Atom–Molecule Coherence in a Bose-Einstein Condensate

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Coherent coupling between atoms and molecules in a Bose-Einstein condensate (BEC) has been observed. Oscillations between atomic and molecular states were excited by sudden changes in the magnetic field near a Feshbach resonance and persisted for many periods of the oscillation. The oscillation frequency was measured over a large range of magnetic fields and is in excellent quantitative agreement with the energy difference between the colliding atom threshold energy and the energy of the bound molecular state. This agreement indicates that we have created a quantum superposition of atoms and diatomic molecules, which are chemically different species.

There is considerable interest in extending the applications of ultracold atoms to ultracold molecules. One route for producing a very cold and possibly Bose-condensed sample of molecules is to create the molecules from an atomic BEC. Wynar et al. created cold $^{87}$Rb$_2$ molecules in a single ro-vibrational state of the ground-state potential from an $^{87}$Rb BEC using a two-photon stimulated Raman transition. The authors could not probe the coherence properties of the molecules in that state, but the prospect of creating a superposition of atomic and molecular condensates initiated a flood of theoretical work on the subject$^{2-6}$. Ultracold molecules have also recently been formed through photoassociation of a sodium BEC$^7$.

Utilizing the natural atom–molecule coupling that arises from a Feshbach resonance is an alternate route for producing ultracold molecules from an atomic BEC, and it is the route we have followed here. A Feshbach resonance is a scattering resonance for which the total energy of two colliding atoms is equal to the energy of a bound molecular state, and atom–molecule transitions can occur during a collision$^8-12$. A schematic representation of the potentials involved is shown in the inset of Fig. 1A. For our $^{85}$Rb resonance, BEC atoms in the $F = 2$, $m_F = -2$ state collide on the open-channel threshold. $F$ and $m_F$ are the total spin and spin-projection quantum numbers. The bound state in the closed channel differs in energy by an amount $\epsilon$ from the open-channel threshold. The bound molecular wave function can be described as a sum of amplitudes of different hyperfine components $(F, m_F)$ having $M_F = m_{F,1} + m_{F,2} = -4^{13}$. Because of their different spin configurations, the atoms and molecules generally have different magnetic moments and the difference depends on magnetic field. Thus $\epsilon$ depends on magnetic field and the degree of atom–molecule coupling is magnetically tunable. The energy difference between the free atoms and the bound molecules is plotted in Fig. 1A. This behavior of the bound-state energy also causes a resonance in the scattering length, $a$, which is shown in Fig. 1B. The scattering length characterizes the mean-field interaction energy of a BEC.

When the magnetic field is tuned to values near the Feshbach resonance, theory predicts coherent oscillations between the atomic and molecular states, but there is significant disagreement on the conversion fraction and the coherence properties$^{16-21}$.

In experiments with a sodium BEC, Stenger et al. observed that inelastic losses were dramatically enhanced when they ramped the magnetic field across the Feshbach resonance. We observed similar results for $^{85}$Rb, but with lower rates$^{23,24}$. It is likely that the formation of molecules played a role in the loss, but there was no experimental evidence for the presence of molecules and the results followed a loss-rate dependence on time. More recently, we measured the time dependence of the loss in an $^{85}$Rb BEC by applying controlled magnetic-field pulses toward but not across the Feshbach resonance$^{25}$. We observed the surprising result that under some conditions, shorter, more rapid pulses actually led to more loss than longer, slower pulses that spent more time near the resonance. The time dependence of the loss was suggestive of a nonadiabatic mixing of states, with the only states within a reasonable energy range being the normal atomic BEC state and the nearby bound molecular state.

In this work we show that much of the loss is likely due to the coherent mixing of atomic and molecular states. To create a superposition and probe its coherence, we applied two short magnetic-field pulses toward the Feshbach resonance, separated by a “free evolution” time during which the magnetic field was held at a constant value some distance from the resonance. We measured the number of atoms in the condensate as a function of time between the two pulses for various values of the steady-state magnetic field between the pulses. We observed dramatic oscillations in the number of atoms remaining in the atomic BEC at frequencies corresponding to the energy splitting between the molecular and the atomic states.

