Energy barriers between metastable states in first-order quantum phase transitions

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A system of neutral atoms trapped in an optical lattice and dispersively coupled to the field of an optical cavity can realize a variation of the Bose-Hubbard model with infinite-range interactions. This model exhibits a first-order quantum phase transition between a Mott insulator and a charge density wave, with spontaneous symmetry breaking between even and odd sites, as was recently observed experimentally [Landig et al., Nature (London) 532, 476 (2016)]. In the present paper, we approach the analysis of this transition using a variational model which allows us to establish the notion of an energy barrier separating the two phases. Using a discrete WKB method, we then show that the local tunneling of atoms between adjacent sites lowers this energy barrier and hence facilitates the transition. Within our simplified description, we are thus able to augment the phase diagram of the model with information concerning the height of the barrier separating the metastable minima from the global minimum in each phase, which is an essential aspect for the understanding of the reconfiguration dynamics induced by a quench across a quantum critical point.

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I. INTRODUCTION

Quantum phase transitions are driven by a competition between different terms in the Hamiltonian, which lead to a rearrangement of the many-body ground state as a certain parameter is varied. Despite being known for many decades, this subject has seen a boom of interest in recent years, chiefly due to experimental progress in condensed-matter physics and ultracold atoms in optical lattices [1–5]. Theoretical and computational advances have also helped improve our understanding of the basic mechanisms underlying these transitions [6]. In addition to understanding the equilibrium properties of quantum phase transitions, considerable effort is currently put into understanding properties related to the relaxation towards equilibrium, for instance, after a quantum quench [7–10]. This requires one to understand how the system reconfigures itself from one phase to another as the critical point is traversed. In classical phase transitions, this reconfiguration is generated by thermal fluctuations, which induce a coarsening dynamics of nucleated domains [11–16]. In the case of quantum phase transitions, on the other hand, one obviously expects that thermal fluctuations can be neglected, the only available mechanism that allows the system to reconfigure from one phase to the other is the local tunneling of atoms between adjacent sites. The tunneling should therefore affect the energy landscape of the system, lowering the barrier separating the system must overcome an extensive energy barrier in order to traverse from one phase to another. A clear demonstration of this behavior was recently given in Refs. [17,18] for a model of interacting bosons in an optical lattice. The model consists of a variation of the standard two-dimensional Bose-Hubbard model [1,6,19,20] that includes a long-range interaction between atoms in the different checkerboard sublattices of a square lattice. It presents in total four quantum phases: superfluid (SF), supersolid (SS), Mott insulator (MI), and charge density wave (CDW). We shall concentrate, in particular, on the MI-CDW transition, which is of first order. Detailed theoretical investigations of this model were recently given in a variety of papers [21–27]. However, these papers were mostly concerned with the equilibrium phase diagram. The mechanisms which allow the transition between the MI and CDW phases and the description of the barrier separating these phases are crucial for further studies of the relaxation (quench) dynamics. The purpose of this paper is to shed light on these mechanisms, which should serve as a first step towards more complex nonequilibrium studies.

The MI and CDW phases [18] are illustrated in the insets in Fig. 1. In the former, the atoms tend to distribute uniformly throughout the square lattice, whereas in the latter they tend to aggregate into either the even or the odd sublattice. Assuming thermal fluctuations can be neglected, the only available mechanism that allows the system to reconfigure from one phase to the other is the local tunneling of atoms between adjacent sites. The tunneling should therefore affect the energy landscape of the system, lowering the barrier separating the
two phases. In this paper, we discuss how this effect occurs. We begin by considering the situation with no tunneling, in which case the phases can be described exactly by a Landau theory for the imbalance order parameter. Next we introduce the effects of tunneling using a variational ansatz based on a set of representative states between the MI and CDW phases. As we show, within this variational setting, we are able to study how the local tunneling affects the barrier between metastable states. Our main result is an equilibrium phase diagram augmented with information on the height of the energy barrier, shown in Fig. 5. The equilibrium phases we obtain match well with other analyses in the literature [23,25] and the information on the energy-barrier heights agree qualitatively with recent experimental results [18].

II. LANDAU THEORY

We consider here a bosonic system on a two-dimensional (2D) square lattice with $K$ sites, each site described by a bosonic operator $b_i$. We write the Hamiltonian as [17]

$$H = H_0 + \mathcal{T},$$

(1)

where

$$H_0 = \frac{K}{2} \sum_{i=1}^K U_i \hat{n}_i (\hat{n}_i - 1) - \frac{U_L}{K} \Theta^2$$

(2)

and

$$\mathcal{T} = -J \sum_{\langle i,j \rangle} (b_i^\dagger b_j + b_j^\dagger b_i).$$

(3)

In these expressions, $U_i$ is the short-range repulsive interaction, $J$ is the hopping parameter, $\hat{n}_i = b_i^\dagger b_i$, and $U_L$ is the long-range interaction. The sum in Eq. (3) is over all nearest neighbors in the square lattice. Moreover, we divide the lattice into a checkerboard structure consisting of even and odd sites, so that the nearest neighbors of an even site are always odd, and vice versa. Then, finally, the operator $\Theta$ appearing in Eq. (2) is defined as

$$\Theta = \sum_{\langle i \in e \rangle} \hat{n}_i - \sum_{\langle i \in o \rangle} \hat{n}_i,$$

(4)

which we refer to as the imbalance operator between the even (e) and odd (o) sublattices. In this paper, we work entirely in the canonical ensemble, with a fixed number of particles $N$. For simplicity, we also consider a spatially homogeneous system.

