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Parity bifurcations in trapped multistable phase locked exciton-polariton condensates

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We present a theoretical scheme for multistability in planar microcavity exciton-polariton condensates under nonresonant driving. Using an excitation profile resulting in a spatially patterned condensate, we observe organized phase locking which can abruptly reorganize as a result of pump induced instability made possible by nonlinear interactions. For $\pi/2$ symmetric systems this reorganization can be regarded as a parity transition and is found to be a fingerprint of multistable regimes existing over a finite range of excitation strengths. The natural degeneracy of the planar equations of motion gives rise to parity bifurcation points where the condensate, as a function of excitation intensity, bifurcates into one of two anisotropic degenerate solutions. Deterministic transitions between multistable states are made possible using controlled nonresonant pulses, perturbing the solution from one attractor to another.

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I. INTRODUCTION

A microcavity exciton-polariton is a bosonic quasiparticle that arises from the strong coupling between a quantum well exciton and a cavity photon [1]. Due to their bosonic statistics and short lifetimes, polaritons can form a nonequilibrium analog of a Bose-Einstein condensate at high nonresonant excitation intensities where stimulated scattering results in large coherent population balanced by gain and decay. Their ability to interact strongly with themselves gives rise to a $\chi^{(3)}$ Kerr-like nonlinearity which in turn, for resonant excitation schemes on scalar condensates, gives rise to optical bistability [2–5], spatial multistability [6], and multistability for vectorial (spin dependent) condensates [7–9].

Recently, ultrafast [10,11], ultralow-energy [12], and nonlinear relaxation [13] switching mechanisms were realized experimentally between bistable states of polariton ensembles and suggest their future application as optical memory elements. Bistability is also a precursor for a number of effects in exciton-polariton systems, including the formation of solitons [14–16], imprinting patterns and spatial images [17–20], screening disorder [21], sustaining superfluid propagation [22], realizing quasicompactons [23], simulating cellular automata [24], realizing emergent descriptions of (classical) Ising models [25], and realizing polaritonic circuits [26].

While optical multistability with resonant excitation schemes has been the subject of intensive research [2–9], optical multistability in nonresonantly excited microcavity polaritons is less well studied. Physically, the nonresonant pump scheme is advantageous since a resonant pump scheme is implemented with an external laser, but nonresonant pumping can be achieved with electrical contacts on both sides of the quantum well cavity to spur the creation of excitons, which is far more compact to implement in optoelectronic circuits. To date, theoretical schemes for incoherently excited bistability based on strong saturated absorption [27], thermally induced changes [28], modulational instability [29], and gain competition between symmetric and antisymmetric condensate modes [30] have been proposed. Fairly recently, nonresonant optical bistability was demonstrated in polariton condensates under electrical injection of charge carriers [31,32] and between spin polarizations in annular pumping geometries [33].

In this paper, we investigate the appearance of optical multistability in a planar two-dimensional (2D) exciton-polariton condensate under spatially patterned nonresonant pumping. In Ref. [30] optical bistability has been shown theoretically to arise due to the modulational instability between the parities of a localized one-dimensional (1D) polariton condensate pattern, each bistable state corresponding either to an antisymmetric or symmetric macroscopic wave function. Consequently, it stands to reason that multistability can be observed in higher dimensional geometries due to the increased degeneracy of the linear part of the equations of motion.

The multistability relies on pump induced parity cross-saturation previously considered in 1D exciton-polariton condensates [30,34]. The symmetry of the dynamical equations results in a condensation threshold belonging to a definite parity of the system due to optimal constructive interference between same-parity modes (phase locked condensates). This result is well studied in the field of nonlinear optics [35] and in agreement with recent experimental observations where different excitation geometries result in exciton-polariton condensates being phase locked either in-phase or antiphase [36–42]. Increasing the pump intensity beyond threshold results in an unstable condensate which rapidly transitions into a new solution of different parity, corresponding to a reorganization of phase lockings within the system. Sweeping backward in pump intensity, several hystereses are recovered between the condensate parities. Within these hystereses we

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find that regimes of multistability between symmetric and antisymmetric condensate solutions exist.

