Numerical Computation of Dynamically Important Excited States of Many-Body Systems

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Dated: May 15, 2012

PACS numbers: 67.85.Hj, 03.75.Kk, 03.75.Lm

I. INTRODUCTION

Properties of many-body quantum systems are difficult to compute, because the eigenstates are usually unknown and the large size of the Hilbert space makes even approximate descriptions difficult to manipulate. Some numerical methods - such as variational methods or mean-field theories - are intrinsically approximate yielding simple expressions describing most of the interesting physical properties, especially in the thermodynamic limit. The so-called “quasi-exact” methods do not have such restrictions, and are supposed to converge to the exact quantum result when some parameter tends to infinity. Those include exact diagonalization methods and quantum Monte-Carlo methods1,2 (the limiting parameter being the statistical sampling). For one-dimensional (1D) systems, particularly effective is the Density Matrix Renormalization Group (DMRG) approach3,4. The states, in modern implementations of the method, are represented by the so-called Matrix Product States (MPS) (for a recent smooth introduction, see5). In numerous quantum systems, the ground state is very well approximated by a MPS, allowing its efficient description with a relatively small number of parameters, which, moreover, does not increase exponentially with the system size. To some extent, DMRG shares the advantages of both variational and quasi-exact methods.

While the original application of these methods considered stationary properties, recent years brought a big progress in addressing the dynamics. Arguably, it was the formulation of the Time Evolving Block Decimation (TEBD) algorithm6, enabling study of real time dynamics of 1D systems of a reasonable size, which triggered a rapid progress in the field. Soon, a time-dependent DMRG (t-DMRG) method was proposed7,8, with a fast realization of a close equivalence between TEBD and t-DMRG. Various applications of the method addressed dynamics of quantum phase transitions9,10, quantum quenches and thermalizations11,12, periodic driving16,17, to give just a few examples.

As soon as the evolution of the system ceases to be adiabatic, the excited states become important in the dynamics. The global properties of whole groups of excited states may be obtained from evaluation of e.g. the spectral function. Following the pioneering work of Hallberg18, a lot of effort has been put in developing efficient schemes for spectral functions evaluations (see e.g.19–23 and references therein).

In the present paper, our aims are somehow complementary. We show how one may obtain selected, dynamically important excited eigenstates of strongly correlated systems in a relatively simple way. Instead of addressing spin chain systems, as typical for tests of various DMRG improvements, we shall immediately turn our attention to cold atoms settings and consider a Bose-Hubbard model as realized by cold atoms in 1D optical lattice potentials. In11, we have shown on such a specific example how to extract the energies of the few lowest excited states of the system, with, however, no access to their properties. In the present paper, we explain how to use the TEBD (or t-DMRG) method in order to extract some of the excited states with quite good accuracy. Those states are expressed also in terms of MPS making the calculations of expectations values and correlation functions easy. While the area law theorem24 assures that the ground state may be efficiently represented in MPS representation this is by no means guaranteed for a generic excited state. Our method will work only for those states for which such an efficient MPS representation exists. We shall show, however, that this restriction leaves plenty of interesting ground to explore as, most often, many excited states up to a reasonably large excitation energy, are also efficiently expressed as MPSs. Interestingly, the method selects states that are important in the systems’ dynamics. It seems more suited for low lying excited states (although we give other examples too). Still, in many-body systems, the density of states increases usually very fast with energy, and even relatively small systems (such as e.g. encountered in ul-
tricold atomic gases) have plenty of excited states lying not far above the ground state energy.

While we use as an example the Bose-Hubbard (BH) model describing an ultra-cold atomic bosonic gas loaded in a one-dimensional optical lattice, the method proposed is quite general and can be used for any system where TEBD (t-DMRG) works for sufficiently long times. The latter restriction is necessary since the method is based on Fourier Transform techniques with resolution being dependent on the integration time. We shall show later that the method has an unexpected self-purifying property allowing sometimes to use much longer integration times than anticipated in standard applications of the algorithm \[6, 7\]. Possible future improvements of the method are discussed in the concluding section.

In the following sections, we first describe the method in some details. Then, we apply it for extracting selected excited states populated either by the periodic driving or by the non-adiabatic dynamics in the disordered Bose-Hubbard system. The illustrative examples of excited states show the power and the limitations of our approach.

II. EXTRACTION OF EIGENSTATES BY FOURIER TRANSFORM

The basic idea is to start from an initial state which is a "wavepacket", that is a superposition of various eigenstates, described by a MPS, to propagate it in real time with a time-independent Hamiltonian, finally to combine the MPS states at various times to reconstruct the excited states. The evolution of a state by a time-independent Hamiltonian with eigenbasis \( |e_i \rangle \) is given by

\[
|\psi(t)\rangle = \sum_i c_i \exp(-iE_it/\hbar)|e_i\rangle, \quad \text{where} \quad c_i = \langle e_i|\psi(0)\rangle.
\]

The Fourier transform (FT) of the autocorrelation function

\[
C(t) = \langle \psi(0)|\psi(t)\rangle = \sum_i |c_i|^2 \exp(-i\epsilon_i t/\hbar)
\]

over a time interval \( T \) is the autocorrelation spectrum:

\[
\tilde{C}_T(E) = \frac{1}{T} \int_{-T/2}^{T/2} e^{iET/\hbar} C(t) dt \approx \sum_i |c_i|^2 \sin((E - E_i)T/2\hbar), \tag{1}
\]

where \( \sin x = \sin x/x \). In the limit of long time \( T \), it yields narrow peaks at the \( E_i \)'s with weights \( |c_i|^2 \), that allows to determine the energy spectrum as shown in \[11\].

