Threefold reduction in thermal conductivity of Vycor glass due to adsorption of liquid $^4$He

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Abstract. We report thermal conductivity measurements of porous Vycor glass when it is empty and when the pores are filled with helium between 0.06 and 0.5 K. The filling of liquid $^3$He and liquid $^4$He inside the Vycor pores brings about two to three-fold reduction of the thermal conductivity as compared with empty Vycor. This dramatic reduction of thermal conductivity, not seen with solid $^3$He and $^4$He in the pores, is the consequence of hydrodynamic sound modes in liquid helium that greatly facilitate the quantum tunneling of the two-level systems in Vycor and enhance the scattering of the thermal phonons in the silica network.

The atoms or molecules in a disordered glassy material have multiple potential energy minima in their spatial configurations. At sufficiently low temperature, typically below 1 K, the thermodynamic properties of the material are dominated by quantum tunneling between two adjacent accessible energy levels. The predictions of this two-level system (TLS) model [1–3], including a specific heat $C$ scaling with temperature $T$ and a thermal conductivity $\kappa$ scaling with $T^2$, have been confirmed by experiments [4–6].

Zeller and Pohl [4] found that the thermal conductivity of vitreous silica below 1 K is much lower than its crystalline counterpart, quartz and scales with $T^2$ instead of $T^3$.

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Heat is conducted in solid by thermally activated phonons and the thermal conductivity is given by $\kappa = (1/3)c_{\text{phonon}}\bar{u}l$, where $c_{\text{phonon}}$ is the contribution to the specific heat per volume by the phonons, $\bar{u}$ the average sound velocity and $l$ the phonon mean free path. Normally $c_{\text{phonon}} \sim T^3$ and $\bar{u}$ is weakly dependent on $T$. At low temperatures, phonons propagating in crystalline solid like quartz are primarily scattered at the defects and boundaries of the sample. Therefore $l$ is temperature independent and $\kappa \sim T^3$. For amorphous solid like vitreous silica, phonons are dominantly scattered by TLS that involve the coordinated motion and rotation of SiO$_4$ tetrahedra groups. The physical dimension of each group, based on simulation and neutron scattering studies, is more than 1 nm [7–10]. TLS scatter phonons by absorbing resonant phonons to tunnel to the excited state and re-emitting phonons to tunnel back to the ground state with a time constant $\tau$. The re-emitted or scattered phonons are generally propagating in a different direction from the incident phonons. TLS-phonon scattering results in an $l$ that scales with $T^{-1}$ [3]. As a result $\kappa \sim T^2$. If there are mechanisms to enhance the TLS tunneling rate $\tau^{-1}$, more phonons will be scattered per unit time and $\kappa$ will be reduced further. We show in this paper that the infusion of liquid helium into Vycor pores dramatically reduces $\kappa$.

In addition to the TLS-phonon scattering, phonons are also scattered by the porous structure when propagating in a porous glass like Vycor. Nitrogen adsorption isotherm measurements at $T = 77$ K show that Vycor has a porosity of 28% and a specific surface area of about 100 m$^2$ g$^{-1}$ [11, 12]. Transmission electron microscopy (TEM) revealed the pore space to be a network of multiply connected cylindrical channels [12]. The diameter of the channel and the thickness of the silica ‘walls’, based on TEM and adsorption studies, are both found to be $\sim 7$ nm [11, 12]. The spatial correlation length of the silica structure (also the pore structure) in Vycor, $\xi$, is found to be distributed between 5 and 60 nm by small angle x-ray and neutron scattering and TEM studies [12–14]. Therefore, phonons with wavelengths near or shorter than 60 nm are expected to be strongly scattered by the Vycor porous structure because of the acoustic mismatch at the silica–pore interface. Phonons with wavelength much longer than 60 nm, on the other hand, will be insensitive to the porous structure and scattered solely by TLS. The spectrum of thermal phonons obeys the Planck’s distribution which shows a peak at $\lambda_0 = \hbar u/4.2k_B T$, the characteristic wavelength [4] and the half maxima of the distribution at $0.5\lambda_0$ and $3\lambda_0$. Here $\hbar$ is the Planck’s constant, $u$ the sound velocity and $k_B$ the Boltzmann constant. Since the temperature range of this experiment is between 0.06 and 0.5 K, the characteristic frequency of thermal phonons, $f_0 = u/\lambda_0 = 4.2k_B T/\hbar$ is between 5 and 40 GHz. Taking the transverse sound velocity in Vycor, $u_{\text{Vycor}}$ to be 2200 m s$^{-1}$ [15], $\lambda_0$ is found to be 420, 170, 50 nm at $T = 0.06, 0.15, 0.5$ K, respectively. At low temperature when $\lambda_0$ is much longer than $\xi$, phonons are mainly scattered by TLS. As a consequence, $\kappa$ should scale with $T^2$. With increasing temperature, $\lambda_0$ decreases toward $\xi$ and the scattering effect by the porous structure becomes prominent. This will reduce $\kappa$ and result in sub-quadratic temperature dependence. This behavior has been experimentally confirmed [16–18].

