

2000-Times Repeated Imaging of Strontium Atoms in Clock-Magic Tweezer Arrays

Jacob P. Covey, Ivaylo S. Madjarov, Alexandre Cooper, and Manuel Endres*

Division of Physics, Mathematics and Astronomy, California Institute of Technology, Pasadena, California 91125, USA

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We demonstrate single-atom resolved imaging with a survival probability of 0.99932(8) and a fidelity of 0.99991(1), enabling us to perform repeated high-fidelity imaging of single atoms in tweezers thousands of times. We further observe lifetimes under laser cooling of more than seven minutes, an order of magnitude longer than in previous tweezer studies. Experiments are performed with strontium atoms in 813.4 nm tweezer arrays, which is at a magic wavelength for the clock transition. Tuning to this wavelength is enabled by off-magic Sisyphus cooling on the intercombination line, which lets us choose the tweezer wavelength almost arbitrarily. We find that a single not retroreflected cooling beam in the radial direction is sufficient for mitigating recoil heating during imaging. Moreover, this cooling technique yields temperatures below 5 μ K, as measured by release and recapture. Finally, we demonstrate clock-state resolved detection with average survival probability of 0.996(1) and average state detection fidelity of 0.981(1). Our work paves the way for atom-by-atom assembly of large defect-free arrays of alkaline-earth atoms, in which repeated interrogation of the clock transition is an imminent possibility.

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Optical lattice clocks of alkaline-earth(-like) atoms (AEAs) have reached record precision [1,2] for which the exploration of fundamental physics, such as geodesy [3], gravitational waves [4], and even dark matter [5], is now a possibility. Yet, despite the precise optical control of AEAs that has been demonstrated in a low-entropy array [2], the ability to address and detect single atoms is currently lacking. Such single-atom control techniques would provide new avenues for optical clock systems. Specifically, they are required for realizing a myriad of quantum computing protocols for AEAs using clock states [6–10] and could provide the foundation for generating and probing entanglement for quantum-enhanced metrology [11,12]. Optical tweezer (OT) techniques have matured into a powerful tool for single-atom control; e.g., they provide the versatility required for atom-by-atom assembly of defect-free arrays [13–17] and they automatically position single atoms at distances such that interaction shifts on the clock transition are expected to be strongly reduced [18]. Further, OT experiments generally have fast experimental repetition rates and, as demonstrated below, enable repeated low-loss clock-state read-out without reloading atoms. Such techniques could provide a pathway for quasicontinuous interleaved clock operation in order to tame the Dick effect [19]. Since state-insensitive magic trapping conditions are required for clock operation [20], tweezers operating at a clock-magic wavelength are highly suited for these directions.

Recently, two-dimensional arrays of AEAs, specifically Sr [21,22] and Yb [23], in optical tweezers have been demonstrated, including single-atom resolved imaging. Cooling during imaging has been performed on the narrow

$^1S_0 \leftrightarrow ^3P_1$ intercombination line (see Fig. 1), similar to quantum gas microscopes for Yb [24]. To this end, trapping wavelengths have been chosen such that the differential polarizability on this transition is small, enabling motional sidebands to be spectrally resolved in the case of Sr [21,22], but precluding the possibility of achieving a magic trapping condition for the optical clock transition. Significantly, a more versatile Sisyphus cooling mechanism [25] has recently been observed for Sr atoms [21,26], providing a general pathway for cooling on narrow lines with strong polarizability mismatch. This observation combined with prior predictions [27,28] should allow for tweezer trapping and cooling of AEAs—and more generally atoms with narrow transitions—in a very wide range of wavelengths.

Here, we demonstrate detection and cooling of single ^{88}Sr atoms in clock-magic optical tweezer arrays of wavelength 813.4 nm [2,12,29–32], where the loss during imaging is suppressed by 2 orders of magnitude compared to previous work with Sr [21,22]. Specifically, we find a survival probability of 0.99932(8) and a fidelity of 0.99991(1) for single-atom detection, enabling us to perform repeated high-fidelity detection thousands of times. We also observe lifetimes under laser cooling of more than seven minutes.

