Synchronization crossover of polariton condensates in weakly disordered lattices

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We demonstrate that the synchronization of a lattice of solid-state condensates when inter-site tunnelling is switched on, depends strongly on the weak local disorder. This finding is vital for implementation of condensate arrays as computation devices. The condensates here are nonlinear bosonic fluids of exciton-polaritons trapped in a weakly disordered Bose-Hubbard potential, where the nearest neighboring tunneling rate (Josephson coupling) can be dynamically tuned. The system can thus be tuned from a localized to a delocalized fluid as the number density, or the Josephson coupling between nearest neighbors increases. The localized fluid is observed as a lattice of unsynchronized condensates emitting at different energies set by the disorder potential. In the delocalized phase the condensates synchronize, and long-range order appears, evidenced by narrowing of momentum and energy distributions, new diffraction peaks in momentum space, and spatial coherence between condensates. Our work identifies similarities and differences of this nonequilibrium crossover to the traditional Bose-glass to superfluid transition in atomic condensates.

I. INTRODUCTION

In a lattice of trapped bosons, disorder inhibits coherent tunneling so localizing condensation in real space. However on-site repulsive interactions combined with a flow of bosons can tune neighboring condensates into resonance, thus enhancing tunneling through the trap barriers. This subtle interplay is crucial for understanding the rich phase diagram composed of Mott-insulator, Bose-glass and superfluid, which acquires further complexity with pumping and dissipation in nonequilibrium situations. This situation is vital for understanding the capabilities of lattices of condensates to act for instance as quantum simulators of complex behavior. Exciton-polaritons (polaritons), which are bosons composed of an admixture of quantum-well exciton and microcavity photon [1], are an attractive system for studying the nonequilibrium Bose-Hubbard system in two-dimensions [2]. A lattice potential can be achieved either structurally by depositing thin metallic films [3] and etching [4] the microcavity or by photo-injecting excitons with a spatially patterned laser [5]. In the latter case, the repulsive on-site interaction can be dynamically tuned through polariton-polariton [6] and polariton-exciton nonlinearities [7]. For single polaritons the polariton nonlinearity is weaker than the disorder potential, but when polaritons condense into a macroscopic state [8–10] the collective nonlinearity can be large enough to exhibit effects such as superfluidity [11, 12] and solitons [13–15]. Polaritons in one condensate ‘puddle’ can tunnel out and drive neighboring condensates [16] described through the Josephson mechanism [17]. The mechanism is more complicated with disorder-induced energy detunings between two nearest neighbors, in the case of repulsive nonlinearity [18] (Fig. 1b). Here there is a critical Josephson tunneling (Jc), below which an extended condensate separates into two localized condensates at different energies, forming an unsynchronized (UNSYNC) phase. However above Jc, the flow of polaritons from the higher-energy to the lower-energy condensate collapses their energy detuning and synchronizes (SYNC) them due to repulsive nonlinearities, as the higher energy condensate becomes less populated and the lower energy condensate more populated than before (see lower panel in Fig. 1a). The equivalent of a Bose-glass to superfluid crossover is thus expected for a nonequilibrium polariton condensate lattice in a disordered potential. This crossover occurs when the phase coherence length, which in a disordered Bose insulator grows with increasing density or strength of the Josephson coupling, [19–21] exceeds the overall size of the lattice. The Bose-glass to superfluid transition was first observed in Josephson junction arrays [22, 23] and it has recently been observed in thermally equilibrated cold atom lattices [24], but different features emerge for the nonequilibrium lattice.

Here, we optically trap 25 polariton condensates inside a 5×5 square lattice within a planar semiconductor microcavity [25, 26] (Fig. 1a). Optical trapping allows rapid and facile tuning of the lattice constant as well as the nearest-neighbor coupling strengths [27]. Due to

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residual weak disorder in the microcavity, polaritons at different sites condense at slightly different energies [28]. When the lattice constant is large, the nearest-neighbor coupling is small and we observe an unsynchronized (UNSYNC) phase evidenced by a broad peak in the momentum space corresponding to that of a single site condensate as well as a distribution of condensate energies, and no long-range order. However, as the nearest-neighbor coupling increases by reducing the lattice constant or increasing the condensate number density (for higher excitation powers), neighboring condensates collapse their energy detunings due to polariton-polariton and polariton-exciton nonlinearities, until they all synchronize (SYNC) resulting in an extended state with long-range order. This appears as progressive narrowing of the momentum space peak and appearance of sets of diffraction peaks signifying a phase-locked fluid with a single delocalized macroscopic wavefunction. Simulations show that disorder plays a key role in the crossover width and the decay of long-range order.

