DISSERTATION

Imaging of Single Barium Atoms in Solid Xenon for Barium Tagging in the NEXO Neutrinoless Double Beta Decay Experiment

Submitted by

Timothy Walton

Department of Physics

In partial fulfillment of the requirements

For the Degree of Doctor of Philosophy

Colorado State University

Fort Collins, Colorado

Fall 2015

Doctoral Committee:

Advisor: William M. Fairbank, Jr.

Robert Wilson Bruce Berger Alan Van Orden Copyright by Timothy Walton 2015 All Rights Reserved

Abstract

The nEXO experiment is designed to search for zero-neutrino double beta decay of the isotope ¹³⁶Xe, in order to better understand the nature of neutrinos. Since the daughter of this decay is barium (¹³⁶Ba), detecting the presence of ¹³⁶Ba at a decay site (called "barium tagging") provides an additional discriminator to reject backgrounds in the search for this decay. This would involve detecting a single barium ion from within a macroscopic volume of liquid xenon. One proposed barium tagging method is to trap the barium ion in solid xenon (SXe) at the end of a cold probe, and then detect the ion by its fluorescence in the solid xenon. William M. Fairbank Jr.'s group at Colorado State University has been working toward this goal with steady success for some time. In this thesis, I demonstrate successful detection of neutral Ba in SXe down to the single atom level, after deposition from vacuum.

ACKNOWLEDGEMENTS

Bill, Chris for all the help in the lab and building things, Shon and Brian for creating the system and training me (Shon) and for the preliminary results, Adam, Cesar, Kendy.

If you want the Leif thing, uncomment it in csuthesis.cls. It says something like "this dissertation is typeset in ... designed by Leif Anderson.

Jamie and LEW, and other fam.

Table of Contents

Abstract	ii
Acknowledgements	iii
Chapter 1. Introduction	1
1.1. Neutrinos	2
1.2. Double Beta Decay	5
1.3. Enriched Xenon Observatory	7
Chapter 2. Theory	18
2.1. Ba/Ba ⁺ Spectroscopy in Vacuum	18
2.2. Matrix Isolation Spectroscopy	19
Chapter 3. Apparatus	22
3.1. Ion Beam	22
3.2. Ba Getter Source	26
3.3. Solid Xenon Matrix Deposition	27
Chapter 4. Results	28
4.1. Fluorescence of Ba in SXe	28
4.2. Bleaching	29
4.3. Imaging	29
Chapter 5. Conclusions	30
5.1. Future Steps	30
Bibliography	31

CHAPTER 1

Introduction

Neutrinos have been at the forefront of discovery since their prediction by W. Pauli, who proposed in 1930 the existence of a neutral, unobserved particle to explain the apparent violation of energy conservation in beta decay [1]. He admitted that neutrinos (then deemed "neutrons" – what we now know as neutrons had not been discovered yet either) should be difficult to observe experimentally, but also that it seemed unlikely that they would never have been noticed before. As it turns out, they are much more difficult to observe than he predicted; they will not be noticed without extreme experimental techniques.

A theory formulated in 1933 by E. Fermi for beta decay [2], including the neutrino, would be the beginnings of weak theory, and the development of the very successful Standard Model (SM) of particle physics. However, the SM assumes that neutrinos are massless. The discovery of non-zero neutrino mass via neutrino oscillations in the late 1990s [3] shows that there is more new physics to be discovered – a path to follow in the further unification of physical theory.

Neutrinos are extremely difficult to study, but the rewards are profound. The favored See-saw Mechanism is a theory which predicts the extreme lightness of neutrinos, while also predicting very heavy neutrinos. The existence of heavy neutrinos in the high-energy environment of the early universe, coupled with the possibility of the violation of CP conservation which could slightly favor the production of matter vs. anti-matter in decays of these heavy neutrinos, could explain why the universe exists as we know it. [4]

If this theory is correct, then neutrinos would be described by the Majorana formulation rather than Dirac, and the process of neutrinoless double beta decay $(0\nu\beta\beta)$ would be allowed. Observation of $0\nu\beta\beta$ would simultaneously demonstrate that neutrinos are Majorana particles, as well as give a measurement of the absolute mass itself [5]. This chapter outlines the current theory for neutrinos, and then describes the $0\nu\beta\beta$ experiments EXO-200 and nEXO, in order to motivate barium tagging for nEXO.

1.1. Neutrinos

Neutrinos are chargeless leptons which only interact via the weak force (and gravity). There are three known "flavors" of neutrinos, each corresponding to one of the three known leptons: ν_e , ν_μ , and ν_τ . These are the eigenstates in the basis of the weak force, so they are the states in which a neutrino will interact via the weak force.

1.1.1. Neutrino Oscillation and Mass. The postulate that neutrinos have an energy basis which is different from the flavor basis predicts the phenomenon of oscillation — that the time evolution of an initially pure flavor state (as a neutrino will be produced) will result in a time-dependent probability of measuring the other two flavors as well.

The very small mass of a neutrino (assumed zero in the SM), specifically relative to its momentum, lets one write its Hamiltonian in terms of mass squared differences $\Delta m_{ij}^2 = m_i^2 - m_j^2$, where i,j=1,2,3, referring to what we then call mass states. The mass basis is really the energy basis with the small mass approximation, along with dropping some constant terms in the Hamiltonian (which do not affect time evolution). Writing the time evolution in terms of mass squared differences means that neutrino oscillation experiments can produce measurements of these differences. In fact, the discovery of neutrino oscillation was the first (and only, so far) demonstration that neutrinos have a non-zero mass. Without

neutrino mass (particularly without differences between the masses of the mass states), neutrinos would not oscillate.

