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Two-photon photoionization of the Ca 4s3d 1D2 level in an optical dipole trap

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We report an optical dipole trap for calcium. The trap is created by focusing a 488-nm argon-ion laser beam into a calcium magneto-optical trap. The argon-ion laser photoionizes atoms in the trap because of a near-resonance with the 4s4f 1F1 level. By measuring the dipole-trap decay rate as a function of argon-ion laser intensity, we determine the 1F1 photoionization cross section at our wavelength to be approximately 230 Mb.

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I. INTRODUCTION

Laser-cooling experiments have expanded in recent years to group-II metals. This is partially due to a growing interest in optical frequency standards [1–3]. Calcium, strontium, and ytterbium all have narrow resonances from the ground state at relatively convenient laser wavelengths. The major isotopes of these elements have no angular momentum in the ground state, making the narrow “clock” transition frequencies less sensitive to external fields. Other experiments, such as metastable-collision studies, photoassociative spectroscopy and quasimolecule formation, Bose-Einstein condensation in simple atomic systems, and ultracold plasma investigations also contribute to the growing interest in laser-cooling alkaline-earth metals.

The atomic density in dipole traps can be much higher than in magneto-optical traps. For some experiments, such as Bose-Einstein condensation, photoassociation, cold plasmas, and collision studies, higher densities can be helpful. Dipole traps for alkaline-earth atoms may also improve the performance of atomic clock experiments. For example, atoms can be held in a dipole trap generated by a laser at the so-called “magic wavelength” where the ac Stark shift is exactly equal for both atomic levels in the clock transition [4,5]. This would make it possible to use trapped neutral atoms for the clock, increasing the maximum interrogation time and therefore increasing the accuracy of the clock.

Only a few experiments have explored optical dipole traps for alkaline-earth atoms. The absence of angular momentum in the ground state prevents sub-Doppler cooling using resonance transitions [6], complicating dipole-trap loading. However, advanced cooling techniques can reduce the atomic temperature to a few microkelvin, and dipole traps in Sr [7,8] and Yb [9,10] have been reported in the literature. We are also aware of a ground-state calcium dipole trap reported in a Ph.D. thesis [11].

In this paper we report an optical dipole trap for calcium. Our trap captures \( T \approx 1 \) mK atoms in the excited 4s3d 1D2 metastable state. Because the atomic temperature is relatively high, the dipole trap operates in the regime where the light shift is several times larger than the natural atomic-transition linewidth [12]. The high intensity required to capture these atoms is also high enough to photoionize them. By measuring the photoionization production rate and the trap lifetime as a function of dipole-trap laser intensity, we determine the effective 1D2 lifetime in our system and the photoionization cross section.

II. MAGNETO-OPTICAL TRAP

The calcium magneto-optical trap (MOT) is formed by three pairs of counterpropagating laser beams that intersect at right angles in the center of a magnetic quadrupole field [13]. The 423-nm laser light required for the calcium MOT is generated by frequency-doubling an infrared laser in KNbO3, and has been described previously [14]. A diode laser master-oscillator-power-amplifier (MOPA) system delivers 300 mW single-frequency power at 846 nm, as shown in Fig. 1. This laser is phase locked to a buildup cavity using the Pound-Drever-Hall technique [15], giving a power enhancement of 30. A 10-mm-long a-cut KNbO3 crystal in the small waist of the buildup cavity is used to generate 45 mW output power at 423 nm via noncritical phase matching at a temperature of −12 °C [14].

![FIG. 1. A schematic drawing of the MOT laser system and frequency-stabilization electronics used in these experiments. ACM: acousto-optical modulator. EOM: electro-optical modulator.](image-url)

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The laser is further stabilized by locking the 423-nm light to the calcium resonance transition using saturated absorption spectroscopy in a calcium vapor cell [16]. An acousto-optic modulator (AOM) in one arm of the saturated absorption laser beams shifts the laser frequency so that the laser beam sent to the MOT is 35 MHz (one natural linewidth) below the atomic resonance. We also use the AOM to chop this beam and use a lock-in amplifier to eliminate the Doppler background in the saturated absorption signal. Because the 846-nm laser is already locked to the frequency-doubling cavity, the feedback from this second lock controls the frequency-doubling cavity length.