We classify a point of instability where an isotropic condensate transitions into one of two anisotropic degenerate condensate solutions with equal probability, a feature which can be regarded as parity bifurcation. Very recently, bifurcating points were observed in spinor polariton condensates where, also as a function of pump intensity, a linearly polarized condensate transitions to either of two circularly polarized states [43].

After characterizing the possible multistable regimes and states therein, we proceed to demonstrate a method to induce controlled, reversible transitions within a group of stable states based on the application of nonresonant excitation pulses.

II. CONDENSATE PHASE LOCKING

Eigensolutions of spatially symmetric Hamiltonians are classified as either symmetric (even) or antisymmetric (odd). For nonlinear equations of motion this is no longer necessarily the case. Regardless, phase locking between spatially separate condensates [37–39] still takes place, made possible through their interactions, where condensates form either symmetric or antisymmetric states due to constructive interference of polaritons traveling from one condensate to another. Due to the short polariton lifetime, there naturally exists only a finite range of distances between the two condensates to coherently phase lock before traveling polaritons decay away.

We consider a nonequilibrium condensate of 2D planar exciton-polaritons described by the driven-dissipative nonlinear Schrödinger equation. Standard heuristics dictate that the feeding of polaritons into the condensate is controlled by active-exciton reservoir rate equations [44]. Under the assumption of a fast active-exciton reservoir dephasing rate and low polariton densities the equation of motion for lower condensate solutions with equal probability, a feature which points reported are not sensitive to naturally occurring disorder in planar cavities, driving field inhomogeneities [34], and energy relaxation due to condensate interactions with the exciton reservoir.

In order to illustrate the phase locking of different condensates we begin by simulating an open system with \( V(r) = 0 \) and \( P(r) \propto P_0 \) corresponding to a hexagonal arrangement of Gaussian pump spots with average intensity \( P_0 \) (see Fig. 1). If the distance between the spots is large enough, no phase locking takes place and each condensate forms with its own phase distribution. When the spots are brought close together, a new condensation threshold of the system belongs to phase locked condensates (see Fig. 2). Here we define a phase locking order parameter, or parity intensity, as follows,

\[
n_e = \left| \int_A \psi(r) dr \right|^2, \tag{2}
\]

\[
n_o = \left| \int_A \psi(r) \frac{x_1 x_2 x_3}{|x_1 x_2 x_3|} dr \right|^2, \tag{3}
\]

where \( A \) is the planar system size and \( x_1, x_2, \) and \( x_3 \) are anticlockwise rotated \( x \) axes by an angle \( \pi/6, \pi/2, \) and \( 5\pi/6 \), respectively (see Fig. 1). Here “\( e \)” and “\( o \)” stand for even and odd parity respectively. As pump intensity is increased the condensates reorganize their phase locking. This type of instability can be regarded as a parity transition where a condensate with a well defined parity is overcome by an opposite sign parity state. High density condensates in such geometric configurations also display vortex-antivortex lattice formation [37] which can be seen in the central region of Fig. 2(g). In what follows we will attempt to shed light on the mechanism of these sudden transitions.

In order to simplify the numerical experiment, and get a clearer understanding on the mechanism causing reorganization of phase locked condensates, we choose an infinite

![FIG. 1. Normalized pump profile P(r) of six Gaussian spots arranged into a hexagon with nearest neighbor distance 20 \( \mu \text{m} \). Three axes of symmetry between pump spots are plotted with white dashed lines and denoted \( x_1, x_2, \) and \( x_3 \).](image-url)
choosing a basis satisfying the following real linear part of $g$

We set the dependent order parameter in the basis of the linear eigenstates:

by arrows in panel (a). Here we set $(e)–(g)$ plots show the corresponding solution at the points indicated of a hexagonal pattern of Gaussian pump spots under slow increasing $0$. Sudden transitions between different parity solutions take place at high pump power. Normalized density $(b)–(d)$ and phase $(e)–(g)$ plots show the corresponding solution at the points indicated of $V(r) = 0$.

quantum well (IQW) of width $L$ as our system:

