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Two-dimensional array of microtraps with atomic shift register on a chip

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Abstract. Arrays of trapped atoms are the ideal starting points for developing registers comprising large numbers of physical qubits for storing and processing quantum information. One very promising approach involves neutral atom traps produced on microfabricated devices known as atom chips, as almost arbitrary trap configurations can be realized in a robust and compact package. Until now, however, atom chip experiments have focused on small systems incorporating single or only a few individual traps. Here, we report experiments on a two-dimensional array of trapped ultracold atom clouds prepared using a simple magnetic-film atom chip. We are able to load atoms into hundreds of tightly confining and optically resolved array sites. We then cool the individual atom clouds in parallel to the critical temperature required for quantum degeneracy. Atoms are shuttled across the chip surface utilizing the atom chip as an atomic shift register and local manipulation of atoms is implemented using a focused laser to rapidly empty individual traps.

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1. Introduction

Of central importance in the quest for a quantum computer is the issue of scalability. In the trapped-ion approach, for example, microfabricated-chip traps [1, 2] have recently been able to accommodate several ions [3, 4] and shuttle them between different zones [5]. Neutral atoms, on the other hand [6], possess the significant advantage of an intrinsically weaker interaction with the environment, yet are routinely manipulated using optical and magnetic trapping techniques. So far, the primary tool used to create registers of neutral atoms has been the optical lattice, which allows near-perfect arrays of atoms with typical separations of half the optical wavelength [7]; however, addressing and locally manipulating atoms on this length scale is a serious challenge. This has motivated the development and use of large-period optical lattices [8, 9] or patterned loading techniques [10], as well as addressable arrays of optical traps produced using holographic techniques [11] or micro-lens arrays [12].

Magnetic microtraps integrated on atom chips [13–16] are a promising alternative to these optical approaches. An atom chip typically consists of a microscopic pattern of wires (microelectromagnets) or permanent magnets [17–19] arranged on a substrate so as to produce tailor-made magnetic potentials for ultracold atoms. Qubits encoded in the ground-state hyperfine levels of the trapped atoms could offer exceptionally long coherence times [20], and quantum-logic gates could be implemented via controlled collisions [21] or by long-range interactions using laser excited Rydberg states [22]. With established micro-/nano-fabrication techniques, it is then feasible to scale-up and integrate many microtraps on a single chip. In the earlier work, one-dimensional periodic potentials [17, 23–25] and small arrays consisting of several traps [13, 26] have been produced using both current-carrying wires and magnetic materials. The present work greatly extends these results to create and study a two-dimensional array of traps accommodating more than 500 atom clouds. This is made possible using two-dimensional structures lithographically patterned into magnetic films [27] that allow the integration of hundreds of tightly confining traps in custom-designed and optically resolvable microtrap arrays.
Figure 1. Magnetic lattice atom chip and experimental setup. The experiments are performed using a FePt magnetic film, lithographically patterned with a two-dimensional lattice with periods of 22 and 36 \( \mu \text{m} \) and coated with a reflective gold layer. The film is uniformly magnetized along \( z \) and combined with an external field \( B_y \) to create an array of magnetic microtraps for atoms positioned 10 \( \mu \text{m} \) from the film surface. For detection, the trapped atoms are illuminated by a retro-reflected probe laser beam that is partly absorbed by the atoms and imaged onto a CCD camera for analysis. Projected on the backplane of the figure is a microscope image of the FePt film with a few trap positions indicated by yellow ellipses.

