Computational Theory of a splitting BEC using a Generalized Wannier basis I: Theory and Statics

Douglas K. Faust∗
Department of Physics, University of Washington, Seattle, Washington 98195-1560, USA

William P. Reinhardt†
Department of Physics, University of Washington, Seattle, Washington 98195-1560, USA and Department of Chemistry, University of Washington, Seattle, Washington 98195-1700, USA

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Abstract

We investigate the behavior of a Bose-Einstein Condensate (BEC) under the influence of a central barrier as the particle number trends towards the thermodynamic limit. In order to perform these studies, we present a novel method which is tractable in the large-$N$ limit. This method employs what may be considered to be a generalized Wannier basis, which successfully incorporates features of previous theoretical and computational assays to the splitting problem, including mean field effects, and has access to the dimensionality, trap parameters, and particle numbers relevant to recent experiments. At any barrier height we are able to discern between a two-mode state and a state which is described sufficiently by mean field theory and, further, give a criterion and technique for matching the two-mode theory to the zero-barrier state. We compare the basis used in this model to the de-localized basis functions underlying alternate models used in recent theoretical work on the double-well splitting problem and show that only the generalized Wannier basis displays the level crossing and emergence of two complex order parameters with overall $U(1) \oplus U(1)$ symmetry as expected from a large-$N$ analogue of the Superfluid to Mott insulator transition. Using this model, we identify a universal structure, independent of $N$, in this phase transition. We also present an analytic and model-independent description of this universal structure and discuss its consequences for realizing true two-mode physics with a BEC which trends towards the thermodynamic limit.

*dfaust@phys.washington.edu
†rein@chem.washington.edu
I. INTRODUCTION

The realization of Bose Einstein condensation in a dilute gas of atoms \([1]\) and the verification that the condensate order parameter both exists, and is characterized by the existence of a well-defined global phase, as witnessed by the observations of solitons \([2]\), vortices \([3, 4]\) and laser-like interference \([5]\) has stimulated a great deal of research in recent years. Specifically, a BEC coherently split in a double-well potential holds promise as a basic tool to study symmetry breaking, decoherence and phase diffusion properties of quantum systems as well as promise for use as an interferometric tool with an efficiency below the shot-noise limit \([6]\). As such, these systems have attracted a lot of attention, with Saba et al. creating the first such BEC interferometer in 2004 \([7]\). These experiments have been refined with atom-chip technology to the point where a deterministic precession of relative phase \([8]\) attributed to differences in mean particle number in each well has been observed as well as a loss of a deterministic interference pattern attributed to phase diffusion \([9]\). At smaller particle number Oberthaler et al. obtained fine enough control over a central barrier to reach the Josephson regime in 2006 \([10]\).

One basic scientific question associated with the “splitting” of a BEC, first raised in \([7]\) is whether, or under what conditions, the two moieties generated after splitting have the same phase and how long such phase-coherence lasts. Since a BEC is a mesoscopic system it is not a priori clear whether a classical or a quantum mechanical description of the order parameter is appropriate. In the former case, two independent condensates split from a common progenitor would share a phase until interaction with the environment destroyed phase coherence similar to the way that the pieces of a cleaved crystal would retain a common orientation. In the latter case, a quantum description of the splitting process suggests, since relative number and relative phase are conjugate variables, phase coherence will be destroyed for independent condensates. Lattice experiments operating at small particle per lattice site exhibit a loss of phase coherence during the so-called Superfluid to Mott insulator transition, but show a restoration of phase-coherence faster than so-called “phase-incoherent” states once the lattice depth is decreased \([11]\). It is not clear, without a theoretical tool to accompany these experiments, what is responsible for these two phenomena, where both the Mott Insulator state and the phase-coherent state are described as “fragmented,” or pure Fock states with perfectly defined particle number. Such a theoretical tool should
also be able to investigate what barrier heights and ramping times are needed to engineer “squeezed” and other exotic states. We introduce such a theoretical method capable of accessing physical data inaccessible to \textit{in situ} imaging in order to compliment the set of BEC splitting experiments currently being performed in hopes of answering these fundamental and technical questions.

