Quantum signatures of charge flipping vortices in the Bose-Hubbard trimer

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(Dated: June 20, 2016)

Abstract

In this work we study quantum signatures of charge flipping vortices, found in the classical discrete nonlinear Schrödinger trimer, by use of the Bose-Hubbard model. We are able to identify such signatures in the quantum energy eigenstates, for instance when comparing the site amplitudes of the classical charge flipping vortices with the probability distribution over different particle configurations. It is also discussed how to construct quantum states that correspond to the classical charge flipping vortices, and which effects that can lead to deviations between the classical and quantum dynamics.

We also examine properties of certain coherent states: classical-like quantum states that can be used to derive the classical model. Several quantum signatures are identified when studying the dynamics of these coherent states, for example when comparing the average number of particles on a site with the classical site amplitude, when comparing the quantum and classical currents and topological charge, and when studying the evolution of the quantum probability amplitudes. The flipping of the quantum currents are found to be an especially robust feature of these states.

PACS numbers:

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I. INTRODUCTION

The Bose-Hubbard model has become a prominent and much studied model in physics through the study of Bose-Einstein condensates in optical lattices. This has been a very active and prolific area of research in the last few decades, and is among other things considered a promising candidate for quantum computers and simulators [1, 2]. In the classical limit of the Bose-Hubbard model, corresponding to a large number of bosons per site, it is well approximated by the discrete nonlinear Schrödinger (DNLS) model [3, 4]. The DNLS model is one of the most studied nonlinear lattice models, and has apart from Bose-Einstein condensates also applications for e.g. optical wave-guide systems [5].

Of fundamental interest here is the quantum-classical correspondence, and especially how structures connected to the nonlinearity of the classical DNLS model emerge from the inherently linear quantum model. Studies of this kind are numerous and include those of discrete breathers [4, 6], compactons [7, 8], and vortices [9], as well as the self-trapping transition [10, 11], modulational instabilities [12, 13] and oscillatory instabilities [14]. More recently, Reference [15] studied quantum signatures of Arnold diffusion in Bose-Hubbard lattices with few sites (especially three and four sites).

The Hilbert space of the Bose-Hubbard model increases very rapidly with the number of sites and particles, and one is typically limited to quite modest system sizes, especially when using exact diagonalization. One beneficial aspect of studying small lattices is therefore that it enables us to use (in this context) many particles, and better connect the classical and quantum world. Smaller lattices are also experimentally accessible nowadays, for instance by painting optical potentials [16].

We will in this paper look for quantum signatures of the charge flipping vortices (CFVs) recently found and studied in the DNLS trimer (symmetric triangular configuration) [17]. A vortex is the rotational flow of a field around a topological phase dislocation; a point where the field amplitude vanishes and around which the field changes its phase by an integer multiple of $2\pi$. This integer is the Topological Charge (TC) of the vortex, and its sign indicating the vortex' direction. CFVs are characterized by a repeated flipping of the TC, and consequently also a reversal of the flow. The TC is conserved in systems with full rotational symmetry, which thus needs to be broken to enable CFVs, for instance by introducing an underlying lattice potential. CFVs have been studied in a wide variety
of lattice geometries and configurations, both theoretically [17–22] and experimentally in photonic lattices [23–25]. The CFVs found in Reference [17] are periodic solutions (in co-rotating frames) that exist in families which can be followed continuously between two stationary solutions: the single depleted well (SDW) solution, \( \psi = (\psi_1, -\psi_1, 0) \), and a dimerlike solution \( \psi = (\psi_1, -\psi_2, -\psi_2) \); \( \psi_1, \psi_2 \in \mathbb{R}^+ (\psi_1 > \psi_2) \) [26]. The TC of these CFVs will naturally have the same periodicity as the CFVs themselves.

The dimerlike solution is the ‘discrete breather’ ground state, with a peak on one site [27].

The outline of the paper is as follows. Sec. II contains some theoretical background, with Sec. II A introducing the classical and quantum model, Sec. II B discussing some aspects of coherent states, and Sec. II C describing how to calculate classical and quantum currents and TC. Sec. III begins by recapitulating some results for the classical CFVs, before examining CFV quantum signatures in the energy eigenstates. Sec. IV focuses on the classical-like coherent states, with Sec. IV A examining the quantum dynamics of these, Sec. IV B studying their projection onto energy eigenstates, and Sec. IV C analyzing how the quantum dynamics changes when using classical CFVs at different times of their evolution as input to the coherent states. The quantum results in Secs. III and IV are for specific number of particles and values of the nonlinear parameter, and Sec. V discusses some effects of changing these. Sec. VI finishes the paper with a summary and conclusions.

II. THEORY

A. Models

Consider the Bose-Hubbard trimer on the following form

\[
\hat{H} = \sum_{j=1}^{3} \left( \hat{a}^\dagger_{j+1} \hat{a}_j + \hat{a}^\dagger_j \hat{a}_{j+1} - \frac{\alpha}{2} \hat{n}_j (\hat{n}_j - 1) \right),
\]

where \( \hat{a}^{(t)}_j \) is the bosonic annihilation (creation) operator for site \( j \), \( \hat{n}_j = \hat{a}^\dagger_j \hat{a}_j \) the number operator for the same site, and \( \alpha > 0 \) determines the interaction strength. Because of the periodicity of the lattice \( \hat{a}^{(t)}_4 = \hat{a}^{(t)}_1 \). This differs from the more standard form of the
Bose-Hubbard model with repulsive interactions:

\[ \hat{H} = \sum_{j=1}^{3} \left( -J (\hat{a}_{j+1}^{\dagger} \hat{a}_j + \hat{a}_j^{\dagger} \hat{a}_{j+1}) + \frac{U}{2} \hat{n}_j (\hat{n}_j - 1) \right), \]

where \( J > 0 \) and \( U > 0 \) are the hopping and (repulsive) interaction parameter, respectively. The two Hamiltonians are however related by \( \hat{H} = -\hat{H}'/J \) and \( \alpha = U/J \), which not only rescales the energies but also flips the energy spectrum. We have chosen to use Hamiltonian (1) to facilitate comparisons with the classical results in Reference [17]. We will also deal with the lowest eigenstates of Hamiltonian (1), which correspond to the highest eigenstates of the standard Hamiltonian (2).

The quantum states of the Bose-Hubbard model are expressed with Fock states, \(|n_1, n_2, n_3\rangle\), where \( n_j \geq 0 \) is the number of bosons on site \( j \). Because Hamiltonian (1) commutes with the total number operator \( \hat{N} = \sum_{j=1}^{3} \hat{n}_j \), its eigenstates have a specified total number of particles \( N = n_1 + n_2 + n_3 \), and we will therefore always work in subspaces with a fixed \( N \). This implies that \( n_j \) for only two sites need to be specified for a Fock state, and that the probability distribution of the different particle configurations \((n_1, n_2, n_3)\) may conveniently be illustrated with two-dimensional triangular plots, similar as in References [11, 15]. We will frequently use the relative number of particles per site, \( \nu_j = n_j/N \), and the associated operator \( \hat{\nu}_j = \hat{n}_j/N \), since it simplifies the comparison of systems with different \( N \).

