Metastable Bose-Einstein condensation in a strongly correlated optical lattice

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We experimentally and theoretically study the peak fraction of a Bose-Einstein condensate loaded into a cubic optical lattice as the lattice potential depth and entropy per particle are varied. This system is well described by the superfluid regime of the Bose-Hubbard model, which allows for comparison with mean-field theories and exact quantum Monte Carlo (QMC) simulations. By correcting for systematic discrepancies between condensate and peak fraction, we find that the QMC simulations and measured peak fraction agree at low entropies per particle.

I. INTRODUCTION

Research efforts using ultracold quantum gases trapped in optical lattices to study models of strongly correlated matter, such as cuprate superconductors, are at a crossroads. Rapid experimental progress has enabled emulation of ever more interesting and complex Hamiltonians, including those that go beyond conventional condensed matter paradigms. However, in most cases, the ultralow temperatures required to study exotic many-particle ground states have been inaccessible to experiments [1]. For example, experiments have prepared a gas at temperatures close to the Néel temperature for fermionic atoms trapped in a cubic lattice [2], which realize the Hubbard model. However, there are no known viable methods for reaching the orders-of-magnitude lower temperature required to achieve the putative $d$-wave superfluid state.

The current approach to achieving low temperatures in experiments is to cool a gas confined in a parabolic trap and then slowly turn on an optical lattice potential. Reaching lower temperatures in the lattice requires achieving lower entropy per particle in the trap. At best, turning on the lattice preserves entropy per particle. Generally, imperfections in the experiment, such as spontaneous scattering of the lattice light, and nonadiabaticity will lead to heating, and the entropy per particle of the gas in the lattice will be higher than the initial state. There are also proposals to cool the relevant degrees of freedom for atoms in the lattice, which have not been realized in experiments. In either approach, understanding and verifying equilibration and measuring temperature are critical to exploring low entropy states.

Understanding relaxation and equilibration dynamics in strongly correlated systems is especially challenging. Exact dynamics can only be calculated reliably for one-dimensional systems and for short times. In two and three dimensions, we are often forced to rely on approximate time-dependent mean-field theories, which ignore correlation effects, or exact simulations of small numbers of particles, which may not reproduce experiments. Ultracold gas experiments introduce an additional complication in that they are isolated from the environment and a thermal bath is absent. Equilibration and thermalization in closed quantum systems via interparticle interactions is an open topic rife with unresolved questions [3,4].

Measuring temperature in closed quantum systems is also challenging. In systems connected to a reservoir, such as electronic solids, thermal contact is maintained with a carefully calibrated thermometer. In contrast, in closed systems, temperature must be measured via its relation to an observable derived from the system itself. Such a procedure can be problematic when the physics and energy spectrum of the system of interest are not fully understood.

In this paper, we use a combination of experiment, quantum Monte Carlo (QMC) simulations, and semianalytic theory to explore equilibration and temperature in a prototypical strongly correlated system: the Bose-Hubbard (BH) model realized using ultracold bosonic $^{87}$Rb atoms trapped in an optical lattice [5,6]. In the BH model, particles tunnel between adjacent lattice sites with energy $t$, and particles interact on the same site with energy $U$. Changing the ratio of $t/U$ realizes a quantum phase transition between superfluid and Mott insulating states [7,8]. An important feature of the BH model is that the equilibrium properties can be computed using a variety of theoretical tools, and thus the experiments and theory can be benchmarked against one another. If $t/U < 1$, strong coupling expansions [9,10], Gutzwiller theory [11], and dynamical mean-field theory [12] can be used. Conversely, if $t/U > 1$, the Hartree-Fock-Bogliubov-Popov (HFBP) approximation can be employed [13–15]. In this paper, we comprehensively test the ability of these approximate methods to predict condensate fraction, which is a primary experimental observable, by comparing to quantum Monte Carlo simulations. QMC [16,17] is statistically exact for the BH model for all ranges of $t/U$, since the sign problem is absent. For all theoretical and...
numerical predictions, we include the effects of the parabolic trapping potential that is present in combination with the lattice in experiments.

To experimentally test equilibration, we compare peak fraction $f_0$ [13,18–20] measured for lattice gases prepared at varied temperature and $t/U$ (in the superfluid regime) to predictions from QMC simulations. Gases are cooled to different temperatures in the trap before the lattice is turned on. Temperature and entropy per particle are straightforwardly determined for the weakly interacting trapped gas using time-of-flight (TOF) imaging and a semi-ideal model [21]. After the lattice is turned on slowly compared with the tunneling and interaction time scales, peak fraction—which we show using QMC simulations is closely related to condensate fraction $n_0$ and can be used as an accurate thermometer—is measured via TOF imaging. If the lattice turn-on is isentropic and the gas remains in thermal equilibrium, then the measured peak fraction should match the QMC prediction for the entropy per particle measured before the lattice is turned on. At fixed entropy per particle, the peak and condensate fraction drops at higher lattice potential depths because of strong interparticle interactions. Generally, the peak and condensate fraction will be lower (and temperature higher) in the lattice than predicted for the adiabatic case because of heating processes.

At low entropy per particle, we observe good agreement with QMC predictions for peak fraction. However, at higher temperatures, $f_0$ is larger than predicted, implying lower entropy per particle in the lattice than the initial state, in apparent contradiction with the second law of thermodynamics. Above the critical entropy per particle for superfluidity in the lattice, a regime that has not been widely explored in experiments, we find that $f_0$ is nonzero. When such disagreement is present, we show that there is no clear adiabatic time scale in the experiment by measuring $f_0$ for varied lattice-turn-on times.

