

Coherent optical association of single molecules

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We report on coherent association of a single weakly-bound NaCs molecule in an optical tweezer through an optical Raman transition without the use of a Feshbach resonance. Our scheme borrows transition dipole moment while reducing photon scattering by selecting a deeply bound electronic excited intermediate state. Starting from two atoms in their relative motional ground state, we achieve optical transfer efficiency of 50% . The molecule has a (change to 8.8G) zero-field binding energy of 770MHz and lifetime up to 1ms . (center of mass motional state population.) This technique is general (does not rely on narrow excited state linewidth) and could allow a wider range of molecular species to be assembled atom-by-atom.

Trapped neutral molecules, assembled in an array of optical tweezers, are a promising platform to study quantum information and quantum simulations. 1. we want to have a diverse species to tailor to different applications. 2. but so far all has been done with Feshbach association (before STIRAP), or in the case of Sr2, a narrow excited state. Previous work on near-threshold Raman transfer has been incoherent. (suggested referees: Florian Shreck and Immanuel Bloch)

In this Letter, we survey theoretically the best electronic intermediate state, detuning, etc that balance coupling/transfer efficiency and photon scattering to demonstrate coherent association of a electronic ground-state molecule from two atoms. Because the electronic excited state we used are generic in most diatomic molecules, this technique is general....

For atoms in the motional ground state in the tweezer, the average distance between the two atoms is about $1000a_0$. However, for the ground rovibrational molecule, the bond length is only around $4a_0$. The size mismatch between the initial and final states cause a very small coupling between them, which poses the biggest challenge for coherent creation of the molecule. Because of this, it is necessary to use an intermediate state with a size in between the initial and final states, and accomplish the transfer in two steps in order to increase the coupling.

The traditional choice for such an intermediate state is a FB molecule . However, the requirement of a suitable FB resonance limits its generality. Instead, we selected a weakly bound molecular state as the intermediate state and we use an optical Raman transition as the first step transfer. This approach minimizes the reliance on system specific properties and can therefore be more generally applied to creating other molecules. Such a transfer using optical transition has been observed in previous experiments . However, they either require the use of a narrow optical transition which limits the generality or were driving the transition incoherently due to

scattering. In this letter, we will present our result on coherent creation of a weakly bond molecule using only optical transitions on a broad optical line for the first time.

Even for the first step transfer to a weakly bound molecular state, it is still a challenge to achieve high Raman Rabi frequency to drive this transition due to the wavefunction size mismatch with the excited molecular state. For this reason, previous experiments also use weakly bound molecular excited states in the Raman transition. However, in addition to a higher Raman Rabi frequency, this choice of state also increases the scattering during the transfer process for two reasons. Firstly, since for a typical molecular potential the spacing between states decreases with increasing vibrational levels, a weakly bound excited molecular state limits maximum usable detuning and therefore the effectiveness of scattering reduction by increasing the detuning for the Raman transition. Secondly, due to size similarity, the weakly bound molecular ground state has a strong coupling with the atomic excited state which contributes significantly to the total scattering rate. The rate of such scattering process is proportional to $1/\delta_{thresh}^2$, where δ_{thresh} is the detuning from the dissociation threshold. This further increases the scattering rate for a weakly bound excited state. Based on these considerations, we calculated both the Raman Rabi frequency and the scattering rate at different detunings. As seen in figure (1....), despite a higher Raman Rabi frequency at the same detuning, the weakly bound state also has a much higher scattering rate and ultimately a smaller Rabi frequency to scattering at the optimal detuning. As a result, we use a deeply bound molecular excited state to drive the Raman transition.

(goes to SM?)

(mention scattering/lightshift depends on up/down matrix element ratio?)

In addition to the final and the excited state, it is also important to select the an initial atomic state in

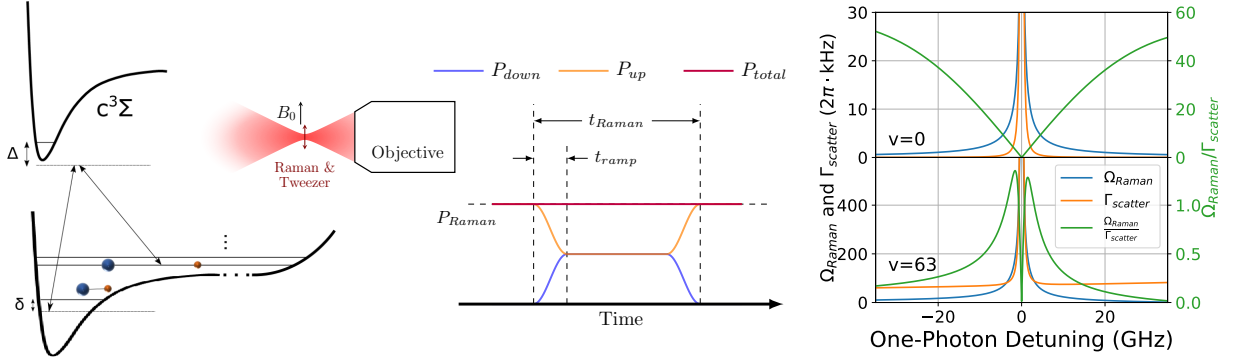


FIG. 1. Optical creation of single molecule from single atoms in tweezer. (A) Schematics of the Raman transition. (B) Geometry and polarization of trap and Raman beam relative to the bias magnetic field. (C) Molecule formation pulse sequence. The tweezer initially consists of only up leg power. This power is smoothly ramped down and the down leg power ramped up over $10\mu\text{s}$ while maintaining the total power of the tweezer.