Here, separation of variables allows us to write the eigenstates of $\hat{H}_0$ in the form $|\phi_{nx,ny}\rangle = |\phi_{nx}\rangle \otimes |\phi_{ny}\rangle$ where $|\phi_{nx}\rangle$ and $|\phi_{ny}\rangle$ are the eigenstates of the 1D IQW. The solutions in the coordinate basis are written

where

Similarly to the analysis in Refs. [30] and [34] we write our time dependent order parameter in the basis of the linear eigenstates:

Here, $\hbar \omega_{nx,ny}$ are the eigenenergies of Eq. (5). For brevity we will write the indexing as $n = (n_x, n_y)$ where $n$ is a unique quantum number for each state. Inserting Eq. (9) into Eq. (1) and integrating out the spatial dependence we arrive at a coupled set of dynamical equations:

Here,

and

It becomes immediately clear that if $P(x, y) = P(x, y)$ then only states $\phi_{nx,ny}$ of the same parity structure along $x$ and $y$ coordinate are coupled by the pump gain mechanism [Eq. (11)]. As an example, the state $\phi_{22}$ does not contribute to the gain of $\phi_{11}$ but does however affect its decay through the nonlinear saturation $R$. It then becomes evident that the condensate threshold belongs to a superposition of same-parity eigenstates $c_x$. Furthermore, it is clear that degenerate states such as $\phi_{12}$ and $\phi_{21}$ evolve equivalently. For large separation distances between pump spots the condensate formation can be said to be uncoupled since the polariton decay rate $\Gamma$ will destroy any coherent overlap between the two condensates. Numerically, the coherence of weak polariton waves is destroyed by introducing stochastic terms to the equations of motions (not shown explicitly here). In this paper we consider only small distances where condensates do interfere coherently.

The last term of Eq. (10) is analogous to polarization coupling of electromagnetic waves in media with $\chi^{(3)}$ Kerr nonlinearity. The elements $M_{nkjl}$ only depend on the choice of basis to represent the condensate order parameter. Evidently, since $\phi_{nx,ny}$ are representable as real functions, the order of the indices in $M_{nkjl}$ does not matter but it does matter for the evolution of the amplitudes $c_m$ [Eq. (10)] which are coupled through the sum of products $\sum_{m} M_{nkjl}^{*} c_m^* c_k c_l$, mixing the phases of different parities. It becomes clear that without nonlinearity the condensate cannot be stable above threshold as it will diverge; it is also clear that the nonlinear term is the only one responsible for causing the instability points shown in Fig. 2(a) where one condensate solution is replaced by another. More interestingly, only the imaginary nonlinear term ($R$) is enough to produce these unstable points [34].

The nonlinear problem is therefore highly nontrivial and searching for multistable solutions relies heavily on numerical modeling. However, a qualitative result on pump-induced, parity-dependent instability was given in Ref. [34] where the critical instability point was demonstrated analytically for a two-mode system. The full quenching (parity transition), as shown in Fig. 2(a), where the green colored curve is replaced by red and vice versa, is then made possible when the critical point gives rise to a solution capable of driving the previous solution to zero, a feature made possible through the nonlinear saturation. The reorganization of phase locked condensates
shown in Fig. 2 can therefore be regarded as parity transitions within a single spatially modulated condensate. The transition is not only characterized by a flip in parity but also a shift in both real and imaginary energies of the condensate which manifests itself in sudden shifted chemical potential, and total polariton population defined as $N = \int_A |\psi|^2 d\mathbf{r}$. The transition does therefore not occur between degenerate solutions. In Ref. [30] the shift in $N$ was too small to be observed and remained unreported.

Furthermore, since the gain of the condensate depends on the condensed mode in question through the saturation term $R|\psi|^2$, it has been shown that a hysteresis can be recovered when the pump intensity is swept below the point of instability, evidencing parity bistability [30]. Here, in contrast to bistable 1D states, the degeneracy of the 2D linear equations of motion results in multistability.