While a full modeling of the dynamics present in the experiment is beyond the scope of this paper, we obtain a rough estimate for the time scales for equilibration and the lifetime of the condensate by calculating the Landau damping time in the lattice. We find that the time scale for thermalization can be much longer than the tunneling and interaction time scales at high entropy per particle. Furthermore, the condensate lifetime is comparable to the heating time scale, making discrimination of adiabaticity in the experiment difficult. Our measurements in combination with these calculations imply the disagreement between QMC and the experiment is likely due to a lack of equilibrium and metastability of the condensate in the lattice, making thermal equilibrium difficult to achieve and challenging to distinguish from heating.

This paper is organized as follows. In Sec. II, we discuss the experimental procedure along with the measurement protocols used to obtain $f_0$ as well as the entropy per particle $S/N$. In Sec. III, we describe how the equilibrium $n_0$ and $f_0$ are computed using full-scale QMC ab initio simulations and mean-field theories. We show that the mean-field theory results are generally in poor agreement with exact QMC predictions. In Sec. IV, we discuss our results and show a comparison between the experimentally measured $f_0$ and QMC predictions. The discrepancy with experimental measures of $f_0$ inevitably points to a lack of adiabaticity during loading and the need for a better understanding of the time scales of dynamics in this system. Subsequently in Sec. V, we attempt to obtain a rough estimate of the time scales for relaxation of the peak fraction by calculating the Landau damping time in the lattice for the experimentally measured entropies. Our calculations suggest that the time scale for equilibration is comparable to the time scale for heating, implying that the lack of agreement between the theoretical and experimental results may be due to a nonequilibrium effect. In Sec. VI, we present concluding remarks and discuss future research directions.

II. EXPERIMENT

The details of our $^{87}$Rb Bose-Einstein condensate (BEC) lattice apparatus are described in Ref. [22]. We prepare approximately 200,000 $^{87}$Rb $|F = 1, m_F = −1⟩$ atoms confined in a harmonic trap that is formed from a single-beam optical dipole trap at 1064 nm traveling perpendicular to gravity and a magnetic quadrupole trap balancing gravity and providing additional harmonic confinement in the two horizontal directions. We control the entropy per particle by evaporating to different depths of the dipole trap, after which we ramp the depth to a constant value with a mean harmonic trap frequency of $\omega_0 = 2\pi(35.78 ± 6)$ Hz. To keep the atom number roughly fixed for the different temperatures, we selectively adjust the efficiency of the evaporation.

A cubic lattice is formed by exponentially ramping on three sets of retroreflected lattice beams at $\lambda = 812$ nm over 100 ms using a 200 ms time constant (for most of the data in this paper). The lattice potential depth is characterized as $sE_R$, where $E_R = (\hbar/\lambda)^2/2m = 2.31 \times 10^{-30}$ J is the recoil energy. We use Kaptiza-Dirac diffraction to calibrate $s$ to within 7%. The lattice beams add additional harmonic confinement, and the overall confinement is well described by $\omega = \sqrt{\gamma_0^2 + \frac{S_0 E_R}{m(2\pi/18\mu m)^2}}$. After turning on the lattice and waiting for 10 ms, we turn off all potentials and allow the atoms to expand for 20 ms time of flight. We optically pump atoms into the $F = 2$ state, and take an absorption image on the $F = 2$ to $F' = 3$ cycling transition.

The optical depth (OD) in absorption images saturates for OD $> 3$ (i.e., high column density) because of nonabsorbable light in the imaging laser beam. This is a complication for imaging condensed gases at nonzero temperature, which possess high-density peaks and a low-density thermal component. To increase the dynamic range of imaging and obtain high signal-to-noise ratio data for the condensate and noncondensate atoms, we take a series of two images in separate measurements. We produce high OD images by transferring all the atoms into $F = 2$, and low OD images are acquired by pumping a fraction of the atoms so that the OD $< 2$. High OD images are used to measure the atoms outside of the condensate peaks, and low OD images are employed to measure the atoms in the peaks (which are saturated in the high OD images). We calibrate the relative number between high and low OD images and verify that quantities such as condensate fraction agree between the two modalities using a gas released from the trap.

A typical experimental sequence at a given lattice depth and entropy per particle involved seven sets of four images: a high OD and low OD image in the harmonic trap, and a high
OD saturates at approximately 3 because of nonabsorbable light in the starting conditions were almost identical for all lattice depths. The iterations. The top line shows the gas released from the harmonic trap; lattice depths (in units of $k_B$) and initial entropies averaged over seven OD and low OD image in the lattice. All quantities presented in this paper are averages over these seven sets of images. Figure 1 illustrates the resulting averaged high OD images.

To determine $S/N$ for the gas before turning on the lattice, images for gases released from the harmonic trap were fit using a multistep scheme similar to that employed in Ref. [23] to determine the condensate fraction. The high OD image was fit to a Thomas-Fermi (TF) profile combined with an independent Gaussian function, taking into account the line-of-sight integration. The pixels within a radius 10% greater than the TF radius were masked, and the remaining image was fit to a Bose-Einstein distribution to determine the number of thermal atoms $N_{th}$. In images with sufficient signal, the low OD image was fit to a combined TF-thermal profile, and the condensate number $N_{BEC}$ was determined from the TF component. The condensate fraction was determined as $N_{BEC}/(N_{th} + N_{BEC})$ in this case. If the signal-to-noise ratio was too small to resolve the thermal component in the low OD image, a fit to a simple TF profile was used to determine the number of atoms $N_{TF}$ within the TF radius. The total number of atoms $N$ was determined in this case by adding the number of atoms outside the TF radius in the high OD image. The condensate fraction was then $1 - N_{th}/N$. Entropy per particle was determined from the condensate fraction and total number using the semi-ideal model [21].

