Stable, Strongly Attractive, Two-State Mixture of Lithium Fermions in an Optical Trap

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We use an all-optical trap to confine a strongly attractive two-state mixture of lithium fermions. By measuring the rate of evaporation from the trap, we determine the effective elastic scattering cross section $4\pi a^2$ to show that the magnitude of the scattering length $|a|$ is very large, in agreement with predictions. We show that the mixture is stable against inelastic decay provided that a small bias magnetic field is applied. For this system, the s-wave interaction is widely tunable at low magnetic field, and can be turned on and off rapidly via a Raman $\pi$ pulse. Hence, this mixture is well suited for fundamental studies of an interacting Fermi gas.

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Trapped, ultracold atomic vapors offer exciting new opportunities for fundamental studies of an interacting Fermi gas in which the temperature, density and interaction strength can be independently controlled. Recently, a degenerate gas of fermionic $^{40}$K has been produced by using a two-state mixture to enable s-wave scattering and evaporation in a magnetic trap [1]. By removing one species, the properties of the noninteracting degenerate gas were measured, demonstrating that the momentum distribution and the total energy obey Fermi-Dirac statistics [4]. However, the properties of interacting two-state fermionic vapors have not been explored experimentally.

Theoretical treatments of an interacting Fermi gas have focused extensively on $^6$Li [2–9]. Certain two-state $^6$Li mixtures are predicted to be strongly attractive, i.e., they have anomalously large and negative scattering lengths [1] arising from a near-zero energy resonance in the triplet state [11]. It has been predicted that these strongly attractive mixtures can undergo a transition to a superfluid state at a relatively high transition temperature [21]. In addition, the two-state effective interaction potential is widely tunable in a magnetic field, permitting systematic studies of fundamental phenomena such as collective oscillations for both the normal and superfluid phases [16,17], as well as new tests of superconductivity theory [18].

Unfortunately, magnetically trappable mixtures in $^6$Li with large s-wave scattering lengths are not stable, since there are correspondingly large spin-exchange and dipolar decay rates [11,21]. Hence, the methods employed to study degenerate $^{40}$K are not applicable. For this reason, we developed an ultrastable CO$_2$ laser trap to confine a stable mixture of the two lowest $^6$Li hyperfine states [12]. However, attaining a large and negative scattering length in this mixture requires high magnetic fields $B \geq 800$ G to exploit either a Feshbach resonance or the triplet scattering length $\pi a$ [11].

In this Letter, we show that there exists another stable hyperfine state mixture in $^6$Li which has the following unique properties. First, we predict that the scattering length $a$ is large, negative, and widely tunable at low magnetic field $B$. By monitoring the rate of evaporation from the CO$_2$ laser trap at a fixed well depth, we measure $|a| = 540^{+216}_{-199} a_0$ at $B = 8.3$ G. This result confirms for the first time that very large scattering lengths exist in $^6$Li mixtures. The predicted scattering length is $-490 a_0$ at $B = 8.3$ G, consistent with our observations, and is expected to increase to $-1615 a_0$ as $B \to 0$. Second, we find that this system is stable against spin exchange collisions provided that $B \neq 0$. In addition, the dipolar decay rate is predicted to be very small [13], consistent with our observations. Finally, in the experiments, a Raman $\pi$ pulse is employed to abruptly create an interacting mixture from a noninteracting one, a desirable feature for studies of many-body quantum dynamics [4].

![Figure 1](image)

**FIG. 1.** $^6$Li hyperfine states, labeled $|1\rangle$ to $|6\rangle$ in order of increasing energy in a magnetic field. The magnetic quantum number of each state is denoted by $m$. The hyperfine constant $a_{hf} = 152.1$ MHz.