### III. RESULTS

We continue our analysis in the IQW system. We work with the following pump profile:

$$P(\mathbf{r}) = P_0 |\phi_{22}(\mathbf{r})|^2.$$  \hspace{1cm} (13)

Here, $P_0$ represents the average nonresonant driving field intensity. Similarly to Eqs. (2) and (3) we define new parity intensities for each possible parity structure along the $x$ and $y$ axes of the system:

$$n_{ee} = \left| \int_A \psi(\mathbf{r}) d\mathbf{r} \right|^2,$$

$$n_{eo} = \left| \int_A \psi(\mathbf{r}) \frac{\partial \psi(\mathbf{r})}{\partial y} d\mathbf{r} \right|^2,$$

$$n_{oe} = \left| \int_A \psi(\mathbf{r}) \frac{\partial \psi(\mathbf{r})}{\partial x} d\mathbf{r} \right|^2,$$

$$n_{oo} = \left| \int_A \psi(\mathbf{r}) \frac{\partial^2 \psi(\mathbf{r})}{\partial x \partial y} d\mathbf{r} \right|^2.$$  \hspace{1cm} (14)

When $\psi$ is in a superposition of modes of only one parity structure then only one of the above integrals is nonzero. We note that the choice of driving field $P(\mathbf{r})$ is nonexhaustive and Eq. (13) is only one example of many to create a spatially patterned condensate and to produce similar results in this paper.

#### A. Multistability

In Fig. 3(a) we show the results of sweeping adiabatically forward in pump intensity. As expected, we observe the clear phase locking upon condensation corresponding to $n_{ee}$ (symmetric state) as the initial dominant parity structure. Increasing the pump intensity further we observe recurrent drops in the parity intensities of the condensate quickly replaced by another solution of different parity, corresponding to parity cross-saturation analogous to the observed drops in Fig. 2. In Fig. 3(b) we sweep backwards in pump intensity (colored markers) and find several hysteresis intervals indicating the bistable nature of the system.

As an example, at $P_0/P_{\text{cond}} = 2.75$, the system can exist in at least two steady states, separately characterized by having $n_{ee}$ (green circles) and $n_{oo}$ (red whole curve) as the dominant parity intensities. These states are analogous to the observations of Ref. [37] except without vortex lattices [see Figs. 4(a), 4(d)], classifying them as ferromagnetic (FM) and antiferromagnetic (AFM) ordered states in phase. However, two degenerate anisotropic states can also exist which were reported as 90° compass states [46] due to their two possible perpendicular orientations, and correspond to the blue and magenta colored curves and markers in Fig. 3 [also Figs. 4(b), 4(c)]. It is clear that these states are interchangeable since the $n_{eo}$ and $n_{ee}$ parity structures are one and the same under $\pi/2$ rotation of the system. Nonadiabatic statistics (instant pump activation) on the formation of the reported solutions are given in Appendix B.

Sweeping forward and backward in pump intensity reveals bistable intervals but does not elucidate the multistability of the system. By the virtue of symmetry between $x$ and $y$ it is clear that if $n_{ee}$ and $n_{oo}$ can exist at the same pump power, then $n_{eo}$ and $n_{oe}$ are also possible stable states. This is verified numerically by activating $P_0$ instantaneously and rapidly condensing the system from a stochastic initial condition, resulting in one of the four different possible stable states (see Fig. 4). Density and phase plots of the four different multistable states are shown in the upper and lower rows of panels in Fig. 4, respectively.

A fifth state appears when slowly sweeping backward in pump intensity (see Fig. 5) showing a vortex in the system center and equal population in both $n_{eo}$ and $n_{ee}$ parity intensities (overlapping blue and magenta markers in Fig. 3). Vortex states, being one of the fundamental solutions of the nonlinear Schrödinger equation, naturally occur in driven-dissipative condensates.

#### B. Parity bifurcation

Earlier it was stated that the $n_{ee}$ and $n_{oo}$ solutions evolve in an indistinguishable manner due to the symmetry of the
system. We verify this by averaging over 1000 stochastic trials (Monte Carlo) simulating the evolution of the condensate as a function of pump power up to the point of instability (bifurcation point) where the condensate transitions to either an \( n_{eo} \) or \( n_{oe} \) state. The trials are stochastic by introducing white noise field to the order parameter at much smaller time steps than the polariton lifetime, keeping any uncondensed states completely stochastic in evolution. In Fig. 6 the average parity intensity drops from the \( n_{ee} \) state and populates either an \( n_{eo} \) or \( n_{oe} \) state with approximately equal probability. Such process can be classified as parity bifurcation.