Theoretical descriptions of certain aspects of degenerate bosonic systems in one or more modes have been contributed from a variety of disciplines within physics over the last decade. The description of a BEC in terms of an order parameter with definite phase as an example of spontaneous symmetry-breaking is inherited from the condensed matter literature, as is the use of model many-body wavefunctions and Fock-space expansion coefficients to describe particle distributions in between multiple wells, aka the Bose-Hubbard model. When describing a two-mode or splitting BEC, Bose-Hubbard type models use tunneling rates and site energies, which are typically input as parameters or approximate wavefunctions\cite{12} but may actually have a complicated dependence on underlying many-body wavefunctions and trap geometry as well as having dynamics of their own during the splitting process. More realistically, in the context of a highly-restricted “quantum optics” type quasi-Gaussian ansatz, Zoller et al.\cite{13} investigated general properties of the dynamical splitting of a BEC. Perhaps most importantly, they noted that because there are two different types of dynamical variables of interest in the system: the spatial variables which govern how the density of the atomic cloud(s) evolve and the Fock space variables which govern whether particles are localize into a single well, there are two different types of adiabaticity, each associated with a characteristic timescale. We further note that since \textit{in situ} imaging of a BEC only probes the atomic density of the cloud, the distribution of the Fock space variables of the system must therefore be inferred from the density, for instance by the loss and revival of an interference pattern. The detailed analysis of one number-squeezing lattice experiment\cite{14}, however, has shown that mean-field effects may mimic the experimental signature of number-squeezing\cite{15} indicating that \textit{in situ} imaging should not be used as a stand-alone diagnostic of the Fock-space distribution of a split or squeezed BEC. However, an \textit{ab initio} computational method allows this variable, the primary component in engineering quantum degenerate states beyond mean-field theory, to be investigated directly, by construction.

Recently, more exact, “multi-configurational” schemes\cite{16,17,18,19,20}, derived from methods familiar to quantum chemists and many-body nuclear theorists have been developed.
which self-consistently include both mean-field and particle number dynamics for a set of indistinguishable bosons with access to more than one mode. In spite of all of these efforts, a single theory which synthesizes the relevant aspects of these approaches and calculates useful quantities like phase diffusion rates and the effect of barrier raising times on the subsequent interference patterns has not yet been presented.

There are two essential difficulties in correctly simulating the splitting of an initially-coherent BEC into two distinct moieties. First, is the fact that as the barrier raises, the system acquires an additional degree of freedom which was absent at $t = 0$. Namely, at some point in the splitting, a breadth of Fock space expansion coefficients corresponding to the possible distributions of particles in either well, is needed. Conversely, mean field theory, as described by the Gross-Pitaevskii equation [21][22] - which is the correct theory at low barrier heights - assumes that all particles are contained in a single quantum state. A successful theory, therefore must correctly discriminate in between a one-mode (GP like) and two-mode states in the course of a time-dependent and potentially non-adiabatic process. The method described in this paper applies the Penrose-Onsager criterion to a form of the reduced density operator following [13] in order to do this. The second difficulty is that, if one takes a condensed matter theorist’s viewpoint, the complete fragmentation of a BEC into two independent BECs should also be an example of symmetry breaking, in which a system described by a single order parameter with a global $U(1)$ symmetry, becomes a system with $U(1) \oplus U(1)$ symmetry. That is, each of the two wells should contain an independent condensate such that an arbitrary phase shift on one should not affect the total system energy. In order to motivate our choice of spatial basis functions, we show that this latter requirement is not satisfied by a naïve implementation of even the sophisticated self-consistent methods such as in [23]. In response to these difficulties, we present a basis and method which smoothly transitions from a single totally-occupied zero-temperature ground state to a fragmented state in such a way that the initial density as well as the final densities and Fock-space distributions have physical meaning.

In Section II we develop such a theory which is correct in both the low and high-barrier limits. First, we develop equations of motion for two spatial mode functions and the possible partitions of particles in between them which are tractable and separable in the limit that $N \gg 1$. We give a criterion for when the system is described by a single totally-occupied state despite using an explicitly two-mode basis and give a further means of checking/correcting
all necessary physical data of the system in that case. In Section III we briefly describe the numerical implementation of the method described in II.