Fig. 1 shows the hyperfine states for $^6$Li labelled $|1\rangle$ – $|6\rangle$, in order of increasing energy in a magnetic field. At low field, the states $|1\rangle$ and $|2\rangle$ correspond to the $|F = 1/2, m\rangle$ states, while states $|3\rangle$ through $|6\rangle$ correspond to states $|F = 3/2, m\rangle$. At nonzero magnetic field, only the magnetic quantum number $m$ is conserved. The subject of this paper is the $|3\rangle$ – $|1\rangle$ mixture.
out at low temperature, the dominant dipolar process is a small $s \to d$ rate in which $\{3,1\} \to \{1,2\}$.

In the experiments, the CO$_2$ laser trap is initially loaded from a magneto-optical trap (MOT) [13]. At the end of the loading period, the MOT laser beams are tuned near resonance and the intensity is lowered to decrease the temperature. Then, optical pumping is used to empty the $F = 3/2$ state to produce a 50-50 mixture of the $|1\rangle - |2\rangle$ states. These states are noninteracting at low magnetic field, i.e., the scattering amplitude vanishes as a result of an accidental cancellation [10]. With a CO$_2$ laser trap depth of 330 $\mu$K, up to $4 \times 10^7$ atoms are confined in the lowest-lying hyperfine states at an initial temperature between 100 and 200 $\mu$K. A bias magnetic field of 8.3 G is applied to split the two-particle energy states by $\approx 16$ MHz. This is twice the maximum attainable energy at our largest well depth of 400 $\mu$K = 8 MHz.

After a delay of 0.5 second relative to the loading phase, a pair of optical fields is pulsed on to induce a Raman $\pi$ pulse. This pulse transfers the population in state $|2\rangle$ to state $|3\rangle$ in two microseconds, initiating evaporative cooling in the resulting $|3\rangle - |1\rangle$ mixture. The optical fields are detuned from resonance with the D2 transition by $\approx 700$ MHz to suppress optical pumping. If the Raman pulse is not applied, the trapped atoms remain in the noninteracting $|1\rangle - |2\rangle$ mixture and exhibit a purely exponential decay with a time constant $\approx 300$ seconds.

An acousto-optic modulator (A/O) in front of the CO$_2$ laser controls the laser intensity, which is reduced to yield a shallow trap depth of 100 $\mu$K. By using a shallow well, we avoid the problem that the elastic cross section becomes independent of the scattering length at high energy, as described below. In addition, the shallow well greatly reduces the number of loaded atoms and makes the sample optically thin, simplifying calibration of the number of trapped atoms. To determine the trap parameters, the laser power is modulated and parametric resonances [15] are observed at drive frequencies of $2\nu$ for three different trap oscillation frequencies $\nu$: At 100 $\mu$K well depth, $\nu_x = 2.4$ kHz $\nu_y = 1.8$ kHz and $\nu_x = 100$ Hz, where the trap laser beam propagates along $z$. Using the measured total power as a constraint, we obtain the trap intensity $1/e^2$ radii, $w_x = 50 \mu$m and $w_y = 67 \mu$m, and the axial intensity $1/e^2$ length, $z_f \approx 1.13$ mm, where $z_f$ is consistent with the expected Rayleigh length within 15%.

The number of atoms in the trap $N(t)$ is estimated using a calibrated photomultiplier. The detection system monitors the fluorescence induced by pulsed, retroreflected, $\sigma_\pm$ probe and repumper beams which are strongly saturating ($I/I_{sat} = 26$ for the strongest transition). To simplify calibration, only the isotropic component of the fluorescence angular distribution is measured: The collecting lens is placed at the magic angle [13] of $55^\circ$ ($P_3(\cos \theta) = 0$) with respect to the propagation direction of the probe beams. The net efficiency of the detection system is determined using laser light of known power. The primary uncertainty in the calibration arises from
the excited state population fraction, which we estimate lies between 1/4 and 1/2.

\[\sigma(k) = \frac{8\pi a_3^2}{1 + k^2 a_3^2}, \quad (1)\]

where \(\hbar k\) is the relative momentum. For \(k|a_3| \ll 1\), the cross section is maximized. When \(k|a_3| \gg 1\), the cross section approaches the unitarity limit \(8\pi/k^2\) which is independent of \(a_3\). Note that \(k|a_3| = 1\) corresponds to a relative kinetic energy of \(\epsilon = \hbar^2/(2\mu a_3^2)\), where \(\mu = M/2\) is the reduced mass. For \(|a_3| = 500 a_0\), \(\epsilon = 115 \mu K\).

