COLD ATOMS

Atom-by-atom assembly of defect-free one-dimensional cold atom arrays

Manuel Endres, 1,2* Hannes Bernien, 1* Alexander Keesling, 1* Harry Levine, 1* Eric R. Anschuetz, Alexandre Krajenbrink, Crystal Senko, Vladan Vuletic, Markus Greiner, Mikhail D. Lukin 1

The realization of large-scale fully controllable quantum systems is an exciting frontier in modern physical science. We use atom-by-atom assembly to implement a platform for the deterministic preparation of regular one-dimensional arrays of individually controlled cold atoms. In our approach, a measurement and feedback procedure eliminates the entropy associated with probabilistic trap occupation and results in defect-free arrays of more than 50 atoms in less than 400 milliseconds. The technique is based on fast, real-time control of 100 optical tweezers, which we use to arrange atoms in desired geometric patterns and to maintain these configurations by replacing lost atoms with surplus atoms from a reservoir. This bottom-up approach may enable controlled engineering of scalable many-body systems for quantum information processing, quantum simulations, and precision measurements.

he detection and manipulation of individual quantum particles, such as atoms or photons, is now routinely performed in many quantum physics experiments (1, 2); however, retaining the same control in largescale systems remains an outstanding challenge. For example, major efforts are currently aimed at scaling up ion-trap and superconducting platforms, where high-fidelity quantum computing operations have been demonstrated in registers consisting of several qubits (3, 4). In contrast, ultracold quantum gases composed of neutral atoms offer inherently large system sizes. However, arbitrary single-atom control is highly demanding, and its realization is further limited by the slow evaporative cooling process necessary to reach quantum degeneracy. Only in recent years has individual particle detection (5, 6) and basic singlespin control (7) been demonstrated in low-entropy optical lattice systems.

Here, we demonstrate atom-by-atom assembly of large defect-free one-dimensional (1D) arrays of cold neutral atoms (8, 9).

We use optical microtraps to directly extract individual atoms from a laser-cooled cloud (10-12) and employ recently demonstrated trapping techniques (13-16) and single-atom position control (17-20) to create desired atomic configurations. Central to our approach is the use of single-atom detection and real-time feedback (17, 20) to eliminate the entropy associated with the probabilistic trap loading (10) [currently limited to 90% loading probability even with advanced techniques

"Maxwell's demon" (8, 9), this method allows us to rapidly create large defect-free arrays and, when supplemented with appropriate atom-atom interactions (15, 16, 24-30), provides a potential platform for scalable experiments with individually controlled neutral atoms.

(21-23)]. Related to the fundamental concept of

The experimental protocol is illustrated in Fig. 1A. An array of 100 tightly focused optical tweezers is loaded from a laser-cooled cloud. The collisional blockade effect ensures that each individual tweezer is either empty or occupied by a single atom (10). A first high-resolution image vields single-atom-resolved information about the trap occupation, which we use to identify empty traps and to switch them off. The remaining occupied traps are rearranged into a regular, defect-free array, and we detect the final atom configuration with a second high-resolution image.

Our implementation relies on fast, real-time control of the tweezer positions (Fig. 1B), which we achieve by employing an acousto-optic deflector (AOD) that we drive with a multitone radiofrequency (RF) signal.

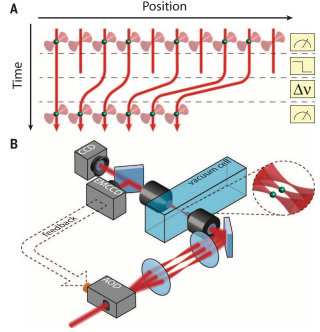
This generates an array of deflected beams, each controlled by its own RF tone (15, 16). The resulting beam array is then focused into our vacuum chamber and forms an array of optical tweezers, each with a Gaussian waist of ≈ 900 nm, a wavelength of 809 nm, and a trap depth of $U/k_B \approx 0.9$ mK [Boltzmann constant (k_B)] that is homogeneous across the array within 2% (31).