In Section IV we show results from a set of calculations which verify that the basis we have employed, the generalized Wannier basis, correctly reproduces the Superfluid to Mott insulator transition (SF-MI). In order to do this, we define what it means within this theory to be in a Superfluid state and within a Mott insulator state, then we show that a curve crossing exists in which the energy of the Mott insulator state eventually falls below that of the Superfluid state as a function of barrier height. Conversely, we show that in the same physical system treated with a de-localized basis, the Superfluid state always lies energetically below what is defined as the “fragmented” state. Therefore, no such curve crossing exists within the analogous “single-” and “double-macroscopically” occupied gerade/ungerade states. Finally, in order to understand how these specific computations will generalize to other systems, we present estimates of how the band-gap of a de-localized basis scales with particle number, trap frequency and interaction strength. We show that this is strictly positive for repulsive condensates and, as such, the SF-MI transition cannot be simulated in models employing such a de-localized basis. On the other hand, an analogous calculation for the generalized Wannier basis model shows that, at high barriers, a Mott insulator phase is always the ground state for repulsive interactions. Therefore, the curve crossing, and hence the SF-MI transition, is a universal feature of this model.

Our initial physical results from this theory, described in V follow from a set of numerical and analytical investigations of the splitting process as the number of trapped particles trends towards the large-$N$ limit. We are able to delineate the regime in which a two-mode model is appropriate and this analysis indicates that there is a very narrow region, characterized by a universal mathematical structure, in which two-mode models are applicable to splitting process.

In Appendix A a method to generate an effective 1D equation which self-consistently incorporates trap and mean-field data from the transverse directions when there is only one principal (splitting) axis of interest is described.
II. THEORY

Here we develop the basic dynamical laws which we use to describe a set of \( N \) identical bosons in an external trapping potential \( V_{\text{ext}}(r) \) and give the means to interpret the state vector.

A. State Vector and Equations of Motion

Starting from the second-quantized Hamiltonian in the contact-approximation for quasi-1D (see \[\text{A}\] for a discussion of how we self-consistently include transverse trap data when only one splitting axis, \( r \), is of interest):

\[
\hat{H} = \int dr [\hat{\Psi}^\dagger(r)(\hat{T} + \hat{V}_{\text{ext}}(r))\hat{\Psi}(r) + \frac{g}{2} \hat{\Psi}^\dagger(r)\hat{\Psi}(r)\hat{\Psi}(r)\hat{\Psi}(r)],
\]

(1)

where \( g \) gives the interaction strength through the s-wave scattering length, \( a_s \), as \( g = \frac{4\pi a_s^2}{\hbar^2} \) as is appropriate for a dilute low-temperature gas and the \( \hat{\Psi} \) are the bosonic field operators satisfying the usual commutation relations \( \{\hat{\Psi}(r),\hat{\Psi}(r')\} = \{\hat{\Psi}^\dagger(r),\hat{\Psi}^\dagger(r')\} = 0 \), and \( \{\hat{\Psi}(r),\hat{\Psi}^\dagger(r')\} = \delta(|r-r'|) \). Because we are including the possibility that the state is a correlated two-mode or fragmented state, we must use two-mode field operators \( \hat{\Psi}(r) = \hat{a}_1\phi_1(r) + \hat{a}_2\phi_2(r) \).

To describe all possibilities for the occupation of these modes, the full state vector is

\[
|\Phi\rangle = \sum_{\alpha=0}^N C_\alpha (\hat{a}_1^\dagger)^\alpha (\hat{a}_2^\dagger)^{N-\alpha}|\text{vacuum}\rangle \equiv \sum_{\alpha=0}^N C_\alpha |\alpha\rangle.
\]

(2)

At this point the standard two-mode operator algebra for bosons \([12]\) gives

\[
E = \langle \Phi | \hat{H} | \Phi \rangle = \sum_{j,k=1}^2 \rho_{jk} \epsilon_{jk} + \frac{g}{2} \sum_{j,k,l,m=1}^2 \rho_{jklm} \Gamma_{jklm},
\]

(3)

with the auxiliary definitions \( \rho_{jk} \equiv \langle \Phi | \hat{a}_j^\dagger \hat{a}_k | \Phi \rangle \), \( \rho_{jklm} \equiv \langle \Phi | \hat{a}_j^\dagger \hat{a}_k^\dagger \hat{a}_l \hat{a}_m | \Phi \rangle \),

\( \epsilon_{jk} \equiv \int dr \phi_j^* \left( -\frac{\hbar^2}{2m} \nabla^2 + V_{\text{ext}} \right) \phi_k \), and \( \Gamma_{jklm} \equiv \int dr \phi_j^* (r) \phi_k^* (r) \phi_l (r) \phi_m (r) \).