Furthermore, both \( \hat{H} \) and \( \hat{N} \) commute with the translation operator \( \hat{T} |n_1, n_2, n_3\rangle = |n_3, n_1, n_2\rangle \). The energy eigenstates will therefore also be labeled with a \( k \)-value, which is related to the eigenvalues of \( \hat{T} \) by \( \hat{T} |\psi\rangle = e^{ik} |\psi\rangle \), and may take the values \( k = 0, \pm 2\pi/3 \) for the trimer. We will denote the \( m \)-th lowest energy in subspace \( k \), and the associated eigenstate, with \( E_m(k) \) and \(|E_m(k)\rangle\). Since there is no preferred direction in the model, the spectra for \( k = \pm 2\pi/3 \) are identical, and the \( \pm \)-sign is hereafter omitted.

In the classical limit of a large number of particles, the Bose-Hubbard model is well approximated by the DNLS model. Formally, this can be derived by using the 'time-dependent variational principle' (TDVP) with SU(3) coherent states as trial states (generally SU(\( f \))
coherent states for $f$ number of sites)\[28\]. The SU(3) coherent states are defined as

$$|CS[\psi]\rangle = \frac{1}{\sqrt{N!}} \left( \sum_{j=1}^{3} \psi_j a_j^\dagger \right)^N |\text{vac}\rangle,$$

(3)

$$\mathcal{N} = \sum_{j=1}^{3} |\psi_j|^2 = 1,$$

(4)

$|\text{vac}\rangle$ being the vacuum state. Other trial states can also be used, e.g. the Glauber coherent states familiar from quantum optics [29, 30], but the advantage of the SU(3) coherent states is that they have a fixed total number of particles $N$ [28]. Note that we let the classical state vector $\psi = (\psi_1, \psi_2, \psi_3)$ be normalized to one, unlike in Ref. [28] where it is normalized to $N$ (see Reference [8] for further discussions). The actual quantum state is however the same. Let us also point out the difference between $|CS[\psi(t)]\rangle$ and $|CS[\psi](t)\rangle$. The state $|CS[\psi(t)]\rangle$ corresponds to $\psi(t)$ inserted into Eq. (3), where $\psi(t)$ has been obtained by integrating the DNLS equation (5) with the initial condition $\psi(0)$. $|CS[\psi](t)\rangle$ on the other hand is the quantum time evolution, calculated with the Bose-Hubbard Hamiltonian (1), of the coherent state obtained when inserting $\psi = \psi(0)$ into Eq. (3). Note that the quantum time evolution of a coherent state generally does not produce another coherent state.

Using SU(3) coherent states in the TDVP means that $\langle CS[\psi]|i\partial/\partial t - \hat{H}|CS[\psi]\rangle = 0$, leading to

$$i\dot{\psi}_j - (\psi_{j-1} + \psi_{j+1}) + \alpha(N - 1) |\psi_j|^2 \psi_j = 0,$$

(5)

which we identify as the DNLS model with nonlinear parameter $\beta = \alpha(N - 1)$ [5]. Eq. (5) indicates that $\alpha$ must scale as $\sim N^{-1}$ with the number of particles for the DNLS to be the proper classical limit. The classical Hamiltonian associated with Eq. (5) is

$$\mathcal{H} = \sum_{j=1}^{3} \left( \psi_j \psi_j^* + \psi_{j+1} \psi_j^* - \frac{\beta}{2} |\psi_j|^4 \right),$$

(6)

with $\{\psi_j\}$ and $\{i\dot{\psi}_j^*\}$ as canonical coordinates and momenta, respectively. By scaling $\alpha \sim N^{-1}$, the eigenvalues of the Hamiltonian (1) scale as $\sim N$. For a classical state with energy $\mathcal{H}$, the energy of a corresponding eigenstate can be approximated by $N\mathcal{H}$.

Reference [17] uses a DNLS model like Eq. (5) but with $\beta$ fixed to unity and a variable norm $\mathcal{N} = \sum_{j=1}^{3} |\psi_j|^2$. Rescaling $\psi$ shows that this is equivalent to Eq. (5) (i.e. with normalized $\psi$) with the nonlinear parameter $\beta = \mathcal{N}$. The norm of a CFV in Reference [17]
will thus be the nonlinearity for the corresponding normalized CFV in Eq. (5), and we will only use the latter type in the rest of the paper.

B. Coherent States

Coherent states are generally thought to be the quantum states which best describe classical systems, and they have several useful properties to this end [31]. For instance, the input vector \( \psi \) of a coherent state (3) becomes the dynamical variable in the classical description (5), and there is thus a one-to-one correspondence between classical states and coherent states. Another important property is that 
\[
\langle \text{CS}[\psi] | \hat{\nu}_j | \text{CS}[\psi] \rangle = |\psi_j|^2
\]
From Bohr’s correspondence principle we expect that 
\[
\langle \text{CS}[\psi](t) | \hat{\nu}_j | \text{CS}[\psi](t) \rangle \to |\psi_j(t)|^2
\]
in the limit \( N \to \infty \), while the variance of \( \hat{\nu}_j \) approaches zero. We will examine how well 
\[
\langle \text{CS}[\psi](t) | \hat{\nu}_j | \text{CS}[\psi](t) \rangle
\]
follows \( |\psi_j(t)|^2 \) for finite \( N \) in different regimes, and how the classical dynamics emerges from the quantum states. Consider therefore

\[
\langle \hat{\nu}_j \rangle = \langle \text{CS}[\psi](t) | \hat{\nu}_j | \text{CS}[\psi](t) \rangle = \sum_{k,k',m,m'} d_m(k)(d_{m'}(k'))^* \langle E_{m'}(k') | \hat{\nu}_j | E_m(k) \rangle \exp(-i[E_m(k) - E_{m'}(k')]t),
\]
where \( |\text{CS}[\psi](0)\rangle = \sum_{k,m} d_m(k) |E_m(k)\rangle \) \( (\hbar = 1) \). The time evolution of \( \langle \hat{\nu}_j \rangle \) in Eq. (7) is (like any observable) determined by the structure of the occupied eigenstates, and how their energies are related. Deviations between \( \langle \hat{\nu}_j \rangle \) and \( |\psi_j|^2 \) can, as we will see later on, be associated with the structure of the eigenstates and energy spectrum.