The long-range term $U_L$ introduces a long-range checkerboard interaction which favors an imbalanced occupation of the even and odd sublattices. When $U_L = 0$, we recover the usual Bose-Hubbard model [1,6,19,20], which presents a quantum phase transition between a SF and a MI. The presence of the checkerboard interaction introduces, in addition to these phases, a SS and a CDW phase. The MI and CDW phases appear when $J/U_L \ll 1$ and are thus essentially determined by the competition between $U_L$ and $U_L$. For $J \to 0$, the Hamiltonian becomes diagonal in the Fock basis and the partition function may be computed exactly at zero temperature, in both the canonical and the grand-canonical ensembles (ensemble inequivalence in long-range interacting systems forces us to explicitly specify the ensemble we work with in [28]).

The partition function in the canonical ensemble is

$$Z = \sum_{\text{states}(N)} e^{-\beta H_0},$$

(5)

where the sum is over all states $(n_1, \ldots, n_K)$ with the constraint that $\sum_i n_i = N$. Note that in this case, all operators in $H_0$ commute, so we do not need to differentiate between operators and their eigenvalues. We may write $Z$ as

$$Z = \sum_{\Theta = -N}^N \left\{ \sum_{\text{states}(N,\Theta)} e^{-\beta \mathcal{T}} \sum_{\langle i \in e \rangle} n_i (n_i - 1) \right\}. $$

(6)

Next we note that fixing $N$ and $\Theta$ is tantamount to fixing the occupations $N_e$ and $N_o$ of the even and odd sublattices, viz.,

$$N_e = \frac{N + \Theta}{2}, \quad N_o = \frac{N - \Theta}{2}.$$  

(7)

Thus we may write $Z$ as

$$Z = \sum_{\Theta = -N}^N e^{\beta U_L \Theta^2/K} \zeta(N_e) \zeta(N_o),$$

(8)

where

$$\zeta(N) = \sum_{\text{states}_e(N_e)} e^{-\beta U_L \sum_{\langle i \in e \rangle} n_i (n_i - 1)}.$$  

(9)

Here the notation $\text{states}_e(N_e)$ means a sum only over sublattice $x = e,o$ with a fixed number of particles $N_e$. Next we define $\theta = \Theta/K, \rho = N/K$, and $\rho_e = 2N_e/K$ and also the quantity $\phi(\rho_e)$ according to $\zeta(N_e) = e^{-\beta U_L / 2} \phi(\rho_e)$. Then, in the thermodynamic limit, the partition function may be written as

$$Z = \text{const} \int_{-\rho}^{\rho} d\theta \, e^{-\beta U_L \phi(\theta)}.$$  

(10)

with the Landau free energy

$$f(\theta) = -U_L \theta^2 + \frac{\phi(\rho + \theta) + \phi(\rho - \theta)}{2}.$$  

(11)
where we used the fact that $\rho_x = \rho + \theta$ and $\rho_o = \rho - \theta$. Thus, we conclude that in the limit of zero hopping ($J = 0$), the system is described exactly by a Landau theory, both at zero and at finite temperatures. This is, of course, not surprising given the mean-field character of the long-range interaction.

The quantities $\phi(\rho_i)$ may be determined exactly in the limit of zero temperature,

$$\phi(\rho_i) = \frac{U_i}{K} \min_{N_i} \left\{ \sum_i n_i (n_i - 1) \right\}. \quad (12)$$

The minimization is equivalent to a Bose-Hubbard model with no hopping, so the configuration which minimizes this quantity will be the one which is as close as possible to the Mott insulator. Let

$$N' = N - \frac{K}{2} |\rho_i|, \quad (13)$$

where $|x|$ is the floor function. Then there will be $N'$ sites with occupation $|\rho_i| + 1$ and the remainder with occupation $|\rho_i|$. Thus we get

$$\phi(\rho_i) = U_i |\rho_i| \left( \rho_i - \frac{1}{2} \right) \left( 1 + |\rho_i| \right). \quad (14)$$

Substituting this in Eq. (11) then yields the free energy as a function of the filling $\rho$ and the imbalance $\theta$.

Equation (11) simplifies considerably when $\rho = 1$. In this case, for $\theta \in [-1,1]$, it follows that

$$\phi(1 + \theta) + \phi(1 - \theta) = U_i |\theta|.$$

Thus we get

$$f(\theta) = -U_i \theta^2 + \frac{U_i}{2} |\theta|, \quad (15)$$

which is a remarkably simple expression. This result is shown in Fig. 1 for different values of $U_i / U_s$. As can be seen, the system presents three minima at $\theta = 0$ (MI) and $\theta = \pm 1$ (CDW). The global minimum changes at $U_i = U_s / 2$, signaling a first-order quantum phase transition.

An interesting point, which has important consequences for the phase reconfiguration, is that the MI minimum at $\theta = 0$ always exists, whereas the CDW minima at $\theta = \pm 1$ cease to exist when $U_i < U_s / 4$, as illustrated in Fig. 1. This asymmetry can manifest in physical observables, e.g., during a quench or a hysteresis loop protocol, as was indeed observed experimentally in [17]. The basic idea is explained diagrammatically in Fig. 2. Suppose the system is initially prepared in the MI state and is then suddenly quenched to the CDW phase. In this case, the MI state will continue to be a local minimum. As long as there is a small energy barrier to surmount, there will be a natural resistance for the system to reconfigure (this is analogous to the Stoner and Wolfarth model in magnetism [29]). Conversely, if we prepare the system in the CDW phase and then quench to the MI, as in Fig. 2(b), then the CDW state will no longer be a local minimum and there will be no energetic cost for the system to reconfigure to the MI phase. As a consequence, if one measures the hysteresis loop as $U_i$ is varied between the two phases, the result will be inherently asymmetric. Of course, this analysis does not take into account thermal or quantum fluctuations, nor the important fact that a quench implies injecting excitations into the system. Notwithstanding, it shows the existence of an inherent asymmetry between the two phases.

