Parallel Low-Loss Measurement of Multiple Atomic Qubits
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Parallel low-loss measurement of multiple atomic qubits

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We demonstrate low-loss measurement of the hyperfine ground state of Rubidium atoms by state dependent fluorescence detection in a dipole trap array of five sites. The presence of atoms and their internal states are minimally altered by utilizing circularly polarized probe light and a strictly controlled quantization axis. We achieve mean state detection fidelity of 97% without correcting for imperfect state preparation or background losses, and 98.7% when corrected. After state detection and correction for background losses, the probability of atom loss due to the state measurement is < 2% and the initial hyperfine state is preserved with > 98% probability.

Experiments with qubits encoded in hyperfine states of neutral atoms are being actively developed as a route towards scalable quantum information processing[1]. Several research groups have demonstrated preparation and control of order 50 qubits in 1D[2], 2D[3, 4] and 3D[5] optical lattices. Quantum computation requires qubit state measurements to determine the result of a computation, and for measurement based quantum error correction[6]. Measurement of the quantum state of an atomic hyperfine qubit is most often performed by using a cycling, or near cycling, transition which repetitively transfers the qubit between a bright state $|B\rangle$ and an excited state $|e_B\rangle$. Detection of scattered photons due to illumination with light that is near resonant with the cycling transition projects the qubit into state $|D\rangle$. Conversely, if no photons are detected, the qubit is projected into the dark state $|D\rangle$. This idealized picture breaks down if the cycling transition is not perfectly closed, in which case an atom in state $|B\rangle$ may suffer a Raman transition to $|D\rangle$ thereby giving a measurement error.

Measurements that use a cycling transition rely on the availability of a metastable qubit dark state $|D\rangle$, or on shelving one of the qubit levels into a metastable dark state, as is done in trapped ion experiments[7]. In alkali atom experiments with qubits encoded in ground hyperfine levels the availability of a cycling transition generally relies on an angular momentum selection rule that is enforced by using probe light with a well defined polarization. This implies that the probe light propagates along a single axis in space which results in atomic heating due to the random direction of scattered photons. For a lossless measurement either the potential confining the atom should be sufficiently deep for the heating to be tolerable, as in experiments with ions[8], or the detection system should allow for a state measurement after scattering only a small number of photons to minimize heating. This latter approach was demonstrated with optically trapped atomic qubits[9–11] using single photon detectors.

There are several possible alternative measurement approaches including coupling of an atom to a high finesse cavity which enables state detection with minimal heat-

![FIG. 1. (color online) Experimental setup around the hexagonal vacuum cell. For state readout $\sigma_+$ polarized light propagates along $\pm \vec{k}_{RO}$ which is parallel to the quantization axis defined by the bias magnetic field $\vec{B}$. The trap light propagates along $\vec{k}_{ODT}$ and a dichroic beamsplitter separates the trap light and fluorescence light which is imaged onto the camera. Inset a) shows the levels used for readout including the bright $|B\rangle$ and dark $|D\rangle$ hyperfine levels, and the corresponding excited state levels $|e_B\rangle$, $|e_D\rangle$. Inset b) shows the experimental timing diagram.](image-url)
high quantum efficiency they suffer from excess readout noise which has hitherto rendered parallel lossless state detection infeasible. To circumvent this limitation previous array experiments used a "blow away" technique where atoms in $|B\rangle$ are ejected from the array using a single unbalanced beam, followed by detection of the presence or absence of an atom. Atom detection is performed using a 3D light field that cools the atoms, but does not prevent state changing Raman transitions during the measurement. This approach provides state measurements, but requires that a new atom has to be reloaded, half the time on average, which severely impacts the experimental data rate.

In this letter we show that low-loss state detection of multiple atomic qubits is possible in parallel using an EMCCD camera. This requires a careful choice of parameters to minimize both the motional heating rate (which is lower at large detuning) and the Raman depumping rate (which is lower at small detuning). The enabling advances include use of a moderately high numerical aperture (NA= 0.4) collection lens, deep optical traps, and careful preparation of the polarization state of the probe light to minimize Raman transitions from $|B\rangle \rightarrow |D\rangle$.