C. Topological Charge

A continuity equation can be obtained for the classical site amplitudes from Eq. (5),

\[
\frac{d|\psi_j|^2}{dt} = J_{j-1} - J_j,
\]
where

\[
J_j = -2 \text{Im}\{\psi_j^* \psi_{j+1}\} = -2 |\psi_j||\psi_{j+1}| \sin(\theta_{j+1} - \theta_j)
\]
gives the current from site \( j \) to \( j + 1 \), with \( \psi_j = |\psi_j|e^{i\theta_j}, -\pi < \theta_{j+1} - \theta_j \leq \pi \). Related to this is the Topological Charge (TC), which for classical vortices indicates the direction of
the energy flow. The TC is for the DNLS trimer given by

\[
TC = \frac{1}{2\pi} \sum_{j=1}^{3} \arg(\psi_j^* \psi_{j+1}) = \frac{1}{2\pi} \sum_{j=1}^{3} (\theta_{j+1} - \theta_j),
\]

which is restricted to \(0, \pm 1\) for the trimer. Turning to quantum mechanics, we can derive a corresponding continuity equation for \(\hat{n}_j\) by using the Heisenberg equation, giving

\[
\frac{d\hat{n}_j}{dt} = \hat{J}_j - \hat{J}_j, \tag{11}
\]

with

\[
\hat{J}_j = -i(\hat{a}^\dagger_{j+1}\hat{a}_j - \hat{a}_{j+1}\hat{a}^\dagger_j) \tag{12}
\]

being the current operator from site \(j\) to \(j + 1\). Comparing Eqs. (9) and (12) hints how we may calculate the quantum phase difference between sites, and thus the quantum TC. We will calculate the average value of the phase difference between two sites with [9]

\[
\left\langle \cos\left(\theta_{j+1}^{QM} - \theta_j^{QM}\right) \right\rangle = \frac{\left\langle \hat{a}^\dagger_{j+1}\hat{a}_j + \hat{a}_{j+1}\hat{a}^\dagger_j \right\rangle}{\sqrt{2(2\hat{n}_{j+1}\hat{n}_j + \hat{n}_j + \hat{n}_{j+1})}}, \tag{13a}
\]

\[
\left\langle \sin\left(\theta_{j+1}^{QM} - \theta_j^{QM}\right) \right\rangle = \frac{\left\langle i(\hat{a}^\dagger_{j+1}\hat{a}_j - \hat{a}_{j+1}\hat{a}^\dagger_j) \right\rangle}{\sqrt{2(2\hat{n}_{j+1}\hat{n}_j + \hat{n}_j + \hat{n}_{j+1})}}. \tag{13b}
\]

This will in practice be done by taking arcsin of Eq. (13b), with the range determined from the expectation value of the cosine function (13a). Replacing \(\theta_{j+1} - \theta_j\) with \(\left\langle \theta_{j+1}^{QM} - \theta_j^{QM} \right\rangle\) in Eq. (10) gives the formula used for the quantum TC. Due to quantum fluctuations in the phase operators (13), the quantum TC is not restricted to integer values, unlike the classical TC (compare with the melting of stationary vortices studied in Reference [9]).

### III. QUANTUM SIGNATURES IN EIGENSTATES

In this section we look for signatures of the classical CFVs in the energy eigenstates, and let us therefore begin by recapitulating some results for the classical CFVs [17]. The CFVs are periodic solutions (in a co-rotating frame) with period \(T\), that exist in continuous families which can be followed between the stationary dimerlike solution and the stationary SDW solution. We therefore introduce the relative Hamiltonian for the CFVs:

\[
\Delta \mathcal{H}_{rel} = (\mathcal{H}_{CFV} - \mathcal{H}_{dimerlike})/(\mathcal{H}_{SDW} - \mathcal{H}_{dimerlike}), \tag{14}
\]
FIG. 1: (Color online) First column: $|\psi_j(t)|^2$ for classical CFVs with $\beta = 1.00$. From top to bottom: $\Delta H_{\text{rel}} = 0.024$, $0.141$, $0.670$, and $0.996$. Second column: time evolution of $\langle \hat{\nu}_j \rangle = \langle CS[\psi]|\hat{\nu}_j|CS[\psi]\rangle$ for CFV coherent states with $\psi = \psi(0)$ of the classical CFV on the same row, $N = 100$ and $\alpha = \beta/(N - 1) = 1/99$. The insets show $\langle \hat{\nu}_j \rangle$ over a longer time. Third column: average currents $\langle \hat{J}_j \rangle$. Fourth column: quantum TC. Both the currents and TC are calculated from the CFV coherent state on the same row. $\langle \hat{J}_2 \rangle$ follows $\langle \hat{J}_3 \rangle$ quite closely and is hard to distinguish in the plots.
with the Hamiltonians calculated for the same nonlinear parameter $\beta$. The left column of Fig. 1 shows $|\psi_j(t)|^2$ of four CFVs that belong to the same subfamily for $\beta = 1.00$. The initial conditions are chosen so that $|\psi_1|^2$ and $|\psi_3|^2$ have minima, while $|\psi_2|^2$ has a maximum, at $t = 0$. Close to the dimerlike solution, the CFVs perform small oscillations in $|\psi_j(t)|^2$ [cf. Fig. 1(a1)], which essentially corresponds to a perturbation of the linearly stable dimerlike solution. By following the CFV subfamily towards the SDW solutions (going down along the left column of Fig. 1) the oscillations in $|\psi_j(t)|^2$ increase, so that close to the SDW solution, the CFVs essentially consist of a ‘hole’ (an empty site) that goes back and forth between two sites [cf. Fig. 1(d1)]. These CFVs can be viewed as the result of perturbing the linearly unstable SDW solution. The SDW solution is linearly unstable, through a non-oscillatory instability, for all positive values of $\alpha$ in (5). This is in contrast to when $\alpha$ is negative, where the SDW solution becomes oscillatory unstable in a parameter window, but is linearly stable otherwise [32]. Note that the initial conditions in the left column of Fig. 1 are chosen to be as close as possible to a SDW in this limit.

A different perspective on the classical CFVs is given in terms of localized discrete breathers. The dimerlike solution is then the on-site centered breather ground state, while the SDW solution is an inter-site centered breather. The quantity $H_{SDW} - H_{dimerlike}$ is interpreted as the Peierls-Nabarro (PN) potential barrier that needs to be overcome to transfer the on-site breather to its neighbor site [33]. The CFVs can thus be considered to be breather states that do not overcome the PN-barrier ($\Delta H_{rel} < 1$), but instead oscillate back and forth in the PN potential around the minimum of the on-site centered ground state. Let us point out that Ref. [26] uses a DNLS model with opposite signs on both the neighbor coupling and nonlinear parameter, meaning that the dimerlike solution (on-site centered breather) is the highest eigenstate rather than the ground state. A corresponding single-peaked solution exists also for the DNLS trimer with opposite sign only on the coupling. This solution exists above a certain threshold value for the nonlinear parameter, where it bifurcates from another solution [27], and it also becomes the ground state above another parameter value [34].

