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New Evidence for Zero-Temperature Relaxation in a Spin-Polarized Fermi Liquid

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Spin-echo experiments are reported for ${}^3\text{He}$ - ${}^4\text{He}$ solutions under extremely high B/T conditions, $B = 14.75$ T and $T \geq 1.73$ mK. The ${}^3\text{He}$ concentration x_3 was adjusted close to the value $x_c \approx 3.8\%$ at which the spin rotation parameter μM_0 vanishes. In this way the transverse and longitudinal spin diffusion coefficients D_\perp , D_\parallel were measured while keeping $|\mu M_0| < 1$. It is found that the temperature dependence of D_\perp deviates strongly from $1/T^2$, with anisotropy temperature $T_a = 4.26 \pm 0.18$ mK. This value is close to the theoretical prediction for dilute solutions, and suggests that spin current relaxation remains finite as the temperature tends to zero.

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A fundamental result of Fermi liquid theory is that the quasiparticle scattering time and hence the transport coefficients diverge as the temperature tends to zero. Recently, there has been much interest in the possibility that spin polarization could remove this divergence for transverse spin currents by creating scattering phase space between spin-up and spin-down Fermi surfaces [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11]. Thus the transverse spin diffusion coefficient D_\perp would remain finite at zero temperature in a *partially spin-polarized* Fermi liquid, while the other transport coefficients (longitudinal spin diffusion, viscosity, thermal conductivity) would diverge as in an unpolarized system. The *existence* of zero-temperature spin relaxation raises key questions about the applicability of conventional Fermi-liquid theory to transverse spin dynamics, even for weakly polarized systems provided the temperature is sufficiently low [4].

An initial round of theoretical studies introduced the idea of zero-temperature spin relaxation and computed its magnitude for very dilute systems [1, 2, 3, 4, 5, 6]. Spin-echo experiments in polarized liquid ${}^3\text{He}$ and ${}^3\text{He}$ - ${}^4\text{He}$ solutions found large deviations of D_\perp from T^{-2} temperature dependence, supporting the existence of zero-temperature relaxation [7, 8]. Experimentally, the apparent magnitude of the effect was stronger than expected from dilute-solution calculations [2, 3]. However, the theoretical basis for zero-temperature relaxation has been questioned by Fomin [9], and a recent experiment based on spin waves rather than spin echoes found that the effect is much weaker than previously measured, if indeed it exists at all [10]. The origin of the theoretical disagreement is unclear. In Ref. 3 a kinetic equation for dilute systems was solved to deduce the existence of zero-temperature attenuation, while in Ref. 4 field-theory methods were used to reach a similar conclusion. Conversely, in Ref. 9 it was argued that a proper separation of hydrodynamic variables shows that long-wavelength transverse spin currents are not relaxed at zero temperature.

In this paper we present new experimental evidence

that polarization-induced relaxation does indeed occur in ${}^3\text{He}$ - ${}^4\text{He}$, although observation of the effect requires higher fields and/or lower temperatures than previously thought necessary. Our results for a 3.8% ${}^3\text{He}$ - ${}^4\text{He}$ solution are consistent with the theoretical prediction for extremely dilute solutions, unlike the earlier results. Although we use NMR spin echoes, in common with earlier experiments that showed polarization-induced relaxation, we have picked ${}^3\text{He}$ concentrations x_3 very near the critical concentration $x_c \approx 3.8\%$ at which the spin rotation parameter $\mu M_0 = -\Omega\tau_\perp$ vanishes [12, 13]. This is significant because effects that destroy spin echo coherence such as restricted diffusion [14] and spin-wave instabilities [15] can limit the apparent magnitude of μM_0 , mimicking a departure from $D_\perp \propto 1/T^2$. The earlier experiments were all in the regime $|\mu M_0| \gg 1$ apart from one experiment that used a field/temperature ratio B/T ten times lower than that employed in the present work [8]. By adjusting x_3 to within 0.02% of x_c we achieved the condition $|\mu M_0| < 1$ at our highest $B/T = (14.75 \text{ T})/(1.73 \text{ mK})$. Thus, we have carried out an experiment showing significant spin-diffusion anisotropy ($D_\parallel/D_\perp > 5$) which is robust against possible effects of large $|\mu M_0|$.