We stress that the critical transition in Fig. 6 depends on the adiabaticity of the pump. It is clear that if the \( n_{eo} \) and \( n_{oe} \) solutions become unstable and condense then \( n_{oo} \) should also be unstable. Stochastic fluctuations can not only seed the odd parity along one of the system axes, resulting in an \( n_{eo} \) or \( n_{oe} \) state, but also seed odd-parity states along both \( x \) and \( y \) axes close enough in time that an \( n_{oe} \) state is recovered. Such a process is however only likely if the adiabaticity of the pump is relaxed (faster ramping rates). The parity bifurcation naturally vanishes when disorder is introduced, corresponding to breaking of the \( \pi/2 \) invariance of the system, making one of the two states deterministically dominant. We stress that the influence of disorder only breaks the equivalence of \( n_{oe} \) and \( n_{oe} \) states without affecting multistable regimes between \( n_{ee} \), \( n_{eo,oe} \), and \( n_{oo} \) states.

We have only demonstrated the bifurcating point as a function of pump intensity which does not distinguish between the paths taken going to either \( n_{eo} \) or \( n_{oe} \) states. There are however other methods which can be used to control the path from the \( n_{ee} \) state to either \( n_{eo} \) or \( n_{oe} \) states by, e.g., changing the geometry of the real and complex potential landscapes. In an animation in the Supplemental Material [47] this is is demonstrated by changing the square well potential and pump shape from square to rectangular.

C. Controlled transitions between multistable states

A sweep of the pump intensity can be utilized to switch between different states as is evident in Fig. 3 but a more pragmatic method relies on controllably perturbing the system to deterministically change between states. By applying short nonresonant pulses to create pressure gradients within the condensate, controlled transitions between the four steady

![Figure 4](image1)

**FIG. 4.** Four steady states within the square trap achieved using Eq. (13) at instantaneous activation of the excitation power \( P_0/P_{\text{cond}} = 2.75 \).

(a)–(d) Normalized condensate density and (e)–(h) phase corresponding to \( n_{ee}, n_{eo}, n_{oe}, \) and \( n_{oo} \) steady states, respectively.

![Figure 5](image2)

**FIG. 5.** (a) Normalized density and (b) phase color maps showing a vortex state appearing in the center of the trap when sweeping backward in excitation intensity (overlapping blue and magenta markers in Fig. 3).

![Figure 6](image3)

**FIG. 6.** Average normalized parity intensity of the trapped condensate over 1000 stochastic trials. As the \( P_0 \) is adiabatically increased the condensate bifurcates into one of two possibilities corresponding to states \( n_{eo} \) and \( n_{oe} \).
states can be induced. We will demonstrate that the transitions (or steps for brevity) based on these nonresonant pulses can be made from any of the four steady states to any of the other steady states. Therefore any path made up of these steps is completely reversible.

We present here three possible steps (not counting backward steps), namely $n_{ee} \leftrightarrow n_{eo}$, $n_{ee} \leftrightarrow n_{oe}$, and $n_{ee} \leftrightarrow n_{oo}$. With this set of steps, it is possible to transit between all of the four states with at most two steps. The pump pulse is simulated by replacing the $iP(r)$ term with $i[P(r) + P'(r,t)]$ where $P'$ is a short pulse superimposed upon the static excitation,

$$P'(r,t) = P_0' e^{-(\frac{t - t_0}{\delta t})^2} P(r)_{(A,B,C,D)},$$

(15)

where $P_0'$ determines the intensity of the pulse, $t_0 = 300$ ps is the pulse delay, and $\delta t = 20$ ps determines the temporal length of the pulse. For simplicity, we choose the following scheme of pulsing:

$$p(r)_{(A,B,C,D)} = A \frac{(1 - x/|x|)(1 + y/|y|)}{4} + B \frac{(1 + x/|x|)(1 + y/|y|)}{4} + C \frac{(-1 + x/|x|)(-1 + y/|y|)}{4} + D \frac{(1 + x/|x|)(1 - y/|y|)}{4}. \quad (16)$$

Here $x,y \in [-L/2,L/2]$ and $A,B,C,D \in \{0,1\}$. The function $P'(r,t)$ corresponds then to a short duration of added uniform gain in the chosen quadrants of the quantum well. The choice of which quadrants are to be excited is given by the binary control variables $A,B,C,$ and $D$ corresponding to a pulse being
activated in the top right, top left, bottom left, and bottom right quadrants of the square well, respectively. We stress that the number of methods of perturbing the system is nonexhaustive and the one given by Eq. (16) is just one of many possibilities.