Here we show that it is actually possible to extract from the dynamics also the excited eigenstates with large overlap - those that contribute most to the dynamical wavepacket. We perform a FT directly on the many-body wavefunction \( |\psi(t)\rangle \):

\[
|\phi_T(E_i)\rangle = \frac{1}{T} \int_{-T/2}^{T/2} e^{iET/\hbar} |\psi(t)\rangle dt = \sum_i c_i \sin((E - E_i)T/2\hbar) |e_i\rangle \tag{2}
\]

For long \( T, |\phi_T(E_i)\rangle \to c_i |e_i\rangle \), providing us with the targeted eigenstate. The method selects excited states relevant for the dynamics, although myriads of other states may be present in the same spectral region. This paves the way to study the properties of individual excited many-body states.

To evaluate \( |\phi_T(E_i)\rangle \), the integral is approximated by a discrete series. To sum the series, a procedure to perform the sum of two MPSs as a MPS is required, which we now briefly sketch. Any state \( |\psi\rangle \) of a 1D system of \( M \) sites may be expressed in the MPS representation \[6, 23\]:

\[
|\psi\rangle = \sum_{a_1 \ldots a_M} \Gamma^{[1]}_{a_1} \cdots \Gamma^{[M]}_{a_M} |i_1, \ldots, i_M\rangle =: \mathcal{M}(\Gamma, \lambda)
\]

where \( \Gamma^{[i]}_{a_i} \) are matrices and \( \lambda^{[i]} \) vectors and \( |i_j\rangle \) span a local Hilbert space on site \( j \). When the \( \Gamma \)'s and the \( \lambda \)'s satisfy simple orthogonality relations detailed in \[26\], their physical interpretation is simple. For example, the spectrum of the reduced density matrix associated with bipartite splitting \( L : R = \{1, \ldots, l\} : \{l+1, \ldots, M\} \), \( \text{Tr}_R|\psi\rangle\langle\psi| \) is nothing but \( \left( \lambda^{[i]}_{a_i} \right)^2 \) for \( a_1 = 1, 2, \ldots \). The corresponding entanglement entropy given by

\[
S_l = -\sum_\alpha (\lambda^{[\alpha]}_{a_\alpha})^2 \ln(\lambda^{[\alpha]}_{a_\alpha})^2, \tag{3}
\]

is an important tool to characterize the strongly correlated systems, e.g. topological phases \[27\]. The collection of \( \lambda \) values forms the entanglement spectrum.

To describe exactly a generic state in terms of a MPS, a large number (exponentially increasing with \( M \)) of \( \alpha_i \) values is needed. However, quite often, physical states created in many experiments are only slightly entangled so that \( \lambda^{[\alpha]}_{a_\alpha} \) are rapidly decaying numbers, which allows for introduction of a rather small cutoff \( \chi \) in all sums above, resulting in tractable numerical computation \[3\]. Our approach is limited to such states only.

Following \[28\] one has that: \( \mathcal{M}(\Gamma, \lambda) + \mathcal{M}(\Gamma', \lambda') = \mathcal{M}(\Gamma \oplus \Gamma', \lambda \oplus \lambda') \). The result does not satisfy orthogonality relations that are vital for achieving efficient MPS algorithms. The algorithm to restore them \[26\] ensures that the sum is stored in a memory-efficient way. Applying the above two-step algorithm to perform elementary addition suffices to obtain \( |\psi_T(E_i)\rangle \). The technical details are presented in the Appendix.

Our goal is to obtain an eigenstate as pure as possible, not merely to detect its presence. This requires long time integration which may affect the accuracy of the TEBD algorithm \[6, 7\]. The integration time may be significantly shortened using few simple ideas which we now describe.

For simplicity let us restrict to a subspace spanned by two eigenstates \( |\psi\rangle = \alpha|e_0\rangle + \beta|e_1\rangle \) as a special case. Performing the Fourier Transform as in eq. \[2\] gives:

\[
|\psi_T(E_0)\rangle = \alpha|e_0\rangle + \beta(T)|e_1\rangle, \tag{4}
\]

with \( \beta(T) = \beta \sin(TE_1/\hbar) \Delta E = E_1 - E_0 \). Clearly \( \beta(+\infty) = 0 \), therefore large enough \( T \) guarantees convergence of \( |\psi_T(E_0)\rangle \) to \( |e_0\rangle \). Another possibility is to choose \( T \) such that \( \beta(T) = 0 \), i.e. a multiple of \( 2\pi/\Delta E \). This ensures
that \( |\psi_T(E_0)\rangle = |e_0\rangle \) despite relatively short evolution time. When other eigenstates contribute significantly to the wavepacket, there is no possibility to choose such a time \( T \) that contributions from other eigenstates vanish simultaneously. However, if for some state \( |\psi\rangle \), and its eigenstate \( |e_{i_0}\rangle \) there exists an eigenstate \( |e_{i_1}\rangle \) with nonzero overlap on \( |\psi\rangle \), such that \( |E_{i_0} - E_{i_1}| \ll |E_{i_0} - E_j| \) if \( j \neq i_0, i_1 \), then choosing \( T \) as a multiple of \( \frac{\pi}{2E_{i_1}} \) is an optimal choice.