The tweezer array is loaded from a laser-cooled cloud of Rubidium-87 atoms in a magneto-optical trap (MOT). After the loading procedure, we let the MOT cloud disperse and we detect the occupation of the tweezers with fluorescence imaging. Fast, single-shot, single-atom-resolved detection with 20-ms exposure is enabled by a sub-Doppler laser-cooling configuration that remains active during the remainder of the sequence (31) (see Fig. 2, A to C). Our fluorescence count statistics show that individual traps are either empty or occupied by a single atom (10, 31), and we find probabilistically filled arrays with an average single-atom loading probability of $p \approx 0.6$ (see Figs. 2A and 3A).

The central part of our scheme involves the rearrangement procedure for assembling defectfree arrays (31) (see Fig. 1A). In the first step,

Fig. 1. Protocol for creating defect-free arrays. (A) A first image identifies optical microtraps loaded with a single atom, and empty traps are turned off. The loaded traps are moved to fill in the empty sites, and a second image verifies the success of the operation.

(B) The trap array is produced by an acousto-optic deflector (AOD) and imaged with a 1:1 telescope onto a 0.5-NA microscope objective, which creates an array of tightly focused optical tweezers in a vacuum chamber. An identical microscope objective is aligned to image the same focal plane. A dichroic mirror allows us to view the trap light on a charge-coupled-device camera (CCD) while simultaneously detecting the atoms via fluorescence imaging on an electron-multiplied-CCD



02138, USA. ²Division of Physics, Mathematics, and Astronomy, California Institute of Technology, Pasadena, CA 91125, USA. ³Department of Physics and Research Laboratory of Electronics, Massachusetts Institute of Technology, Cambridge, MA 02139, USA.

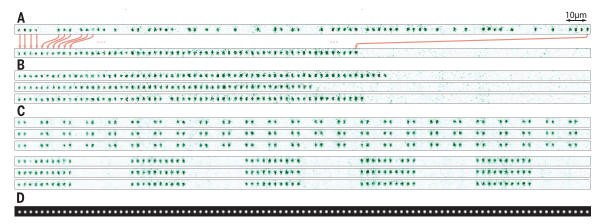
¹Department of Physics, Harvard University, Cambridge, MA

*These authors contributed equally to this work. †Corresponding Laboratoire de Physique Theorique de l'Ecole Normale Superieure, 24 Rue L'Homond, 75231 Paris Cedex, France,

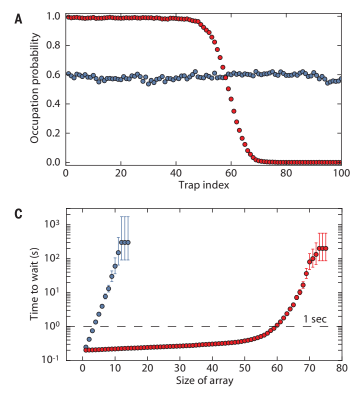
camera (EMCCD). The rearrangement protocol is realized through fast feedback onto the multitone radio-frequency (RF) field driving the AOD.

Fig. 2. Assembly of regular atom arrays.

(A) Single-shot, singleatom-resolved fluorescence images recorded with the EMCCD before (top) and after (bottom) rearrangement. Defects are identified, and the loaded traps are rearranged according to the protocol in Fig. 1, indicated by arrows for a few selected atoms. (B) Two instances of successfully rear-



ranged arrays (first two pictures), and one instance where a defect is visible after rearrangement (last picture). (C) The final arrangement of atoms is flexible, and we generate, e.g., clusters of 2 (top) or 10 (bottom) atoms. Nonperiodic arrangements and adjustable lattice spacings are also possible. (**D**) High-resolution CCD image of trap array. Our default configuration for loading atoms consists of an array of 100 tweezers with a spacing of 0.49 MHz between the RF tones, corresponding to a real-space distance of 2.6 µm between the focused beams (31).



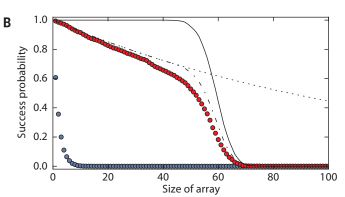


Fig. 3. Quantifying the rearrangement performance. (A) The initial loading (blue circles) results in an occupation probability of ≈ 0.6 for each trap in the array. After rearrangement (red circles), close to unity filling is reached on the left side of the array. (B) In the initial image, the probability of finding a defect-free length-N array (starting from the leftmost trap) falls off exponentially with N (blue circles). After the rearrangement of all loaded traps to form the largest possible array, we demonstrate strongly enhanced success probabilities at producing defect-free arrays (red circles). Theory curves show limits set by the total initial atom number (solid line), the backgroundlimited lifetime of τ = 6.2 s (dotted line), and the product of both (dashed dotted line) (31). (C) Expected amount of time to wait, on average, to produce a defect-free array of a given size, taking into account the experimental cycle time of 200 ms (150 ms without rearrangement). Without

