign the systematic errors. In order to investigate any data to Monte Carlo differences that may originate from either the higher energy ring or lower energy ring, two samples are built; a sample in which the electron constitutes the higher energy ring from the π^0 decay (called the primary sample) and another one in which it constitutes the lower energy ring (called the secondary sample). The standard T2K ν_e fitQun event selection criteria are used to select events.

Final contributions to the covariance matrix are determined by supplementary uncertainties obtained by comparing stopping muon data to Monte Carlo

prediction, as first introduced in section 5.2. The efficiency of tagging decay electrons is estimated by the stopping muon data to Monte Carlo differences by comparing the number of one decay electron events to the number of events with one or fewer decay electrons. Similarly, the rate at which fake decay electrons are reconstructed by `f iTQun` is estimated by comparing the number of two decay electron events to the number of events with one or two reconstructed decay electrons. The two sources of systematics are added in quadrature weighted by the number of events with one true decay electron yielding a 0.2% systematic uncertainty. A fiducial volume systematic of $\pm 2.5\text{cm}$ which corresponds to a 0.5% shift in the normalisation of events is also applied. Additional normalisation uncertainties based on neutrino flavour and interaction mode are also defined in [183, 205, 206].

Two additional sources of uncertainty are included: secondary and photo-nuclear interactions. These are estimated by varying the underlying parameters are building a distribution of sample event rates. These contributions are then added in quadrature to the above covariance matrix.

DB: Matrix Diagonals from SK Det Matrix

6.4.5.2 Atmospheric Samples

The detector systematics for atmospheric samples, documented in [86], are split into two sub-groups: those which are related to particle identification and ring counting systematics, and those which are related to calibration, separation, and reduction uncertainties.

The particle identification systematics consist of five parameters. The ring separation systematic enforces an anti-correlated response between the single-ring and multi-ring samples. This is implemented as a fractional increase/decrease in the overall normalisation of each sample, depending on the distance to the nearest wall from an event's vertex. The coefficients of the normalisation are estimated prior to the fit and depend on the particular atmospheric sample. Two electron-muon separation systematics are included within this model which

anti-correlate the response of the electron-like and muon-like samples: one for single-ring events and another for multi-ring events.

The multi-ring electron-like separation likelihood, discussed in section 6.1, encodes the ability of the detector to separate neutrino from anti-neutrino events. Two normalisation parameters vary the relative normalisation of multi-ring ν_e and $\bar{\nu}_e$ samples whilst keeping a consistent overall event rate.

There are 22 systematics related to calibration measurements, including effects from backgrounds, reduction, and showering effects. They are documented in [86] and are briefly summarised in Table 6.8. They are applied via normalisation parameters, with the separation systematics requiring the conservation of event rate across all samples.

6.4.5.3 Correlated Detector Model

A complete uncertainty model of the SK detector would be able to determine the systematic shift on the sample spectra for a variation of the underlying parameters, e.g. PMT angular acceptance. However, this is computationally intensive, requiring Monte Carlo predictions to be made for each plausible variation. Consequently, an effective parameter model has been utilised for a correlated detector model following from the T2K-only model implementation documented in subsubsection 6.4.5.1. It correlates the detector systematics between the far-detector beam and subGeV atmospheric samples due to their similar energies and interaction types. As there are no equivalent beam samples, the multiGeV, multiring, PC, and Up- μ samples will be subject to the particle identification systematics implementation as described in subsubsection 6.4.5.2 rather than using this correlated detector model. The calibration systematics also described in the aforementioned chapter still apply to all atmospheric samples.

The correlated detector model utilises the same smear and shift parameters documented in subsubsection 6.4.5.1, split by final state topology. Beyond this, the shift and smear parameters are split by visible energy deposited within the detector, with binning specified in Table 6.9. This is because atmospheric events

Index	Description
0	Partially contained reduction
1	Fully contained reduction
2	Separation of fully contained and partially contained events
3	Separation of stopping and through-going partially contained events in top of detector
4	Separation of stopping and through-going partially contained events in barrel of detector
5	Separation of stopping and through-going partially contained events in bottom of detector
6	Background due to cosmic rays
7	Background due to flasher events
8	Vertex systematic moving events into and out of fiducial volume
9	Upward going muon event reduction
10	Separation of stopping and through-going in upward going muon events
11	Energy systematic in upward going muon events
12	Reconstruction of the path length of upward going muon events
13	Separation of showering and non-showering upward going muon events
14	Background of stopping upward going muon events
15	Background of non-showering through-going upward going muon events
16	Background of showering through-going upward going muon events
17	Efficiency of tagging two rings from π^0 decay
18	Efficiency of decay electron tagging
19	Background from downgoing cosmic muons
20	Asymmetry of energy deposition in tank
21	Energy scale deposition

Table 6.8: Sources of systematic errors specified within the grouped into the “calibration” systematics model.

are categorised by subGeV and multiGeV events based on visible energy, so this splitting is required when correlating the systematic model for beam and atmospheric events. Alongside the technical requirement, higher energy events will be better reconstructed due to fractionally less noise within the detector. As a result of the inclusion of visible energy binning, Equation 6.4 becomes

$$L_{jk}^i \rightarrow \bar{L}_{jk}^i = \alpha_{jk}^i L + \beta_{jk}^i, \quad (6.5)$$

where k is the visible energy bin.

The implementation of this systematic model takes the events reconstructed values of the cut parameters, modifies them by the particular shift and smear

Index	Range (MeV)
0	$30 \geq E_{vis} > 300$
1	$300 \geq E_{vis} > 700$
2	$700 \geq E_{vis} > 1330$
3	$E_{vis} \geq 1330$

Table 6.9: Visible energy binning for which the correlated SK detector systematics are based

parameter for that event, and then re-applies event selection. This causes event migration, which is a new feature incorporated into the MaCh3 framework which is only achievable due to the event-by-event reweighting scheme.

Particular care has to be taken when varying the ring counting parameter. This is because the number of rings is a finite value (one-ring, two-rings, etc.) which can not be continuously varied through this shift and smear technique. Consequently a continuous ring counting parameter, RC_i , is calculated for the i^{th} event, following the definition in [182]: the preferred likelihoods from all considered one-ring (L_{1R}) and two-ring (L_{2R}) fits are determined. The difference is computed as $\Delta_{LLH} = \log(L_{1R}) - \log(L_{2R})$. The ring counting parameter is then defined as

$$RC_i = \text{sgn}(\Delta_{LLH}) \times \sqrt{|\Delta_{LLH}|}, \quad (6.6)$$

where $\text{sgn}(x) = x/|x|$. This ring counting parameter corresponds to an intermediate likelihood value used within the `fitQun` algorithm to decide the number of rings associated with a particular event. However, fake-ring merging algorithms are applied after this likelihood value is used. Consequently, this ring counting parameter does not always exactly correspond to the number of reconstructed rings. This can be seen in Figure 6.11.

As the `fitQun` algorithm does not provide a likelihood value after the fake-ring algorithms have been applied, the ring counting parameter distribution is correlated to the final number of reconstructed rings through “maps”. These are two-dimensional distributions of the ring counting parameter and the final number of reconstructed rings. An example is illustrated in Figure 6.12. In principle,

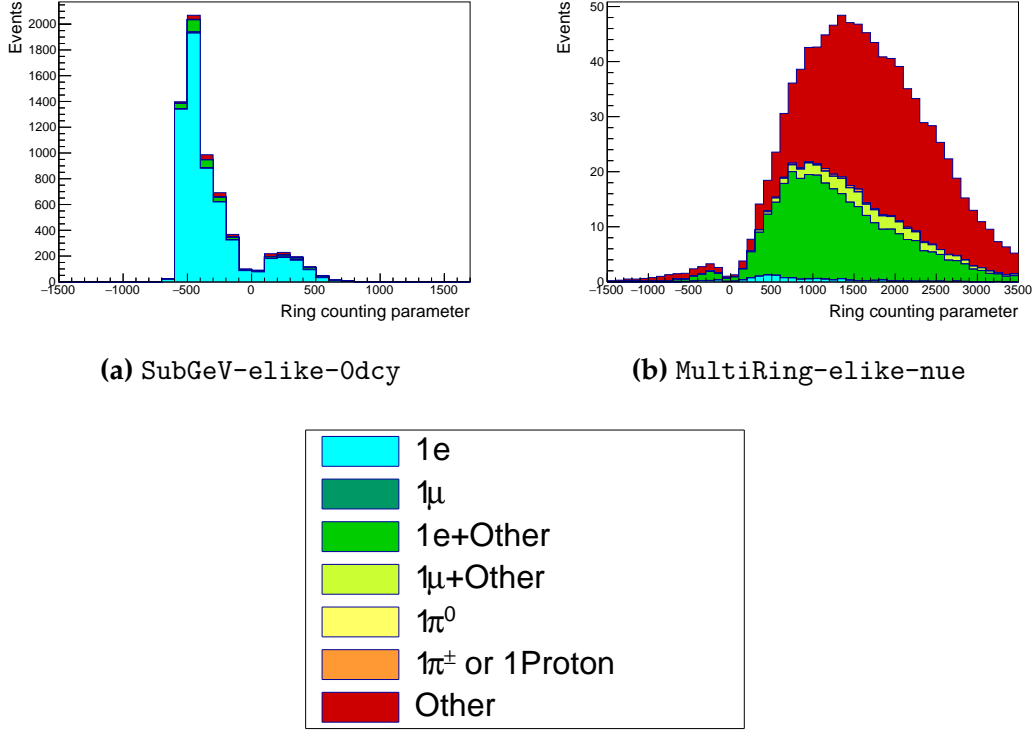


Figure 6.11: The ring counting parameter as defined in Equation 6.6 for the SubGeV-else-like-0dcy and MultiRing-else-like-nue samples.

the `fitQun` reconstruction algorithm should be re-run after the variation in the ring counting parameter. However, this is not computationally viable. Therefore the “maps” are used as a reweighting template.

The maps are split by final state topology and true neutrino flavour and all `fitQun`-reconstructed Monte Carlo events are used to fill them. The maps are row-normalised to represent the probability of X rings for a given RC_i value. Prior to the oscillation fit, an event’s nominal weight is calculated as $W^i(N_{Rings}^i, L_{jk}^i)$, where N_{Rings}^i is the reconstructed number of rings for the i^{th} event and $W^i(x, y)$ is the bin content in map associated with the i^{th} event, where x number of rings and y is ring counting parameter. Then during the fit, the value of $R = W^i(N_{Rings}^i, \bar{L}_{jk}^i)/W^i(N_{Rings}^i, L_{jk}^i)$ is calculated as the event weight for the i^{th} event. This is the only cut variable that uses a reweighting technique rather than event migration.

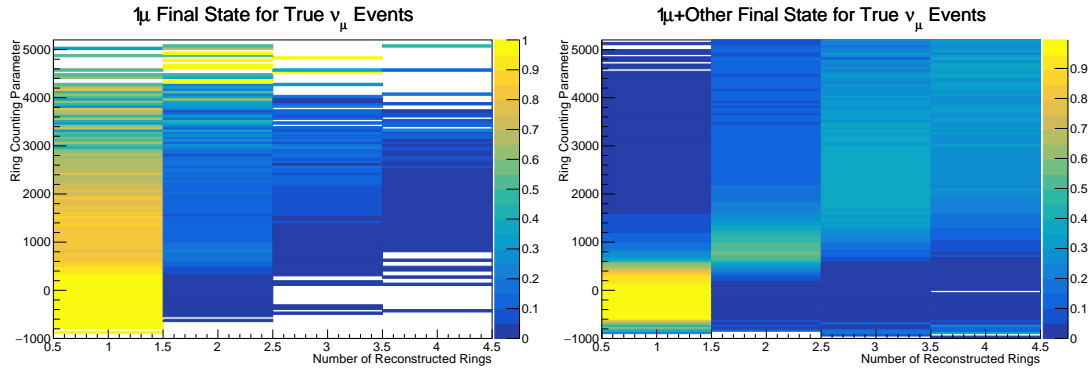


Figure 6.12: The ring counting parameter, defined in Equation 6.6, as a function of the number of reconstructed rings as found by the `fitQun` reconstruction algorithm. Left: true ν_μ events with only one muon above the Cherenkov threshold in the final state. Right: true ν_μ events with one muon and at least one other charged particle above the Cherenkov threshold in the final state.

The π^0 systematics introduced in subsection 6.4.4 are applied via a covariance matrix. This is not possible in the alternative model as no covariance matrix is used. Thus, the implementation of the π^0 systematics has been modified. The inputs from the hybrid π^0 sample are included via the use of “ χ^2 maps”, which are two-dimensional histograms in α_{jk}^i and β_{jk}^i parameters over some range. Illustrative examples of the χ^2 maps are given in Figure 6.13. Due to their nature, the shift and smear parameters are typically very correlated. A map is produced for each cut parameter given in Table 6.6 and for each visible energy bin given in Table 6.9.

The maps are filled through the χ^2 comparison of the hybrid π^0 Monte Carlo and data in the particle identification parameters documented in Table 6.6. The Monte Carlo distribution is modified by the α_{jk}^i and β_{jk}^i scaling, whilst cross-section and flux nuisance parameters are thrown from their prior uncertainties. The χ^2 between the scaled Monte Carlo and data is calculated and the relevant point in the χ^2 map is filled.

The implementation within this alternative detector model is to add the bin contents of the maps, for the relevant values of the α_{jk}^i and β_{jk}^i parameters, to the likelihood penalty. Only 1 π^0 final state topology shift and smear parameters use this prior uncertainty.

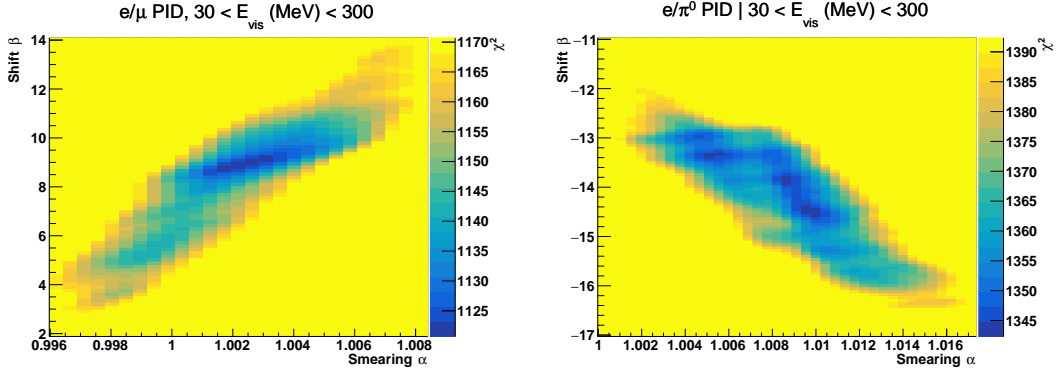


Figure 6.13: The χ^2 between the hybrid- π^0 Monte Carlo and data samples, as a function of smear (α) and shift (β) parameters, for events which have $1\pi^0$ final state topology. Left: Electron-muon separation PID parameter for events with $30 \leq E_{\text{vis}}(\text{MeV}) < 300$. Right: Electron- π^0 separation PID parameter for events with $30 \leq E_{\text{vis}}(\text{MeV}) < 300$.

Similarly, the implementation of the supplementary systematics documented in subsubsection 6.4.5.1 needs to be modified. A new framework [207] was built in tandem with the T2K-SK working group [183] so the additional parameters can be incorporated into the MaCh3 framework. These are applied as normalisation parameters, depending on the particular interaction mode, number of tagged decay electrons, and whether the primary particle generated Cherenkov light. They are assigned Gaussian uncertainties with widths described by a covariance matrix. Furthermore, the secondary interaction and photo-nuclear effects need to be accounted for in this detector model using a different implementation than that in subsubsection 6.4.5.1. This was done by including a shape parameter for each of the secondary interactions and the photo-nuclear systematic parameters.

There are a total of 224 α_{jk}^i and β_{jk}^i parameters, of which 32 have prior constraints from the hybrid π^0 samples.

One final complexity of this correlated detector model is that the two sets of samples, beam and subGeV atmospheric, use slightly different parameters to distinguish electron and muon-like events. The T2K samples use the value of $\log(L_e/L_\mu)$ whereas the atmospheric samples use the value of $\log(L_e/L_\pi)$, where L_X is the likelihood for hypothesis X. This is because the T2K fits use single-ring `f1TQun` fitting techniques, whereas multi-ring fits are applied to the atmospheric samples where only the electron and pion hypothesis are considered.

The correlation between the two likelihood ratios is illustrated in Figure 6.14. As discussed in section 5.2, the pion hypothesis is a very good approximation of the muon hypothesis due to their similar mass. Consequently, using the same shift and smear parameters correlated between the beam and subGeV atmospheric samples is deemed a good approximation.

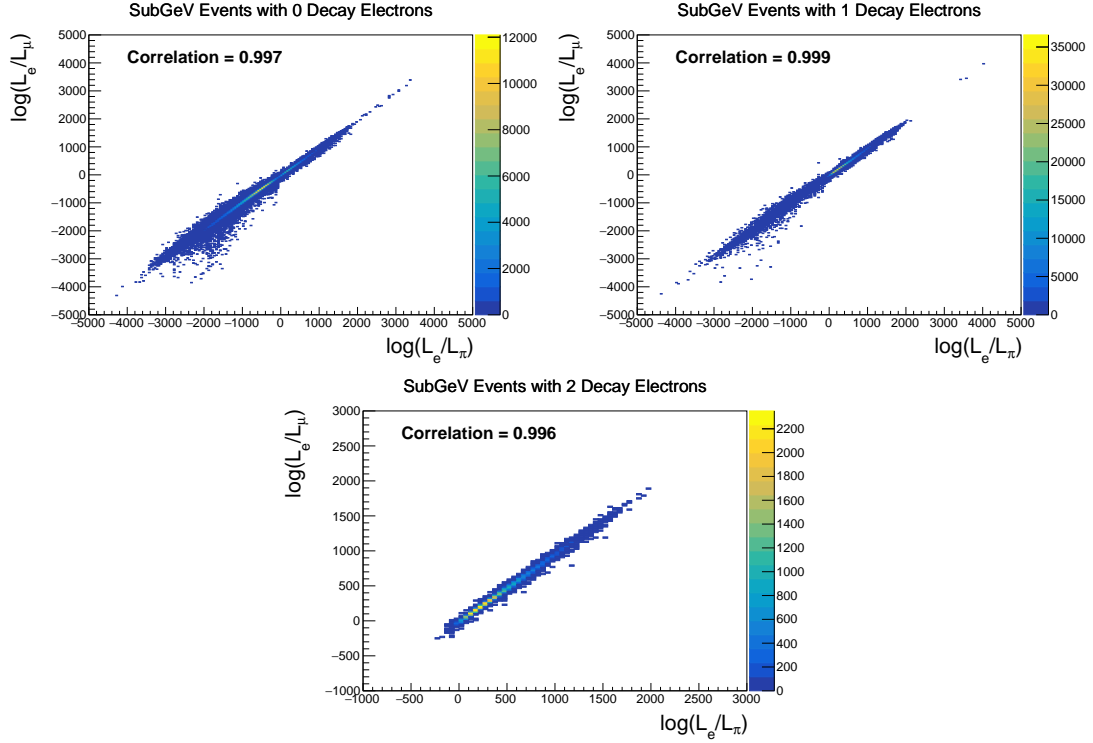


Figure 6.14: The distribution of $\log(L_e/L_\mu)$ compared to $\log(L_e/L_\pi)$ for subGeV events with zero (top left), one (top right) or two (bottom) decay electrons. The correlation in the distribution is calculated as 0.997, 0.999 and 0.996, respectively.

7

Oscillation Probability Calculation

It is important to understand how and where the sensitivity to the oscillation parameters comes from for both atmospheric and beam samples. An overview of how these samples observe changes in δ_{CP} , Δm_{23}^2 , and $\sin^2(\theta_{23})$ affect these samples is given in section 7.1. This section also explains the additional complexities involved when performing an atmospheric neutrino analysis as compared to a beam-only analysis.

Without additional techniques, atmospheric sub-GeV upward-going neutrinos ($E_\nu < 1.33\text{GeV}, \cos(\theta_Z) < 0.$) can artificially inflate the sensitivity to δ_{CP} due to the quickly varying oscillation probability in this region. Therefore, a “sub-sampling” approach has been developed to reduce these biases ensuring accurate and reliable sensitivity measurements. This technique ensures that small-scale unresolvable features of the oscillation probability have been averaged over whilst the large-scale features in the oscillation probability are unaffected. The documentation and validation of this technique are found in section 7.2. The oscillation probability calculation is computationally intensive due to the large number of matrix multiplications needed. Consequently, the CUDAProb3 implementation choice made within the fitting framework, as detailed in section 7.3, ensures that the analysis can be done in a timely manner.

Whilst the beam neutrinos are assumed to propagate through a constant density slab of material, the density variations through the Earth result in more complex oscillation patterns for atmospheric neutrinos. Furthermore, the uncertainty in the electron density can modify the oscillation probability for the denser core layers of the Earth. The model of the Earth used within this analysis is detailed in section 7.4. This includes information about the official SK-only methodology as well as improvements that have been made to remove some of the approximations used in that analysis. Another complexity of atmospheric neutrinos oscillation studies is that the height of production in the atmosphere is not known on an event-by-event basis. An analytical averaging technique that approximates the uncertainty of the oscillation probability has been followed, with the author of this thesis being responsible for the implementation and validation. This implementation of an external technique is illustrated in section 7.5.

7.1 Overview

DB: Should this be moved into an earlier chapter? The selections chapter references the matter resonance which has not yet been explained at that point

The analysis presented within this thesis focuses on the determination of oscillation parameters from atmospheric and beam neutrinos. Whilst subject to the same oscillation formalism, the way in which the two samples have sensitivity to the different oscillation parameters differs significantly.

Atmospheric neutrinos have a varying baseline, or “path length”, L , such that the distance each neutrino travels before interacting is dependent upon the zenith angle, θ_Z . As primary cosmic rays can interact anywhere between the Earth’s surface and $\sim 50\text{km}$ above that, the height, h , in the atmosphere at which the neutrino was generated also affects the path length,

$$L = \sqrt{(R_E + h)^2 - R_E^2 (1 - \cos^2(\theta_Z))} - R_E \cos(\theta_Z). \quad (7.1)$$

Where $R_E = 6,371\text{km}$ is the Earth's radius. Consequently, the oscillation probability is dependent upon two parameters, $\cos(\theta_Z)$ and E_ν .

The oscillation probability used within this analysis is based on [21]. The neutrino wavefunction in the vacuum Hamiltonian evolves in each layer of constant matter density via

$$i \frac{d\psi_j(t)}{dt} = \frac{m_j^2}{2E_\nu} \psi_j(t) - \sum_k \sqrt{2} G_F N_e U_{ej} U_{ke}^\dagger \psi_k(t), \quad (7.2)$$

where m_j^2 is the square of the j^{th} vacuum eigenstate mass, E_ν is the neutrino energy, G_F is Fermi's constant, N_e is the electron number density and U is the PMNS matrix. The transformation $N_e \rightarrow -N_e$ and $\delta_{CP} \rightarrow -\delta_{CP}$ is applied for antineutrino propagation. Thus, a model of the Earth's density is required for neutrino propagation. Following the official SK-only methodology [208], this analysis uses the Preliminary Reference Earth Model (PREM) [209] which provides piecewise cubic polynomials as a function of the Earth's radius. This density profile is illustrated in Figure 7.1. As the propagator requires layers of constant density, the SK methodology approximates the PREM model by using four layers of constant density [208], detailed in Table 7.1.

Layer	Outer Radius [km]	Density [g/cm ³]	Chemical composition (Z/A)
Inner Core	1220	13	0.468 ± 0.029
Outer Core	3480	11.3	0.468 ± 0.029
Lower Mantle	5701	5.0	0.496
Transition Zone	6371	3.3	0.496

Table 7.1: Description of the four layers of the Earth invoked within the constant density approximation of the PREM model [209].

The atmospheric neutrino oscillation probabilities can be presented as two dimensional “oscillograms” as illustrated in Figure 7.2. The distinct discontinuities, as a function of $\cos(\theta_Z)$, are due to the discontinuous density in the PREM model.

Atmospheric neutrinos have sensitivity to δ_{CP} through the overall event rate. Figure 7.3 illustrates the difference in oscillation probability between CP-conserving ($\delta_{CP} = 0$) and a CP-violating ($\delta_{CP} = -1.601$) value taken from

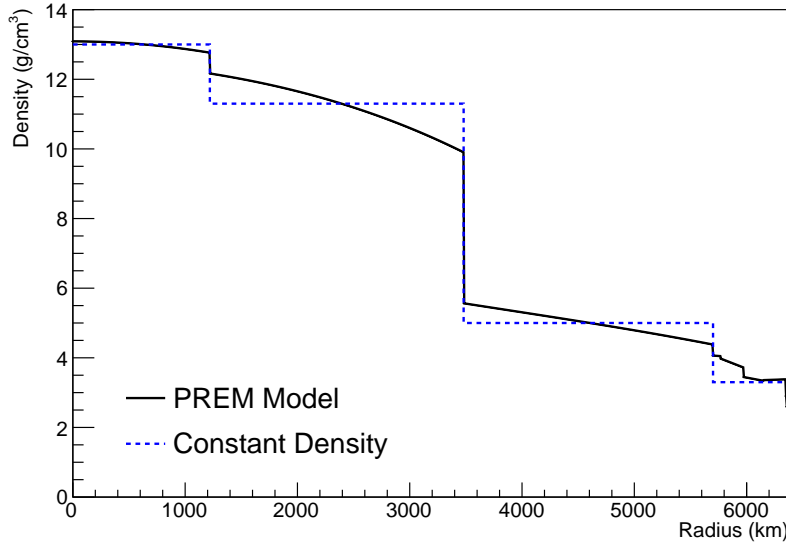


Figure 7.1: The density of the Earth given as a function of the radius, as given by the PREM model (Black), and the constant density four-layer approximation (Blue), as used in the official SK-only analysis.

Asimov A oscillation parameter set (Table 2.2). The result is a complicated oscillation pattern in the appearance probability for sub-GeV upgoing neutrinos. The detector does not have sufficient resolution to resolve these individual patterns so the sensitivity to δ_{CP} for atmospheric neutrinos comes via the overall normalisation of these events.

The presence of matter means that the effect δ_{CP} has on the oscillation probability is not equal between neutrinos and antineutrinos. Furthermore, the interaction cross-section for neutrinos is larger than antineutrinos so the two effects have to be disentangled. These effects are further convoluted by detector efficiencies as SK cannot distinguish neutrinos and antineutrinos well. Furthermore, the sample selections discussed in section 6.1 have difference efficiencies for neutrino-antineutrino selections. All of these effects lead to a difference in the number of neutrinos detected compared to antineutrinos. This changes how the δ_{CP} normalisation term is observed, resulting in a very complex sensitivity to δ_{CP} .

