Probing the Superfluid to Mott Insulator Transition at the Single Atom Level


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Quantum gases in optical lattices offer an opportunity to experimentally realize and explore condensed matter models in a clean, tunable system. We investigate the Bose-Hubbard model on a microscopic level using single atom–single lattice site imaging; our technique enables space- and time-resolved characterization of the number statistics across the superfluid–Mott insulator quantum phase transition. Site-resolved probing of fluctuations provides us with a sensitive local thermometer, allows us to identify microscopic heterostructures of low entropy Mott domains, and enables us to measure local quantum dynamics, revealing surprisingly fast transition timescales. Our results may serve as a benchmark for theoretical studies of quantum dynamics, and may guide the engineering of low entropy phases in a lattice.

Microscopic measurements can reveal properties of complex systems that are not accessible through statistical ensemble measurements. For example, scanning tunneling microscopy has allowed physicists to identify the importance of nanoscale spatial inhomogeneities in high temperature superconductivity [1], and single molecule microscopy [2] has enabled studies of local dynamics in chemical reactions revealing e.g. the importance of multiple reaction pathways [3]. While previous ultracold quantum gas experiments have focused primarily on statistical ensemble measurements, the recently introduced single atom-single lattice site imaging technique in a Quantum Gas Microscope (QGM) [4] opens the door for probing and controlling quantum gases on a microscopic level.

Here we present a microscopic study of an atom-lattice system that realizes the bosonic Hubbard model and exhibits a quantum phase transition from a superfluid to a Mott insulator [5–7]. In the weakly interacting superfluid regime, the many-body wavefunction factorizes into a product of states with well-defined phase on each lattice site, known as coherent states, with Poissonian number fluctuations. As the strength of the interaction increases, the number distribution is narrowed, resulting in a fixed atom number state on each site deep in the Mott insulator regime. We study this change in the number statistics across the transition; these microscopic studies are complementary to previous experiments that have focused on measuring ensemble properties such as long range phase coherence, excitation spectra or compressibility [7–9]. Local properties such as onsite number statistics [10] were accessible only indirectly [8, 11, 12] and averaged over several shells of superfluid and Mott insulating domains in the inhomogeneous system, complicating quantitative interpretation. More recently, the shell structure was imaged through tomographic [13], spectroscopic [14], and in-situ imaging techniques, coarse-grained over several lattice sites [15].

We start with a two-dimensional $^{87}$Rb Bose-Einstein condensate of a few thousand atoms confined in a single well of a standing wave, with a harmonic oscillator length of 130nm [16]. The condensate resides 9µm from an in-vacuum lens that is part of an imaging system with a resolution of $\sim 600$nm. This high resolution system is used to project a square lattice potential onto the pancake cloud with a periodicity of $a = 680$nm, as described in previous work [4]. The lattice depth is ramped exponentially with a time constant of 81ms up to a maximum depth of $16E_r$, where $E_r$ is the recoil energy of the effective lattice wavelength given by $\hbar^2/8ma^2$, with $m$ being the mass of $^{87}$Rb and Planck’s constant $\hbar$.

![FIG. 1. Single-site imaging of atom number fluctuations across the superfluid-Mott insulator transition.](image)
a homogeneous system in two dimensions, the transition to a Mott insulator with one atom per site occurs at a ratio of interaction energy to tunneling of $U/J = 16.7^{17}$, corresponding to a lattice depth of $12.2E_r$. During this ramp, the initial transverse confinement of $9.5\text{Hz}$ is increased such that the cloud size remains approximately constant. After preparing the many-body state, we image the atoms by increasing the lattice depth several hundred-fold, and then illuminate the atoms with an optical molasses that serves to localize the atoms while fluorescence photons are collected by the high resolution optics. As a result of the imaging process, the many-body wavefunction is projected onto number states on the lattice. In addition, light-assisted collisions immediately eject atoms in pairs from each lattice site, leaving behind an atom on a site only if its initial occupation was odd $^{20}$. Remaining atoms scatter several thousand photons during the exposure time and can be detected with high fidelity. By preparing the sample repeatedly under the same conditions, we deduce the probability $p_{\text{odd}}$ of having an odd number of atoms on a site before the measurement.

For a coherent state on a lattice site with mean atom number $\lambda$, $p_{\text{odd}}$ is given by $1/2(1 - e^{-2\lambda}) < 1/2$. In a Mott-insulating region in the zero temperature and zero tunneling limit, $p_{\text{odd}} = 1 \ (0)$ for shells with an odd (even) atom number per site. Fig. 1 shows fluorescence images in a region of the cloud as the final depth of the lattice is increased. The initial superfluid density is chosen to obtain an insulator with two shells on the Mott side of the transition, and the region shown is in the outer shell containing one atom per site. For high filling fractions, the lattice sites in the images are barely resolved, but the known geometry of the lattice and imaging system point spread function obtained from images at sparser fillings allow reliable extraction of site occupations $^{16}$.