2. Magnetic microtrap arrays

At the core of our experiment (figure 1) is an atom chip (for design and fabrication details see [27]), which incorporates a single 300 nm-thick FePt film, patterned using optical lithography, then gold-coated and magnetized in the direction perpendicular to the film surface (\( M_z = 670 \text{ kA m}^{-1} \)). Combined with an external field (\( B_y = 22.5 \text{ G} \)), this simple set-up produces an array of magnetic field minima (magnetic lattice), which is used to trap ultracold \(^{87}\text{Rb} \) atoms in the \( |F = 2, m_F = 2 \rangle \) state. In the following experiments, each trap is produced 10 \( \mu \text{m} \) from the surface and provides tight harmonic confinement to the atoms (with trapping frequencies \( \omega_\parallel /2\pi = 5.5 \text{ kHz} \) and \( \omega_\perp /2\pi = 13.4 \text{ kHz} \) in the axial and transverse directions, respectively). The distances between neighbouring traps are 22 and 36 \( \mu \text{m} \) (1250 traps \( \text{mm}^{-2} \)), chosen to facilitate optical addressing. The barriers between traps, the trapping frequencies and the trap positions are precisely controlled in parallel through the external field strength and orientation. We believe magnetic-film atom chips are best suited for high-density integration, as for example, an equivalent potential produced using currents through micron-sized gold wires would require a power dissipation of \( \sim 30 \text{ W mm}^{-2} \).
We first describe the loading of the microtrap array. A cloud of $^{87}\text{Rb}$ atoms is collected and cooled in a mirror-magneto-optical trap and then transferred to a single magnetic trap created by a current-carrying wire and the external field [15]. A two-dimensional view of the atomic distribution is obtained using reflective resonant absorption imaging (figure 1). The cloud contains $N = 1 \times 10^6$ atoms at a temperature of $\sim 30 \mu\text{K}$ before it is transferred to the film surface by ramping off the wire current. The atom cloud first becomes corrugated as the trap merges with the field of the lattice, until for zero wire current the atoms are confined in the microtrap array alone (figure 2). We determine the number of atoms in each trap by integrating the absorption over each unit cell. During merging, the atom cloud expands over an area of $\sim 1.4 \times 0.4 \text{mm}^2$, populating more than 500 traps, each with $N > 200$ and up to 2500 atoms, showing that it is possible to transfer atoms from a single magnetic trap to hundreds of microtraps. After loading, we observe atom loss over two characteristic time scales: minor initial loss over the first 100 ms, caused by residual evaporation of atoms over the potential barriers, then background loss, occurring with a time constant of around 2 s.

### 3. Evaporative cooling and site-resolved thermometry

#### 3.1. Radio-frequency (rf) evaporation and spectroscopy

For storing quantum information, it is desired to initialize the register to a well-defined state with negligible thermal excitations. One powerful feature of magnetic lattices is the availability of rf cooling techniques, which we show can be applied to cool hundreds of atom clouds in parallel. We perform forced evaporative cooling using a two-part rf sweep. Rf spectroscopy [19] is then used to characterize more than 150 microtraps near the centre of the array (figure 3).

Sweeping the rf from high to low frequency continuously couples the more energetic atoms to untrapped Zeeman states (that subsequently leave the traps), whereas the remaining atoms

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**Figure 2.** Loaded array of magnetic microtraps. An absorption image of the two-dimensional atomic distribution (shown in false colour) after transfer to the magnetic lattice. The optical resolution is approximately 7 $\mu\text{m}$. The inset shows a magnified region of the central part of the array as indicated by the outlined box. Atoms populate more than 500 array sites over a region of $1.4 \times 0.4 \text{mm}^2$ of the chip surface; the scale bar corresponds to a distance of 250 $\mu\text{m}$. The solid arrow indicates the direction of transport for the atomic shift register experiment (figure 4).

Figure 3. Radio-frequency evaporative cooling and site-resolved thermometry. Forced rf evaporation is used to cool the individual atom clouds to the critical temperature for quantum degeneracy. Site-resolved rf spectroscopy and three-body decay measurements are performed to determine the distribution of cloud temperatures. (a) Two representative rf spectra and fits (solid lines) for one selected site for final evaporation frequencies of $\nu_f = 13.5$ MHz (red squares) and $\nu_f = 8.50$ MHz (green diamonds), with fitted values for $N$ and $T$ labelled. The fraction of atoms remaining in a well after the application of an rf outcoupling pulse is plotted along the vertical axis. (b) Two representative three-body decay curves and fits (solid lines) for the same array site as in (a) for $\nu_f = 8.50$ and 8.15 MHz (blue circles). The dashed line indicates the expected loss due to background vapour collisions with $\tau = 2$ s. (c) The full set of data $(N, T)$, for $>150$ array sites and five different final evaporation frequencies. The error bars represent statistical uncertainties estimated from several repetitions of the experiment. The blue-dashed line indicates a constant evaporation efficiency of $d(\ln T)/d(\ln N) = 1.2$ calculated for one site. The dotted line marks the Bose–Einstein condensation transition (degeneracy temperature) in the thermodynamic limit neglecting interactions. The numerically calculated boundary to the grey region is for a constant phase-space density of $\rho_0 = 2.612$, corresponding to the onset of Bose–Einstein condensation including the effects of finite atom number and interactions.