We now construct the Action in order to generate equations of motion for all of the quantities declared to be dynamical variables

\[
S = \int dt \{ \langle \Phi | \hat{\Pi} | \Phi \rangle - \frac{\hbar}{i} \frac{\partial}{\partial t} - \sum_{j,k=1}^2 (\mu_{jk} \int dr \phi_j^* \dot{\phi}_k - \delta_{jk}) |\Phi\rangle \}.
\]

(4)
where the Lagrange multipliers $\mu_{jk}$ are introduced to enforce the constraint $\int dr \phi_j^* \phi_k = \delta_{jk}$

Using the above

$$S = \int dt \{ E - i \hbar \langle \Phi | \frac{\partial}{\partial t} | \Phi \rangle - \sum_{j,k=1}^2 (\mu_{jk} \int dr \phi_j^* \phi_k - \delta_{jk}) \}, \quad (5)$$

we now declare the set of Fock-space coefficients $\{ C_\alpha \}$ and the spatial functions $\{ \phi_j(r) \}$ to be dynamical variables. Employing the Dirac-Frenkel variational principle, as is done for the case of arbitrary modes \[18\], the equations of motion for the Fock-space coefficients are given by the condition

$$\frac{\partial S}{\partial C_\gamma^*} = 0 \iff \frac{\partial E}{\partial C_\gamma^*} - i \hbar \sum_{\alpha,\beta=0}^N \frac{\partial C_\alpha^*}{\partial C_\gamma^*} \langle \alpha | \beta \rangle = 0, \quad (6)$$

$$i \hbar \dot{C}_\gamma = \sum_{j,k=1}^2 \frac{\partial \rho_{jk}}{\partial C_\gamma^*} + \frac{g}{2} \sum_{j,k,l,m=1}^2 \frac{\partial \rho_{jklm}}{\partial C_\gamma^*} \Gamma_{jklm} \quad (7)$$

or, in a more compact notation

$$i \hbar \dot{C}_\gamma = \sum_{\beta=0}^N \mathcal{H}_{\gamma\beta} C_\beta \quad (8)$$

where

$$\mathcal{H}_{\gamma\beta} \equiv \langle \gamma | \left\{ \sum_{j,k=1}^2 \epsilon_{jk} (\hat{a}_j^\dagger \hat{a}_k) + \frac{g}{2} \sum_{j,k,l,m=1}^2 \Gamma_{jklm} (\hat{a}_j^\dagger \hat{a}_l^\dagger \hat{a}_k \hat{a}_m) \right\} | \beta \rangle. \quad (9)$$

Similarly, the equations of motion for the two mode functions are given by the condition

$$\frac{\partial S}{\partial \phi_q(r')} = 0. \quad (10)$$

Now, using the fact that $\frac{\partial \phi_q(r)}{\partial \phi_q(r')} = \delta_{jq} \delta(|r - r'|)$, this condition yields:

$$i \hbar \sum_{j,k=1}^2 \rho_{jk} \delta_{jq} \dot{\phi}_k = \sum_{j,k=1}^2 \rho_{jk} \delta_{jq} (\hat{T} + \hat{V}_{\text{ext}}) \phi_k + \frac{g}{2} \sum_{j,k,l,m=1}^2 \rho_{jklm} (\delta_{jq} \phi_k^* \phi_l \phi_m + \delta_{j}^* \delta_{kq} \phi_l^* \phi_m) - \sum_{j,k=1}^2 (\mu_{jk} \delta_{jq} \phi_k) \quad (10)$$

or

$$i \hbar \sum_{k=1}^2 \rho_{qk} \dot{\phi}_k = \sum_{k=1}^2 \rho_{qk} (\hat{T} + \hat{V}_{\text{ext}}) \phi_k + g \sum_{k,l,m=1}^2 \rho_{qklm} (\phi_k^* \phi_l \phi_m) - \sum_{k=1}^2 (\mu_{qk} \phi_k). \quad (11)$$