rearrangement, the wait time grows exponentially (blue circles). Employing the rearrangement procedure, we can produce arrays of length 50 in less than 400 ms (red circles). Error bars denote 68% confidence intervals, which are smaller than the marker size in (A) and (B).

unoccupied traps are switched off by setting the corresponding RF amplitudes to zero. In a second step, all occupied tweezers are moved to the left until they stack up with the original spacing of 2.6 µm. This movement is generated by sweeping the RF tones to change the deflection angles of the AOD and lasts 3 ms (31). Finally, we detect the resulting atom configuration with a second high-resolution image. These steps imple-

ment a reduction of entropy via measurement and feedback. The effect is immediately visible in the images shown in Fig. 2, A and B. The initial filling of our array is probabilistic, whereas the rearranged configurations show highly ordered atom arrays. Our approach also allows us to construct flexible atomic patterns (Fig. 2C).

The rearrangement procedure creates defect-free arrays with high fidelity. This can be quantified by considering the improvement of single-atom occupation probabilities (Fig. 3A) and the success probabilities, p_N , for creating defect-free arrays of length N (Fig. 3B). The single-atom occupation probability in the left-most 40 traps increases from ≈ 0.6 before rearrangement to 0.988(3) after rearrangement, demonstrating our ability for high-fidelity single-atom preparation. Furthermore, the success probabilities for creating defect-

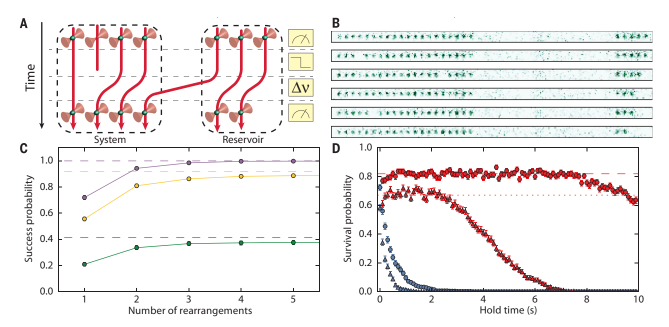


Fig. 4. Creating and maintaining regular arrays using an atomic reservoir. (**A**) For a given target array size, surplus atoms are kept in a reservoir and used for repetitive reloading of the array. (**B**) A 20-atom target array with a reservoir of atoms on the right. Defects occasionally develop in the target array and are replaced by atoms in the reservoir. The reservoir depletes as it is used to fill in defects. (**C**) By performing repeated rearrangements (once every 50 ms), the probability to successfully produce a defect-free array in any of these attempts increases and approaches the limit set by the number of initially loaded atoms (dashed lines). We show data for targeting 40 (purple), 50 (yellow), and 60 (green) atom arrays. Solid lines are guides to

the eye. (**D**) Probing for defects and filling them once every 100 ms from the reservoir extends the lifetime of a defect-free array. Shown is the success probability of maintaining arrays of 20 (circles) and 40 (triangles) atoms with (red) and without (blue) replenishing atoms from the reservoir. With replenishing, the probability to maintain a defect-free array remains at a fixed plateau for as long as we have surplus atoms in the reservoir. The initial plateau value is set by the probability that no atoms in the array are lost in 100 ms (calculated value for 10-s single-atom lifetime shown as the dotted line). Error bars denote 68% confidence intervals, which are smaller than the marker size in (C).

free arrays show an exponential improvement. Before rearrangement, the probability of finding a defect-free array of length N is exponentially suppressed with $p_N = p^N$, where $p \approx 0.6$ (blue circles, Fig. 3B). After rearrangement, we find success probabilities as high as $p_{30} = 0.75(1)$ and $p_{50} = 0.53(1)$ (red circles, Fig. 3B).