The Landau free energy presented in Fig. 1 corresponds to the case $\rho = 1$ [cf. Eq. (15)] and shows one or three minima. For noninteger fillings, other local minima appear. Examples of Eq. (11) for different fillings $\rho$ are shown in Fig. 3. As can be seen, noninteger fillings produce local minima at intermediate values of $\theta$. The existence of these degenerate configurations can be traced back to the results in Eq. (13) for the occupations that minimize the energy. We conjecture that these intermediate

FIG. 2. Description of the asymmetric behavior in the hysteresis of the MI-CDW transition. (a) The free energy as a function of $\theta$ when the system was initially prepared in the MI state ($U_i / U_s = 0.2$) and then suddenly quenched to the CDW phase ($U_i / U_s = 0.7$). In this case, there is a stronger resistance for the system to reconfigure towards the CDW minimum since the MI state continues to be a local minimum. Conversely, (b) shows the reverse effect. Now, after the quench, the system is found in a configuration which is no longer a local minima and, therefore, can easily reconfigure itself to the MI phase.

FIG. 3. The Landau free energy (11) for different values of $U_i$. Each image corresponds to a different filling $\rho$: (a) 1.15, (b) 1.25, (c) 1.35, and (d) 1.5. The values of $U_i$ and the color scheme are the same as in Fig. 1.
configurations might explain some of the plateaus observed in Fig. 4 of Ref. [18].

III. VARIATIONAL TREATMENT OF THE HOPPING TERM

When \( J \neq 0 \), the Hamiltonian (1) can no longer be diagonalized exactly. Instead, we approach the problem using a variational Schrödinger-Lagrangian ansatz,

\[
\mathcal{L} = \langle \psi | (i \partial_t - H) | \psi \rangle,
\]

with a choice of variational wave function \( |\psi\rangle \) that conveniently captures the physics of the MI-CDW transition. For concreteness, we fix \( \rho = 1 \) so that the most relevant states are of the form \(|\text{MI}\rangle = |1, \ldots, 1; 1, \ldots, 1\rangle\), \(|\text{CDW}_e\rangle = |2, \ldots, 2; 0, \ldots, 0\rangle\), and \(|\text{CDW}_o\rangle = |0, \ldots, 0; 2, \ldots, 2\rangle\), where the semicolon separates the Fock occupations of the even and odd sublattices.

Our goal is to describe the idea of an energy barrier forming between the MI and CDW phases. In order to do so, we need to choose a representative set of states which describes relevant intermediate configurations. Since we are interested in the case of small tunneling, we discard states with more than double occupation, restricting for each site to the occupations \( n_i = 0, 1, 2 \). States with higher occupation will have a much higher \( U_i \) repulsion, with no gain in \( U_i \) attraction, and hence should be energetically suppressed as long as \( J \) is small.

This truncation of the total Hilbert space is still insufficient to render the problem tractable, and we still need to find an appropriate choice of suitable intermediate states between MI and CDW. A natural choice is the states of definite imbalance, which are, eigenstates of the imbalance operator \( \Theta \) in Eq. (4). Since \( \rho = 1 \), the eigenvalues \( Q \) of \( \Theta \) can vary from \(-K\) to \( K\) in steps of two, with MI being \( Q = 0 \) and CDW being \( Q = \pm K \). Of course, each eigenvalue is highly degenerate and may thus be labeled as \(|Q, v\rangle\), for some additional index \( v \). We then have

\[
\Theta |Q, v\rangle = Q |Q, v\rangle.
\]

The MI and CDW states are the only elements of the \( Q \) states which are not degenerate. In particular, \(|\text{MI}\rangle = |Q = 0\rangle\) and \(|\text{CDW}_e, o\rangle = |Q = \pm K\rangle\).

In order to make the problem tractable, we must choose a representative set of states for intermediate values of \( Q \), which interpolates smoothly between these configurations. These states shall be denoted simply by \(|Q\rangle\). The principles that will guide our specific choice will be discussed below. With this choice, the Schödinger-Lagrangian will then produce a reduced dynamical description within this truncated subspace. The main results of the remainder of this section will be the main results of the Hamiltonian (1) in this \( Q \) basis. The Hamiltonian \( H_0 \) in (2) will be diagonal, while the hopping term will only connect neighboring \(|Q\rangle\) states. With these matrix elements at hand, we will then construct the effective Hamiltonian,

\[
\langle \psi | H | \psi \rangle = \sum_Q \left( \epsilon_Q \psi_Q^\dagger \psi_Q + \gamma_Q^+ \psi_{Q+2}^\dagger \psi_Q + \gamma_Q^- \psi_{Q-2}^\dagger \psi_Q \right),
\]

with coefficients \( \epsilon_Q, \gamma_Q^\pm \) to be found in the following and with

\[
|\psi\rangle = \sum_Q \psi_Q |Q\rangle.
\]

Hence, within this variational picture, the Hamiltonian is tri-diagonal and the hopping term produces an effective tight-binding model within the space of imbalances \( Q \).