The vacuum and matter oscillation probabilities for $P(\nu_e \rightarrow \nu_e)$ and $P(\bar{\nu}_e \rightarrow$

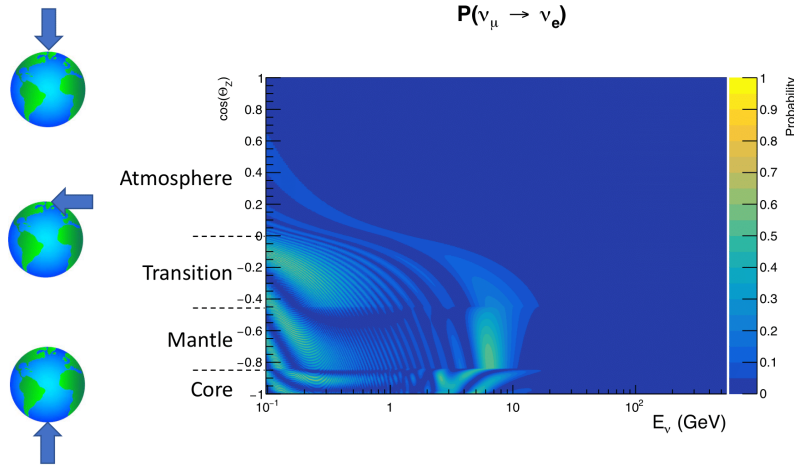


Figure 7.2: An “oscillogram” that depicts the $P(\nu_\mu \rightarrow \nu_e)$ oscillation probability as a function of neutrino energy and cosine of the zenith angle. The zenith angle is defined such that $\cos(\theta_Z) = 1.0$ represents neutrinos that travel from directly above the detector. The four-layer constant density PREM model approximation is used and Asimov A oscillation parameters are assumed (Table 2.2).

$\bar{\nu}_e$) are presented in Figure 7.4, where the PREM model has been assumed. The oscillation probability for both neutrinos and antineutrinos is affected in the presence of matter. However, the resonance effects around $O(5)\text{GeV}$ only occur for neutrinos in normal mass hierarchy and antineutrinos in inverse mass hierarchy. The exact position and amplitude of the resonance depend on $\sin^2(\theta_{23})$, further increasing the atmospheric neutrinos’ sensitivity to the parameter.

As the T2K beam flux is centered at the first oscillation maximum ($E_\nu = 0.6\text{GeV}$), the sensitivity to δ_{CP} is predominantly observed as a change in the event-rate of e-like samples in $\nu/\bar{\nu}$ modes. Figure 7.5 illustrates the $P(\nu_\mu \rightarrow \nu_e)$ oscillation probability for a range of δ_{CP} values. A circular modulation of the first oscillation peak (in both magnitude and position) is observed when varying throughout the allowable values of δ_{CP} . The CP-conserving values of $\delta_{CP} = 0, \pi$ have a lower(higher) oscillation maximum than the CP-violating values of $\delta_{CP} = -\pi/2(\delta_{CP} = \pi/2)$. A sub-dominant shift in the energy of the oscillation peak is also present, which aids in separating the two CP-conserving values of δ_{CP} .

T2K’s sensitivity to $\sin^2(\theta_{23})$ and Δm_{23}^2 is observed as a shape-based variation

$$\mathbf{P}(\nu_\mu \rightarrow \nu_e; \delta_{CP} = -1.601) - \mathbf{P}(\nu_\mu \rightarrow \nu_e; \delta_{CP} = 0.)$$

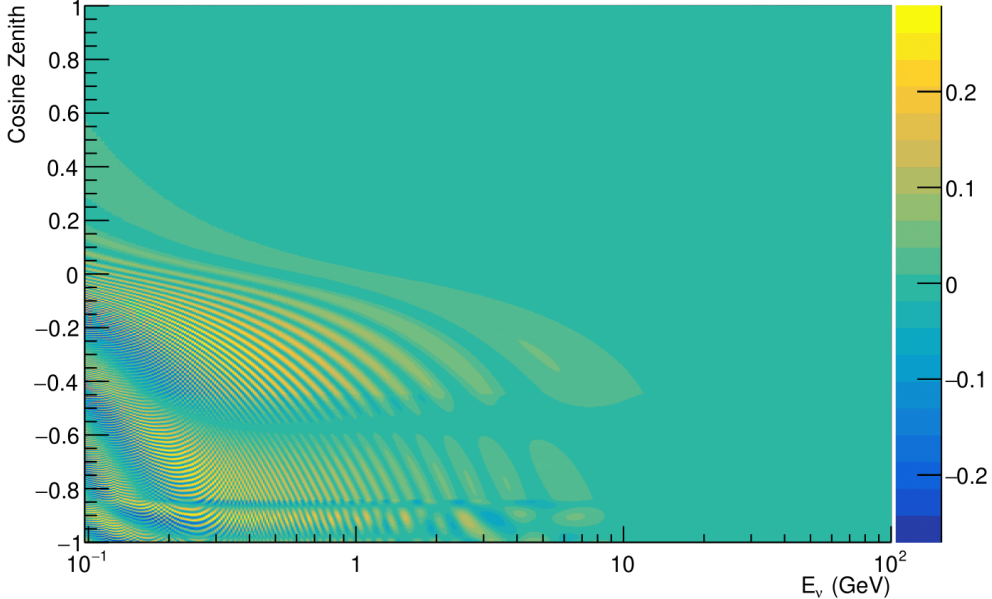


Figure 7.3: The effect of δ_{CP} for atmospheric neutrinos given in terms of the neutrino energy and zenith angle. This oscillogram compares the $P(\nu_\mu \rightarrow \nu_e)$ oscillation probability for a CP conserving ($\delta_{CP} = 0.0$) and a CP violating ($\delta_{CP} = -1.601$) value taken from the Asimov A parameter set. The other oscillation parameters assume the Asimov A oscillation parameter set given in Table 2.2.

of the muon-like samples, as illustrated in Figure 7.5. The value of Δm_{32}^2 laterally shifts the position of the oscillation dip (around $E_\nu \sim 0.6\text{GeV}$) in the $P(\nu_\mu \rightarrow \nu_\mu)$ oscillation probability. A variation of $\sin^2(\theta_{23})$ is predominantly observed as a vertical shift of the oscillation dip with second-order horizontal shifts being due to matter effects. The beam neutrinos have limited sensitivity to matter effects due to the relatively shorter baseline as well as the Earth's mantle being a relatively low-density material (as compared to the Earth's core). For some values of δ_{CP} , the degeneracy in the number of e-like events allows the mass hierarchy to be broken. This leads to a δ_{CP} -dependent mass hierarchy sensitivity which can be seen in Figure 7.6.

Whilst all oscillation channels should be included for completeness, the computational resources required to run a fit are limited and any reasonable approximations which reduce the number of oscillation probability calculations

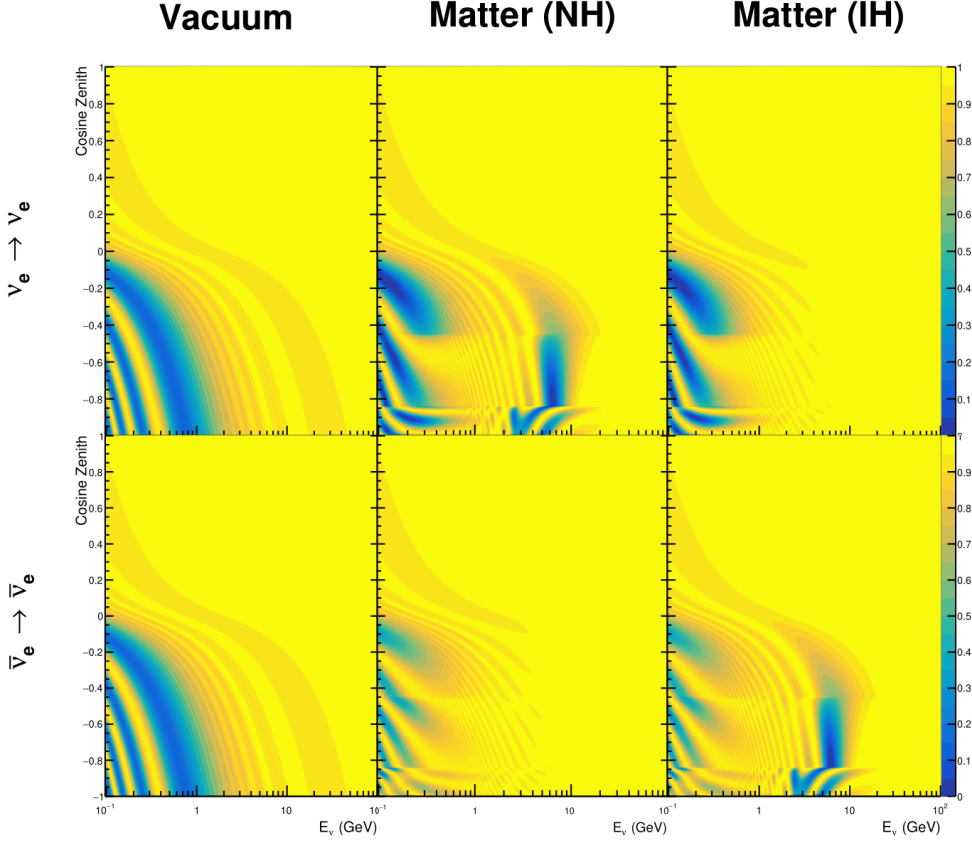


Figure 7.4: An illustration of the matter-induced effects on the oscillation probability, given as a function of neutrino energy and zenith angle. The top row of panels gives the $P(\nu_e \rightarrow \nu_e)$ oscillation probability and the bottom row illustrates the $P(\bar{\nu}_e \rightarrow \bar{\nu}_e)$ oscillation probability. The left column highlights the oscillation probability in a vacuum, whereas the middle and right column represents the oscillation probabilities when the four-layer fixed density PREM model is assumed. All oscillation probabilities assume the “Asimov A” set given in Table 2.2, but importantly, the right column sets an inverted mass hierarchy. The “matter resonance” effects at $E_\nu \sim 5\text{GeV}$ can be seen in the $P(\nu_e \rightarrow \nu_e)$ for normal mass hierarchy and $P(\bar{\nu}_e \rightarrow \bar{\nu}_e)$ for inverted hierarchy.

that need to be made should be applied. The $\nu_e \rightarrow \nu_{e,\mu,\tau}$ (and antineutrino equivalent) oscillations can be ignored for beam neutrinos as the $\nu_e/\bar{\nu}_e$ fluxes are approximately two orders of magnitude smaller than the corresponding $\nu_\mu/\bar{\nu}_\mu$ flux. Furthermore, as the peak neutrino energy of the beam is well below the threshold for charged current tau production ($E_\nu = 3.5\text{GeV}$ [50], only a small proportion of the neutrinos produced in the beam have the required energy. For the few neutrinos that have sufficient energy, the oscillation probability is very

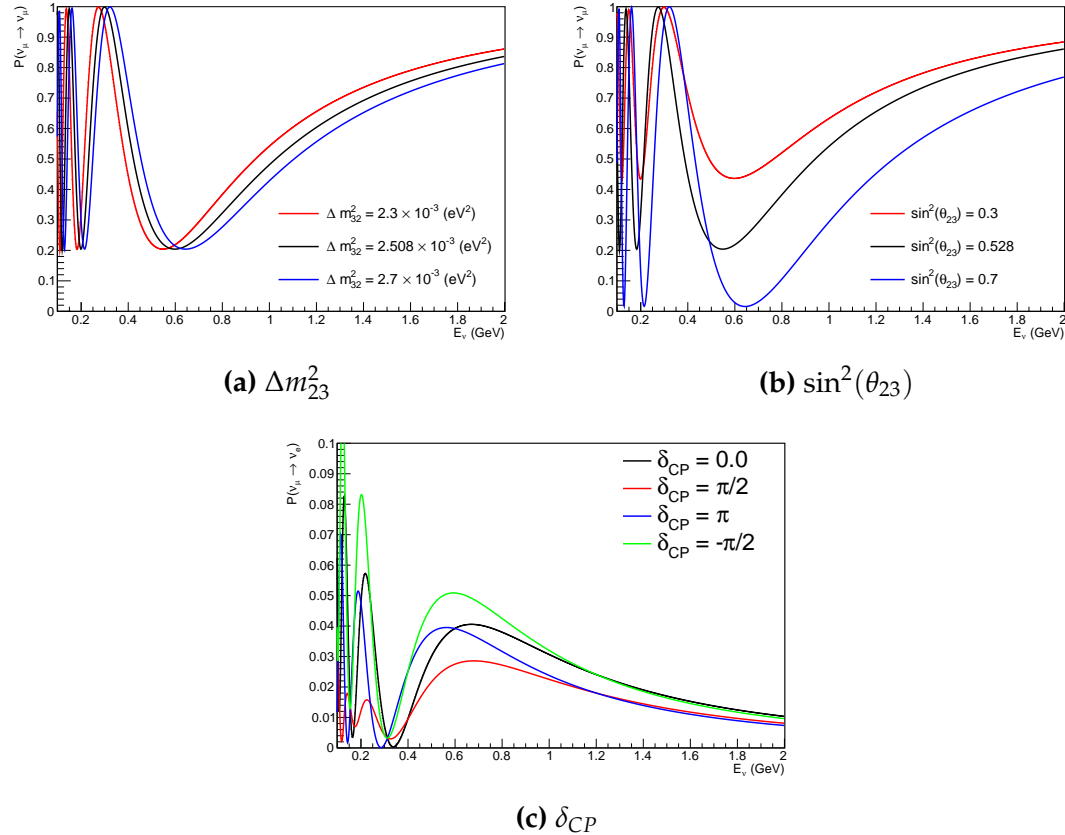


Figure 7.5: The oscillation probability for beam neutrino events given as a function of neutrino energy. All oscillation parameters assume the “Asimov A” set given in Table 2.2 unless otherwise stated. Each panel represents a change in one of the oscillation parameters whilst keeping the remaining parameters fixed.

small due to the short baseline. Whilst these approximations have been made for the beam neutrinos, the atmospheric flux of ν_e is of the same order of magnitude as the ν_μ flux and the energy distribution of atmospheric neutrinos extends well above the tau production threshold.

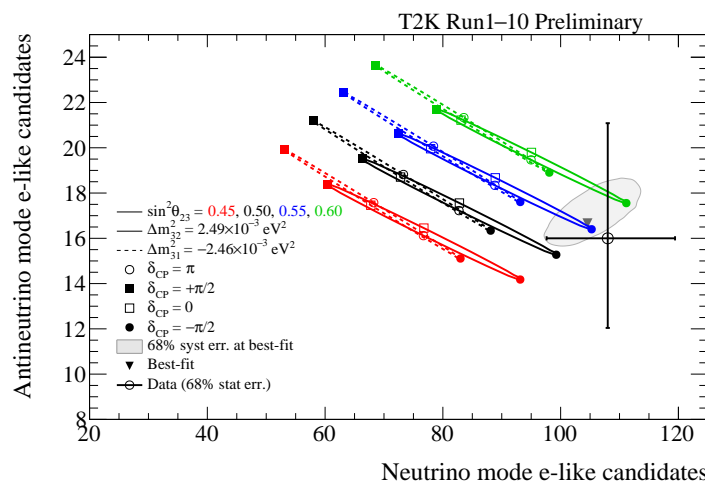


Figure 7.6: The number of electron-like events in the FHC and RHC operating mode of the beam, as a function of the oscillation probabilities. Both normal hierarchy (Solid) and inverse hierarchy (Dashed) values of Δm_{23}^2 are given.

7.2 Treatment of Fast Oscillations

As shown in Figure 7.7, atmospheric neutrino oscillations have a significantly more complex structure for upgoing neutrinos with energy below 1GeV. This is because the L/E dependence of the oscillation probability in this region induces rapid variations for small changes in L or E . As discussed in section 7.1, this is also the region in which atmospheric neutrinos have sensitivity to δ_{CP} . In practice, the direction of the neutrino is inferred from the direction of the final state particles traveling in the detector, which can be poor for low-energy neutrino interactions. This creates a distinct difference from the beam neutrinos where the position of the source is very precisely known.

As a consequence of the unresolvable structure, an event rate consistent with the averaged oscillation probability is observed in the subGeV upgoing region. This creates a computational problem: A significantly large amount of Monte Carlo statistics would be required to accurately predict the number of events if Monte Carlo averaging was the only technique used. This section describes the ‘sub-sampling’ approach developed for this analysis and compares it to the methodology used within the SK-only analysis.

The official SK-only analysis uses the osc3++ oscillation parameter fitter [208]. To perform the fast oscillation averaging, it uses a ‘nearest-neighbour’ technique. For a given Monte Carlo neutrino event, the nearest twenty Monte Carlo neighbours in reconstructed lepton momentum and zenith angle are found and a distribution of their neutrino energies is built. The RMS, σ , of this distribution is then used to compute an average oscillation probability for the given neutrino Monte Carlo event.

For the i^{th} event, the oscillation weight is calculated as

$$W_i = \frac{1}{5}P(E_i, \bar{L}_i) + \frac{1}{5}\sum_{\beta=-1,-0.5,0.5,1} P(E_i + \beta\sigma_i, L_\beta), \quad (7.3)$$

where $P(E, L)$ is the oscillation probability calculation for neutrino energy E and path length L and the two path lengths, \bar{L}_i and L_β are discussed below.

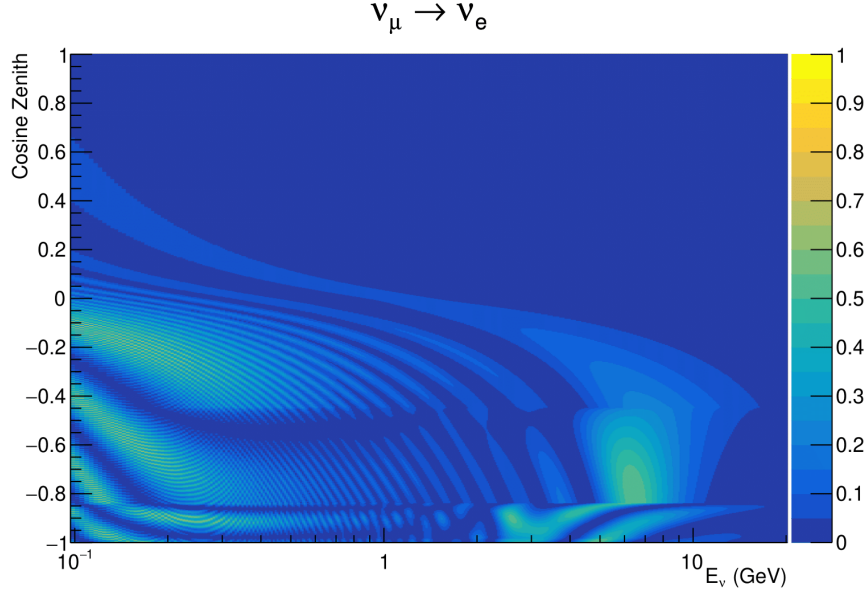


Figure 7.7: The oscillation probability $P(\nu_\mu \rightarrow \nu_e)$, given as a function of neutrino energy and zenith angle, which highlights an example of the “fast” oscillations in the sub-GeV upgoing region.

All of the oscillation probability calculations are performed with a fixed zenith angle such that the same density profile is used.

The uncertainty in the production height is controlled by using an “average” production height, \bar{L}_i , which represents the average path length computed using twenty production heights taken from the Honda flux model’s prediction [44]. For a given event, the production heights are sampled in steps of 5% of their cumulative distribution function. L_β values are similarly calculated but instead use different combinations of four production heights,

$$\begin{aligned}
 L_{-1.0} &= \frac{1}{4}L(45, 50, 55, 60), \\
 L_{-0.5} &= \frac{1}{4}L(35, 40, 65, 70), \\
 L_{+0.5} &= \frac{1}{4}L(25, 30, 75, 68), \\
 L_{+1.0} &= \frac{1}{4}L(15, 20, 85, 89).
 \end{aligned} \tag{7.4}$$

This averaging technique works because of the inference between the zenith angle and the reconstructed direction of final state particles in the detector. For

low-energy neutrinos, where the resolution of the true neutrino direction is poor, σ_i will be large, resulting in significant averaging effects. Contrary to this, the inferred direction of high-energy neutrinos will be much closer to the true value, meaning that σ_i will be smaller, culminating in small averaging effects.

In practice, these calculations are performed prior to the fit as only oscillation parameters at fixed points are considered. The MCMC technique used in this thesis requires oscillation probabilities to be evaluated at arbitrary parameter values, not known *a priori*. Calculating the five oscillation probabilities per event required by the SK technique is computationally infeasible, so a different averaging technique is used. However, the concept of the averaging technique can be taken from it.

To perform a similar averaging as the SK analysis, a sub-sampling approach using binned oscillograms has been devised. A coarsely binned oscillogram is defined in $\cos(\theta_Z)$ and E_ν . For a given set of oscillation parameters, a single oscillation probability will be assigned to each coarse bin. This value will then apply to all Monte Carlo events which fall into that bin. To assign these oscillation probabilities, the probability is calculated at $N \times N$ points on a grid within a particular bin. This ensemble of oscillation probabilities is averaged to define the coarse bin's oscillation probability, assuming a flat prior in E_ν and $\cos(\theta_Z)$. Figure 7.8 illustrates the $N = 2$ example where the assigned value to a coarse bin is the average of the four fine bins which fall in that coarse bin. Whilst the coarse bin edges do not have to be linear on either axis, the sub-division of the fine bins is linear over the range of a coarse bin.

The coarse binning is defined with 67×52 bins in true neutrino energy \times cosine zenith. It is picked to be identical to that provided in [210]. In general, the binning is logarithmically spaced in neutrino energy but has some hand-picked bin edges around the matter resonance to smoothly increased the bin density. This is to avoid smearing this region which can be well sampled by the Monte Carlo. The cosine zenith binning is approximately linearly spaced across the

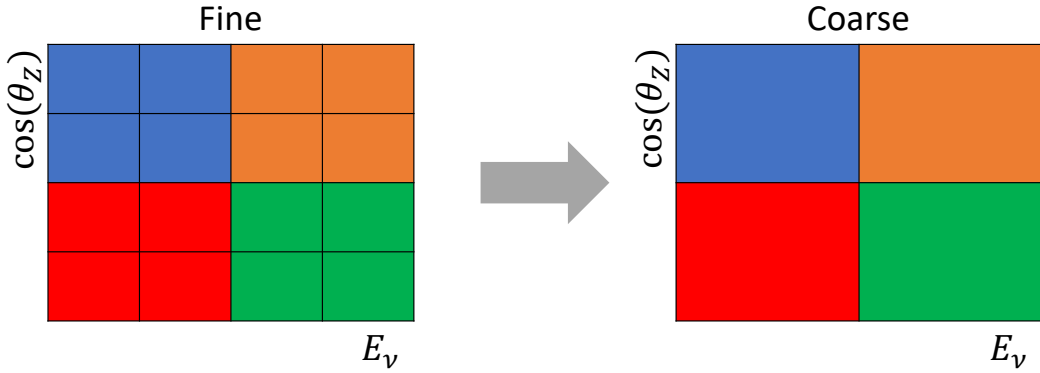


Figure 7.8: Illustration of the averaging procedure for $N = 2$. The oscillation probabilities calculated on the finer left binning are averaged to obtain the oscillation probabilities in the coarser right binning. These averaged oscillation probabilities with the coarser binning are then applied to each event during the fit.

allowable range but the values of layer transitions are hit precisely: -0.8376 (core-mantle) and -0.4464 (mantle/transition zone). Bins are spread further apart for downgoing events as this is a region unaffected by the fast oscillation wavelengths and reduces the total number of calculations required to perform the calculation.

The choice of N is justified based on two studies. Firstly, the variation of event rates of each sample is studied as a function of N . For a given set of oscillation parameters thrown from the PDG prior constraints (detailed in Table 2.1), the oscillation probabilities are calculated using a given value of N . Each sample is re-weighted and the event rate is stored. The value of N is scanned from 1, which corresponds to no averaging, to 19, which corresponds to the largest computationally viable subdivision binning. The event rate of each sample at large N is expected to converge to a stationary value due to the fine binning fully sampling the small-scale structure. Figure 7.9 illustrates this behaviour for the SubGeV_elike_0dcy sample for 9 different throws of the oscillation parameters.

Denoting the event rate for one sample for a given throw t at each N by λ_t^N , the average over all considered N values ($\bar{\lambda}_t = \frac{1}{24} \sum_{N=1}^{24} \lambda_t^N$) is computed. The variance in the event rate at each N is then calculated as

$$\text{Var}[\lambda^N] = \frac{1}{N_{\text{throws}}} \sum_{t=1}^{N_{\text{throws}}} (\lambda_t^N - \bar{\lambda}_t)^2 - \left[\frac{1}{N_{\text{throws}}} \sum_{t=1}^{N_{\text{throws}}} (\lambda_t^N - \bar{\lambda}_t) \right]^2. \quad (7.5)$$

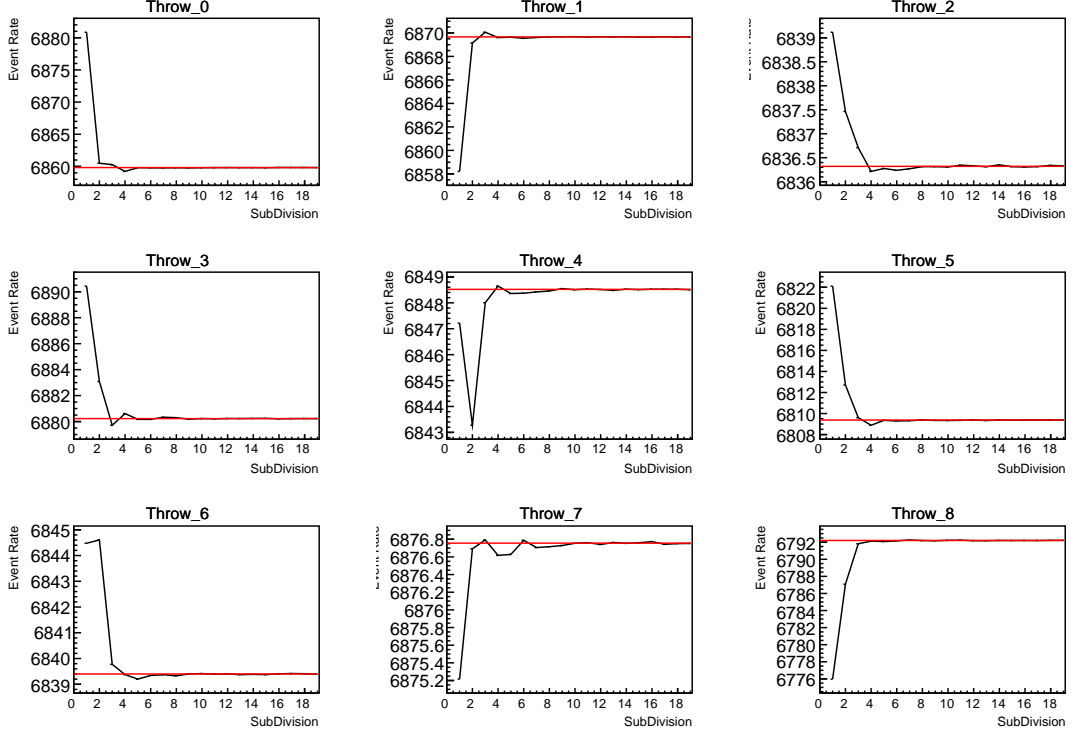


Figure 7.9: Event rate of the SubGeV_elike_0dcy sample as a function of the number of sub-divisions per coarse bin. Each subplot represents the event rate of the sample at a different oscillation parameter set thrown from the PDG priors detailed in Table 2.1. The red line in each subplot represents the mean of the event rate over the different values of sub-divisions for that particular oscillation parameter throw.