These variational calculations have been performed elsewhere in the context of identical and distinguishable bosons \[18\] up to this point. Typically, the derivation continues by multiplying either side of the above equation by $(\rho_{qk})^{-1}$ in order to decouple the two
time derivatives, however, this produces singular equations of motion when \( \rho_{qk} \) has a zero eigenvalue (i.e. when only one state is occupied). Since this is precisely the initial state we wish to consider in the case of a single well deformed into two wells, another procedure is employed. We will observe and comment on some ramifications of using schemes which become singular in the case of only one occupied mode in Section IV.

In deciding how to proceed we need to know what approximations to the state vector are relevant in the large \( N \) limit. We do not parametrize, truncate or approximate the Fock-space variables keeping the entire expansion (2) and the exact equation of motion (8), since the degree of coherence and fundamental interpretation of the system will depend vitally on both the breadth and relative phases of the distribution of the variables \( \{C_{\alpha}\} \).

In order to decouple the equations of motion for the \( \{\phi_j\} \), we note that the approximate form of the distribution of the \( \{C_{\alpha}\} \) is binomial for the case of any ground state configuration of the symmetric double-well if the mode corresponding to \( \phi_1 \) (\( \phi_2 \)) is approximately left (right) localized.

More strictly, in the case of repulsive interactions \( (g > 0) \), the following bound holds:

\[
|C_\alpha| < \frac{1}{4} \frac{N/2}{\sqrt{N!/(\alpha!)(N-\alpha)!}}.
\] (12)

Therefore, the standard deviation of the distribution of the \( \{C_\alpha\} \) should scale as \( \sim \frac{1}{\sqrt{N}} \) and the numerical prefactors on (11) which derive from the quantities \( \rho_{jk}, \rho_{jklm} \) will be well characterized by their Fock-state values when \( N \gg 1 \). Consequently, in the thermodynamic limit, the equations of motion for the lobes are given, to leading order in \( 1/N \), by the “diagonal” contributions: \( \rho_{jj}, \rho_{jkk}, \rho_{jkk} \).

Using this large-\( N \) approximation, (11) becomes, explicitly writing out the components:

\[
i\hbar \begin{pmatrix} \rho_{11} & 0 \\ 0 & \rho_{22} \end{pmatrix} \begin{pmatrix} \dot{\phi}_1 \\ \dot{\phi}_2 \end{pmatrix} = \begin{pmatrix} \rho_{11}(\hat{T} + \hat{V}_{\text{ext}}) & 0 \\ 0 & \rho_{22}(\hat{T} + \hat{V}_{\text{ext}}) \end{pmatrix} \begin{pmatrix} \phi_1 \\ \phi_2 \end{pmatrix} - \begin{pmatrix} 0 & \mu_{12} \\ \mu_{21} & 0 \end{pmatrix} \begin{pmatrix} \phi_1 \\ \phi_2 \end{pmatrix} + g \begin{pmatrix} \rho_{1111}|\phi_1|^2 + \rho_{1221}|\phi_2|^2 \\ \rho_{2121}\phi_1^*\phi_2 \end{pmatrix} \begin{pmatrix} \phi_1^* \\ \phi_2^* \end{pmatrix} \begin{pmatrix} |\phi_1|^2 + \rho_{2222}|\phi_2|^2 \\ \rho_{2121}\phi_1^*\phi_2 \end{pmatrix} \begin{pmatrix} \phi_1 \\ \phi_2 \end{pmatrix}. \] (13)

In the above, we also use the fact that, for unitary time evolution, the norm-preserving Lagrange multipliers, \( \mu_{11} \) and \( \mu_{22} \), are unnecessary.

As the art of variational science is to find a variational space that gives the desired limiting cases automatically, we show that, by using as \( \phi_j \) what may be considered to be
generalized Wannier functions satisfying the non-linear equations \([13]\), we get the correct high barrier limit of two fragmented condensates. We discuss this in detail in Section [IV] giving a comparison to other assays at the splitting problem.