We now proceed to derive these results in detail. In Sec. IV, they will be used to study the resulting properties of the system.

A. The choice of \( Q \) states

Recall that we are focusing on \( \rho = N/K = 1 \) and restricting to occupations \( n_i = 0, 1, 2 \). Next, consider the following states, each of which has an imbalance \( Q = 4 \):

\[
|1, 2, 1, 2; 0, 1, 0, 1\rangle,
\]

\[
|1, 2, 1, 2; 0, 0, 0, 2\rangle.
\]

However, the second state will be energetically more unlikely since it will have a higher \( U_i \) repulsion. Hence, the most likely states will be those where, for a given \( Q \), the atoms are as distributed as possible within each sublattice. That is, if \( Q > 0 \), then the even sublattice should only have 1’s and 2’s and the odd sublattice should only have 0’s and 1’s (and vice versa for \( Q < 0 \)). We can also reach the same conclusion by imposing that in the limit \( J \to 0 \), our variational method should reproduce exactly the Landau free energy (15). That is, we should impose that given the Hamiltonian \( H_0 \) in Eq. (2), our states must produce diagonal elements of the form

\[
\epsilon_Q := \langle Q | H_0 | Q \rangle = K f(Q/K) = \frac{U_i |Q|}{2} - \frac{U_i Q^2}{K}.
\]

It is straightforward to show that this will only take place if the above conditions are met.

The previous analysis still leaves a large ambiguity on which the physically reasonable \( Q \) states are. In order to gain further insight, let us write the hopping Hamiltonian (3) as

\[
\mathcal{T} = -\sqrt{2} J (\mathcal{T}_e + \mathcal{T}_o),
\]

where

\[
\mathcal{T}_e = \frac{1}{\sqrt{2}} \sum_i \sum_{j \not= i} b_i^\dagger b_j,
\]

and \( \mathcal{T}_o = \mathcal{T}_e^\dagger \). Here, \( \Gamma_i \) stands for the neighborhood of site \( i \). A direct calculation then shows that

\[
[\Theta, \mathcal{T}_o] = 2 \mathcal{T}_o, \quad [\Theta, \mathcal{T}_e] = -2 \mathcal{T}_o.
\]

Thus, \( \mathcal{T}_e \) and \( \mathcal{T}_o \) function as lowering and raising operators for the imbalance operator. The dynamics produced by the particle hopping can thus be interpreted as transitions between neighboring \( Q \) states; i.e., as a tight binding in the space of even-odd imbalances.

This analysis suggests that a reasonable set of \( Q \) states is precisely given by those states that are generated by the successive application of \( \mathcal{T}_e \) and \( \mathcal{T}_o \) onto the MI state. This process therefore describes the nucleation of CDW 0–2 pairs within the lattice, which has a similar picture to the usual nucleation of domains in thermal phase transitions [12,30]. Of
course, when we apply these operators, we will also generate states with more than double occupation. To amend for this we define

$$\tilde{T}_e = \mathcal{P} T_e \mathcal{P},$$

(24)

where $\mathcal{P}$ is a projection operator onto the subspace of states which are at most doubly occupied. Note that $\tilde{T}_e$ no longer satisfies the closed algebra of Eq. (23), as is customary for projected operators. We may then finally define, for $Q > 0$,

$$|Q\rangle = \frac{1}{\sqrt{A(Q)}} (\tilde{T}_e)^{Q/2} |\text{MI}\rangle,$$

(25)

where $A(Q)$ is a normalization constant, to be determined. For $Q < 0$, we replace $\tilde{T}_e$ with $\tilde{T}_{e0}$, defined in an analogous manner.

In order to construct the Schrödinger Lagrangian (16), we need the matrix elements of the total Hamiltonian (1) in the $Q$ basis. The diagonal part $H_0$ is given in Eq. (20). Next, we must turn to the matrix elements of the hopping term $T$ in Eq. (21). This operator connects $Q$ with $Q \pm 2$. In particular, the connection with $Q + 2$ will be given by

$$\langle Q + 2|T_e|Q\rangle = \langle Q + 2|\tilde{T}_e|Q\rangle = \sqrt{\frac{A(Q+2)}{A(Q)}},$$

(26)

Hence,

$$\langle Q + 2|\mathcal{P}T_e\mathcal{P}|Q\rangle = -\sqrt{2} \sqrt{\frac{A(Q+2)}{A(Q)}},$$

(27)

which is the general formula relating this matrix element with the normalization constant $A(Q)$ of Eq. (25). Analogous formulas can be obtained for $Q < 0$, noting that for symmetry reasons, $A(Q) = A(-Q)$ and $\langle Q + 2|\mathcal{P}T_e\mathcal{P}|Q\rangle = -\langle Q + 2|\mathcal{P}T_e\mathcal{P}|Q\rangle$.

### B. Distortion of the hopping term

Unfortunately, computing the normalization constant $A(Q)$ in Eq. (25) is in general a very difficult task since it depends on the structure of the underlying lattice, which appears in the generator $\tilde{T}_e$ in Eq. (22). In fact, as shown in the Appendix, this calculation maps exactly to the matching problem, which not only has no analytical solution, but is also a $\mathcal{NP}$-hard problem to solve numerically (even if we attempt to estimate these coefficients by Monte Carlo methods, the state-of-the-art algorithms have a complexity of $O(K^7 \log^4 K)$ [31,32]). We also mention in passing that this turns out to be the same problem that one encounters in computing the partition function of the monomer-dimer model or the three-dimensional Ising model [33].