If \( \beta \gg \alpha \) in eq. (5), a realization of \( \beta(T) \ll 1 \) is more difficult. One solution is to restart the Fourier Transform evaluation from a partially converged result. Let us first define a map \( f_{E,T}(\psi) = \psi_T(E) \). One may verify that for a two-level system, the \( n \) fold composition of \( f_{E,T} \) satisfies:

\[
f_{E,T}^n(\psi) = f(f_{E,T}^{-1}(\psi)) = \alpha |e_0\rangle + \beta \left( \sin \frac{\Delta E T}{2\hbar} \right)^n |e_1\rangle
\]

This requires performing evolution for time \( n T \) and results in a power-\( n \) decay of \( |e_1\rangle/|e_0\rangle \). This observation remains true in the full Hilbert space with almost no modification.

Another way to obtain the power-\( n \) decay of other eigenstates is to use a proper window function [24]. Instead of performing FT as in [23], one calculates

\[
|\phi^n(E)\rangle = \frac{1}{T} \int_{-T/2}^{T/2} e^{i\frac{\Delta E t}{\hbar}} \psi(t) \psi(t) dt
\]

with \( \psi(t) = 0.5 (1 - \cos \left( \frac{\pi t}{T} \right) ) \), called a Hahn window, being one possibility. This window gives a \( 1/T^2 \) convergence (instead of \( 1/T \) without a window) of \( \beta(T) \).

Further possible improvements will be apparent when discussing specific examples below.

### III. BOSE-HUBBARD MODEL

We now illustrate the method using the Bose-Hubbard (BH) model [29][30][31] as realized in a gas of ultracold atoms in an optical lattice. We are aware of limitations of the tight-binding Bose Hubbard description of the system for deep optical lattices [32][33]. The BH model provides, however, a realistic and commonly used simplification of the problem. A pioneer experiment [33] observed a quantum phase transition (QPT) from a superfluid (SF) phase to a Mott insulator (MI), and stimulated research on interacting many-body systems with external perturbations, for a review see [32]. A similar QPT was observed in a 1D realization of the BH model [33]. A \(^{87}\)Rb condensate was loaded in a deep two-dimensional "transverse" optical lattice potential realizing an array of one-dimensional atomic tubes. A shallower optical potential along the tubes coming from a pair of counter-propagating beams \( V/E_R = s \sin^2(2\pi x/\lambda) \) (in units of the recoil energy \( E_R = \hbar^2/2m_{\text{Rb}}\lambda^2 \)) was superimposed on the system. After preparing the system at different potential depth, the lattice was modulated periodically at different frequencies measuring the absorbed energy (for details see [30]). In another realization of this essentially 1D experiment, the effects created by disorder were analyzed [33]. We shall consider both these situations in the following.

#### A. Periodic lattice oscillations

Let us consider first the excitations created by a periodic modulation of the lattice. This has been addressed in [16] where the energy gain due to such a modulation was studied using a 1D Bose-Hubbard model in a parabolic trap and TEBD for the time evolution, obtaining results which were in a qualitative agreement with the experimental spectra [16]. Recently, a novel extended study appeared [17] which analyzes in detail different possible excitations in the same model using both t-DMRG and a perturbative approach. This later calculation considers smaller and simpler system so, for pedagogical reasons, we start our analysis for cases discussed in [17].

The 1D Bose-Hubbard Hamiltonian reads:

\[
H = -J \sum_j b_j^\dagger b_{j+1} + \text{h.c.} + \frac{U}{2} \sum_j n_j (n_j - 1) + \sum_j \epsilon_j n_j
\]

where \( M \) is the number of sites, \( b_j (b_j^\dagger) \) is the destruction (creation) operator of an atom at site \( j \), \( n_j = b_j^\dagger b_j \) the number operator. The tunneling amplitude \( J \) and the interaction energy \( U \) depend on the lattice depth \( s \) (see below), \( \epsilon_j = c(j - r_0)^2 \) is the energy offset of a given site, due to the harmonic trap with \( r_0 \) the position of its minimum. When the lattice depth is modulated, \( s(t) = s_0[1 + \delta \cos(\omega t)] \), the energy absorption is enhanced at frequencies leading to resonant excitation of some excited states.

The time-dependence of \( s(t) \) maps onto the corresponding time dependence of the tunneling amplitude \( J(t) \) and interaction strength \( U(t) \) using, as in [16][17], the approximate formulae due to Zwerger [38]:

\[
J(t) = \frac{4}{\sqrt{\pi}} s(t)^2 \exp(-2\sqrt{s(t)})
\]

and

\[
U(t) = 4\sqrt{2}\pi a_s \lambda^{1/4} s_1(3/2) s_1^{1/2}
\]

where recoil energy units are used and \( s_1 \) is the depth of "transverse" potential creating the 1D tubes. Following [17], we denote by \( U_0 \) the value of \( U \) in the absence of modulation \( (\delta = 0) \) and by \( J_0 \) the corresponding tunneling amplitude. Also, we take \( s_1 = 30 \), \( a_s = 5.45 \)nm, \( \lambda = 825 \)nm and the curvature of the harmonic trap \( c = 0.0123 \) as in [17].
Consider first a system of $N = 36$ particles (corresponding to Fig. 2 of [17]) in a deep lattice with $s_0 = 15$. The corresponding characteristic interaction energy is $U_0 \approx 0.714$, while the tunneling amplitude is $J_0/U_0 \approx 0.01$.