Consider now Fig. 2(a), which plots the trajectories of some classical solutions from the same subfamily as in the left column of Fig. 1. The black triangle is the border of the accessible area defined by $\sum_j |\psi_j|^2 = 1$. The black crosses mark the dimerlike solutions and the black circles the SDW solutions, while the orange (dark gray) and yellow (light gray) curves are the same CFVs as in Figs. 1(b1) and 1(d1), respectively. The CFVs
FIG. 2: (Color online) (a) Trajectories of classical solutions for $\beta = 1.00$. The black crosses mark dimerlike solutions while the black circles mark the SDW solutions. The orange (dark gray) and yellow (light gray) lines are the CFVs of Figs. 1(b1) and 1(d1), respectively. For (b)-(j) $N = 100$ and $\alpha = \beta/(N-1) = 1/99$. (b)-(g): $|\langle n_1, n_2, n_3|E_m(k)\rangle|^2/\max_{n_1, n_2, n_3}(|\langle n_1, n_2, n_3|E_m(k)\rangle|^2)$, where (b)-(f): $k = 0$ and $m = 1, 2, 3, 7, 11$ (energy increasing from (b) to (f)), and (g): $k = 2\pi/3$ and $m = 11$. (h) Lower part of the energy spectrum for $k = 0$ (crosses) and $k = 2\pi/3$ (circles). $N\mathcal{H}_{SDW}$ and $N\mathcal{H}_{dimerlike}$ are include as the top and bottom dashed lines, respectively. (i) $\Gamma_m$ of Eq. (15). (j) $2\pi/|E_m(0) - E_m(2\pi/3)|$.

are thus represented as arcs in Fig. 2(a), with CFVs close to the dimerlike solution being small arcs close to the black cross, similar to the orange (dark gray) one. Moving along the CFVs subfamily towards the SDW solution means that the arc grows, until it touches the two circles of the SDW solution, similar to the yellow (light gray) arc. A larger arc thus corresponds both to larger oscillations in $|\psi_j|^2$, and to a larger energy (Hamiltonian). In this discussion, we have assumed that the third site is the most occupied, but the CFVs can be translated in the lattice, corresponding essentially to a rotation of the triangle.

Turning now to the quantum eigenstates, Figs. 2(b)-(f) display $|\langle n_1, n_2, n_3|E_m(0)\rangle|^2/\max_{n_1, n_2, n_3}(|\langle n_1, n_2, n_3|E_m(0)\rangle|^2)$ for $m = 1, 2, 3, 7, 11$ with $N = 100$ and $\alpha = \beta/(N-1) = 1/99$, with the energy increasing from Fig. 2(b) to 2(f). Fig. 2(g) shows $|\langle n_1, n_2, n_3|E_{11}(2\pi/3)\rangle|^2/\max_{n_1, n_2, n_3}(|\langle n_1, n_2, n_3|E_{11}(2\pi/3)\rangle|^2)$ for the same $N$ and
As can be seen in these plots, the eigenstates spread out more over the triangle when increasing the energy. The size of the bright lobes in the plots indicate the quantum uncertainty of $\nu_j$, which shrinks as $N$ is increased. Fig. 2(h) shows the energy spectrum of the 13 lowest eigenstates in each $k$-space, with $NH_{\text{dimerlike}}$ and $NH_{\text{SDW}}$ included as dashed, red lines. The line for the dimerlike solution does not coincide with the ground state, but it should approach it as $N$ is increased further. The region between the dashed lines indicate (approximately) which eigenstates that will contribute to the quantum CFVs.

Due to the Bloch theorem, all eigenstates have a translational invariance, which is manifested in the similarity of the triangles in Figs. 2(b)-(g) when rotated $\pm 2\pi/3$. More classical-like (symmetry-broken) states can be obtained by adding together eigenstates from different $k$-spaces with suitable coefficients. The basic idea is to cancel out certain bright lobes in the probability distribution plots, leaving them only in one third of the triangle. In order for this to work well, it is necessary that we combine eigenstates with similar occupation of Fock states, i.e. similar triangular probability-distribution plots. A plot corresponding to Fig. 2 for $k = 2\pi/3$ would reveal that the $m$-th lowest eigenstates in each $k$-space look similar, especially for low $m$. As a quantitative measure of how similar they are, we introduce

$$
\Gamma_m = \sum_{n_1,n_2,n_3} \left| \langle n_1,n_2,n_3|E_m(0)\rangle \right|^2 - \left| \langle n_1,n_2,n_3|E_m(2\pi/3)\rangle \right|^2.
$$

(15)

Note that this quantity completely disregards the phase between the Fock states. This quantity is shown for $m = 1, \ldots, 13$ in Fig. 2(i), and $\Gamma_m$ is very small for $m < 9$, but increases for larger $m$. Figs. 2(f)-(g) are for $m = 11$, and $k = 0$ and $k = 2\pi/3$, respectively, and it is evident that these plots differ quite significantly. Note also that the larger $\Gamma_m$ for $m > 9$ is associated with an increased energy splitting in Fig. 2(j). The 'lifetime' of the localization, obtained from the $m$-th lowest eigenstates, is associated with the inverse of the energy splitting $\Delta E_m = E_m(0) - E_m(2\pi/3)$, plotted in Fig. 2(j).

This type of translational symmetry breaking is also relevant for e.g. quantum discrete breathers, and has been studied extensively in this context [4]. The arguably simplest example is for the Bose-Hubbard dimer, where the quantum breather (or the 'self-trapped' state) consists of a superposition of the two lowest quasi-degenerate eigenstates, one being symmetric and one anti-symmetric under site-permutation (a 'tunneling pair'). The lifetime of the localization is given by the inverse energy splitting[6].

The eigenstate in Fig. 2(b) can be identified with the stationary dimerlike solution, noting
that the three lobes in Fig. 2(b) and the black crosses in Fig. 2(a) have approximately the same positions. \( \Gamma_1 \) in Fig. 2(h) is very small, and so is the corresponding energy splitting, which means that a very long lived localized quantum state can be created, consisting essentially of one bright spot in a probability distribution plot. Reference [11] studied the ground state of the Bose-Hubbard Hamiltonian in more detail, however with a positive hopping term in Eq. (1). The results in [11] are however connected to our work in the strong nonlinearity limit, and the reader is directed there for further details on this ground state. We note only that Fig. 2(b) is qualitatively similar to the plots in Reference [11] for the ground state above the self-trapping transition.

The SDW solution can on the other hand be associated with the eigenstate in Fig. 2(f), which has bright spots at the border of the triangle, at positions corresponding to the classical SDWs (easiest to see at the bottom of the triangle). There is however no eigenstate with \( k = 2\pi/3 \) that ‘matches’ this eigenstate well, as can seen both in the energy spectrum, and that \( \Gamma_{11} \) is rather big. The stationary SDW solution is thus more difficult to recreate quantum mechanically compared to the dimerlike solution, which is expected since the SDW is unstable and the dimerlike solution stable.