The same exponential improvement is observed by considering the average wait time for producing defect-free arrays, given by T/p_N , where T=200 ms is the cycling time of our experiment (see Fig. 3B). For example, we are able to generate defect-free arrays of 50 atoms with an average wait time of less than 400 ms (red circles, Fig. 3C).

The success probabilities can be further enhanced through multiple repetitions of the rearrangement protocol. Figure 4 illustrates the procedure in which we target an atomic array of fixed length and create a reservoir from surplus atoms in a separate zone. After the initial arrangement of atoms into the target and reservoir zones, we periodically take images to identify defects in the target array and pull atoms from the reservoir to fill in these defects. This enhances our initial success probabilities at producing defect-free arrays within one MOT-loading cycle to nearly the ideal limit (Fig. 4C).

Finally, a similar procedure can be used for correcting errors associated with atomic loss. This becomes a substantial limitation for large

arrays because for a given lifetime of an individual atom in the trap τ , the corresponding lifetime of the N atom array scales as τ/N . To counter this loss, we repeatedly detect the array occupation after longer time intervals and replenish lost atoms from the reservoir. This procedure leads to exponentially enhanced lifetimes of our arrays (Fig. 4D).

These results demonstrate the ability to generate and control large, defect-free arrays at a fast repetition rate. The success probabilities are limited by two factors: the initial number of loaded atoms and losses during rearrangement. For example, the average total atom number in our array is 59 ± 5 (31), which results in the cutoff in the success probability in Fig. 3B starting from $N \approx 50$ (solid line). For shorter arrays, the fidelity is mostly limited by losses during rearrangement. These losses are dominated by our finite vacuum-limited lifetime, which varies from $\tau \approx 6$ s to $\tau \approx 12$ s (depending on the setting of our atomic dispenser source), and are only minimally increased by the movement of the atoms (31). The single-atom occupation probability is correspondingly reduced by a factor $\exp(-t_r/\tau)$, where $t_r = 50$ ms is the time for the whole rearrangement procedure (31). This results in the success probabilities of creating length-N arrays scaling as $\exp(-t_r N/\tau)$, which dominates the slope for N = 50 in Fig. 3B (dotted line). Currently, we reach vacuum-limited lifetimes only with sub-Doppler cooling applied throughout the sequence. However, the lifetime without cooling could be improved—for example, by using a different trapping laser and trapping wavelength (31).

The size of the final arrays can be considerably increased by implementing a number of realistic experimental improvements. For example, the initial loading probability could be enhanced to 0.9 (21-23) and the vacuum-limited lifetime could be improved to $\tau \approx 60$ s in an upgraded vacuum chamber. Increasing the number of traps in the current configuration is difficult because of the AOD bandwidth of ≈ 50 MHz and strong parametric heating introduced when the frequency spacing of neighboring traps approaches ≈ 450 kHz (31). However, implementing 2D arrays could provide a path toward realizing thousands of traps, ultimately limited by the availability of laser power and the field of view of high-resolution objectives. Such 2D configurations could be realized by either directly using a 2D-AOD or by creating a static 2D lattice of traps [using spatial light modulators (14) or optical lattices (12)] and sorting atoms with an independent AOD (31). With increased loading efficiencies (21-23), realistic estimates for the rearrangement time t_r in such 2D arrays indicate that the robust creation of defect-free arrays of hundreds of atoms is feasible (31). Finally, the repetitive interrogation techniques,

in combination with periodic reservoir reloading from a cold atom source (such as a MOT), could be used to maintain arrays indefinitely.

Atom-by-atom assembly of defect-free arrays forms a scalable platform with unique possibilities. It combines features that are typically associated with ion-trapping experiments, such as single-qubit addressability (32, 33) and fast cycling times, with the flexible optical trapping of neutral atoms in a scalable fashion. Furthermore, in contrast to solid-state platforms, such atomic arrays are highly homogeneous (31) and mostly decoupled from their environment. The homogeneity of our array should also allow for cooling of the atomic motion via simultaneous sideband cooling in all tweezers at once (34, 35).

These features provide an excellent starting point for multiqubit experiments, for studies of quantum many-body effects, and for exploring future applications. The required interactions between the atoms can be engineered using several approaches. Even without sideband cooling, exciting the atoms into high-lying Rydberg states would introduce strong dipole interactions that can be used for fast entangling gates (24, 25, 27). The parallelism afforded by our flexible atom rearrangement enables efficient diagnostics of such Rydberg-mediated entanglement. These interactions may also enable approaches to quantum simulations that involve both coherent coupling and engineered dissipation (26, 27), as well as large-scale entangled quantum states for applications in precision measurements (36).