For any state in MPS representation - be it the ground state, an excited state or a wavepacket - it is easy to compute expectation values of simple operators, such as the average occupation number $\langle n \rangle$ at site $i$, as well as its standard deviation $\Delta n_i = \sqrt{\langle n_i^2 \rangle - \langle n_i \rangle^2}$. The ground state of the system is obtained by the TEBD algorithm with imaginary time evolution. The average occupation number of sites, displayed in the inset of Fig. 1, shows two Mott plateaus with single and double occupancy. By simulating a periodic driving of the lattice using a real time TEBD evolution, we find the absorption spectrum. We take the modulation to be 8%, that is 8 times bigger than in [17]: this creates bigger excitation in the system, making extraction of the excited states easier. For the same purpose, we shift slightly the lattice and the bottom of the harmonic trap taking $r_0 = 18.62345$. This breaks the reflection symmetry of the system with respect to the center of the trap, making the analysis and visualization of the excitations easier. Still, the spectrum for shorter evolution time, $t = 100$ (for $\hbar = 1$, the unit of time is the inverse of the recoil energy) resembles the perturbative theory curve in Fig. 2 of [17]. Longer integration time $t = 300$ gives a better frequency resolution and several sharp peaks emerge. The absorption for time $t = 100$ multiplied by a factor of 3 seems to reproduce quite nicely the global amplitude of the broad structure around $\omega = U_0$, showing that, for such times, absorption in this frequency region is linear. This is not true for isolated peaks at low frequency as well as for the peak around $\omega = 1.75U_0$, suggesting that these peaks correspond to isolated single levels (with the corresponding Rabi-type evolution). The integration is performed with time step $\Delta t = 0.02$ using a standard second order Trotter approach assuming maximal occupation at the sites $i_{\text{max}} = 6$ and $\chi = 50$ (those values are highly sufficient for such a deep lattice and mean occupation of sites less than four).

The absorption spectrum now serves as a guide for extracting the excited states. Suppose we are interested in excited states contributing to the first small peak around $\omega/U_0 = 0.24$. We take the wavepacket obtained at $t = 300$ with the modulation at frequency $\omega/U_0 = 0.24$ as a starting wavepacket for the analysis. We first evolve this wavepacket for $T = 10000$ with time step $\Delta t = 0.05$ using a *time independent* Hamiltonian (i.e. $\delta = 0$) and find the autocorrelation function $C_T(E)$. The latter shows a doublet around energy $E/U_0 \approx 0.24$, corresponding to two local excitations, one on the left side and one on the right side of the trap center. From this autocorrelation function, we obtain accurate eigenenergies. In order to extract the eigenstate with the chosen eigenenergy $E$, we perform a Fourier transform on MPS states during a time sufficiently long for all states outside the peak at $E/U_0 \approx 0.24$ to be significantly damped (in this specific case, $T_1 = 3000$ is convenient). This provides us with a wavepacket $|\psi_i\rangle$ with a large overlap with the desired eigenstate (typically 0.7-0.9, while the overlap of the eigenstate with the initial wavepacket was in the range 0.0001-0.01). The last step is to “purify” the wavepacket $|\psi_i\rangle$ recursively: the wavepacket $|\psi_i\rangle$ is propagated for a relatively long time ($T_i = 6000$ in our specific case). The Fourier transform of the time series at energy $E$ again which yields a new wavepacket $|\psi_{i+1}\rangle$, where the weight of all other eigenstates has been decreased (a Hahn window is used to speed up this decrease). This leads to an exponentially fast convergence of the wavepacket $|\psi_i\rangle$ towards the desired eigenstate. We observed that numerical inaccuracies may cap the exponential convergence of this procedure after 15-30 steps. In all propagating steps, we have again used time step $\Delta t = 0.05$. In general, exact computational setup is adjusted taking into account the initial overlap, details of the spectrum and numerical stability.

Altogether, one iteration takes about 4 hours, using 8 threads on 4 core Intel i7 processors (the program is multi-threaded). We have performed 6 iterations to check the accuracy; a single additional iteration suffices for a 1% accuracy on the occupation numbers.

As mentioned above, two closely spaced in energy states correspond to the absorption peak at $\omega/U_0 = 0.24$, they differ from the ground state by the hopping of one particle between two Mott domains with $n = 1$ and $n = 2$ occupations occurring either on the left or on the right side of the trap center. Had the trap be placed symmet-
They corre-

The corresponding excitation energy lies not in the broad Mott domain, but not between neighboring sites. Thus, it resembles a typical particle hole excitation in the central Mott domain as identified by [17].

In Fig. 1) corresponds to single particle-hole excitations shown in Fig. 2a. Fig. 2b shows a similar excitation where a particle hops from an occupied to an empty side (this time shown for a “right” excitation). The corresponding energy is larger since such a situation occurs further from the center, where the harmonic trap is steeper. Two other excitations shown in Fig. 2c and Fig. 2d are the “left” and “right” excitations associated to the high frequency peak at $\omega/U_0 = 1.75$. Panels (c) and (d) present two states associated to the “high” frequency peak at $\omega/U_0 = 1.75$.

The black line with circles is the ground state, while red lines with squares show selected excited states. One of the two states corresponding to the absorption peak at $\omega/U_0 = 0.24$ is shown in panel (a) (the other state is symmetric w.r.t. the trap center). Panel (b) gives the density for one of the states associated to the very small peak around $\omega/U_0 = 0.37$. Panels (c) and (d) present two states associated to the “high” frequency peak at $\omega/U_0 = 1.75$. The black line with circles is the ground state, while red lines with squares show selected excited states.

**Figure 2:** (color online) Average occupation numbers per site for some selected excited states of the system analyzed in figure 1. The black line with circles is the ground state, while red lines with squares show selected excited states. One of the two states corresponding to the absorption peak at $\omega/U_0 = 0.24$ is shown in panel (a) (the other state is symmetric w.r.t. the trap center). Panel (b) gives the density for one of the states associated to the very small peak around $\omega/U_0 = 0.37$. Panels (c) and (d) present two states associated to the “high” frequency peak at $\omega/U_0 = 1.75$.