In practice the following procedure is undertaken. For a particular throw, the difference between the event rate at a particular choice of N and the mean of the distribution is calculated. This is illustrated in Figure 7.10. This value is then calculated for all the 2000 throws, generating a distribution of $\lambda_t^N - \bar{\lambda}_t$. This is repeated for each of the values of N considered within this study. The distributions of this value, for $N = \{1, 5\}$, are given in Figure 7.11. As expected, the distribution gets narrower and tends towards zero for the higher values of N .

The aim of the study is to find the lowest value of N such that this variance is below 0.001. This utilises the width of the distributions given in Figure 7.11. This is the typical threshold used by T2K fitters to validate systematic implementation so has been set as the same criteria. The results of this study for each atmospheric sample used within this thesis are illustrated in Figure 7.12 for 2000 throws of the oscillation parameters. As can be seen, the variance is below

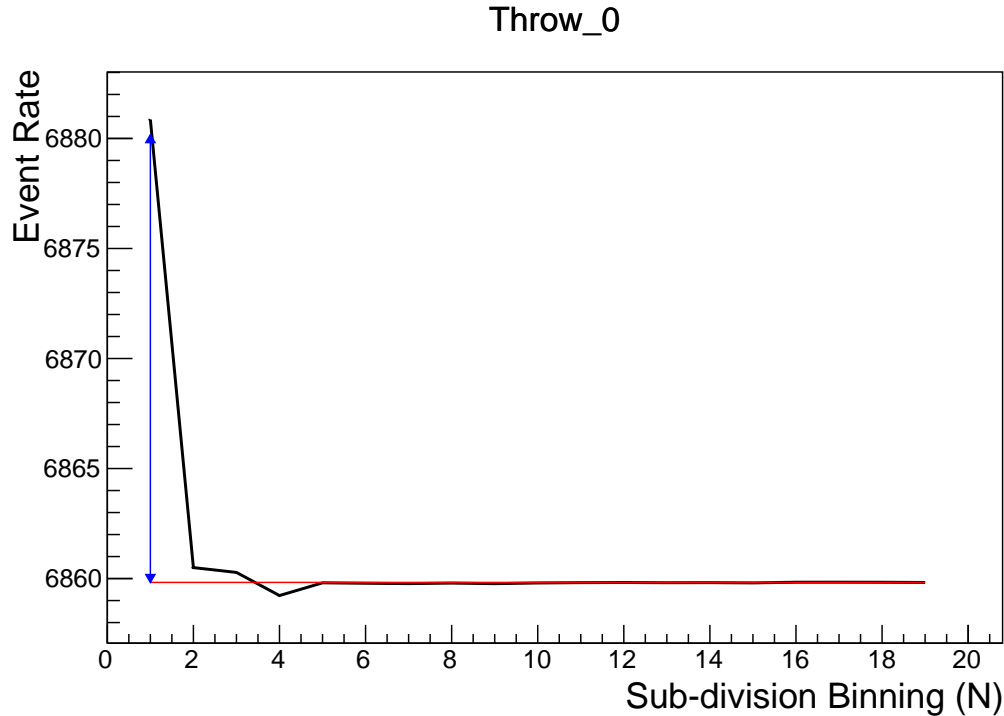


Figure 7.10: Event rate of the SubGeV_elike_0dcy sample, for a particular oscillation parameter throw, as a function of the number of sub-divisions, N , per coarse bin. The difference between the mean event rate (red), $\bar{\lambda}$, and the event rate at $N = 1$, $\lambda^{N=1}$ is defined as $\lambda^N - \bar{\lambda}$ and illustrated by the blue arrow.

the threshold at $N = 10$, and is driven primarily by the SubGeV_mulike_1dcy and SubGeV_elike_0dcy samples.

The second study to determine the value of N is as follows. The likelihood for each sample is computed against an Asimov data set created with Asimov A oscillation parameters (Table 2.2). Following Equation 7.5, the variance of the log-likelihood over all considered N is computed. The results are shown in Figure 7.13.

A choice of $N = 10$ sub-divisions per coarse bin has a variance in both event rate and log-likelihood residuals less than the required threshold of 0.001. The largest value of the likelihood variance is of order 10^{-7} , corresponding to an error on the log-likelihood of about 3×10^{-4} which is small enough to be negligible for the oscillation analysis.

Figure 7.14 illustrates the effect of the smearing using $N = 10$. The fast oscil-

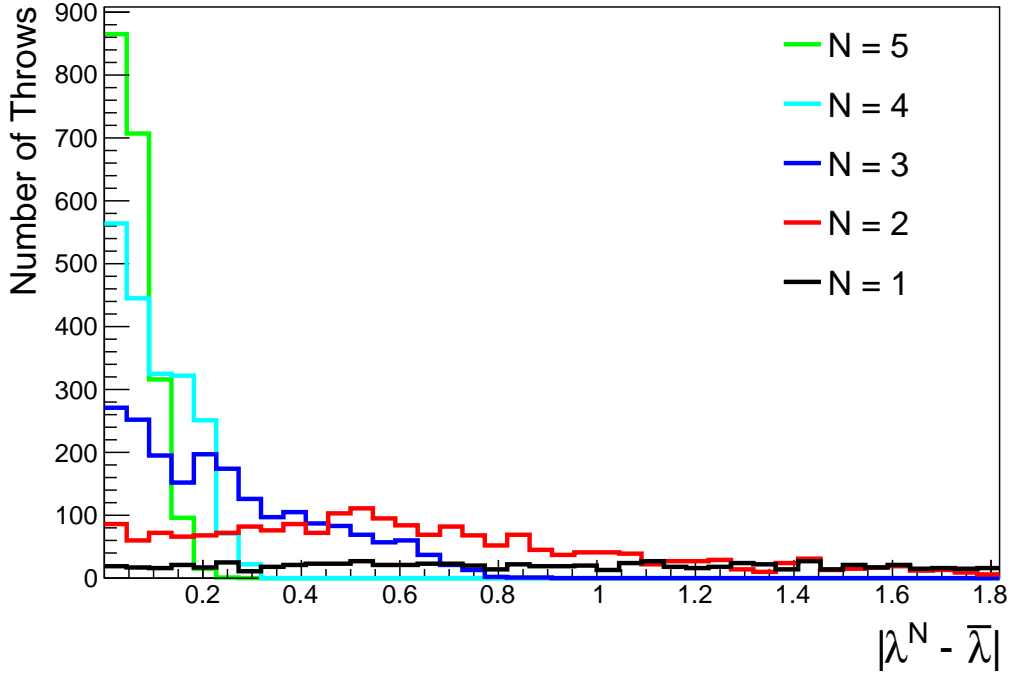


Figure 7.11: The distribution of $\lambda^N - \bar{\lambda}$ for various values of N . As expected, the distribution gets narrower for larger values of N .

lations in the sub-GeV upgoing region have been replaced with a normalisation effect whilst the large matter resonance structure remains.

7.3 Calculation Engine

As previously discussed in section 7.2, the calculation of oscillation probabilities is performed at run-time. Consequently, the time per calculation is crucial for fit performance. The initial fitting framework used for this analysis was developed with ProbGPU [211]. This is a GPU-only implementation of the prob3 engine [212]. It is primarily designed for neutrino propagation in a beam experiment (single layer of constant density) with the atmospheric propagation code not being used prior to the analysis in this thesis.

Another engine, CUDAProb3 [213], has been interfaced with the fitting framework used in this analysis. It has been specifically optimised for atmospheric neutrino oscillation calculation so does not contain the code to replace the

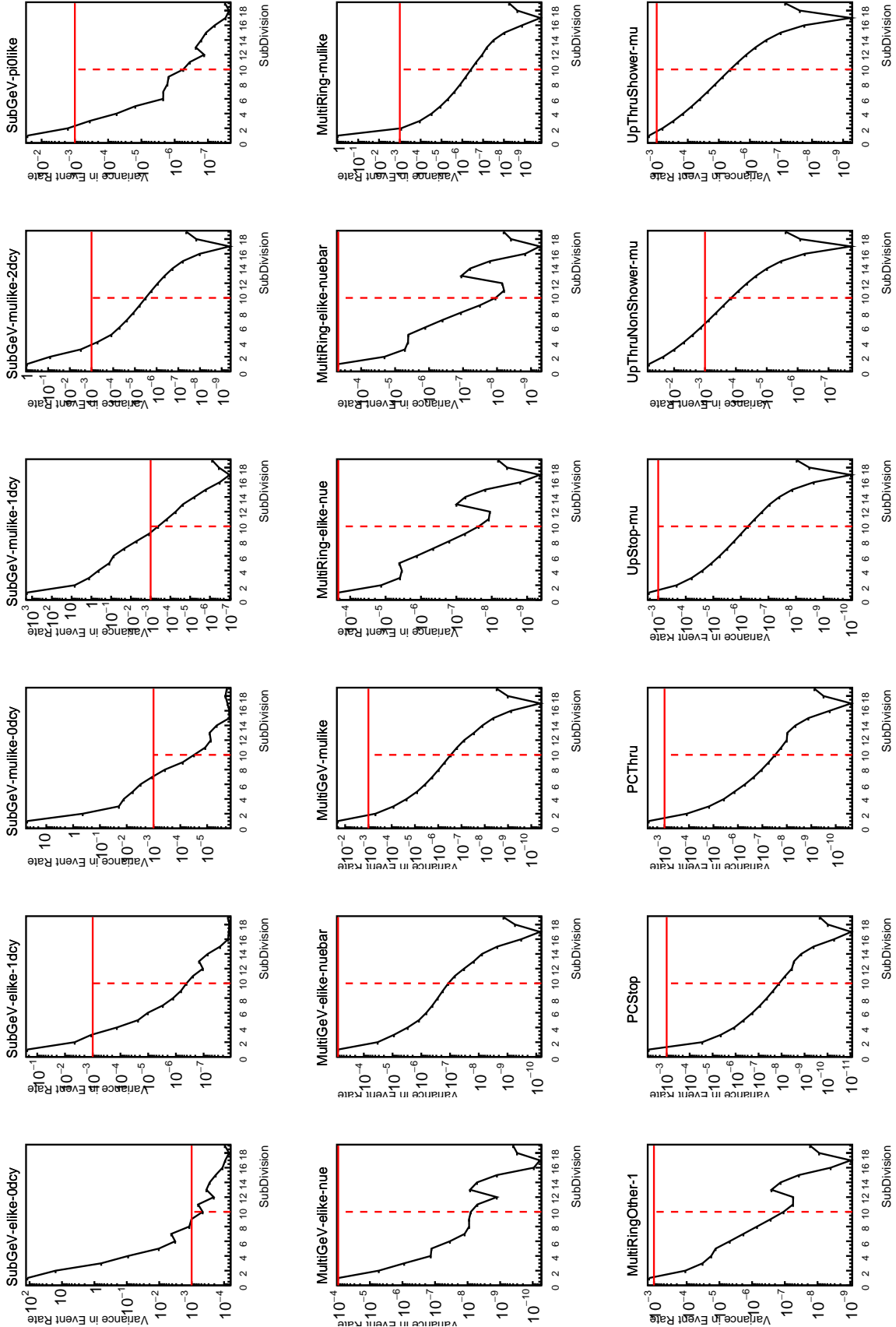


Figure 7.12: Variance of event rate for each atmospheric sample as a function of the number of sub-divisions per coarse bin. The solid red line indicates the 0.1% threshold and the dashed red line indicates the variance at a sub-division $N = 10$.

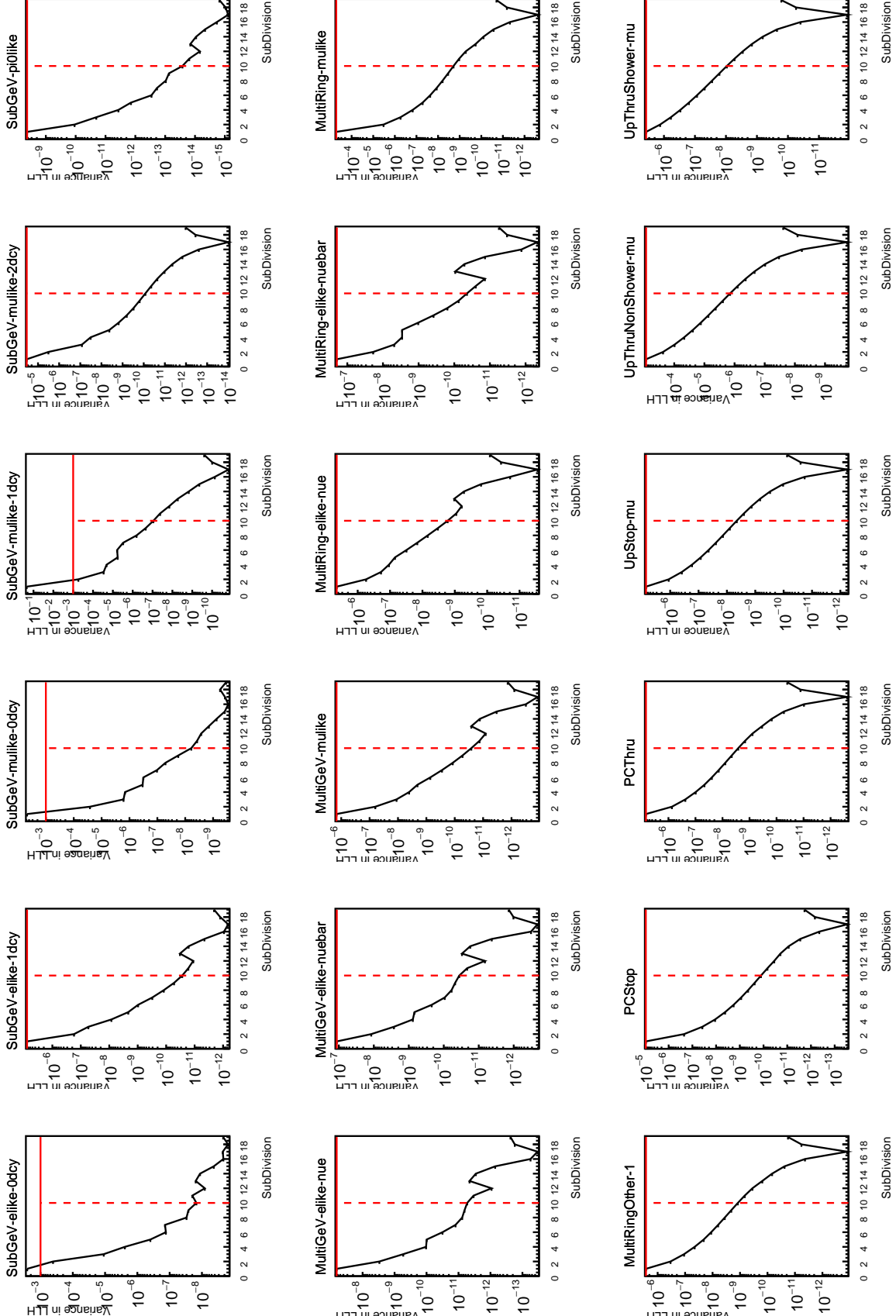


Figure 7.13: Variance of sample likelihood, when compared to ‘Asimov data’ set at Asimov A, for each atmospheric sample as a function of the number of sub-divisions per coarse bin. The solid red line indicates the 0.1% threshold and the dashed red line is a graphical indication of the variance at a sub-division $N = 10$.

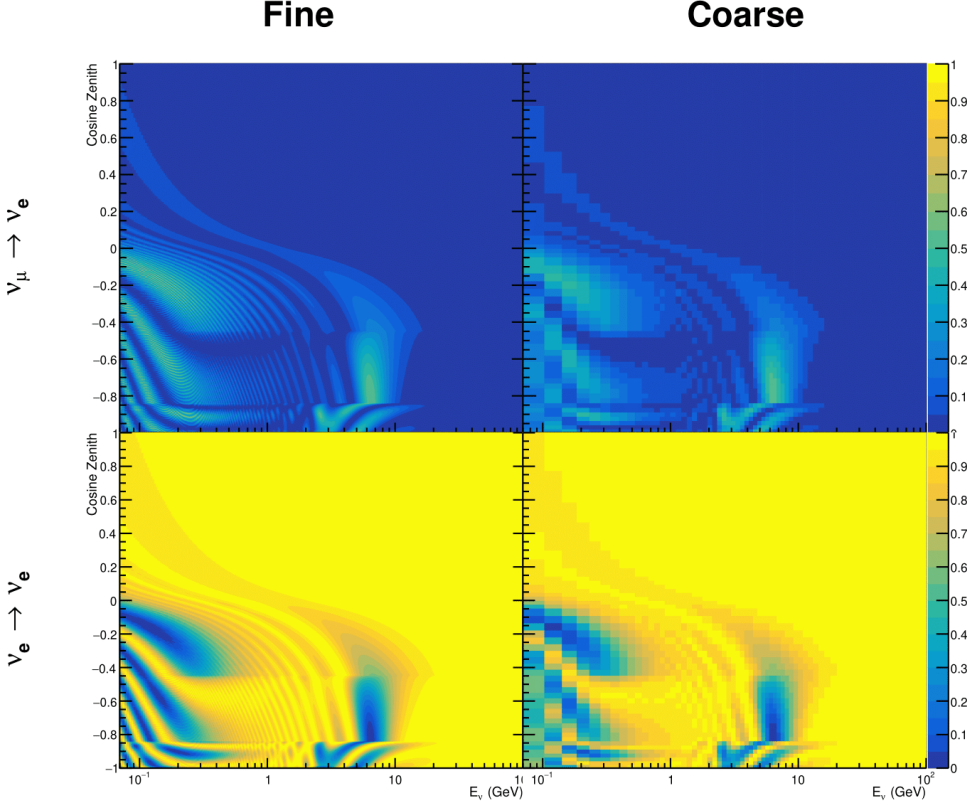


Figure 7.14: The oscillation probability, $P(\nu_\mu \rightarrow \nu_e)$ (top row) and $P(\nu_e \rightarrow \nu_e)$ (bottom row), given as a function of neutrino energy and zenith angle. The left column gives the “fine” binning used to calculate the oscillation probabilities and the right column illustrates the “coarse” binning used to reweight the Monte Carlo events. The fine binning choice is given with $N = 10$, which was determined to be below the threshold from Figure 7.12 and Figure 7.13.

beam oscillation calculation. The engine utilises object-orientated techniques as compared to the functional implementation of ProbGPU. This allows the energy and cosine zenith arrays to be kept on GPU memory, rather than having to load these arrays onto GPU memory for each calculation. Reducing the memory transfer between CPU and GPU significantly reduces the time required for calculation. This can be seen in Figure 7.15, where the GPU implementation of CUDAProb3 is approximately three times faster than the ProbGPU engine.

Another significant advantage of CUDAProb3 is that it contains a CPU multithreaded implementation which is not possible with the ProbGPU or prob3

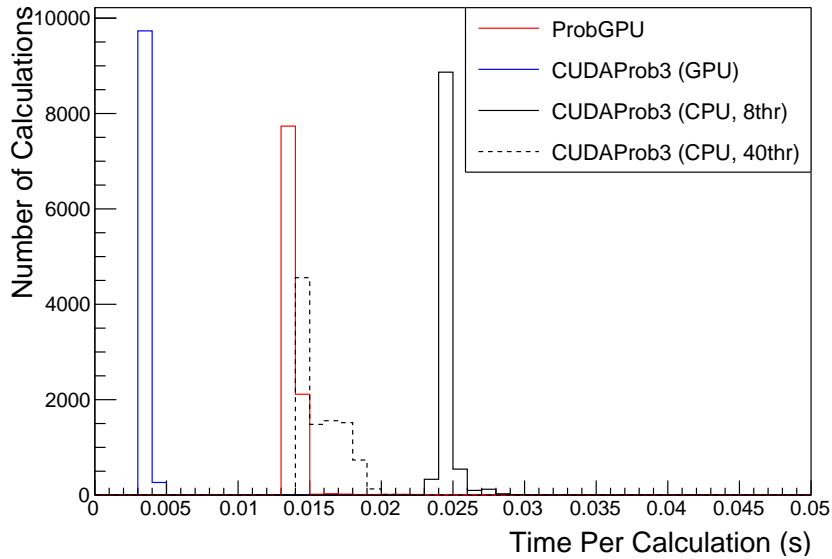


Figure 7.15: The calculation time taken to both calculate the oscillation probabilities and fill the “coarse” oscillograms, following the technique given in section 7.2, for the CUDAProb3 and ProbGPU (Red) calculation engines. CUDAProb3 has both a GPU (Blue) and CPU (Black) implementation, where the CPU implementation is multithreaded. Therefore, 8-threads (solid) and 40-threads (dashed) configurations have been tested. Prob3, which is a CPU single-thread implementation has a mean step time of 1.142s.

engines. This eliminates the requirement for GPU resources when submitting jobs to batch systems. As illustrated in Figure 7.15, the calculation speed depends on the number of available threads. Using 8 threads (which is typical of the batch systems being used) is approximately twice as slow as the ProbGPU engine implementation, but would allow the fitting framework to be run on many more resources. This fact is utilised for any SK-only fits but GPU resources are required for any fits which include beam samples due to the ProbGPU requirement. Based on the benefits shown by the implementation in this section, efforts are being placed into including linear propagation for beam neutrino propagation into the engine [214].

7.4 Matter Density Profile

For an experiment observing neutrinos propagating through the Earth, a model of the Earth’s density profile is required. The model used within this analysis is

based on the Preliminary Reference Earth Model (PREM) [209], as illustrated in Figure 7.1. Table 7.1 documents the density and radii of the layers used within the constant density approximaton used by the SK-only analysis [208]. The density measurements provided in the PREM model are provided in terms of mass density, whereas neutrino oscillations are sensitive to the electron number density. This value can be computed as the product of the chemical composition, or the Z/A value, and the mass density of each layer. Currently, the only way to measure the chemical composition value for layers close to the Earth's core is through neutrino oscillations. The chemical composition of the upper layers of the Earth's Mantle and the Transition zone is well known due to it being predominantly pyrolite which has a chemical composition value of 0.496 [215]. The chemical composition dial for the core layers is set to a value of 0.468, as calculated in [216]. As this value is lesss well known, it is assigned a Gaussian error with a standard deviation equivalent to the difference in chemical composition in core and mantle layers. Figure 7.16 illustrates the effect of moving from the $Z/A = 0.5$ method which is used in the official SK-only analysis to these more precise values.

The beam oscillation probability in this thesis uses a baseline of 295km, density $2.6\text{g}/\text{cm}^3$, and chemical composition 0.5 as is done by the official T2K-only analysis [217].

For a neutrino with given E_ν , $\cos(\theta_Z)$, the oscillation probability calculation engine must be passed a list of the matter regions that the neutrino traversed, with the path length and fixed density in each region. However, a neutrino passing through the earth experiences a range of radii, and thus a range of densities, in each region. In the SK-only analysis, the earth density model used is piecewise-constant, thereby ignoring this effect. For this thesis, the density values for the calculation engine are found by averaging the earth density along the neutrino's path,

$$\langle \rho \rangle_i = \frac{1}{t_{i+1} - t_i} \int_{t_i}^{t_{i+1}} \rho(t) dt \quad (7.6)$$

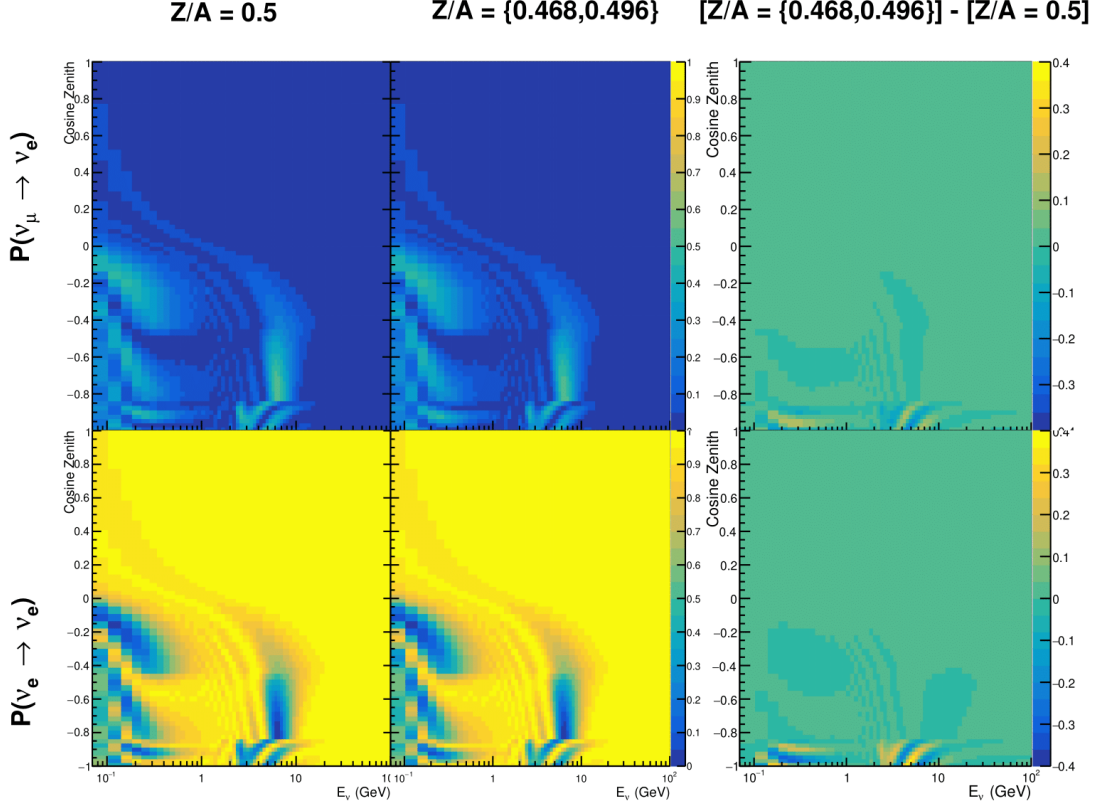


Figure 7.16: The oscillation probability, $P(\nu_\mu \rightarrow \nu_e)$ (top row) and $P(\nu_e \rightarrow \nu_e)$ (bottom row), given as a function of neutrino energy and zenith angle. The left column gives probabilities where the constant $Z/A = 0.5$ approximation which is used in the official SK-only analysis. The middle column gives the probabilities where $Z/A = [0.468, 0.498]$ values are used, as given in Table 7.1. The right column illustrates the difference in oscillation probability between the two different techniques.

where t_i are the intersection points between each layer and t is the path length of the trajectory across the layer. This leads to an improved approximation. For this averaging, the simplification of the PREM model developed in [218] is used. The layers of the prem model are combined into four to reduce calculation time, with a quadratic fit to each section. This fit was not performed by the author of the thesis and is documented in [210]. The coefficients of the quadratic fit to each layer are given in Table 7.2 with the final distribution illustrated in Figure 7.17. The quadratic approximation is clearly much closer to the PREM model as compared to the constant density approximation.