The detection method is compatible with an ideal, projective measurement of the quantum state that leaves the atom in an eigenstate of $\hat{\sigma}_z$ (for neutral atom qubits a state of definite $|F, M_F\rangle$). Observation of the dark state $|D\rangle$ leaves the atomic state unchanged. If the bright state is encoded in $|F_+, M_F\rangle$, with $F_+$ the upper hyperfine state, observation of the bright state $|B\rangle$ leaves the atom in $|F_+, F_+\rangle$ for any initial $M_F$. The qubit can then be restored to $|F_+, M_F\rangle$ by following the detection of $|B\rangle$ with a sequence of rotations using stimulated Raman transitions or microwave pulses. The measurement can also be made crosstalk free, a requirement for quantum error correction[18]. We emphasize that the experimental approach is compatible with quantum gate experiments in qubit registers[1], with no changes to the experimental apparatus. Since EMCCD cameras are available with up to $10^6$ pixels the method demonstrated here has the potential for scaling to large arrays with thousands of atomic qubits. Similar results to ours have been independently reported in [19].

The experimental geometry and measurement sequence are shown in Fig. 1. Atoms are prepared in the $|F = 1\rangle$ or $|F = 2\rangle$ hyperfine levels of the $^{87}\text{Rb} \ 5\sigma_{1/2}$ electronic ground state, corresponding to $|D\rangle$ and $|B\rangle$ respectively. Although qubits are defined in terms of specific hyperfine-Zeeman states $|1\rangle = |F_1, M_{F_1}\rangle$ and $|0\rangle = |F_0, M_{F_0}\rangle$ we use $|B\rangle$ and $|D\rangle$ here to represent random mixtures of the $M_F$ states of the $F = 2$ and $F = 1$ hyperfine levels, respectively. The demonstration of measurement of states $|B\rangle$ and $|D\rangle$ also applies, without modification, to any pair of hyperfine-Zeeman states as long as they are attached to different hyperfine levels.

To prepare states of single atoms we start with a standard magneto-optical trap (MOT) that is then overlapped with a 1D array of five optical dipole traps (ODTs) formed by focusing 1040 nm light to a waist of $w \approx 2.5 \mu$m. The traps are (2.8, 4.4, 5.6, 3.9, 3.4) mK deep and are spaced by $\sim 9 \mu$m. The traps are pencil shaped with atomic density distributions of size (standard deviations) $\Delta_x, \Delta_y \sim 7.0, 0.7 \mu$m, with the long axis along the optical axis of the collection optics. Single atoms are loaded with probability 20-30% at a temperature of $\sim 100 \mu$K.

In order to measure the initial trap populations, the atoms are probed using 6 MOT beams with components near-resonant with $|B\rangle \leftrightarrow |e_B\rangle$ and $|D\rangle \leftrightarrow |e_D\rangle$ simultaneously, where $|e_B\rangle$ is the $F' = 3$ level and $|e_D\rangle$ is the $F' = 2$ level of the $5\sigma_{3/2}$ excited state. Atom fluorescence is collected by a NA = 0.4 lens, and imaged onto an EMCCD camera (Andor iXon EM+ DU-860). The magnification was chosen such that the site separation is 2 pixels, and the signal from each ODT is integrated over a region of interest (ROI) defined by 5 camera pixels, as shown in Fig. 2a). We image each atom onto only a few pixels to minimize the electronic background noise incurred during camera readout. The excited states, $|e_B\rangle, |e_D\rangle$, are