Neutrino oscillation experiments also provide measurements on the amount of mixing between the flavor basis and the mass basis. We define the mixing between them by a rotation in terms of three mixing angles, θ_{12} , θ_{23} , and θ_{13} . Transformation between the flavor and mass bases is done with the following unitary matrix, called the Pontecorvo–Maki-Nakagawa-Sakata (PMNS) matrix:

$$U = \begin{pmatrix} 1 & 0 & 0 \\ 0 & c_{23} & s_{23} \\ 0 & -s_{23} & c_{23} \end{pmatrix} \begin{pmatrix} c_{13} & 0 & s_{13}e^{-i\delta} \\ 0 & 1 & 0 \\ -s_{13}e^{i\delta} & 0 & c_{13} \end{pmatrix} \begin{pmatrix} c_{12} & s_{12} & 0 \\ -s_{12} & c_{12} & 0 \\ 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} 1 & 0 & 0 \\ 0 & e^{i\alpha_{1}/2} & 0 \\ 0 & 0 & e^{i\alpha_{2}/2} \end{pmatrix}$$

$$= \begin{pmatrix} c_{12}c_{13} & s_{12}c_{13} & s_{13}e^{-i\delta} \\ -s_{12}c_{23} - c_{12}s_{23}s_{13}e^{i\delta} & c_{12}c_{23} - s_{12}s_{23}s_{13}e^{i\delta} & s_{23}c_{13} \\ s_{12}s_{23} - c_{12}c_{23}s_{13}e^{i\delta} & -c_{12}s_{23} - s_{12}c_{23}s_{13}e^{i\delta} & c_{23}c_{13} \end{pmatrix} \begin{pmatrix} 1 & 0 & 0 \\ 0 & e^{i\alpha_{1}/2} & 0 \\ 0 & 0 & e^{i\alpha_{2}/2} \end{pmatrix}$$

where $c_{ij} = \cos \theta_{ij}$ and $s_{ij} = \sin \theta_{ij}$. δ is a phase factor related to lepton CP violation, and α_i are Majorana phases.

Studying oscillations of neutrinos from different kinds of sources, with different energies and path lengths, can isolate sensitivities to the different parameters. For example, the study of solar neutrinos (neutrinos emanating from nuclear fusion reactions in the core of the sun) provides sensitivity to θ_{12} and Δm_{12}^2 . The oscillation parameters so far measured shown in Table 1.1:

TABLE 1.1. Best-fit values for neutrino oscillation parameters, from a global fit to oscillation experiment data. Parameters which depend on the mass hierarchy have values for NH (IH). The atmospheric parameter Δm^2 is defined as $\Delta m^2 = \Delta m^2 = \Delta m^2_{31} - \Delta m^2_{21}/2 > 0 (\Delta m^2 = \Delta m^2_{32} + \Delta m^2_{21}/2 < 0)$. [6]

Parameter	Measurement $(\pm 1\sigma)$
Δm_{12}^2	$7.54_{-0.22}^{+0.26} \ 10^{-5} \ eV^2$
$ \Delta m^2 $	$2.43 \pm 0.06 \ (2.38 \pm 0.06) \ 10^{-3} \ eV^2$
$\sin^2 heta_{12}$	0.308 ± 0.017
$\sin^2 heta_{23}$	$4.37^{+0.033}_{-0.023} (4.55^{+0.039}_{-0.031})$
$\sin^2 heta_{13}$	$0.0234_{-0.0019}^{+0.0020} (0.0240_{-0.0022}^{+0.0019})$
δ/π (2 σ range)	$1.39_{-0.27}^{+0.38} \ (1.31_{-0.33}^{+0.29})$

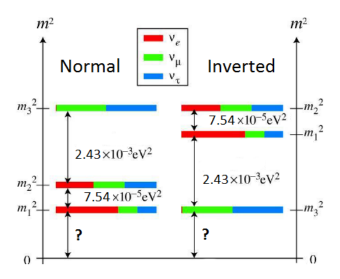


FIGURE 1.1. The two possible hierarchies of neutrino masses. The colors depict the mixing between the mass and flavor bases.

Note that only the absolute value of the atmospheric neutrino oscillation parameter Δm^2 is known. As a consequence, there are two possibilities for the hierarchy of the three neutrino masses. These are called the Normal and Inverted Hierarchies, as shown in Fig. 1.1. The correct mass hierarchy remains unknown, but next-generation neutrino experiments, possibly including nEXO, will be able to discern this.

Neutrino oscillation demonstrates that neutrinos have non-zero mass, and though oscillation experiments measure the mass squared differences, we still do not have a measurement of the absolute masses of the three neutrinos. Cosmology can put limits on the sum of the

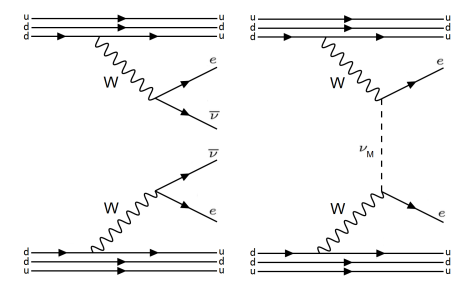


FIGURE 1.2. Two-neutrino (left) and neutrinoless (right) double beta decay.

three neutrino masses. The Planck collaboration reports an upper bound on this sum at $\sum_i m_i < 0.23$ eV [7]. The KATRIN experiment will search for absolute neutrino mass with limiting capability of $m_{\bar{\nu}_e} < 0.2$ eV (90% CL) by measuring the spectrum of tritium beta decay near the Q-value, searching for deviation from the spectrum with zero neutrino mass [8].

1.2. Double Beta Decay

Double beta decay is the simultaneous decay of two neutrons in a nucleus into two protons and two electrons. Two-neutrino double beta decay $(2\nu\beta\beta)$, shown in Fig. 1.2(left), is allowed by the Standard Model and has been observed in eleven isotopes with half-lives between 10^{19} and 10^{21} years. Similar to beta decay, a neutrino accompanies each electron in this decay, broadening the spectrum of the summed electron energy. This is a second-order process, making it a rare decay, and requiring low backgrounds to measure.

 $0\nu\beta\beta$, shown in Fig. 1.2(right), is a postulated mode of double beta decay. In this case, the neutrino is exchanged as a virtual particle (which would require that it is a Majorana

particle), and there are no neutrinos in the final products. If discovered, not only would neutrinos be determined Majorana particles, but their absolute mass could also be measured as it relates to the $0\nu\beta\beta$ half-life according to Eqn. 2:

(2)
$$T_{1/2}^{0\nu} = (G^{0\nu}(Q, Z)|M^{0\nu}|^2 \langle m_{\nu} \rangle^2)^{-1}$$

where $T_{1/2}^{0\nu}$ is the $0\nu\beta\beta$ half-life, $G^{0\nu}$ is a known phase space factor, and $M^{0\nu}$ is a model-dependent nuclear matrix element. The effective electron neutrino mass $\langle m_{\nu} \rangle$ is the expectation value of the mass for a pure electron neutrino:

$$\langle m_{\nu} \rangle = \sum_{i} U_{ei}^{2} m_{i}.$$

The sum of the energies of the emitted electrons in double beta decay will serve as the distinction between the two-neutrino and zero-neutrino modes, shown in Fig. 1.3. In the two-neutrino mode, the total decay energy is shared probabilistically between the electrons and the neutrinos (the nucleus recoil energy is negligible), resulting in a broad distribution in the summed electron energy. (Recall the similarly broad electron energy in single beta decay, which ultimately led to discovery of the neutrino involved.) But in the zero-neutrino mode, all of the decay energy is carried away by the two electrons, resulting in only a single allowed value for the summed electron energy – a peak in the summed electron energy spectrum at the Q-value.