**Experimental methods.** The apparatus has been described in detail elsewhere$^{23,25}$. We first created $^{85}$Rb condensates typically containing 16,500 atoms, with fewer than 1,000 uncondensed thermal atoms. The initial number $N_{\text{init}}$ fluctuated from shot to shot by $\sim$500 atoms (3% number noise). After producing the condensate at a field of $\sim$162 G, we ramped the magnetic field adiabatically to $\sim$166 G, corresponding to an initial scattering
FIG. 1: Feshbach resonance bound-state energy and scattering length. (A) Energy splitting versus magnetic field. The resonance is centered at ∼155 G. The solid curve is a theoretical estimate of the energy found with a coupled-channel calculation, and the dotted line indicates $\epsilon = 0$. The inset schematically shows the collision channels involved in the resonance. $\epsilon$ depends on magnetic field because the atoms and molecules have different magnetic moments and thus the potentials have different Zeeman shifts. (B) Scattering length versus magnetic field for fields above the Feshbach resonance. The scattering length $a_{\text{init}} \simeq 10 a_0$, where $a_0 = 0.053$ nm. The spatial distribution of the atoms was Gaussian with a peak atom density of $n_0 = 5.4 \times 10^{13}$ cm$^{-3}$, and the trap frequencies were (17.4×17.4×6.8) Hz. After preparing the condensate we applied a selected fast magnetic-field pulse sequence by sending an appropriate time-dependent current through an auxiliary magnetic-field coil. A typical pulse sequence is shown in Fig. 2. It is composed of two nearly identical short trapezoidal pulses separated by a region of constant (but adjustable) magnetic field. Upon completion of the fast-pulse sequence in Fig. 2, we ramped the magnetic field from ∼166 G to ∼157 G in 5 ms and held at that field for an additional 7 ms to allow the repulsive mean-field energy to expand the condensate. Then we turned off the magnetic trap and used destructive absorption imaging 12.8 ms later to observe the atomic condensate and measure the number of remaining atoms. This detection scheme was neither sensitive to atoms with kinetic energies larger than ∼2 µK nor to atoms in off-resonant molecular states. We determined the value of the magnetic field between the pulses, $B_{\text{evolve}}$, by measuring the resonance frequency for transitions from the $F = 2, m_F = -2$ to the $F = 2, m_F = -1$ spin state by applying a 10 µs RF pulse to a trapped cloud of atoms.

FIG. 2: Magnetic field pulse shape. Fields shown for pulses #1 and #2 correspond to scattering lengths of ∼2500 $a_0$, and the free precession field $B_{\text{evolve}}$ corresponds to a scattering length of ∼570 $a_0$. The dashed line indicates the position of the Feshbach resonance. In the text, we refer to the free precession time as $t_{\text{evolve}}$. The rise/fall time for all of the pulses that we used was 14 µs.

Burst and remnant oscillations. As we observed for single pulses toward the Feshbach resonance, there were two distinct components of atoms observed in the absorption images and a third “missing” component that we could not detect. One of the observed components was a cold remnant BEC which was not noticeably heated or excited by the fast-pulse sequence, while the other component was a relatively hot (∼150 nK) “burst” of atoms that remained magnetically trapped during the BEC expansion time. Using a variational approach to model the mean-field expansion that we applied to the BEC remnant to measure its number, $N_{\text{remnant}}$, we found that we should impart $\leq 3$ nK worth of energy to the remnant before imaging. This estimate agrees well with the expansion velocity that we observed after the trap turn-off. Thus the remnant BEC was nearly 50× colder than the burst.

The missing component contained atoms that were in the initial sample but were not detected after the trap turn-off. To find the number of atoms in the remnant BEC and the number of burst atoms, we allowed the magnetic trap to “focus” the burst cloud before imaging. A typical image is shown in Fig. 3. We fit the focussed...
Plank's constant divided by 2π and a\text{B}

burst (which had a much larger spatial extent than the remnant) with a two-dimensional Gaussian surface, excluding the central region of the image that contained the remnant. This fit yielded the number of burst atoms, \(N_{\text{burst}}\). Subtracting this fit from the image and performing a pixel-by-pixel sum of the central region of the image gave \(N_{\text{remnant}}\).

\(N_{\text{remnant}}\) versus \(t_{\text{evolve}}\) is plotted in Fig. 4 for two different values of \(B_{\text{evolve}}\). The number clearly oscillates. Changing the value of \(B_{\text{evolve}}\) affected the oscillation frequency dramatically (note the change in scale from Fig. 4A to Fig. 4B). After only pulse #1 and the subsequent constant field but with no pulse #2, \(N_{\text{remnant}}\) showed no variation except for a slow decay consistent with the loss rate expected for a single pulse to that field\textsuperscript{29}.

