Measurements of Ultracold-Neutron Lifetimes in Solid Deuterium


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We present the first measurements of the survival time of ultracold neutrons (UCNs) in solid deuterium (SD$_2$). This critical parameter provides a fundamental limitation to the effectiveness of superthermal UCN sources that utilize solid ortho-deuterium as the source material. These measurements are performed utilizing a SD$_2$ source coupled to a spallation source of neutrons, providing a demonstration of UCN production in this geometry and permitting systematic studies of the influence of thermal up-scatter and contamination with para-deuterium on the UCN survival time.

Neutrons with kinetic energies less than 340 neV can be trapped in material bottles and are referred to as ultracold neutrons (UCNs) [1–3]. UCN densities at reactor sources have gradually increased with reactor power and improved techniques for extracting the UCN flux. The highest bottled densities reported in the literature, 41/cm$^3$, have been obtained at the Institut Laue-Langevin (ILL) reactor in Grenoble [4].

Measurements of the neutron electric dipole moment [5,6] and the neutron lifetime [7–9] attest to the utility of bottled UCNs for fundamental experiments with neutrons. UCNs may also prove useful in improved measurements of angular correlations in neutron beta decay [10,11]. All of these experimental programs are limited by the available densities of UCNs.

A superthermal UCN source was first proposed in 1975 by Golub and Pendlebury [12] in superfluid $^4$He and experimentally investigated shortly thereafter [13,14]. In this process, phonon creation in the liquid is used to down-scatter cold neutrons to the UCN regime. Up-scattering is suppressed by maintaining the superfluid at sufficiently low temperature. Because $^4$He has no nuclear absorption, the limitations to the density of UCNs accumulated are wall losses and neutron beta decay. The production of UCNs by this process agrees with theoretical expectations [14–18].

While superfluid $^4$He is an excellent superthermal converter, a few other materials, such as solid deuterium (SD$_2$), satisfy the criteria for superthermal production. The limiting UCN density, $\rho_{UCN}$, one can obtain using a SD$_2$ source is given by the product of the rate of UCN production, $R$, and the lifetime of UCNs in the solid, $\tau_{SD}$: $\rho_{UCN} = R\tau_{SD}$. A storage bottle opened to such a source will come into density equilibrium with the density in the solid. This idea led to the concept of a source in which the inside of a neutron bottle is coated with a thin layer of SD$_2$ and the bottle is embedded in a cold-neutron flux [19,20]. The volume comes into equilibrium with the UCN density with a time constant for the coupled system, $\tau$, given by

$$\tau = \tau_{SD} \frac{V}{V_{SD}},$$

where $V_{SD}$ is the volume of SD$_2$ in the source and $V$ is the total volume of the storage bottle. A valuable summary of SD$_2$ thin-film sources is presented in [21]. The effects of gravity and of the potential of the solid as well as UCN losses other than absorption in the film, neglected in this expression, do not fundamentally alter this picture. The limit to the UCN density is established by the tradeoff between the cold-neutron flux intensity and energy distribution (which determine the production rate) and heating by neutrons and gamma rays in the SD$_2$ and bottle walls, because $\tau_{SD}$ is a strong function of temperature. However, the predicted production rates in SD$_2$ [20] and lifetimes [22] have not yet been quantitatively verified. Efforts to utilize SD$_2$ sources at reactors identified possible gains but suffered from problems with cooling the solid at full reactor power [13].

Pokotilovski pointed out the advantages of UCN production in SD$_2$ at pulsed neutron sources [23] and showed that UCN densities 2–3 orders of magnitude greater than existing reactor-based UCN sources might be possible.
The use of spallation as a pulsed source has been suggested [24,25]. In a spallation UCN source cold neutrons, produced by moderating spallation neutrons produced in a heavy target by a medium-energy pulsed proton beam, are used to drive a SD$_2$ superthermal converter. The amount of heating for each neutron is lower than in a reactor allowing higher neutron densities in the vicinity of a spallation target to be achieved. Even higher neutron densities can be obtained by pulsing the proton beam and valving off the UCN storage volume from the production volume when the beam is off, using this time to remove heat from the deuterium. The maximum UCN density that is produced is limited by the impulse heating of the SD$_2$. Experiments with the stored UCNs can be performed while the beam is off with low backgrounds.