The magnitude of polarization-induced relaxation is characterized by an “anisotropy temperature” T_a [4, 7], defined by fitting the transverse diffusion coefficient to

$$D_\perp(T) \propto 1/(T^2 + T_a^2). \quad (1)$$

This form implies that D_\perp tends to a constant value as the temperature is lowered well below T_a , hence the term “zero temperature” relaxation. For extremely dilute ${}^3\text{He}$ - ${}^4\text{He}$ solutions, it is predicted that $T_a = \mu_3 B / 2\pi k_B = (248 \mu\text{K/T})B$ where μ_3 is the ${}^3\text{He}$ nuclear magnetic moment [2, 3]. Therefore, very high B/T ratios exceeding 4000 T/K are required to measure T_a , unless nonequilibrium spin polarization is used as in Ref. 10.

To reach these conditions in equilibrium, we have employed a nuclear demagnetization cryostat that incorporates a 15 T NMR-grade sample magnet, which was oper-

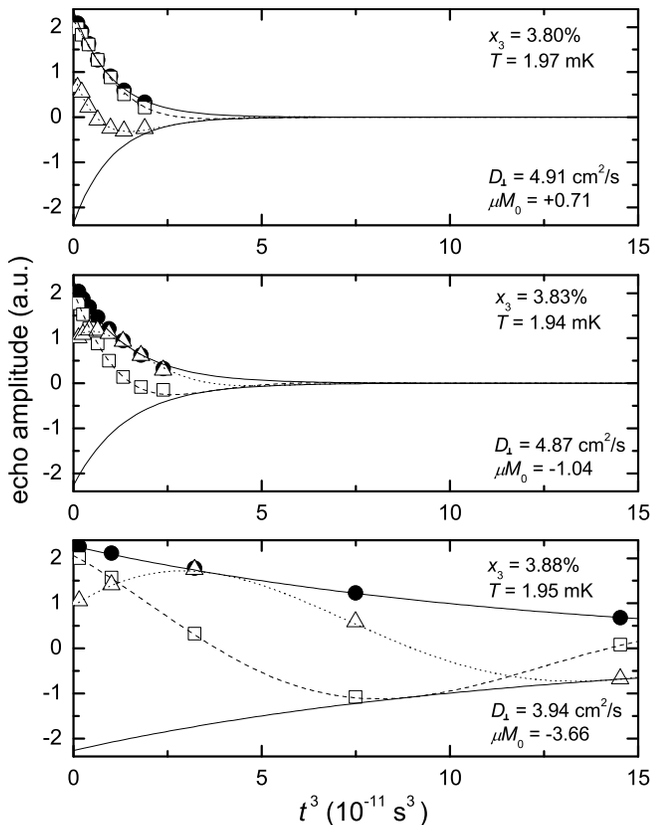


FIG. 1: Spin-echo amplitude as a function of time at temperatures near 2 mK for three slightly different values of the ^3He concentration x_3 . Circles, triangles, and squares show the magnitude, real, and imaginary parts respectively of the measured echo amplitude. The curves show fits to Eq. 2, which result in the values shown on each graph for the transverse spin diffusion coefficient D_{\perp} and the spin rotation parameter μM_0 . Even though μM_0 and thus the echo decay time varies rapidly with x_3 , for all of these data D_{\perp} is within 12% of the average value we measure at this temperature, $D_{\perp} = 4.46 \text{ cm}^2/\text{s}$ (Table I). This should be compared with our measured value of the *longitudinal* diffusion coefficient at the same temperature, $D_{\parallel} = 16.6 \text{ cm}^2/\text{s}$.

ated at $B = 14.75 \text{ T}$ for the experiments reported here. The sample cell has an epoxy NMR tube that extends into but does not touch a 478 MHz NMR resonator [16] thermally anchored to the mixing chamber. With this arrangement we are able to apply 50 W NMR pulses (180° pulse duration = $5.5 \mu\text{s}$) with negligible sample heating.

The NMR tube consists of a cylindrical sample cavity 2.4 mm dia. \times 2.5 mm high, connected to the main sample cell by a channel 0.7 mm dia. \times 6 mm high. The main sample cell contains a sintered heat exchanger (40 m^2 area) and three vibrating-wire viscometers. One of the viscometers includes an 0.82 mm diameter epoxy rod to reduce slip effects [17]. This viscometer retains temperature sensitivity down to our base temperature of 1.7 mK, and serves as the sample thermometer after calibration

at higher temperatures against a ^3He melting-pressure thermometer outside the high-field region. It is important to note that the viscometer directly measures the sample temperature in the main cell with no intervening thermal resistance. The sample inside the NMR cavity is in excellent thermal contact with the sample in the main cell (measured time constant $\approx 150 \text{ ms}$ at the base temperature), due to the T^{-1} temperature dependence of the sample thermal conductivity. The mechanical analysis for this composite viscometer and its calibration as a high B/T thermometer will be detailed elsewhere [18].