Peak fraction $f_0$ was determined from images of the gas released from the lattice using the procedure from Ref. [24]. Several complications make interpreting fits to TOF images for lattice gases difficult. Unlike the trapped case, the functional form of the density distribution after TOF is unknown. Furthermore, atoms that appear outside of the condensate peaks are a combination of superfluid atoms (expelled from single-particle low-momentum states by strong interactions) and thermal atoms. A final challenge is specific to our apparatus—there is no spherical symmetry in a lattice momentum distribution, and our imaging direction is not along a lattice direction. This makes even the noninteracting distribution difficult to fit [see Eq. (23) in Ref. [22]].

In the face of these difficulties, we have adopted a heuristic approach that is fast and independent of experiment details. In the high OD image, we mask the peaks and fit the distribution to a single Gaussian. The number of atoms from this fit is $N_{Gauss}$. Next, the Gaussian fit is subtracted from the image and we fit each peak to a TF profile. These fits are used to mask the peaks, and the remaining signal is summed to determine the number of atoms $N_{\Sigma}$ outside of the peaks and not included in the Gaussian fit. These steps capture all the nonpeak atoms, except those that are part of the broader distribution and coincident with (or “under”) the peaks in the image. Counting these atoms accurately would require knowledge of the density distribution after TOF, including the effects of interactions [25–28]. Since analytical expressions for the TOF distribution are unknown, we make the simple assumption that the nonpeak distribution that overlaps with the peak is uniform [see (a) and (b) of Fig. 2]. We determine the average nonpeak OD coincident with the peak by averaging the OD around the perimeter of the peak. The number of nonpeak atoms $N_{nonpk}$ coincident with the peak is estimated using a uniform distribution with this average OD. The total nonpeak number of atoms is $N_{nonpk} = N_{Gauss} + N_{\Sigma} + N_{nonpk}$.

We fit only the peaks of the low OD image to determine the peak number $N_{pk}$ using a series of TF profiles (subtracting off a Gaussian if there is a sufficient signal) [see (c) and (d) of Fig. 2]. The total number is then $N = N_{pk} + N_{nonpk}$, and the peak fraction is $f_0 = 1 - N_{nonpk}/N$. In the next section we demonstrate that $f_0$ is related to the condensate fraction $n_0$ by applying this fitting procedure to simulated TOF images produced using QMC calculations.

III. EQUILIBRIUM THEORY

To assess whether equilibrium is achieved in the lattice, we determine peak fraction $f_0$ and entropy per particle $S/N$ for equilibrium gases using QMC simulations. We also compute condensate fraction $n_0$ in order to establish the relationship between condensate and peak fraction. We match the parameters of the experiment as closely as possible. The particle number is kept constant within 5% to $N = 200,000$ for optical lattice depths of $s = 6, 8, 10$, and $12$. Furthermore, the confining potential is accounted for exactly in the QMC simulations.

QMC simulations are carried out using the stochastic series expansion method (SSE) [29]. SSE is an exact method suffering from no biases or approximations. The condensate fraction $n_0$ is obtained directly from the single-particle density matrix in adherence to its rigorous definition as the ratio of the occupation number of the macroscopically occupied single-particle mode to the total particle number. The general prescription used to calculate $n_0$ and the momentum distribution $n(k)$ that includes finite TOF effects has been discussed elsewhere [30].

We use the momentum distribution $n(\vec{k})$ from QMC simulations to generate line-of-sight integrated images. Finite TOF effects—which lead to additional overlap between the condensate and noncondensate distributions—are fully accounted for as in Ref. [30], and the integration direction is
matched to the experiment. These images are then used to extract the peak fraction using the fitting procedure applied to the experiment modified to work with a single image. Figure 3 shows \( n_0 \) and \( f_0 \) computed at the same temperature for the \( s \) sampled in the experiment. It is evident that \( n_0 \) and \( f_0 \) are monotonically related, and that peak fraction can be used to determine temperature in the experiment (assuming that equilibrium is achieved). The relationship between \( f_0 \) and \( n_0 \) also does not depend strongly on lattice potential depth. When \( n_0 < 0.05 \), the fitting procedure cannot identify the presence of condensate even with virtually noiseless QMC data.

Generally, \( f_0 \) is less than \( n_0 \). This discrepancy between \( f_0 \) and \( n_0 \) arises from inaccurate accounting of the condensate and noncondensate distributions. In part, this is due to the high-momentum tails of the condensate that extend into the broad Gaussian-like part of the momentum distribution, and that are mistakenly accounted for as noncondensate by the fitting procedure [30]. Apart from these systematic errors in the fitting protocol, there may also be additional interaction effects that arise during TOF that we cannot account for in the QMC simulations.

The QMC simulations are carried out at temperatures that span the range explored by the experiment, but that are chosen independently of the measurements. In order to compare with experiments that have access to the entropy per particle but not temperature, we also use QMC simulations to compute the temperature dependence of \( S/N \). This is done by integrating the internal energy per particle \( [u(\beta) \equiv U/N(\beta)] \) shown in Fig. 4, obtained from QMC simulations, where \( \beta = 1/k_BT \).