The trap is loaded from a thermal beam of calcium atoms that passes through the center of the MOT. The thermal beam is formed by heating calcium in a stainless steel oven to 650 °C. The beam is weakly collimated by the 1-mm-diameter, 10-mm-long aperture in the oven wall. As the beam passes through the MOT, the slowest atoms in the velocity distribution are cooled and trapped. An additional laser beam passes through the MOT, the slowest atoms in the trap increases as the trap depth increases. The trap number falls to a flat line. At low powers, the number of atoms in the dipole trap, but there is no dissipative cooling mechanism to capture them.

Some of the $^1D_2$ atoms are ionized by the 488-nm laser beam via a two-photon transition to the continuum. This photoionization pathway is enhanced by a near resonance with the $^1F_3$ level. As discussed below, it is probably further enhanced by a near resonance in the Rydberg series leading up to the Ca II 3d ionization limit.

We measure the ion production rate for different 488-nm laser-beam intensities [22]. Sample data are plotted in Fig. 3(a). These data are replotted in Fig. 3(b) with the ion production rate divided by the laser power squared. This ratio is proportional to the number of $^1D_2$ atoms in the laser focus, and in the absence of a dipole trap, this signal should be a flat line. At low powers, the number of atoms in the dipole trap increases as the trap depth increases. The trap number maximizes when the average well depth is a few times the Doppler temperature. At higher power, the trap number falls off because the 488-nm laser beam shifts the $4s^2\,^1S_0$ and $4s4p\,^1P_1$ levels out of resonance with the MOT laser, reducing the efficiency of the optical-pumping loading mechanism. We performed a Monte Carlo simulation of trap loading and found agreement with our data, as plotted in Fig. 3(b).

At higher laser powers, the ion production rate is somewhat higher than expected. This occurs when the light shift of the $^1S_0$ and $^1P_1$ levels due to the 488-nm laser beam exceeds the natural linewidth. This is precisely the condition under which ground-state atoms can be captured in the dipole trap.

IV. DIPOLE-TRAP LOADING

We load the dipole trap while the MOT light is on. The dipole trap fills up with atoms optically pumped into the $^1D_2$ state in the region of the 488-nm laser-beam focus. Other $^1D_2$ atoms from outside the focal region pass through the dipole trap, but there is no dissipative cooling mechanism to capture them.

Dipole traps for heavier group-II atoms have been reported. Those experiments cooled on the intercombination lines, which have a much lower Doppler limit. This could be done in calcium, especially in conjunction with channeled cooling [19,20] or two-photon cooling [21]. However, no reports have been published to our knowledge.
ground $^1D_2$ atoms. But the background atoms roll through the trap in a few microseconds, and the trapped atoms remain in the trap approximately 100 times longer. The photoionization probability increases with the time spent in the 488-nm laser focus, so the ionization signal is predominantly from the trapped atoms.

V. TWO-PHOTON PHOTOIONIZATION RATE

We measure the lifetime of the dipole trap by blocking the MOT laser beams, and measuring the decay of the ion signal. The three most important decay mechanisms are radiative decay of the $^1D_2$ level, collisional decay due to hot atoms from the thermal atomic beam, and two-photon ionization of the $^1D_2$ atoms. Because we do not have a suitable method for turning off the thermal beam, we cannot reliably extract the $^1D_2$ radiative lifetime. Published values of the lifetime are around 2 ms [23–27], somewhat longer than measured in our experiment. While our experiment cannot determine the radiative lifetime, by measuring the decay rate as a function of 488-nm laser intensity we can determine the photoionization cross section.

A rate equation for $^1D_2$ level decay in the dipole trap after the loading has turned off is

$$\frac{dN_D}{dt} = -N_D \left( \frac{1}{\tau_{\text{eff}}} + A F^2 \right),$$

where $A$ is the two-photon ionization rate coefficient. This has the well-known solution

$$N_D(t) = N_D(0) \exp \left( \frac{1}{\tau_{\text{eff}}} + A F^2 \right).$$

In second-order perturbation theory, two-photon ionization is written as an overlap of the initial and final states summed over all possible intermediate states, divided by an energy denominator. For near-resonant ionization, the energy denominator makes the near-resonant term dominant, collapsing the sum to just one term. This one term looks like the product of the probability that an atom is excited into the $^1F_3$ state multiplied by the probability of photoionizing out of that state. We can write this term as