The ^3He concentration x_3 was determined to within $\pm 0.08\%$ (i.e. relative uncertainty of 2%) by measuring the quantities of gas added to the sample cell. To precisely adjust x_3 to x_c , small quantities of ^4He were added to the cell between temperature scans. Thus, the *differences* between x_3 values are known to within $\pm 0.005\%$.

The transverse spin diffusion coefficient D_{\perp} and spin-rotation parameter μM_0 were measured by observing the amplitude h and phase ϕ of the spin echoes formed by the two-pulse sequence $\theta-t/2-180^\circ-t/2$ -echo. Here $\theta = 8^\circ$ is the tipping angle of the first pulse and $t/2$ is the time between pulses. To measure D_{\perp} and μM_0 at each temperature, a series of spin echo experiments with different delay times $t/2$ were carried out. The echo amplitude was fit to the following form, valid for $\theta \ll 90^\circ$ [19]:

$$h e^{i\phi} = h_0 \exp[-\mathfrak{D}(\gamma G)^2 t^3/12], \quad \mathfrak{D} = \frac{D_{\perp}}{1 + i\mu M_0}. \quad (2)$$

The vertical static field gradient $G = 29.5 \pm 2.5 \text{ G/cm}$ applied to the sample was accurately measured by a least-squares fit to the shape of a single spin echo. The *longitudinal* spin diffusion coefficient D_{\parallel} was measured by the recovery of the longitudinal magnetization following its modification by a pulse of tip-angle $\theta' = 30^\circ$ and subsequent decay of the transverse components (pulse sequence $30^\circ-t-8^\circ$) [20].

Figure 1 shows spin echo data for three closely spaced values of x_3 , along with fits to Eq. 2. Despite the wide variation of μM_0 and the echo decay time, the fitted D_{\perp} values agree to within $\pm 12\%$. This is further evidence that effects that might limit the apparent magnitude of μM_0 have not significantly affected the measured value of D_{\perp} . Figures 2 and 3 show the data for D_{\perp} and D_{\parallel} respectively for several concentrations x_3 near the critical concentration x_c . Table I shows the results of fitting these data to

$$D_{\perp,\parallel}(T) = C_{\perp,\parallel}/(T^2 + T_{\perp,\parallel}^2). \quad (3)$$

Ideally we would find $C_{\perp} = C_{\parallel}$, $T_{\parallel} = 0$ and T_{\perp} would be the experimental estimate of T_a . In fact, the difference between the fitted values for C_{\perp} and C_{\parallel} is within their combined uncertainties, and the fitted value for T_{\parallel} is consistent with zero.

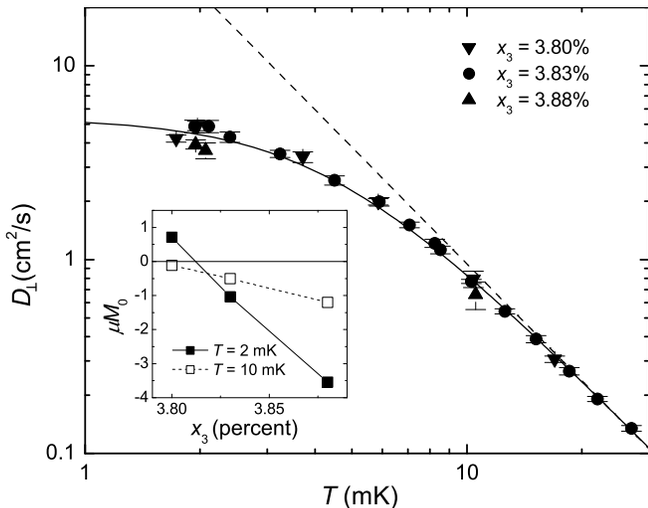


FIG. 2: Transverse spin diffusion coefficient D_{\perp} measured for three values of the ^3He concentration x_3 . The solid line shows a fit of the data to Eq. 3, which gives $T_a = 4.26 \pm 0.18$ mK. The dashed line shows a $1/T^2$ temperature dependence, corresponding to $T_a = 0$. Inset: Fitted spin-rotation parameter μM_0 as a function of x_3 for two temperatures.