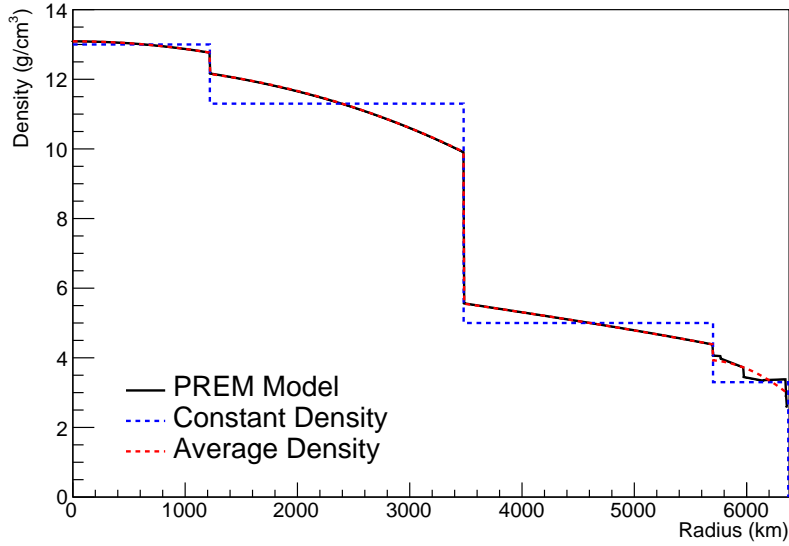


Figure 7.17: The density of the Earth given as a function of the radius, as given by the PREM model (Black), the constant density four-layer approximation (Blue), as used in the official SK-only analysis, and the quadratic approximation of the PREM model (Red).

Layer	Outer Radius [km]	Density [g/cm ³]
Inner Core	1220	$13.09 - 8.84x^2$
Outer Core	3480	$12.31 + 1.09x - 10.02x^2$
Lower Mantle	5701	$6.78 - 1.56x - 1.25x^2$
Transition Zone	6371	$-50.42 + 123.33x - 69.95x^2$

Table 7.2: The quadratic polynomial fits to the PREM model for four assumed layers of the PREM model. The fit to calculate the coefficients is given in [210], where $x = R / R_{\text{Earth}}$.

The effect of using the quadratic density per $\cos(\theta_Z)$ model is highlighted in Figure 7.18. The slight discontinuity in the oscillation probability around $\cos(\theta_Z) \sim -0.45$ in the fixed density model, which is due to the transition to mantle layer boundary, has been reduced. This is expected as the difference in the density across this boundary is significantly smaller in the quadratic density model as compared to the constant density model. Whilst the difference in density across the other layer transitions is reduced, there is still a significant difference. This means the discontinuities in the oscillation probabilities remain but are significantly reduced. However, as the quadratic density approximation matches the PREM model well in this region, these discontinuities are due to the Earth model rather than an artifact of the oscillation calculation.

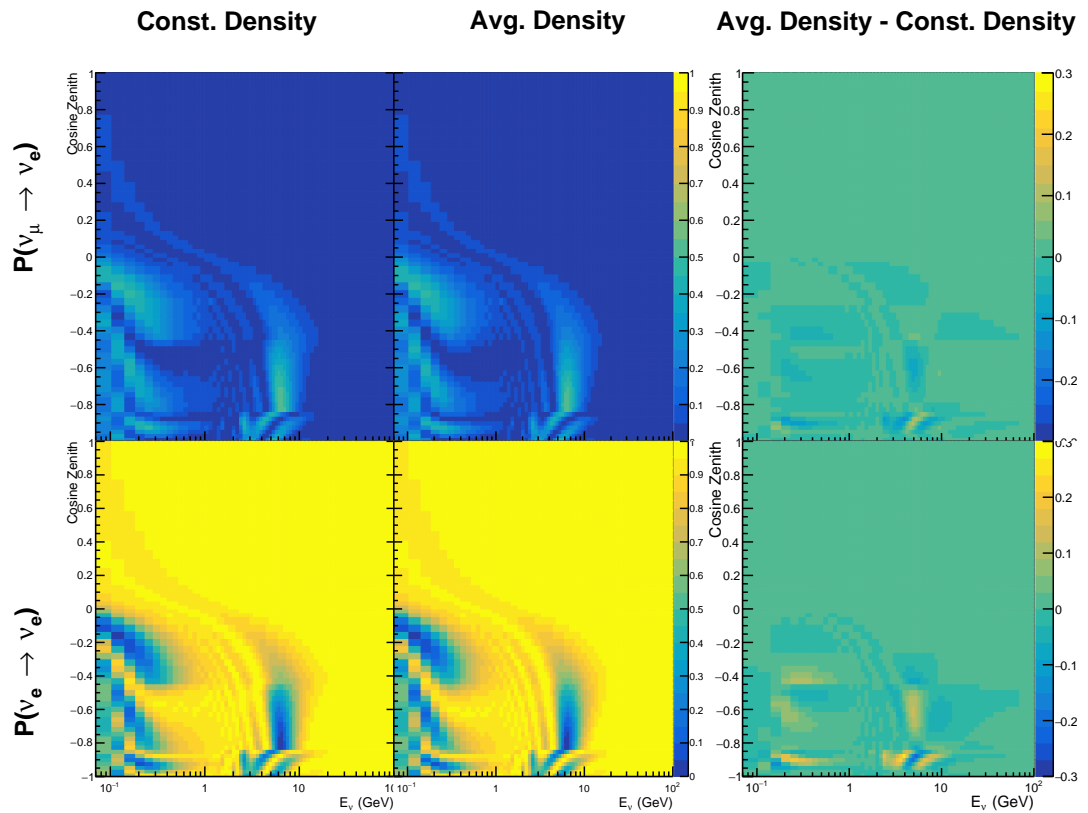


Figure 7.18: The oscillation probability, $P(\nu_\mu \rightarrow \nu_e)$ (top row) and $P(\nu_e \rightarrow \nu_e)$ (bottom row), given as a function of neutrino energy and zenith angle. The left column gives probabilities where the four-layer constant density approximation is used. The middle column gives the probabilities where the density is integrated over the trajectory, using the quadratic PREM approximation, for each $\cos(\theta_Z)$ is used. The right column illustrates the difference in oscillation probability between the two different techniques.

7.5 Production Height Averaging

As discussed in section 7.1, the height at which the cosmic ray flux interacts in the atmosphere is not known on an event-by-event basis. The production height can vary from the Earth's surface to $\sim 50\text{km}$ above that. The SK-only analysis methodology (described in section 7.2) for including the uncertainty on the production height is to include variations from the Honda model when pre-calculating the oscillation probabilities prior to the fit. This technique is not possible for this analysis which uses continuous oscillation parameters that can not be known prior to the fit. Consequently, an analytical averaging technique was developed in [210]. The author of this thesis was not responsible for the derivation of the technique but has performed the implementation and validation of the technique for this analysis alone.

Using the 20 production heights per Monte Carlo neutrino event, provided as 5% percentiles from the Honda flux model, a production height distribution $p_j(h|E_\nu, \cos \theta_Z)$ is built for each neutrino flavour $j = \nu_e, \bar{\nu}_e, \nu_\mu, \bar{\nu}_\mu$. In practice, a histogram is filled with 20 evenly spaced bins in production height h between 0 and 50km. The neutrino energy and cosine zenith binning of the histogram is the same as that provided in section 7.2. The average production height, $\bar{h} = \int dh \frac{1}{4} \sum_j p_j(h|E_\nu, \cos(\theta_Z))$, is calculated. The production height binning of this histogram is then translated into $\delta t(h) = t(\bar{h}) - t(h)$, where $t(h)$ is the distance travelled along the trajectory.

For the i^{th} traversed layer, the transition amplitude, $D_i(t_{i+1}, t_i)$, is computed. The time-ordered product of these is then used as the overall transition amplitude via

$$A(t_{n+1}, t_0) = D_n(t_{n+1}, t_n) \dots D_1(t_2, t_1) D_0(t_1, t_0), \quad (7.7)$$

where,

$$\begin{aligned} D_n(t_{n+1}, t_n) &= \exp[-iH_n(t_{n+1} - t_n)] \\ &= \sum_{k=1}^3 C_k \exp[i a_k (t_{n+1} - t_n)] \end{aligned} \quad (7.8)$$

is expressed as a diagonalised time-dependent solution to the Schrodinger equation. The 0th layer is the propagation through the atmosphere and is the only term that depends on the production height. Using the substitution $t_0 = t(\bar{h}) - \delta t(h)$, it can be shown that

$$D_0(t_1, t_0) = D_0(t_1, \bar{h}) D_0(\delta t). \quad (7.9)$$

Thus Equation 7.7 becomes

$$\begin{aligned} A(t_{n+1}, t_0) &= D_n(t_{n+1}, t_n) \dots D_1(t_2, t_1) D_0(t_1, \bar{h}) D(\delta t) \\ &= A(t_{n+1}, \bar{h}) \sum_{k=1}^3 C_k \exp[i a_k \delta t], \\ &= \sum_{k=1}^3 B_k \exp[i a_k \delta t]. \end{aligned} \quad (7.10)$$

The oscillation probability averaged over production height is then calculated as

$$\begin{aligned} \bar{P}(\nu_j \rightarrow \nu_i) &= \int d(\delta t) p_j(\delta t | E_\nu, \cos \theta_Z) P(\nu_j \rightarrow \nu_i) \\ &= \int d(\delta t) p_j(\delta t | E_\nu, \cos \theta_Z) A(t_{n+1}, t_0) A^*(t_{n+1}, t_0) \\ &= \sum_{km} (B_k)_{ij} (B_m)_{ij}^* \int d(\delta t) p_j(\delta t | E_\nu, \cos \theta_Z) \exp[i(a_k - a_m)\delta t] \end{aligned} \quad (7.11)$$

In practice, implementation in CUDAProb3 [213] is relatively straightforward as the majority of these terms are already calculated in the standard oscillation calculation. Figure 7.19 illustrates the results of the production height averaging. As expected, the main effect is observed in the low-energy downward-going and horizontal-going events. Upward-going events have to travel the radius of the Earth, $R_E = 6371\text{km}$, where the production height uncertainty is a small fraction of the total path length.

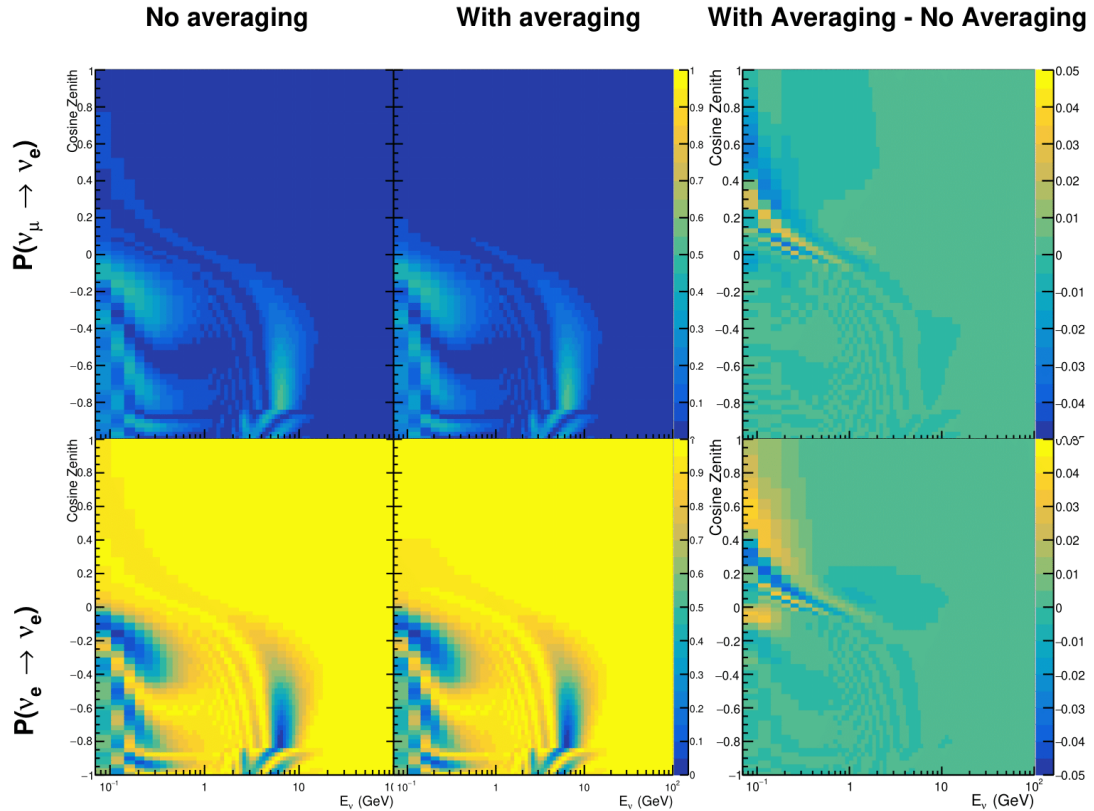


Figure 7.19: The oscillation probability, $P(\nu_\mu \rightarrow \nu_e)$ (top row) and $P(\nu_e \rightarrow \nu_e)$ (bottom row), given as a function of neutrino energy and zenith angle. The left column gives probabilities where a fixed production height of 25km is used. The middle column gives the probabilities where the production height is analytically averaged. The right column illustrates the difference in oscillation probability between the two different techniques.

8

Oscillation Analysis

The MaCh3 framework used throughout this thesis has been validated through many tests. The code which handles the beam far detector samples was validated by comparison to the 2020 T2K analysis. The sample event rates and log-likelihood scans generated by the framework used within this thesis were compared to those from the T2K analysis by the author of this thesis. Variations of the sample spectra were compared at $\pm 1, 3\sigma$ and good agreement was found in all cases. A similar study, led by Dr. C. Wret was used to validate the near detector portion of the code. The implementation of the atmospheric samples within MaCh3 was cross-checked by the author of this thesis against the P-Theta framework. Both fitters are provided the same inputs and thus act as self-validation. These validations compared the event rate and log-likelihood scans. Documentation of all the above validations can be found in [203].

8.1 Monte Carlo Prediction

Using the three sets of dial values defined in subsection 6.4.3, the predicted event rates for each sample are defined in Table 8.1. Both the oscillated event rates assuming Asimov A oscillation parameters (defined in Table 2.2) and the un-oscillated event rates are given.

Sample	Total Predicted Events					
	Generated		Pre-fit		Post-fit	
	Osc	UnOsc	Osc	UnOsc	Osc	UnOsc
SubGeV- <i>elike</i> -0dcy	7121.0	7102.6	6556.8	6540.0	7035.2	7015.7
SubGeV- <i>elike</i> -1dcy	704.8	725.5	693.8	712.8	565.7	586.0
SubGeV- <i>mulike</i> -0dcy	1176.5	1737.2	1078.6	1588.1	1182.7	1757.1
SubGeV- <i>mulike</i> -1dcy	5850.7	8978.1	5351.7	8205.1	5867.0	9009.9
SubGeV- <i>mulike</i> -2dcy	446.9	655.2	441.6	647.7	345.9	505.6
SubGeV- <i>pi0like</i>	1438.8	1445.4	1454.9	1461.1	1131.1	1136.2
MultiGeV- <i>elike</i> -nue	201.4	195.6	201.1	195.3	202.6	196.7
MultiGeV- <i>elike</i> -nuebar	1141.5	1118.3	1060.7	1039.5	1118.5	1095.7
MultiGeV- <i>mulike</i>	1036.7	1435.8	963.1	1334.1	1015.2	1405.9
MultiRing- <i>elike</i> -nue	1025.1	982.2	1026.8	984.3	1029.8	986.4
MultiRing- <i>elike</i> -nuebar	1014.8	984.5	991.0	962.0	1008.9	978.5
MultiRing- <i>mulike</i>	2510.0	3474.4	2475.6	3425.8	2514.6	3480.4
MultiRingOther-1	1204.5	1279.1	1205.8	1280.3	1207.4	1281.0
PCStop	349.2	459.2	338.4	444.7	346.8	456.1
PCThrus	1692.8	2192.5	1661.5	2149.8	1689.2	2187.8
UpStop-mu	751.2	1295.0	739.7	1271.6	750.4	1293.0
UpThruNonShower-mu	2584.4	3031.6	2577.9	3019.4	2586.8	3034.0
UpThruShower-mu	473.0	488.6	473.2	488.7	473.8	489.4
FHC1Rmu	328.0	1409.2	301.1	1274.7	345.1	1568.0
RHC1Rmu	133.0	432.3	122.7	396.2	135.0	443.9
FHC1Re	84.6	19.2	77.4	18.2	93.7	19.7
RHC1Re	15.7	6.4	14.6	6.1	15.9	6.3
FHC1Re1de	10.5	3.2	10.3	3.1	8.8	2.9

Table 8.1: The Monte Carlo prediction of each sample observed at SK used within this analysis. Three model parameter tunes are considered, as defined in subsection 6.4.3. The oscillated predictions assumed Asimov A oscillation parameters provided in Table 2.2.

Generally, samples which target CCQE interaction modes observe a decrease in prediction when using the pre-fit dial values. This is in accordance with the Monte Carlo being produced assumed $M_A^{QE} = 1.21\text{GeV}$ whilst the pre-fit dial value should be $M_A^{QE} = 1.03\text{GeV}$, as suggested by [201]. Furthermore, the predicted event rates of samples which target CCRES interaction modes is significantly reduced when considering the post-ND fit. This follows the observations in subsection 6.4.3. The strength of the accelerator neutrino experiment can also be seen in the remarkable difference between the oscillated and unoscillated predictions in the FHC1Rmu and RHC1Rmu samples. There is a very obvious decrease in the expected event rate between the two predictions which is not as clearly

observed in the atmospheric samples. This is due to the fact that the beam energy is tuned to the maximum disappearance probability, which is not the case for the naturally generated atmospheric neutrinos.

8.2 Likelihood Calculation

This analysis performs a joint oscillation parameter fit of the ND280, and the SK atmospheric samples.

Once the Monte Carlo predictions of each beam and atmospheric sample has been built, following from chapter 6, a likelihood needs to be constructed. This is done by comparing the Monte Carlo prediction to “data”. The data can consist of either an Asmiov Monte Carlo prediction, which is typically used for sensitivity studies, or real data. The Monte Carlo prediction is calculated at a particular point, $\vec{\theta}$, in the model parameter space, $N_i^{MC} = N_i^{MC}(\vec{\theta})$. Both the data and Monte Carlo spectra are binned, where the i^{th} bin content is represented by N_i^D and N_i^{MC} , respectively. The bin contents for the beam near detector, beam far detector and atmospheric samples are denoted with *ND*, *FD* and *Atm*, respectively. The binning index, i , runs over all the bins within the sample and all samples with that set. Taking the beam far detector samples as example, it would run over all the reconstructed neutrino energy bins in all samples (FHC1R μ , RHC1R μ , etc.). The likelihood calculation between data and Monte Carlo for a particular bin follows a Poisson distribution, where the data is treated as a fluctuation of the simulation.

Following the T2K analysis presented in [76], the likelihood contribution from the near detector also includes a Monte Carlo statistical uncertainty term, derived from the Barlow and Beeston statistical treatment [219, 220]. In addition to treating the data as a fluctuation of the Monte Carlo prediction, it includes a contribution from the likelihood that the generated simulation is a statistical fluctuation of the actual true simulation assuming infinite statistics. The technical

implementation of this additional likelihood term is documented in [192]. The term is defined as,

$$\frac{(\beta_i - 1)^2}{2\sigma_{\beta_i}^2}, \quad (8.1)$$

where β_i represents a scaling parameter for each bin i , which is a value based on the amount of Monte Carlo statistics in a bin [192]. $\sigma_{\beta_i} = \sqrt{\sum_i w_i^2 / N_i^{MC}}$, and $\sqrt{\sum_i w_i^2}$ represents the sum of the square of the weights of the Monte Carlo events which fall into bin i .

Additional contributions to the likelihood come from the variation of the systematic model parameters. For those parameters with well-motivated uncertainty estimates, a covariance matrix, V describes the prior knowledge of each parameter as well as any correlations between the parameters. Due to the technical implementation, a single covariance matrix describes each “block” of model parameters, e.g. beam flux systematics. For simplicity, the covariance matrix associated with the k^{th} block is denoted V^k . This substitution results in $\vec{\theta} = \sum_k^{N_b} \vec{\theta}^k$ and $V = \sum_k^{N_b} V^k$, for N_b number of blocks describing: oscillation parameters, beam flux, atmospheric flux, neutrino interaction, near detector, beam far detector and atmospheric far detector systematics detailed in section 6.4. The number of parameters in the k^{th} block is defined as $n(k)$.

The final likelihood term is defined as,

$$\begin{aligned} -\ln(\mathcal{L}) = & \quad (8.2) \\ & \sum_i^{\text{NDbins}} N_i^{\text{ND},MC}(\vec{\theta}) - N_i^{\text{ND},D} + N_i^{\text{ND},D} \times \ln \left[N_i^{\text{ND},D} / N_i^{\text{ND},MC}(\vec{\theta}) \right] + \frac{(\beta_i - 1)^2}{2\sigma_{\beta_i}^2} \\ & + \sum_i^{\text{FDbins}} N_i^{\text{FD},MC}(\vec{\theta}) - N_i^{\text{FD},D} + N_i^{\text{FD},D} \times \ln \left[N_i^{\text{FD},D} / N_i^{\text{FD},MC}(\vec{\theta}) \right] \\ & + \sum_i^{\text{Atmbins}} N_i^{\text{Atm},MC}(\vec{\theta}) - N_i^{\text{Atm},D} + N_i^{\text{Atm},D} \times \ln \left[N_i^{\text{Atm},D} / N_i^{\text{Atm},MC}(\vec{\theta}) \right] \\ & + \frac{1}{2} \sum_k^{N_b} \sum_i^{n(k)} \sum_j^{n(k)} (\vec{\theta}^k)_i (V^k)_{ij}^{-1} (\vec{\theta}^k)_j. \end{aligned}$$

This is the value determined at each step of the MCMC to build the posterior distribution, as discussed in chapter 4.

8.2.1 Likelihood Scans

Using the defintion of the likelihood presented in section 8.2, the response of each sample to a variation particular parameter can be studied. Figure 8.1 presents the variation of all the samples (beam and atmospheric) at SK. Each plot represents a “scan”, where a particular parameter is scanned in some range. The “data” being used within the definition of the likelihood equation is built using the Asimov A oscillation parameter values defined in Table 2.2 alongside the pre-fit dial values as discussed in subsection 6.4.3. Due to the caveat of fixed systematic parameters and correlations between oscillation parameters being ignored within these likelihood scans, the value of $\chi^2 \sim 1$ does not equate to the typical 1σ sensitivity. However, it does give an indication of which samples response the strongest to a variation in the oscillation parameters. The point at which the likelihood tends to zero illustrates the value of the parameter used to build the Asimov data prediction. The likelihood scans only include the sample response and ignore the penalty contribution term from the variation of the parameter.

The sensitivity to $\sin^2(\theta_{23})$ is mostly dominated by the beam samples although the atmospheric sample contribution is certainly non-negligible. The summed response over all atmospheric samples becomes comparable to that of the muon-like beam samples. Consequently, one would expect that the joint fit would become more sensitivity to $\sin^2(\theta_{23})$ than just T2K experiment alone. For this particular choice of asimov point, the only samples which respond to the $\sin^2(\theta_{13})$ parameter are the electron-like beam samples. Consequently, no increase in sensitivity beyond that of the T2K-only analysis is expected. Furthermore, the sensitivity of the beam sample is significantly weaker than the reactor constraint so the ‘reactor constraint’ prior will dominate any measurement it is included within. The Δm_{12}^2 and $\sin^2(\theta_{12})$ are not considered as there is simply no sensitivity in any sample considered within this analysis. The response

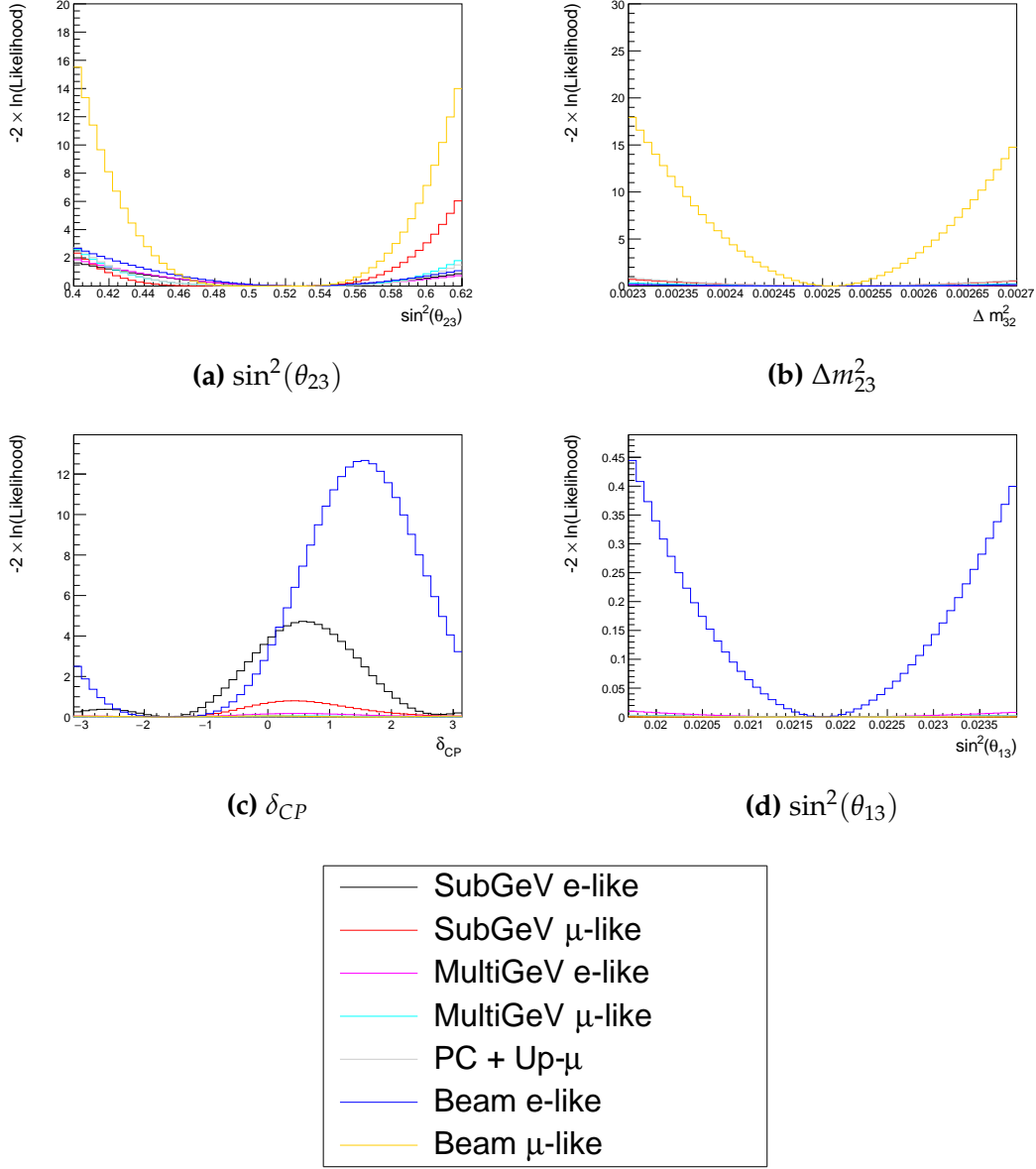


Figure 8.1: The response of the likelihood, as defined in section 8.2, illustrating the response of the samples to the oscillation parameters. Δm_{12}^2 and $\sin^2(\theta_{12})$ are negated because these samples have no sensitivity to those parameters. The Asimov data set is built using the pre-fit dial values assuming Asimov A oscillation parameters defined in Table 2.2.

to Δm_{23}^2 is much larger in beam samples, specifically for the μ -like samples, compared to atmospheric samples. This is to be expected as the beam neutrino energy can be specifically tuned to match the maximal disappearance probability.