Three pulse profiles are employed, \( p_{(100)} \), \( p_{(110)} \), and \( p_{(101)} \) for the \( n_{ee} \leftrightarrow n_{oo}, n_{ee} \leftrightarrow n_{eo}, \) and \( n_{ee} \leftrightarrow n_{oo} \) transitions, respectively. The results are presented in Fig. 7. Let us analyze one transition as an example. The steps \( n_{ee} \rightarrow n_{oa} \) and \( n_{eo} \rightarrow n_{ee} \) [see Figs. 7(a), 7(b)] are both induced using a \( p_{(100)} \) profile. This is a logical choice of pump pulse profile since the condensate is being perturbed in the \( x > 0 \) quadrants, evolving differently from the rest of the condensate at \( x < 0 \). The perturbing dynamics are thus symmetric about the \( x \) axis but encourage asymmetry about the \( y \) axis. In Fig. 7(a) we see that the \( n_{ee} \) state decays much slower as opposed to the decay of the \( n_{oo} \) state in Fig. 7(b) indicating that the transition dynamics are distinguishable. Numerical simulations over long times reveal that the \( n_{ee} \) intensity eventually decays down to the white noise level of the simulation.

The other two transition pairs (\( n_{ee} \leftrightarrow n_{oo} \) and \( n_{ee} \leftrightarrow n_{oo} \)) can be induced in a fashion similar to that of the \( p_{(1100)} \) and \( p_{(1010)} \) pump profiles, respectively. Our system therefore possesses a cyclical transition scheme that allows for controlled transitions between all four steady states.

**D. Effects of pump induced blueshift**

We shift our attention to the case when \( g \neq 0 \). The real potential \( gP(\mathbf{r}) \) alters the energy landscape of the system but is determined by the symmetry of the pump, so only same-parity structures are still coupled together in both energy and gain coming from the pump [see Eq. (11)]. Mixing between different parity structures is therefore still only realized through the system nonlinearity which is unchanged. Previous results are thus only quantitatively changed with all parity related effects still present. This is demonstrated in Fig. 8 where for realistic values of pump blueshift the points of parity transition are shifted along the \( P_0 \) axis. We define the blueshift parameter as \( g_F = \max(g P(\mathbf{r})) \) for \( P_0 = 2P_{\text{cond}} \), that is, the maximum energy shift at twice the threshold intensity.

**FIG. 8.** Forward sweep of \( P_0 \) for different values of \( g \). Green, blue, and red curves correspond to \( n_{ee}, n_{oo,ee}, \) and \( n_{oo} \) parity intensities, respectively. The value \( g_F \) corresponds to maximum blueshift at \( P_0 = 2P_{\text{cond}} \) where \( P_{\text{cond}} \) is defined as the condensation value for \( g = 0 \).

**E. Generalization to symmetric pump and trapping geometries**

The above results are naturally not exclusive to the IQW and can mostly be replicated using other symmetric trapping geometries \( V(\mathbf{r}) \) or driving-field profiles \( P(\mathbf{r}) \) such as the hexagonal pattern shown in Fig. 1. In fact, any real potential and driving field in the planar system which are symmetric about the \( xy \) coordinates will never mix together the different parity structures. As an example, the harmonic oscillator is a cylindrically symmetric potential and does not distinguish the \( x \) and \( y \) coordinates. When the system is driven by a symmetric nonresonant field \( P(\mathbf{r}) = P_0|\psi_{22}(\mathbf{r})|^2 \) parity transitions analogous to those shown in Fig. 3 are observed in the polariton condensate (see Fig. 9).

