Solid deuterium ($sD_2$) is of great importance for a whole class of new sources of ultracold neutrons (UCN). Theoretically [1-3] and experimentally [4-6] it was shown that $sD_2$ at sufficiently low temperature (around 5K), with high enough purity (less than 0.2 % ordinary hydrogen) and with high ortho concentration ($c_o > 0.98$) offers the possibility for ultra-cold neutron sources with about two orders of magnitude higher UCN densities as compared to the present best sources. Several $sD_2$ based sources are currently under construc-
tion worldwide. Their common principle is to expose sD$_2$ to a high flux of cold neutrons in order to produce UCN by down-scattering and to extract the UCN from sD$_2$ into vacuum and to guide them to storage volumes and experiments. Source performance obviously depends crucially on a high extraction efficiency of UCN from sD$_2$. Efficient extraction allows the equilibrium UCN density to build up faster and, if desired, to deliver a larger continuous UCN flux to experiments. One can expect D$_2$ crystal properties to have an influence on the extraction efficiency. Indications for considerable changes in scattering of very cold neutrons exist [7], depending on the procedures used for D$_2$ crystal preparation. Crystal properties which could be manipulated and potentially influence the extraction efficiency of UCN are of special interest and are the focus of the present studies.

2. The Experimental Setup

Solid D$_2$ samples can be frozen from the liquid in a cryogenic cell. The cell is mounted on a 4He flow cryostat and allows for neutron transmission and scattering experiments with simultaneous optical access. The sample thickness in the neutron beam direction is 10 mm, the optical path length perpendicular to it is 72 mm. The neutron beam windows are made out of AlMg$_3$ alloy machined to 150 $\mu$m thickness, the cold optical windows are 3 mm thick sapphire. The thermal shield (80 K to 100 K) surrounds the target cell but leaves the optical access open. Outside of the vacuum, the two optical windows are equipped for optical photography and Raman spectroscopy, respectively. The sample can be easily cooled to 5K and by pumping on the He to below 3 K. A dedicated D$_2$ gas system is used for purification and para-to-ortho conversion. Ortho-D$_2$ is produced at temperatures around the triple point using OXISORB$^{\text{b2}}$ [8,9] as a catalyst. The target cryostat and cell, the gas system and the optical systems are described in detail in [10]. Three neutron transmission and scattering setups have been used at the Paul Scherrer Institut, Villigen, Switzerland (cold neutrons, CN, SANS-I instrument [11]) and at the Institute Laue-Langevin, Grenoble, France (very cold, VCN, and UCN, PF2 instrument [12]), always using essentially the same strategy: preparation of the respective neutron beam using a velocity selector (PSI) or a chopper (ILL) in connection with adequate collimation; detection of transmitted and, in part, of scattered neutrons in 2D-detectors.

3. Raman Spectroscopy, Light and Neutron Transmission

Rotational Raman spectroscopy is done for two major reasons (see Fig. 1): a) it allows direct monitoring of the ortho-D$_2$ concentration of the sample by measuring the intensity ratio of the lines belonging to ortho- [S$_0$(0)] and para-D$_2$ [S$_d$(1)]; b) it yields information about the (hcp-) crystallite orientation in the sample by measuring the intensity distribution between the multiplet lines $\alpha$, $\beta$, and $\gamma$ which belong to the angular momentum substrates $m = \pm 1$, $\pm 2$, and 0, respectively, of the $J = 2$ final state of the S$_0$(0) transition [13]. Earlier, we used vibrational Raman spectroscopy for the investigation of gaseous D$_2$ samples at 300 K [14], however, with only $J = 0$ and $J = 1$ states populated at low temperature, the purely rotational transitions yield more information.

Fig. 1. Rotational Raman spectrum of solid ortho-D$_2$ (98.6%).

Another important reason for optical monitoring of the samples is to make sure that the neutron beam volume is filled with a known amount of material. Besides, from this information the images of sample crystals can be analyzed with respect to their light transmission. The sample is illuminated by the Raman laser from one side and photographs are taken from the opposite side. Figure 2 shows the development of the image brightness as a function of time during which the sample crystal undergoes thermal cycling. The initially high
light transmission of a 5 K crystal reduces slightly during cycling the sample between 5 K and 10 K, but becomes very small after seven cycles between 5 K and 18 K. Figure 3 shows preliminary results for UCN transmission through a sD₂ sample under the same thermal treatment as before. The initial transmission is only slightly affected by thermal cycling between 5 K and 10 K while the effect of cycling up to 18 K is dramatic.

4. Conclusions and Outlook

Cross sections for CN, VCN, and UCN on sD₂ have been measured. As an example the influence of thermal cycling on the (elastic) scattering of UCN was shown. The cross sections are only slightly affected by thermal cycling between 5 K and 10 K: A pulsed UCN source can thus be operated without deterioration of the sD₂ converter, as long as the temperature stays below 10 K, as planned for the PSI UCN source. The analysis of all the data from the experiments is under way. The experimental investigations will be extended to freezing from the gas phase, to UCN production cross section measurements on D₂ and to a comparison with O₂ and CD₄ as converter materials.

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