For a two-state mixture of fermions, the effective cross section is reduced from that of Eq. (1) by a factor of 2 since pairs of colliding atoms are in an antisymmetric hyperfine state with \(I = 1/2\). This effective cross section is used in a Boltzmann collision integral for each state \(i = 1, 3\). A decay term \(-N_i(t)/\tau\) with \(\tau = 370\) sec is added to account for the measured trap lifetime. A detailed description of our coupled Boltzmann equation model will be published elsewhere.

The coupled s-wave Boltzmann equations for the two states are numerically integrated to determine \(N(t)\) using the well parameters as fixed inputs. From the calibrated photomultiplier signal, assuming that 1/3 of the atoms are in the excited state, we obtain an initial total number \(N_0 = 44,000\). For this case, the initial collision rate in Hz is estimated to be \(1/(2\pi \tau_c) \approx N_0 M_0 \sigma_0/\hbar k_B T\), where \(\nu^3 = \nu_x \nu_y \nu_z\), \(\sigma_0 = 8\pi a_3^2\), and \(M\) is the \(^6\)Li mass. Assuming \(|a_{31}| = 500 a_0\), \(\tau_c = 30\) ms. Hence, for \(t > 0.3\) seconds, when on average 10 collisions have occurred, the sample should be thermalized as assumed in the theory.

The best fit to the data starting with 22,000 atoms in each state is shown as the solid curve in Fig. 3. The \(\chi^2\) per degree of freedom for this fit is 1.4 and is found to be very sensitive to the initial temperature \(T_0\) of the atoms in the optical trap. From the fit, we find \(T_0 = 46\mu K\), which is less than the well depth. We believe that this low temperature is a consequence of the MOT gradient magnet, which is turned off after the MOT laser beams. The effective well depth of the optical trap is therefore reduced until the gradient is fully off, allowing hotter atoms to escape before the Raman pulse is applied to create the \(|3\rangle - |1\rangle\) mixture. The fit is most sensitive to data for \(t > 0.5\) second, where the thermal approximation is expected to be valid. From the fit, we obtain the scattering length \(|a_{31}| = 540 \pm 25 a_0\), which is within 10\% of the predictions of Fig. 2. The quoted error corresponds to a change of 1 in the total \(\chi^2\).

We determine the systematic errors in \(a_{31}\) due to the uncertainties in the calibration and in the population imbalance as follows. The data is fit for an initial number of atoms \(N_0\) of 58,000 and 29,000, corresponding to an excited state fraction of 1/4 and 1/2. This yields \(|a_{31}| = 440 \pm 20 a_0\) and \(|a_{31}| = 750 \pm 42 a_0\), respectively. Note that for the larger scattering lengths, the cross section given by Eq. (1) approaches the unitarity limit and the error increases. We assume that the initial population imbalance for states \(|3\rangle\) and \(|1\rangle\) is comparable to that of states \(|2\rangle\) and \(|1\rangle\) in the optically pumped MOT. To estimate the latter population imbalance, we use state-selective Raman \(\pi\) pulses to excite \(|2\rangle \rightarrow |3\rangle\) or \(|1\rangle \rightarrow |6\rangle\) transitions in the MOT. Probe-induced fluorescence signals from states \(|3\rangle\) or \(|6\rangle\) show that the initial \(|1\rangle\) and \(|2\rangle\) populations are equal within 10\%. Note that residual population in state \(|2\rangle\) is expected to be stable and weakly interacting, since we estimate \(|a_{32}| < 30 a_0\) for \(0 \leq B \leq 50 G\) using the ABC method, and \(a_{12} \approx 0\). Using the parameters for the fit shown in Fig. 3 but changing the initial mixture from 50-50 to 60-40, we find a slight increase in the fitted scattering length from 540 \(a_0\) to 563 \(a_0\). Thus, the uncertainty in the calibration of the number of atoms produces the dominant uncertainty and \(|a_{31}| = 540^{+210}_{-100} a_0\).