Looking now at Figs. 2(c)-(g), a clear signature of the classical CFV trajectories of Figs. 2(a) is that the probability distributions are spread out over a region corresponding to where the arcs of the classical CFVs are located. The probability distributions also widen out more for increasing energy, in agreement with the increasing span of the arcs in Fig. 2(a) for higher energy. Reference [15] made a similar comparison between eigenstates and non-flipping vortices. Let us however emphasize that CFVs, unlike the dimerlike solution, are not stationary solutions, and will therefore not correspond to a single eigenstate in each \( \mathbf{k} \)-space, but rather a superposition of several. It might be tempting to identify an arc in Fig. 2(a) with an eigenstate with a similar width, but we should remember that the arc represents the whole trajectory of the CFV. Dynamically, a classical CFV is represented by a single point in Fig. 2(a) which moves back and forth along the arc. The dynamics of the corresponding quantum CFV will thus be a blob that moves in the probability distribution plot, which is a result of the interference of several eigenstates. As \( N \) is increased, the extension of the bright spots will decrease. Since the total number of eigenstates also increases with \( N \), and there are more eigenstates available that can contribute to the CFVs, the resulting dynamics also becomes richer, so that the blob can follow the classical point more accurately. We will

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see in the next section that our expectations on quantum CFVs, which have been discussed in this paragraph, are confirmed for the appropriate coherent states.

When calculating the average currents and quantum TC for the eigenstates, the currents are zero for \( k = 0 \) due to symmetry. The classical TC would in this case be equal to zero, which the quantum TC sometimes is, but due to fluctuations in the cosine operator (13a), it may also take the value \(-1.5\). For \( k = 2\pi/3 \) on the other hand, the average currents can take non-zero values. All three currents are the same, since the states are stationary, but the sign of \( k \) does not indicate their direction. The quantum TC will generally not take the classically allowed integer values, but end up in an interval either between \(-0.5\) and \(0.5\), or between \(\pm 1\) and \(\pm 1.5\), this again due to quantum fluctuations. We will return to a discussion on the validity of the quantum TC later on.

IV. CFV COHERENT STATES

A. Dynamics

In this section we study quantum time evolution of initial CFV coherent states: coherent states obtained by inserting a classical CFV state vector \( \psi \) into Eq. (3). Consider again Fig. 1, and recall that the first column shows \( |\psi_j(t)|^2 \) for four classical CFVs that belong to the same continuous CFV subfamily with \( \beta = 1.00 \). The second column of Fig. 1 displays the fully quantum time evolution of \( \langle \hat{\nu}_j \rangle = \langle CS[\psi]|\hat{\nu}_j|CS[\psi] \rangle \), with \( \psi \) the initial condition of the classical CFV on the same row, for \( N = 100 \) and \( \alpha = \beta/(N-1) = 1/99 \). Since the classical CFVs are non-stationary solutions, using \( \psi = \psi(t) \) in Eq. (3) with different \( 0 \leq t < T \) results in different coherent states, and some effects of this will be discussed in Sec. IV C. The insets in Figs. 1(b2) and 1(c2) show \( \langle \hat{v}_1 \rangle \) and \( \langle \hat{v}_2 \rangle \) over a longer time span. The third column shows the average value of the current operators (12) while the fourth column shows the quantum TC, both calculated with the quantum time evolved CFV coherent state on the same row. Note that the average value of all three current operators are plotted in the third column, but that they follow each other quite closely (especially \( \langle \hat{J}_2 \rangle \) and \( \langle \hat{J}_3 \rangle \)), and therefore are difficult to distinguish on this scale.

From these plots it is evident that the closer \( \psi \) are to the dimerlike solution, the longer and more accurately do \( \langle \hat{\nu}_j \rangle \) follow \( |\psi_j|^2 \). There is also a clear similarity in the top three
rows between the oscillations in $\langle \hat{\nu}_j \rangle$ and $\langle \hat{J}_j \rangle$. Note though that the changes in $\langle \hat{\nu}_j \rangle$ are connected to the difference between the average currents through Eq. (11), which may be difficult to distinguish in the large plots, but is visible in the inset of Fig. 1(b3). The arguably most important aspect of the third column is however that the average of all three currents follow each other closely, even in the bottom row, and that currents repeatedly change direction. This is a clear signature of the CFVs, since all currents (at least their averages) are going in the same direction, and also flip at approximately the same time.

The flipping is very close to periodic in the top three rows, with the exception of a period $100 \lesssim t \lesssim 150$ in Fig. 1(c3) when the oscillations in $\langle \hat{J}_j \rangle$ are small and rapid, and the charge flipping breaks down. This is during the time in Fig. 1(c2) when the amplitude of the oscillations in $\langle \hat{\nu}_j \rangle$ are small. It is remarkable that even in the bottom row, when $\langle \hat{\nu}_j \rangle$ deviates strongly from $|\psi_j|^2$, we may still discern a charge flipping in $\langle \hat{J}_j \rangle$, and that it also is somewhat regular.

Turning to the quantum TC in the column to the right, there is a periodic flipping between $\approx \pm 1$ for the top three rows. The exception is once again the region $100 \lesssim t \lesssim 150$ in Fig. 1(c4). Note though that the TC is generally not equal to exactly $\pm 1$ due to quantum fluctuations in the angles. The fluctuations in TC are of the order $10^{-3}$ in Figs. 1(a4) and 1(b4), while it is between $10^{-3}$ and $10^{-2}$ for Fig. 1(c4). In Fig. 1(d4) the quantum TC jumps to $\approx 1.5$ and $\approx 0.5$ at certain points, which is mainly due to large quantum fluctuations in the cosine function Eq. (13a). We are in fact in a parameter regime where the fluctuations and uncertainty in $\cos\left(\theta^{(QM)}_j - \theta^{(QM)}_{j+1}\right)$ are large (compare again with the melting of stationary vortices in Reference [9]), even for the top two rows, and one may argue that the angles and the quantum TC therefore are not well defined quantities. We note however that by using this particular convention for the quantum TC, it does indicate the direction of the currents and thus the vortex (even in Fig. 1(d4), where the quantum TC takes values different from $\pm 1$, the sign still indicates the currents direction).