Finally, we point out an external criterion with which we can ensure that our theory is consistent with correct theory in the low barrier limit (i.e. mean field theory as encompassed in the condensate order parameter and Gross-Pitaevskii equation). For a zero-temperature Bose-Einstein condensate, all physical data resides in the complex order parameter \(\Phi_{OP}(r,t)\) which satisfies the relationship \(\Phi_{OP}(r,t) = \sqrt{\rho_S(r,t)}e^{i\Theta(r,t)}\), where \(\rho_S(r,t)\) is the spatial density of bosons \([25]\). As such, even in the case when a compound or parametrized computational basis is used, the above relation can be used to check the consistency of our ansatz.

We further use the sufficiency of the complex order parameter to determine a renormalized interaction strength as a means to compensate for any errors introduced by the possibly “spurious” kinetic energy terms in the generalized Wannier basis or from using the uncoupled Fock-state equations for the \(\{\phi_k\}\). The procedure we adopt is to use a diagonal and off-diagonal interaction strength, \(g\) and \(g_{12}\) respectively, in order to describe the strength of the particle-particle interactions. To keep consistency with the high-barrier limit of two independent condensates, the \(|\phi_k|^2\phi_k\) terms use the prefactor \(g\), however, we mediate the strength of the \(|\phi_j|^2\phi_k (k \neq j)\) by a \(g_{12}\) chosen to give the correct density, and therefore order parameter, at zero barrier. While this procedure introduces an \(ad \; hoc\) modification to the equations of motion, we note that, because of the sufficiency of the order parameter, this does generate the exact ground state of the system when correctly interpreted. In practice, we find that the modulation of \(g_{12}\) is on the order of a few to tens of percent and has no significant effect on the basic scientific conclusions derived from this computational theory.

With a final rearrangement of terms and introduction of the renormalized parameter \(g_{12}\), Our final equations of motion, a coupled system of equations for the Fock and spatial dynamical variables, therefore, appear as:

\[
i\hbar \dot{C}_\gamma = \sum_{\beta=0}^{N} \mathcal{H}_{\gamma\beta}C_\beta
\]

for the \(\{C_\alpha\}\), with \(\mathcal{H}_{\gamma\beta}\) defined in \([9]\) and
Where in the above the renormalized parameter $g_{12}$ has been introduced and the identity $ho_{1221} = ho_{1212}$ has been used to allow a useful rearrangement of terms.

### B. Interpretation of State Vector

Despite explicitly using two spatial basis functions $\{\phi_1, \phi_2\}$ in this theory and a full breadth of Fock-space variables $\{C_\alpha\}$, we are interested in determining whether the system is characterized as two-mode vs. one-mode. In order to do this, we follow [13] and use the Onsager-Penrose criterion. By tracing over the spatial variables of the density operator one gets a $2 \times 2$ matrix.

$$
\rho_F = \begin{pmatrix}
\langle a_1^\dagger a_1 \rangle & \langle a_1^\dagger a_2 \rangle \\
\langle a_2^\dagger a_1 \rangle & \langle a_2^\dagger a_2 \rangle 
\end{pmatrix}
$$

This reduced density operator has two eigenvalues $\Lambda_+/-$ which sum to $N$. In the case that $\Lambda_+ = N$ and $\Lambda_- = 0$, the system could be sufficiently described by the GP equation. Should the Fock-space density operator yield eigenvalues $\Lambda_+ = \Lambda_- = N/2$, then the system is “fragmented” into two independent condensates. Finally, a correlated two-mode model, such as has been used to describe Josephson junctions [26][27] and BEC atom interferometers [28][29], is only appropriate when $N > \Lambda_+ > N/2$.