We have operated a test source with single pulses of protons produced by the LANSCE 800 MeV proton accelerator at Los Alamos National Laboratory. In this Letter, we report the measurements of $\tau_{\text{SD}}$, the lifetime of UCNs in SD$_2$ made using this source. These measurements demonstrate the influences of heating and para-deuterium contamination on the UCN lifetime, and provide a quantitative foundation for the development of SD$_2$ superthermal sources.

A schematic view of our apparatus is shown in Fig. 1. Spallation neutrons were produced in a tungsten target with less than 160 ns long pulses of 800 MeV protons provided by the LANSCE accelerator. The fast neutron flux was amplified using $(n, 2n)$ reactions in a beryllium reflector surrounding the spallation target. The spallation neutrons were moderated and cooled in a thin layer of polyethylene surrounding a $^{58}$Ni-coated stainless steel guide tube with an inner diameter of 7.8 cm. The polyethylene and the bottom of the guide were cooled with liquid helium, and a layer of deuterium was frozen on the inside of the guide. UCNs produced in the SD$_2$, confined by the guide tube, could be directed through a series of valves to the UCN detector. Neutrons were detected in a 5-cm-thick multiwire chamber detector filled with a mixture of $^3$He at 5 mbar and CF$_4$ at 1 bar. The low $^3$He pressure and the large bend angle in the guide resulted in a high degree of selectivity for detecting UCNs in the apparatus. Data were acquired using a multiscalar that was started by the proton beam passing through a toroidal pickup coil and that scaled the count rate from the detector.

Up-scatter from phonons in the solid [20], up-scatter from para-deuterium molecules in the solid [22], absorption on deuterium, and absorption on hydrogen impurities limit the lifetime of UCNs in SD$_2$. All of these effects have been calculated. The total loss rate is a sum of contributions from these sources:

$$1/\tau_{\text{SD}} = 1/\tau_{\text{phonon}} + 1/\tau_{\text{para}} + 1/\tau_{\text{D, abs}} + 1/\tau_{\text{H, abs}}$$  

with the loss rate due to phonon up-scatter having different contributions from the ortho- and para-deuterium in the solid. Establishing the experimental basis to validate these models is essential for the design of a UCN source based on SD$_2$.

SD$_2$ was frozen in the lower part of the cryostat using a helium transfer refrigeration. The temperatures of the lower guide walls and of the liquid-helium cryostat were monitored with an array of silicon diodes. The temperature of the solid was obtained by averaging the temperatures of two diodes mounted on the outside of the guide wall. These tracked the vapor pressure curve of SD$_2$ well at higher temperatures. Later measurements made with diodes embedded in the solid indicate these measurements are accurate to 1 K.

Both the hydrogen contamination and the para-fraction in the SD$_2$ were measured by means of rotational Raman spectroscopy on a gaseous sample taken by warming the deuterium after the measurement [22]. These measurements yielded values for the HD concentrations in the gas that varied from 0.2%–0.3% with an uncertainty of about 0.1% (all uncertainties quoted at the 67% confidence level). Other contamination was removed by passing the D$_2$ through a palladium membrane. The para-fraction was controlled by converting the D$_2$ to a near thermal equilibrium ortho/para ratio in an iron-hydroxide-filled cell [26] cooled to a temperature at or slightly below the triple point. In this way, the para-fraction was reduced from a room-temperature equilibrium value of 33% to 2%–4%. Intermediate values were obtained by mixing deuterium at room-temperature equilibrium with converted deuterium before freezing. The precision of the para-fraction measurements was about 1%.

The SD$_2$ volume was measured by integrating the flow of gas while growing the solid. The volume was checked.
when the solid was warmed and the gas was returned to a buffer volume. The overall uncertainty in the volume measurement is approximately 15%.