rethermalize to lower temperatures. In our experiments, the field minima of the potentials is around $11.41 \pm 0.05$ G corresponding to a Zeeman energy splitting of $\sim 7.99$ MHz. We first ramp from 13.5 to 8.5 MHz in 200 ms (corresponding to effective trap depths of around $48 \mu$K), then from 8.5 to 8.15 MHz in 35 ms ($15 \mu$K). In rf spectroscopy, an rf field is pulsed on for $\sim 1$ ms with a variable frequency to selectively outcouple atoms depending on their energy. We record an absorption image to determine the number of atoms remaining in each trap after

the pulse and repeat the experimental cycle for each value of the spectroscopy frequency to construct a collection of energy distributions (one for each site (figure 3(a))). The amplitude of the rf field is adjusted for each dataset to ensure a sufficient number of atoms are outcoupled for detection (typically 30–50% at peak), with typical Rabi frequencies of $\Omega \approx 2\pi \times 15$ and $2\pi \times 4$ kHz for $\nu_f = 13.50$ and 8.50 MHz datasets, respectively. We fit the measured spectra to a spatial outcoupling model [19] based on the calculated atomic distributions (see below) and including finite spectral resolution, to determine the cloud temperatures and the magnetic field strengths at the trap minima. For the spectra shown in figure 3(a), we observe low-energy tails associated with broadening widths of 30 and 10 kHz for $\nu_f = 13.50$ and 8.50 MHz datasets, respectively.

Rf spectroscopy is performed at several intervals during the evaporation sweep, and analysed independently for each microtrap to determine the cloud temperature $T$ and the magnetic field strength at the trap minimum (figure 3(a)). Before evaporation, we find the cloud temperatures are nearly equal for all traps (red-squares in figure 3(c)) with $\langle T \rangle = 43 \pm 4(3)$ µK (most likely constrained by the trap depth of around 400 µK). Here, we quote the rms spread among the traps with the estimated contribution from statistical uncertainties per trap in parentheses.

3.2. Magnetic field variations

The precise values of the magnetic field strengths at the trap minima are also determined using rf spectroscopy performed after the evaporation sweep. We find site-to-site variations of $\Delta B = 46(5)$ mG $\text{rms}$, which appear randomly distributed over the lattice, but with some long-range correlations observed on the 100 µm length scale. Compared with the total field produced by the film the figure of merit for trap variations is $\Delta B/B_y = 2 \times 10^{-3}$ for a trap height of 10 µm. This value compares favourably with previous studies of permanent magnet atom chips for trap heights between 50 and 100 µm [28, 29], but is a factor of $\gtrsim 10$ larger than experiments using thin Au wires at comparable surface distances [30], and approximately 100 times larger than for broad wires [31]. The precise origin of the variations is unknown, but could be associated with small imperfections in the patterning process or slight demagnetization of the magnetic film during vacuum bakeout. We note that the requirements on the homogeneity of the applied bias fields are not very stringent ($\sim 10^{-3}$ over 100 µm) and are easily satisfied by the external Helmholtz configuration coils. Although inhomogeneity typically causes fragmentation of atom clouds in elongated trapping potentials, in our two-dimensional array each trapped cloud is much smaller than the characteristic length scale of the variations and experiences a single harmonic potential well. Furthermore, the small site-to-site variations affect our prospective qubit states equally (minimizing dephasing [20]), and any residual effects could be precisely calibrated using site-resolved rf spectroscopy.

3.3. Density-dependent decay

During rf evaporation, we find high cooling rates and high evaporation efficiencies ($d[\ln T]/d[\ln N]$ between 1.2 and 1.7), attributed to the extremely high elastic collision rates present in our stiff traps (between 2500 and 5000 s$^{-1}$), despite the relatively small number of atoms per array site. At low temperatures (<5 µK), we find that rf spectroscopy becomes unreliable due to unavoidable spectral broadening caused by a combination of power...
broadening, rapid decay from the untrapped state and depletion. Additionally, the influence of the rf probe on the atomic energy distribution cannot be neglected at these temperatures, where collision rates are high, resulting in the redistribution of energy during the spectroscopy pulse. Instead, towards the end of the evaporation sweep we observe high decay rates ($\gamma > 20 \text{ s}^{-1}$) due to three-body recombination loss [32, 33], which are accurately measured to characterize the array (figure 3(b)).