The correlations between oscillation parameters induce marginalisation effects

within the response of the likelihood. That is to say, the response to δ_{CP} is affected by the choice of $\sin^2(\theta_{13})$ or $\sin^2(\theta_{23})$. The two-dimensional scans of the appearance ($\sin^2(\theta_{13}) - \delta_{CP}$) and disappearance ($\sin^2(\theta_{23}) - \Delta m_{23}^2$) parameters are illustrated in Figure 8.2 and Figure 8.3, respectively. The caveat of fixed systematic parameters and correlations between other oscillation parameters being neglected still apply.

The appearance log-likelihood scans show the distinct difference in how the beam and atmospheric samples respond. The beam samples have an approximately constant width of the 2σ and 3σ contours, throughout all ranges of δ_{CP} . The atmospheric samples response to δ_{CP} is very strongly correlated to the choice of $\sin^2(\theta_{13})$, with the strongest constraints around $\delta_{CP} \sim 1$. Consequently, this difference allows some of the degeneracy in a beam-only fit to be broken. Comparing the beam and joint fit log-likelihood scans, the 2σ continuous contour in δ_{CP} for beam samples is broken when the atmospheric samples are added. Furthermore, the width of the 3σ contours also becomes dependent upon the value of δ_{CP} . Whilst these are encouraging results for the joint fit, these are not sensitivity measurements as the systematic parameters are fixed and the correlation between oscillation parameters is neglected.

The disappearance log-likelihood scans in $\sin^2(\theta_{23}) - \Delta m_{23}^2$ space show the expected result when considering the one-dimensional scans already discussed. The uncertainty on the width of $|\Delta m_{32}^2|$ is mostly driven by the beam-only sensitivities. However, the width of this contour in the inverted mass region ($\Delta m_{32}^2 < 0$) is significantly reduced due to the ability of the atmospheric samples to select the correct mass hierarchy (these log-likelihood scans use the Asimov A oscillation probabilities which assumes true normal hierarchy). The width of the uncertainty in $\sin^2(\theta_{23})$ is also reduced compared to a beam-only analysis, with a further decrease in the inverted hierarchy region due to mass hierarchy determination.

The log-likelihood scans illustrated thus far only give the sensitivity of this analysis at a fixed asimov point, namely Asimov A defined in Table 2.2. Whilst

computational infeasible to run many fits at different asimov points, it is possible to calculate the log-likelihood response to different asimov data sets. Figure 8.4 and Figure 8.5 illustrate how the sensitivity changes for differing values of δ_{CP} and $\sin^2(\theta_{23})$, respectively, whilst the other oscillation parameters are fixed at Asimov A. Consequentially, the caveat of fixed systematic parameters and correlations between other oscillation parameters being neglected still applies.

To explain how these plots are made, consider Figure 8.4. This plot is built by considering multiple one-dimensional log-likelihood scans, each using the Asimov A oscillation parameter set but having a slightly differing value of δ_{CP} . Consequently, the results can be interpreted as vertical slices of the log-likelihood response made at different asimov points. The procedure starts by building an asimov ‘data’ prediction for a particular value of δ_{CP} taken from the x-axis. Then a likelihood value is calculated at every δ_{CP} point taken from the y-axis. This is then repeated for each point on the x-axis, such that a series of one dimensional likelihood scans are displayed in sequential order of δ_{CP} .

Figure 8.4 illustrates how the sensitivity to δ_{CP} is offset between the beam and atmospheric samples. This offset agrees with the one dimensional scan illustrated in Figure 8.1. Notably for the 1σ intervals, there are regions in the off-diagonal for which the beam and atmospheric samples have broken and discontinuous contours. For example, for the asimov point $\delta_{CP} = 0.$, the beam samples sensitivity would include two discontinuous regions which would be preferred: $\delta_{CP} = 0, \pi$. However the offset in δ_{CP} between these beam and atmospheric samples allow the joint fit to have increased sensitivity in these regions, thus mitigating the degeneracy. However, the 2σ intervals from the joint fit are more similar to the two independent sensitivities and the off-diagonal degeneracies mostly remain. This indicates that the joint fit has the strength to aide parameter determination but still can not entirely break the degeneracies in δ_{CP} at higher confidence levels.

Figure 8.5 illustrates the same study as above, although the value of δ_{CP} is fixed to the Asimov A parameter set whilst the value of $\sin^2(\theta_{23})$ is varied.

Due to the beam parameters and baseline being tuned to specifically target this oscillation parameter, the average sensitivity of the beam samples is stronger than the atmospheric samples. However, the degeneracy around maximal mixing ($\sin^2(\theta_{23}) = 0.5$) is significantly more peaked in the beam samples compared to the atmospheric samples. This behaviour is strengthened when considering the 2σ intervals, to the point where two distinct discontinuous regions of the 2σ intervals exist around the asimov point $\sin^2(\theta_{23}) \sim 0.41, 0.6$. Given the caveat of only considering log-likelihood scans, the joint analysis would mostly eliminate the discontinuous intervals in these regions.

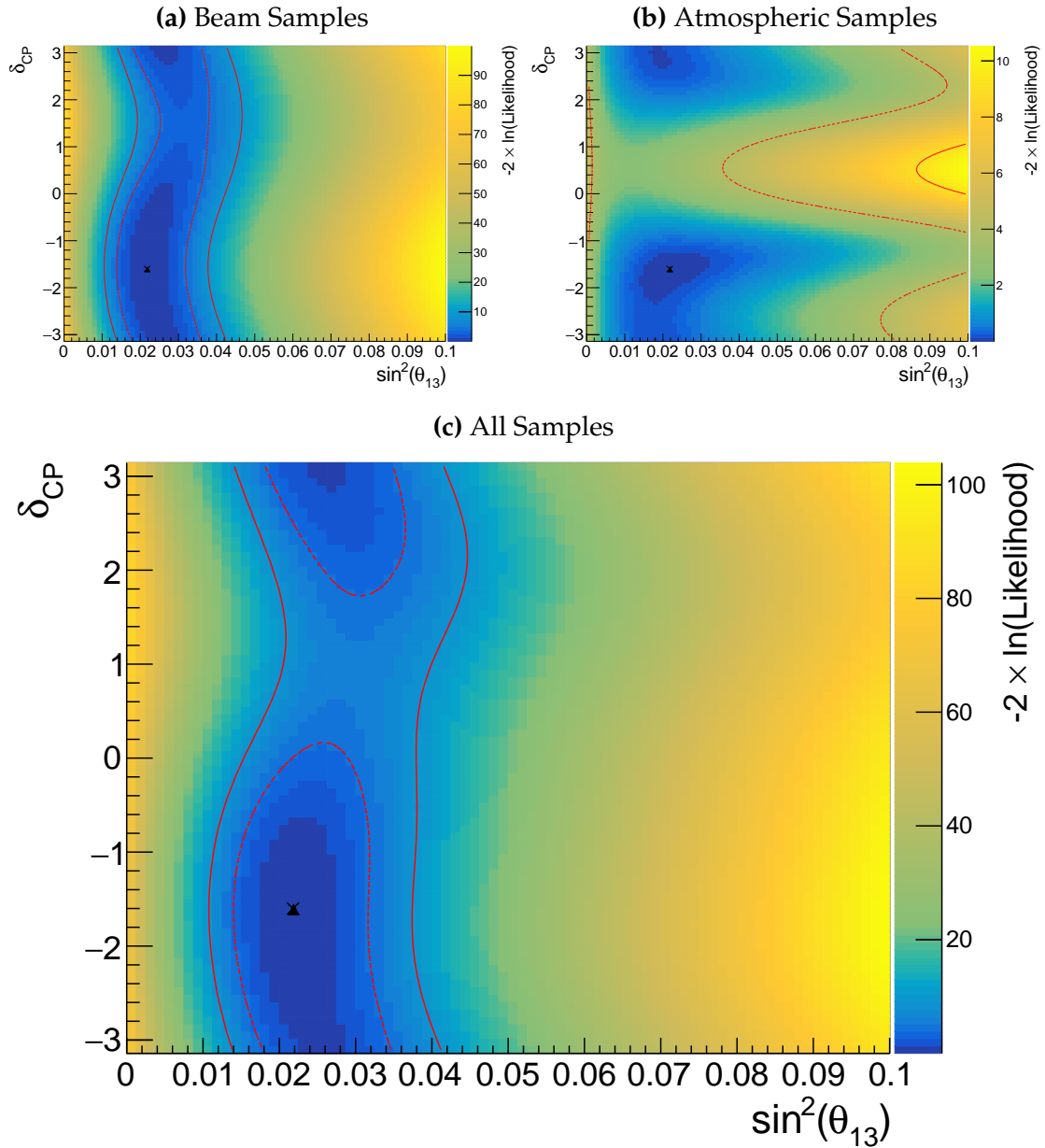


Figure 8.2: Two-dimensional log-likelihood scan of the appearance ($\sin^2(\theta_{13})$ - δ_{CP}) parameters showing the response of the beam samples (top), atmospheric samples (middle) and the summed response (bottom). The Asimov A oscillation parameters, defined in Table 2.2, are assumed to be the true point (Black Cross). The position of the smallest log-likelihood is highlighted with the triangle. Prior uncertainty terms of the oscillation parameters are neglected. The two(three) sigma contour levels are illustrated with the dashed(solid) red line.

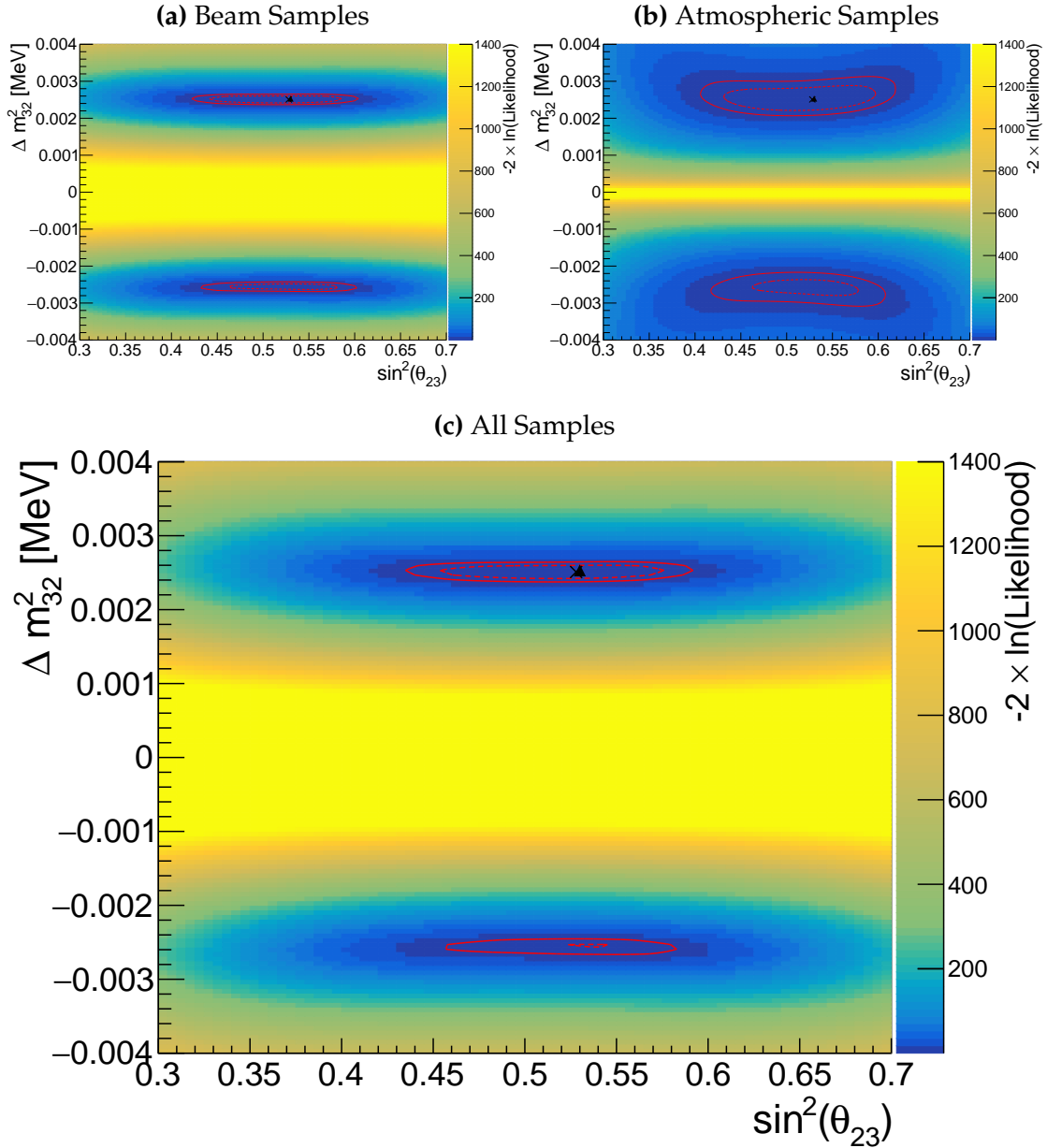


Figure 8.3: Two-dimensional log-likelihood scan of the disappearance ($\sin^2(\theta_{23})$ - Δm_{32}^2) parameters showing the response of the beam samples (top), atmospheric samples (middle) and the summed response (bottom). The Asimov A oscillation parameters, defined in Table 2.2, are assumed to be the true point (Black Cross). The position of the smallest log-likelihood is highlighted with the triangle. Prior uncertainty terms of the oscillation parameters are neglected. The two(three) sigma contour levels are illustrated with the dashed(solid) red line.

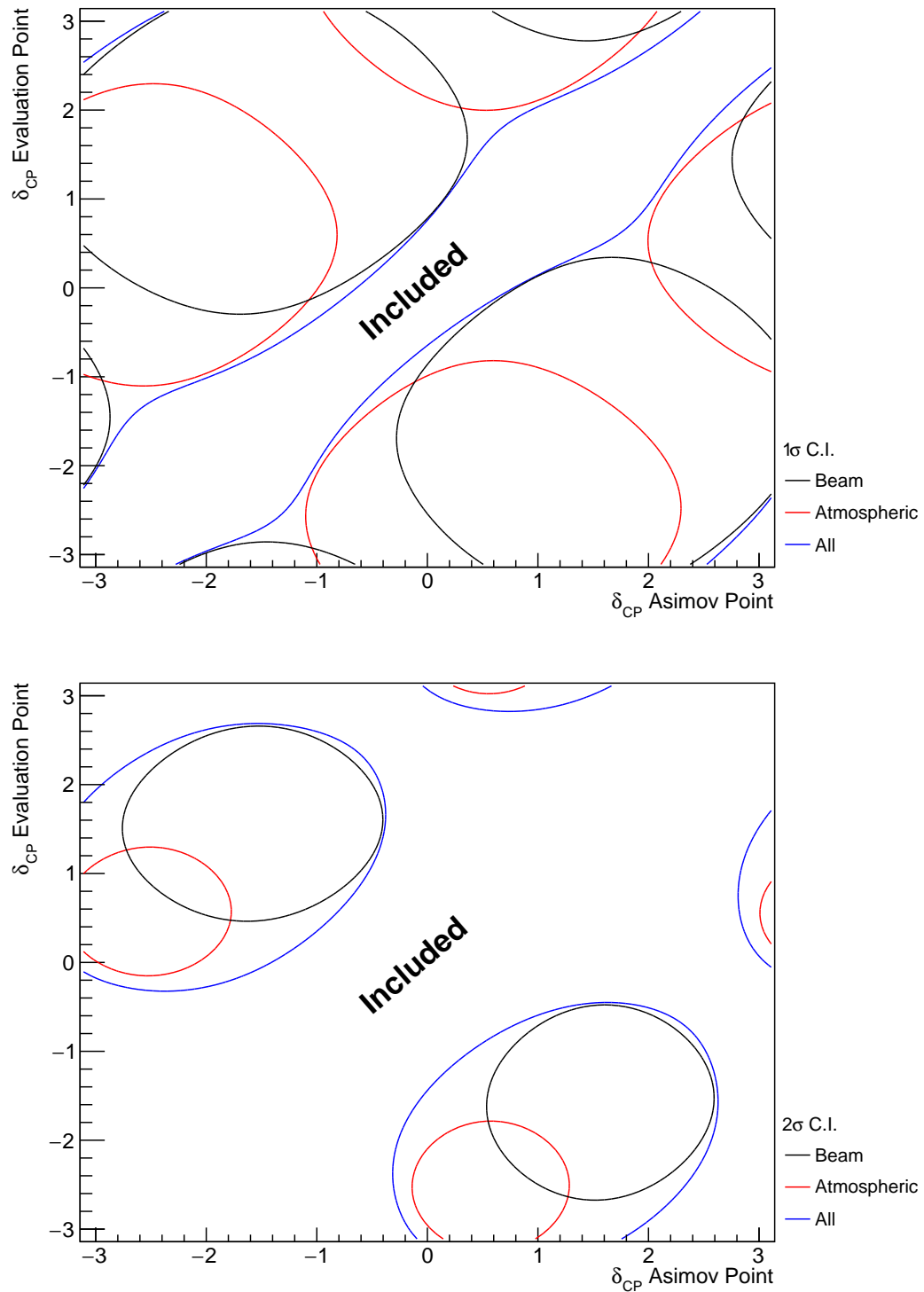


Figure 8.4: A series of one-dimensional log-likelihood scans over δ_{CP} , where an asimov data set is built for each value of δ_{CP} on the x-axis and the log-likelihood is evaluated for each value on the y-axis. The diagonal represents the minimum log-likelihood and defines the region included within the 1σ (Top) and 2σ (Bottom) confidence intervals. The beam (black) and atmospheric (red) samples are individually plotted and the joint fit (blue) is the sum of the two.

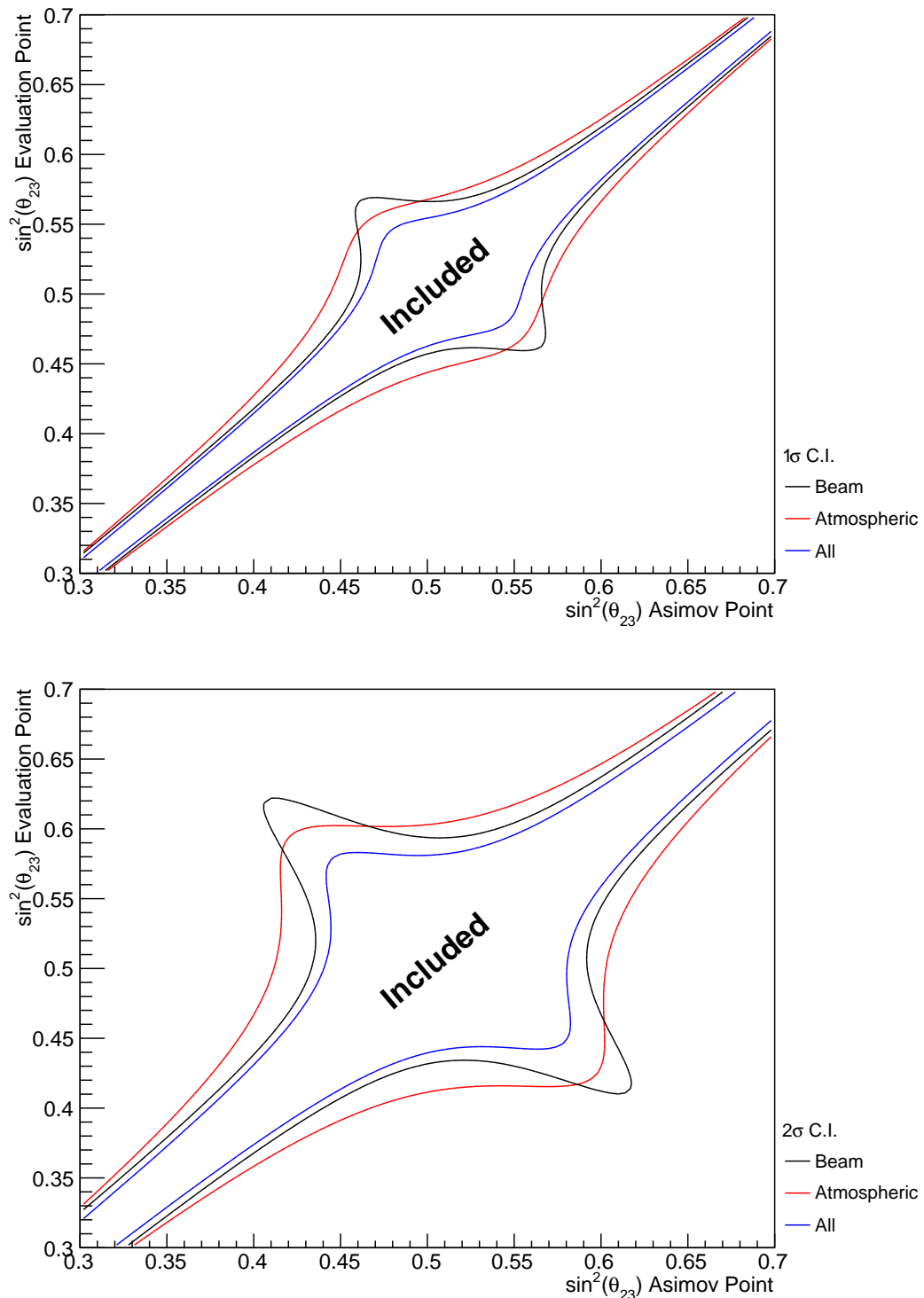


Figure 8.5: A series of one-dimensional log-likelihood scans over $\sin^2(\theta_{23})$, where an asimov data set is built for each value of $\sin^2(\theta_{23})$ on the x-axis and the log-likelihood is evaluated for each value on the y-axis. The diagonal represents the minimum log-likelihood and defines the region included within the 1σ (Top) and 2σ (Bottom) confidence intervals. The beam (black) and atmospheric (red) samples are individually plotted and the joint fit (blue) is the sum of the two.

Alongside oscillation parameters, the sensitivity to systematic parameters can also be studied. As some of these parameters are correlated between the beam and atmospheric events, the addition of the atmospheric samples can modify the near detector constraint producing results which could have differing constraints than a T2K-only analysis. Consequently, the relative strength of the response between beam and atmospheric samples has been compared for various systematic parameters in Figure 8.6. For example, the systematic parameter controlling the effective axial mass coupling in CCQE interactions, MAQE, is clearly dominated by the ND constraint. An example where the atmospheric samples response is approximately similar to the near detector constraint is the 2p2h_Cto0 normalisation systematic. There are also systematics which have no near detector constraint, for example the systematic parameters which describe the normalisation of the NC1Gamma and NCOther interaction modes. The atmospheric samples are significantly more sensitive to these systematics than the beam samples. As an example of how the atmospheric samples can help constrain systematic parameters used within the T2K-only analysis, the neutral current background events in beam electron-like samples will be considerably more constrained with the additional sensitivity observed in Figure 8.6. This would be expected to reduce the overall uncertainty on the beam electron-like event rates in the joint analysis compared to the beam-only studies.

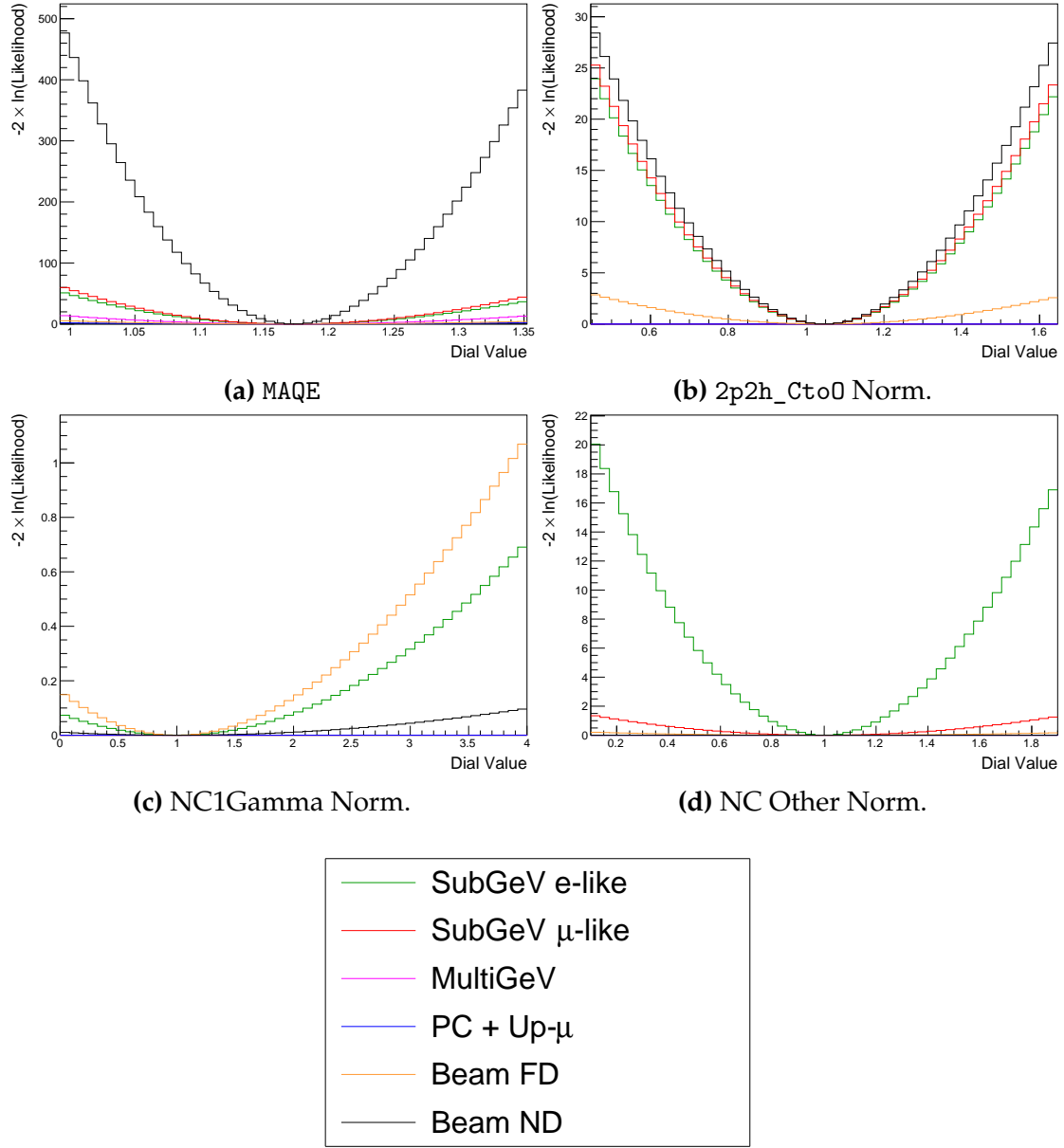


Figure 8.6: The response of the likelihood, as defined in section 8.2, illustrating the response of the samples to the various cross-section systematic parameters. The Asimov data set is built using the pre-fit dial values assuming Asimov A oscillation parameters defined in Table 2.2.

