Metastability and Superfluid Fraction of the A-like and B Phases of ³He in Aerogel in Zero Magnetic Field[¶]

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Received March 18, 2004

We report the low-frequency sound measurements of the metastable A-like (A*) phase of superfluid ³He confined within a 98% open aerogel matrix in zero magnetic field. The second soundlike (slow) mode provides an accurate determination of the superfluid fraction of (and the transition between) the A* and B phases. The A* and B phases exhibit stable coexistence in the presence of disorder, the ratio of their superfluid fractions

 $(\rho_a^{A^*}/\rho_s^B)$ is much smaller than that of the bulk A and B phases, and argues that the A* and bulk A phases are distinct. © 2004 MAIK "Nauka/Interperiodica".

PACS numbers: 67.57.Bc; 64.60.My

³He is an ultrapure system that undergoes a transition from a normal Fermi liquid to the *p*-wave paired superfluid state. Strong confinement of ³He within regular geometries of characteristic size $\sim \xi_0$ (the superfluid ³He coherence length) had been predicted to significantly alter the bulk phase diagram [1], but this was never manifested in any experiments. The introduction of silica aerogel, a network of a few nm diameter SiO₂ strands, with fractal correlations of a few to 100 nm [2] provides a route by which superfluid ³He can experience correlated disorder because ξ_0 is larger than the strand diameter and of the same order of magnitude as the correlations.

The introduction of disorder also significantly alters the resulting phase diagram [3-6] beyond suppressing the superfluid transition temperature, T_{ca} . Recent work finds that a metastable A-like phase appears in both zero and finite magnetic fields, the polycritical point vanishes and the A \longrightarrow B transition, \overline{T}_{AB} and T_{ca} exhibit a width (partially attributed to temperature control and thermometry) [4, 7]. T_{AB} was explored in strong magnetic fields [8]; low-field NMR results definitively identified a hysteretic A \leftrightarrow B transition at high pressure [9] and identified the equilibrium low-temperature phase of ³He in aerogel as the B phase [10]. However, there is controversy as to the metastability of the A-like phase in zero field [7, 11]. The only other *p*-wave paired system, the heavy fermion UPt₃, has a rich phase diagram that is modified by both pressure and magnetic fields [12], though the influence of disorder (other than the suppression of T_c) by elastic scattering [13] has yet

[¶]This article was submitted by the authors in English.

to be mapped. Thus, there is an incomplete understanding of the effects of disorder on the phase diagrams of unconventionally paired systems.

Theoretical effort to understand the role of disorder has concentrated on the B phase [14, 15]. Only very recently, work by Fomin [16, 17], drawing on earlier general results of Volovik [18], *specifically excludes* the A phase as a stable phase in the presence of disorder and proposes other equal spin paired states as candidates for the A-like (A*) phase. However, properties that distinguish A* states from the A phase are not explicitly identified.

The A* superfluid phase that is initiated when cooling from the normal state exhibits a variety of timedependent behavior (in magnetic fields) quite different from that of bulk A to B phase conversion. A negative frequency shift (usually associated with the ³He A phase) and a positive frequency shift (associated with the B phase) were observed to be simultaneously present while cooling by Barker et al. [9] and the Moscow group [19]. For aerogel preplated with ⁴He, the weight of A* decreased continuously over a broad temperature range [19] before abruptly vanishing below a temperature, T_{A*B} [9, 19]. The results are different without ⁴He preplating. The A* phase was unstable with a lifetime that decreased depending on the proximity to T_{A*B} [19]. Pinning [11] has been invoked to explain the finite thermal and temporal width of the $A^* \longrightarrow B$ conversion process, in contrast to bulk ³He, where once initiated, the A \longrightarrow B transition proceeds rapidly to completion.