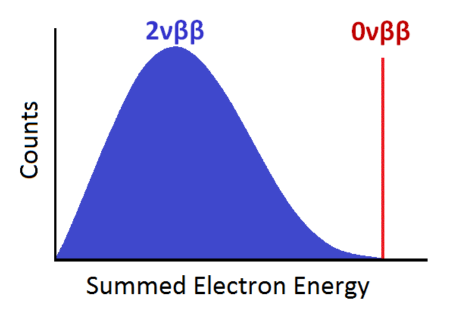


FIGURE 1.3. Conceptual two-neutrino (blue) and zero-neutrino (red) double beta decay spectra.

The rarity of double beta decay requires very low backgrounds, especially around the Q-value for the $0\nu\beta\beta$ search. The next sections describe the experiments EXO-200 and it's next-generation successor, nEXO.

1.3. Enriched Xenon Observatory

EXO-200 and nEXO (EXO standing for Enriched Xenon Observatory) are a progression of two experiments, each a LXe time projection chamber (TPC) designed to study the double beta decay of the isotope ¹³⁶Xe, and ultimately to search for the zero-neutrino mode. Xe is fairly unique among the double beta decay isotopes in that it can be studied in a gas or liquid TPC instead of solid crystals or foils. The 3D event position reconstruction abilities of a TPC have advantages in background reduction, described in section 1.3.1. Purification of Xe is straightforward and can be done continuously in a detector. LXe is transparent, and produces substantial ionization and scintillation at 178 nm when energy is deposited in the LXe [9]. A liquid TPC approach also offers the opportunity to identify, or "tag", the

daughter Ba⁺⁺ at the site of the double beta decay event, which would provide a backgroundfree identification of $0\nu\beta\beta$ [10]. Barium tagging is the focus of our group at CSU and is the subject of this thesis. The following sections decribe the EXO-200 experiment, as well as nEXO, the next-generation tonne-scale LXe TPC which is now in the design stages. EXO-200 does not have barium tagging implemented, but it is hoped that nEXO will.

1.3.1. EXO-200. EXO-200 has been operational since 2011. It is a TPC using 110 kg of active LXe enriched to $80.672\% \pm 0.14\%$ ¹³⁶Xe [9], designed to probe Majorana neutrino masses down to around 109-135 meV (the range covers different matrix element calculations) [11], and is located about half a mile underground in the Waste Isolation Pilot Plant (WIPP) near Carlsbad, NM. This mine is in a salt basin, which contains lower levels of Uranium and Thorium than a more typical mine in rock.

A schematic of the TPC in the class 100 cleanroom is shown in Fig. 1.4. Several layers of lead wall surround the copper cryostat, which is filled with HFE-7000, a cryogenic fluid which keeps the TPC cooled to LXe temperatures, as well as aids in shielding. The TPC vessel is made of low-radioactivity copper, and is kept as thin as possible to minimize backgrounds. Scintillating panels on the outside of the cleanroom provide muon veto.

A cut-away view of the EXO-200 detector is shown in Fig. 1.5. It is two mirrored TPCs which share a cathode. The detection planes are a combination of ionized charge induction/collection wires and large area avalanche photodiodes (LAAPDs) which detect scintillation light [12].

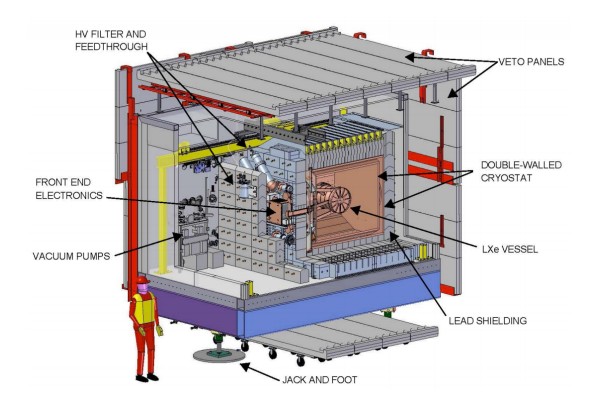


FIGURE 1.4. Drawing of cleanroom laboratory in the WIPP drift.

A photograph of the detection plane is shown in Fig. 1.6, and a schematic of event detection is shown in Fig. 1.7. When a double beta decay event occurs in the LXe, the energetic electrons ionize many surrounding Xe atoms, as well as produce a scintillation signal. Scintillation light provides a time stamp for the event as well as an energy measurement.

The cathode is set to -8 kV, providing an electric field of 374 V/cm across the 20 cm drift length of each TPC. Ionized electrons drift from the decay site, first passing the v-wires, which receive an induction signal, and are then collected by the u-wires, which are set at a 60° angle from the v-wires. An electric field of 778 V/cm between the u- and v-wires ensures 100% v-wire transparency. The charge collection provides an energy measurement complimentary to that of the scintillation. Together, the u- and v-wires give an x/y position measurement for the event. The time between the initial scintillation detection and the charge collection give a

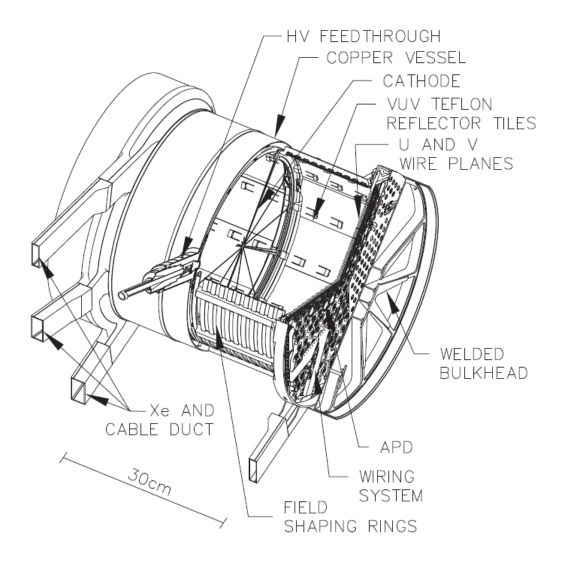


FIGURE 1.5. EXO-200 TPC cutaway [9].

z position, and a 3D position can be reconstructed for the event. [9] Additional scintillation resulting from recombination of ionized electrons leads to a well-known microscopic anticorrelation between ionization and scintillation signals [13]. Applying this to the combination of those signals improves the energy resolution. This correction defines a combined energy axis, called the rotated energy. Energy resolution is important in a $0\nu\beta\beta$ search, as it distinguishes those events from $2\nu\beta\beta$ events in the tail of their spectrum.