The other reduced density matrix, i.e. the density matrix traced over the Fock-space variables, returns the spatial density of the system and is given explicitly by

$$
\rho_S = |\langle a_1^\dagger a_1 \rangle|^2 + 2\Re\{\langle a_1^\dagger a_2 \rangle \phi_1^* \phi_2\} + |\langle a_2^\dagger a_2 \rangle|^2
$$

this is the quantity we use, along with the Landau’s identification, which becomes:

$$
\Phi_{OP}(r,t) = \sqrt{\rho_S(r,t)} e^{i\Theta(r,t)}
$$

in order to verify that our two-mode basis correctly reproduces all the physical data of the mean-field state when the mean-field description of the condensate is appropriate.
III. IMPLEMENTATION

In this section we describe how we solve and implement the theory described above. Because this paper, the first of a two-part series, will be concerned with comparing the stationary states of various theories, this amounts to diagonalizing the equations of motion described above for the various basis sets under consideration in Section IV. Time evolution and applications of this formalism to barrier raising and ballistic expansion will be the subject of Part II. We find the ground state of the system by employing a relaxation method to the complex time version of the equations discussed above. This is done by taking the Wick-rotated equations of motion $t \rightarrow \tau = -it$ and adding self-consistent estimates of the chemical potential and system energies from the spatial and Fock equations of motion respectively. The Wick-rotated equation of motion for either variable are:

$$-\hbar \begin{pmatrix} \rho_{11} & 0 \\ 0 & \rho_{22} \end{pmatrix} \begin{pmatrix} \frac{\partial \phi_1}{\partial \tau} \\ \frac{\partial \phi_2}{\partial \tau} \end{pmatrix} = \begin{pmatrix} \rho_{11}(\hat{T} + \hat{V}_{\text{ext}}) & 0 \\ 0 & \rho_{22}(\hat{T} + \hat{V}_{\text{ext}}) \end{pmatrix} \begin{pmatrix} \phi_1 \\ \phi_2 \end{pmatrix} - \begin{pmatrix} \mu_{11} & \mu_{12} \\ \mu_{21} & \mu_{22} \end{pmatrix} \begin{pmatrix} \phi_1 \\ \phi_2 \end{pmatrix} + \begin{pmatrix} g|\phi_1|^2 \rho_{1111} & 2g_{12} \phi_2^* \phi_1 \rho_{1212} \\ 2g_{12} \phi_1^* \phi_2 \rho_{2121} & g|\phi_2|^2 \rho_{2222} \end{pmatrix} \begin{pmatrix} \phi_1 \\ \phi_2 \end{pmatrix}$$

$$\hbar \frac{\partial C_\gamma}{\partial \tau} = E_0 - \sum_{\beta=0}^{N} \mathcal{H}_{\gamma\beta} C_\beta$$

Now, instead of the evolution characteristic of the Schrödinger equation $\phi_j(t) \propto e^{-it\mu/\hbar}$ and $C_\alpha(t) \propto e^{-itE/\hbar}$, the equations (19) (20) evolve as a form of exponential decay when the state has higher energy/chemical potential than the true ground state: $\phi_j(\tau) \propto e^{(\mu_{jj} - \mu)/\hbar}$ and $C_\alpha(\tau) \propto e^{(E_0 - E)/\hbar}$. So that, as the quantities $\mu_{jj}$ and $E_0$ iteratively converge to the ground state of the chemical potential and system energy, respectively. This is done by updating these quantities by the identifications $\mu_{jj} = i\hbar \int dx \phi_j^* \dot{\phi}_j$ and $E_0 = \sum_{\alpha,\beta} C_\alpha^* \mathcal{H}_{\alpha\beta} C_\beta$, for the provisional values of $\phi$ and $C$, renormalized after a few e-foldings of the relaxation process.

When considering the excited states of the gerade, ungerade basis discussed in Section IV, the configuration of $C_\alpha$ for each such excited state is known a priori, and in this case, for a fixed distribution of $C_\alpha$, the system (19) will unambiguously converge to the functions and chemical potential for that excited state and not the true ground state of the system.
One technical point is that since the system of equations comprised of (19), and (20) is non-linear, there is no guarantee that a stationary state produced by this method is a true ground state of the system. Stated another way, non-linearity means that there is no single energy surface for these variables since the energy surface depends on the \( \{ \phi_j \} \) themselves. In order to ensure that we have not converged to such a “non-linear local minimum,” we perform this relaxation method on several different guesses for the initial conditions and verify that they converge to the same answer.