Using the definition \( u(\beta) = u_0 + f(\beta) \), where \( u_0 \equiv \lim_{\beta \to \infty} u(\beta) \) is the ground-state energy and \( f(\beta) \) is a monotonically decreasing function that characterizes the temperature dependence with \( \lim_{\beta \to \infty} f(\beta) \to 0 \), we obtain

\[
\frac{S}{N(\beta)} = \beta f(\beta) |_{\beta=0}^{\beta=\infty} + \int_{\beta}^{\infty} f(\beta) d\beta = \beta f(\beta) + \int_{\beta}^{\infty} f(\beta) d\beta.
\]

(1)

Here the limit \( \lim_{\beta \to \infty} \beta f(\beta) \) can be taken provided \( f(\beta) \) decays faster than linearly—a condition that is easily met since the data fit well to an exponential function with a reduced \( \chi^2 \sim 1 \). Our strategy is to integrate \( u \) using a fit to all sampled points. We fit the high \( \beta \) tail to an exponential decay, since we cannot use a finite temperature method to access the true ground-state energy at \( \beta \to \infty \). The high temperature points are fit using a cubic spline. Error bars for \( S/N \) are computed using a resampling procedure. For each \( u(\beta) \) that we have computed, we sample a Gaussian of mean \( u(\beta) \) and standard

FIG. 2. (Color online) Illustration of the fitting procedure used to determine peak fraction from a high OD image [(a) and (b)] and from a low OD image [(c) and (d)]. Here we use an averaged image for \( s = 10 \) and \( S/Nk_B \approx 1.35 \) (middle image in Fig. 1). For all these images, we have first subtracted off a Gaussian fit as described in the main text. In (a), we show the high OD image with the peaks masked and the nonpeak distribution coincident with the peak assumed to be a uniform distribution. Note that we have masked the nine peaks that correspond to zero momentum and the first- and second-order diffraction peaks. We ignore the higher-order peaks, which contain a negligible number of atoms. A slice along the red dashed line is displayed in (b). The black line is the nonpeak distribution, and the red line is the complete image including the peaks. In (c), we show the low OD image used to determine the peak number. A slice along the red dashed line is displayed in (d). The black line is the TF fit to the peaks and the red line is the data from the image. Note that there is a scale factor between the high and low OD images of approximately 5.75.
deviation $\sigma(u)$ to generate a sample $u_i(\beta)$. A collection of
$u_i$ describe the vector $U_i \equiv \{u(\beta)\}$ that is fitted to obtain an
estimate of $(\langle r_i^2 \rangle_0)^{1/2}$. This procedure is repeated multiple times
to obtain an estimate of $(S/N(\beta))$ and $\sigma(S/N)$.

We compare the QMC results to a variety of approximate
theories in order to evaluate the accuracy of these less
time-consuming methods. We consider mean-field theories
(MFT) viz., the Hartree-Fock (HF), HFBP, and Gutzwiller
approximations. While these approaches have been used to
analyze experiments as benchmarks for more sophisticated
theories [14,19,20,31–34], their validity has not been compre-

sively examined. For calculations involving MFTs, the trap
is accounted for using the local density approximation.

The HF and HFBP approaches rely on a perturbative ex-
ansion of the local field operator $\hat{\psi}(\vec{r}) = \psi_0(\vec{r}) + \delta \hat{\psi}(\vec{r})$
that accounts for the macroscopically occupied scalar condensed
mode $\{\psi_0(\vec{r})\}$ together with the fluctuations $\{\delta \hat{\psi}(\vec{r})\}$. The
former treats the kinetic energy terms of the BH Hamiltonian
exactly, while treating the interactions via the HF approxima-
tion, which leads to the HF spectrum

$$\epsilon_{HF}(\vec{q}) = \epsilon(\vec{q}) - \mu + 2U(\rho_0 + \rho_h). \tag{2}$$

where $\vec{q}$ is the quasimomentum, $\rho_0$ is the condensate
density, $\rho_h$ is the noncondensate density, $\mu$ is the chemi-
cal potential, and $\epsilon(\vec{q}) = 2(3 - \cos(q_d d_h) - \cos(q_d d_h) - \cos(q_d d_h))$ is the tight-binding dispersion for single particles
in the lowest energy band. The HFBP approach allows for
mixing of partikelike and holelike excitations, which produces a
dispersion

$$\epsilon_{HFBP}(\vec{q}) = \sqrt{[\epsilon_{HF}(\vec{q})]^2 - (U \rho_0)^2}. \tag{3}$$

We obtain the noncondensed density via

$$\rho_h = \frac{1}{2} \sum_{\vec{q} \neq 0} \left[ \frac{\epsilon_{HF}(\vec{q})}{\epsilon(\vec{q})} \coth[\beta \epsilon(\vec{q})/2] - 1 \right]. \tag{4}$$

where $\epsilon(\vec{q}) = \epsilon_{HF}(\vec{q}) [\epsilon_{HF}(\vec{q})]$ for the HF (HFBP) calculation. Due to the appearance of $\rho_h$ on both sides of (4),
it must be solved self-consistently, starting with a guess
for $\rho_0$ and utilizing the relation $\mu = U(\rho_0 + 2\rho_h)$. Using
the spectrum determined using this procedure, we calculate
relevant thermodynamic observables, such as the entropy

$$S(\beta) = \sum_{\vec{q} \neq 0} \frac{\beta \epsilon(\vec{q})}{e^{\beta \epsilon(\vec{q})} - 1} - \ln(1 - e^{-\beta \epsilon(\vec{q})}). \tag{5}$$

where $\beta = 1/k_B T$.