The superfluid fraction, ρ_s/ρ , measured with both torsional oscillators [5, 20] and sound [21] is signifi-



Fig. 1. Slow mode (SM) spectra at 21.01 bar offset vertically by temperature (the ordinate displays temperature and the received signal). The positive peak that merges into the weak "edge mode" below 100 Hz at $T_{ca} = 1.585$ mK is the fundamental SM resonance [21]. The broad feature above 170 Hz is a Helmholtz mode combined with SM harmonics that cross close to T_{ca} . The upper panel shows the evolution of the SM on cooling with the A* \longrightarrow B transition at $T_{A*B} = 1.415$ mK. The lower panel depicts the SM behavior on warming.

cantly suppressed from unity as $T \longrightarrow 0$. However, the A* phase superfluid density, ρ_s^{A*} has not been measured to date in zero field [5, 20, 21], because data were always obtained on warming after entering the B phase (due to temperature control difficulties while cooling).

In this letter, we report on zero-field low-frequency sound measurements as a function of temperature and pressure. Zero-field results are significant because of their lack of time dependence and also because the characteristic fields that alter the phase diagram and orient textures of superfluid ³He can be quite small. The magnetization of the surface ³He that may affect the time dependence is also field-dependent. In our experiments, the stability (*no time dependence*) of the A* and B phases and mixtures of these phases against conversion to the B phase allows us to reliably evaluate ρ_s . The sequence of phase transitions observed during warming and cooling show the following:

(1) The A* and B phases can coexist in a long-lived metastable state.

(2) The ratio of the A* and B phase superfluid fractions $\rho_s^{A^*}/\rho_s^B$ is very different from the bulk ρ_s^A/ρ_s^B . This ratio will be useful in establishing the nature of the A* phase.

(3) The transitions from A* to B (cooling) and B to normal (*N*) (warming) have widths of 20 and 25 μ K, respectively, and could be partially penetrated to produce coexistent A* and B phases after warming from T_{A^*B} or cooling from T_{ca} . The A* phase reproducibly supercools below T_{ca} before conversion to the B phase along a smooth curve T_{A^*B} on the *PT* diagram.

The experimental cell's cylindrical aerogel filled resonator has length l = 1.64 cm and diameter d =1.27 cm. Sound is coupled to and from the ³He via piezoceramic wafers attached to coin silver "speaker" and "microphone" membranes that are in contact with the 98% open aerogel sample's ends [22]. The spectrum of the acoustic resonator from which the sound velocities can be obtained is determined by sweeping the excitation frequency. A susceptibility thermometer monitored the ³He temperature, and a ³He melting curve thermometer monitored the PrNi₅ demagnetization stage.

For a superfluid in the interstices of a compliant solid such as aerogel, the interpenetrating normal and superfluid components [23] move in phase with the aerogel to give rise to a first soundlike "fast" mode whose frequency is continuous across the superfluid transition. The onset of the "slow" mode (SM) (in which ρ_s and ρ_n move out of phase) analogous to fourth sound in rigid porous media or second sound in the bulk is the signature of superfluidity. These modes were first observed by McKenna *et al.* [24] for ⁴He in aerogel, and the B phase modes were mapped (on warming) in ³He by Golov *et al.* [21]. The SM velocity is used to determine ρ_s/ρ .

Examples of the SM's evolution while cooling (upper panel) and warming (lower panel) at 21.01 bar are shown in Fig. 1. The A* \longrightarrow B transition (width ~20 μ K) is seen in the cooling trace and its onset, T_{A^*B} , and the superfluid transition T_{ca} are marked with arrows. For $T \ge T_{ca}$, the SM converts into the "edge mode" that involves counterflow of the normal fluid in the aerogel and the layer of bulk superfluid ³He around the aerogel [21]. Similar features were noted at all pressures between 28.6 and 13.16 bar and used to map out the phase diagram (Fig. 2).

In contrast to bulk ³He, where the A phase is the equilibrium high-pressure, high-temperature phase and vanishes below the polycritical point (~21 bar, see Fig. 2), our studies show no evidence for a B \longrightarrow A* transition (in agreement with [3, 4]). Unless the B phase

superheats (unlike the bulk), it is likely that there is no region where the A* phase is the lowest free-energy (equilibrium) phase in zero magnetic field. T_{A*B} was observed down to 13.16 bar and has marked similarities to the bulk ³He results of Schiffer *et al.* [25] obtained with ultraclean surfaces (Fig. 2). In traversing T_{A*B} , we varied the cooling rates between 9.5 and 100 µK/h with no effect on T_{A*B} or the transition's width (~20 µK), and T_{A*B} reproduces precisely upon cycling above T_{ca} .