$$A F^2 = \frac{s^2}{1 + s + (2\Delta/\gamma)^2} \frac{I}{\sigma} \exp \left( \frac{1}{\tau_{\text{eff}}} + A F^2 \right),$$

where $\sigma$ is the $^1F_3$ one-photon photoionization cross section. The approximation in Eq. (6) assumes that $s \ll (2\pi\Delta/\gamma)^2$ and $\gamma \ll \Delta$. In our experiment, the intensity of the 488-nm laser has a Gaussian spatial profile. Averaging the square of the intensity over the laser profile allows us to relate the photoionization rate to the total laser power. In this case, it can be written as

$$A F^2 = \frac{\lambda^4 \sigma \gamma}{2\pi h c^3 \Delta^2 w^2} \frac{I}{P^2}. $$
FIG. 4. Dipole-trap decay versus power squared. The decay signal from the dipole trap. (a) shows the ion signal versus time with 1 W focused to a 90-μm waist. The effective decay rate for these data is 1.02 kHz. (b) shows the apparent decay rate for a range of powers, all focused to a 90-μm waist. The decay rate is fitted to a line proportional to the square of the power, as suggested by Eq. (3).

The dipole-trap decay rate as a function of the square of the 488-nm laser power is shown in Fig. 4. For each power level, we measured the dipole-trap decay. At sufficiently low power levels, this decay is approximately exponential, and we extract the decay rate using a least-squares fitting routine [28]. The decay rate depends on power. The zero-extrapolated decay rate is $\tau_{dl}^{-1}=0.93$ kHz. This rate is approximately twice the radiative decay rate. We see evidence in our experiment that our rate is significantly influenced by collisions with atoms in the thermal atomic beam. This is not surprising because the dipole trap sits in the middle of the thermal atomic beam. Without detailed characterization of the thermal beam, we cannot reliably extract a thermal-atom–$1\text{D}_2$-atom collision cross section.

The slope of the decay rate with power is 80 Hz/W². Using Eq. (7), we can determine the $^{1}F_3$ photoionization cross section. This gives a photoionization cross section for the $^{1}F_3$ level of $\sigma=230\times10^{-18}$ cm². This extraordinarily large cross section suggests that the final state lies near a Rydberg state in the continuum. The final state is 133 cm⁻¹ below the Ca ii $^2\text{D}_{3/2}$ ionization limit. The principal quantum number of hydrogenic Rydberg levels in this region is $n \sim 29$, and the separation between levels is 9 cm⁻¹. Our measurements are carried out in the presence of an electric field, further increasing the probability of finding a nearby Rydberg level. The NIST database tabulates only odd-parity levels in this energy region. We are unaware of applicable quantum defect calculations or measurements.

VI. CONCLUSION

We have demonstrated an optical dipole trap for neutral calcium atoms. These atoms are nonadiabatically loaded into the trap by an optical pumping mechanism. The lifetime of our trap is limited by the ~2 ms lifetime of the $^1\text{D}_2$ and by collisions with atoms in the thermal atomic beam.

Our initial interest in the calcium optical dipole trap was its potential application to our ultracold plasma research. It may be possible to use the dipole trap as the beginning point for ultracold plasma expansion studies. Because of the high aspect ratio, a plasma generated from a dipole trap would be a two-dimensional ultracold neutral plasma at early times. Correlation heating would be reduced compared to the three-dimensional case [29], bringing the two-dimensional plasma closer to the strongly coupled regime. Furthermore, the plasma expansion depends on the exact density distribution of the initial cloud. Compared to standard MOT traps, a dipole trap has a well-defined density profile, meaning that these new plasmas could greatly improve the reproducibility of plasma expansion experiments. We plan to explore these possibilities in future work.

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The trap depth is proportional to $I$, and the photoionization signal is proportional to $I^2$. At high laser powers, when atoms can be trapped even at large radii, it is possible that kinetic effects will skew the photoionization signal. Atoms trapped at the bottom of the potential will be ionized first, and atoms trapped at the edge of the potential will be ionized at a later time. For low enough laser powers, when the potential is too shallow to trap significant numbers of atoms, this effect should not influence the photoionization signal.