An alternative approach to engineering interactions involves the integration of atom arrays with nanophotonic platforms as demonstrated previously (28, 29). These enable photonmediated interactions that can be employed to couple the atoms within a local multiqubit register or for efficient communication between the registers using a modular quantum network architecture (3).

Finally, our platform could enable new bottomup approaches to studying quantum many-body physics in Hubbard models (15, 16, 30), where atomic Mott insulators with fixed atom number and complex spin patterns could be directly assembled. This requires atom temperatures close to the ground state, coherent tunneling between the traps, and sizable on-site interactions. With side-band cooling, ground-state fractions in excess of 90% have already been demonstrated (34, 35) and can likely be improved via additional optical trapping along the longitudinal tweezer axes, which would also increase on-site interaction strengths. Coherent tunneling of Rb atoms between similarly sized tweezers has been observed before by reducing the tweezer distance (15, 16). The parametric heating, currently limiting the minimal distance between our traps, could be reduced by working with shallower traps, as needed for tunneling, and by employing fewer traps to increase the frequency separation between neighboring traps. Eventually, this approach could be applied to create ultracold quantum matter composed of exotic atomic species or complex molecules (37, 38) that are difficult to cool evaporatively.

REFERENCES AND NOTES

- 1. S. Haroche, Ann. Phys. 525, 753-776 (2013).
- D. J. Wineland, Rev. Mod. Phys. 85, 1103-1114 (2013).
- C. Monroe, J. Kim, Science 339, 1164-1169 (2013).
- M. H. Devoret, R. J. Schoelkopf, Science 339, 1169-1174 (2013).
- W. S. Bakr et al., Science 329, 547-550 (2010). 6. J. F. Sherson et al., Nature 467, 68-72 (2010).
- C. Weitenberg et al., Nature 471, 319-324 (2011).
- D. S. Weiss et al., Phys. Rev. A 70, 040302 (2004).
- J. Vala et al., Phys. Rev. A 71, 032324 (2005)
- 10. N. Schlosser, G. Reymond, I. Protsenko, P. Grangier, Nature 411, 1024-1027 (2001).
- 11. M. Weber, J. Volz, K. Saucke, C. Kurtsiefer, H. Weinfurter, Phys. Rev. A 73, 043406 (2006).
- 12. K. D. Nelson, X. Li, D. S. Weiss, Nat. Phys. 3, 556-560 (2007).
- 13. M. J. Piotrowicz et al., Phys. Rev. A 88, 013420 (2013).
- 14. F. Nogrette et al., Phys. Rev. X 4, 021034 (2014).
- 15. A. M. Kaufman et al., Science 345, 306-309 (2014).
- 16. A. M. Kaufman et al., Nature 527, 208-211 (2015).
- 17. Y. Miroshnychenko et al., Nature 442, 151 (2006).
- 18. J. Beugnon et al., Nat. Phys. 3, 696-699 (2007) 19. M. Schlosser et al., New J. Phys. 14, 123034 (2012).
- 20. H. Kim, et al., Nat. Commun. 7, 13317 (2016). 21. T. Grünzweig, A. Hilliard, M. McGovern, M. F. Andersen,
- Nat. Phys. 6, 951-954 (2010).
- 22. B. J. Lester, N. Luick, A. M. Kaufman, C. M. Reynolds, C. A. Regal, Phys. Rev. Lett. 115, 073003 (2015).
- 23. Y. H. Fung, M. F. Andersen, New J. Phys. 17, 073011 (2015).
- 24. D. Jaksch et al., Phys. Rev. Lett. 85, 2208-2211 (2000).
- 25. M. Saffman, T. G. Walker, K. Mølmer, Rev. Mod. Phys. 82, 2313-2363 (2010).
- 26. H. Weimer, M. Müller, I. Lesanovsky, P. Zoller, H. P. Büchler, Nat. Phys. 6, 382-388 (2010).
- 27. A. Browaeys, D. Barredo, T. Lahaye, J. Phys. B 49, 152001
- 28. J. D. Thompson et al., Science 340, 1202-1205 (2013).