We have taken data similar to those in Fig. 4 for a variety of different \(B_{\text{evolve}}\) values. As in Fig. 4, we fit each curve to the function \(y = y_0 + A \exp(-t/\tau) \sin(2\pi\nu t + \phi)\) to find the oscillation frequency \(\nu\) and decay time constant \(\tau\). The measured frequencies are plotted versus \(B_{\text{evolve}}\) in Fig. 5 along with theoretical predictions for the bound-state energy relative to the atomic state.

In the regime where the scattering length is much larger than the radius of the interatomic potential well, the bound state energy for an arbitrary attractive potential can be approximated by \(\epsilon = -\hbar^2/ma^2\textsuperscript{29}\). \(\hbar\) is Plank's constant divided by 2\(\pi\), \(m\) is the atomic mass, and \(a\) is the scattering length. The same equation relates the bound state energy to the effective scattering length, which is calculated from the Feshbach resonance parameters through the relation \(a = a_{bg} \times (1 - \frac{\Delta}{B - B_0})\), where \(a_{bg}\) is the background scattering length, \(\Delta\) is the width of the Feshbach resonance, and \(B_0\) is the resonant magnetic field\textsuperscript{30}.

The quantity |\(e|/\hbar\) is plotted with no adjustable parameters in Fig. 5. The measured oscillation frequencies are in excellent agreement with this simple model over the range of magnetic fields where the model is expected to be valid. The theoretical results found with a much more sophisticated coupled-channels scattering calculation\textsuperscript{14} in Fig. 5 are in excellent agreement with the data over the entire range.

The fact that the oscillations occurred at exactly the frequency corresponding to the bound-state energy clearly indicates that we are creating a coherent superposition of atoms and molecules with the sudden magnetic-field pulses\textsuperscript{11,12}. Although we do not have a detailed understanding of how the field pulses couple atoms and molecules, by choosing the shapes of the perturbing
pulses such that a single pulse results in roughly 50% loss, we observe high-contrast oscillations in the number of atoms in the atomic BEC.

From the amplitude of the oscillations, one can put a lower bound on the number of molecules being created. Take, for example, the data in Fig. 4A. The amplitude of the atom oscillations was 1800(300) atoms. Assuming the fringes are coming from interference with molecules, there must be at least 1800/2 = 900(200) molecules on average. Assuming that the missing atoms are molecules that fail to convert back into atoms gives an upper bound of 3200(100) molecules for the conditions of Fig. 4A.

The damping time for the oscillations, \( \tau_{\text{decay}} \), was more difficult to measure with high precision than the oscillation frequency. To within our measurement precision, \( \tau_{\text{decay}} \) did not depend on \( B_{\text{evolve}} \), but our uncertainties in \( \tau_{\text{decay}} \) were as large as 100% for some fields. We had the highest precision measurements for frequencies around 200 kHz where the oscillation period was long compared to our experimental timing jitter but short compared to \( \tau_{\text{decay}} \). At frequencies near 200 kHz, we measured \( \tau_{\text{decay}} = 38(8) \mu s \) for \( n_0 = 1.3 \times 10^{14} \text{ cm}^{-3} \) and \( \tau_{\text{decay}} = 91(33) \mu s \) when we decreased \( n_0 \) to 1.1 \times 10^{13} \text{ cm}^{-3}.

\( N_{\text{burst}} \) also had interesting dependencies. For the conditions under which most of the data were collected, the burst contained \(~5000\) atoms on average, which is \(~30\%\) of \( N_{\text{init}} \). \( N_{\text{burst}} \) depended on density and varied from one-half of the atoms lost from the condensate for our typical peak density of \( n_0 = 5.4 \times 10^{13} \text{ cm}^{-3} \) to nearly all of the atoms lost from the condensate for \( n_0 = 1.1 \times 10^{13} \text{ cm}^{-3} \). \( N_{\text{burst}}, N_{\text{remnant}}, \) and total number of atoms detected are plotted in Fig. 6 for \( B_{\text{evolve}} = 159.84(2) \text{ G} \) and \( n_0 = 1.1 \times 10^{13} \text{ cm}^{-3} \) (5\% lower density than was used for the data shown in Fig. 4). All three components oscillated at the same frequency. The burst oscillation lagged behind the remnant oscillation by 155(4)\(^\circ\). Since the relative phase shift is nearly 180\(^\circ\), the oscillation amplitude for the total number was smaller than either the burst or the remnant oscillation amplitudes. The relative phase depended sensitively on the fall time of pulse #2. For example, when we increased the fall time from 11 \( \mu s \) to 159 \( \mu s \), the burst oscillation then lagged behind the remnant oscillation by 68(7)\(^\circ\) and the peak-to-peak amplitude of the total number oscillation was 5600(400).