II. FORMATION OF TRAPPED CONDENSATES IN A DISORDERED POTENTIAL

We create polariton condensates by the nonresonant optical excitation of a microcavity composed of GaAs quantum-wells sandwiched between two distributed Bragg reflectors (DBR). The cavity top (bottom) DBR is made of 32 (35) pairs of Al$_{0.15}$Ga$_{0.85}$As/AlAs layers of 57.2 nm/65.4 nm. Four sets of three 10 nm GaAs quantum wells separated by 10 nm thick layers of Al$_{0.3}$Ga$_{0.7}$As are placed at the maxima of the cavity light field. The 5λ/2 (583 nm) cavity is made of Al$_{0.3}$Ga$_{0.7}$As. Photons in the cavity are strongly coupled to quantum-well excitons to form mixed light-matter bosonic polaritons. The quasi continuous wave pump is a single-mode Ti:Sapphire laser tuned to the first Bragg mode ∼100 meV above the polariton energy. A spatial light modulator is used to spatially pattern the pump beam into a square lattice. A 0.4 NA objective is used for imaging the pattern onto the sample. A cooled CCD and a 0.55 m spectrometer is used for imaging and energy resolving the emission.

The nonresonant excitation patterned in a square geometry [29] initially creates a plasma of hot electrons which then relax in energy and form bound excitons. The excitons in this ‘reservoir’ eventually relax to form polaritons. Polaritons are repelled from reservoirs generated at each pump spot due to repulsive exciton-polariton and polariton-polariton interactions. The resulting repulsive potential causes polaritons to roll off and gather at the minima of the square potential. Polaritons condense in the ground state by stimulated scattering once the density at each center surpasses a critical threshold and form a macroscopic state in each optical trap [25, 26, 30].

As the trapped condensate is moved across the semiconductor sample, the energy of the condensate varies by a standard deviation of ∼30 µeV (see Appendix A). This ∼30 µeV disorder potential is approximately 10% of the confining potential from the optical trap [26], which

![FIG. 1. Below threshold, unsynchronized and synchronized phases. (a) Schematic of condensation in optical lattice potentials (top panel), and synchronization of two energy-detuned condensates in a double-well potential (bottom panel). There is a nonzero energy detuning $\delta E$ at the UNSYNC phase but this vanishes at the SYNC phase as the Josephson coupling increase. (b-d) Real-space and momentum space emission of (b) uncondensed polaritons, (c) unsynchronized (localized), and (d) synchronized (delocalized) condensate lattice. Pump spots are marked by grey circles in (c,d) and visible by dark spots in (b). The schematic diagrams above the panels show the trapping potential $V(x)$, Josephson coupling $J$ and energies of the condensates $E_n$. Momentum space images are in logarithmic scale.](image-url)
remains relatively constant after condensation.

III. NARROWING OF MOMENTUM

Condensate lattices are created by patterning the optical excitation into a square lattice of \(6 \times 6\) Gaussian spots (marked by grey circles in Fig. 1c,d) using a spatial light modulator. The lattice constant \(a\) of the trapping potential can be continuously varied while monitoring the total emission of the condensate lattice in momentum space (see also Supplemental Movie 1). At low excitation powers incoherent polaritons confined by the highest points of the lattice potential (dark spots in Fig. 1b) are created with a very broad momentum distribution (full width half max = \(\delta k \simeq 2.1\ \mu m^{-1}\), \(k\) is the in-plane polariton momentum). At a threshold power \(P = P_{th}\) polaritons condense in each lattice site and \(\delta k\) drops sharply. At threshold there is a condensate at each site and the zero-momentum peak has a width similar to that of the spatially filtered single condensate, \(\delta k_1\). This is because there is no long-range phase correlation and the lattice is in the unsynchronized phase. However, as power increases further, \(\delta k\) gradually decreases until it plateaus to \(\delta k \approx \delta k_1/4\), at \(P \approx 2P_{th}\). This value again matches with a 5-slit grating (see SI.B), indicating long-range phase correlation of a delocalized fluid across the entire lattice.