Fig. 3 shows a different perspective on the CFV coherent states’ quantum time evolution, displaying the probability distribution for $|CS[\psi](t)|$ at different $t$ (cf. end of Sec. III). The plots marked with (ax), (bx), and (cx) are the time evolved CFV coherent states shown in Figs. 1(b2), 1(c2), and 1(d2), respectively. The time $t$ is shown in the top right corner of each plot, and the location of the corresponding classical CFV $\hat{\psi}(t)$ (with $\nu_j \rightarrow |\psi_j|^2$) is marked with a blue (gray) cross.
FIG. 3: (Color online) Time evolution of the probability distribution $|\langle n_1, n_2, n_3 | CS[\psi](t) \rangle|^2$ of the CFV coherent states corresponding to: (a1)-(a5) ↔ Fig. 1(b2); (b1)-(b10) ↔ Fig. 1(c2); (c1)-(c10) ↔ Fig. 1(d2). The time $t$ is shown in the top right corner of each plot. The corresponding position of the classical associated CFVs are marked with blue (gray) crosses. In (c1), (c4), and (c8) the probability distribution is localized close to the triangle border and may be difficult to distinguish.
Figs. 3(a1)-(a5) show the probability distribution over a time when the evolution of $\langle \hat{\nu}_j \rangle$ in Fig. 1(b2) follows the corresponding $|\psi_j|^2$ well. The probability distribution in Figs. 3(a1)-(a5) therefore forms a localized bright spot that follows the classical solution well without much spreading over the time shown.

There is however a clear deviation between Figs. 1(c1) and 1(c2) over the time span of Figs. 3(b1)-(b10). This deviation is also reflected in Figs. 3(b1)-(b10), where we can see a spreading of the probability distribution over a region that corresponds to the classical trajectory. This spreading is related to the reduced amplitude for the oscillations in $\langle \hat{\nu}_j \rangle$, seen in Fig. 1(c2). The blob in the probability distribution is however well localized around the classical solution initially. The CFV is in some sense still preserved (which also is indicated by Fig. 1(c3)), but with quantum mechanics introducing an uncertainty about where along the trajectory of the classical CFV the quantum state is. The probability distribution shown in Figs. 3(a1)-(a5) will spread out in a similar way at $t \approx 200$, as can be seen from the reduced amplitude in Fig. 1(b2). This type of spreading is oscillatory, meaning that the blob in the probability distribution alternates between spreading out and contracting, which can be seen for the amplitude modulations in the insets of Figs. 1(b2) and 1(c2). This type of spreading and contracting, and thus also the amplitude modulation of $\langle \hat{\nu}_j \rangle$, occurs in a more regular way for Fig. 1(b2) than for Fig. 1(c2), which also can be seen in the insets.

The deviation between Figs. 1(d1) and 1(d2) is almost instant, and the probability distribution is not clearly centered around the classical CFV in any of Figs. 3(c2)-(c10). Comparing Figs. 3(c1)-(c10) with the three top rows, we can see that there is a spreading of the probability distribution over the whole triangle (along a circular shape), whereas the probability distribution is restricted to the top of the triangle in the top three rows. The charge flipping is however not completely destroyed by this type of spreading either, which is indicated also by Fig. 1(d3), but it rather delocalizes the CFV over the trimer.

A crucial difference between the classical and quantum CFVs, is that for classical CFVs the same site always is the mostly populated. This is also true even if the exactly periodic CFVs are slightly perturbed, since isolating KAM-tori strictly forbid classical spreading (cf. [32]). For the quantum CFVs on the other hand, quantum fluctuations inevitably causes dynamical tunneling across the classical KAM-tori, and the probability distributions will eventually spread over the whole triangle (along a circular shape) also for the top three rows in Fig. 3, however on a much longer time scale. For these CFV coherent states the spreading
FIG. 4: (Color online) $\alpha = 1.00/(N - 1)$ and $N = 100$. (a),(b) $|\langle E_m(k)|CS[\psi]\rangle|^2$ for the lowest eigenstates with (a) $k = 0$ and (b) $k = 2\pi/3$, with $\psi$ following the CFV subfamily with $\beta = 1.00$ from the dimerlike to SDW solution. All $\psi$ are chosen so that site 1 has an amplitude minimum, similar to the initial conditions in the first column of Fig. 1. (c)-(f) show these projections in more detail for the four CFV coherent states in Fig. 1(a)-(d), respectively, with crosses for $k = 0$ and circles for $k = 2\pi/3$.

is related to the energy splitting between the $k$-spaces [cf. Fig. 2(j)].

**B. Projection on eigenstates**

Let us now look at how the CFV coherent states project onto the energy eigenstates. Figs. 4(a)-(b) displays $|\langle E_m(k)|CS[\psi]\rangle|^2$ for $N = 100$, where $\psi$ follows the same continuous family of classical CFVs as in Figs. 1 and 2, running from the dimerlike to the SDW solution. The configurations of all $\psi$ are chosen so that site 1 and 3 have amplitude minima, while site 2 has a maximum, similar to the initial conditions in the first column of Fig. 1. As the CFV family is followed from the dimerlike to the SDW solution, the projection onto higher
eigenstates increases, and the CFV coherent states project on several eigenstates, as was anticipated in Sec. III.

In Figs. 4(c)-(f) we take a closer look at $|\langle CS[\psi]\mid E_m(k)\rangle|^2$ for the four CFV coherent states in Fig. 1. Fig. 4(c) has a large projection for $m = 1$, a smaller one for $m = 2$, and practically zero projection for $m \geq 3$. The frequency of the oscillations of $\langle \hat{\nu}_j \rangle$ in Fig. 1(a2) is given by $E_2(k) - E_1(k)$ (the energy splittings between the $k$-spaces for $m = 1, 2$ are very small in comparison, see Fig. 2(j)). Inserting numerical values gives $2\pi/(E_2(k) - E_1(k)) \approx 21.9$, while $T \approx 21.7$ for the $|\psi_j|^2$-oscillations in Fig. 1(a1).

In Fig. 4(d) the largest projection is no longer on the lowest eigenstates, but has moved up to $m = 2$. There is also a significant population on more than two eigenstates for each $k$, namely for $m = 1, 2, 3$. The amplitude modulation seen in Fig. 1(b2) is related to the difference between $E_3(k) - E_2(k)$ and $E_2(k) - E_1(k)$, and is thus essentially the familiar beating phenomenon, caused by the interference of two closely lying frequencies. The time period for the amplitude modulation is given by $2\pi/|E_1(k) - 2E_2(k) + E_3(k)| \approx 472$, which agrees well with the inset of Fig. 1(b2). Note that the quantity $|E_1(k) - 2E_2(k) + E_3(k)|$ is a measure of how much these three energy levels deviate from lying on a straight line, see Fig 2(h) (cf. the energy spectrum of the harmonic oscillator). Thus, the development of this modulation can be seen as a quantum signature of the increasing anharmonicity of the classical CFV oscillations, when moving away from the regime of linear internal-mode oscillations around the breather ground state. The time period of the classical-like oscillations in $\langle \hat{\nu}_j \rangle$ is in this case related to $4\pi/(E_3(k) - E_1(k)) \approx 22.4$, which can be compared with $T \approx 22.6$ in Fig. 1(b1).