In this paper, we present the thermal conductivity of Vycor infused by solid helium, liquid helium and helium films. Remarkably and counter-intuitively, a significant reduction in $\kappa$, up to a factor of 3, is observed when Vycor is infused with superfluid $^4$He films, liquid $^4$He and liquid $^3$He.

Figure 1 shows our experimental setup. The Vycor rod is 3 mm in diameter and 22.8 mm long. It is secured mechanically and thermally into a copper base with epoxy resin (Stycast 2850FT) and attached to the mixing chamber of a dilution refrigerator. Outside the copper base, the Vycor rod is sealed only by a thin layer of epoxy resin less than 0.07 mm thick.
Fig. 1. Experimental setup of thermal conductivity measurements.

Painted on its outer surface. The thin epoxy layer, impregnating the pores on the surface, is able to hold pressure up to at least 85 bar below 4 K but does not measurably contribute to the thermal conductance of the sample. Such an experimental configuration was recently used in studies of supersolidity in solid helium [19, 20]. Helium is introduced into Vycor through the copper base with a thin Cu–Ni capillary. A heater is attached at the top of the Vycor rod. A germanium thermometer is secured 12.7 mm away from the copper base reading $T_1$ and another thermometer is attached directly onto the copper base reading $T_2$. The copper base is at a uniform temperature because the thermal conductivity of copper is $10^5$ times higher than Vycor [21]. Thermal conductivity of the Vycor rod is given by $\kappa = L \dot{Q} / A \Delta T$, where $L$ is the distance between the germanium thermometer and the copper base, $A$ is cross section area of Vycor, $\dot{Q}$ is the steady state heat power and $\Delta T = T_1 - T_2$. Measurements are made by imposing a dc power between 0.1 and 1.6 nW to maintain a small but experimentally significant $\Delta T$ (typically 2 mK). The measured $\kappa$ of empty Vycor from 0.06 to 0.5 K is shown in both figures 2 and 3. As noted above, the deviation below the $T^2$ dependence for $T > 0.15$ K is clearly evident. Our results are consistent with previous experiments within a few per cent [16–18].

After the measurements of empty Vycor, solid, liquid and adsorbed films of both helium isotopes were introduced into the Vycor pores. For ease of description, these samples will be identified as solid, liquid and film samples. For clarity, we plot $\kappa$ of the solid and atomically thin film samples in figure 2 and $\kappa$ of liquid and superfluid film samples in figure 3. When Vycor is infused with liquid or solid helium, the samples can be considered as composites consisting of the two intertwining (silica and helium) networks. However, the heat is still conducted primarily by the silica network. This is the case because the sound velocities (and hence phonon wavelengths) in liquid and solid helium are $\sim 10$ times smaller than that in silica. Therefore the phonon wavelengths in helium are always shorter than 60 nm for $T > 0.06$ K. As a result, phonon propagation along the helium network, if exists, is scattered by the porous structure much more strongly than that along the silica network.
Figure 2. Thermal conductivity of empty Vycor and Vycor with solid $^3$He and $^4$He at 50 bar. $\kappa$ of solid $^3$He sample at 47 bar, not shown in the figure, was found to lie between the two data sets shown in the figure over the entire temperature range. Inset shows thermal conductivity of Vycor with inert helium films.

Figure 3. Thermal conductivity of empty Vycor and Vycor with liquid $^4$He, liquid $^3$He and superfluid $^4$He films with superfluid transition temperatures well above 1 K.