IV. BUILDUP OF LONG-RANGE COHERENCE AND COLLAPSE OF ENERGY DETUNING

The gradual decrease in momentum width \(\delta k\) with power implies that phase correlations in the lattice continuously increase with power as the coherence length of the order parameter \(\psi\) gradually increases. It has been shown that in a driven-dissipative condensate with disorder, the phase correlation length \(\mathcal{L}_\phi\), over which the first order correlation function drops by \(1/e\), increases with Josephson coupling and condensates density \(|\omega|\). To confirm this we measure the spatial first-order coherence function \(g^{(1)}\) at zero time delay. The lattice emission is sent to a modified Mach-Zehnder interferometer with a retroreflector in one arm. This interferes emission from opposite points relative to the central condensate (placing the origin at the white dashed circle in Fig. 3a) so that emission from \(r\) interferes with \(-r\). The contrast of these interferograms then gives

\[|g^{(1)}(r, -r)| = |g^{(1)}(-r, r)|\]

with \(\mathcal{L}_\phi\) being the phase co-

FIG. 2. Synchronization phase diagram. Dependence of \(\delta k \cdot a\) versus (a) lattice constant, and (b) power. Top panel in (a) shows real-space and momentum-space intensity (log scale) at \(a = 7.8\ \mu m\) (left) and \(a = 11.8\ \mu m\). (b) Power dependence comparison of \(\delta k \cdot a\) for full lattice (dark blue) and a spatially filtered single condensate (light blue). Real-space scale is 10 \(\mu m\) and momentum space scale is 1 \(\mu m^{-1}\).
FIG. 3. First order correlation function. (a) Real-space interferograms for the unsynchronized (top) and synchronized (bottom) phases. (b,c) Power dependence of $g^{(1)}(r,−r)$ at various lattice position $r$, as a function of total lattice power in (b) experiment and (c) simulations. In the experiment, an average is taken over six site separations marked A–F and their corresponding mirrors in (a), with separations of $(0.5, 2, \sqrt{2}, 4, 2\sqrt{2}, 4\sqrt{2})a$. In simulations, the average is extracted for all site separations in the lattice. Lines are exponential fits. Panels to the right of (c) are the time-averaged momentum space emission for UNSYNC ($P = P_{th}$) and SYNC ($P = 2P_{th}$) phases. The simulated disorder ratio is 10%. (d) Coherence length ($\xi$) vs power in simulations (with and without 10% disorder) and in experiment. Simulation curves are averages of 25 randomly generated disorder potentials.

FIG. 4. Condensate energies as a function of power. (a) Energy spectra of a column of 5 condensates crossing the center of the lattice for $P = P_{th}$ (top panel) and $P = 2P_{th}$ (bottom panel). (b) Standard deviation of condensate energies ($\sigma(E)$) as a function of power (light blue) and the corresponding $\delta k$-a (dark blue). Site energies of condensates in the synchronization crossover reveals the reason behind this behavior. At threshold, condensates form at slightly different energies (Fig. 4a). The energy distribution observed in the UNSYNC phase is a result of spatial inhomogeneities in the sample. However, as the power increases the condensate energies converge as long-range order propagates across the lattice, and at the SYNC phase they condense to one energy (Fig. 4b).

Our system can be modelled using the stochastic driven-dissipative Gross-Pitaevskii (GP) equation which is a nonlinear Schrödinger equation including pumping, dissipation, energy relaxation and Langevin noise to describe fluctuations:

$$i\hbar \frac{d\psi}{dt} = \left( -\frac{\hbar^2 \nabla^2}{2m} + V_0 + g_R n + g_P P + \alpha |\psi|^2 \right) \psi,$$

$$+ \frac{i}{2}(h R n - h\gamma) \psi - i\hbar \Lambda \psi + h\sqrt{\gamma + R\hbar} \frac{dW}{dt}$$

Here, $\psi$ is the wavefunction, $n$ is the reservoir density, $m$ is the polariton mass, $R$ is the scattering rate from the reservoir to the condensate, $\gamma$ is the polariton decay rate, $g_R$ and $g_P$ describe energy repulsion due to the presence of the reservoir and pump, $\alpha$ is the strength of polariton-polariton interactions, $P$ is the non-resonant pumping rate, $\gamma_R$ is the reservoir decay rate and $\Lambda$ is the damping rate.