In Fig. 4(e) there are significant projections onto even more eigenstates than in Fig. 4(d), with the peak at $m = 7$. The dynamics in Fig. 1(c2) also shows an amplitude modulation, but not in an as regular and periodic manner as in Fig. 1(b2) (compare the insets). This can again be understood as a manifestation of the beating phenomenon, but this time with more contributing frequencies. Remember also that this type of amplitude modulation is associated with a spreading of the probability distribution over a region that corresponds to the trajectory of the classical CFV, as seen in Fig. 3(b1)-(b10).

The time evolution of all three CFV coherent states above will eventually be affected by the energy splitting between the different $k$-spaces. The probability distribution will spread out over all thirds of the triangle in plots like Fig. 3, and the values of all $\langle \hat{\nu}_j \rangle$ will thus
FIG. 5: (Color online) $\alpha = 1.00/(N - 1)$ and $N = 100$. First row: $|\langle E_m(0)|CS[\psi(t)]\rangle|^2$ for the four classical CFVs shown in Fig. 1 for different $t$, with $T$ being the period of $|\psi(t)|^2$. Going from left to right corresponds to going from top to bottom in Fig. 1. The time $t$ only needs to run over one quarter of a period, because of symmetries in the classical CFVs dynamics. (e),(f) Time evolution of $\langle \hat{\nu}_j \rangle$ for the CFV coherent states of (c) and (d), respectively, with $t = T/4$. The line coloring is the same as in Fig. 1. (g),(h) Average currents, with (g) connected with (e), and (h) with (f).

approach 1/3. The time scale for this is determined by which eigenstates that are occupied and the $k$-splitting for these [see Fig. 2(j)]

Turning now to Fig. 4(f), the dynamics of $\langle \hat{\nu}_j \rangle$ in Fig. 1(d2) shows an almost instant deviation from the corresponding $|\psi_j|^2$, and $\langle \hat{\nu}_j \rangle$ also shows a more complex time evolution compared to the other three considered CFV coherent states. The obvious difference between Fig. 4(f) and Figs. 4(c)-(e) is that the circles do not overlap with the crosses for the populated eigenstates, meaning that the occupations of the $m$-th lowest eigenstates with $k = 0$ and $k = 2\pi/3$, respectively, differ. This is also connected to that the occupied eigenstates have a large $\Gamma_m$ [cf. Fig. 2(i)].
C. CFV coherent states with other $t$

We have up until now only considered CFV coherent states with a specific choice of $\psi$, namely $\psi = \psi(0)$, with $t = 0$ set so that $|\psi_1(0)|^2$ and $|\psi_3(0)|^2$ have minima, and $|\psi_2(0)|^2$ has a maximum. Fig. 5 shows how the projection of the CFV coherent states onto eigenstates changes when altering $t$ of $\psi = \psi(t)$ in Eq. (3). The four plots (going from left to right) use $\psi(t)$ from the four classical CFVs in the first column in Fig. 1 (going from top to bottom). It is only necessary that the time $t$ runs from 0 to $T/4$ in Fig. 5, due to symmetries in the classical CFVs (cf. first column of Fig. 1).

The CFV coherent states are more spread out over the eigenstates for $t \approx T/4$ and it is around $t \approx T/4$ that $|\psi_j|^2$ change most rapidly for the classical CFVs ($|\psi_3|^2$ is stationary at $t = T/4$, but changes quite rapidly in the vicinity), especially compared to $t = 0$. The difference between $t = 0$ and $t = T/4$ is not so pronounced in Figs. 5(a)-(b), and using CFV coherent states with $\psi = \psi(T/4)$ from these plots would result in a similar evolution of $\langle \hat{\nu}_j \rangle$ as in Figs. 1(a2) and 1(b2), respectively. For Figs. 5(c)-(d) on the other hand, there is a clear difference between $t = 0$ and $t = T/4$, and the time evolution of $\langle \hat{\nu}_j \rangle$ for the CFV coherent states with $\psi = \psi(T/4)$ from these plots are shown in Figs. 5(e)-(f). The oscillations in $\langle \hat{\nu}_j \rangle$ decay faster, especially compared to Fig. 1(c2), and it is also difficult to discern the classical oscillations after some time. The associated average currents are plotted in Figs. 5(g)-(h), and even though there are oscillations in the currents, they do not change the direction. Changing $t$ in $\psi(t)$ for the CFV coherent states will thus destroy the charge flipping of the vortex, for these $\psi$. Note finally that it is at $t = nT/2$, $n \in \mathbb{Z}$, that Fig. 1(d1) is closest to the configuration of a SDW solution, and also where Fig. 5(d) is most concentrated on one eigenstate.

V. CHANGING $N$ AND NONLINEAR PARAMETER

We will in this section analyze how certain results from Sec. IV depend on the number of bosons $N$, focusing primarily on $N = 10$ and $N = 30$. Consider Fig. 6, where the top row is for $N = 10$ and the bottom for $N = 30$. From Fig 6(a)-(b) we see that $\Gamma_m$ is noticeably larger than zero for all $m$ when $N = 10$, and that the corresponding energy splitting is significant for $m \geq 2$ ($|E_1(0) - E_1(2\pi/3)| \approx 0.02$). From Figs. 6(c)-(d) we see that the CFV
coherent states occupy mainly the three lowest eigenstates for each $k$ for $N = 10$.

From the bottom row we can see that $\Gamma_m$, and the corresponding energy splitting, is rather small for $m < 3$, while the CFV coherent states occupy the five lowest eigenstates in each $k$-space.

Consider now Fig. 7, which in the top row shows $\langle \hat{\nu}_j \rangle$ for the CFV coherent states corresponding to those in the second column of Fig. 1, but with $N = 30$ (going left to right in Fig. 7 corresponds to top to bottom in Fig. 1). The second row shows the associated average currents, and the third row the CFV coherent states projection onto energy eigenstates. From this plot we can see that many of the results seen earlier for $N = 100$ also hold for $N = 30$. There is an amplitude modulation in Fig. 7(c1) which is related to the significant occupation of more than two eigenstates in each $k$-space. In Figs. 7(c3) and 7(d3) we can also discern differences in the occupation between some of the $m$-th lowest eigenstates in either $k$-space, which is connected to non-vanishing $\Gamma_m$ seen in Fig. 6(f). This difference also becomes more distinct for $\psi$ closer to the unstable SDW solution, i.e. more distinct in Fig. 7(d3) than in Fig. 7(c3). Note also the difference between Figs.
FIG. 7: (Color online) $N = 30$, $\alpha = 1.00/(N - 1)$. Top row: Time evolution of $\langle \hat{\nu}_j \rangle$ for CFV coherent states corresponding the those in Fig. 1 but with $N = 30$. The line coloring is the same as in Fig. 1. Second row: Associated average currents. Bottom row: Projection of CFV coherent states onto energy eigenstates.