NMR features have characteristics of a combination of A and B phases [9, 19], but their coexistence cannot be studied in zero field. Brussaard et al. [8] showed that the $A^* \longrightarrow B$ transition could be arrested and reversed by altering the magnetic field. We show evidence for the metastable coexistence of the A* and B phases of ³He in aerogel in zero magnetic field (Fig. 3). The upper and the lower traces represent the resonant frequency in the B and A^{*} phases, respectively. The intermediate trace was obtained by cooling into the T_{A^*B} band by only ~7 µK (i.e., not traversing the transition to completion). Following this partial transition, the sample was warmed to just below the T_{ca} band (defined below) and cooled while taking data. The frequency followed a path intermediate between that of the A* and B phases, implying a partial conversion from A* to the B phase. The coexistence of the A* and B phases in aerogel is evidence for strong pinning of the A*-B interface. The mixed state (without preplating with ⁴He) was stable at any temperature (including within 5 μ K of T_{A*B} for a period of a day), in contrast to the Moscow group's findings [19] of instability for pure ³He, or a broad transition for ⁴He covered surfaces (both in a magnetic field). The stability against conversion of coexistent A* to B phases in our experiment is similar to that for the ⁴He coated case [9, 19]. This suggests that the combination of a magnetic field and the surface solid ³He may play a role in mediating the conversion from $A^* \longrightarrow B$.

The superfluid transition, T_{ca} , occurs over a temperature band ~25 µK wide in accord with the heat capacity results of He et al. [26]. By carefully warming the sample from the B phase, we partially entered the T_{ca} band. After thermal equilibration and subsequent cooling at a rate of $\sim 40 \,\mu$ K/h, we observed a trace that was intermediate between the pure A* and pure B phase. Thus, if the cell was warmed to a temperature within the T_{ca} band, the result was a partial conversion from the $B \longrightarrow A^*$ phases. We never initiated a partial or complete $B \longrightarrow A^*$ transition on warming unless we entered the T_{ca} band which we could approach to within $2 \,\mu\text{K}$ (see inset to Fig. 3) [27]. The evolution of the SM was very similar to that depicted in Fig. 3, indicating that only part of the ³He B sample was converted back into A*. Different proportions of A* and B phase could be created depending on the depth of penetration into the T_{ca} or T_{A^*B} bands. The pressure-independent widths of T_{ca} and T_{A^*B} may be related to one another and due to inhomogeneities induced by compression of the ends of

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Fig. 2. The metastable ³He A* phase in 98% open aerogel in zero magnetic field is bounded by squares that mark the onset of superfluidity (T_{ca}) and open circles that mark the A* \longrightarrow B transition observed while cooling. The symbols' width is comparable to the transitions' width. Solid lines show the bulk ³He phase diagram in zero field, the dotted line shows the thermodynamically stable bulk B \longrightarrow A transition in a 24.9 mT field, and the A \longrightarrow B transitions observed in bulk ³He in a specially prepared cell (filled circles) [25]. A similar (less pronounced) T_{A*B} suppression was seen at Northwestern [3].



Fig. 3. The evolution of the peak frequency of the SM at 27.97 bar. The upper (warming) and lower (cooling) traces depict the B and A* phase results. The intermediate curve is a history-dependent trace obtained by initiating but not carrying to completion the A* \longrightarrow B transition. The inset shows a similar trace obtained by partial penetration into the T_{ca} band. The intermediate frequency SM is evidence for the stable coexistence of the A* and B phases of ³He in 98% aerogel.

the aerogel during cell assembly. However, if the width were solely due to inhomogeneities and not pinning, we would expect the T_{A^*B} width to diminish at high pressures, where the pressure dependence of T_{A^*B} is weak (Fig. 2).