V. THEORETICAL MODEL AND NUMERICAL SIMULATIONS
FIG. 5. Time-averaged momentum space, real space and phase of an unsynchronized lattice (a-c) and synchronized lattice (d-f). (g) Static potential accounting for sample inhomogeneity. (h) \(g^{(1)}\) versus length for UNSYNC (blue) and SYNC phase (red). (i) Power dependence of density (light blue) and momentum width (dark blue) for a synchronized lattice (solid line) and a single site (dotted line). (j) Power dependence of \(g^{(1)}\) for different lattice lengths.

A is a spatial dependent function matching the pump profile, which accounts for a phenomenological energy relaxation. Langevin noise is given by \(dW\) which is a complex Gaussian random variable characterized by correlation function \(\langle dW^* dW \rangle = dt\). Here, we also add a static disorder potential \(V_0\) accounting for the sample inhomogeneity. Disorder ratio is defined as \(\text{DOR} = V_0/(g_{\text{r}} n + g_P + \alpha |\psi|^2)\). The first order correlation function is given by:

\[
g^{(1)}(\mathbf{r}, -\mathbf{r}) = \frac{\langle \psi^*(\mathbf{r}) \psi(-\mathbf{r}) \rangle}{\sqrt{\langle |\psi(\mathbf{r})|^2 \rangle \langle |\psi(-\mathbf{r})|^2 \rangle}}. \tag{3}
\]

Simulating the condensate lattice in a randomly generated disorder potential demonstrates the crucial role of disorder in the synchronization crossover, as shown in Fig. 5. [31] Here, the disorder ratio (DOR) defined as the mean ratio of a Gaussian-distributed disorder potential (Fig. 5g) to the confinement potential by the pump pattern is \(\sim 10\%\) (as in the experiment). The condensate lattice crosses from unsynchronized to synchronized as the lattice constant is shrunk, with similar signatures to the experiment. In the UNSYNC phase when \(a = 15\ \mu\text{m}\) [Fig. 5(a-c)], momentum-space peak is broad, condensates phases are unlocked and \(\mathcal{L}_\phi \simeq a\) (blue line in Fig. 5h). This changes in the SYNC phase as the lattice is shrunk to \(a = 9\ \mu\text{m}\): the momentum-space peak narrows and new diffusion peaks appear, condensates phase-lock [Fig. 5(d-f)], and \(\mathcal{L}_\phi\) goes beyond the lattice dimensions (red line in Fig. 5h). The power dependence of the SYNC lattice shows gradual narrowing of the momentum peak and buildup of long-range coherence [Fig. 5(i,j)].

We find that the power dependence of correlation function strongly depends on the spatial profile of the disorder potential. For this reason, \(g^{(1)}\) simulations are averaged for 25 randomly-generated disorder potentials, as shown in Fig. 3c. The general trend stays the same: in the UNSYNC phase \(g^{(1)}\) does not expand beyond a single site, whereas in the SYNC phase \(g^{(1)}\) builds up throughout the lattice with an exponential drop off (Fig. 3c,d) (see Appendix C for larger lattices). In the synchronized phase, the Josephson coupling is strong enough to overcome the disorder potential and phase-lock the condensates to form a single macroscopic state. Without static disorder this synchronization crossover is sharp and the coherence length is 3 orders of magnitude longer than in the experiment (Fig. 3d). The minimization of the width of this synchronization crossover with power can thus be used to search for an optical potential that minimizes the static disorder potential, which has applications in enabling using such arrays as simulators. We note that disorder is inherent to any supported array of condensates, suggesting these observations will be universal to all implementations.

VI. CONCLUDING REMARKS

We demonstrated a controllable crossover from unsynchronized to synchronized in a driven-dissipative polaron condensate lattice with a background disorder potential. The crossover occurs either by increasing the density of all condensates or the nearest-neighbor Josephson coupling. The synchronized (delocalized) phase is accompanied by the appearance of long-range order and narrowing of the central peak in momentum space, whereas in the unsynchronized phase order remains local, and the momentum space resembles the emission from a single condensate. Using simulations of driven-dissipative GP
equations, we find that in the absence of disorder a sharp synchronization transition with long-range spatial coherence is observed. However, the introduction of disorder results in a softer crossover from unsynchronized to synchronized regimes with finite range for the spatial coherence.