For the conditions of Fig. 6, \( N_{\text{init}} = 17,100 \) exceeded the time-averaged total number of atoms counted after the pulse sequence by (3\%) on average. For the higher-density measurements in Fig. 4, 39(4)\% of the atoms were missing. Experiments with longer pulses #1 and #2 also had a higher fraction of missing atoms. For example, when we used 50 \( \mu s \) pulses with \( n_0 = 5.4 \times 10^{13} \text{ cm}^{-3} \), 56(3)\% were missing.
We have carried out double-pulse measurements with a variety of widths and amplitudes for pulses #1 and #2 and a variety of different densities and initial magnetic fields. Although the oscillation frequency was unchanged, the phase, contrast, and damping of the oscillations did vary. The contrast was very sensitive to the pulse length, and was lower for longer pulses that created more missing atoms. Defining the contrast as the oscillation amplitude divided by the time averaged number of remnant atoms detected, we observed an optimum contrast of 0.42(7) for 15 µs pulses to 156.6(1) G. A single such pulse removed about half of the atoms from the BEC. When the pulse length was comparable to $\tau_{\text{decay}}$, the contrast was reduced by about a factor of two. Under those conditions, $\sim 3/4$ of the atoms were lost after pulse #1. The phase was shifted, but we did not observe a change in the contrast when we varied the amplitudes of pulses #1 and #2 from $B = 156.6(1)$ G (2,400 $a_0$) to $B = 155.1(1)$ G (24,000 $a_0$). The contrast did depend on the intermediate level, however, and was reduced for $B_{\text{evolve}}$ values closest to the resonance, for which the magnetic-field jumps between $B_{\text{evolve}}$ and pulses #1 and #2 were shortest.

We also looked for a temperature dependence of both the damping and the frequency at $\sim 200$ KHz and did not see any. The high-temperature data was much noisier than the data for pure condensates, due to unexplained enhanced noise in the number of thermal atoms after the magnetic-field pulse, but when the initial thermal fraction was increased from $<5\%$ to $30\%$, the data still fit to oscillations with frequency, amplitude, and damping consistent with what was observed with low temperature data.

Conclusions and Outlook. Our interpretation of our observations is that the first magnetic-field pulse provides a sufficiently rapid perturbation to result in nonadiabatic mixing between atomic and molecular states. The superposition then evolves according to the energy difference between the states, which is determined by the magnetic field during the free evolution stage, $B_{\text{evolve}}$. The second pulse mixes atom and molecule states again, such that the final state of the system depends on the relative phase of atomic and molecular fields at the time of the second pulse. This is somewhat analogous to Ramsey’s method of separated oscillating fields. Under very limited conditions (a near 1700 $a_0$), we could also observe Rabi-like oscillations with a single pulse towards the Feshbach resonance. This narrow window results from the conflicting needs for both strong coupling and condensate loss time$^{20}$ long compared to a Rabi oscillation period.

After pulse #2, a fraction of the coherent molecular component is converted into the energetic but still spin-polarized burst atoms through a yet to be determined process. Another mystery is the missing atoms. Are they molecules that are not converted back into atoms and are not detected in the burst or the remnant signals? If so, why do we not see them as atoms after the field is turned off and the corresponding molecular state is no longer bound? Why are there fewer missing atoms for lower-density condensates and quicker pulses towards the Feshbach resonance? What is the actual conversion efficiency from atoms to molecules and how could we maximize it? Very near the Feshbach resonance, the molecular state has a magnetic moment nearly the same as that of the free atoms, and hence will remain magnetically trapped. A major remaining question concerns the nature of the molecules. Could they be considered a molecular BEC? Clearly there is much to be learned about this curious system.

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