8.3 Sensitivities

The sensitivities of the joint T2K and SK oscillation analysis are presented in the form of Asimov fits. This technique builds a fake ‘data’ prediction of each sample’s spectra from the Monte Carlo, reweighted to a particular set of oscillation and systematic parameters. This prediction is then used as data in which to fit against. Whilst this results in unphysical non-integer data predictions, it eliminates statistical fluctuations from the data. Therefore, the results of a fit to Asimov data should not include any biases from statistical fluctuations. Furthermore, these results should produce posterior probability distributions consistent with the parameters which were used to make the data prediction. That is to say, the fit results should return the known parameters. Any biases seen would be attributed to correlations between each oscillation parameter and correlations between oscillation and systematic parameters. Consequently, Asimov fit results present the maximum precision at which the oscillation parameters could be measured to.

In practice, the asimov fits presented within this analysis are modified from the above definition. An asimov prediction of both beam and atmospheric far detector samples is fit whilst the true data is used for near detector samples. The asimov predictions at the far detector are built using the BANFF tuning (as discussed in section 3.2). This is equivalent to performing a far detector asimov fit using inputs from the BANFF data fit. Consequently, this allows the results to be cross-checked to the P-Theta analysis.

8.3.1 Atmospheric-Only Sensitivity Without Reactor Constraint

This section presents the results of an asimov fit using samples from the near detector and only atmospheric samples from the far detector. The results are marginalised over the nuisance parameters using the technique outlined in subsection 4.3.1. Each histogram displays the posterior probability density at each point within the parameter space. One dimensional histograms illustrate

the 1 , 2 and 3σ credible intervals, calculated using the technique discussed in subsection 4.3.2. For these fits, a flat penalty term is used for $\sin^2(\theta_{13})$ (i.e. the reactor constraint is not applied). The asimov data is generated assuming the AsimovA oscillation parameter set defined in Table 2.2 and the post-BANFF systematic parameter tune.

The posterior probability density in Δm_{23}^2 is given in Figure 8.7. This distribution includes steps in both the normal hierarchy (NH, $\Delta m_{32}^2 > 0$) and the inverse hierarchy (IH, $\Delta m_{32}^2 < 0$). The highest posterior density is found within the NH, which agrees with the asimov point. However, all of the credible intervals span both hierarchies. This is the result of marginalising over both hierarchies. If instead, only steps in the normal hierarchy were considered, the shape of the contours would change. The known oscillation parameter is 2.509×10^{-3} eV 2 , which is contained within the 1σ credible interval.

Without Reactor Constraint, Both Hierarchies

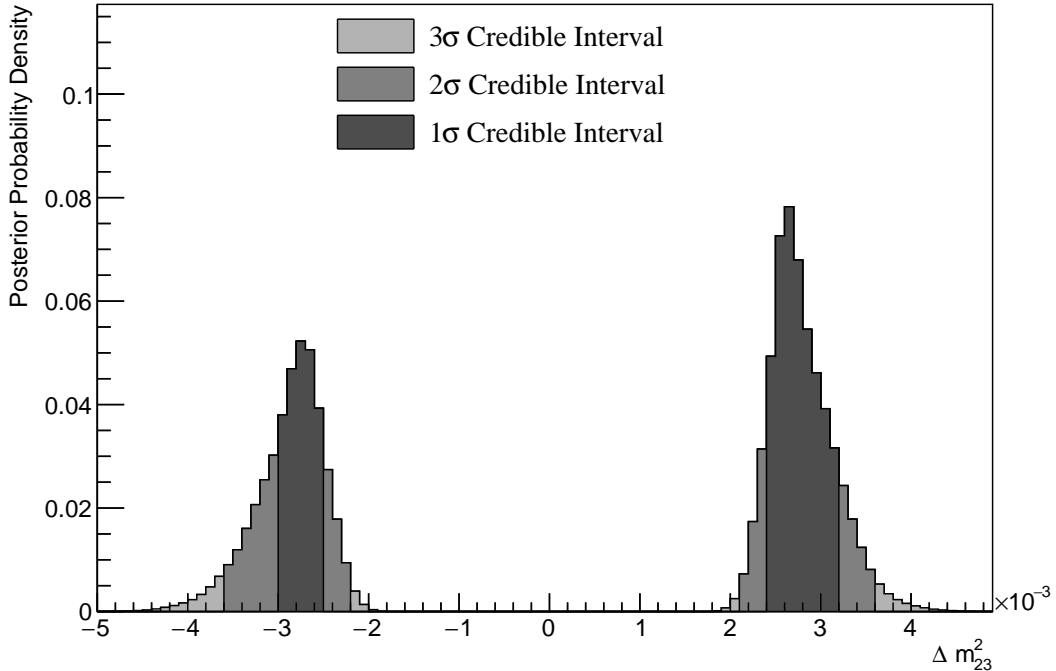


Figure 8.7

Following the discussion in subsection 4.3.3, the Bayes factor for hierarchy preference can be calculated by determining the fraction of steps which fall into

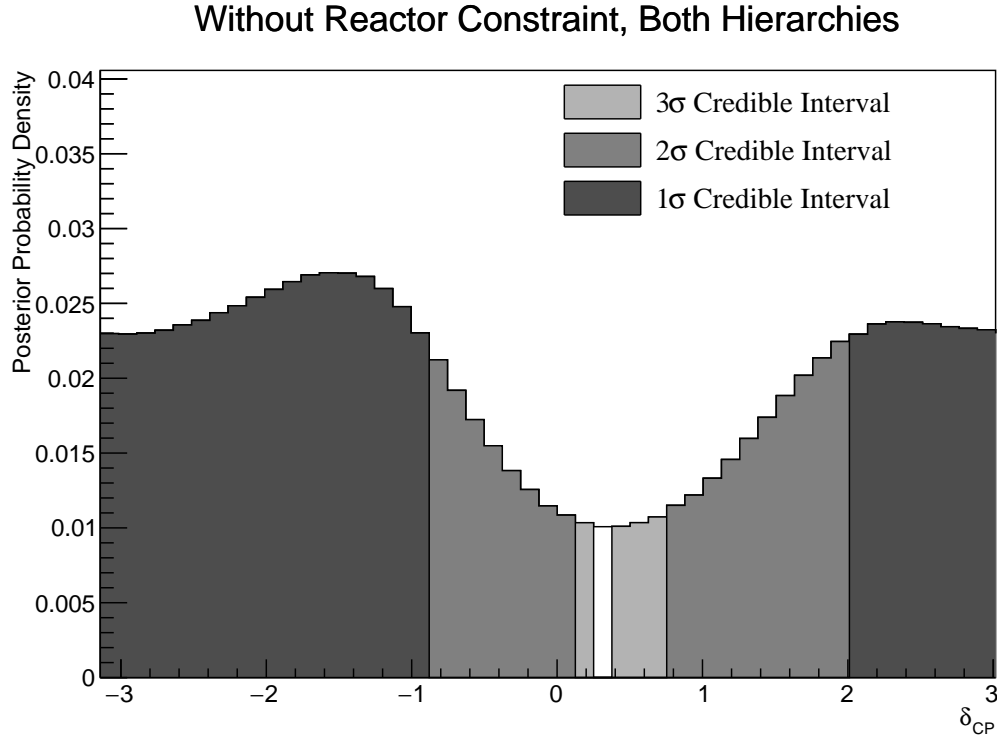
the NH and the IH, as an equal prior is placed on both models. A similar analysis can be performed by calculating the fraction of steps which fall in the lower octant (LO, $\sin^2 \theta_{23} < 0.5$) or upper octant (UO, $\sin^2 \theta_{23} > 0.5$). The fraction of steps, broken down by hierarchy and octant, are given in Table 8.2. The Bayes factor for preferred hierarchy model is $B(\text{NH}/\text{IH}) = 1.37$. Table 4.1 states this value of the Bayes factor indicates a weak preference for the normal hierarchy model. The Bayes factor for choice of octant is $B(\text{UO}/\text{LO}) = 2.24$. This is also classified as a weak preference for the UO. Both of these show that the fit is returning the correct choice of models (NH and UO) for the known Asimov A oscillation parameters defined in Table 2.2.

	LO ($\sin^2 \theta_{23} < 0.5$)	UO ($\sin^2 \theta_{23} > 0.5$)	Sum
NH ($\Delta m_{32}^2 > 0$)	0.17	0.40	0.58
IH ($\Delta m_{32}^2 < 0$)	0.13	0.29	0.42
Sum	0.31	0.69	1.00

Table 8.2

Figure 8.8 illustrates the posterior probability density for δ_{CP} , marginalised over both hierarchies. The fit does favour the asimov position ($\delta_{CP} = -1.601$) although the posterior probability is very flat through the of $2 < \delta_{CP} < \pi$ and $-\pi < \delta_{CP} < -1$. There is also a region around $\delta_{CP} \sim 0.4$ which is disfavoured at 2σ . This indicates that the SK samples can rule out some parts of the CP conserving parameter space reasonably well, near $\delta_{CP} \sim 0$, when the true value of δ_{CP} is CP violating.

The 1σ credible intervals, broken down by hierarchy, and position in parameter space of the highest posterior density is given in Table 8.3. These are taken from the one dimensional projections of the oscillation parameters, marginalised over all other parameters within the fit. For the known asimov value of $\delta_{CP} = -1.601$, the 1σ credible interval rules out a region between $\delta_{CP} = -0.86$ and $\delta_{CP} = 1.96$, when marginalising over both hierarchies. Interestingly, when considering the width of the interval when only considering steps in the NH,

**Figure 8.8**

Parameter	Interval	HPD
δ_{CP} , (BH)	$[-\pi, -0.86], [1.96, \pi]$	-1.57
δ_{CP} , (NH)	$[-\pi, -0.86], [1.88, \pi]$	-1.57
δ_{CP} , (IH)	$[-\pi, -0.94], [1.96, \pi]$	-1.57
Δm_{32}^2 (BH) [$\times 10^{-3}\text{eV}^2$]	$[-3.00, -2.50], [2.35, 3.15]$	2.65
Δm_{32}^2 (NH) [$\times 10^{-3}\text{eV}^2$]	$[2.39, 3.04]$	2.64
Δm_{32}^2 (IH) [$\times 10^{-3}\text{eV}^2$]	$[-3.15, -2.45]$	-2.70
$\sin^2(\theta_{23})$ (BH)	$[0.476, 0.59]$	0.542
$\sin^2(\theta_{23})$ (NH)	$[0.476, 0.59]$	0.554
$\sin^2(\theta_{23})$ (IH)	$[0.476, 0.59]$	0.542

Table 8.3

the intervals become narrower and the results excludes a large region of phase space. Thus, if the hierarchy model is known before the fit, the constraint would be stronger. The 1σ credible intervals for $\sin^2(\theta_{23})$ were found to be the same in

all three hierarchy choices (marginalised over both, NH and IH). This illustrates that the distribution of $\sin^2(\theta_{23})$ is symmetric across the hierarchy discontinuity.

As expected, the width of the credible intervals in Δm_{23}^2 is smaller when only the NH is considered compared to when both models are marginalised over. This follows from the fit weakly preferring the NH model over the IH model. Conversely, when the credible intervals are built using only IH steps, the credible intervals are wider than when both hierarchies are considered.

The sensitivity of the atmospheric samples to $\sin^2(\theta_{13})$ is presented in Figure 8.9. The likelihood scans presented in Figure 8.1 suggest that the sensitivity to $\sin^2(\theta_{13})$ will be small. This is seen from the fit results, where the width of the 1σ credible intervals span the region of $\sin^2(\theta_{13}) = [0.008, 0.08]$. This is an order of magnitude larger than the reactor constraint.

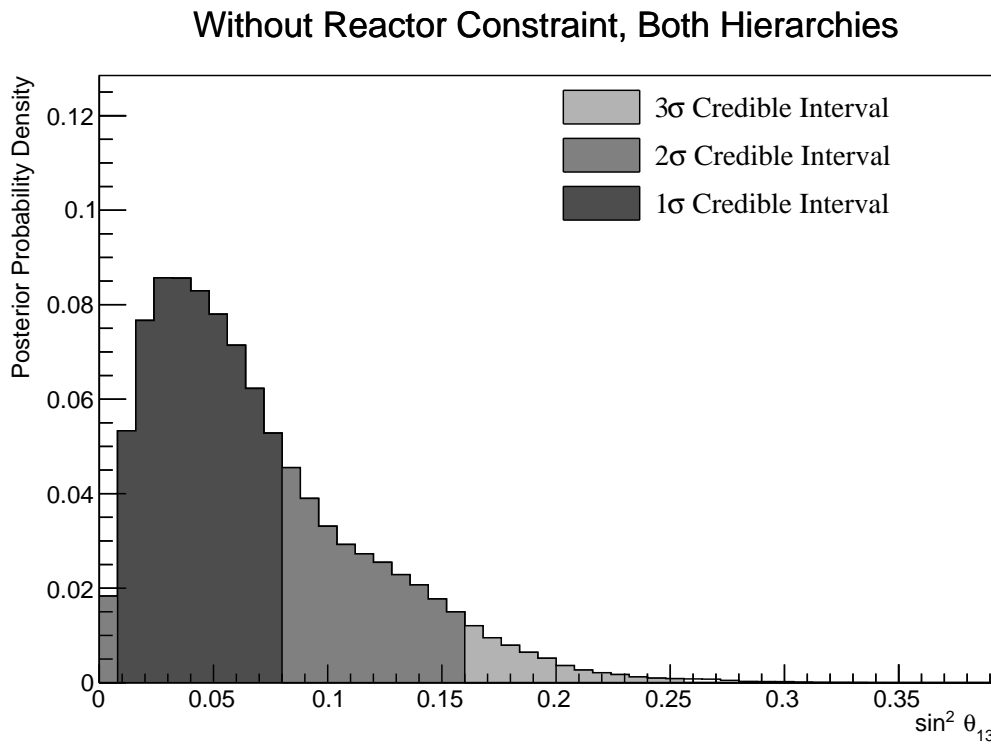


Figure 8.9

As previously discussed, the correlations between oscillation parameters are also important to understand how the atmospheric samples respond. Figure 8.10 illustrates the two dimensional $\sin^2(\theta_{13}) - \delta_{CP}$ projection of the full posterior

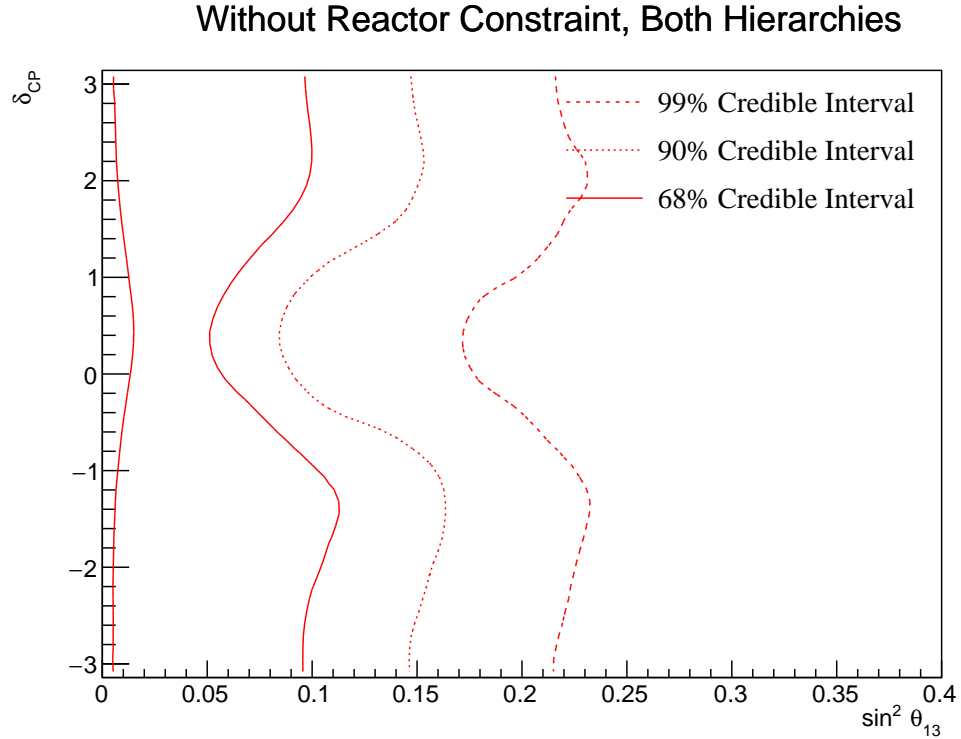


Figure 8.10

probability distribution, marginalised over both hierarchies. The shape of the 1σ credible interval shows that the constraining power of the fit on δ_{CP} is dependent upon the choice of $\sin^2(\theta_{13})$. As expected from Figure 8.1, the atmospheric samples do not strongly constrain the value of $\sin^2(\theta_{13})$. However, the choice of preferred value of $\sin^2(\theta_{13})$ does impact the atmospheric sensitivity to δ_{CP} . The 1σ credible interval is consistent with the reactor constraint.

The $\sin^2(\theta_{23}) - \Delta m_{32}^2$ disappearance contours are illustrated in Figure 8.11. As expected, the distribution in the inverted hierarchy is slightly smaller than that in the normal hierarchy. This follows from the Bayes factor showing weak preference for NH meaning that more of the steps will exist in the $\Delta m_{32}^2 > 0$ region. The asimov points of $\sin^2(\theta_{23}) = 0.528$ and $\Delta m_{32}^2 = 2.509 \times 10^{-3} \text{ eV}^2$ are clearly contained within the 1σ credible interval.

Figure 8.12 highlights the two dimensional projections for each combination of oscillation parameters which this analysis is sensitive too. The contours are built when marginalising over both hierarchies but only the NH is illustrated

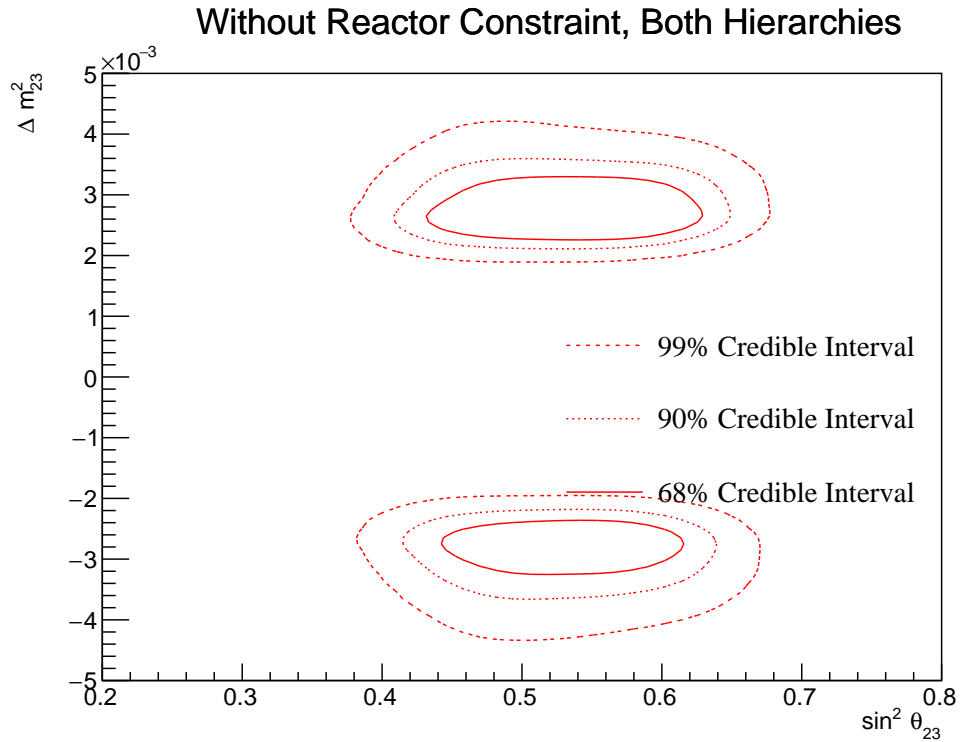


Figure 8.11

when considering Δm^2_{23} . As expected the correlations play a significant role in these sensitivity measurements, especially the choice of $\sin^2(\theta_{13})$ constraint. The application of reactor constraint would be expected to alter both the width and position of the Δm^2_{23} credible interval and the width of the δ_{CP} and $\sin^2(\theta_{23})$ constraints. The majority of the octant model preference comes from the region of $\sin^2(\theta_{13}) \sim 0.03$. Consequently, the application of the reactor constraint would not be expected to significantly change the octant preference. Interestingly, the distribution of steps in the δ_{CP} - $\sin^2(\theta_{13})$ plot is slightly flatter in the region of the reactor constraint. This follows from Figure 8.2 where the likelihood scan shows a region of low negative log-likelihood extending out towards higher values of $\sin^2(\theta_{13})$ in the $\delta_{CP} = -1.601$ region. Consequently, the reactor constraint could feasibly reduce the sensitivity of the atmospheric samples to δ_{CP} , due to these marginalisation effects.

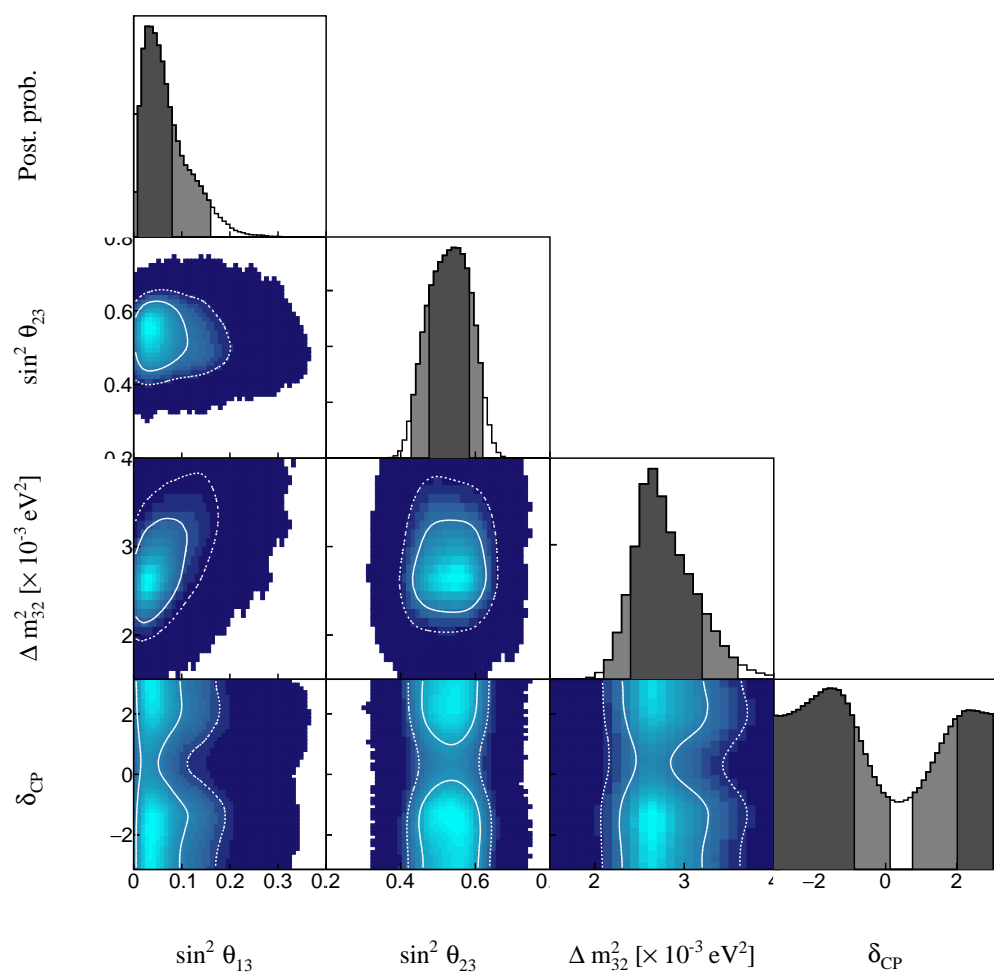


Figure 8.12

8.3.2 Atmospheric-Only Sensitivity With Reactor Constraint

The results in subsection 8.3.1 discuss the atmospheric sensitivity when the reactor constraint is not applied. However, the correlations illustrated in Figure 8.12 indicate that the marginalisation effects could contribute to differing sensitivities when the external constraint is applied. Using the technique discussed in subsection 4.1.1, the posterior distribution can be reweighted to include the reactor constraint of $\sin^2(\theta_{13}) = (2.18 \pm 0.08) \times 10^{-2}$.

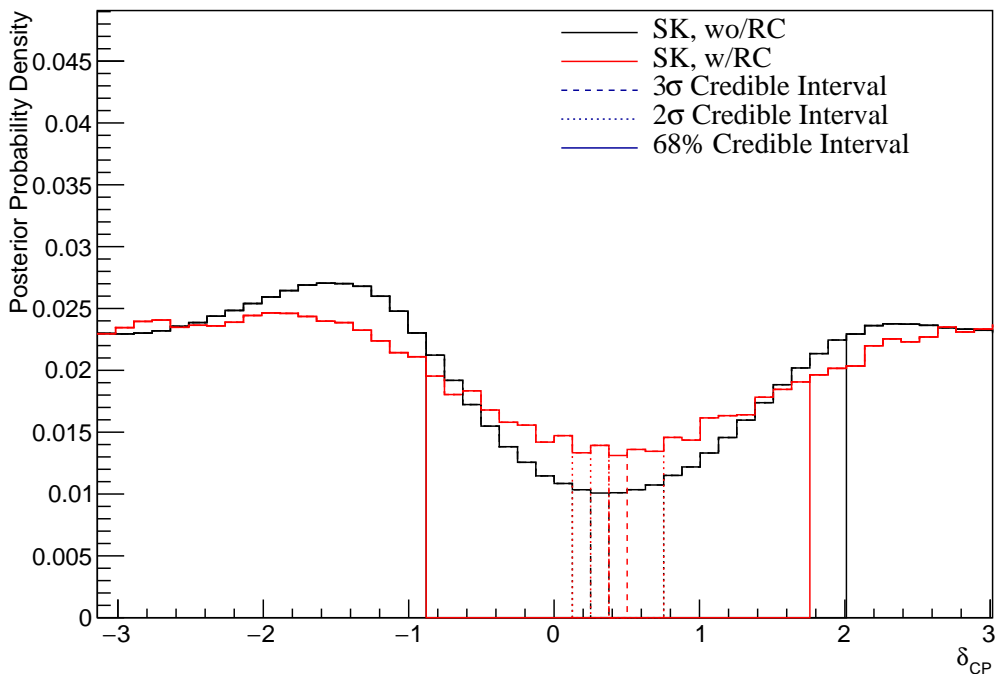


Figure 8.13

Figure 8.13 illustrates the sensitivity to δ_{CP} . The posterior distribution for the ‘with Reactor Constraint’ fit is less peaked than the previous results. This is due to the expected marginalisation effect observed in the likelihood scan in Figure 8.2. The width of the 1σ credible interval is increased when the reactor constraint is applied, indicating less sensitivity in that region of $\sin^2(\theta_{13})$.