![FIG. 2. (color online) (a) Regions of interest are five pixels enclosed by red borders with the relative photon counts on each pixel shown by the green shading. Each 5 pixel ROI receives (76, 88, 89, 92, 76)% of the light from the corresponding trapped atom. Neighboring site fluorescence crosstalk is $\sim 2\%$. Each pixel represents a 4 $\mu$m × 4 $\mu$m area and the site-to-site separation is $\sim 9 \mu$m. (b) Histograms of non-destructive readout in the central region (#2) for initial states |B⟩ and |D⟩. (c) The same data set post-selected on the presence of an atom in the ROI in the third measurement, leaving only Raman depumping and state preparation as sources of error. Signals in histograms are background-subtracted.](Image)
anti-trapped in the ODT, so to avoid heating the atom we toggle the ODT and the probe beams out of phase with a 50% duty cycle at 1.25 MHz. The photon detection efficiency is estimated to be 1.6 – 2.0%, accounting for the lens solid angle and dipole emission pattern (3.9%), transmission through optics (74%), EMCCD quantum efficiency ($\eta = 75\%$), and fluorescence lying outside of the camera pixels used to define regions of interest (76-92%).

Upon completion of the population measurement, there is a 100 ms delay for image transfer to the computer, after which the atoms are initialized in a random superposition of the Zeeman substates of one of the hyperfine levels, chosen by leaving either $|D\rangle \leftrightarrow |e_D\rangle$ or $|B\rangle \leftrightarrow |e_B\rangle$ on to depopulate the coupled state. To prevent leakage light from disrupting the initial substate mechanical shutters block unwanted light after initialization is completed. We estimate the state preparation fidelity for both states to be $>99.5\%$, limited by the fidelity of blow away measurements that are performed at reduced ODT depth.

After state initialization, a bias magnetic field $B_z \sim 20$ G making an angle of 60° from $\vec{k}_{\text{ODT}}$, the long axis of the ODTs, is switched on. The probe beams propagate long and counter to $\vec{k}_{\text{ROI}}$, which is set to be parallel to $\vec{z}$ with a possible small alignment error $\theta$. We use counter-propagating probe beams to mitigate the effect of heating due to near-resonant radiation pressure. In order to suppress Raman transitions both readout beams are $\sigma^+$ polarized which optically pumps the atoms into the lower state of the $|2,2\rangle \leftrightarrow |3',3\rangle'$ cycling transition. The counter-propagating probe beams are generated from separate lasers with a relative frequency offset of 500 kHz. This technique avoids standing wave patterns, which can cause a time dependent drift in the single atom scattering rate thereby broadening the camera signal distribution. During the state measurement sequence the trap depths are temporarily doubled to enhance retention of the atoms. The probe beams are set to saturation parameter $s_0 = 1$ (summed over both beams) and detuning $\delta = -\frac{2}{\gamma}$ red of the Zeeman shifted $|2,2\rangle \leftrightarrow |3',3\rangle'$ transition to provide maximal motional damping[20] with $\gamma$ the excited state linewidth. The atoms are illuminated for 6 ms with the same 50% duty cycle as is used for the population measurement and fluorescence light is collected by the EMCCD for analysis. The resulting data are shown in Fig. 2. The hyperfine state is determined on the basis of a simple threshold condition relative to the vertical dashed lines in Fig. 2b,c). Although more extensive analysis that utilizes information gained from the temporal or spatial distribution of light in each ROI can further reduce uncertainties[7, 19] our results show that the threshold condition alone is adequate for high fidelity measurements.

After an additional 100 ms delay for image transfer, a third readout sequence probes the atoms again. Depending on the experiment, the third readout is either a second population measurement for probing atom loss or a destructive state selective measurement using a blow away beam for measuring the number of atoms depumped from $|B\rangle$ to $|D\rangle$. Full characterization of the non-destructive measurement requires 4 experiments: 2 (state preparation $|B\rangle$ or $|D\rangle$) $\times$ 2 (blow away on or off). The results of the 4 experiments are summarized in Table I for the center site and Table II for the other sites. We note that the results marked with a) include 2% atom loss between each camera readout due to the finite trap lifetime $\tau \sim 5$ s and the 100 ms gap between each measurement. The background collision loss is not a fundamental limitation, and could be reduced by decreasing the chamber pressure or by shortening the image transfer time.