The Gutzwiller approach utilizes a perturbative expansion
of the kinetic term (thereby treating it approximately) while
keeping the interaction term intact. We use $\hat{a}_i^\dagger \hat{a}_i \approx \langle \hat{a}_i^\dagger \hat{a}_i \rangle + \langle \hat{a}_i \rangle \hat{a}_j - \langle \hat{a}_i \rangle \hat{a}_j$ to decouple the Hamiltonian, where $\hat{a}_i$ and
$\hat{a}_j$ are particle creation operators on neighboring sites $i$ and $j$.

In this approximation, the Hamiltonian becomes independent
The energy per particle for varying lattice depths $s = \{6, 8, 10, 12\}$ [(a)–(d), respectively] for a system of $N \sim 200\,000$ particles in a trap. These points were fitted using a procedure outlined in the text to obtain the entropy per particle that can be compared with experiment. Temperatures only below the critical temperature for superfluidity are sampled.

The energy per particle is given by:

$$H \approx -zt(\alpha \hat{a}^\dagger + \alpha^* \hat{a} - |\alpha|^2) + \frac{U}{2} (\hat{n} - 1) - \mu \hat{n}, \quad (6)$$

where $z = 6$ is the coordination number and $\hat{n} = \hat{a}^\dagger \hat{a}$. Using a truncated occupation number basis, we construct a matrix for $H$ that is diagonalized utilizing the relation $\alpha = \sum_i \langle i | \hat{a} | i \rangle e^{-\beta E_i}$, needed to account for the finite temperature occupation of states. We solve this system iteratively, starting with a guess for $\alpha$ which is updated until convergence occurs. All thermodynamic quantities are computed using $\alpha$ at a specific $\beta$, $\mu$, and $U/t$.

In order to account for the parabolic trap used in experiments, we rely on the local density approximation (LDA) for MFT results. The LDA treats the slowly varying external confining potential as a local chemical potential shift. Thermodynamic quantities are computed at different $\mu(r)$ (corresponding to a homogeneous system) and then integrated over all $r$ to determine observables for the trapped lattice gas. The LDA ignores correlations across the trap.

We show predictions for condensate fraction $n_0$ at different entropies per particle and lattice potential depths in Fig. 5 for QMC simulations and MFTs. In each case, $T$ and $\mu$ are varied to match $S/N$ and $N$ as computed by the method of interest. While all approaches show the same qualitative trend, it is clear that HF theory is, in general, not in good agreement with the QMC approach. This disagreement is expected because HF theory does not account for the mixing of particle and holelike excitations that result from interactions.

The comparisons of QMC against Gutzwiller and HFBP results suggest that for moderate interaction strengths (at $s = 6$ to 8) these MFTs generally work well. For stronger interactions (at $s = 10$), HFBP calculations fail to capture finite temperature effects, whereas the Gutzwiller approximation is able to capture qualitative features. At the highest lattice potential depths we sample ($s = 12$), the Gutzwiller method can capture the high temperature regime, but cannot accurately compute the correlations at low temperature, as is evidenced by the 15%–25% error in $n_0$ for $S/N < 0.5k_B$.

While the validity of the Gutzwiller approximation over a wide range of interaction strengths and temperatures is suggestive, this is only likely true for the high filling limit (a central density of approximately three particles per site) considered in this work. QMC simulations for low filling systems show that the differences with the Gutzwiller approximation can be quite large, as evident for the results for a central density of one particle per site shown in Fig. 6. The discrepancy apparent in Fig. 6 may also arise from quantum critical behavior, as the lattice potential depth is set close to the Mott-insulator to superfluid phase transition.

We conclude that the HF and HFBP approximations should not be used to compute observables such as $n_0$ and thermodynamic variables like $S/N$. While the Gutzwiller approximation can provide quantitatively accurate results at high filling and for moderate lattice potential depths, it fails at low $S/N$ and for strong interactions and for low densities. We therefore use only QMC simulation results to compute $S/N$ and $f_0$ for comparisons with the experimental data.
IV. RESULTS AND DISCUSSION

To assess thermal equilibrium in the experiment, we show a comparison between the experimentally measured $f_0$ (determined from images shown in Fig. 1 and the fitting procedure described in Sec. I) and the equilibrium QMC prediction at the same $S/N$ in Fig. 7. A strong discrepancy is evident: there is an alarming degree of disagreement between the QMC results and the experiment at high temperatures (i.e., high $S/N$). The experimentally observed peak fraction (and therefore condensate fraction) is systematically high compared with the QMC prediction. What is more troubling is that the condensate survives beyond the transition temperature, which is marked by $f_0$ (and $n_0$) vanishing in the QMC predictions.

We do not believe that this discrepancy can be explained by systematic errors in the analysis of the experimental data. One source of error might be that binary collisions between atoms in different peaks eject condensate atoms during TOF [35,36]. While this effect may play a role in suppressing peak fraction for the lowest entropies at $s = 6$, its impact should be reduced at higher lattice potential depths and temperatures as the condensate density decreases. Furthermore, this effect would lead to lower, not higher, peak fractions than the equilibrium QMC prediction.

Another possibility is that we erroneously identify peaks in the TOF image. It has been previously discussed that sharp peaks in momentum distributions may not be uniquely associated with a condensate [37,38]. However, calculations focusing on trapped systems for realistic experimental conditions and experiments have demonstrated that the visibility, which is a quantitative measure of peak sharpness, can distinguish between condensed and noncondensed states [20,39,40]. The features we observe at high $S/N$ are narrow and quantitatively similar in extent to the peaks observed at low temperatures. The sharpness of the peaks we measure at high $S/N$ is most

FIG. 5. (Color online) Comparison between different types of mean-field theories against QMC results. Panels (a)–(d) correspond to $s = 6$, 8, 10, and 12, respectively. The particle number was kept constant at $N \sim 200,000$. All other parameters were matched to experimental specifications presented in Fig. 7. Green squares (dashed line) correspond to HF, orange triangles (dotted line) to HFBP, red circles (solid line) to Gutzwiller, and black diamonds (with error bars) to QMC. The error bars show the statistical uncertainty in the QMC results.