As shown in the inset of figure 2, $\kappa$ of the atomically thin $^4$He and $^3$He film samples are slightly (~3%) higher than that of the empty Vycor over the entire temperature range. The helium atoms in these thin films are immobile and tightly bound to the random silica surface with a typical adsorption potential higher than 25 K for the $^4$He film (surface coverage...
The sound modes conform to and also extend to the entire multiply connected pore space. Frequency restriction for the hydrodynamic sound modes because helium atoms are mobile. If the pore dimension exceeds the wavelength, there is no such wavelength and hence no such excitation in solid helium. Van der Waals interaction at the silica–helium interface. But in contrast to phonons in solid helium, the hydrodynamic sounds are also excited by TLS via the modulation of the van der Waals interaction at the silica–helium interface. With increasing temperature, the TLS–helium coupling facilitates the TLS tunneling and enhances the tunneling rate. As a result TLS-phonon scattering in silica becomes stronger and reduced. However phonon excitation in solid helium is strongly suppressed in the low temperature limit when the phonon wavelength $\lambda_{\text{He}}$ exceeds the pore dimension. This is why an enhancement of $\kappa$ is seen in the solid samples for $T < 0.2$ K due to the softening of acoustic mismatch across the silica–solid helium interface. With increasing temperature, $\lambda_{\text{He}}$ decreases to become comparable and then shorter than the pore dimension, thus opening the TLS–helium coupling channel and resulting in a reduction of $\kappa$. The difference in the crossover temperatures of the solid $^3\text{He}$ (0.2 K) and solid $^4\text{He}$ (0.36 K) samples reflects the different transverse sound velocities $u_t$ and hence $\lambda_{\text{He}}$ of the two solid helium samples. $u_t$ of 50 bar solid $^3\text{He}$ is 180 m s$^{-1}$ and $u_t$ of $^4\text{He}$ is 270 m s$^{-1}$ which translates to $\lambda_{\text{He}}$ of $\sim$10 nm near 0.36 K. The modest difference in $\kappa$ between the solid samples and the empty Vycor over the entire temperature range of the experiment confirms the reasoning stated above that heat conduction is primarily along the silica network and perturbed by the presence of helium.

Although the density and (first) sound velocity of liquid helium are similar to those of solid helium, the measured $\kappa$ of the liquid samples are dramatically different. Instead of a modest change, $\kappa$ of full pore liquid $^4\text{He}$ sample is reduced by a factor between 2 and 3 as compared with empty Vycor over the entire temperature range as shown in figure 3. A smaller but still sizable (a factor of 1.5) reduction of $\kappa$ is also seen in unsaturated superfluid $^4\text{He}$ film samples. $n_4 = 51.7$, 65.3 $\mu$mol m$^{-2}$). Interestingly, an approximately twofold reduction of $\kappa$ is found when the pores are filled with non-superfluid liquid $^3\text{He}$. This suggests the dramatic reduction of $\kappa$ is a consequence of fluidity and not superfluidity of liquid helium. The fluidity of the liquid helium enables hydrodynamic sound modes to be excited in the porous structure. Biot showed that such excitation requires the viscous penetration depth of the liquid $\delta = \sqrt{\eta/\pi\rho f}$ to be smaller than the pore radius $D/2$, so that a fraction of liquid can be decoupled and in relative motion with respect to the solid (in our case the silica) matrix. Here $\eta$ and $\rho$ are viscosity and density of the liquid and $f$ is the sound frequency. Similar to phonons in solid helium, the hydrodynamic sounds are also excited by TLS via the modulation of van der Waals interaction at the silica–helium interface. But in contrast to phonons in solid helium in the pores which must satisfy the boundary condition that their wavelengths should be comparable or shorter than the pore dimensions, there is no such wavelength restriction for the hydrodynamic sound modes because helium atoms are mobile. The sound modes conform to and also extend to the entire multiply connected pore space.
The frequency spectrum of this hydrodynamic sound is continuous and is given by the Planck’s distribution of thermal phonons in silica with the additional Biot condition: $f \geq f_c = 4\eta/\pi \rho D^2$. Therefore the hydrodynamic sound is much more efficient in enhancing $\tau^{-1}$ of TLS in the silica over the entire temperature range as compared with phonons in solid helium. This is the key reason for the dramatic reduction of $\kappa$ in the liquid samples. The hydrodynamic sound is often named as slow wave or slow sound [27–31] because its velocity is slower than the sound velocity in the solid matrix. Johnson pointed out that the slow sound in superfluid $^4$He confined in porous media is the well-known fourth sound [28]. Fourth sound can be excited at any finite frequency within the Planck’s distribution because the zero viscosity of superfluid gives $f_c = 0$. In the case of unsaturated superfluid films where viscosity is also zero, third sound plays the role of slow sound in enhancing $\tau^{-1}$. The fourth and third sounds themselves do not contribute to thermal conductivity because superfluid carries no entropy.