The primary limitation to the non-destructive measurement is the mean number of photons, $N_\gamma$, that can be scattered before the atom is depumped from $|B\rangle$ to $|D\rangle$. When using random polarization $N_\gamma = \frac{38340}{1 + 4\delta^2/\gamma^2 + s_0}$, where $s_0 = I/I_{\text{sat},\gamma}$ and $I_{\text{sat},\gamma} = 3.6$ mW/cm$^2$ is the saturation parameter for randomly polarized light, see [18] for a derivation. With typical experimental parameters 10$^4$ photons could be scattered which would lead to approximately 100 photo-electrons, which is technically enough to clearly resolve the $|B\rangle$ and $|D\rangle$ photon histograms. However the $|B\rangle$ state histogram would leave a long tail from depumping events during the exposure that would overlap with the $|D\rangle$ state distribution. Therefore,

### Table I

Results in the central site (#2) averaged over 2000 measurements. Data marked (a) are without correction, and data marked (b) are post-selected on the presence of an atom in the ROI in the third measurement, leaving only Raman depumping and state preparation as sources of error. The final state results are found from a third, state-selective measurement using a blow away beam.

<table>
<thead>
<tr>
<th>initial state</th>
<th>detected state (%)</th>
<th>final state (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$</td>
<td>B\rangle$</td>
<td>$</td>
</tr>
<tr>
<td>(a) 95.6(6)</td>
<td>98.6(1.9)</td>
<td>98.6(1.9)</td>
</tr>
<tr>
<td>(b) 98.0(4)</td>
<td>99.6(1.6)</td>
<td>99.6(1.6)</td>
</tr>
<tr>
<td>$</td>
<td>D\rangle$</td>
<td>0.4(1.6)</td>
</tr>
</tbody>
</table>

### Table II

Loss-corrected detection fidelities for the outer four traps. $|\psi\rangle_i$ is the prepared state.

<table>
<thead>
<tr>
<th>ROI</th>
<th>#0</th>
<th>#1</th>
</tr>
</thead>
<tbody>
<tr>
<td>$</td>
<td>\psi\rangle_i$</td>
<td>$</td>
</tr>
<tr>
<td>B</td>
<td>97.1(5)</td>
<td>2.9(5)</td>
</tr>
<tr>
<td>D</td>
<td>0(0)</td>
<td>100(0)</td>
</tr>
</tbody>
</table>

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in order to obtain clearly distinguishable photo-electron statistics we need the additional constraint that atoms scatter $\sim 10^4$ photons with minimal depumping, a condition that isotropic polarization does not satisfy.

To suppress the depumping we have used $\sigma_+-$polarized light along the quantization axis, to pump the atoms into $\{2,2\}$ the lower level of the cycling transition as described above. In an actual experiment there is still a finite depumping rate due to polarization impurity or a small angular mismatch $\theta$ between $\hat{z}$, the direction of the magnetic field, and $\hat{k}_\text{RO}$, the axis of the readout beams. The figure of merit is the number of photons that the bright state can scatter before it falls into the dark state, as shown in Fig. 3. We can quantify the probability of depumping by summing the rates over Raman depumping channels and comparing to the scattering rate on the cycling transition[18]. We estimate that we are able to scatter $N_{\gamma,\sigma} = 3.7 \times 10^5$ photons corresponding to an enhancement factor of $\sim 20$ over the unpolarized case with parameters $s_0 = 1$, $\delta = -\gamma/2$ and measured polarization purity $I_{\sigma_+}/I_{\sigma_-} = 1600$[18]. There is also a small transient contribution to the depumping probability as the atoms are pumped from the initial state towards $\{2,2\}$, which is estimated in [18].

It is also necessary to account for depumping due to the vector and tensor light-shifts imposed by the ODT. Circular polarization of the ODT light results in a vector shift on the atoms which adds a fictitious magnetic field, $B_\text{fict}$, along $k_{\text{ODT}}$. The 60$^\circ$ angle between $k_{\text{ODT}}$ and $k_\text{RO}$ drives Larmor precession, which reopens the depumping channels. In terms of the trap depth $U_0$ the fictitious field is $B_\text{fict}/U_0 = 29.77 A^{(1)}_{\{2,2\}} k_{\text{ODT}}/A^{(0)}_{\{2,2\}}$ (G/mK)

\[ \gamma = 3 \times 10^5 \text{ photons} \]