We determine $p_{\text{odd}}$ for each site using 24 images at each final lattice depth. The transverse confining potential varies slowly compared to the lattice spacing and the system is to good approximation locally homogeneous. We make use of this to improve the error in our determination of $p_{\text{odd}}$, by averaging over a group of lattice sites, in this case 51 (30) sites for regions in the first (second) shell (Fig. 2). In the $n = 1$ shell, we detect an atom on a site with probability $94.9 \pm 0.7\%$ at a lattice depth of $16E_r$. We measure the lifetime of the gas in the imaging lattice and determine that $1.75 \pm 0.02\%$ of the occupied sites are detected as unoccupied due to atoms lost during the imaging exposure time of 1s because of background gas collisions. We correct for this effect in the given average occupation numbers and errorbars.

Measuring the defect density in the Mott insulator provides sensitive local thermometry deep in the Mott regime. Thermometry in the Mott state has been a long-standing experimental challenge $^{21, 22}$ and has acquired particular significance as experiments approach the regime of quantum magnetism $^{23, 24}$, where the temperature scale should be on the order of the superexchange interaction energy. We directly image excitations of the $n = 1$ Mott insulator, holes and doublons, as they both appear as missing atoms in the images. Similarly, for Mott insulators with higher fillings $n$, sites with excitations $(n + 1, n - 1)$ can be detected through their opposite parity signal. For finite tunneling rate $J$ much smaller than the interaction energy $U$, the admixture fraction of coherent hole-doublon pairs excitations is $\sim (J/U)^2$, whereas any other excitations are due to incoherent thermal fluctuations and are suppressed by a Boltzmann factor $e^{-U/T}$.

The theory curves presented in Fig. 2 are the predicted $p_{\text{odd}}$ in the two shells for different values of $T/U$. The curves are obtained using a quantum Monte-Carlo “worm” algorithm $^{26, 27}$, and the average temperature extracted using the data points at the three highest $U/J$ ratios is $T/U \sim 0.16 \pm 0.03$. At the transition point for $n = 1$, this corresponds to a temperature of 1.8nK. Assuming this value of $T/U$ to be the overall temperature, the thin layer between the Mott shells should be superfluid, and the transition to a normal gas is expected around a critical temperature of $zJ = 2.8nK$, where $z$ is the number of nearest neighbors in the lattice $^{28}$.

Next we study the global structure of the Mott insulator. The high resolution images provide an atom-by-atom picture of the concentric shell structure, including the transition layers in between the insulating shells. In Fig. 3A to D, the formation of the various shells, up to

![FIG. 2. Measured value of $p_{\text{odd}}$ vs. the interaction to tunneling ratio $(U/J)$. Data sets, with statistical error bars, are shown for regions that form part of the $n = 1$ (squares) and $n = 2$ (circles) Mott shells in a deep lattice. The lines are based on finite temperature Monte-Carlo simulations in a homogeneous system at constant temperature to interaction ratio $(T/U)$ of 0.20 (dotted red), 0.15 (solid black) and 0.05 (dashed blue). The axis on the right is the corresponding odd-even variance given by $p_{\text{odd}}(1 - p_{\text{odd}})$.](attachment:image.png)
FIG. 3. Single-site imaging of the shell structure in a Mott insulator. (A – D) The images show \( p_{\text{odd}} \) on each site determined by averaging 20 analyzed fluorescence images. The lattice depth is \( 22E_r \) and the transverse confinement is 45Hz. As the atom number is increased from (A) to (D), the number of shells in the insulator increases from one to four. The value of \( p_{\text{odd}} \) for odd (even) numbered shells is close to one (zero). The atom numbers, determined by in-situ imaging of clouds expanded in the plane, are (A), 120 \( \pm \) 10, (B), 460 \( \pm \) 20, (C), 870 \( \pm \) 40 and (D), 1350 \( \pm \) 70. (E – F) Long wavelength disorder can be corrected by projecting an appropriate compensation light pattern onto the atoms, resulting in nearly circular shells. (E) shows \( p_{\text{odd}} \) (average of 20 analyzed images) and (F) is a single shot raw image (arbitrary units).

the fourth, is shown as the atom number in the trap is increased. Slowly varying optical potential disorder causes deviation from circular symmetry in the shells. The contour lines of the potential are directly seen in the images in Fig. 3. In Fig. 3E and F, we have compensated this disorder by projecting a light pattern generated using a digital micromirror device through the objective, resulting in a nearly circular shell structure.