We also demonstrate certain excitation schemes which do not show the sudden change in parities of the system. It is clear that the competition between the parities to dominate the condensate ultimately depends on how the real and complex potentials couple together same-parity modes, and subsequently how the modes interact through the nonlinearity of the condensate pattern. Clearly some potentials will favor this conflict between the parities as shown above, whereas others show no sudden transitions in the condensate parity. As an example, annular pump shapes are used to create cylindrically symmetric condensates [48] to investigate the onset of spontaneous currents [49], vorticity [50], and petal formation [51–53].

We point out that in Ref. [51] spatial ellipticity of the excitation beam breaks the degeneracy of symmetric and antisymmetric superpositions of clockwise and anticlockwise currents and stabilizes a standing wave (petal) pattern. In the same manner here, if the equivalence between
the circulating current [see Figs. 10(a)–10(c)]. In the absence of a single condensate with a well-defined phase corresponding to parity arrangements involving nearest neighbor bonds which, in the hexagonal geometry, can only form a steady state with all bonds being antisymmetric or symmetric to avoid frustration when accounting for only nearest neighbor interactions. For example, in Fig. 4 every solution has the same configuration of symmetric and antisymmetric bonds in each quadrant of the system, making them energetically equivalent. The symmetric and antisymmetric bonds can be regarded as a type of FM (0) and AFM (π) bond ordering among scalar [37] and spinor polariton condensates [54]. Figure 11 then shows an example of a lattice favoring a periodic arrangement of FM and AFM bonds. We note that a spin wave state was recently reported theoretically in a hexagonal graph of polariton condensates [55] which elucidates that nearest neighbor frustration can be balanced through interactions with more neighbors.

We point out that, as expected, a lattice of vortices forms within the center of the octagon just like in the hexagonal geometry.

**IV. CONCLUSIONS**

We have shown through extensive numerical simulation of the complex nonlinear Schrödinger equation that multistability in planar microcavity exciton-polariton condensates, with spatially patterned nonresonant pumping, can be achieved between symmetric and antisymmetric solutions of the condensate pattern. Solutions of different parity structures compete for the system gain with ultimately one winner driving all other solutions to zero through the condensate saturation mechanism. Most notably, doubly degenerate condensate solutions give rise to bifurcating points where the condensate transitions to one of either degenerate state with equal probability at a critical pump power. Most of the results focus on π/2 symmetric linear equations of motion, such as the planar infinite quantum well, where the linear physics of the system is decomposable onto the Cartesian basis. But systems with a symmetry axis at angles < π/2 such as hexagons and octagons also display the same competition and quenching between symmetric and antisymmetric bonded states.

Within the multistable regime, the use of controlled nonresonant pulses allows deterministic switching between different forming symmetric and antisymmetric bonds along the lattice (see Fig. 11), a state not observed when using hexagonal lattices.

The reason for this observation stems from the fact that the energy and gain of each condensate depends on the number of nearest neighbor symmetric and antisymmetric bonds which, in the hexagonal geometry, can only form a steady state with all bonds being antisymmetric or symmetric to avoid frustration when accounting for only nearest neighbor interactions. For example, in Fig. 4 every solution has the same configuration of symmetric and antisymmetric bonds in each quadrant of the system, making them energetically equivalent. The symmetric and antisymmetric bonds can be regarded as a type of FM (0) and AFM (π) bond ordering among scalar [37] and spinor polariton condensates [54]. Figure 11 then shows an example of a lattice favoring a periodic arrangement of FM and AFM bonds. We note that a spin wave state was recently reported theoretically in a hexagonal graph of polariton condensates [55] which elucidates that nearest neighbor frustration can be balanced through interactions with more neighbors.

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The reason for this observation stems from the fact that the energy and gain of each condensate depends on the number of nearest neighbor symmetric and antisymmetric bonds which, in the hexagonal geometry, can only form a steady state with all bonds being antisymmetric or symmetric to avoid frustration when accounting for only nearest neighbor interactions. For example, in Fig. 4 every solution has the same configuration of symmetric and antisymmetric bonds in each quadrant of the system, making them energetically equivalent. The symmetric and antisymmetric bonds can be regarded as a type of FM (0) and AFM (π) bond ordering among scalar [37] and spinor polariton condensates [54]. Figure 11 then shows an example of a lattice favoring a periodic arrangement of FM and AFM bonds. We note that a spin wave state was recently reported theoretically in a hexagonal graph of polariton condensates [55] which elucidates that nearest neighbor frustration can be balanced through interactions with more neighbors.