FIG. 6. (Color online) Comparison between Gutzwiller (red triangles) and QMC (black squares) results for a strongly interacting system ($s = 13.6$) at low filling. The central density for these results is approximately one particle per site with $\omega = 2\pi \times 55.6$ Hz and $N \sim 60,000$. The line is a guide to the eye.
s = is the mean of the number at each lattice depth (results. The uncertainty in the QMC results is too small to be visible. The error bars in the experimental data show the standard error of the mean for the average across seven measurements. The horizontal error bars for the measurements also include the influence of the uncertainty in the trap frequency and number of atoms and do not impact the QMC predictions and the experimental data agree well in this regime. Another issue is how the lattice turn-on time compares with the Mott-insulator regime, which we do not sample here. Furthermore, we estimate that overall rms radius of the gas changes by less than 20% at t = 10 ms, which vary from 6 ms and 2 ms for (a) s = 6, N = [2.38 ± 0.03, 2.08 ± 0.06, 1.64 ± 0.03, 2.04 ± 0.07, 2.06 ± 0.09] x 10^5, (b) s = 8, N = [2.36 ± 0.06, 2.05 ± 0.11, 1.66 ± 0.07, 2.02 ± 0.09, 1.73 ± 0.05] x 10^5, (c) s = 10, N = [2.41 ± 0.08, 2.15 ± 0.11, 1.68 ± 0.09, 2.03 ± 0.16, 1.85 ± 0.09] x 10^5, and for (d) s = 12, N = [2.55 ± 0.12, 1.98 ± 0.10, 1.65 ± 0.04, 2.06 ± 0.11, 1.74 ± 0.11] x 10^5. QMC curves are constrained to a constant number, which is the mean of the number at each lattice depth (\(N_s \approx 200\) 000). The tunneling, interaction energies, and trap frequencies for s = \{6, 8, 10, 12\} are, respectively, \(t = \{51, 55.71, 58.12, 59.4\}\) x 10^{-11}s, \(U = [3.92, 5.23, 6.48, 7.67]\) x 10^{-31}J, and \(\omega = 2\pi \times [51.27, 55.48, 59.4, 63.07]\) Hz. The error bars in the experimental data show the standard error of the mean for the average across seven measurements. The horizontal error bars for the measurements also include the influence of the uncertainty in the trap frequency and number of atoms and do not impact the QMC results. The uncertainty in the QMC results is too small to be visible.

FIG. 7. Experimentally determined peak fraction at each lattice depth as a function of the initial entropy per particle in the harmonic trap (circles). Corresponding peak fraction from QMC simulations determined by fitting the momentum distribution using the same procedure used in experiments in the lattice are also shown (triangles). The number of atoms at each experimental point (from lowest to highest entropy) is \(s = 6, N = [2.38 ± 0.03, 2.08 ± 0.06, 1.64 ± 0.03, 2.04 ± 0.07, 2.06 ± 0.09] \times 10^5\). The experimental observations imply that equilibrium is not established and the condensate is metastable during the lattice turn-on. A natural question is whether the lattice is turned on before the condensate appears present at temperatures exceeding the critical temperature for superfluidity in the lattice. Condensate and peak fraction should always decrease in equilibrium as the lattice is turned on to higher s because of quantum depletion resulting from strong interparticle interactions. The experimental observations imply that equilibrium is not established and the condensate is metastable during the lattice turn-on. A natural question is whether the lattice is turned on before the condensate appears present at temperatures exceeding the critical temperature for superfluidity in the lattice. Condensate and peak fraction should always decrease in equilibrium as the lattice is turned on to higher s because of quantum depletion resulting from strong interparticle interactions.

A further potential source of systematic error is heating by the lattice light [1], which would have the largest impact at low entropy. However, we find that the QMC equilibrium predictions and the experimental data agree well in this regime. For higher entropies, heating should reduce peak fraction below the QMC prediction, which is opposite to the observed behavior.

We conclude that the measured peak fraction is systematically higher than that allowed in equilibrium by the second law of thermodynamics—the measured \(f_0\) implies that the gas is at lower \(S/N\) after the lattice is turned on. Furthermore, a condensate appears present at temperatures exceeding the critical temperature for superfluidity in the lattice. Condensate

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The exponential time constant used for ramping on the lattice light is set to 2τ. Data and theoretical predictions are shown for \( S/N = 0.44 \) (triangles, solid), 1.2 (squares, dashed), and 2.2 (circles, dotted) \( k_B \). The lines are the equilibrium \( f_0 \) predicted by QMC simulations taking into account heating from the lattice light. At the highest \( S/N \), QMC predicts that a condensate should be absent in the lattice.

In all cases, no adiabatic time scale is apparent—the peak fraction continuously decreases with the lattice turn-on time. This behavior is characteristic for all the \( s \) sampled in this work. At low and intermediate \( S/N \), the decrease in peak fraction is consistent with the heating from the lattice light. We compute the predicted equilibrium peak fraction including heating for different lattice turn-on times \( \tau \) using QMC results for \( u \). The rate of increase in \( u \) from the lattice light and the corresponding increase in \( S/N \) and decrease in \( f_0 \) are determined using the results from Ref. [1]. We conclude that the adiabatic time scale is comparable to or longer than the time scale for heating, and thus discriminating between the two processes is not possible.