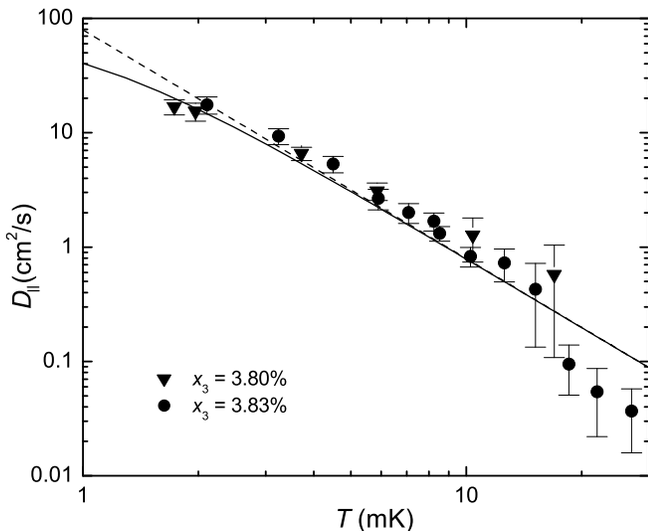


FIG. 3: Longitudinal spin diffusion coefficient D_{\parallel} measured for two of the same x_3 values used to measure D_{\perp} . The solid and dashed lines show fits as in Fig. 2. For D_{\parallel} , the deviation from $1/T^2$ (difference between solid and dashed lines) is not statistically significant.

An important consideration for both D_{\perp} and D_{\parallel} measurements is sample heating due to irreversible spin diffusion [21]. We have calculated the temperature rise of a polarized free Fermi gas following a tipping pulse and diffusive decay of the transverse magnetization. As in Ref. 21, we find that the initial and final temperatures $T_{i,f}$ are related approximately by $T_f^2 = T_i^2 + T_b^2$, and we have calculated T_b numerically as a function

TABLE I: Results of least-squares fits of the NMR data to Eq. 3. In addition to the fit uncertainties listed in this table, there is a 17% uncertainty in C_{\perp} due to the uncertainty in the gradient G , and a 17% uncertainty in C_{\parallel} due to uncertainties in the cell dimensions.

Parameter	Value	Units
C_{\perp}	$9.79 \pm 0.33 \times 10^{-5}$	$\text{cm}^2\text{K}^2/\text{s}$
T_{\perp}	4.26 ± 0.18	mK
C_{\parallel}	$7.9 \pm 1.9 \times 10^{-5}$	$\text{cm}^2\text{K}^2/\text{s}$
T_{\parallel}	0.98 ± 0.96	mK

of tipping angle and initial polarization and temperature. Here T_b is the upper bound for an apparent (false) anisotropy temperature that would be due to this type of heating, in the absence of true spin diffusion anisotropy ($T_a = 0$). For the conditions of our experiment, we calculate $T_b = 1.23$ mK for $\theta = 8^\circ$ as used for the D_{\perp} measurements, and $T_b = 3.3$ mK for $\theta' = 30^\circ$ as used for the D_{\parallel} measurements. Similar spin-diffusion heating occurs due to imperfections in the 180° pulse used to form spin echoes. However, we compute that the RMS deviation of the magnetization from a perfect 180° rotation is only 9° in our experiments, and any such heating would occur well after the main spin echo decay.

We have checked for several other possible conditions that might affect the D_{\perp} measurements: (1) The smaller of the spin mean free path and the spin rotation distance is never greater than 8% of the magnetization pitch for an echo decay of $1/e$, consistent with the requirements for the applicability of Leggett's spin dynamic equation [19]. (2) Similarly, we find that for the conditions of our experiment, relaxing the "steady state" approximation $\partial\mathbf{J}/\partial t = 0$ used in Ref. [19] never changes the apparent value of D_{\perp} by more than a few percent. (3) The fitted values of μM_0 are always much less than the apparent saturation value due to restricted diffusion found in Ref. [14], $(\mu M_0)_{\text{sat}} \approx 0.3b_L^{1/2}$. Here $b_L = L^3(\gamma G)|\mu M_0|/D_{\perp}$ where L is the cell height.

In Figure. 4 we show our data for the quantity $\mu M_0/D_{\perp}$. In Fermi-liquid theory, $\mu M_0/D_{\perp}$ is expected to be temperature independent, as both μM_0 and D_{\perp} are proportional to the transverse spin current relaxation time τ_{\perp} [19]. The vanishing of μM_0 at $x_3 = x_c$ can be viewed as a result of cancellation between positive and negative portions of the quasiparticle interaction potential. Therefore, it is perhaps not surprising that this cancellation is upset by thermal excitation of the system, leading to a variation with temperature of $\mu M_0/D_{\perp}$. The observed variation is approximately linear in temperature (Fig. 4). We have checked that the temperature variation is the same on warming and cooling.