- 29. A. Goban et al., Nat. Commun. 5, 3808 (2014).
- 30. S. Murmann et al., Phys. Rev. Lett. 115, 215301 (2015).
- 31. See supplementary materials on Science Online.
- 32. T. Xia et al., Phys. Rev. Lett. 114, 100503 (2015).
- 33. Y. Wang, X. Zhang, T. A. Corcovilos, A. Kumar, D. S. Weiss, Phys. Rev. Lett. 115, 043003 (2015).
- 34. A. M. Kaufman, B. J. Lester, C. A. Regal, Phys. Rev. X 2, 041014 (2012)
- 35. J. D. Thompson, T. G. Tiecke, A. S. Zibrov, V. Vuletić, M. D. Lukin, Phys. Rev. Lett. 110, 133001 (2013).
- 36. P. Kómár et al., Phys. Rev. Lett. 117, 060506 (2016).
- 37. J. F. Barry, D. J. McCarron, E. B. Norrgard, M. H. Steinecker,
- D. DeMille, Nature 512, 286-289 (2014).
- 38. N. R. Hutzler, L. R. Liu. Y. Yu. K.-K. Ni. https://arxiv.org/abs/1605.09422 (2016).
- 39. D. Barredo, S. de Léséleuc, V. Lienhard, T. Lahaye,
 - A. Browaeys, Science 354, 1021-1023 (2016).

ACKNOWLEDGMENTS

We thank K.-K. Ni, N. Hutzler, A. Mazurenko, and A. Kaufman for insightful discussion. This work was supported by NSF, Center for Ultracold Atoms, National Security Science and Engineering Faculty Fellowship, and Harvard Quantum Optics Center. H.B. acknowledges support by a Rubicon Grant of the Netherlands Organization for Scientific Research (NWO). During the completion of this work, we became aware of a related approach (39),

SUPPLEMENTARY MATERIALS

www.sciencemag.org/content/354/6315/1024/suppl/DC1 Materials and Methods Figs. S1 to S5 Movies S1 to S3 References (40, 41)

17 June 2016; accepted 17 October 2016 Published online 3 November 2016 10.1126/science.aah3752

GEOPHYSICS

Mega-earthquakes rupture flat megathrusts

Quentin Bletery, 1x Amanda M. Thomas, 1 Alan W. Rempel, 1 Leif Karlstrom, 1 Anthony Sladen,2 Louis De Barros2

The 2004 Sumatra-Andaman and 2011 Tohoku-Oki earthquakes highlighted gaps in our understanding of mega-earthquake rupture processes and the factors controlling their global distribution: A fast convergence rate and young buoyant lithosphere are not required to produce mega-earthquakes. We calculated the curvature along the major subduction zones of the world, showing that mega-earthquakes preferentially rupture flat (low-curvature) interfaces. A simplified analytic model demonstrates that heterogeneity in shear strength increases with curvature. Shear strength on flat megathrusts is more homogeneous, and hence more likely to be exceeded simultaneously over large areas, than on highly curved faults.

ast mega-earthquakes, such as the magnitude (M) 9.6 Chile earthquake in 1960 and the M 9.3 Alaska earthquake in 1964, occurred in areas where the subducting lithosphere was relatively young (and buoyant) and the plate convergence rate was relatively high (I). These observations led some authors to hypothesize that maximum earthquake

¹Department of Earth Sciences, University of Oregon, 1272 University of Oregon, Eugene, OR 97403, USA. ²Université Côte d'Azur, CNRS, OCA, IRD, Géoazur, 250 rue Albert Einstein, Sophia Antipolis, 06560 Valbonne, France. *Corresponding author. Email: gbletery@uoregon.edu

size is controlled by these two geological parameters (2, 3). The development of space-based geodesy enabled refined measurements of plate motion that challenged the role of convergence rate (4-6). Additionally, the moment magnitude $(M_{\rm w})$ 9.0 Tohoku-Oki earthquake (7) ruptured lithosphere that is over 120 million years old (8), ruling out lithospheric age as the dominant control. Weak correlations appear in recent data sets among a variety of parameters, including forearc structure (9, 10); age, density, and buoyancy of the slab (6); upper plate motion (11); upper plate strain (12); long-term trench migration (11); trench sediment thickness (12); and width of the seismogenic