Ultimately, interparticle interactions via binary collisions are the dominant mechanism by which equilibrium is established and condensate fraction decreases as the lattice potential is turned on. Despite extraordinary advances in methods such as QMC, computing exact dynamics in higher than one dimension for more than a few tens of particles and short times remains infeasible for strongly interacting systems such as the BH model. In the next section, we use a hybrid of equilibrium QMC simulations and approximate theory to compute the time scale associated with the simplest mechanism—Landau damping—for relaxation of \( n_0 \).

V. LANDAU DAMPING THEORY FOR RELAXATION OF THE CONDENSATE

In this section, we outline a model for relaxation of the condensate atom number via Landau damping. In a closed system, the condensate number can change by collisions with quasiparticles, whose number is not conserved. The dominant collision processes involve two-body scattering, where a condensate atom collides with a quasiparticle to produce two new quasiparticles. In equilibrium, the rates for this process and its reverse (whereby two quasiparticles collide to produce a condensate atom and a quasiparticle) are identical but opposite in sign, and the condensate number is conserved. However, out of equilibrium, these processes do not cancel, and induce changes in the condensate atom number.

In general, the theoretical problem of how the condensate interacts with the thermal gas is an extremely challenging one, but an understanding of the precise mechanisms for equilibration is paramount to current experiments in optical lattices. Briefly, a condensate can relax by colliding with thermal atoms in two ways: At the mean-field level, the condensate introduces fluctuations in the thermal cloud which in turn lead to a damping of the condensate. This process is termed Landau damping, which is what we calculate. There is a second process where the condensate atoms exchange energy with the thermal atoms via collisions [42,43]. Although we do not calculate this process here, it is roughly of the same order as the Landau damping mechanism we compute [43]. A third process is called Beliaev damping, whereby a noncondensed atom collides with a condensate atom and decays into two noncondensed atoms with lower energy. This mechanism is dominant at very low temperatures and its rate decreases rapidly with temperature. Calculating the Landau damping rate thus suffices to estimate the time scales for equilibration in lattices.

Our treatment of Landau damping directly follows that of Tsuchiya and Griffin [44], who computed the damping rate for a homogeneous lattice Bose gas. Physically, this rate corresponds to the annihilation of a Bogoliubov quasiparticle and a condensate atom to produce two Bogoliubov quasiparticles (Fig. 9). By conservation of total particle number, integrating the damping rate over all momenta yields the rate at which the condensate fraction changes via Landau damping. For a homogeneous gas, the Landau damping rate for a quasiparticle...
at momentum $\vec{q}$ is given by \[ \Gamma(\vec{q}) = \pi n_0 \sum_{\vec{p}, k} |M_{\vec{q}, \vec{k}; \vec{p}}|^2 \delta(\epsilon(\vec{q})) \]

\[-[\epsilon(\vec{p}) - \epsilon(\vec{k})])[f^0[\epsilon(\vec{k})] - f^0[\epsilon(\vec{p})]],\]

\[M_{\vec{q}, \vec{k}; \vec{p}} = \frac{2U}{\sqrt{L^D}} \sum_{\vec{G}} [(u_{\vec{q}}u_{\vec{p}} + v_{\vec{q}}v_{\vec{k}} - u_{\vec{q}}u_{\vec{k}})\delta(\vec{q} + \vec{k} - \vec{p} - \vec{G}),\]

(7)

where $D$ is the dimension, $L$ is the length on one side of the system, $n_0$ is the condensate density, the $G$ are reciprocal lattice vectors (which incorporate umklapp scattering), \[ f^0(\epsilon) = \frac{1}{\exp(\beta \epsilon) - 1} \] is the Bose distribution at temperature $k_B T = 1/\beta$, and $\epsilon(\vec{k})$ are the usual Bogoliubov quasiparticle dispersions in a lattice of the form in Eq. (3). The Bogoliubov coefficients $u_{\vec{k}}$ and $v_{\vec{k}}$ are given by $u_{\vec{k}} = \sqrt{2\epsilon^2(\vec{k}) + 1}$ and $v_{\vec{k}} = \sqrt{2\epsilon^2(\vec{k}) - 1}$.

The detailed derivation of this result can be found in Ref. \cite{44} and is not repeated here. Note that a key assumption in this theory is that thermal (i.e., noncondensed) gas is in local equilibrium, which is enforced on short time scales by collisions between thermal atoms. We emphasize here that we are not modeling the full far-from-equilibrium problem of relaxation of condensed and noncondensed atoms following a lattice ramp. Rather, we compute the time scales for relaxation when the condensate density departs slightly from its equilibrium value at a given lattice depth. This near-equilibrium computation is the first step in gaining a systematic understanding of relaxation times in generic nonequilibrium situations.

We numerically compute the damping rate for all momenta $\vec{q}$ ranging from $-\pi/d \leq q_i \leq \pi/d$, where $i = \{x, y, z\}$, and $d$ is the lattice constant. We discretize momenta in steps of $\pi/L$, where $L$ is the system size, defined as where the total density vanishes. The finite system size also sets an infrared cutoff, $q_{\text{min}} = \pi/L$, which renders the one-dimensional integrals convergent. We have checked numerically that this cutoff is small enough that changing $q_{\text{min}}$ does not introduce significant errors.