To demonstrate that evaporative cooling is occurring, rather than just trap loss, we have also measured the...
final temperature of the mixture using release and recapture \[18\] from the CO$_2$ laser trap. We obtain 9.8 ± 1 µK, which is within 10% of the final temperature of 8.7 µK predicted by the Boltzmann equation model. An excellent fit to the data is obtained for the final temperature, which describes a thermal distribution. However, the initial temperature is not so readily measured, as it is nonthermal before evaporation is initiated, and is rapidly changing during evaporation, unlike the final temperature, which stagnates.

Good fits to the evaporation data are obtained neglecting inelastic collisions, suggesting that the dipolar rate for the |3⟩ − |1⟩ mixture is small, in contrast to the scattering length. A limit on the dipolar loss rate for the |3⟩ − |1⟩ mixture can be estimated from the τ = 370 second lifetime of the mixture after evaporation stagnates. For equal populations in both states, dipolar decay results in an initial loss rate \( \dot{n} = -Gn^2/4 \), where \( G \) is the dipolar rate constant and \( n \) is the total density. To obtain a high density, the trap is loaded at a well depth of 330 µK and the temperature of the atoms is reduced by evaporation to \( T \approx 30 ± 1 \mu K \). The number of atoms remaining in each state after evaporation is estimated to be \( N = 6.5 ± 2.2 \times 10^3 \), where the uncertainty is in the calibration. We cannot rule out the possibility that one state is depleted on a long time scale, since we do not directly measure the individual state populations. However, we believe that, after evaporation stagnates in the deep well, a |3⟩ − |1⟩ mixture remains, since subsequent reduction of the well depth yields final temperatures consistent with evaporative cooling. Note that the mixture ratio is not critical: An 80:20 mixture yields an initial loss rate \( \dot{n} = -0.16 G n^2/23 \) that of a 50/50 mixture. For a fixed 330 µK trap depth, \( \nu^3 = 2.6 ± 0.3 \text{kHz}^3 \), and the phase space density for one state in the harmonic approximation is then \( \rho_K = N/(K_B T/\nu)^3 = 7 \times 10^{-4} \). This corresponds to a maximum total density of \( n = 2 \rho_K/\lambda_B^3 = 6.4 \times 10^{13} / \text{cm}^3 \), where \( \lambda_B \equiv h/(2 \pi M K_B T) \). Since the exponential decay time of the |3⟩ − |1⟩ mixture is similar to that obtained in the noninteracting |1⟩ − |2⟩ mixture, we assume the loss is dominated by background gas collisions. Thus, we must have \( G n/4 << 1/\tau \), which yields \( G << 2 \times 10^{-14} \text{cm}^3/\text{sec} \). This result is consistent with the value \( G \approx 2 \times 10^{-15} \text{cm}^3/\text{sec} \) predicted for the dipolar rate constant at 30 µK by van Abeelen and Verhaar \[13\].