In a second series of experiments, we use on-site number statistics to probe the adiabaticity timescale for the transition, focusing on the local dynamics responsible for the narrowing of the number distribution. We start by increasing the lattice depth adiabatically to \( 11E_r \), still in the superfluid regime, using the same ramp described previously. Next the depth is ramped linearly to \( 16E_r \) where, for an adiabatic ramp, a Mott insulator should form. The ramp time is varied from 0.2ms to 20ms, and \( p_{\text{odd}} \) is measured in the first and second shells as before (Fig. 4); we find that the data fits well to exponential curves that asymptote to the value of \( p_{\text{odd}} \) obtained in the adiabatic case. The fitted time constant in the first (second) shell is \( 3.5 \pm 0.5 \text{ms} \) (3.9 \( \pm \) 1.3ms).

Compared to the critical value of the tunneling time \( h/J_e = 68\text{ms} \) for the first shell, the observed dynamics are counter-intuitively fast. This can be understood using a simple picture of two atoms in a double well. In this system, as the tunneling is varied, the minimal gap between the ground state and the first excited state is \( U \), which sets the adiabaticity timescale. It is an open question whether this argument can be generalized to a lattice. In an infinite system, the appearance of Goldstone modes in the superfluid regime leads to a vanishing gap at the transition point, but the density of states is low for energies much less than \( U \). In fact, the \( 1/e \) timescale observed experimentally is comparable to \( h/U_c = 4.1\text{ms} \), where \( U_c \) is the critical interaction energy for an \( n = 1 \) insulator.

Although the local number statistics change on a fast timescale of \( h/J \), entropy redistribution in the inhomogeneous potential should occur on a much slower timescale of \( h/J \). Because superfluid and normal domains have a larger specific heat capacity than Mott domains, in an inhomogeneous system, entropy is expelled from the Mott domains and accumulates in the transition regions after.
A nearly pure Bose-Einstein condensate of $5 \times 10^4$ $^{87}\text{Rb}$ atoms is prepared in the $F = 1, m_f = -1$ state in a magnetic trap by radiofrequency (rf) evaporation. The Thomas-Fermi radii of the condensate are $(3.1, 3.1, 27)\mu m$. The condensate is transferred into a single well of a 1D standing wave with periodicity $9.2\mu m$ created by a beam reflected from the flat glass surface of an in-vacuum hemispheric lens. The light for this standing wave is centered at 755nm, has a 3nm spectral width and is incident at an angle of 2.3° relative to the surface. The condensate is loaded into the first nodal plane from the surface. The harmonic oscillator width of the condensate at full lattice depth along the direction perpendicular to the surface is 360nm. By increasing the bias field the confinement in the 2D plane is relaxed, resulting in an elliptic cloud with Thomas-Fermi radii $(18, 36)\mu m$ in the 2D plane.

In order to obtain a suitable initial density for creating a Mott insulator, the atom number in the 2D plane must be reduced to a few thousand atoms in a reproducible way. For this, a red-detuned (840nm) beam with an 8µm waist is focused through the objective onto the center of the pancake and creates a “dimple” potential in the magnetically confined cloud. The magnetic confinement is then removed and the number of atoms remaining in the dimple trap is proportional to its depth, with a residual RMS fluctuation of 6%. A second collinear 840nm beam with a 27µm waist is then turned on, and the dimple is adiabatically ramped down to expand the cloud into the larger beam. The transverse confinement of the condensate provided by this beam is 9.5Hz. The 840nm light source used for creating these beams has a spectral width of 12nm. The short coherence length eliminates unwanted interferences which would corrugate the confining potential.

The interaction between the atoms is then further enhanced by increasing the axial trapping frequency by a factor of eight. This is achieved by turning on a second 1D standing wave at an angle of 14° to the glass surface, with a lattice spacing of 1.54µm. The maximum axial trapping frequency is 7.1kHz. The condensate resides in the sixth well from the surface. At this point, the 9.2µm standing wave is ramped down. The surface provides a reproducible way to overlap the nodes of these two standing waves. In addition, the proximity of the atoms to the glass surface enhances the resolution of the imaging system by the index of refraction of glass, resulting in a measured point spread function (PSF) with full width at half maximum of $\sim 600$nm.