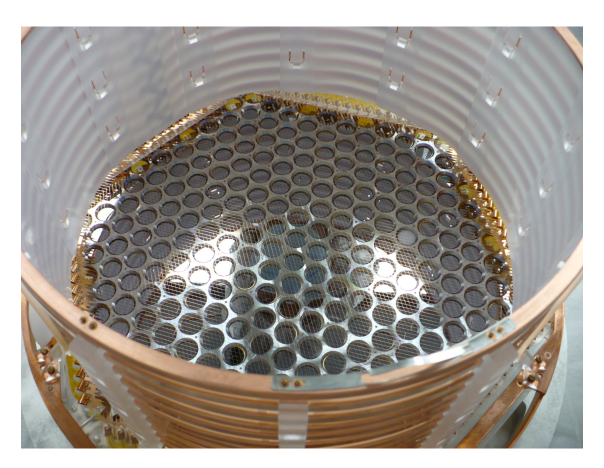


FIGURE 1.6. View of the detection plane in one of the two EXO-200 TPCs.

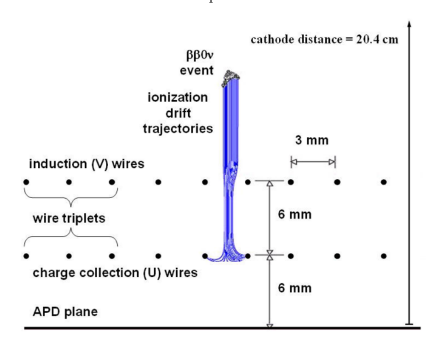


FIGURE 1.7. EXO-200 event detection.

Having a reconstructed 3D event position is important in several ways. Firstly, positionbased corrections on scintillation and charge collection can be applied. For charge, electrongative impurities in the LXe will absorb the drifting charge, requiring a drift-length (z-position) correction. High purity levels, measured in terms of electron lifetime, of 2- 3×10^3 µs are maintained in EXO-200, resulting in a small correction of few % for maximal drift lengths. For scintillation, a 3D correction is applied, as some regions have more efficient light collection by the LAAPDs. A 3D position also allows a fiducial volume to be defined. The effect of radio-impurities on detector surfaces, e.g. radon daughters collecting on the cathode, is mitigated by defining a stand-off distance required for events used in analysis. Finally, 3D reconstruction allows the distinction between single-site (SS) and multi-site (MS) events. A MS event is one where two spatially separated events occur in the same $2048-\mu s$ time window. These are mostly caused by gamma rays interacting in the LXe, which can Compton-scatter several times. Rejecting MS events strongly separates gamma events from double beta decay events. [9] Of course, barium tagging will also require a 3D reconstructed position in nEXO.

The final data set is fit using a combination of probability distribution functions (PDFs) for $0\nu\beta\beta$, $2\nu\beta\beta$, and all possible backgrounds. Fits to the final energy spectrum data for (a) SS events, and (b) MS events are shown in Fig. 1.8 for the most recent $0\nu\beta\beta$ search analysis. The green bands beneath each plot show the residuals vs. energy. The $2\nu\beta\beta$ spectrum, in gray, dominates the backgrounds in the SS spectrum. The vertical red lines in the SS spectrum outline the $\pm 2\sigma$ region of interest around the Q-value, where the $0\nu\beta\beta$ peak will lie. The insets are a zoom into this region. The fit value for $0\nu\beta\beta$ in this dataset is non-zero, but it is consistent with the null hypothesis at 1.2σ . This sets an upper limit on the

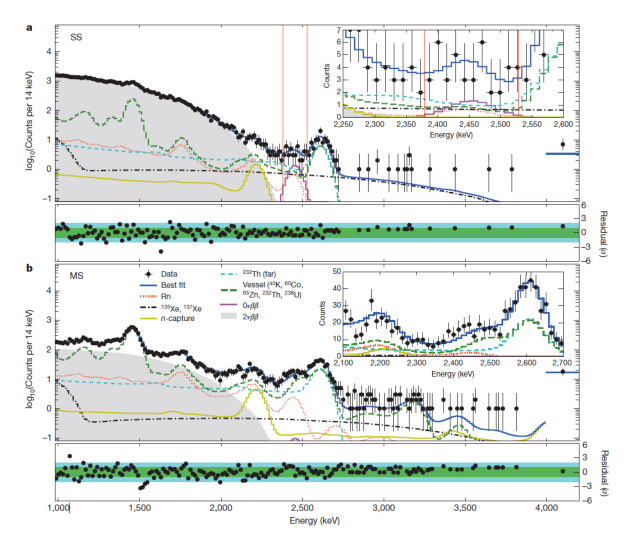


FIGURE 1.8. EXO-200 energy spectrum from $0\nu\beta\beta$ search analysis [14].

 $0\nu\beta\beta$ half-life at $T_{1/2}^{0\nu\beta\beta}$ < 1.1 × 10²⁵ yr (90% CL), which corresponds to $\langle m_{\nu_e} \rangle$ <190-450 meV depending on nuclear matrix element calculations [14]. EXO-200 reports the $2\nu\beta\beta$ half-life measurement in [9] of $T_{1/2}^{2\nu\beta\beta}$ = 2.165 ± 0.016(stat)±0.059(sys)×10²¹ yr. This is consistent with previous measurements by EXO-200 in 2011 [15] and KamLAND-ZEN in 2012 [16].