Future experiments will employ continuous evaporation by slowly reducing the well depth \[19\]. In this case, very large scattering lengths can be obtained at low temperatures and small well depths by using a reduced bias magnetic field \( B \). By adiabatically recompressing the well, experiments can be carried out with the precooled atoms in a deep trap to obtain high density as well. In such experiments, the final low temperature limits the number of atoms in the high energy tail of the energy distribution, exponentially suppressing spin-exchange collisions for \( B \neq 0 \). For example, if a total of \( 3 \times 10^5 \) atoms were contained in our trap at a well depth of 400 µK, the Fermi temperature \( T_F = 7 \mu K \) and the Fermi density is \( 4 \times 10^{13} / \text{cm}^3 \). At a temperature of \( T = 0.1 T_F = 0.7 \mu K \), a bias field of \( B = 0.16 \text{G} \) would split the two-particle hyperfine states by \( |k_B T_F + 12 k_B T| \), suppressing the spin exchange rate by \( \exp(-12) \), and giving \( a_{31} \approx -1200 a_0 \). Alternatively, as shown in Fig. 2, large \( a_{31} \) can be obtained at moderate \( B \approx 300 \text{G} \).

In conclusion, we have observed that an optically trapped |3⟩ − |1⟩ mixture of ²⁶Li atoms has a very large scattering length at low magnetic field. This mixture is stable against spin-exchange collisions provided that a small bias magnetic field is applied. The evaporation curves measured for this mixture are in good agreement with a model based on an s-wave Boltzmann equation which neglects inelastic processes. We have predicted that the scattering interactions are strongly attractive and widely tunable at low magnetic field. If the parameters described above for deep wells can be attained, the system will be close to the threshold for superfluidity \[3\] and ideal for investigating frequency shifts and damping in collective oscillations \[3,4\]. Further, since s-wave interactions can be turned on and off in a few microseconds, this system is well suited for studies of many-body quantum dynamics.

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\[1\] B. DeMarco, and D. S. Jin, Science \textbf{285}, 1703 (1999).
\[2\] H. T. C. Stoof, et al., Phys. Rev. Lett. \textbf{76}, 10 (1996); See also, M. Houbiers et al., Phys. Rev. A \textbf{56}, 4864 (1997).
\[3\] L. Vichi and S. Stringari, Phys. Rev. A \textbf{60}, 4734 (1999).
\[4\] R. Combescot, Phys. Rev Lett. \textbf{83}, 3766 (1999).
\[5\] G. M. Bruun, and C. W. Clark, Phys. Rev. Lett. \textbf{83}, 5415 (1999).
\[6\] G. M. Bruun and C. W. Clark, cond-mat/9906392.
\[7\] M. Houbiers and H. T. C. Stoof, Phys. Rev. A \textbf{59}, 1556 (1999).
\[8\] G. Bruun, et al., Eur. Phys. J. D \textbf{7}, 433 (1999).
\[9\] M. Houbiers and H. T. C. Stoof, cond-mat/9808171.
\[10\] M. Houbiers, et al., Phys. Rev. A \textbf{57}, R1497 (1998).
\[11\] E. R. I. Abraham, et al., Phys. Rev A \textbf{55}, R3299 (1997).
\[12\] K. M. O’Hara, et al., Phys. Rev. Lett. \textbf{82}, 4204 (1999).
\[13\] We are indebted to F. A. van Abeelen and B. J. Verhaar who calculated the inelastic |{3,1}| → |{1,2}| dipolar rate and confirmed our calculations of the magnetic field dependence of \( a_{31} \).
\[14\] P. Törmä and P. Zoller, Phys. Rev. Lett. \textbf{85}, 487 (2000).
\[15\] S. Friebel, et al., Phys. Rev. A \textbf{57}, R20 (1998).
\[16\] Atomic, Molecular, and Optical Physics Handbook, ed. G. W. Drake, (AIP Press, New York, 1996), p. 176.
\[17\] O. J. Luiten, et al., Phys. Rev. A \textbf{53}, 381 (1996).
\[18\] S. Chu, et al., Phys. Rev. Lett. \textbf{55}, 48 (1985).
\[19\] C. S. Adams, et al., Phys. Rev. Lett. \textbf{74}, 3577 (1995).