A collection of decay curves is obtained by holding the atoms for a variable duration (one value per experimental cycle) before taking an absorption image and integrating the number of atoms in each site. During the hold time we apply a rf ‘shield’ tuned to the final evaporation frequency $\nu_f$ to limit the trap depth and prevent heating. Three-body decay curves are then modelled with the following differential equation:

$$\dot{N} = -L \left[ \int n_0^3 d^3r + 9 \int n_0^2 n_{\text{th}} d^3r + 18 \int n_{\text{th}}^2 n_0 d^3r + 6 \int n_{\text{th}}^3 d^3r \right] - N/\tau,$$

where we have taken into account wavefunction symmetrization [34], $L = 1.8 \times 10^{-29} \text{ cm}^6 \text{ s}^{-1}$ is the three-body decay coefficient taken from [33] and $\tau = 2 \text{ s}$ is the $e^{-1}$ decay time for one-body (background) loss calibrated against dilute clouds trapped within the array. The atomic density distributions for a thermal fraction, $n_{\text{th}}(r)$ and a Bose-condensed fraction, $n_0(r)$ for a given $N$ and $T$ are obtained using a self-consistent mean-field model. The model accounts for quantum statistics and the effect of interactions for both the thermal and condensed components but neglects the kinetic energy of the condensed fraction via the Thomas–Fermi approximation.

We approximate the trap potential as isotropic $V(r) = m \tilde{\omega}^2 r^2/2$ with mean trap frequency $\tilde{\omega} = (\omega_1 \omega_2^3)^{1/3}$, and iteratively solve the following set of equations [35]:

$$n_0(r) = \max \left[ 1 \quad (\mu - V(r)) / 2g n_{\text{th}}(r), \quad 0 \right],$$

$$n_{\text{th}}(r) = \frac{1}{\Lambda^3} g^{3/2} \left[ \text{e}^{(\mu - V_{\text{eff}}(r))/k_B T} \right],$$

$$V_{\text{eff}}(r) = V(r) + 2g \left( n_{\text{th}}(r) + n_0(r) \right)$$

with $N = \int (n_0 + n_{\text{th}}) d^3r$ and using the standard definitions: $g$ is the coupling strength, $\Lambda$ is the thermal de Broglie wavelength, $g_n[z]$ is the polylogarithm function and $\mu$ is the chemical potential. The numerical solutions are then used to predict the spectral energy distribution and three-body decay rate for fitting data as a function of $N$ and $T$ over the full range probed in our experiments. The solutions are also used to numerically determine the line $n(0)\Lambda^3 = 2.612$ in figure 3(c). For a more complete description of these models see [35].

We assume that thermal equilibrium is maintained due to the high elastic collision rate and we treat the cloud temperature as a constant parameter during decay. Fitting the numerical solution of equation (1) to each decay curve provides the initial $N$ and $T$. For all sets of experimental parameters, the three-body decay curves are measured between 6 and 15 times and fit independently to reduce noise and estimate statistical uncertainties. For the intermediate temperatures ($\nu_f = 8300$ and 8500 kHz datasets), we compare the results of both three-body decay and rf-spectroscopy measurements to calibrate systematic uncertainties such as the effective absorption cross-section used to determine $N$ from absorption images.

The measurements taken after the evaporation sweep indicate peak atomic densities $> 1 \times 10^{15} \text{ cm}^{-3}$ and temperatures around $\langle T \rangle = 2.0 \pm 0.3 \mu \text{K}$ (blue circles in figure 3(c))
corresponding to 2.6 and 7.1 mean vibrational quanta in the transverse and axial modes, respectively. From this, we infer that each cloud is smaller than the optical wavelength (Lamb–Dicke regime), with $\text{rms}$ position spreads of $0.17 \pm 0.01$ and $0.4 \pm 0.03 \mu m$. Our data are consistent with $\sim 70$ of the 156 analysed clouds having a phase-space density ($\rho > \rho_0 \simeq 2.612$) corresponding to the onset of Bose–Einstein condensation. However, of these clouds, we infer an average condensate fraction of less than 0.1. Around this point the cooling efficiency sharply decreases due to increased three-body loss for partially condensed clouds, where preferential loss of atoms from the dense, Bose-condensed components during evaporation limits the final temperatures to around $T_c$ (solid line in figure 3(c)). The observations are supported by our model that predicts three-body decay times, $\gamma^{-1}$, of around 20 ms (7 ms), for $N_{\text{BEC}} = 50$ (200), respectively. This effect can be circumvented by using the weaker-confined $|F = 1, m_F = -1\rangle$ state which also has a lower three-body-decay coefficient [32]. Alternatively, three-body decay could be used to our advantage to produce a well-defined number of atoms in each microtrap and to approach the few-atom regime for quantum information processing.