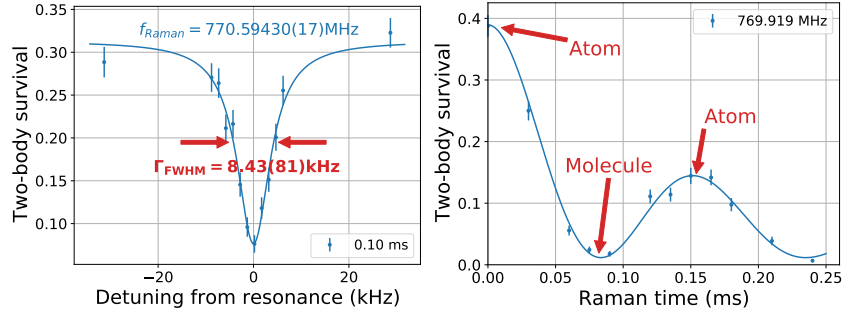


FIG. 2. (A) Raman resonance from atomic state to molecular state, showing Fourier limited linewidth. (B) Rabi oscillation on resonance

order to improve the coupling. Due to the small size of the molecular wavefunction, the coupling between the ground atomic state and the excited molecular state is approximately proportional to the value of the relative atomic wavefunction at short distance. In addition to the confinement, this value is related to the interaction between the two atoms. States with a large scattering length (positive or negative), the phase shift in the relative wavefunction between the atoms can significantly increase the short range wavefunction (add figure?). The increase in the coupling is proportional to (quote/cite Olive's equation?). For our system, among the stable spin combinations, 4422 and 3311 has a small scattering length of $|a| = \dots$ respectively and 3322 has a large and negative scattering length. (interaction shift \approx binding?)

Our experiment begins by loading a single ^{23}Na atom and a single ^{133}Cs atom into an optical tweezer from a dual-species MOT into separate optical tweezers. The atoms are imaged to distinguish between loading of two atoms, one atom (Na or Cs), or no atom during post selection. We then perform simultaneous Raman side-band cooling (RSC) to cool both atoms into a single 3-dimensional motional ground state in the tweezers. After

RSC, the Na tweezer is moved by sweeping the frequency on an acoustical optical beam deflector (AOBD) to overlap with the Cs tweezer before smoothly ramping off so that the Na and Cs atoms are merged into the same tweezer. The spin states for the Na and Cs atoms after RSC and during the merge process are $|F = 2, m_F = 2\rangle$ and $|F = 4, m_F = 4\rangle$ respectively. This states combination has a low scattering length of $\dots a_0$ which allows the two atoms to be merged into the same tweezer with minimum perturbation on each other and remains in their motional ground state after the merge.

After preparing the Na and Cs atoms in the same tweezer in a single quantum state, we perform interaction shift spectroscopy using a Cs Raman transition to drive the Cs into the $|F = 3, m_F = 3\rangle$ state. The new spin state combination has a larger scattering length of $\dots a_0$ which generates a interaction shift of $\dots k\text{Hz}$ in the tweezer. This interaction shift is larger than the differential axial trapping frequency between Na and Cs atoms, which decouples the relative and center of mass motional state and improves the robustness of our preparation of relative motional ground state. The stronger interaction in this spin state also enhances the atomic wavefunction

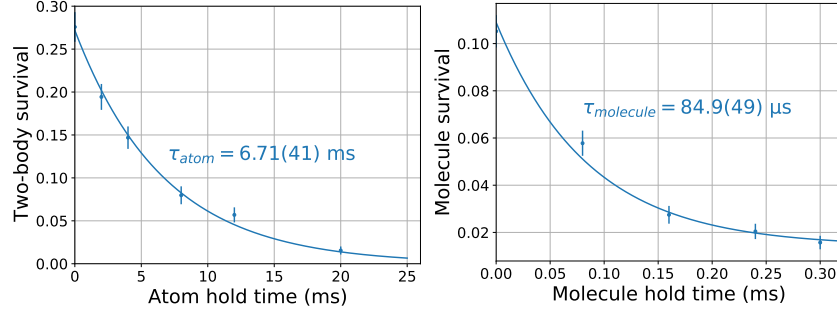


FIG. 3. (A) Two-body atom lifetime in 15 mW of trap depth (B) Molecule lifetime in 15 mW of trap depth

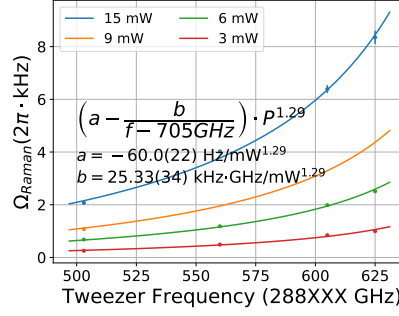


FIG. 4. Raman Rabi frequency vs detuning

at short range and increases its overlap with the intermediate molecular state used for our Raman transfer.