**Figure 3:** (color online) Average occupation numbers for another set of excited states of the system analyzed in Fig. 1. The black line with circles represent the ground state while red lines with squares show selected excited states.
The absorption spectrum for a system of 65 particles on 40 sites, in an harmonic trap with curvature $c = 0.006$, centered around $r_0 = 20.62345$. The mean lattice depth is $s_0 = 12$ (corresponding to $U_0 = 0.6752$ recoil energy). It is modulated sinusoidally with 1.67% relative amplitude. The smooth black line with open circles correspond to a modulation duration $t = 50/E_R$; for longer modulation time $t = 200/E_R$, sharper peaks become resolved (red thicker line). The absorbed energy is divided by 4 in the later case, to compensate for a longer perturbation. The thick (green online) dashed line shows the absorption curve for a much stronger 20% modulation for $t = 50/E_R$, rescaled by the intensity factor $12^2$.

Figure 4: (color online) Absorption spectrum for a system of 65 particles on 40 sites, in an harmonic trap with curvature $c = 0.006$, centered around $r_0 = 20.62345$. The mean lattice depth is $s_0 = 12$ (corresponding to $U_0 = 0.6752$ recoil energy). It is modulated sinusoidally with 1.67% relative amplitude. The smooth black line with open circles correspond to a modulation duration $t = 50/E_R$; for longer modulation time $t = 200/E_R$, sharper peaks become resolved (red thicker line). The absorbed energy is divided by 4 in the later case, to compensate for a longer perturbation. The thick (green online) dashed line shows the absorption curve for a much stronger 20% modulation for $t = 50/E_R$, rescaled by the intensity factor $12^2$.

The wavepacket obtained for a given frequency, at the end of the modulation, is a starting wavepacket for our procedure. The strong 20% modulation, as used in [10], leads to a "too excited" wavepacket for the TEBD algorithm to work effectively over the long time required for a high-resolution FT (see, however, a discussion below). Of course, the excited states do not depend on the strength of the modulation (while the shape of the absorption curve in the nonperturbative regime does). For that reason, we use the wavepacket obtained after a "gentle" modulation with a smaller (1.67%) amplitude.

Let us concentrate first on the wavepacket modulated with frequency $\omega/U_0 = 2.3$. We perform a FT of the autocorrelation function over a long time and observe several partially overlapping peaks. As shown in the inset of fig. 5 we identify a doublet of states, relatively well separated from the others, significantly excited from the ground state by the modulation. Fig. 5 shows the average occupation numbers $\langle n_l \rangle$ at site $l$ for the two excited states, extracted by our method, together with the ground state. Observe that, indeed, these states correspond to a particle-hole excitation in the SF regime (with average occupation numbers between 2 and 3) confirming the suggestion of [10]. The two states are quasi-symmetric images; they have slightly different energies, because of the choice $r_0 = 20.62345$ which breaks parity w.r.t. the trap center.

We have found that, in order to obtain a fully converged excited state (e.g. the excitation appearing on the right hand side of the center of the trap represented as crosses in Fig. 5), another technique is useful. Knowing from the preliminary runs where the local (in space) excitation should be created, we prepare a new wavepacket by a local modulation, i.e. local chemical potential is modulated for a few nearby sites (around site 25 for this particular case). Explicitly, $\mu_i(t) = \mu_i(1 + \delta \exp[-(i - i_0)/w^2] \sin(\omega t))$ where $w$ is the spatial width of the modulation. Experimentally, such a modulation could be in principle created by an additional focused laser. Here it is used in the numerical...
simulation to speed up the convergence as well as to isolate excitations quasi-degenerate in energy but spatially well separated.

The data in Fig. 5 show some interesting differences when compared to the Mott regime previously discussed. While, in Figs. 2, and 21, the excited states display a transfer of particles towards the trap center - explaining why they have an excess energy (compared to the ground state) smaller than 2U0 - it is the opposite for states in Fig. 5 where particles are transferred further from the trap center, and the associated excess energy is larger than 2U0. The MPS representation of the excited states makes it easy to calculate several observables, not only the average occupation number of sites. The lower plot in Fig. 5 shows the standard deviation of the occupation of sites ∆nl, both for the ground state and the excited one. A small (resp. large) standard deviation is a signature of a Mott insulator (resp. superfluid region). The excited state has lower ∆nl in the region of excitation, proving that it is more an “excited Mott”. This is confirmed by the occupation numbers themselves which are close to integer values for the excited state, with larger fractional part for the ground state.

To explore other excited states contributing to the structure around ω/U0 = 2, we prepare the wavepacket with modulation frequency ω/U0 = 2. The first attempt to calculate the autocorrelation function by propagating such an excited wavepacket with χ = 70 (and a local Hilbert space on each site containing occupation numbers up to imax−7) failed, as e.g. the energy in the system was not conserved. We chose, however, to simply run the algorithm for a much longer time and observed an unexpected behavior depicted in Fig. 6.