These values provide a benchmark for simultaneous low-loss and high-fidelity imaging as well as trapping lifetimes for single neutral atoms, including work with alkalis [13–17,33–35]. We expect this development to be important for improved scalability of atom-by-atom assembly schemes [13–17,33] and for verifying high-fidelity quantum operations with neutral atoms [34,36]. For example, the success probability in atom-by-atom assembly is fundamentally limited by p_s^M , where p_s is the combined

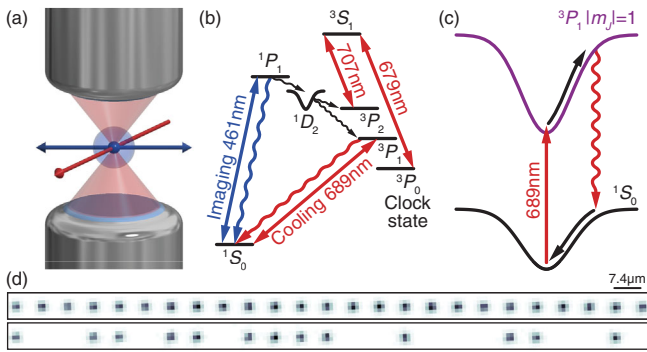


FIG. 1. Low-loss imaging scheme for ^{88}Sr . (a) Our single-atom imaging scheme requires only a single *not* retroreflected cooling beam at 689 nm for narrow-line Sisyphus cooling (red single arrow) to compensate recoil heating from fluorescence generated by exciting atoms with a single retroreflected 461 nm beam (blue double arrow). Microscope objectives with $\text{NA} = 0.5$ are used to generate 813.4 nm tweezers and to collect the fluorescence light. (b) Previous studies found an imaging loss channel via the decay of 1P_1 to 1D_2 with a branching ratio of $\sim 1:20000$, where atoms left the tweezers since 1D_2 was strongly antitrapped [21,22]. Crucially, the 1D_2 state is now trapped in 813.4 nm tweezers and our results show that atoms are recovered into the 3P_J manifold with very high probability. Two lasers (679 and 707 nm) repump atoms to 3P_1 , which decays back into the ground state, thus closing the 3P_J loss channel. (c) We use narrow-line *attractive* Sisyphus cooling on the $m_J = \pm 1$ states [26], originally proposed in Refs. [27,28]. This mechanism is based on the excited state being more strongly trapped than the ground state (in contrast to *repulsive* Sisyphus cooling demonstrated recently by us [21]). Atoms at the bottom of the trap are excited and have to climb up a steeper potential than they would in the ground state, leading to an average reduction in energy after spontaneous emission. Cooling results from the trapping potential mismatch, and not from photon recoil, thus requiring only a single cooling beam. (d) Average image (top) and single-shot image (bottom) of atomic fluorescence from twenty-five uniformized tweezers with an imaging time of one second.

survival probability for two images and hold time for rearrangement, and M is the final array size [14]. Our work improves this fundamental limitation of $p_S^M \sim 0.5$ to $M \gtrsim 1000$, enabling in principle assembly of arrays with thousands of atoms in terms of imaging- and vacuum-limited lifetimes. Finally, we demonstrate single-shot clock-state resolved detection with average fidelity of 0.981(1) and average atom survival probability of 0.996(1), which could be used for repeated clock interrogation without reloading.

Experimental techniques.—Single atoms are loaded stochastically from a narrow-line magneto-optical trap into an array of tweezers as described in detail in Ref. [21]. In contrast to our previous work, we use 813.4 nm light to generate tweezers [Fig. 1(a)]. While providing a magic wavelength for the clock transition, this wavelength also closes a previously observed loss channel, providing the basis for the low-loss detection demonstrated here [Fig. 1(b)]

[21,22]. Further, the imaging scheme is simplified to a single *not* retroreflected cooling beam at 689 nm and a retroreflected imaging beam at 461 nm. Both beams propagate in the plane orthogonal to the tweezer propagation axis. The cooling (imaging) beam is polarized parallel (perpendicular) to the tweezer propagation axis. We modulate the retroreflector of the imaging beam to wash out interference patterns [14]. The tweezers are linearly polarized and have a depth of $\approx 450 \mu\text{K}$ and waist of $\approx 700 \text{ nm}$. The array of 25 tweezers has a spacing of $\approx 7.4 \mu\text{m}$ and is uniformized to within $\approx 2\%$ [21]. Tweezer arrays are generated with a bottom objective, while a second, top objective is used to image the fluorescence light on an electron multiplied charge-coupled device (EMCCD) camera.