The optical transfer scheme we used is shown in figure . We use an optical Raman transition to drive the system from the atomic initial state to a ground electronic molecular state. In order to reduce the size mismatch between the atomic and molecular states, we selected the first bound state for the 3322 spin state as our final state. . For similar reason, the natural choice for the intermediate excited molecular state is one with highly excited motional level. However, from our calculation (and experiment?), the smaller level space for high vibrational state and the smaller detuning from the atomic threshold increases the the scattering rate of the molecular state which causes a reduced Raman Rabi frequency to decoherence rate ratio and a lower transfer efficiency. Therefore, in our experiment, we selected the $v=0$ state as the intermediate state for our Raman transition.

The pulse sequence for the experiment can be seen in figure (). Instead of adding another beam to drive the Raman transition on the atoms in the tweezer, we use the tweezer itself to achieve this goal. After the total tweezer power is set to the desired value, we smoothly ramp down the power of one frequency in the tweezer while simultaneously ramping up the power of a different frequency so that the total tweezer power remains unchanged. Both frequencies are kept on for a variable length of time before the process is reversed and we re-

turn to having a single frequency in the tweezer. The dual use of the tweezer beam ensures that there is not any undesired laser frequency that can interfere with the Raman transition. Using the tweezer beam for the Raman transition also allow us to maximize the intensity of the Raman beams and minimize the transfer time.

Figure () shows the geometry of the experiment setup. A 8.8G B field was used to defines the quantization axis in the experiment and is applied perpendicular to the tweezer axis. The tweezer beam (both frequencies for the Raman transition) has a π polarization relative to the quantization axis, which allows us to selectively drive the Raman transition to a final state with the same total m_F quantum number as our initial state.

This excited state used in the Raman transition was measured in our previous experiment using photoassociation to be at 288...GHz from our atomic state. The ground molecular state has not been observed previously in experimentally. Based on our measurement of FB resonance, interaction shift and the binding energy of the 4422 bound state. Theory prediction was at 770...MHz.

We locate the Raman resonance for the atom to molecule transition at 770...MHz (figure) with a 15mW tweezer at 288...GHz which corresponds to a ...GHz single photon detuning. The background level of ...% corresponds to the propability of preparing the two atoms in the relative motional ground state using interaction shift spectroscopy. When the atoms are transferred into

molecule by the Raman transition, there is a decrease in the two body survival since the resulting molecule is not directly detected by our image step.

We observed the narrowest linewidth of $\dots kHz$ for the Raman resonance at a pulse time of $\dots ms$, which corresponds to a linewidth-pulsetime product of \dots . This is consistent with the expected value of \dots for an ideal π pulse which is an evidence that the transfer is coherent. In order to verify the coherence of the transfer directly, we fixed the Raman frequency on the resonance and scanned the pulse time. figure ... shows the observed Rabi oscillation between the atomic and molecular states. Fitting the data with a decaying Rabi oscillation suggests that $\dots\%$ of initial ground state atoms are transferred into the molecular state.

In order to understand the fidelity of molecule formation, we fit our measurements to a model that includes a Raman Rabi frequency and a finite lifetime for the final molecular state. We also account for atomic state loss by measuring the single and two body lifetime for the atomic state (figure ...). The result suggests that our transfer has a Raman Rabi frequency of \dots and the molecule we form has a lifetime of \dots . We can also directly verify this result by measuring the lifetime of the molecule directly. This is done by applying a reverse Raman pulse to dissociate the molecule back to atoms after a variable wait time. The result of the direct measurement shows a molecular life-

time consistent with our fitting of the Raman transition data.

The molecule lifetime measured in the experiment is ... larger than the expected value of ... calculated from By measuring the lifetime for different tweezer powers, we observed a linear dependency of the lifetime (fig ...). This confirms that the short lifetime is mainly caused by scattering from the tweezer light rather than the intrinsic lifetime of the molecule itself. Additionally, we also measured the loss rate as a function of the tweezer frequency (fig ...) which shows only a very weak dependency for a $\dots x$ change in the detuning from the $v = 0$ excited state. This result is significantly different from the $1/\delta^2$ dependency expected from off-resonant scattering and suggests that the main source of the scattering is due to a broadband process that is not directly related to the $v = 0$ excited state.

(ASE filter?)

B field dependency $\dots kHz/G$ which agrees with theory prediction of $\dots kHz/G$. Dependency on tweezer power $\dots kHz/mW$, extrapolated to obtain the bare resonance at 0 tweezer power to be at $\dots MHz$.

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