Slow sound can also be excited in liquid $^3$He with a finite viscosity as long as $f > f_c$. The density of liquid $^3$He at 3 bar (pressure of our sample) is 0.089 g cm$^{-3}$. The viscosity ranges from 150 to 50 μP between 0.15 and 0.5 K [32]. As a result $f_c$ ranges from 4.4 to 1.5 GHz, substantially lower than the characteristic frequency of the slow sound $f$, equivalent to $f_0$, ranging from 12 to 40 GHz in the same temperature range. Therefore, there is a significant fraction of liquid $^3$He in the pores of Vycor that supports the slow sound to enhance $\tau^{-1}$. The slow sound in liquid $^3$He is expected to vanish below 0.07 K when the viscosity of liquid $^3$He is large enough to completely lock the liquid to the silica matrix. Unfortunately, the large heat capacity of $^3$He and the low thermal conductivity of the Vycor prevent us from cooling the composite below 0.15 K.

The TLS tunneling rate in an amorphous system has been shown to be [3, 23]

$$\tau^{-1} = \frac{M^2 E^3}{2\pi \hbar^4 \rho u^5} \coth \frac{E}{2k_B T}. \quad (1)$$

Here $E$ is TLS energy separation, $\rho$ and $u$ are the density and sound velocity of the medium where TLS release energy and $M$ is the deformation potential that characterizes the coupling strength between TLS tunneling and the resultant strain in the medium. For empty Vycor, the deformation occurs at Si–O bonds and $M_{Si-O}$ is on the order of 1 eV [3]. In this case the TLS tunneling rate is labeled as $\tau^{-1}_{Si-O}$. With liquid helium in the pores, the TLS on the pore surface also couple with helium via modulating the silica–helium van der Waals interaction and $M_{SiO2-He}$ is $\sim$2 meV [23]. In this case the total TLS tunneling rate is given by $\tau^{-1} = \tau^{-1}_{Si-O} + \tau^{-1}_{SiO2-He}$. Since the density and sound velocity of liquid helium are $\sim$10 times smaller than those of silica, (1) shows that $\tau^{-1}_{SiO2-He}$ is larger than $\tau^{-1}_{Si-O}$ by a factor of 4. As a result, $\tau^{-1}$ is enhanced by a factor of 5. Since the physical dimension of the SiO$_4$ tetrahedra groups that make up the TLS is more than 1 nm and the thickness of silica ‘walls’ is $\sim$7 nm, at least 30% of the TLS reside on the pore surface. The average $\tau^{-1}$ of the entire sample is therefore enhanced at least by 2.2 times, close to the reduction in $\kappa$ found in full pore liquid $^4$He sample. This agreement supports the model we are proposing for the observed reduction of $\kappa$. Superfluid films cause less reduction of $\kappa$ than full pore liquid $^4$He because less helium are coupled to TLS. In the liquid $^3$He sample, the slow sound propagates with finite damping because of the finite viscosity. The damping may dissipate energy to the silica network in the form of phonons. This may explain why liquid $^3$He causes less reduction in $\kappa$ than liquid $^4$He.

Beamish et al [24] and Mulders et al [15] also reported an enhancement of TLS tunneling rate in Vycor due to the adsorption of liquid and solid helium in ultrasound experiments.

New Journal of Physics 15 (2013) 063030 (http://www.njp.org/)
between 0.08 and 5 K. However, these experiments were not probing thermal phonons since the ultrasound frequencies between 5 and 200 MHz correspond to characteristic temperature between 0.3 and 10 mK. Schubert et al [33] observed that the transmission of 25 GHz phonons from quartz to amorphous paraffin film is enhanced by the adsorption of liquid helium film on the paraffin at 1.8 K. They attributed this to the enhancement of TLS tunneling in the amorphous paraffin film.

In summary, we observe a dramatic reduction of thermal conductivity of Vycor when the pores are filled with liquid helium and superfluid films. The reduction is a consequence of hydrodynamic slow sounds in liquid $^4$He, superfluid $^4$He films and liquid $^3$He that facilitate the TLS tunneling in silica and enhance the TLS-phonon scattering. It is reasonable to suggest that the reduction in $\kappa$ found here should be general to all amorphous solids infused with liquid helium in the low temperature range where thermal transport is dominated by TLS-phonon scattering.

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New Journal of Physics 15 (2013) 063030 (http://www.njp.org/)
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