Cooling during imaging is based on a narrow-line *attractive* Sisyphus cooling scheme on the 7.4 kHz transition at 689 nm [Fig. 1(c)], following the original proposals in Refs. [27,28], which has been observed only very recently in continuous beam deceleration [26]. In contrast to the *repulsive* Sisyphus cooling scheme demonstrated recently by our group [21], the attractive scheme relies on the excited state experiencing a significantly stronger trapping potential than the ground state. For linearly polarized trapping light, this situation is realized for the $m_J = \pm 1$ sublevels of 3P_1 in wavelengths ranging from ≈ 700 to $\approx 900 \text{ nm}$. (For longer wavelengths, including 1064 nm, repulsive Sisyphus cooling can be used.) This enables us to fine-tune the wavelength to 813.4 nm, which is magic for the clock transition to 3P_0 , while providing cooling conditions for the transition to 3P_1 .

Imaging results.—Our results show simultaneous high-fidelity and low-loss detection of single atoms. First, we observe a histogram of photons collected in 50 ms with clearly resolved count distributions corresponding to cases with no atom and a single atom [Fig. 2(a)]. Taking a second image, after 29 ms hold time under Sisyphus cooling alone, we find a survival probability of 0.99932(8). At the same time, the fidelity of the scheme (defined by the accuracy of distinguishing no atom from a single atom [21]) reaches a value of 0.99991(1), demonstrating simultaneous low-loss and high-fidelity imaging. These values are enabled by a lifetime under imaging conditions of more than two minutes [Fig. 2(b)], while requiring only tens of milliseconds to acquire enough photons. We also find lifetimes under Sisyphus cooling (without 461 nm light) that are longer than seven minutes.

Without cooling or imaging the atoms, we observe lifetimes in the 1S_0 ground state between several seconds and 20 seconds, depending on the number of tweezers and their spacing—a similar behavior to earlier tweezer array experiments with acousto-optic deflectors [14]. The lifetime in the 3P_0 upper clock state is discussed below.

These results enable us to repeatedly image single atoms thousands of times. Specifically, we alternate between 50 ms long imaging blocks and 29 ms pure cooling blocks