Our most important result is that D_{\perp} follows Eq. 1 with a nonzero anisotropy temperature $T_a = 4.26 \pm 0.18$ mK, as shown by the solid line fit in Fig. 2. We be-

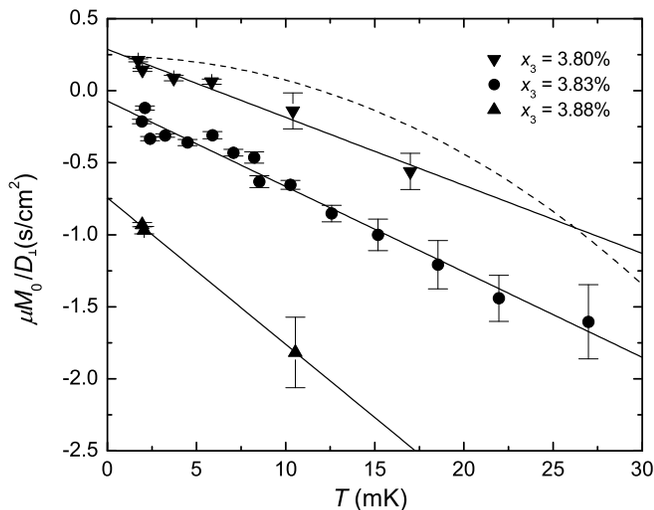


FIG. 4: Measured ratio of spin rotation parameter μM_0 to transverse spin diffusion coefficient D_\perp (data points), and linear fits (lines). In lowest-order Fermi liquid theory, this ratio should be independent of temperature, depending only upon the Fermi velocity and Fermi-liquid interaction parameters F_0^a and F_1^a . The dashed line shows the prediction of a calculation based on a phenomenological quasiparticle potential $V(q)$, evaluated for $x_3 = 3.65\%$.

lieve this result is robust against the effect of spin heating ($T_a > T_b = 1.23$ mK, and the latter temperature would only be reached after the echoes fully decayed [21]). It is possible that the spin dynamics are modified from Eq. 2 at the special point $x_3 = x_c$ [4, 22], although we see no experimental evidence of this. Interestingly, our measured T_a is nearly consistent with the theoretical value $\mu_3 B / 2\pi k_B = 3.73$ mK predicted for very dilute solutions, and is considerably smaller than the value 22 ± 3 mK that is obtained by extrapolating earlier results for this x_3 to our higher field [8]. Recently Buu, et al. reanalyzed NMR data taken at $x_3 = 6.1\%$ taking into account restricted diffusion effects, and concluded that T_a is considerably smaller than previously thought, although still 2.2 times larger than the dilute-solution value [11].

From a completely microscopic point of view, it appears impossible at present to predict T_a for ^3He concentrations outside the s-wave regime, which is roughly $x_3 < 10^{-3}$ [1, 4]. To provide some comparison with the present results, we have carried out numerical calculations of D_\perp and μM_0 using an effective quasiparticle scattering “potential” $V(q)$, along the lines of Ref. 3. Details of these calculations, which must be regarded as semi-phenomenological, will be reported elsewhere [23] and only the results are given here. Using the $V(q)$ proposed in Ref. [24] the calculated μM_0 for $T \rightarrow 0$ crosses zero at $x_3 = 3.65\%$, close to the experimental x_c (Fig. 4). At this x_3 (and in fact nearly independent of x_3), the calculated anisotropy temperature is $T_a = 3.7$ mK. Thus, the calculation predicts that the anisotropy temperature retains

its s-wave value for x_3 far outside the s-wave regime, in agreement with the present experiments. Unlike T_a the diffusion coefficients $D_{\parallel,\perp}$ and the spin rotation parameter μM_0 calculated from $V(q)$ are drastically modified from their s-wave values for this same range of concentrations; they agree well with our experimental values. As shown in Fig. 4, the calculations predict a temperature variation for $\mu M_0 / D_\perp$ over the range 2-30 mK that is similar in magnitude to the variation observed experimentally. However, the calculated variation is approximately quadratic in temperature, unlike the linear variation seen experimentally. This is the only significant discrepancy we find between the experimental results presented here and these calculations based on $V(q)$.

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