Fig. 4. Values of ρ_s/ρ in 98% aerogel at 27.97 bar. The upper curve shows data obtained in the B phase on warming. The lower curve was recorded on cooling; both were carried out at ~40 μ K/h. The lower curve shows the signature of the A* \longrightarrow B transition. The cooling and warming traces overlap below the $T_{A^* \rightarrow B}$ "band."



Fig. 5. The $\rho_s/\rho_s^{\rm B}$ ratio for P = 27.97, 22.53, and 16.05 bar (from left to right). Below $T_{\rm A*B}$ (once the transition is complete), only the B phase is present; thus, $\rho_s/\rho_s^{\rm B} = 1$ ($\rho_s^{\rm B}$ is a function fitted to the B phase data obtained while warming).

Analysis of the velocity of the slow mode allows us to determine the superfluid fraction through the equation $\rho_s / \rho = \rho / \rho_a (c_s / c_a)^2$, where c_s is the slow mode velocity and c_a is the longitudinal sound velocity in aerogel. Using this equation, we calculate the ρ_s / ρ for the A* and B phases, after we subtract the frequency shift of the edge mode (described in Golov *et al.* [21]). The edge-mode velocity is $\sim (\rho_s^{\text{bulk}} / \rho)^{1/2}$ (ρ_s^{bulk} is the bulk ³He superfluid fraction). In Fig. 4, we plot the ρ_s/ρ found for the A* and B phases at 28 bar. Errors due to the extrapolation of the edge-mode frequency shift ~±1% are comparable to the scatter in the data in Fig. 4.

The B-phase superfluid fraction ρ_s^B / ρ in Fig. 4 is similar to that measured by the torsional oscillator technique [5], while the supercooled A* data fall below. The superfluid density of the A* and B phases (28 bar, Fig. 4) is well fitted by the equation $\rho_s/\rho = A(1 - T/T_{ca})^b$. The (A, b) coefficients at 28, 22.53, and 16.05 bar are (1.2, 2.1), (1.4, 2.2), and (1.5, 3.5) and (0.83, 1.5), (1.5, 1.9), and (0.97, 2.1) for the A* and B phases, respectively. The higher exponent at 16.05 bar manifests the relatively rapid growth of $\rho_s^{A^*}$ near T_{ca} seen in Fig. 5 but is confined to a limited temperature range (Fig. 2). The ratio $\rho_s^{A^*} / \rho_s^{B}$ diminishes near T_{ca} (see Fig. 5), and the ratio $\rho_s^{A^*} / \rho_s^{B} \le 0.5$ is smaller than that expected for bulk ³He A even when **l** is oriented parallel to the superflow [28], a situation that is impractical to achieve due to wall orientation effects [29]. It is surprising that this strong suppression of $\rho_s^{A^*}$ is manifested in a medium with no preferred orientation and in the absence of a magnetic field, unless the A* phase gap is much smaller than that of the B phase.

The strong reduction of $\rho_s^{A^*}$ (and A* phase gap) compared to ρ_s^{B} raises the following question: Why is the A* phase so reproducibly initiated from the *N* state even in the presence of B phase within the aerogel and in the surrounding bulk fluid [4]? It is possible that the *N* state may transform more readily into A* rather than the pseudoisotropic B phase, or the region near T_{ca} may possibly contain precursor nonsuperfluid states [18, 30] that transform more readily into the A* phase.

In summary, the slow mode of ³He in 98% aerogel in zero field was used to quantify the A* and B phase superfluid fractions, which differ by more than a factor of two, strongly implying that the A* and bulk A phases are distinct. We mapped the metastable A* \longrightarrow B transition in pressure and temperature. The A* phase persists with lifetimes greater than a day well below T_{ca} and can coexist on similar timescales with the B phase. We measure and quantify the width of the T_{A*B} and T_{ca} transitions. It is hoped that these measurements will spur theoretical efforts to understand the effects of disorder on the phase diagram.

J.M.P. acknowledges helpful conversations with I.A. Fomin, V.V. Dmitriev, T.L. Ho, and D. Einzel. Support was provided by the NSF (grant nos. DMR-0202113 and 9970817), by the CRDF, and by NATO under SA (grant no. (PST.CLG.979379)6993/FP).

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