1.3.2. NEXO. The next-generation successor to EXO-200 is nEXO, a tonne-scale LXe TPC which will probe Majorana neutrino masses down to the 10 meV scale. The sensitivity projections for nEXO are shown in Fig. 1.9, along with those of EXO-200. nEXO will reach phase space where the two possible mass hierarchies begin to split; if nEXO successfully

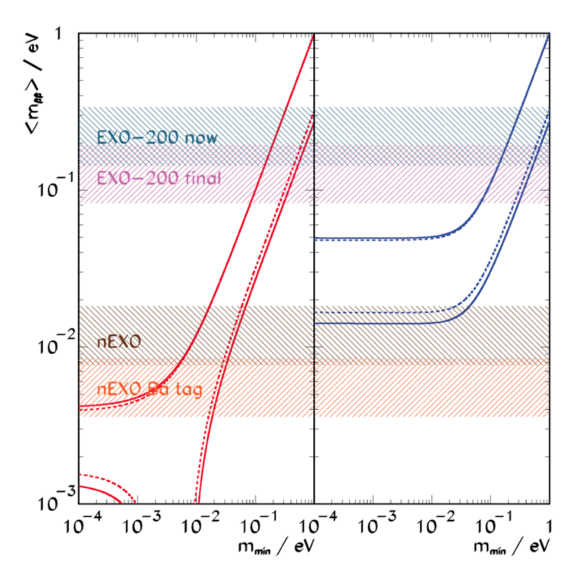


FIGURE 1.9. nEXO projected sensitivity to the Majorana neutrino mass vs. the minimum of the three masses m_{min} . check for updated version

observes $0\nu\beta\beta$ in these regions, it may be able to also determine the mass hierarchy. Barium tagging will push the sensitivity further into the region allowed only by the normal hierarchy [ref for this?].

A schematic of the experimental setup is shown in Fig. 1.10 in one of the possible locations for nEXO, the SNOLAB cryopit. Similar to EXO-200, the copper-housed TPC will be submerged in HFE fluid, inside a copper cryostat. The cryostat will be insulated and submerged in a large volume of water shielding, in which photo-multiplier tubes could

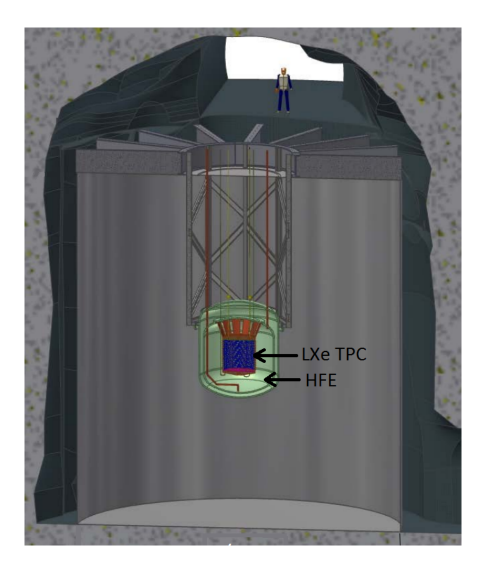


FIGURE 1.10. nEXO TPC in the SNOLAB cryopit.

provide muon veto by observing Cherenkov radiation. nEXO will be a single TPC with charge and light readouts at opposing ends of the TPC. Rather than wires, nEXO will use tile electrodes for charge readout, shown in Fig. 1.11.

1.3.3. Barium Tagging. The backgrounds observed around the Q-value in EXO-200 can be expected to scale up with the size of nEXO. But with the veto power of barium tagging, nEXO's sensitivity to $0\nu\beta\beta$ scales with the total ¹³⁶Xe mass in the detector, vs. the square root of that mass without barium tagging.

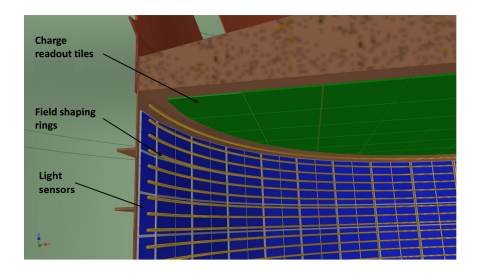


FIGURE 1.11. nEXO readout.

Several possible barium tagging techniques have been proposed. Perhaps the most natural concept is to direct one or more lasers at the decay site to induce fluorescence of the barium daughter. This technique was explored thoroughly by Fairbank's group, and was abandoned when reliable fluorescence of Ba⁺ in LXe could not be observed [ref Kendy's thesis, after kamland in list].

Three barium tagging techniques continue to be explored. One of these is to grab the daughter on a surface, brought to the decay site by a probe, and then move it to a location where it can be desorbed from that surface by an infrared laser, and subsequently resonantly ionized by two lasers in order to detect it by time-of-flight spectroscopy. The apparatus for the study of this method is described in [17], along with some initial results. Another concept is to also grab the daughter on the surface of a probe, but then to thermally desorb it for subsequent trapping and detection [ref upcoming Liang paper, after Twelker].

The other remaining technique for barium tagging, now the concentration of Fairbank's group and the subject of this thesis, is called tagging in SXe. Here, a cold probe would go to the site of the candidate $0\nu\beta\beta$ event in order to trap the Ba/Ba⁺ in a small amount of SXe,

and then observe it by its laser-induced fluorescence in the SXe, a technique called matrix isolation spectroscopy.

A concept for a probe is shown in Fig. [ref fig cryoprobe]. The SXe forms on a sapphire window or plug. Sapphire is a good candidate for a substrate; it has good thermal conductivity at low temperature, and is extremely transparent. An excitation laser would be brought into the probe through a fiber, and aimed through the sapphire to excite the Ba/Ba⁺ in the SXe. A return fiber could collect the laser reflections to measure the SXe thickness via interference fringes. [in the fig., label things like the fibers like (A), (B), and just say what those are in the caption] The Ba/Ba⁺ fluorescence would then be collected by a lens/filter system and imaged onto a CCD. Additional components, not shown, would be required for cooling the sapphire, either by liquid He or by a Joule-Thompson nozzle, as well as a thermometer.

Whether the daughter Ba will neutralize or remain ionized is not yet known. It is however expected that a Ba⁺⁺ ion will neutralize once to Ba⁺ in lXe, since the lXe conduction band gap is slightly less than the ionization potential for Ba⁺ [10]. Further neutralization could also occur. A study of neutralization of alpha decay daughters in EXO-200 has observed a fairly equal balance of neutralized daughters vs. ionized daughters [ref alphaion]. The feasibility of detecting single Ba ions as well as atoms is still of interest.