4. Atomic shift register

In addition to trapping and cooling the atom clouds, we have transported the atoms using the atom chip as an atomic shift register. This parallelized version of atomic [13, 36] and molecular [37] conveyor belts is the cold-atom analogue of an electronic CCD (charge-coupled device). The atoms are trapped in local minima of the total magnetic field strength, which is the superposition of fields from the chip and the external field. Thus the position of field minima depends also on the external field orientation. By applying time-varying, external uniform fields in a cyclic fashion we can move the minima across the chip surface. Rotating the external magnetic field continuously displaces the traps, shifting the array along one lattice direction as shown in figure 4. We have computed the optimized field components (figure 4(a)) required to keep the traps at a fixed height of $10 \mu m$ and with the field minima at a constant 3.23 G. The shifting direction is reversed by simply reversing the order of the applied fields. Details of this calculation, as well as restrictions to where one can or cannot move the atoms can be found in [38].

In the experiment, we first cool the atom clouds to a temperature of $\sim 10 \mu K$ and then linearly ramp between six settings of the external field per period, in $6 \times 3$ ms intervals. We investigate the operation of our atomic shift register using thermal clouds that provide a good signal to noise in a regime where interactions can be neglected and three-body loss is reduced. By repeating the shifting sequence, we have transported the atoms over a round-trip distance of $360 \mu m$, with five shifts in the $(2, -3)$ direction (figure 2) followed by five reverse shifts (figure 4(b)). During the experiment, we observe about 40% atom loss. This is consistent with expectations for three-body loss over the 183 ms experiment duration. We have also measured the effect of the shift register on the temperature of the clouds using rf spectroscopy. In the experiment, we keep the total trapping time after the evaporation sweep constant, but in the first measurement we transport the atoms over five lattice periods, whereas in the second measurement they remain trapped in their original positions. We measure mean temperatures of $10.9 \pm 0.7$ and $11.5 \pm 0.6 \mu K$ with and without transport, respectively, consistent with no observed additional heating. The demonstrated ability to smoothly shuttle atoms across the chip is an important ingredient for a scalable architecture for quantum computing and is an inherent feature of our magnetic-film atom chip.

Figure 4. Transport of atoms over the chip surface: atomic shift register. An external homogeneous magnetic field with rotating orientation smoothly shuttles the array of atom clouds in one direction and back. (a) Calculated optimal time sequence for the external magnetic fields $B_x$ (red), $B_y$ (green) and $B_z$ (blue) applied to carry out a single cycle shift operation. The field values shown as open symbols are applied in the experiment and interpolated by linear field ramps. (b) Line scans of optical density images along the shifting direction (arrow indicated in figure 2). Data for an integer number of shift cycles are shown, each with a vertical offset of $-0.1$ for clarity. A total of five shifts, each covering one lattice period, and five in the reverse direction to the original position are shown. The full sequence takes 183 ms and the atoms are transported over a distance of 360 µm. The inset shows intermediate steps within the first shift cycle for each 3 ms interval.

5. Single-site addressing

Finally, we show the addressable nature of our microtrap array by selectively emptying a single site near the centre of the array using a focused addressing laser pulse. The laser is frequency stabilized to the $F = 2 \rightarrow F' = 2$ optical pumping transition of the D2 line, which has a high probability of changing the spin of the atoms to an untrapped Zeeman state (0.5 per scattered
Figure 5. Single-site addressability and transport of an empty site. Sequence of images demonstrating single-site addressability using a focused laser pulse to empty a selected trap and transport using the atomic shift register. (a) Atoms are loaded to the lattice at a temperature of \( \sim 10 \mu \text{K} \). The center of the red circle indicates the position of the focused addressing laser spot. (b) Optical pumping light focused on one lattice site is pulsed on for 1 ms to drive transitions to untrapped Zeeman states, completely emptying the microtrap. The effect of the addressing pulse on neighbouring traps is negligible. (c) All the traps along with the emptied site are shifted by three lattice periods to a new position (bottom right). (d) The addressing laser is pulsed on again to empty a second trap, and finally (e) the array is shifted back to the original position.