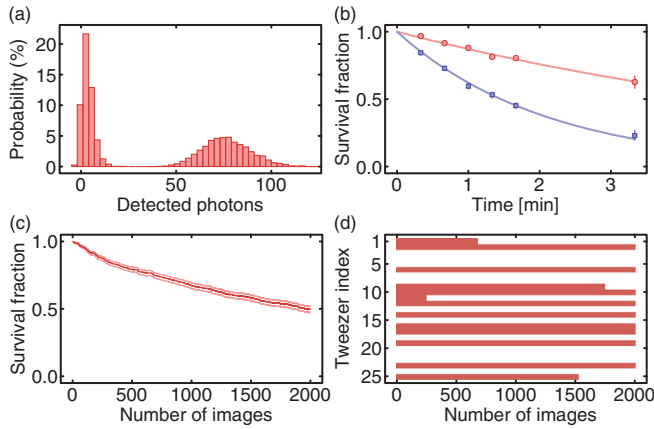


FIG. 2. Low-loss high-fidelity imaging. (a) Histogram of fluorescence counts from a single representative tweezer. We find a detection fidelity of 0.99991(1) and an average survival probability of 0.99932(8), demonstrating simultaneous high-fidelity and low-loss imaging. Results are for an imaging time of $t = 50$ ms under simultaneous repumping and Sisyphus cooling. (b) The survival fraction as a function of hold time in minutes under these imaging conditions (blue squares and fitted line). Importantly, we find a lifetime of $\tau = 126(3)$ s, while only needing $t \lesssim 50$ ms imaging time for reaching high detection fidelity, leading to small loss fractions consistent with $\exp(-t/\tau)$. Moreover, we find a lifetime of 434(13) s under Sisyphus cooling alone (without 461 nm), demonstrating a vacuum-limited lifetime greater than seven minutes (red circles and fitted line). (c) Survival fraction versus image number for 2000 repeated images. The dark red line represents the mean over 40 realizations, with the lighter red lines showing the standard error of the mean. Atoms are imaged with high fidelity for 50 ms followed by a 29 ms cooling block. (d) A representative realization of atom detection over the course of the 2000 images. Detected atoms are plotted in red versus the image number, where the rows represent the 25 tweezers. Note that we find no occurrences of atoms returning after they are lost. A video of the 2000 images is available in the Supplemental Material (SM) [37].

2000 times, and collect photons on the EMCCD camera in each imaging block. Recording the survival probability as a function of the number of images N , we find that even after 2000 high-fidelity images, the survival fraction stays above ≈ 0.5 . The decay follows an approximate exponential trend with p_1^N , where the single image survival probability is $p_1 \approx 0.9997$, slightly higher than the above quoted value measured with only two images. We note that the success probability in atom-by-atom assembly schemes is also limited by p_1^{2N} as a function of the final array size. (The factor of 2 appears because two consecutive images with interleaved hold times are needed for successful rearrangement.) Our results indicate that assembly of systems with thousands of atoms could be possible in terms of imaging fidelity, loss probability, and lifetime during the assembly step.

Cooling results.—These low-loss high-fidelity results are achieved by optimizing the Sisyphus cooling frequency

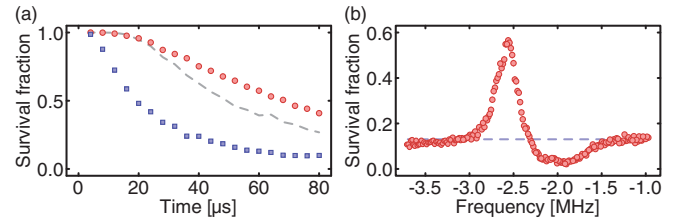


FIG. 3. Sisyphus cooling to low temperatures (a) The survival fraction in an array in a release-and-recapture measurement performed by diabatically turning off the traps for a variable time followed by a sudden switch-on [38]. We show data after imaging (blue squares) and after adding a dedicated cooling block with Sisyphus cooling alone (red circles). Results are compared with classical Monte Carlo simulations for a three-dimensional thermal distribution at $5 \mu\text{K}$ (dashed line). Note that the release-and-recapture method is mostly sensitive to the energy distribution in the radial direction. (b) Survival fraction in an array after release-and-recapture for $60 \mu\text{s}$ off-time versus the red frequency during Sisyphus cooling for 25 ms with an intensity of $I/I_s \approx 200$. The dashed line represents the case without a dedicated cooling block. We find that atoms are cooled for appropriately chosen red detunings, and heated for detunings further to the blue. This is consistent with an understanding of Sisyphus cooling as an *attractor* in energy space [27,28]. Data in (a) are at -2.6 MHz detuning.

and picking a conservative 461 nm scattering rate of ≈ 41 kHz as detailed in the Supplemental Material (SM) [37].

In addition, attractive Sisyphus cooling without the 461 nm beam results in a radial temperature below $5 \mu\text{K}$ (Fig. 3), which we quote as a conservative upper bound based on a release-and-recapture technique [38,39]. This technique is primarily sensitive to radial temperatures, and is compared to classical Monte Carlo simulations to extract a temperature. However, comparison to classical simulation would overestimate actual temperatures that are close to or below the energy scale of the radial trapping frequency ($T \lesssim (\hbar\omega/2k_B)$), which for us is roughly $2.4 \mu\text{K}$. More precise measurement of lower temperatures could be done via resolved sideband spectroscopy, which we leave for further work. An open question in this context is whether cooling to the motional ground state can be achieved in the strongly off-magic cooling configuration used here.