In thermalized cold atoms in a spatially disordered Bose-Hubbard potential, there is the compressible Bose-glass (insulator) crossover phase between the Mott insulator and superfluid phases [23, 24, 32, 33]. There, although the condensates share a global chemical potential there is no long-range spatial coherence. Instead, the system forms puddles or domains of coherent condensates, but there is no coherence between neighboring puddles. As the tunneling rate exceeds the disorder potential, the superfluid puddles coalesce until there exists a global superfluid with a spatial coherence length that exceeds the size of the system.

By contrast, in the nonequilibrium case studied here, in the crossover regime the puddles or domains have different chemical potentials, or emission energies. Furthermore, in the superfluid phase the spatial coherence has a finite range that increases with pump power and tunneling rate. Hence driven-dissipative generalization of superfluidity can only be observed in a system of finite size, as discussed theoretically in ref. [20]. Unless actively compensated [28], this may have consequences for the scale-up of simulators based on lattices of exciton-polariton condensates.

With our capability to optically compensate for disorder [28], it would be interesting to study the interplay between on-demand disorder strengths and the synchronization crossover, and their relation to the characteristic length-scales of the first-order spatial correlation. In addition to fundamental interest, synchronization of arrays of polariton lasers for example, would be advantageous for creating high-power density coherent sources at low pump density. Conventional laser diodes locked by injection coupling can synchronize only in a narrow range of parameters [34], due to their large carrier induced red-shift. By contrast, we demonstrate that with a carrier induced blue-shift, synchronization readily occurs, suggesting that polariton laser approaches may be highly profitable.

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Appendix A: Disorder potential

Background disorder potential $V_0$ is measured experimentally by moving a single trapped condensate over the sample and measuring its energy, as shown in Fig. 6. Standard deviation of disorder potential is $\sim 29 \, \mu eV$.

Appendix B: Phenomenological model

Let us assume a coherent superposition of Gaussian wavefunctions localized at each lattice point:

$$A \sum_{n,m} e^{-((x-na)^2+(y-ma)^2)/(2L^2)},$$

where $A$ is an overall amplitude, $a$ is the lattice constant, and $L$ defines the width of each Gaussian. In reciprocal space the wavefunction is given by the Fourier transform:

$$\tilde{\psi}(k_x, k_y) = \frac{1}{2\pi} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \psi(x, y) e^{i(k_x x + k_y y)} dx dy = AL^2 e^{-(k_x^2 + k_y^2)L^2/2} \sum_{n,m} e^{i\alpha(k_n n + k_m m)}.$$
The intensity in reciprocal space is obtained taking the Fourier transform of each Gaussian spot separately, and summing the intensities rather than the amplitudes:

$$\tilde{I}(k_x, k_y) = NM|A|^2 L^4 e^{-(k_x^2 + k_y^2)/L^2}.$$  \hspace{1cm} (B4)

Since there is no coherence between Gaussian spots, the interference term appearing in Eq. B2 is no longer present in Eq. B4. The sum over n and m has been reduced to the total number of spots, NM, in the square lattice. The corresponding intensity of the wavefunctions in the momentum space is shown in Fig. 7c. This model obviously cannot describe why the crossover is not sharp and why the coherence length decays over length. For that we model the system using a stochastic driven-dissipative GP equation.

We note here the similarities between the synchronized lattice intensity profile, and Fraunhofer diffraction from many slits given by [35]:

$$I = I_0 \left( \frac{\sin \beta}{\beta} \right)^2 \left( \frac{\sin N\alpha}{\sin \alpha} \right)^2,$$  \hspace{1cm} (B5)

where $\beta = \frac{1}{2} k_0 b$, $\alpha = \frac{1}{2} k_0 a$. Here, $a$ is the slit separation, $b$ is the slit width, $N$ is the number of slits and $I_0$ includes all the constants to give maximum intensity from a single slit (Fig. 8). From Eq. B5, we can see that the first intensity minimum occurs at $ka = 2\pi/N$. For 5 slits, full width at half maximum of center peak is $\delta k \cdot a \simeq 1.13$.

### Appendix C: Large lattice simulations with disorder

The dependence of first order correlation function with the lattice constant for a $15 \times 15$ lattice in a random disorder potential ($DOR = 10\%$) shows an exponential decay with a coherence length $L_{\parallel}$ which grows as the lattice constant $a$ reduces, as shown in Fig. 9.

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