CHAPTER 2

THEORY

Theory relevant to the spectroscopy and detection of single Ba/Ba⁺ in SXe matrices is discussed.

2.1. Ba/Ba⁺ Spectroscopy in Vacuum

The lowest-lying energy levels in vacuum for Ba and Ba⁺ are shown in Fig. 2.1. For Ba, the main transition is between the ground $6s^2$ 1S_0 to the excited 6s6p 1P_1 state. Spin(?)-suppressed transitions between the P state and three metastable D states results in a decay in to a D state after about 350 excitations. For Ba⁺, two strong transitions exist between the ground 6s $^2S_{1/2}$ is the doublet correct?? and the 6p $^2P_{1/2}$ and 6p $^2P_{3/2}$ excited states. Transitions to the two metastable D states are higher than for the atom, resulting in a decay into a D state after about 4 excitations.

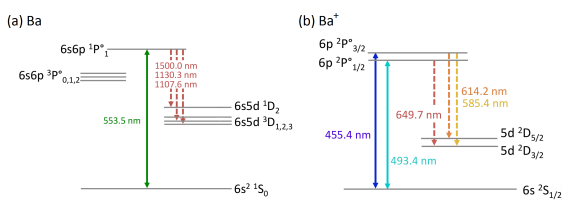


FIGURE 2.1. Energy level diagrams for (a) Ba and (b) Ba⁺

These energy levels and their transition rates are well known, and are documented in the NIST tables [ref]. Single atom/ion detection by spectroscopy generally requires lasers to provide transitions out of the metastable D states once the atom/ion decays into one of them, in addition to the main excitation laser. For the atom, this requires three additional infrared lasers, and single atom trapping/detection in a magneto-optical trap (MOT) is achieved in [ref Ba MOT]. Single Ba⁺ observation requires only two lasers if the ${}^{2}P_{1/2}$ excited state is used. This is demonstrated in [ref something that does this ... is there something? How about the Carleton group? I think MOE has a reference to it maybe – yeah, its [19]].

2.1.1. 5- OR 6-LEVEL MODEL? This could be for just the vacuum, and the in-matrix could reference it with changes (so I guess this would just be 5-level).

2.2. Matrix Isolation Spectroscopy

The spectroscopy of a species trapped in an inert solid matrix is called matrix isolation spectroscopy, the concept of which was pioneered around 19?? by [that one guy] [ref ... maybe [1] of ba spec]. Though absorption and emission are significantly broadened and shifted, a species can retain properties of its vacuum counterpart, such as quantum numbers.

Studies of some matrix isolation systems have been made. The most thoroughly studied has been Na(?) in solid matrices of (Ar, Kr, Xe?)... (Na? Mg?)...[refs]

We are of course interested in the spectroscopy of Ba and/or Ba⁺ in SXe matrices, and this particular system has not been studied until recently. The first report of the spectroscopy of neutral Ba in SXe, along with candidate fluorescence peaks for Ba⁺ in SXe, was published by our group in [ref ba spec], and conversations following this publication with ???'s group in ??? have confirmed our basic observations of the absorption spectrum of Ba in SXe.

From here forward I will refer to the host species as Xe and the guest as Ba/Ba⁺, specific to our system. The leading interaction between [put this here if the van der waals is only necessarily the force for Ba in particular] the Ba/Ba⁺ and a neighboring Xe atom is a (the) Van Der Waals (sp?) force, an induced dipole-dipole interaction. Xe is the most polarizable (sp?) of the noble gases.

The forms for this force(?) is shown in Eqn. [ref van der ba] for Ba and Eqn. [ref van der ba+] for Ba⁺. Talk about it.

This binding between the Ba/Ba⁺ and its surrounding Xe atoms results in vibrational modes, and the Franck Condon (sp?) principle applies. Fig. [ref fig franckcondon] helps illustrate this effect. In a cold matrix, the system will be in the ground vibrational state before excitation. The distribution of the wavefunction, even for the ground state alone, overlaps in space in general with more than one of the excited state vibrational modes, resulting in a broadening in energy of the absorption.

Rapid decay occurs to the lowest vibrational mode in the excited state before electronic decay can occur [ref] [is this just for Ba?]. Then a similar broadening in emission energy occurs as several overlapping vibrational mode wavefunctions exist in the ground electronic state, and a redshift is also observed as some energy was dissipated into phonons in the crystal in vibrational mode decays. (How can you get a blueshift?)

Shifts and broadening in absorption and emission depend in general on the distribution of Xe atoms around the Ba/Ba⁺. The fcc crystal structure of the Xe restricts these environments to discrete number of so-called matrix sites, defined by (a) whether the Ba/Ba⁺ is [intersituational or whatever – is that still a thing?], and (b) the number of vacancies in the Xe matrix surrounding the Ba/Ba⁺. Experimental observation of emission peaks from different matrix sites of [Na] in solid noble gases is reported in [ref(s)], with theoretical calculations in [ref(s)] attributing observed peaks to specific vacancy distributions defining the matrix sites (maybe this has been done?).

Energy level transition probabilities can also be affected in a matrix. Distortions in electron wavefunction shapes by asymmetric matrix sites can affect radiative transitions by altering parity [ref]. If electronic potential energy curves cross each other, nonradiative transitions can also become allowed for otherwise forbidden transitions [ref]. (Is a phonon emission called a nonradiative transition / does this actually happen?)

 ${\it Jahn-Teller?}$

The detectability of a single atom/ion can depend greatly on altered transition rates. For example, in the Ba atom, matrix-allowed decay of the ¹P₁ to the ³P states could be much stronger than the main transition back to ground, which would suppress fluorescence and make single-atom detection impossible. But the matrix could also help by strengthening decays from the D states back to ground, eliminating any need for re-pump lasers to keep the transition cycle going.

CHAPTER 3

Apparatus

This chapter describes the apparatus at Colorado State University, which we have used for all described studies of Ba/Ba⁺ fluorescence in SXe after deposition in vacuum. Our main barium source, the Ba⁺ ion beam, is first described, as well as a purely Ba neutral source. The co-deposit of Ba/Ba⁺ with Xe gas onto a cold sapphire window, subsequent laser excitation, and finally the collection optics for the fluorescence are described.