The total damping rate is given by $\Gamma_L = \sum_{\vec{q}} \Gamma(\vec{q})/\Omega$, where $\Omega$ is the system size. Using a simple rate equation $d\delta n_0/dt = \Gamma_L \delta n_0$, where $\delta n_0(t) = n(t) - n_0$, we obtain an estimate for the relaxation time $\tau_{LD} = 1/\Gamma_L$ of the condensate.

To compute the damping rate, we require three quantities: the condensate density $n_0$, lattice depth $s$, and the temperature $T$. For a given lattice depth $s$ and temperature $T$, we use $n_0$ from exact QMC simulations discussed in Sec. III to compute the different Landau damping rate. Using the relation between temperature and entropy per particle, we obtain the damping rate as a function of entropy per particle for different lattice depths.

Our derivation of the damping rate assumes that all of the condensate atoms are at $\vec{q} = 0$. While this is true in a weakly interacting homogeneous system at low temperature, the presence of a trap and strong correlations modifies this picture significantly, as momentum is no longer a good quantum number. The condensate therefore develops a spread in momentum, which we have computed numerically using QMC for a given temperature and lattice depth.

To incorporate trap and interaction effects into our damping calculations, we produce the Landau damping rate in two cases: (1) We assume all the condensate atoms are at $\vec{q} = 0$. This yields an upper bound on the damping rate. (2) We obtain a lower bound on the damping rate by assuming that the total condensate number is equal to only the contribution to the momentum distribution at $\vec{q} = 0$. The experimentally observed damping rate must be between these two bounds.

The results of our calculations presented in Fig. 10 paint a fascinating picture. At low entropy, the lower bound on the Landau damping time ranges from 200 ms to 10 ms from the lowest to highest lattice depth $s = 6E_R$ to $s = 12$, while the upper bound varies roughly from 1 s to 100 ms. At the highest $S/N$, the Landau damping time is at least 20 times larger than the Hubbard tunneling time $h/t$. The decrease in the damping time for higher lattice potential depth is consistent with the increase in interaction strength. For $s = 6$, where the system can be treated as weakly interacting, the two bounds coincide at low entropy because the condensate largely resides at zero momentum. Interactions induce a significant broadening of the condensate momentum distribution, which leads to a large difference between the upper and lower bounds at higher $s$. This is particularly stark at high entropies per particle, because the condensate is almost entirely absent. Note that for $n_0 = 0$, $\Gamma_L \rightarrow \infty$. Strictly speaking, this theory is not accurate when the condensate number is small, either because of nonzero temperature or strong interactions. At high temperatures, interactions between noncondensate atoms play an important role that is not captured by our approach. For the strongly interacting case, quantum depletion reduces $n_0$ and the Bogoliubov spectrum, which is the underlying assumption in this theory, is not valid. Therefore the upper bounds in the damping time at $S/Nk_B = 1$ for $s = 10$ and 12 may not be correct.

Comparing the predicted Landau damping times to the 100 ms lattice turn-on used for the data in Fig. 8 is complicated by the lack of equilibrium in the lattice. Generally, the damping time exceeds 100 ms at the highest $S/N$, where the discrepancy between the QMC and experimental $f_{el}$ is greatest. This implies that the lattice turn-on time used for Fig. 7 was nonadiabatic, since times much longer than $\tau_{LD}$ are required to achieve equilibrium in the lattice. The predicted Landau damping time is also long at low $S/N$, but $n_0$ changes less in this regime as the lattice is turned on, and thus the impact of nonadiabatic behavior is less apparent. Diabaticity with respect to Landau damping at high $S/N$ explains the experimental observation that the peak and condensate fraction in the lattice is higher than the equilibrium prediction, since the condensate does not have sufficient time to decay as the lattice is turned on.

The upper bound on the damping time in combination with the heating rate from the lattice light implies that adiabaticity may be impossible to achieve under certain conditions. At $s = 6$, for example, the upper bound on the Landau damping time at $S/N > 1$ is close to 1 s. This time is longer than the heating time scale observed for the data shown in Fig. 8. Differentiating between heating and thermalization becomes more challenging at higher $s$, since the characteristic energy
scale $t$ drops rapidly, but the heating rate from the lattice light grows linearly with $s$.

VI. CONCLUSIONS

In conclusion, we have experimentally and theoretically investigated peak fraction in the superfluid regime of a cubic lattice for varied entropy per particle and lattice potential depth. Qualitatively, the experimentally observed peak fraction behaved as theoretically expected: lower initial entropies led to higher peak fractions, and higher lattice depths (lower $t/U$) resulted in lower peak fractions. However, we did not find quantitative agreement with exact QMC simulations at high entropy per particle. We observe that the condensate is metastable, and that nonzero peak fraction persists above the critical temperature for superfluidity for times much longer than the Hubbard time scales. We have investigated the time scales for relaxation of the condensate by calculating the Landau damping time in the lattice. Our calculations indicate that the relaxation time is comparable or greater than typical lattice turn-on times. The Landau relaxation time is also longer than the time scale associated with heating from the lattice light, which may make adiabaticity impossible to achieve.

Going forward, our theory-experiment comparison points to the urgent need to carefully understand time scales for dynamics in strongly correlated optical lattice experiments. Furthermore, using this model as a starting point, one can envision branching out to situations where competing theories disagree, break down, or do not even exist. A canonical example is that of strongly correlated systems far from equilibrium, where the very paradigms for how to think about these systems are only now being developed. In addition to being of fundamental importance, questions of nonequilibrium dynamics serve a practical purpose in cold atom experiments, which are largely isolated from the environment. Understanding the mechanisms for driving these systems to equilibrium is thus essential in order to understand time scales for maintaining adiabaticity.

ACKNOWLEDGMENTS

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