Materials and Methods

Preparation of the two-dimensional condensate

In addition to the number statistics studied in this work, single-site imaging could be applied to study spatial correlations in strongly correlated quantum gases, and to directly measure entanglement in a quantum information context. The low defect Mott states we detect would provide an ideal starting point for quantum magnetism experiments; if the low entropy in the Mott domains can be carried over to spin models, it should be possible to realize magnetically ordered states such as antiferromagnets, which could be directly detected with single-site imaging.
Preparation and imaging of the Mott insulator

To bring the cloud into a strongly-correlated regime, a two-dimensional square lattice with 680nm periodicity is created in the plane by projecting a mask through the objective onto the atoms, as described in [4]. The lattice light, like the light used for producing the 1D standing waves, is centered at 755nm and has a spectral width of 3nm to reduce disorder in the potential.

The lattice depth is increased linearly to 0.4$E_r$ in 50ms, and from there ramped exponentially to its final value ($16E_r$ for most experiments) with a time constant of 81ms. During the lattice ramp, the transverse confinement is increased so as to keep the size of the cloud constant, compensating for the increasing inter-atomic interaction and deconfinement due to the blue lattice. This allows for faster ramps while maintaining adiabaticity, because the density redistribution during the lattice ramp is minimized. Lattice depths are calibrated to 5% accuracy using Kapitza-Dirac scattering, and the tunneling matrix element and interaction energy at different depths are obtained from a band structure calculation. The dominating loss process in the lattice is three-body collisions in Mott shells with $n > 2$, as observed in other experiments [31]. The rate for such losses is $\gamma n(n-1)(n-2)$, with $\gamma = 2 \times 10^{-3}$Hz for a lattice depth of $22E_r$ [34]. Due to the relatively large lattice spacings, such losses are negligible in our lattice even for the fourth Mott shell ($< 1\%$).

For imaging the atoms, the same procedure as described in [3] is used. Briefly, the lattice depth is increased over three hundred-fold by changing the light source illuminating the mask to a monochromatic source detuned 50GHz to the blue of the D1 line. The frozen atom distribution, now projected onto number states in each well, is illuminated with a cooling molasses on the D2 line. Within the first 100$\mu$s, light assisted collisions eject atoms in pairs, leaving behind an atom only if the initial atom number on the site was odd. The remaining atoms are imaged in fluorescence by collecting the scattered molasses photons during a 1s exposure, resulting in $\sim 2,000$ photons registered by the camera for each atom.

Atom numbers are measured by switching off the transverse confinement and letting the cloud expand in the 2D plane for 5ms before turning on the deep lattice used for fluorescence imaging. This ensures that the probability of two atoms being on the same site is negligible, avoiding photo-assisted losses for accurate atom number determination.

Image analysis

A sparse atom cloud image is used to extract the PSF. The geometry of the lattice is then extracted from such an image. First, the lattice spacing is obtained and then the region of interest is fitted in blocks of 10 by 10 lattice sites. The block centers are allowed to vary to extract any distortions of the lattice pattern due to imaging aberration over the field of view. A histogram of atom brightness is used to set a threshold that identifies the presence or absence of an atom on a site. The information about the PSF, lattice geometry and threshold obtained from these sparse images is then used to fit other images with much higher lattice filling, only allowing for a single global offset in the lattice phase determined by fitting atoms at the edges of the cloud.

During imaging, a small fraction of the atoms are lost due to background gas collisions. If this occurs before they scatter enough photons to surpass the detection threshold, they are not counted. The mean fraction of such uncounted atoms is $1.75 \pm 0.02\%$, determined from 15 movies (30 frames, 0.5s exposure per frame) of the atom population decay in the near-resonant lattice.

Correction of disorder in optical potentials

Scatterers on optical surfaces produce ring-like patterns on the optical potentials used to trap the atoms. The spatial pattern of the disorder is static in time and for a lattice depth of $22E_r$, has an RMS gradient of $(0.13 \pm 0.01)U$ per lattice site and a characteristic length scale of 10 lattice sites. The contour lines of the potential are directly extracted from the shell structure boundaries in the Mott regime. Different contour lines are obtained by varying the atom number. The ability to observe these contour lines enables us to correct the potential by projecting an appropriate light pattern through the objective. This pattern is produced by illuminating a digital micromirror device (DLP Discovery 4100, Texas Instruments) with incoherent light of spectral width 1nm, centered at 840nm. A block of $14 \times 14$ mirrors maps onto a single lattice site in the plane of the atoms, allowing the creation of grayscale patterns, with the aperture of the objective providing Fourier filtering. An error diffusion algorithm [35] is used to convert the desired grayscale image to a binary pattern. Potential corrections of either sign are possible by operating the micromirror device with a bias light level produced by flattening the profile of the Gaussian illumination beam.

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[16] Materials and methods are available as supporting material on Science Online.