3.1. Ion Beam

The full ion beam is shown in Fig. 3.1. This is a clean source of Ba⁺ which can do a very wide range of deposit sizes, from billions of ions in a focused laser region all the way down to the single-ion level, and even deposits sparse enough to scan a laser over spatially separated ions (may only want to say that if we have those scans).

3.1.1. Ion Source/Acceleration. Ba⁺ ions are produced in a Colutron [type?] ion gun system [reference], shows Fig. [ref fig with ion source, showing Ar gas as well as Ba charge, and lens with ion path]. A solid barium charge is placed into the hollowed end of a stainless steel rod, which is then inserted into the discharge chamber, near the hot filament. The heated barium vaporizes, and is allowed to escape the hollowed rod around a loosely threaded set screw at the end of the rod.

The source is designed to produce a discharge between the anode plate and the filament cathode, through an argon buffer(?) gas leaked into the source chamber. This controlled discharge would then also ionize atoms, vaporized from the solid charge, to produce the desired ion beam, and Ar ions would be filtered out. However, to avoid contamination of

our solid matrix with residual Ar gas, we do not use the buffer gas in the ion source. We are still able to maintain a discharge between the filament and anode circuits. This discharge produces a plasma, containing barium ions, which escapes the chamber through a small hole in the anode, where it enters the acceleration potential.

The longevity of our ion current from a single charge (at least several 10s of hours) suggests that Ba is coating the inner walls of the chamber and is depleted slowly. This is supported by the observation of white oxidation of the inner source parts after a few minutes of exposure to air when opening the system.

The acceleration potential is 2 kV, between the ion source anode and an aperture, which is the first element of the Einzel lens 1. This lens approximately collimates the ion beam for passage through the E×B velocity filter.

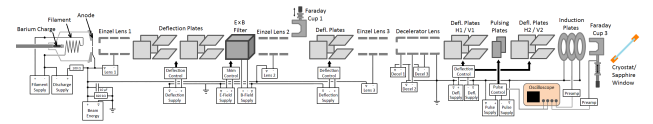


FIGURE 3.1. Ba⁺ ion beam. landscape this

3.1.2. E×B VELOCITY FILTER. The E×B velocity filter selects Ba⁺ by creating perpendicular electric and magnetic fields, which produce opposing forces on charged particles moving straight through the filter. The opposing forces will be equal for ions with velocity $v = \frac{E}{B}$. Since ion velocity is determined by mass (m), charge (q) and beam potential (V), the filter selects ions satisfying Eq. 4:

$$\frac{m}{q} = \frac{2VB^2}{E^2}$$

where B and E are the magnetic and electric fields, respectively. Those fields are chosen such that the forces are equal for Ba^+ , while other ions will be deflected.

The E×B filter is shown in Fig. 3.2. Electromagnets provide the vertical magnetic field. Electrode plates and field-shaping guard rings provide the horizontal electric field. The guard rings prevent a lensing and astigmatism effect from fringe fields of the plates [20].

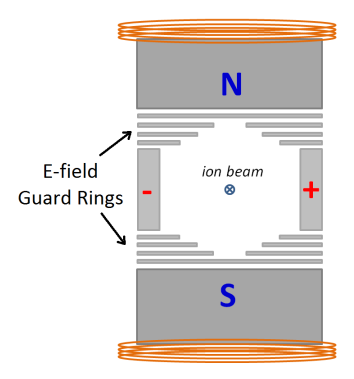


FIGURE 3.2. Colutron E×B ion velocity filter.

To determine the mass components of the beam, the electric field can be scanned. Fig. [ref mass scan fig] shows a scan of the Ba⁺ beam, as well as of an Ar⁺ beam. We found it necessary to run with the known mass of Ar in order to calibrate the magnetic field, which came to differ from the calculation given by Colutron [ref like Shon has?], likely due to hysteresis in the magnet. The Ba⁺ peak agrees with the Ba mas s...

3.1.3. Other Beam Components. The first three sets of deflection plates can be used for beam diagnostics, and are set to 0 V during normal operation. The set just before the

pulsing plates, called H1 and V1, are set to constant values of 50 V and 0 V, respectively, which have been selected such that the beam, in both pulsing and continuous modes, can be deposited at the sapphire window for reasonable settings on the final deflection plates, H2 and V2. As described in 3.1.5, different settings in H2/V2 are required for peak ion current in cup 3 vs. peak deposit at the window.

Einzel lens 2 focuses the beam the pass through the aperture in the first element of the decelerator lens. Einzel lens 3 is not used in this setup. The decelerator lens can be used to vary Ba⁺ deposit energy, but it in this work it acts as an Einzel lens with only the second element at voltage, and it focuses the beam at Faraday cup 3 (there is no Faraday cup 2 in this setup). Faraday cup 3 measures the ion current during experiments, and is retracted when deposits are being made. Its use in deposit calibration is described in 3.1.5. Faraday cup 1 is used for beam diagnostics, and is usually retracted on its bellows.

3.1.4. ION BEAM PULSING. When running in pulsing mode, the pulsing plates are first set to 200 V and -200 V to deflect the beam, and are pulsed to 0 V for 2 μ s for each pulse. The pulsing circuit is shown in Fig. [ref fig pulsing circuit, ref Shon thesis if this is same figure]. Square waves, triggered by LabVIEW at 50 Hz, enter the circuit at [x]. [describe how the pulse happens]

The induction plates observe pulses during a deposit. Pulses just prior to a deposit can be observed by cup 3 as well as the induction plates, for a local measurement of ion current in the pulses. An example of an oscilloscope readout of 16 averaged pulses is shown in Fig. [ref fig scope pulse – in caption, explain v-divided plate signals].

[x brand, or whatever] pre-amplifiers convert the ... [put this before scope picture?] [have a figure of the shaped plots too]

Pulsing data also provides additional confirmation that the beam is composed of Ba⁺. [figure and data (may need to average Ar+ as well as Ba+ stuff, w/ some kind of informed pulse data feature to determine velocity) showing velocity measurement]

3.1.5. Calibration of Ion Deposit. To calibrate the signal at cup 3 to an ion density at the sapphire window, another Faraday cup (called cup w) is attached to the cold finger in place of the sapphire window. Firstly, the ratio in $\frac{fC}{pulse}$ between cup 3 and cup w is measured. Then, knowing the radius of cup w lets one determine the ion density per pulse at the sapphire window:

$$\frac{ions}{pulse \times m^2} = \frac{\sigma}{Ae}$$

where σ is

The required settings of the final deflection plates H2 and V2 are also determined by cup w. These typically differ from the peak values for cup 3 by about ???, corresponding to ??? mm.