Fig. 6a presents the energy of the wavepacket versus time (in recoil units, multiplication by U0 = 0.6752 give time in units of 1/U0). After the initial decrease, we observe stabilization of the energy at some value above the ground state. Fig. 6b presents the corresponding time dependence of the entropy of entanglement in the system defined as S = max Sl with Sl given by Eq. 4. It also decays and stabilizes. This behavior can be understood by performing the FT of the wavepacket autocorrelation function (shown in Fig. 6c). While the FT over the time interval [0,5000] displays broad erratic structures, the FT over the time interval [15000,20000] gives sharp peaks at some frequencies. The heuristic understanding of this behavior is the following. The initial excited wavepacket is a linear combination of eigenstates of the system. Some of them are well represented as an MPS, some other ones are represented poorly. During the time evolution the “well represented” part evolves accurately while the remaining part produces signatures of TEBD algorithm breakdown. During the course of evolution, a self-purification of the wavepacket occurs with the non-converged fraction seemingly disappearing.

The information on the non-converged fraction is actually lost on the well represented excited states. Thus, unconverged TEBD time evolution filters out information which anyway cannot be obtained, purifying the wavepacket.

A further confirmation comes from the fact that the peaks of the FT provide the energies of excited states which can now be extracted. Fig. 6b shows examples of excited states involving excitations in the superfluid regime. Contrary to the previous examples, these excited states show increased particle number fluctuations, see lower panel of Fig. 6.

States presented in Fig. 6a–Fig. 6c are other examples, well localized on the edge between the ⟨nl⟩ = 1 and ⟨nl⟩ = 2 zones. The corresponding standard deviations shown in Fig. 6b show that these excitations, nevertheless, lead to an increase of local particle number fluctuations. Fig. 6c shows a state with much larger excitation energy, almost 8U0. It shows that our method may produce states with quite high and complicated excitations. This state displays several particle transfers as well as almost a transfer of a part of the right hand side Mott ⟨nl⟩ = 2 plateau into a superfluid region.

In effect a vast majority of states identified by the FT of the autocorrelation function can be extracted. Naturally, most of these states correspond to local excitations and not too high entanglement entropy.
Figure 7: (color online) Average occupation numbers of exemplary excited states showing a significant superfluid-type excitation, when compared to the ground state (black open circles). Red squares connected by the dashed line correspond to the excited state with excess energy $\Delta E/U_0 = 2.2418$ (lower energy peak of the doublet in the inset of Fig. 5). Green crosses connected by dotted lines correspond to a state (excess energy $\Delta E/U_0 = 1.9504$) with significant excitation around the trap center. The lower panel shows the standard deviation of the occupation of sites for the ground state (open circles) and two excited states. The latter show regions of increased superfluid-like fluctuations.

Figure 8: (color online) Average occupation numbers of exemplary excited states showing localized particle-hole excitations, compared to the ground state (black open circles). Red squares connected by dashed lines correspond to the excited states. (a) Excitation similar to those discussed for the smaller system, i.e. particle-hole excitation on the transition region between two Mott plateaus (excess energy $\Delta E/U_0 = 1.8417$). (b) and (c) show two particles being transferred between sites, the excess energies are $\Delta E/U_0 = 1.9988$ and $\Delta E/U_0 = 2.0539$. (d) gives an example of higher lying excited state $\Delta E/U_0 = 7.5280$ with a partially melted, right $\langle n_i \rangle = 2$ Mott plateau.

Figure 9: (color online) Standard deviations of the occupation number, for the states presented in Fig. 8. Local excitations lead to an increase of particle number fluctuations.

B. Florence experiment

A second important example is the 1D experiment of the Florence group [37] where a second, weaker lattice potential (incommensurate with the primary lattice) is added to simulate a disorder. Its strength is denoted by $s_2$. Theoretically, the addition of a disordered potential reveals a new insulating, compressible and gapless phase, called Bose glass (BG) [31]. Experimental results confirm its existence [37, 39], but are not fully conclusive: the system is prepared from a low temperature BEC by ramping up an optical lattice, starting from a SF initial state. Preparing a MI or a BG requires, generally non-adiabatic, passage through a quantum phase transition [40, 41], thereby exciting the system.

The Florence experiment is described in [37]. We simulated the experiment as detailed in [11]: we integrate numerically the evolution of the system when $s$ increases using the TEBD algorithm [6], starting from the SF ground state at $s = 4$. At the same time, a secondary shallow lattice creating the disorder is turned on. Its effect is to modify the energy offset of the harmonic wells at different sites yielding now:

$$\epsilon_j = c(j - r_0)^2 + s_2 E_{R2} \sin^2 \left( \frac{\pi j \lambda}{\lambda_2} + \phi \right). \quad (10)$$

The curvature of the trap, $c$ depends on the harmonic trap and on the additional trapping potential created by the transverse profile of the laser beams [37].

As $s$ increases, we found [11] that the overlap of the dynamically evolved wave packet on the ground state for that $s$, rapidly decreases in the region of the SF-MI transition (around $s = 8$ for the experimental parameters) down to less than 10% for the final $s = 14$, a signature of broken adiabaticity. The situation is even worse (with the final overlap of the order of $10^{-6}$) for the disordered system. Disorder enriches the energy level structure around
the ground state, killing adiabaticity both for shallow and deep lattices.

To identify the induced excitations, we compute the various excited states building the dynamically created wavepacket, by evolving it further at the constant final value and calculating the FT in Eqs. ([12]), as explained in the previous Section. In particular we characterize the states obtained using similar observables (easily obtained in the MPS representation ([28])): the average occupation number $\langle n_l \rangle$ at site $l$ and its standard deviation $\Delta n_l$. As a by-product, we can also read off the average number of particles in the left part of the system $N_l = \sum_{i=1..l} n_i$ and its standard deviation $\Delta N_l$. This quantity is relatively easy to measure in an experiment. In our system it qualitatively resembles the entanglement entropy, as shown below ([13]).