The addressing laser is applied by imaging of the output facet of a single-mode optical fibre to the atom chip. We make double use of the optical imaging system to focus the light: positioning the fibre in a second image plane (at the same distance as the CCD camera from the chip) created with a beamsplitter. We estimate a focused spot corresponding to a \( \sim 7 \mu \text{m} \) Airy disc radius at the atom chip, which is highly homogeneous over the extent of a single cloud but is small compared with the distance between neighbouring traps. The spot is then easily aligned on any single array site by controlling the fibre position with an \( x-y \) translation stage and imaging the back-reflected light on the CCD camera for position reference.

Figure 5(a) shows part of an absorption image taken of the array after an evaporative cooling sweep to \( T \approx 10 \mu \text{K} \). The focused addressing laser is aligned on the site at the centre of the image as indicated by the red circle. The addressing laser is then pulsed on for 1 ms to completely empty this trap of atoms (figure 5(b)). We do not measure any atom loss or heating caused by the addressing pulse on neighbouring atom clouds, even for pulse durations as long as 10 ms, indicating excess light incident on neighbouring traps is negligible. We then demonstrate how optical addressing can be combined with the atomic shift register in order to write patterns to the array. Figure 5(c) shows the lattice after emptying the central microtrap and shifting the lattice by three periods (lower right). Subsequently, we can pulse on the addressing laser to empty another trap (figure 5(d)). Finally, the lattice is shifted again three periods in the reverse direction back to the original position. The addressed traps remain empty for the
full duration of the experiment of 126 ms. The same techniques could be used to create and readout spin coherence between quantum states of atoms in a single trap. These experiments clearly demonstrate local manipulation of individual atom clouds in our array and point to the possibility of encoding and shifting quantum information. It is also possible to use focused resonant imaging light for the addressing laser, to detect atoms in one trap without affecting neighbouring traps of the array. Focused lasers are particularly powerful for addressing two-dimensional arrays, especially for our system where atoms are confined to regions much smaller than the distance between traps. Straightforward modifications to our set-up using acousto-optical or spatial light modulators to control the position of the addressing laser will allow for sequential or simultaneous writing to the quantum state of atoms trapped in each microtrap.

We expect this system to be the ideal platform for studying controlled many-particle entanglement and quantum information processing with neutral atoms on an atom chip. Specifically, qubits could be encoded in two magnetically trappable states $|F = 1, m_F = -1\rangle$ and $|F = 2, m_F = +1\rangle$ of the $^{87}$Rb electronic ground state. With focused Raman lasers, it is possible to coherently drive transitions between logical states in targeted traps. Focused lasers could also be used to state-selectively drive atoms to highly excited Rydberg states. In these small traps, the dipole blockade mechanism [22, 39] would limit the process to a single Rydberg excitation per trap and could mediate interactions on demand over distances of tens of micrometres, between neighbouring traps [40, 41]. If required, single-atom detection sensitivity could be achieved through subsequent field ionization of the excited atoms. Alternatively, entangled mesoscopic ensembles of atoms could be produced and used for quantum information processing using Rydberg excitations [39, 42]. Future studies will be aimed at investigating the effect of magnetic and electric field noise near the chip surface on the lifetime and phase coherence of ground state and excited atoms.

6. Conclusion

To conclude, we have reported experiments on a two-dimensional array of ultracold atom clouds prepared near the surface of a magnetic-film atom chip. This promising new architecture for scalable quantum information processing combines optically resolved trap separations with tight magnetic confinement. We have loaded atoms into more than 500 array sites and performed rf evaporative cooling of these atom clouds to the quantum degeneracy temperature. In the process, we have demonstrated preparation, cooling, detection and analysis of hundreds of atom clouds in parallel; highlighting magnetic microtrap arrays as a unique experimental test bed for studying ultracold atomic systems and quantum matter. Using existing microfabrication technology we anticipate further scaling, to lattice dimensions of a few micrometres, in order to accommodate thousands of individual atomic qubits. An atomic shift register is used for parallel transport of atoms across the chip surface, allowing the separation of storage, interaction and readout areas of a future, fully integrated device. Finally, we demonstrate local manipulation of trapped atoms by selectively emptying individual traps using a focused optical pumping laser.

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