Talk about beam size?

3.2. BA GETTER SOURCE

Ba getters are neutral Ba sources typically used to grab reactive gas molecules in purifiers and vacuum tubes. We can use a getter as a source of neutral Ba. It is very helpful to have a completely different Ba source, especially a neutral-only source, in identifying observed fluorescence peaks. This is discussed further in 4.1.1.

The getter used briefly in this work is an endothermic ... getter wire ... [describe it] ... show diagram, maybe also of it in the system at same position as cup 3.

3.3. SOLID XENON MATRIX DEPOSITION

The final destination of the barium ions is of course in the solid xenon matrix, which is deposited onto a cold sapphire window. Sapphire has good thermal conductivity, good optical transparency in the visible, and does not fluorescese in the wavelength region where barium fluoresces.

Xenon freezes around 73 K (?) at our pressures $(0.5 - 1 \times 10^{-7} \text{ Torr } ... \text{ it's lower when cold, can you determine what that is?}), so the window is cooled to temperatures below that. The window is held to a cold finger (Fig. 6x, picture of), cooled by a -brand- cryostat.$

3.3.1. Deposition Procedure. Before barium ions are let through, xenon gas is allowed to flow, controlled by a leak valve, onto the cold sapphire window, where it freezes and begins growing the solid matrix. The Faraday cup is then retracted, to clear the path for barium ions. The cup serves as a shutter for DC deposition, or if pulsing is being used, they are performed at this time. Barium ions land in the solid xenon as the matrix continues to grow. The cup is then replaced, and the xenon leak stopped.

- 3.3.2. Laser Excitation.
- 3.3.3. COLLECTION OPTICS. talk about all that, including spectrometer

CHAPTER 4

RESULTS

Improved studies of several emission peaks of neutral Ba in SXe, which are first reported in the theses of B. Mong [ref] and S. Cook [ref], are reported. The bleaching of several of these peaks is carefully observed (do a correction on p-meter sensitive area and on p-meter quantum efficiency, and also a spherical aberation correction for power??), and a model of atomic transition rates is fit to this data. Improved excitation spectra are achieved for all observed Ba emission peaks. Images of $\leq 10^{3?}$ Ba atoms are achieved using the bleaching peaks. Ultimately, images of Ba atoms at the few-atom level are achieved using the 619-nm fluorescence peak. Improved studies of candidate fluorescence peaks of Ba⁺ in SXe are also reported. Where/how should I reference the ba spec paper?

4.1. Fluorescence of Ba in SXE

The multiple peaks observed from deposits of Ba in SXe are attributed to Ba atoms occupying different Xe matrix sites, the phenomena of which are described in Chapter 3 (ref chap theory doesn't work). Fig. [ref fig spectra for 10K, 50K, annealed 10K] shows spectra of Ba deposits under different conditions. A deposit made at 11 K shows a different ... um how is this described in the paper?

excitspec of 50 K [fig of all, including r110 of 619]... and 10 K? leak rate dependence?

4.1.1. Identifying Peaks as Ba in SXE.

4.2. Bleaching

590 etc, model fit (see results intro paragraph)

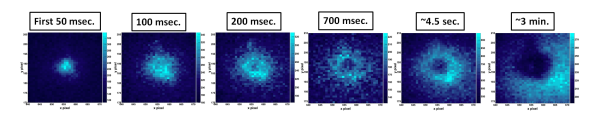


FIGURE 4.1

619, with the changes in time and I

4.3. Imaging

- 4.3.1. Imaging 577- and 591-nm peaks.
- 4.3.2. Imaging 619-nm peak.

$CHAPTER \ 5$

Conclusions

5.1. Future Steps

Talk about whatever Chris will do next ... ions? Other sites? Populations? Talk about Adam's work, show pics and stuff.

BIBLIOGRAPHY

- [1] C.D. Ellis and W.A. Wooster, "The average energy of disintegration of radium e,"

 Proceedings of the Royal Society (London) A117 109-123 (1927).
- [2] F. Wilson, American Journal of Physics 36, 12 (1968).
- [3] Y. Fukuda et al. (Super-Kamiokande Collaboration), Phys. Rev. Lett. 81 1562 (1998).
- [4] S.T. Petcov, W. Rodejohann, T. Shindou, Y. Takanishi, Nuclear Physics B 739, 208233 (2006).
- [5] M. Danilov et al., Physics Letters B 480, 1218 (2000).
- [6] K.A. Olive et al. (Particle Data Group), "14. Neutrino Mass, Mixing, and Oscillations,"
 Chin. Phys. C 38, 090001 (2014) (http://pdg.lbl.gov)
- [7] P.A.R. Ade et al. (Planck Collaboration), Astronomy and Astrophys. 571, A16 (2014).
- [8] N. Steinbrink et al., New Journal of Physics 15, 113020 (2013).
- [9] J. Albert et al. (EXO-200 Collaboration), Phys. Rev. C 89, 015502 (2014).
- [10] M. Moe, Phys. Rev. C 44, R931 (1991).
- [11] M. Auger et al., JINST 7, P05010 (2012).
- [12] R. Nielson et al., Nuclear Intruments and Methods in Physics Research A 608, 68 (2009).
- [13] E. Conti et al., Phys. Rev. B 68, 054201 (2003).
- [14] J. Albert et al. (EXO-200 Collaboration), Nature **510**, 229 (2014).
- [15] N. Ackerman et al. (EXO-200 Collaboration), Phys. Rev. Lett. 107, 212501 (2011).
- [16] A. Gando et al. (KamLAND-Zen Collaboration), Phys. Rev. C 86, 021601 (2012).
- [17] K. Twelker et al., Review of Scientific Instruments 85, 095114 (2014).
- [18] B. Mong et al., Phys. Rev. A 91, 022505 (1954).
- [19] T. Brunner et al., International Journal of Mass Spectrometry 379 (2015) 110-120.

[20] L. Wåhlin, "The Colutron, a Zero Deflection Isotope Separator," Nuclear Instruments and Methods ${f 27}$ 55-60 (1964).