Fig. 10 shows the autocorrelation spectrum, eq. (1), of the dynamically created wavepackets, at increasing disorder strengths. In the absence of disorder ($s_2 = 0$), about ten states are significantly excited. In Fig. 11 we show various relevant quantities for the ground state, few excited states and the wavepacket. The average occupation number $\langle n_l \rangle$ on each site $l$ has the well known “wedding cake” structure, with large MI regions with integer $\langle n_l \rangle$ separated by narrow SF regions. Because the energy excess brought by non adiabatic preparation is small, all significantly populated excited states have similar shapes. Clearly, all excitations take place in or around the SF regions: these are transfers of one atom from the edge of a Mott plateau to the edge of another Mott plateau or to the neighboring SF region ("melting" of the Mott plateau).

The standard excitations in an homogeneous system (without trapping potential) such as a particle-hole excitation in a Mott plateau are absent in the stationary states because they are dynamically unstable. They are also absent in the wavepacket because they are energetically too costly: a particle-hole excitation costs at least one interaction energy $U \approx 0.6 E_R$, much larger than the excess energy of the wavepacket $0.1 E_R$. Observe, therefore, that we are in a regime significantly different from the one discussed in the previous Section where excited states populated by modulation of the lattice depth have been considered. Here we are much "closer" in energy to the ground state.

The description used in ([13]) where the ground state is contaminated by local particle-hole excitations is not compatible with our findings. A description in terms of melting of the MI seems more relevant. The local occupation fluctuation $\Delta n_l$ confirms the existence of large MI regions with low $\Delta n_l$ separated by SF peaks with larger $\Delta n_l$. While the ground state and excited states look very similar, the wavepacket is different, with higher $\Delta n_l$ in the SF peaks. An even greater difference is revealed by the entanglement entropy. For the eigenstates, it is essentially zero in the MI regions and displays sharp peaks in the SF regions. In stark contrast, the $S_l$ of the wavepacket is also non-zero in the Mott plateaus. Sim-
Figure 12: (color online) Same as Fig. 11 but for small disorder ($s_2=0.4375$, $r_0=0$ and $\phi=0.5432$ in Eq. (10)). Brown (blue) lines correspond to excitations A and B indicated in Fig. 10(b). Mott insulating regions are clearly visible for all eigenstates, more vague for the wavepacket.

Similarly, $\Delta N_l$ is much larger in the Mott plateaus for the wavepacket than for the eigenstates. This suggests the existence of long-range entanglement between SF interfaces linking different MI regions caused by long range particle dislocations, absent in the eigenstates.

Disorder strongly modifies the properties of the system. Possible phases are analyzed in details in [46]. We consider first a small disorder $s_2=0.4375$. The breakdown of adiabaticity is stronger in presence of disorder, with twice larger excess energy - still below the particle-hole excitation energy - and more states significantly excited. The properties of various states are shown for small $s_2$ in Fig. 12. Several MI regions - identified by plateaus in $\langle n_l \rangle$ - still exist, separated by intermediate parts which are either SF or a BG. Remarkably, the ground state and the excited states do have very similar structures, with several visible Mott plateaus. In contrast, these plateaus are less visible (only the central one seems to survive, with a reduced size) for the wavepacket. This can be interpreted as a partial melting of the BG and MI phases, producing a thermal insulator [45].

For strong disorder, $s_2 = 2.1875$, no MI state exists and the ground state of the system forms a BG. The exemplary excitations are shown in Fig. 13. The occupations of various sites strongly fluctuate, low lying excitations seem quite similar to the ground state. Excitations are local in character modifying the occupation number and its variance in selected (not always close) sites only. The entanglement entropy is small, except in small SF pockets. Again, the wavepacket has large entanglement entropy and $\Delta n_l$, with numerous peaks indicating melted regions.

In conclusion, we have built an extension of the TEBD algorithm capable of extracting, from a non stationary state (created dynamically), the eigenstates dynamically populated, and the nature of the “defects” created. In the specific case of the BH model realized in a ultracold atomic gas, a quasi-adiabatic quench populates excited states that are essentially similar to the ground state. In contrast, the dynamical wavepacket has a markedly different character, with a significant entanglement across the whole sample. By modulation spectroscopy, higher excited states may also be selectively reached and analyzed.

It is clear that our method will not provide an access to all excited states of a given many body system. By construction, only those excited states which are well represented by a MPS within a given assumed size of the internal Hilbert space $\chi$ may be obtained. On the other hand, the TEBD algorithm seems to create a “self-purification” mechanism which enables to reach integration times much longer than originally expected [6, 7]. Naturally the result of this integration is not a good approximation of the true dynamics but contains information about the well represented excited states. Thus our method seems to have larger range of applicability than expected on the basis of standard limitations of the TEBD algorithm.

In that context, wavepackets with low excess energy
w.r.t. the ground state created via quasi-adiabatic quench simulating the Florence experiment with disorder [37] behave exceptionally well. Higher excitations reached in modulation spectroscopy were more computer intensive and revealed that the method cannot be used as a kind of “black-box” approach but has to be tuned to individual cases. While this is certainly some drawback, the method potential capabilities (as exemplified with the highly excited state shown in fig. [31]) seem quite encouraging.