7(c3) and 4(e), which are for the same $\psi$, where the circles and crosses overlap well in Fig. 4(e), but not in 7(c3). For higher $N$, we may thus follow the CFV family closer to the SDW solution before $k$-splitting becomes important. A notable difference between Figs. 7(b3) and 4(d) is that the peak is at the ground state for $N = 30$, but it has moved up to $m = 2$ for $N = 100$. Increasing $N$ thus 'pushes' the CFV coherent state up along the eigenstates.

Fig. 8 corresponds to Fig. 7 but with $N = 10$. In this case $\Gamma_m$ [Fig. 6(b)] is not close to zero for any $m$, and we can see in the third row that there are noticeable differences for projections onto the $m$-th eigenstate in either $k$-space (for the occupied eigenstates). In Figs. 8(a1), 8(b1) and 8(c1) we can also see the effect of the energy splitting between the $k$-spaces, with the slow, large amplitude oscillations. $2\pi/|E_1(0) - E_1(2\pi/3)| \approx 150$ which agrees well with the oscillations. Figure 9(a) shows $2\pi/|E_1(0) - E_1(2\pi/3)|$ as a function of $N$, and thus indicates the time scale when this type of mixing becomes important, which is relevant for Figs. 1(a2), 7(a1), 8(a1), and 8(b1). This time scale is rather long already for $N = 20$ ($\approx 5 \cdot 10^3$) compared to the time period of the classical oscillations. Figure 9(a) also shows $2\pi/|E_m(0) - E_m(2\pi/3)|$ for $m = 2, 3$, which come into play as these eigenstates
become occupied, e.g. in Fig. 1(b2) (cf. Fig. 4). This time scale is according to Fig. 9(a) decreasing with \( m \) for fixed \( N \).

We can distinguish a charge flipping in the average currents for all the time evolved CFV coherent states in Figs. 7 and 8, which is more regular for \( \psi \) closer to the dimerlike solution.

We have so far only considered a specific family of CFVs, which was for \( \beta = 1.00 \) (and with \( N = 1 \)). Most of the results that we have obtained are however also valid for CFV families with other \( \beta \). For example will \( \langle \hat{\nu}_j \rangle \) follow the corresponding \( |\psi_j|^2 \) better for CFVs closer to the stable dimerlike solution, and it will also be possible to distinguish charge flipping in both the average currents and the quantum TC, even in the regimes where \( \langle \hat{\nu}_j \rangle \) deviates strongly from \( |\psi_j|^2 \). A comments on this is necessary. For \( \beta \gtrsim 1.8 \) it is not possible to follow the CFV subfamilies (with fixed \( \beta \)) from the dimerlike all the way to the SDW solution [17]. CFVs do however exist in the vicinity of the dimerlike solution, and the results discussed above hold also for them.

It is thus generally possible to determine the timescale for when \( \langle \hat{\nu}_j \rangle \) starts to deviate from \( |\psi_j|^2 \) by looking at which eigenstates that are occupied, and how their energies are related, as discussed in Sec. IV. Figures 9(b)-(c) show how the quantity plotted in Fig. 2(j) depends on \( \alpha \) for fixed \( N \), and from these figures we conclude that the timescale where the mixing of eigenstates with different \( k \) becomes important is increasing with both \( N \) and \( \alpha(N-1) \) (at

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FIG. 8: (Color online) Similar to Fig 7, but for \( N = 10 \).
FIG. 9: (Color online) (a) $2\pi/|E_m(0) - E_m(2\pi/3)|$ for $\alpha = 1.00/(N - 1)$. (b) $2\pi/|E_m(0) - E_m(2\pi/3)|$ for (from top to bottom), $m = 1, 2, 3, 4, 5$, and $N = 100$. (c) $2\pi/|E_1(0) - E_1(2\pi/3)|$. (d) $2\pi/|E_m(k) - 2E_{m+1}(k) + E_{m+2}(k)|$ for $N = 100$. The curves for $k = 0$ and $k = 2\pi/3$ are almost identical.

least for the plotted $N > 10$). Figure 9(d) instead shows $2\pi/|E_m(k) - 2E_{m+1}(k) + E_{m+2}(k)|$ for $N = 100$, which indicates that the timescale for the amplitude modulation shown in e.g. Fig. 1(b2) is decreasing with $\alpha$ for fixed $N$, related to the fact that a stronger nonlinearity makes classical oscillations more anharmonic (at given amplitude).

VI. SUMMARY AND CONCLUSIONS

We have in this paper studied the Bose-Hubbard model, and searched for quantum signatures of classical CFVs from the corresponding DNLS trimer. Several such signatures have been identified, which illuminate different aspects of the quantum-classical correspondence.

Some signatures can be seen in the energy eigenstates when comparing the site amplitudes of the classical CFVs with the eigenstates’ probability distributions over the different Fock states $|n_1, n_2, n_3\rangle$. This is conveniently illustrated with the triangular plots in Fig. 2 (a
pedagogical advantage of the trimer) where the eigenstates’ probability distributions occupy
the corresponding region of the classical CFVs’ trajectories.

We have furthermore considered certain CFV coherent states, and especially their dyna-
matic properties. Quantum signatures can then be observed when comparing the evolution
of the classical site amplitudes $|\psi_j|^2$ with the quantum average number of particles $\langle \hat{\nu}_j \rangle$.
The classical CFVs exist in continuous families located between the linearly stable dimer-
like solution (which also is the "breather" ground state) and the unstable SDW solution,
and $\langle \hat{\nu}_j \rangle$ follows $|\psi_j|^2$ better for CFVs that are closer to the dimerlike solution. We have
been able to identify certain deviations between $\langle \hat{\nu}_j \rangle$ and $|\psi_j|^2$ with properties of the energy
spectrum, which also gives the timescales for when these deviations become important. The
time evolution of the coherent states’ probability distribution over the Fock states has also
been considered. Plotting this, the CFV coherent states form blobs in the probability dis-
tribution that follow the trajectory of the associated CFV. The different types of deviations
that occur between $\langle \hat{\nu}_j \rangle$ and $|\psi_j|^2$ is related to a spreading of this blob.

The most prominent quantum CFV signatures is arguably seen in the currents and quan-
tum TC of the CFV coherent states. The average currents follow each other quite closely,
and also flip at approximately the same time, even in the regimes where $\langle \hat{\nu}_j \rangle$ and $|\psi_j|^2$ de-
viate strongly. The quantum CFVs are in this sense rather robust. Related to this is the
quantum TC, where its sign indicates the currents direction, even though it in some regimes
takes classically forbidden non-integer values. It was however found that for certain CFVs,
close to the SDW solution, it matters which $\psi = \psi(t)$ that is used in the CFV coherent
states, and that for some choices the charge flipping is destroyed.

We hope that our work will inspire experimental efforts in realizing charge flipping vor-
tices with BECs in optical lattices, e.g. with the technique of painting potentials [16].
The realization of optical potential rings, discussed in e.g. [35], would also provide a very
promising candidate.