Instead, in order to obtain an analytical treatment, we propose here a deformation of $\tilde{T}_e$ to eliminate the underlying lattice structure. The deformation is similar to the one routinely done in mean-field treatments and consists of allowing an atom to hop from any even site to any odd site. That is, we deform $\tilde{T}_e$ to a new hopping operator $\tilde{T}_e'$ defined as

$$\tilde{T}_e' = \frac{1}{\sqrt{2}} \sum_{i\in e} \sum_{j\in o} \mathcal{P} b_i^\dagger b_j \mathcal{P},$$

(28)

where the two sums in $e$ and $o$ are now unrestricted. We remark that we make this replacement for the definition of the $|Q\rangle$ basis states only, and not in the Hamiltonian of the system.

With this modified hopping generator, we can now compute the matrix element of the hopping term $T$. To illustrate the procedure, it suffices to consider $Q > 0$. Then the even sublattice will only have occupation 1 or 2 and the odd sublattice will only have occupation 1 or 0. Let us define $C_e$ as the list of sites in the even sublattice which are doubly occupied and $C_o$ as the list of sites in the odd sublattice which are empty. We also define the state $|C_e,C_o\rangle$ as the Fock state with the sites in $C_e$ doubly occupied, the sites in $C_o$ empty, and all others singly occupied. Then, the set of $Q$ states generated as in Eq. (25), but with $\tilde{T}_e'$ instead, may be written as

$$|Q\rangle = \frac{1}{\sqrt{A(Q)}} \sum_{|C_e,C_o\rangle} |C_e,C_o\rangle.$$

(29)

The normalization constant $A(Q)$ is now simply the number of states in this list, which is

$$A(Q) = \left(\frac{K/2}{Q/2}\right)^2.$$

(30)

Now let us analyze the action of the operator $\mathcal{P}(b_i^\dagger b_j)\mathcal{P}$ in the $Q$ state (29), assuming $i \in e$ and $j \in o$. This will only contribute if it acts on a pair of sites with $n = 1$. Thus,

$$\mathcal{P}(b_i^\dagger b_j)\mathcal{P}|C_e,C_o\rangle = \left\{ \begin{array}{ll} \sqrt{2}|C_e+i,C_o+j\rangle, & i \notin C_e, \; j \notin C_o \\ 0 & \text{otherwise,} \end{array} \right.$$
which, after simplifying, yields
\[ \langle Q + 2 | b_i^\dagger b_j^\dagger | Q \rangle = \sqrt{2} \frac{(K - Q)(Q + 2)}{K^2}. \] (32)

Note that this result is independent of \((i, j)\), which is a consequence of the distortion introduced in Eq. (28), that washes out all lattice information. Consequently, it is now trivial to find the matrix elements of \( \mathcal{T} \):
\[ \langle Q + 2 | \mathcal{T} | Q \rangle = -J \sqrt{2} \frac{(K - Q)(Q + 2)}{K^2} \sum_{i \in \mathcal{E}} \sum_{j \in \mathcal{N}_i} 1. \] (33)
This sum has \( \frac{z}{2} \times z \) elements, with \( z \) the number of neighbors (\( z = 4 \) in the two-dimensional case we consider), so we finally obtain
\[ \gamma_Q^+ := \langle Q + 2 | H | Q \rangle = -\frac{\alpha}{4K} \left\{ \begin{array}{ll} (K - Q)(Q + 2), \quad Q \geq 0 \\ (K - |Q|)|Q|, \quad Q < 0, \end{array} \right. \] (36)
\[ \gamma_Q^- := \langle Q - 2 | H | Q \rangle = -\frac{\alpha}{4K} \left\{ \begin{array}{ll} (K - Q + 2)Q, \quad Q > 0 \\ (K - |Q|)(|Q| + 2), \quad Q \leq 0, \end{array} \right. \] (37)
where we have defined a rescaled hopping parameter,
\[ \alpha = 2\sqrt{2}\gamma J. \] (38)

The functions \( \gamma_Q^\pm \) represent the effective hopping in the space of imbalances and correspond to the off-diagonal entries in the effective Hamiltonian (18).

We finish this section by emphasizing that the present choice of \( \langle Q \rangle \) states neglects lattice information, such as dimensionality and number of neighbors. One should therefore not expect that this model would reproduce details of the phase diagram depending on these quantities such as, for instance, the location of the MI-SF transition. On the other hand, as far as the MI-CDW transition is concerned, this dependence should be weak since, in the limit of zero hopping, the lattice structure plays no role at all.

IV. ANALYSIS OF THE VARIATIONAL HAMILTONIAN

A. Numerical analysis

In this section, we study, both numerically and analytically, the effective variational Hamiltonian in Eq. (18). We begin with a numerical analysis of the ground state and the excitations, computed by diagonalizing the tridiagonal Hamiltonian (18) using standard routines. The results are shown in Fig. 4 as a function of \( U_l / U_s \), for a fixed \( K = 2000 \) and different values of the effective hopping term \( \gamma = 2\sqrt{2}\gamma J \). We stress that for completeness, we include results corresponding to all values of \( \alpha \). However, our model is only built to describe the MI-CDW transition, i.e., the regime of small tunneling rates.

In Fig. 4(a), we present the average imbalance \( \langle \Theta \rangle / K \). It shows a discontinuous transition at \( U_l = U_s / 2 \), from 0 to 1. This represents the MI-CDW transition in the limit of small hopping. Such a discontinuous transition continues up until \( \alpha = 1 \), above which one enters a compressible phase which falls outside the scope of this variational model. A similar behavior is found in the energy gap between the ground state and the first excited state, illustrated in Fig. 4(b). For \( \alpha < 1 \), the gap is independent of \( U_l \) in the MI phase and grows linearly in the CDW phase. For \( \alpha = 1 \), the gap closes at \( U_l = 1 \), marking the existence of a compressible phase (we stress though that our model is not a good description of this regime).