The sensitivity of the apparatus to UCNs was demonstrated by measuring neutron arrival times with and without a $^{58}\text{Ni}$-coated 0.024-cm-thick aluminum foil in place at location C (in Fig. 1). The $^{58}\text{Ni}$ foil reflects all neutrons with velocities normal to the foil below 8 m/s. The UCN count rate should drop to near zero with the foil in place. These data are shown in Fig. 2. The number of counts, arriving in a time window between 0.5 and 10 s, with the foil closed was 8m

SD2

The UCN count rate should drop to near zero with the foil in place at location C (in Fig. 1). The short dashed curve is the result of a Monte Carlo calculation of UCN arrival times.

fig. 2 - background subtracted spectra with (long dashed line) and without (solid line) the $^{58}\text{Ni}$ barrier in the beam at the location of valve C. The short dashed curve is the result of a Monte Carlo calculation of UCN arrival times.

For a given $V_{SD}$, the surviving UCNs were recorded for a series of storage times, $t_s = 0.5, 1.0, 2.0, \text{ and } 4.0 \text{ s.}$ These data were fitted with the form $c_n e^{-t_s/\tau}$, where the parameters $c_n$ and $\tau$ were varied to produce the best fit to the data (the $\chi^2$ per degree of freedom for the background and the UCN signal fits were all close to 1 and consistent with Poisson statistics). We then systematically varied $V_{SD}$, the temperature of the SD2, and the ortho/para fraction of the SD2 to produce families of experimentally determined UCN bottle lifetimes, $\tau$.

The UCN lifetimes in the solid deuterium, $\tau_{SD}$ were extracted from the measured bottle lifetimes, $\tau$, using a Monte Carlo generated lookup table of $\tau_{SD}$ vs $\tau$. In the Monte Carlo transport, the full experimental geometry, gravitation, the SD2 potential (108 nV), and wall collisions in the guide tube were taken into account. There are a number of parameters that must be determined from the experimental data in order to extract $\tau_{SD}$: the probability of UCN loss and nonspecular reflection for each collision with the guide wall, the mean-free path for elastic scattering in the SD2, $\lambda_{el}$ (which may depend on the grain size of the sample), and the physical configuration of the SD2 frozen on the walls (SD2 can freeze as a flat “pancake” on the bottom of the guide, or coat the walls and bottom of the entire He-cooled surface at the end of the guide, in a “bucket”-shaped configuration). The shape of the SD2 was modeled by a single parameter $r$, which specified the ratio of SD2 volume in a uniform layer coating the walls and bottom and the volume in a pancake at the bottom.

Because samples of 0.4 cm or less of SD2 were used for most of the lifetime measurements, results were quite insensitive to $\lambda_{el}$. We therefore used the theoretical value of $\lambda_{el} = 8 \text{ cm}$ for the results presented here.

The ratio of diffuse to specular reflections for wall collisions was adjusted to fit time-of-arrival spectra (see Fig. 2). The remaining transport and physical parameters were adjusted to reproduce the dependence on $V_{SD}$ for all of the data below 10 K. The volume dependence and time-of-arrival spectra constrained the transport and physical parameters well enough so that the remaining errors in $\tau_{SD}$ are due to our inability to control systematic uncertainties in the ortho/para ratio and the volume. These have been accounted for by adding an uncertainty of 1 ms and a relative uncertainty of 20% in quadrature to all of the extracted deuterium lifetimes. These additions resulted in reduced $\chi^2$ of near one in the fitting described below.
The lifetimes due to absorption on deuterium and hydrogen were calculated using the known hydrogen contamination, tabulated thermal cross sections, and the 1/\nu dependence of the cross sections. The lifetimes in solid ortho- and para-deuterium due to temperature up-scattering were taken from the literature [20,22]. The lifetime in solid para-deuterium due to molecular transitions was treated as a free parameter and found to be 
\[ \tau_{\text{para}} = 1.2 \pm 0.14 \text{ (stat)} \pm 0.20 \text{ (sys)} \text{ ms}, \]
roughly consistent with a calculation that gives 
\[ \tau_{\text{para}} = 1.5 \text{ ms} \] [22].

Results for UCN lifetimes \( \tau_{\text{SD}} \) in SD2 as a function of the SD2 temperature and para/ortho fractions are shown in Fig. 3. The difference between the solid and the dashed line demonstrates the need to include the effect of deuterium vapor in the guide on the lifetime above 10 K. The measured lifetimes agree well with theoretical predictions of the up-scatter rate. The main contributions to the UCN lifetime in SD2 have been measured and are quantitatively understood. These data demonstrate the potential of a UCN source based on a spallation neutron driven SD2 converter.

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