V. ACKNOWLEDGEMENTS

We are grateful to M. Fleischhauer and his colleagues for critical remarks on an early version of this paper. Support within Polish Government scientific funds for 2009-2012 as a research project is acknowledged. ML acknowledges support from Jagiellonian University International Ph.D. Studies in Physics of Complex Systems (Agreement No. MPD/2009/6) provided by Foundation for Polish Science and cofinanced by the European Regional Development Fund. Computer simulations were performed at supercomputer Deszno, purchased thanks to the financial support of the European Regional Development Fund in the framework of the Polish Innovation Economy Operational Program, contract no. POIG.02.01.00-12-023/08 (JZ), ACK Cyfronet AGH as a part of the POIG PL-Grid project (ML) and at ICM UW under Grant No. G29-10 (JZ and ML).

VI. APPENDIX: IMPLEMENTATION OF MPS-ADDITION

The addition of two MPS, an elementary step in the FT, produces \( \lambda \) vectors with increased size. Thus it seems that the \( \chi \) value required for representing successive sums should increase fast, ruining the efficiency of the MPS representation. This is not the case: the \( \chi \)\textsuperscript{[j]} for the partial sums decrease relatively fast so that the smallest components can be safely discarded [25], preventing the cutoff to blow up, a key point for the success of the method. Altogether, several thousands states can be summed, at the price of roughly doubling the \( \chi \) value to preserve the accuracy of the MPS representation: when the series is dynamically created, as in eq. (10), the result is close to an eigenstate, i.e. simpler than the summands which are wavepackets, and the vectors added together are closely related.

To be able to compute the sum of several thousands of MPS-vectors two issues have to be resolved: restoring the canonical form of a MPS vector and performing truncation in \( \chi \). The former has been described in [26]. The truncation from \( \chi = \chi_2 \) to \( \chi = \chi_1 \) can be done in two nonequivalent ways: “blunt” and “canonical”. The canonical truncation is done in parallel with restoring the canonical form. The algorithm described in [26] processes sites sequentially restoring orthogonalities relations at each of them one at a time. The “left” and “right” Schmidt vectors for site \( k \) are:

\[
|\psi_{\alpha_k}^{1\ldots k}\rangle = \sum_{\alpha_1,\alpha_2,\ldots,\alpha_k} \Gamma_{\alpha_1\alpha_k}^{[1]} \chi_{\alpha_1}^{[1]} \ldots \Gamma_{\alpha_{k-1}\alpha_k}^{[k-1]} \chi_{\alpha_{k-1}}^{[k-1]} |i_1,\ldots,i_k\rangle \tag{11}
\]

\[
|\psi_{\alpha_k}^{k+1\ldots M}\rangle = \sum_{\alpha_{k+1},\ldots,\alpha_M} \Gamma_{\alpha_k\alpha_{k+1}}^{[k+1]} \chi_{\alpha_{k+1}}^{[k+1]} \ldots \Gamma_{\alpha_M\alpha_1}^{[M]} |i_{k+1},\ldots,i_M\rangle \tag{12}
\]

\[
|\psi\rangle = \sum_{\alpha_k} \chi_{\alpha_k}^{[k]} |\psi_{\alpha_k}^{1\ldots k}\rangle \otimes |\psi_{\alpha_k}^{k+1\ldots M}\rangle \tag{13}
\]

For a MPS vector in its canonical form, these vectors are orthonormal eigenvectors of the reduced density matrices \( \rho^{[1\ldots k]} \), \( \rho^{[k+1\ldots M]} \). The algorithm in [26] restores orthonormality of left and right Schmidt vectors at site \( k \) assuming it has already been applied for sites \( 1,\ldots,k-1 \). After the orthonormality has been restored at site \( k \), and the \( \chi_{\alpha_k}^{[k]} \) sorted by decreasing magnitude, we zero all \( \chi_{\alpha_k}^{[k]} \) for \( \chi_1 < \alpha_k \leq \chi_2 \). The truncation error is estimated by \( \sum_{\alpha_k=\chi_1+1}^{\chi_2} \chi_{\alpha_k}^{[k]} \chi_{\alpha_k}^{[k]} \). After all sites have been processed, at each site left Schmidt vectors are orthonormal eigenvectors of the appropriate reduced density matrix. Orthonormality of right Schmidt vectors at site \( k \) is lost when truncation of \( \chi_{\alpha_k}^{[k+1]} \) takes place. If the algorithm were applied again (without truncation), both left and right Schmidt vectors would be canonical for any site. We do not perform this second part since we do not use orthonormality of right Schmidt vectors for the addition procedure.

The blunt way to truncate the MPS state is to first use the previous algorithm with no parallel truncation at all, only then perform truncation of the vectors \( \chi_{\alpha_k}^{[k]} \) at all sites. This method is mathematically flawed as, starting from the second site truncation, the canonical form is lost. However, when the \( \chi \) value used for FT evaluation is larger than \( \chi \) for the temporal evolution, both methods agree well.

In that respect, wavepackets with low excess energy w.r.t. the ground state created via quasi-adiabatic quench simulating the Florence experiment with disorder [37] behaved exceptionally well. The converged results could be obtained with \( \chi = 80 \) as tested and compared with \( \chi = 120 \) or \( \chi = 150 \).

In contrast, extraction of selected eigenstates from wavepackets created by harmonic modulation is more demanding. Too strong modulation creates wavepackets containing apparently strongly entangled contributions. In such a case, a long time propagation of such a wavepacket is not possible even for large \( \chi \) values. Lowering the amplitude of the modulation and adjusting its frequency is a key ingredient for successful extraction of a given excited state. The results presented in this paper have been obtained for \( \chi = 120 \) again testing the convergence by comparison with \( \chi = 150 \).
The convergence may be also tested by monitoring the growth of the smallest \( \lambda^{(i)}_\chi \) values.