The simplification afforded by our choice of \( \langle Q \rangle \) states in Eq. (29) allows us to also compute the entanglement entropy

FIG. 4. Ground-state properties of the variational Hamiltonian, computed as a function of \( U_l / U_s \), for \( K = 2000 \) and different values of \( \alpha / U_s \), from 0 to 1 [the curve corresponding to \( \alpha / U_s = 0 \) is the uppermost curve in (b) and the lowermost curve in (c) and (d)]. The quantities plotted in each figure are the following: (a) \( \langle \Theta \rangle \), (b) the energy gap between the ground state and the first excited state, (c) the entanglement entropy between the even and the odd sublattices, and (d) the fidelity susceptibility (plotted logarithmically for clarity).
between the even and the odd sublattices, which is defined as

\[ S_{\text{EN}} = -\text{tr} \rho_e \ln \rho_e, \tag{39} \]

where \( \rho_e = \text{tr}_o |\psi\rangle \langle \psi| \). The \( Q \) basis in Eq. (29) can be split as a tensor product of states in the even and odd sublattices. Consequently, it follows from a straightforward calculation that we will have

\[ S_{\text{EN}} = -\sum_Q |\psi_Q|^2 \ln |\psi_Q|^2. \tag{40} \]

This is shown in Fig. 4(c) for the numerically computed ground-state wave function. The entanglement entropy is taken as a measure of the correlation between the two sublattices. As expected, we see that the hopping increases their correlation, which is also generally larger in the MI than in the CDW phase.

Finally, in Fig. 4(d), we compute the fidelity susceptibility [34,35],

\[ \chi = -\frac{\partial^2}{\partial \delta^2} \ln |\langle \psi(U_\delta) |\psi(U_\delta + \delta) \rangle| \bigg|_{\delta=0}. \tag{41} \]

This quantity is a well-known indicator of criticality, and indeed for \( \alpha < 1 \), it presents a discontinuous jump at the MI-CDW transition, while for \( \alpha \geq 1 \), it presents a peak at the onset of the compressible phases. In fact, we find from our simulations that, quite remarkably, the fidelity can pinpoint all transitions of the model even for sizes as small as \( K = 10 \).

From these results, we reconstruct the equilibrium phase diagram depicted by the blue (solid) and yellow (dotted) lines in Fig. 5. Surprisingly, even though our model was constructed with the intent of describing only the MI-CDW transition, it turns out to capture a similar phase diagram as that obtained in Ref. [23] using the Gutzwiller approximation. Our model does induce, however, some imprecision in the case of the compressible phases. First, it cannot distinguish between superfluid and supersolid, which is a consequence of the reduced Hilbert space employed in our variational wave function. Second, it predicts a MI-SF transition occurring at \( \alpha/U_\delta = 1 \). Since \( \alpha = 2\sqrt{2} \gamma J \), this gives a transition at \( U_\delta/J = 11.31 \), which is approximately half of the value obtained by mean-field approximations or quantum Monte Carlo [1,6,19,20,23,25].

**B. Discrete WKB method**

Next we show that the phase diagram in Fig. 5 can actually be obtained analytically using a discrete WKB approximation [36–38]. Moreover, this approach will allow us to augment the phase diagram with additional information on the height of the energy barrier separating the two phases (red lines in Fig. 5).

The essence of the WKB method is to note that the effective Hamiltonian (18) closely resembles the tight-binding model describing the hopping of particles through a linear lattice. The major difference is that in our case, the hopping terms \( \gamma_Q \) are nonuniform. If the hopping was uniform, then it is well known that the eigenstates of the Hamiltonian would be momentum plane waves \( e^{i\gamma Q} \) with dispersion relation \( E = \epsilon + 2\gamma \cos(p) \), where \( p \) is the momentum and \( E \) is the energy. In our case, even though the hopping is not uniform, in the limit of large \( K \) the hopping parameters \( \gamma_Q \) will change slowly with \( Q \). This motivates us to define \( Q \)-dependent momenta of the form [37]

\[ \cos p(Q) = \frac{E - \epsilon_Q}{2\gamma_Q}, \tag{42} \]

where

\[ \gamma_Q := \frac{\gamma_Q^+ + \gamma_Q^-}{2} = -\frac{\alpha}{4} \left( 1 + |Q| - \frac{Q^2}{K} \right). \tag{43} \]

which, by construction, is always negative. The condition that \( p(Q) \) should be real determines the classically allowed regions where the eigenvectors of the effective Hamiltonian are oscillatory,

\[ \epsilon_Q + 2\gamma_Q \leq E \leq \epsilon_Q - 2\gamma_Q. \tag{44} \]

In Fig. 6, we present these classically allowed regions for different combinations of \( U_\delta \) and \( \alpha \).