As Sisyphus cooling occurs due to a trapping mismatch between the ground and excited state, it is expected that cooling occurs in all directions even for a single radial cooling beam. The low loss we observe during imaging already provides evidence of this mechanism, as fluorescence recoil heating must be mitigated in all directions. Determination of the axial temperature after cooling can be achieved via techniques such as adiabatic rampdown [38,40] or spectroscopy of thermally broadened light shifts [21]. Our preliminary results with such techniques are consistent with three-dimensional temperatures similar to our quoted radial temperatures; however, we leave a

thorough investigation to future work. We note that we have made no explicit attempt to further cool the axial direction, and that doing so is likely possible by applying a beam in that direction. Finally, we note that the clock-magic condition of our tweezers opens the door to well-resolved sideband thermometry on the clock transition, which would be required to see resolved axial sidebands that are otherwise poorly resolved on the intercombination line at our trapping frequencies.

Clock-state resolved detection.—Finally, as an outlook we characterize our ability to perform low-loss state-resolved read-out of the optical clock states $^1S_0 \equiv |g\rangle$ and $^3P_0 \equiv |e\rangle$. Our scheme relies on shelving techniques that are routinely used in ion trap experiments to realize low-loss, high-fidelity state-resolved detection [41–43]. They are also prevalent in optical lattice clocks with alkaline-earth atoms, but have not been extended to single-atom-resolved imaging [31,32]. More generally, low-loss state-resolved detection of single *neutral* atoms has been realized only recently with alkali atoms [34,35,44–46]. Since in this case hyperfine states are used, simultaneous cooling during state-resolved detection is challenging and thus deep traps are required. This will limit scalability, and the approach has so far only been demonstrated in up to five traps [35]. Note that Stern-Gerlach detection of hyperfine spins via spatial separation in a lattice has been performed very recently [34,45], which provides an alternative route for high-fidelity lossless state-resolved detection.

Our scheme consists of two consecutive images (Fig. 4). In the first image, we aim at detecting atoms in $|g\rangle$. To this end, we turn off the 679 nm repump laser such that atoms in $|e\rangle$, in principle, do not scatter any photons. Hence, if we find a signal in the first image, we identify the state as $|g\rangle$. In the second image, we turn the 679 nm repump laser back on to detect atoms in both $|g\rangle$ and $|e\rangle$. Thus, if an atom is not detected in the first image but appears in the second image, we can identify it as $|e\rangle$. If neither of the images shows a signal, we identify the state as “no atom.” We find that the inaccuracy of this scheme is dominated by off-resonant scattering of the tweezer light when atoms are shelved in $|e\rangle$ during the first image. Specifically, by pumping atoms into $|e\rangle$ before imaging, we observe that, at our trap depth of $\approx 450 \mu\text{K}$, they decay back to $|g\rangle$ with a time constant of $\tau_p = 470(30)$ ms. This leads to events in the first image, where $|e\rangle$ atoms are misidentified as $|g\rangle$ atoms. To minimize the probability of misidentification, the first imaging time should be as short as possible. To reduce the imaging time, we compromise slightly on the survival probability in order to work with higher 461 nm scattering rates in the first image (see Supplemental Material [37]). Specifically we use $t = 15$ ms at a scattering rate of ≈ 72 kHz. The second image is performed with the same settings as in Fig. 2.