The reactor constraint increases the sensitivity of the atmospheric samples to Δm_{23}^2 as illustrated in Figure 8.14. Due to the marginalisation effects observed in Figure 8.12, the favoured region of Δm_{23}^2 moves closer to zero for both hierarchies.

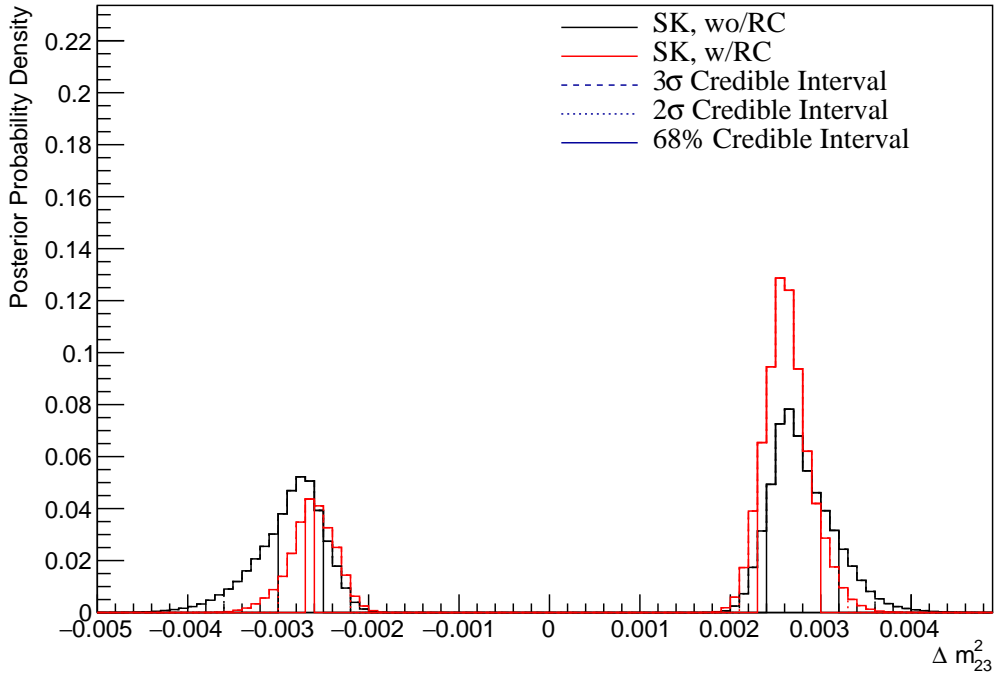


Figure 8.14

A clear explanation of this behaviour is illustrated in Figure 8.15. The correlation between Δm_{23}^2 and $\sin^2(\theta_{13})$ is such that lower values of $\sin^2(\theta_{13})$ tend towards lower values of $|\Delta m_{32}^2|$. The 1σ credible interval for the Δm_{23}^2 parameter is determined to be $[-2.70, -2.55] \times 10^{-3}\text{eV}^2$ and $[2.25, 2.95] \times 10^{-3}\text{eV}^2$. The width of the IH credible interval is reduced by approximately 70% when the reactor constraint is applied.

	LO ($\sin^2 \theta_{23} < 0.5$)	UO ($\sin^2 \theta_{23} > 0.5$)	Sum
NH ($\Delta m_{32}^2 > 0$)	0.21	0.53	0.74
IH ($\Delta m_{32}^2 < 0$)	0.08	0.18	0.26
Sum	0.29	0.71	1.00

Table 8.4

The asymmetry in the number of steps within the NH to IH clearly shows that the reactor constraint increases the NH preference. The fraction of steps in each hierarchy and octant model for this fit are given in Table 8.4. The preference of

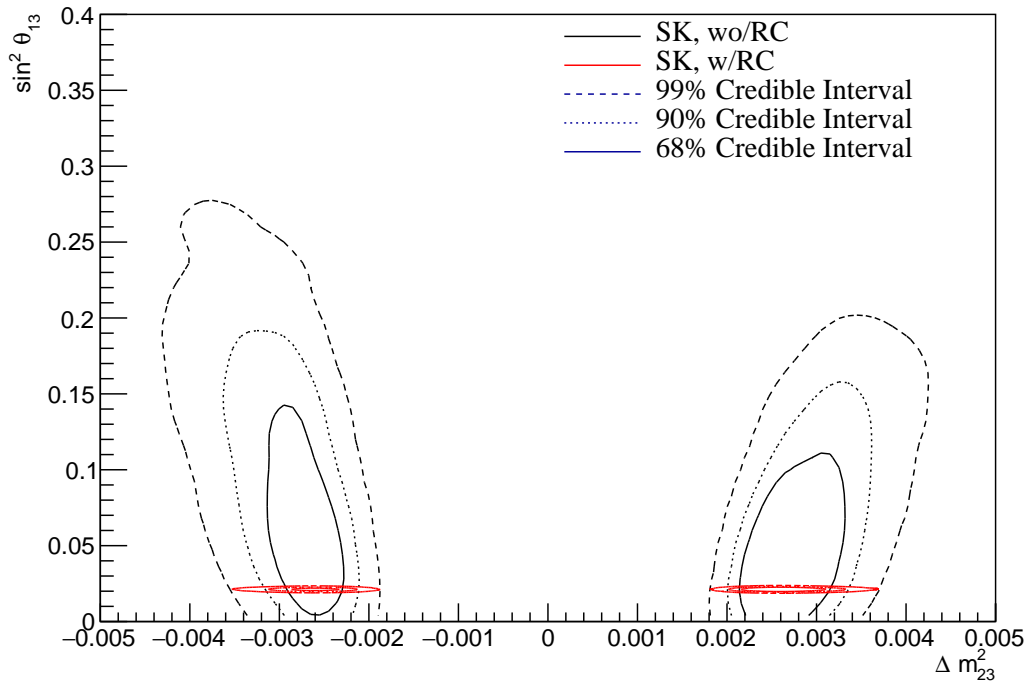


Figure 8.15

octant model is very slightly increased by the application of the reactor constraint which is consistent with expectation. The reactor constraint significantly increases the NH preference, increasing the Bayes factor from $B(\text{NH}/\text{IH}) = 1.37$ to $B(\text{NH}/\text{IH}) = 2.85$ when the reactor constraint is applied. This is still defined as a weak preference for NH according to the Jeffrey's scale (see Table 4.1), however it is a stronger preference than without the constraint. The Bayes factor for octant determination is calculated as $B(\text{UO}/\text{LO}) = 2.45$.

8.3.3 Application of Near Detector Constraints for Atmospheric Samples

The choice to apply the near detector constraints onto the low energy atmospheric samples was introduced in subsection 6.4.3. The effect of this choice can be studied by comparing the sensitivities of the atmospheric only analysis when the near detector constraints are and are not applied. The change in sensitivity on δ_{CP} is given in Figure 8.16. The shape of the posterior is approximately the same

although slightly less peaked at the asimov point. The width of the 1σ credible intervals are approximately the same (identical to within a bin width) and the same conclusion holds for the higher credible intervals. Similar conclusions hold for the other oscillation parameters. As expected, the sensitivities are still statistics dominated and the exact choice of systematic model and constraint does not effect the physics conclusions one would make from this analysis.

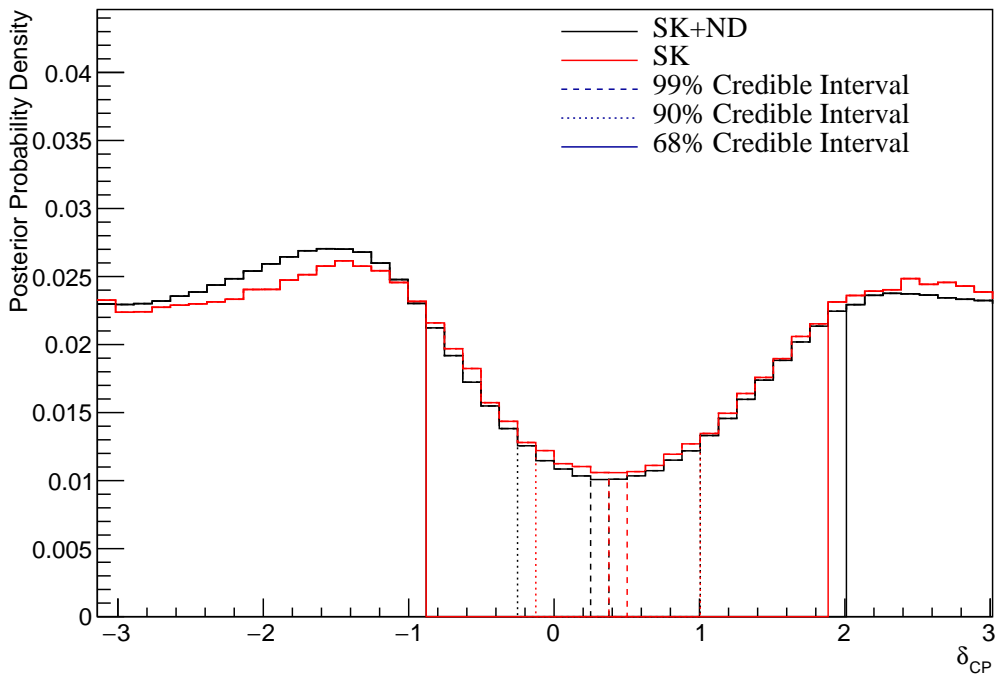


Figure 8.16

8.3.4 Atmospheric and Beam Sensitivity without Reactor Constraint

This section presents the sensitivities of the simultaneous beam and atmospheric analysis without the reactor constraint. The asimov data is built assuming the post-BANFF cross section tune and Asimov A oscillation parameters defined in Table 2.2. This is a fit to all 18 near detector beam samples, 5 far detector beam samples and 18 atmospheric samples. The sensitivity to δ_{CP} , marginalised over both hierarchies is given in Figure 8.17.

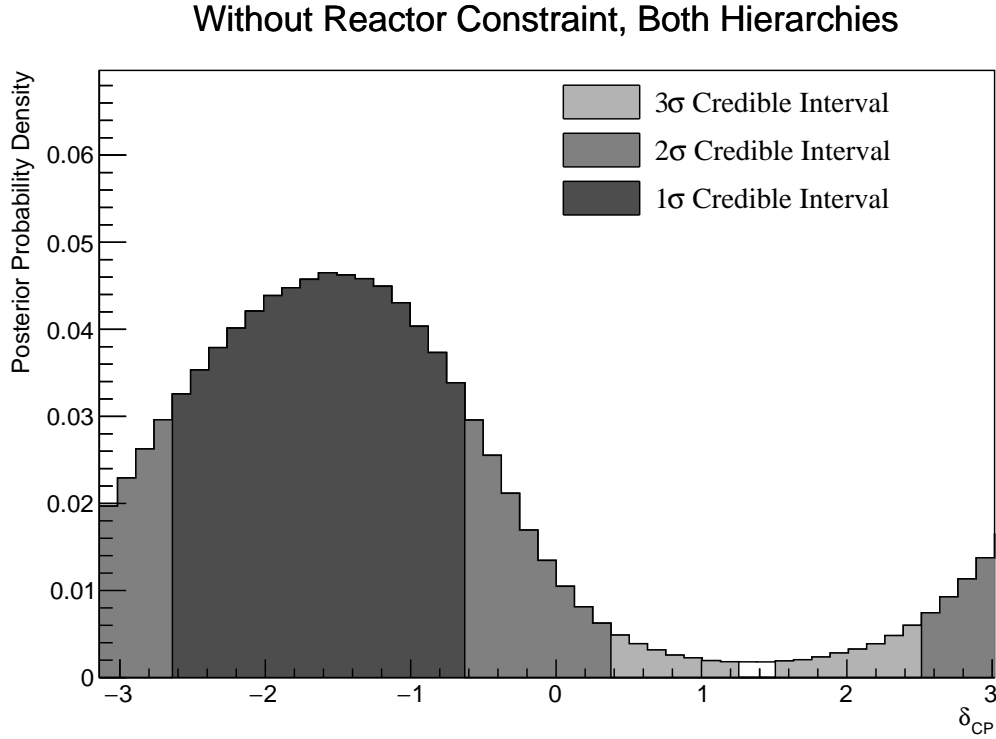


Figure 8.17

The credible intervals and highest posterior distribution for each oscillation parameter is given in Table 8.5. The highest posterior density is $\delta_{CP} = -1.58$ and is clearly compatible with the known Asimov A value of $\delta_{CP} = -1.601$. The CP-conserving values of $\delta_{CP} = 0, \pi, -\pi$ are disfavoured at 1σ credible interval. There is also a region around $\delta_{CP} = 1.4$ which is disfavoured at more than 3σ . Whilst these conclusions can be made at this particular asimov point, it does show that if the true value of δ_{CP} was CP-violating, this joint analysis would be able to disfavour CP conserving values at over 1σ . The highest posterior density does move further away from the asimov point when only steps in the NH region are considered. This is due to the correlations between the value of δ_{CP} and the mass hierarchy.

The sensitivity to Δm_{23}^2 is illustrated in Figure 8.17 for the normal hierarchy model hypotheses. Notably, the 1σ credible interval is entirely contained within the normal hierarchy region, as illustrated in Table 8.5. This is illustrates reasonable sensitivity to the mass hierarchy model. This is also reflected in the 1σ

Parameter	Interval	HPD
δ_{CP} , (BH)	[-2.64, -0.63]	-1.57
δ_{CP} , (NH)	[-2.76, -0.63]	-1.45
δ_{CP} , (IH)	[-2.39, -0.88]	-1.57
Δm_{32}^2 (BH) [$\times 10^{-3}\text{eV}^2$]	[2.46, 2.58]	2.49
Δm_{32}^2 (NH) [$\times 10^{-3}\text{eV}^2$]	[2.48, 2.56]	2.51
Δm_{32}^2 (IH) [$\times 10^{-3}\text{eV}^2$]	[-2.60, -2.52]	-2.55
$\sin^2(\theta_{23})$ (BH)	[0.48, 0.55]	0.509
$\sin^2(\theta_{23})$ (NH)	[0.48, 0.55]	0.509
$\sin^2(\theta_{23})$ (IH)	[0.48, 0.55]	0.521

Table 8.5

credible intervals being approximately the same when they are made considering both hierarchies and when considering only the NH.

The fraction of steps in each of the mass hierarchy regions and octants of $\sin^2(\theta_{23})$ is given in Table 8.6. The Bayes factors are determined to be $B(\text{NH}/\text{IH}) = 3.67$ and $B(\text{UO}/\text{LO}) = 1.73$. Jeffrey's scale (presented in Table 4.1) states that this value of the hierarchy Bayes factor illustrates suubstantial evidence for normal hierarchy. This corresponds the known oscillation parameters input into the asimov data and is a strong statement than the atmospheric-only analysis can provide. The Bayes factor for octant determination represents a weak preference for the upper octant.

	LO ($\sin^2 \theta_{23} < 0.5$)	UO ($\sin^2 \theta_{23} > 0.5$)	Sum
NH ($\Delta m_{32}^2 > 0$)	0.29	0.50	0.79
IH ($\Delta m_{32}^2 < 0$)	0.08	0.13	0.21
Sum	0.37	0.63	1.00

Table 8.6

The Asimov A definition is $\Delta m_{32}^2 = 2.509 \times 10^{-3}\text{eV}^2$. The normal hierarchy distribution clearly favours this region with the highest posterior density of

$\Delta m_{32}^2 = 2.51 \times 10^{-3}$ eV². The highest posterior density contained within the inverse hierarchy posterior distribution is $\Delta m_{32}^2 = -2.55 \times 10^{-3}$ eV².

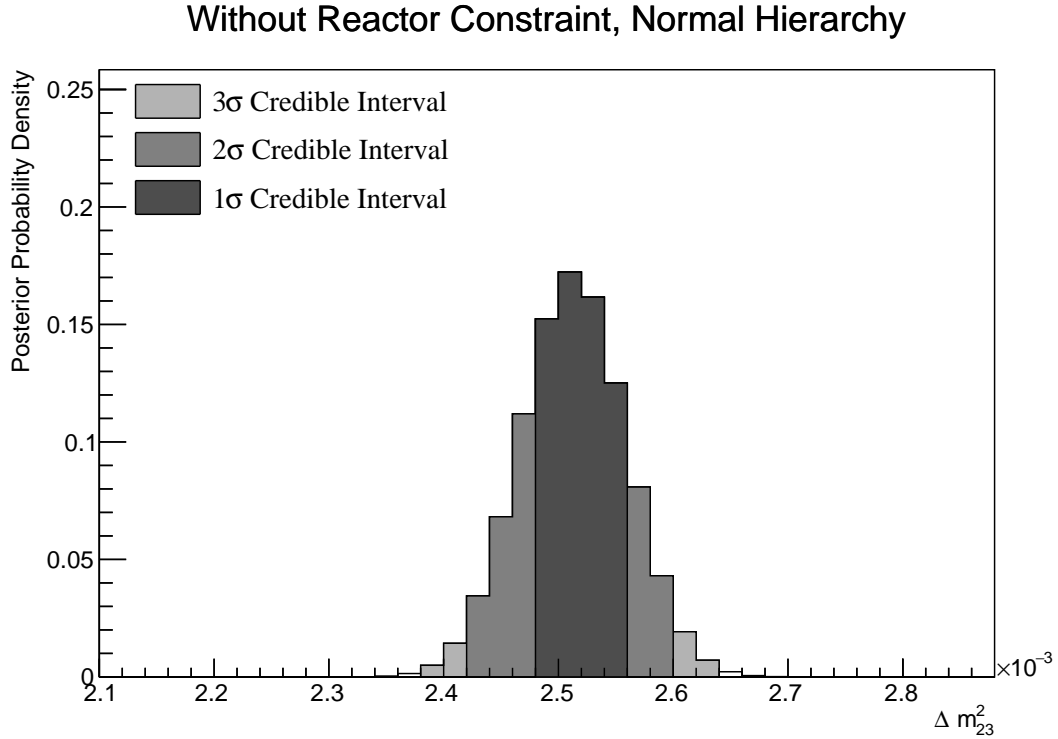


Figure 8.18

The sensitivity to $\sin^2(\theta_{23})$ is presented in Figure 8.19. There is clear preference for the upper octant but the peak of the distribution is relatively flat. It peaks in the region of the expected Asimov value of $\sin^2(\theta_{23}) = 0.528$. The difference in the highest posterior distribution and the width of the credible interval is relatively unchanged when considering different hierarchy models. Consequently, the correlation between Δm_{23}^2 and $\sin^2(\theta_{23})$ is such that it is approximately mirrored across the hierarchy discontinuity.

The sensitivity presented as a function of the appearance parameters ($\sin^2(\theta_{13}) - \delta_{CP}$) is given in Figure 8.20. As expected, the contours follow that given in Figure 8.2, where the 2σ credible intervals have a closed contour excluding the region around $\delta_{CP} \sim 1.2$. The width of the 3σ credible interval is also clearly dependent upon the value of δ_{CP} . Close to the asimov point, $\delta_{CP} = -1.601$, the width of the 3σ credible interval approximately spans $\sin^2(\theta_{13}) = [0.013, 0.04]$.

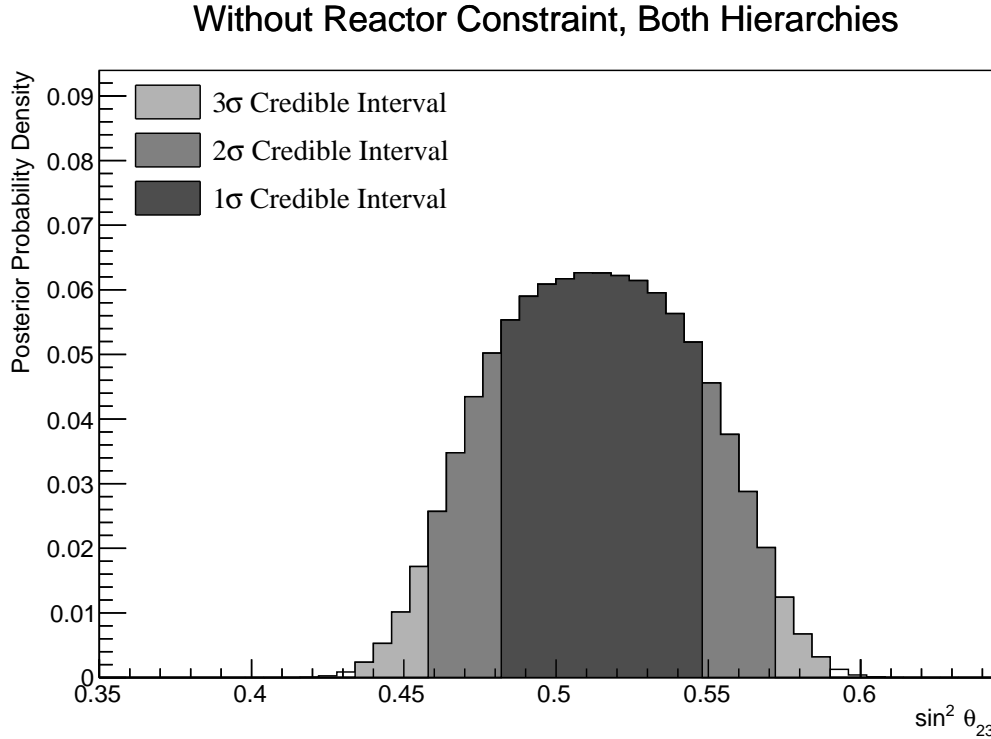


Figure 8.19

This is reduced to a region of $\sin^2(\theta_{13}) = [0.023, 0.042]$ at the most disfavour value of δ_{CP} . The 1σ credible interval is consistent with both the asimov point and the reactor constraint. Application of the reactor constraint would be expected to decreased the width of the 1σ credible intervals of δ_{CP} due to the triangular shape of the posterior probability.

The sensitivity in terms of the ‘disappearance’ parameters marginalised over both hierarchies is given in Figure 8.21. In this two-dimensional projection of the posterior distribution, a small section of the 1σ credible interval is contained within the inverse hierarchy region. That IH region is clearly favouring the upper octant as expected. The 1σ credible region of the NH contour spans both octants but favours the UO.

Without Reactor Constraint, Both Hierarchies

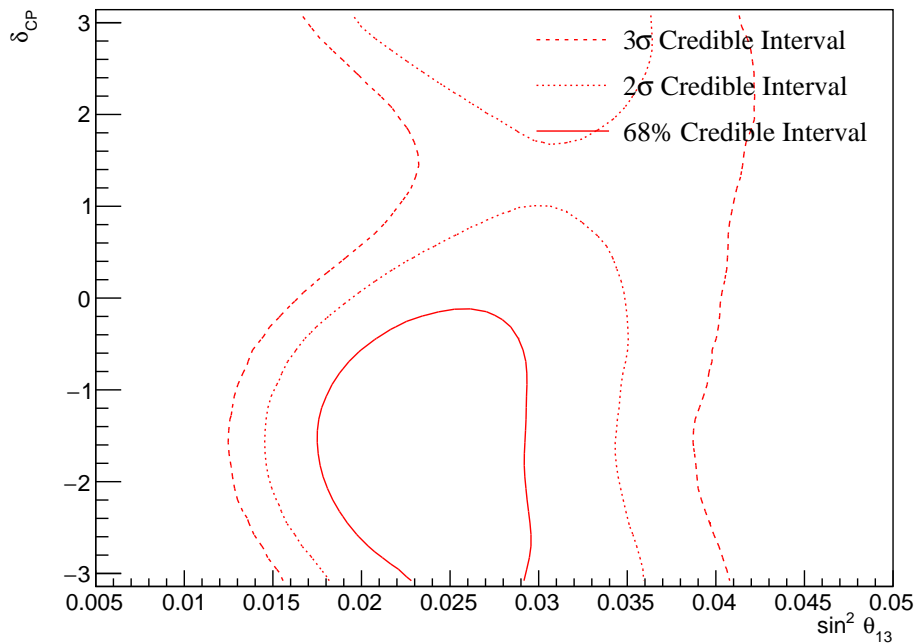


Figure 8.20

Without Reactor Constraint, Both Hierarchies

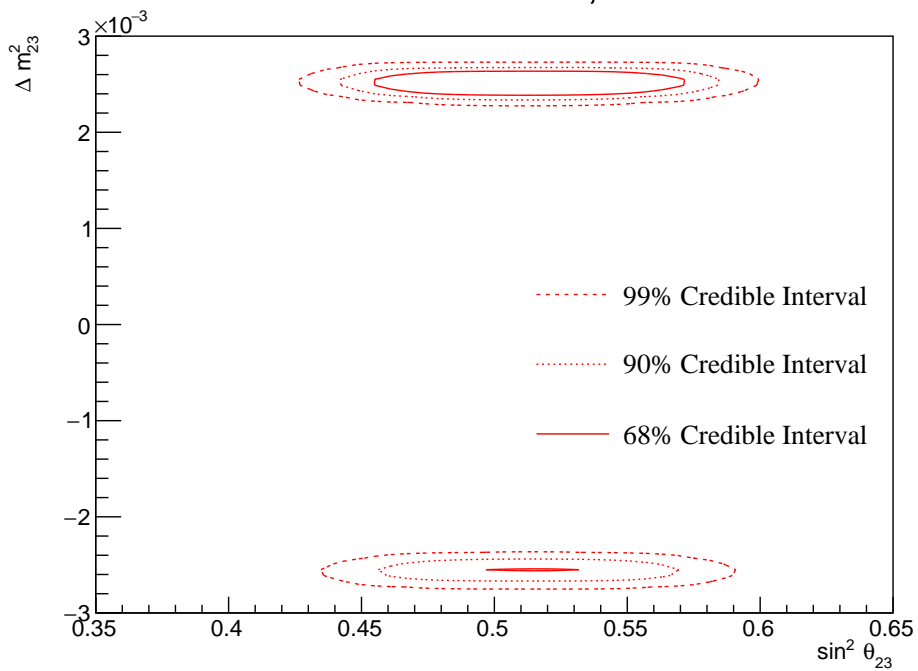


Figure 8.21

The two-dimensional posterior distribution for each permutation of the oscillation parameters of interest is given in Figure 8.22. The most notable observation is that the $\sin^2(\theta_{13})$ and $\sin^2(\theta_{23})$ are anti-correlated. If the value of $\sin^2(\theta_{13})$ was known to be closer to the asimov point, the preferred value of $\sin^2(\theta_{23})$ would increase. That would move the highest posterior probability closer in line with the asimov value. This also means that, the preference for the UO would be increased if the reactor constraint was to be applied.