With the classically allowed regions properly identified, we may now finally make the link with the metastability in Fig. 2 and understand exactly how the particle hopping facilitates the transition between metastable states, which is the primary goal of this paper. In Fig. 6(a), for instance, we present results for \( U_\delta/U_\delta = 0.4 \) and \( \alpha/U_\delta = 0.3 \). In this case, the global minimum is the MI phase, whereas the CDW configurations represent metastable states. In the absence of hopping (\( \alpha = 0 \)), the energy barrier that must be surmounted is represented by the black dashed curve in Fig. 6(a). When hopping is present, however, the entire shaded region within the two red curves [cf. Eq. (44)] becomes classically allowed, meaning that the barrier that the system will actually have to surmount will be that given by the lowest of the two red curves, which is substantially
smaller than the original barrier. In fact, in Fig. 6(b), we present curves for the same value of $U_1/U_s$, but a larger value of $\alpha$. In this case, as can be seen, the hopping has completely destroyed the barrier between the two minima.

We now use this information to construct the phase diagram in Fig. 5 and augment it with information on the barrier height. From Eq. (44), we see that the quantity determining the energy barrier will be the lower bound,

$$\mathcal{E}(Q) = \epsilon_0 + 2\gamma_0$$

$$= \frac{\alpha}{2} + |Q| (U_s - \alpha) - \frac{Q^2}{2K} (2U_s - \alpha), \quad (46)$$

which correspond to the bottom red curves in Fig. 6. Our first goal is to use this to establish in which cases a barrier exists. For instance, in the examples of Fig. 6, it exists only in Figs. 6(a), 6(c), and 6(d), where they are signaled by vertical arrows. First, a necessary (but not sufficient) condition can be easily found by noting that Eq. (46) is a parabola in $Q$, so that for a barrier to exist, the coefficient of the quadratic term must be negative, hence implying $\alpha < 2U_s$. Above this value, we are always in a compressible phase and thus no barrier exists.

Next, focusing on $Q > 0$, we note that the extremum of $\mathcal{E}$ occurs at

$$\frac{Q^*}{K} = \frac{1}{2} (U_s - \alpha)/(2U_s - \alpha). \quad (47)$$

We must then demand that this extremum exists in the interval $Q^*/K \in [0, 1]$; the limiting cases turn out to be

$$\alpha_1 = U_s \quad \text{and} \quad \alpha_2 = 4U_s - U_s. \quad (48)$$

The meaning of these equations changes depending on the value of $U_1$ since the point $Q^*/K$ will only be a maximum ($E'' < 0$) provided $\alpha < 2U_s$.

(i) If $U_1 > U_s/2$ (CDW phase), then the line $\alpha_1 = U_s$ determines the region where the barrier ceases to exist, which is denoted by the red (dashed) vertical line in Fig. 5. Conversely, the line $\alpha_2 = 4U_s - U_s$ denotes the line after which the extremum becomes a minimum, which signals the transition to a compressible phase. This is denoted by the curved blue line in Fig. 5, which is defined for $U_1 > U_s/2$.

(ii) If $U_1 < U_s/2$ (MI phase), then the roles are inverted.

The line $\alpha_1 = U_s$ determines the transition to a compressible phase (blue line in Fig. 5) and the line $\alpha_2 = 4U_s - U_s$ denotes the region below which the barrier vanishes (red line in Fig. 5).

We may also construct the energy-barrier height that must be overcome in order to transition from a metastable minimum to the true global minimum. If we are in the CDW phase, then the barrier height will be given by

$$\Delta \mathcal{E}_{\text{CDW}} = \mathcal{E}(Q^*) - \mathcal{E}(0) = \frac{K (U_s - \alpha)^2}{8 (2U_s - \alpha)}. \quad (49)$$

Conversely, if we are in the MI phase, we will have

$$\Delta \mathcal{E}_{\text{MI}} = \mathcal{E}(Q^*) - \mathcal{E}(0) = \frac{K (\alpha + U_s - 4U_s)^2}{8 (2U_s - \alpha)}. \quad (50)$$

These results are shown in the form of density plots in Fig. 5. Thus, we see that from the discrete WKB method, we may not only compute the entire equilibrium phase diagram, but also determine the regions where barriers between minima exist and the height of these barriers.

Finally, we use the discrete WKB method to address the problem of macroscopic quantum tunneling from the metastable state toward the true global minimum. For instance, in the example of Fig. 6(a), we shall consider the tunneling from the CDW state $Q = K$ toward the MI state $Q = 0$, and vice versa in the case of Figs. 6(c) and 6(d). Following [36–38], we write the tunneling probability as

$$\mathcal{T} = e^{-2\mathcal{A}}, \quad (51)$$

where

$$\mathcal{A} = \frac{1}{K} \int_{Q_1}^{Q_2} |p(Q)| dQ. \quad (52)$$

with $p(Q)$ given in Eq. (42) and $Q_{1,2}$ being the classical turning points of the dynamics. They are indicated in Figs. 6(a), 6(c) and 6(d) by horizontal dashed lines. From $\mathcal{T}$, one may estimate the average lifetime as $\tau \sim 1/\mathcal{T}$, which can unfortunately only be done up to a prefactor that is not easily computed.