Additionally, atoms in $|g\rangle$ can be misidentified as $|e\rangle$ if they are pumped to $|e\rangle$ in the first image. This can occur

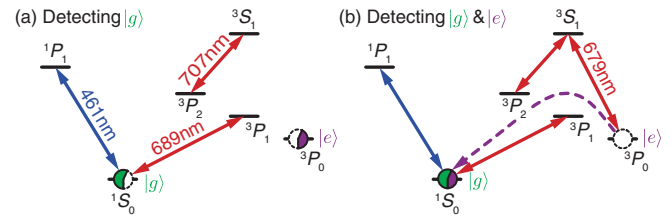


FIG. 4. Low-loss state-resolved detection. (a) A statistical mixture of optical clock qubit states is represented as a circle, where the green section at $^1S_0 \equiv |g\rangle$ represents the population in $|g\rangle$ and the purple section at $^3P_0 \equiv |e\rangle$ represents the population in $|e\rangle$. To measure the population in $|g\rangle$, we image without the 679 nm repumper during which $|e\rangle$ remains dark. The accuracy of measuring the population in $|g\rangle$ is limited both by off-resonant scattering of the tweezer light, which pumps $|e\rangle$ back to $|g\rangle$, and by pathways that pump $|g\rangle$ to $|e\rangle$ such as the 1D_2 decay channel and the off-resonant scattering from 3P_1 during cooling. As a result, the average state detection fidelity is 0.981(1). (b) We perform a second image that includes the 679 nm repumper, which pumps $|e\rangle$ to $|g\rangle$ via the 3S_1 state and the 707 nm repumper, such that both states are detected. The pumping process is illustrated by the purple arrow. This image measures the population in $|g\rangle$ and $|e\rangle$, and informs whether the atom has been lost as a result of the first image. We find that the average survival probability of this double-imaging sequence is 0.996(1).

either via the 1D_2 leakage channel and subsequent scattering of 707 nm photons, or via off-resonant scattering of the trap light from 3P_1 during cooling. We identify this misidentification probability by initializing atoms in $|g\rangle$, and counting how often we identify them as $|e\rangle$ in the state-resolved imaging scheme.

In summary, we place a conservative upper bound for the probability of misidentifying $|e\rangle$ as $|g\rangle$ by $e^{-t/\tau_p} = 0.031(2)$ and we directly measure the probability of misidentifying $|g\rangle$ as $|e\rangle$ yielding 0.008(1). We define the average state detection infidelity for a generic initial state as the mean of these probabilities [34], yielding an average state detection fidelity of 0.981(1). Further, we similarly define the average survival probability of the double imaging scheme in terms of the measured survival probabilities of $|g\rangle$ and $|e\rangle$, for which we obtain 0.996(1).

Our fidelity is comparable to recent measurements with alkali atoms in tweezers [35,44,46], yet our survival probability is substantially higher than any tweezer- or lattice-based schemes to our knowledge [34,35,44–46]. These results constitute an excellent setting for continuous measurement in an optical clock. However, we emphasize that this investigation was not exhaustive, and we expect that further optimization of these imaging parameters is possible. In general, these values could be further improved by either imaging in shallower tweezers or in tweezers at a wavelength further detuned from higher lying states. For instance, tweezers operating at 1064 nm are a promising possibility, and would be a convenient choice for operating a quantum gas microscope. Further, it is possible to switch

between 813.4 nm tweezers or lattices for clock interrogation during which the trap depth can be orders of magnitude lower, and 1064 nm tweezer or lattices for imaging.

In conclusion, we have addressed two major limitations of tweezer arrays for optical clock-based quantum information processing and metrology. By working at the magic wavelength for clock operation, we observe imaging with a fidelity of 0.99991(1) and a survival probability of 0.99932(8), and lifetimes under cooling of more than seven minutes. By employing a double imaging technique with specific combinations of repump lasers, we study low-loss state-resolved detection and observe an average fidelity of 0.981(1) with an average survival probability of 0.996(1). This work provides a setting for continuous measurement in an optical clock that can suppress laser fluctuations due to the Dick effect [19,47]. Clock operation on bosonic isotopes of AEAs such as ^{88}Sr used in this work has been performed with Sr [48] and Yb [49,50]. Moreover, the tools developed in this work enable excitation to highly excited Rydberg states in the 3S_1 series via the clock state 3P_0 . Engineering long-range Rydberg-mediated interactions facilitates the generation of entanglement between optical clock qubits, which can be used for quantum information processing [51], quantum simulation [52,53], and quantum-enhanced metrology via spin squeezing [11].

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*mendres@caltech.edu

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