We point out that, as expected, a lattice of vortices forms within the center of the octagon just like in the hexagonal geometry.

**IV. CONCLUSIONS**

We have shown through extensive numerical simulation of the complex nonlinear Schrödinger equation that multistability in planar microcavity exciton-polariton condensates, with spatially patterned nonresonant pumping, can be achieved between symmetric and antisymmetric solutions of the condensate pattern. Solutions of different parity structures compete for the system gain with ultimately one winner driving all other solutions to zero through the condensate saturation mechanism. Most notably, doubly degenerate condensate solutions give rise to bifurcating points where the condensate transitions to one of either degenerate state with equal probability at a critical pump power. Most of the results focus on π/2 symmetric linear equations of motion, such as the planar infinite quantum well, where the linear physics of the system is decomposable onto the Cartesian basis. But systems with a symmetry axis at angles < π/2 such as hexagons and octagons also display the same competition and quenching between symmetric and antisymmetric bonded states.

Within the multistable regime, the use of controlled nonresonant pulses allows deterministic switching between different forming symmetric and antisymmetric bonds along the lattice (see Fig. 11), a state not observed when using hexagonal lattices.
parity structures of the phase locked condensate pattern. Applying the same pulse repeatedly causes the system to alternate between two states. For such pairs of states, the pulse acts effectively as a NOT gate, paving the way towards controllable optical quantum fluid circuitry. Consequently, a possible future direction would be to consider multiple trapped condensates and attempt to engineer an AND gate. Together, the two gates form a universal set of logic gates for binary processing. Although this relies on a binary logic, a further extension can be to attempt to create a universal set of quaternary logic gates from multiple trapped/localized condensates.

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APPENDIX A: RESERVOIR MODEL

Equation (1) is valid for a static reservoir at low to moderate condensate densities. The above results can be reproduced using the more honest approach which models the saturation of the condensate through the hot excitonic reservoir $n_X$ using the following equations [44]:

\[ i \hbar \frac{\partial \psi}{\partial t} = \left[ -\frac{\hbar^2 \nabla^2}{2m} + V(\mathbf{r}) + \frac{i \hbar}{2} (R n_X - \Gamma) + \alpha |\psi|^2 \right] \psi, \quad (A1) \]

\[ \frac{\partial n_X}{\partial t} = - (\Gamma_R + R |\psi|^2) n_X + P(\mathbf{r}), \quad (A2) \]

where $\Gamma_R$ is the exciton reservoir average decay rate, and $V(\mathbf{r}) = V_0(\mathbf{r}) + g_X n_X$ where $g_X$ is the reservoir interaction strength. In Fig. 12 we show the parity transition taking place using Eqs. (A1) and (A2). Here $P(\mathbf{r})$ is a superposition of two spatially separated Gaussian spots creating two condensate maxima which interfere, creating a standing wave pattern between themselves. Slowly increasing the peak intensity of the Gaussians $P_0$ we observe the same parity transition as with Eq. (1). In Fig. 12(a) a symmetric state appears initially as a final steady state through stochastic initial conditions.

APPENDIX B: ATTRACTOR STATISTICS

The results in Fig. 3 are done under adiabatic conditions where the condensate wave function slowly evolves as a stationary solution of the equations of motion [Eq. (1)] with increasing excitation intensity $P_0$. In the nonadiabatic limit where the external excitation $P(\mathbf{r})$ is activated instantaneously with $\psi$ initially as a weak stochastic field the probability of retrieving a given solution depends on its attractor strength in solution space. Numerically this attractor strength can be extracted by Monte Carlo counts where each count is a new realization of stochastic initial conditions for $\psi$. In Fig. 13 we show the results of 70 counts for 70 different excitation strengths corresponding to Fig. 3. The stacked histogram shows regimes of coexistence between different solutions elucidating the multistability properties of the system. We point out that nonstationary solutions (cyclical) can appear as a result of many energies being occupied within the condensate.