The integral in (52) can be computed analytically, even though the result is somewhat cumbersome. To do so, we separately treat the cases $U_1 < U_s/2$ and $U_1 > U_s/2$ since the turning points $Q_{1,2}$ are different in each case. Let us start with $U_1 < U_s/2$, for which the global minimum is the MI phase. This situation is illustrated in Fig. 6(a). The classical
turning points \( Q_1 \) and \( Q_2 \) are precisely the endpoints of the gray dashed line. Thus \( Q_2 = K \), whereas \( Q_1 \) is the solution of

\[
\mathcal{A} = \text{sech}^{-1} \left( \frac{\alpha}{4U_\ell - U_s} \right) + \frac{U_s - 2U_\ell}{U_s} \ln \left\{ \frac{\alpha(U_s - 2U_\ell)}{2U_\ell(U_s - 8U_\ell^2 + 2\alpha^2 - \sqrt{(4U_\ell^2 - U_s)^2 - \alpha^2})(4U_\ell^2 - \alpha^2)} \right\}.
\]

We proceed similarly for \( U_\ell > U_s/2 \). In this case, the turning points are \( Q_1 = 0 \) and \( Q_2 = K(U_s - \alpha)/(2U_\ell - \alpha) \), which yield

\[
\mathcal{A} = \ln \left[ \frac{U_s + \sqrt{U_s^2 - \alpha^2}}{U_s} \right] + \frac{U_s - 2U_\ell}{U_s} \ln \left\{ \frac{\alpha(U_s - 2U_\ell)}{2\alpha^2 - 2U_\ell U_s + \sqrt{(4U_\ell^2 - \alpha^2)(U_s^2 - \alpha^2)}} \right\}.
\]

These results simplify considerably in the limit \( U_\ell = U_s/2 \), which is when all energy minima have the same energy. In this case, they both tend to

\[
\mathcal{A} = \cosh^{-1} \left( \frac{U_s}{\alpha} \right).
\] (55)

Thus, we see that when \( \alpha \to 0 \), the amplitude \( \mathcal{A} \) diverges and, consequently, the tunneling probability tends to zero, as expected. Conversely, when \( \alpha \geq U_s \), the tunneling amplitude is identically zero, as there is no energy barrier to surmount. The same is also true, for instance, in the example of Fig. 6(b), although that is not so readily seen from Eq. (53).

We can compare these results directly with the lifetime of thermally assisted transitions. From the standard Arrhenius theory, the average lifetime for thermal transitions should be proportional to \( e^{\Delta E/\hbar \tau} \), where \( \Delta E \) is the height of the energy barrier. In the case \( U_\ell = U_s/2 \), the height of the barrier is \( \Delta E = K(U_s - \alpha)/8 \). Thus, by analyzing how the average lifetime of the metastable states scales with the hopping parameter \( \alpha \), it should be possible to infer the relative contribution of quantum tunneling versus thermally assisted transitions. The method used here only allows us to estimate the main exponential dependence, thus making it difficult to directly compare with experiment. A more thorough analysis of the tunneling time will be the subject of a future paper.

\section{V. Conclusions}

In summary, we have discussed the role of quantum fluctuations, represented by the local tunneling of atoms in an optical lattice, in the phase reconfiguration of a first-order quantum phase transition. Within the energy-landscape picture provided by the discrete WKB method, we find that the tunneling acts to lower the energy barrier separating the two phases, hence facilitating the transition between a metastable minimum and the true global minimum. Our approach relied on a choice of variational states describing the relative imbalances between the even and odd sublattices. As a result, we found that the tunneling acts as a local tight binding in the space of imbalances, causing transitions between neighboring states. One may thus interpret the tunneling of atoms as the generators of CDW 0–2 pairs within the lattice. These pairs should then form domains which eventually nucleate into a new phase. It is our hope that our analysis sheds further light on the nonequilibrium mechanisms responsible for phase reconfigurations in quantum critical systems.

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\textbf{APPENDIX: MAPPING TO THE MATCHING PROBLEM}

In this appendix, we show that the task of computing the normalization constants \( A(Q) \) in Eq. (25) can be mapped into a matching problem that is widely studied in computational combinatorics \cite{33}. Let \( c_i \) be the operators

\[
c_i = \begin{cases} b_i, & i \in o \\ \frac{1}{\sqrt{2}}b_i^\dagger, & i \in e, \end{cases}
\] (A1)

and, for any set \( S \) of lattice sites, define an operator \( \mathcal{D}_S \) as

\[
\mathcal{D}_S = \prod_{i \in S} c_i \mathcal{P}.
\] (A2)

States created by the action of \( \mathcal{D}_S \) onto the MI state are orthonormal. That is, if \( |S\rangle \equiv \mathcal{D}_S |MI\rangle \), then \( \langle S|S'\rangle = \delta_{S,S'} \). Moreover, the \( \mathcal{D} \) operators obey the algebra

\[
[D_X, D_Y] = 0, \\
D_X D_Y = 0 \text{ if } X \cap Y \neq \emptyset, \\
D_X D_Y = D_X \delta_{X,Y} \text{ if } X \cap Y = \emptyset.
\]

In terms of these new operations, we may write the hopping generator \( \hat{T}_e \) as

\[
\hat{T}_e = \sum_{\langle i,j \rangle} \mathcal{D}_{i,j}.
\] (A3)

Hence,

\[
\hat{T}_e = \sum_{\mathcal{R}} (n!)^\mu_{\mathcal{R}} D_{\mathcal{R}}.
\] (A4)
are allowed in elements that μ show elements of neighbor pairs in only one way (hence, μ = 1). The three other panels show elements of C3 which can be covered in three different ways so that μ = 3. Note that since only neighboring pairs with no overlaps are allowed in elements R of Cn, it follows that all R cover an equal number of even and odd sites.

Calculating the exact expression (A6) for all the needed values of Q involves being able to calculate the total number of matchings in a square lattice,

\[ \sum_n \sum_{R \in C_n} \mu_R, \]

which is a \( \mathcal{O}(P) \)-hard problem, making it unfeasible even numerically. Even if we relax our requirements and attempt to only estimate these quantities by Monte Carlo methods, the state-of-the-art algorithm is still \( O(K^7 \log^4 K) \) [31,32].