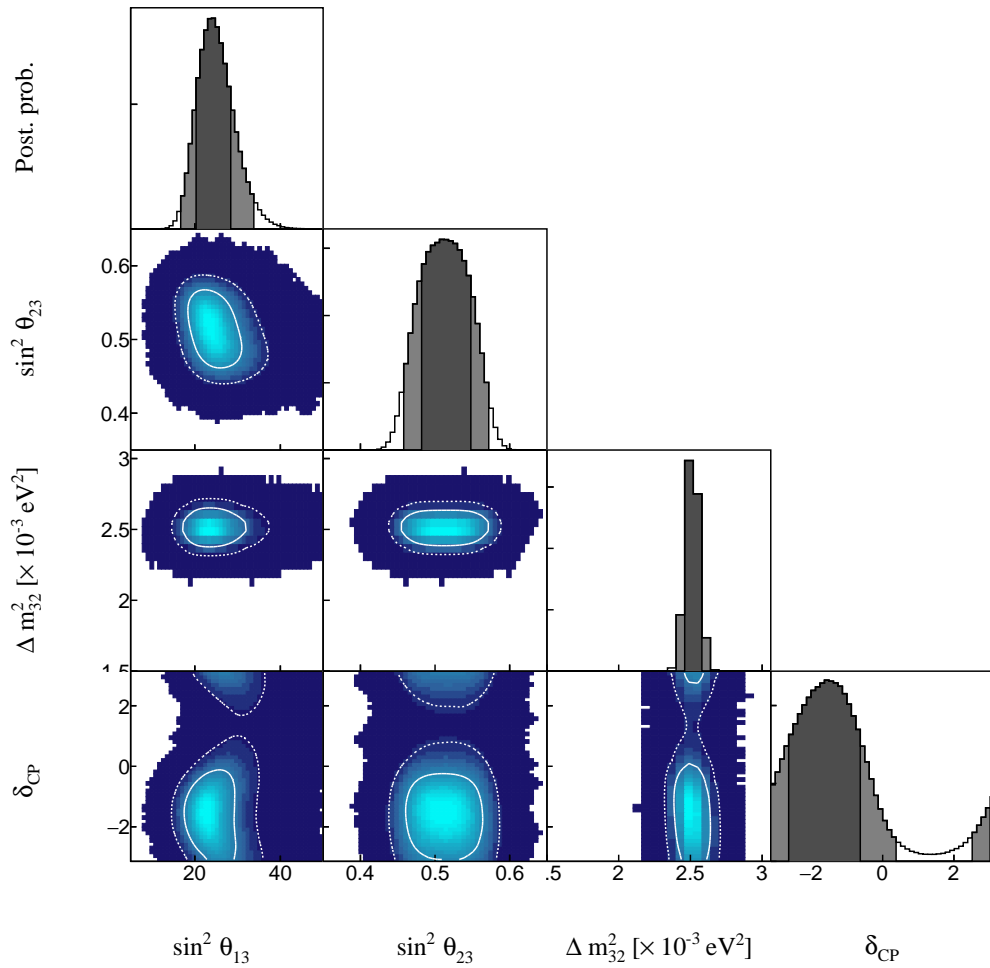


Figure 8.22

The correlation between $\sin^2(\theta_{13})$ and Δm_{23}^2 can be seen in Figure 8.23. A much larger fraction of the posterior distribution is contained in the NH for lower

values of $\sin^2(\theta_{13})$. Consequently, the application of the reactor constraint would be expected to significantly increase the preference for NH.

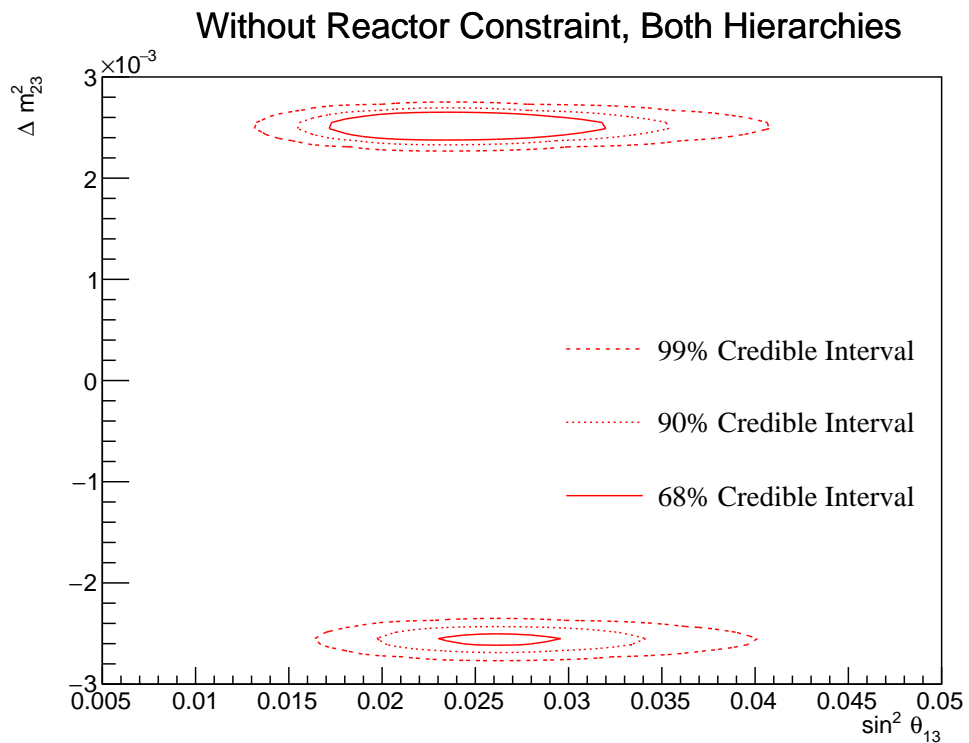


Figure 8.23

8.3.5 Atmospheric and Beam Sensitivity with Reactor Constraint

8.3.6 Comparison to Latest T2K Sensitivities

9

Conclusions and Outlook

Appendices

A

Atmospheric Sample Spectra

This appendix documents the interaction mode breakdown of all the atmospheric samples used within the analysis. The generated tune of the model parameters and the Asimov A oscillation parameter set (defined in Table 2.2) are assumed. The livetime of SK-IV is taken to be 3244.4 days.

A.1 Binning

The lepton momentum and cosine zenith binning edges for the atmospheric samples used within this analysis are defined in Table A.1.

A.2 Fully Contained Sub-GeV Samples

The interaction mode breakdown of the fully contained Sub-GeV samples are shown in Figure A.1 and Figure A.2, for the samples with enriched CC0 π and CC1 π^\pm respectively.

The CC0 π sample are dominated by CCQE events ($\sim 70\%$) with smaller contributions of 2p2h ($\sim 12\%$) and CC1 π ($\sim 10\%$) components. The energy peaks around 300 MeV, which is slightly below that of the T2K samples but still has significant contribution upto 1 GeV which overlaps the T2K sample energy range.

Sample	$\cos(\theta_Z)$ Bins	Momentum Bin Edges ($\log_{10}(P)$ MeV)
SubGeV-elike-0dcy	10	2.0, 2.4, 2.6, 2.8, 3.0, 3.2
SubGeV-elike-1dcy	1	2.0, 2.4, 2.6, 2.8, 3.0, 3.2
SubGeV-mulike-0dcy	10	2.0, 2.4, 2.6, 2.8, 3.0, 3.2
SubGeV-mulike-1dcy	10	2.0, 2.4, 2.6, 2.8, 3.0, 3.2
SubGeV-mulike-2dcy	1	2.0, 2.4, 2.6, 2.8, 3.0, 3.2
SubGeV-pi0like	1	2.0, 2.2, 2.4, 2.6, 2.8, 3.2
MultiGeV-elike-nue	10	3.0, 3.4, 3.7, 4.0, 5.0
MultiGeV-elike-nuebar	10	3.0, 3.4, 3.7, 4.0, 5.0
MultiGeV-mulike	10	3.0, 3.4, 5.0
MultiRing-elike-nue	10	3.0, 3.4, 3.7, 5.0
MultiRing-elike-nuebar	10	3.0, 3.4, 3.7, 5.0
MultiRing-mulike	10	2.0, 3.124, 3.4, 3.7, 5.0
MultiRing-Other1	10	3.0, 3.4, 3.7, 4.0, 5.0
PC-Stop	10	2.0, 3.4, 5.0
PC-Through	10	2.0, 3.124, 3.4, 3.7, 5.0
Upmu-Stop	10	3.2, 3.4, 3.7, 8.0
Upmu-Through-Showering	10	2.0, 8.0
Upmu-Through-NonShowering	10	2.0, 8.0

Table A.1: The reconstructed cosine zenith and lepton momentum binning assigned to the atmospheric samples. The “ $\cos(\theta_Z)$ Bins” column illustrates the number of bins uniformly distributed over the $-1.0 \leq \cos(\theta_Z) \leq 1.0$ region for fully and partially contained samples and $-1.0 \leq \cos(\theta_Z) \leq 0.0$ region for up- μ samples.

The one-ring CC1 π samples, where the pion is tagged via its decay electron, are dominated by CC1 π events ($\sim 75\%$) with a small contribution of CCM π ($\sim 10\%$). The two-ring pion sample is mostly dominated by the NC1 π^0 via resonances, and has several equally-sized contributions from CCQE, NC1 π^\pm via resonances, and NC coherent pion production, where the π^0 likely comes from nucleon and π^\pm final state interactions in the nucleus.

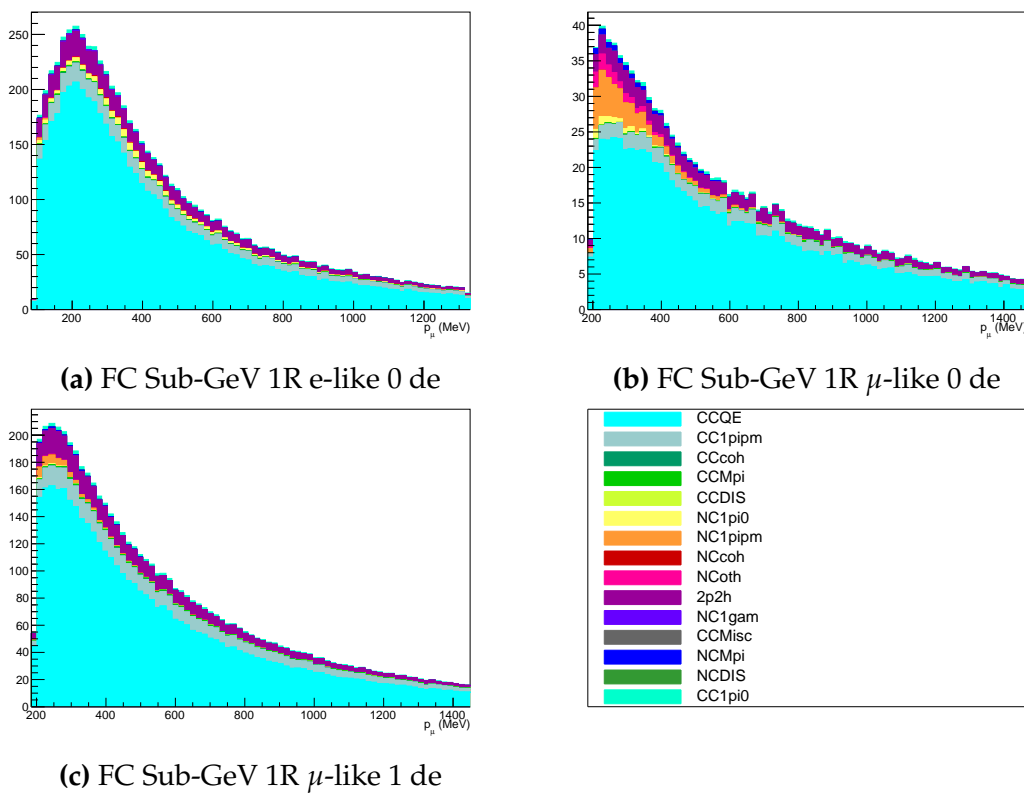


Figure A.1: Breakdown by interaction mode of the FC Sub-GeV atmospheric samples targeting CC 0π events.

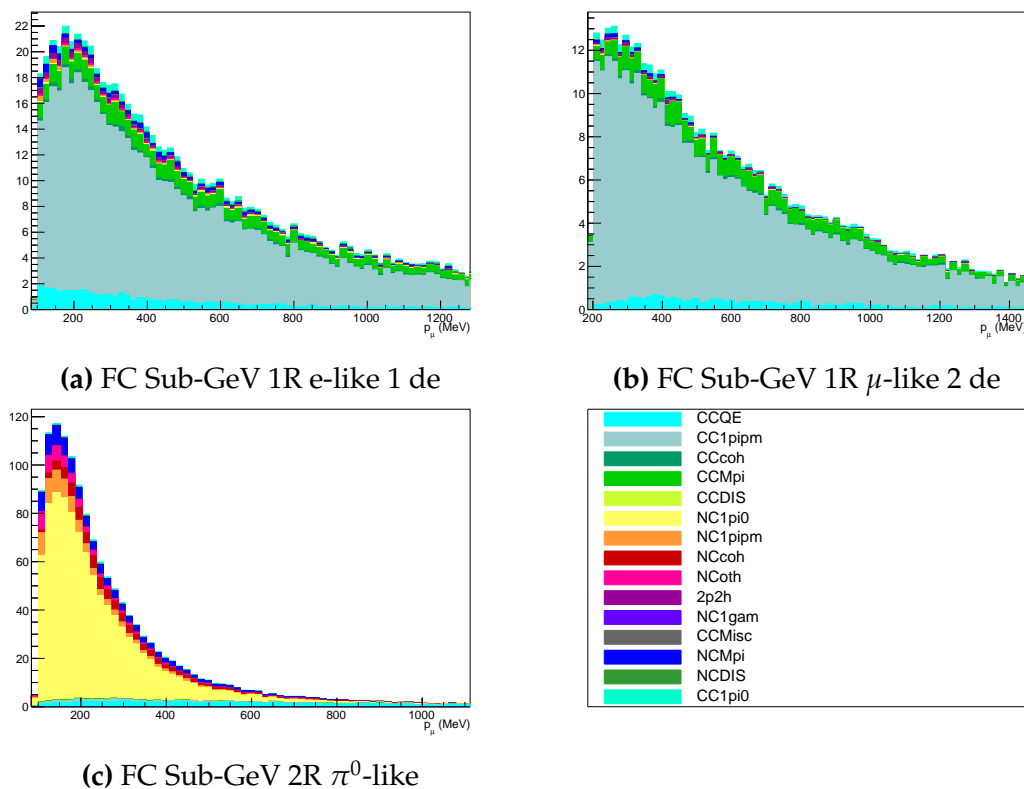


Figure A.2: Breakdown by interaction mode of the FC Sub-GeV atmospheric samples targeting single pion events.

A.3 Fully Contained Multi-GeV Samples

The interaction mode breakdown of fully contained multi-GeV samples is highlighted in Figure A.3. Due to the event selection applied in SK which targets π^+ and π^- separation, the ν_e sample mainly consists of events with pions (single pion production or multi-pion/DIS interactions). The pion separation is explained in Section section 6.1. This reasoning also explains the significant CCQE contribution of the $\bar{\nu}_e$ sample. The muon-like sample is dominated by CCQE interactions with $\sim 10 - 15\%$ 2p2h and CC1 π contribution of events.

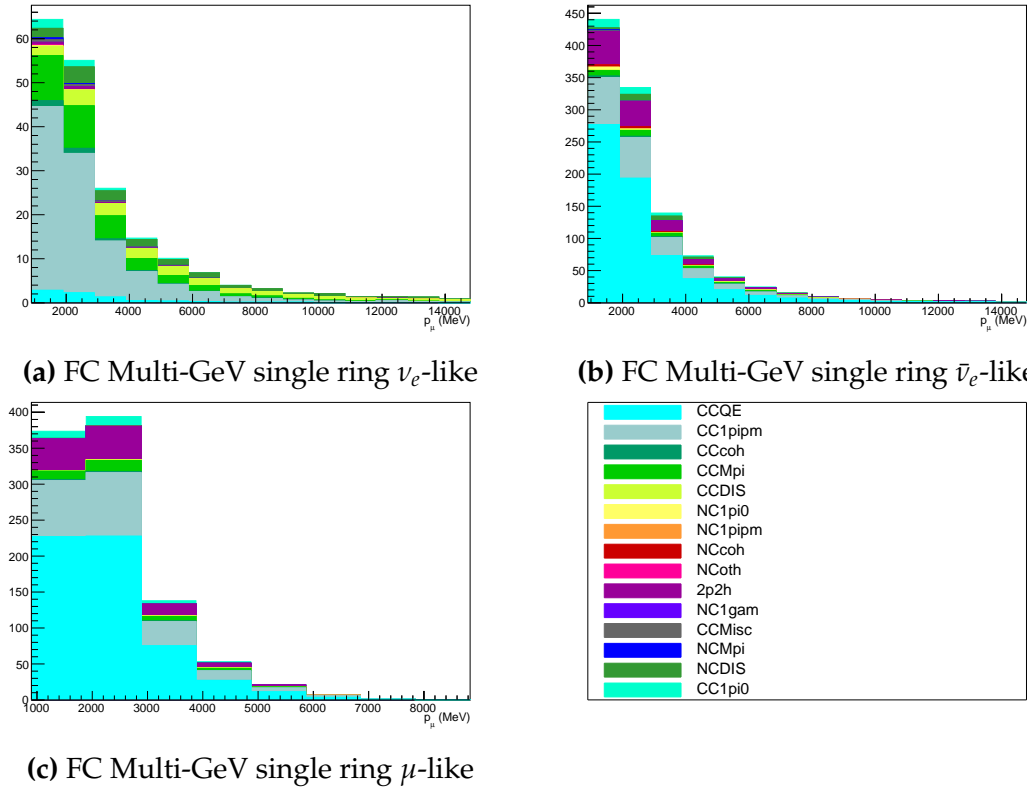


Figure A.3: Breakdown by interaction mode of the FC Multi-GeV single ring atmospheric samples.

A.4 Fully Contained Multi-Ring Samples

The interaction mode breakdown of fully contained multi-ring events is shown in Figure A.4. These samples see more interaction modes contributing in general, and there is a much larger contribution from multi-pion and DIS interaction modes, compared to the other samples.

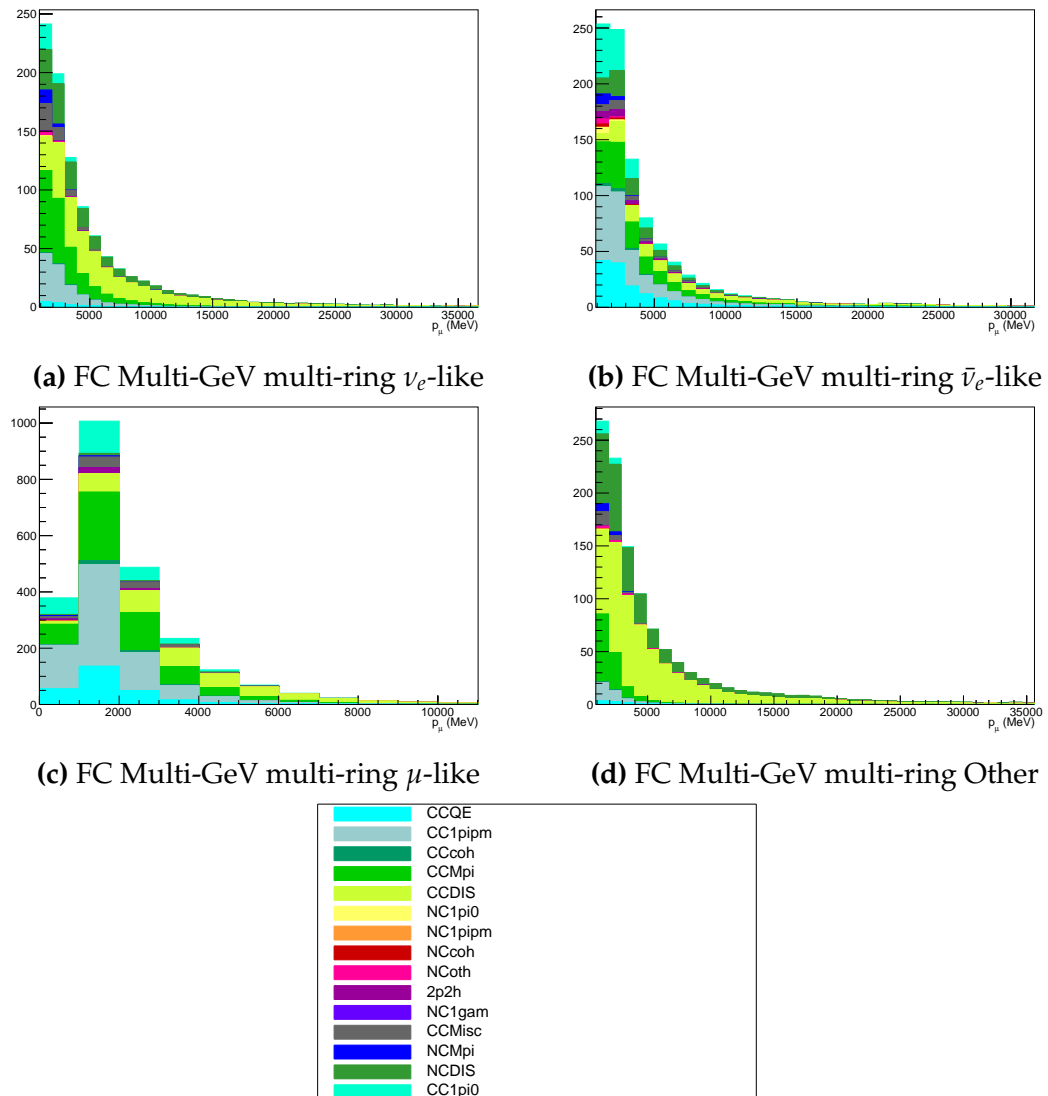


Figure A.4: Breakdown by interaction mode of the FC Multi-GeV multi-ring atmospheric samples.

A.5 Partially Contained Samples

The breakdown for partially contained samples is highlighted in Figure A.5. As with the multi-ring samples, there is no dominating interaction mode. The neutrino energies of events in this sample extend into the tens of GeV and become dominated by DIS interaction modes in the high energy limit.

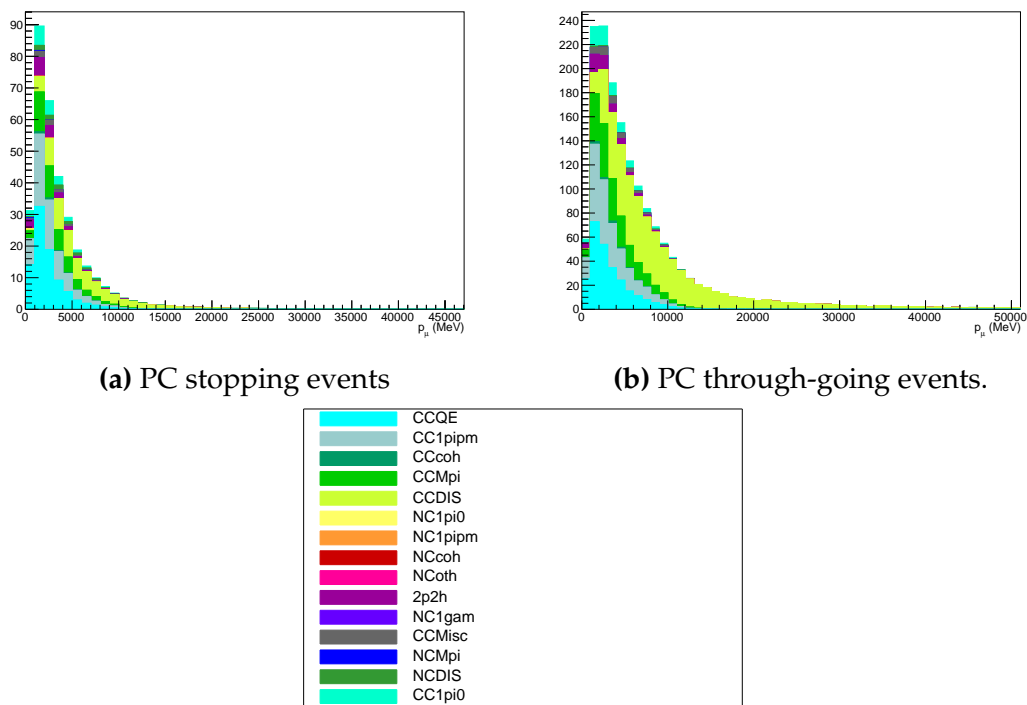


Figure A.5: Breakdown by interaction mode of the PC atmospheric samples.

A.6 Upward-Going Muon Samples

The breakdown for upward-going muons is illustrated in Figure A.6. These samples are significantly dominated by DIS interactions with energies extending up into the hundreds of GeV.

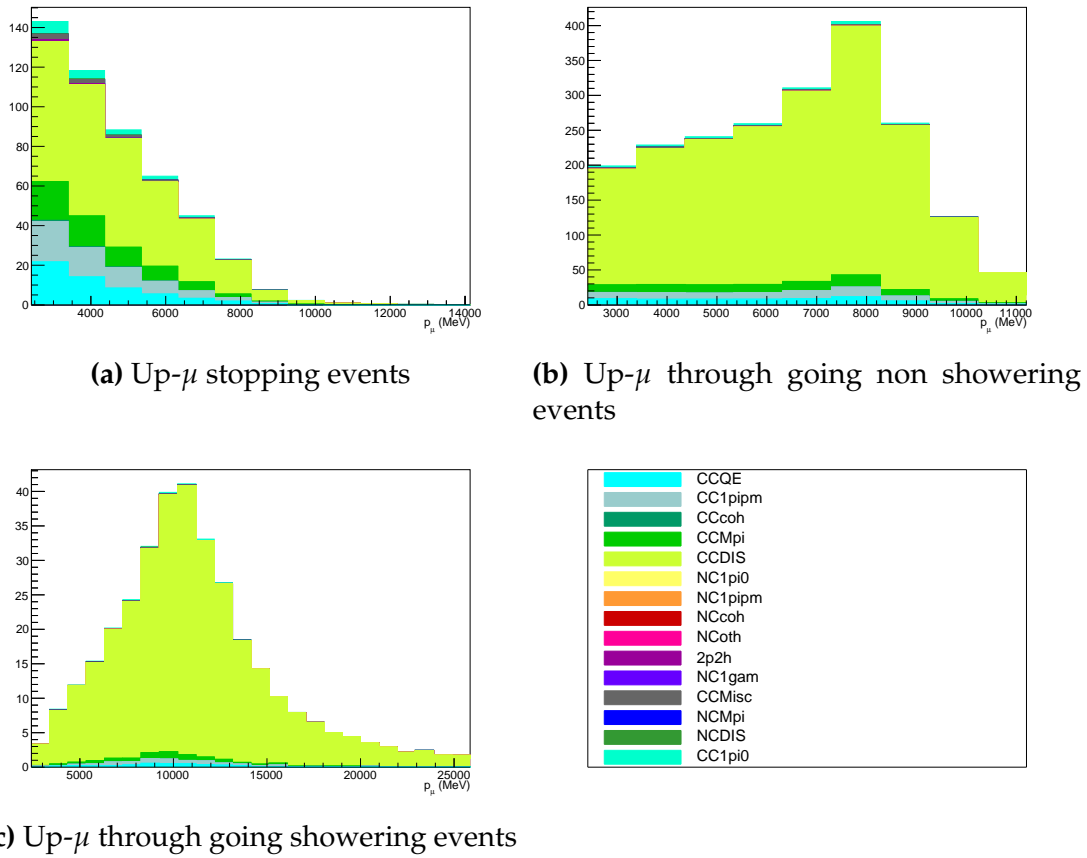


Figure A.6: Breakdown by interaction mode of the atmospheric upward going muon samples.

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