IV. BASIS COMPARISON

A. Numerics

The typical choice in discussing the splitting problem is to select basis functions of gerade and ungerade symmetries and use them to create approximately left and right-localized functions in order to describe the high barrier state. This was first done in the context of the statics of a weakly-interacting gas in a double well potential in [12] using the ground \((\phi_g)\) and first excited state \((\phi_u)\) solutions of the Schrödinger equation and their linear combinations \(\phi_L = \frac{1}{2}(\phi_g + \phi_u)\) and \(\phi_R = \frac{1}{2}(\phi_g - \phi_u)\). This has the appeal that, in the non-interacting, static limit, at zero barrier \((\phi_L + \phi_R)^N = \phi_N\), the true many-body ground state of the system.

More recently, the dynamics of the splitting process has been investigated using the \(\{\phi_g, \phi_u\}\) basis directly with the physical identification that if both gerade and ungerade functions are macroscopically occupied, the system is “fragmented” [23, 30].

We use neither of these methods and, in this section, point out some pathologies associated with either the gerade, ungerade scheme. Instead we show that a basis which can be considered to be the non-linear generalization of Wannier basis functions generates a theory with a level crossing as the central barrier increases and show that this is a generic feature of that basis.

We show that no such level crossing exists in the \(\{\phi_g, \phi_u\}\) basis for the same system and give a criterion for when the ungerade states become important as the system trends towards the thermodynamic limit.

Considering a system of 200\(^{87}\)Rb atoms confined in a trap with frequencies \(\omega_x = 2\pi \times 44.7 \text{Hz}, \omega_y = \omega_z = \omega_\perp = 2\pi \times 1.1 \text{kHz}\) in the presence of a central Gaussian barrier
characterized by a variance of 10µm, we compute the energies of various configurations of interest. The complex time versions of the effective 1D equations of motion developed above in sections III and A are implemented in order to perform this study.

B. Analytic model of energy crossings

We present a discussion, in the context of this model, of generic features of either choice of basis and show that the Wannier basis always has a Mott Insulator ground state at high barriers while the gerade, ungerade does not. Further, it can be shown that the ungerade state becomes irrelevant in the large-\(N\) limit, as the energy gap scales as \(gN\) - an alternate proof of the perturbative result of Huang and Yang [31].

In order to show that the Mott Insulator state is a generic feature of the Wannier basis, we show that the Hamiltonian expressed in the Fock basis [23] has a minimum at \(N_1 = N/2\) and is concave. In the high-barrier limit, as in lower-right panel of Fig II when the basis functions are localized \(\int dr |\phi_1|^2|\phi_2|^2 \to 0\), the (otherwise pentadiagonal) Fock-basis Hamiltonian, becomes diagonal:

\[
\mathcal{H}_{\beta\beta} \equiv \langle \beta | \left\{ \sum_{j=1}^{2} \epsilon_{jj} (\hat{a}_j^\dagger \hat{a}_j) + \frac{g}{2} \sum_{j=1}^{2} \Gamma_{jjjj} (\hat{a}_j^\dagger \hat{a}_j^\dagger \hat{a}_j \hat{a}_j) \right\} |\beta \rangle
\]  

(21)
FIG. 2: (Non-)existence of curve crossings in de-localized and Wannier basis theories. In a) the heights of various excited states (as defined by promoting an additional particle from $g$ to $u$) above the ground state (all particles in $g$) is shown. No crossing is observed in this basis, indicating that observed population of low-barrier “excited” states in $g/u$ models may be a function of equations of motion which require initial population of the ground state. In b), we observe an energy level crossing consistent with the SF-MI transition in the Wannie r basis. In this panel, the two-mode coherent state is defined by a Binomial distribution of $|C_\alpha|^2$ and self-consistently relaxed $\phi_j$, and the “fragmented” state is a single, unit $C_{\alpha=N/2}$ as described in the text.

and the energy becomes

$$\langle \mathcal{H} \rangle = \sum_{\beta=0}^{N} C_\beta^* \mathcal{H} C_\beta = |C_\beta|^2(\epsilon_{11}\beta + \epsilon_{22}(N-\beta) + \frac{g}{2}\Gamma_{1111}(2\beta^2 - 2N\beta + N^2 - N))$$

(22)

Invoking the symmetry of the well, $\Gamma_{1111} = \Gamma_{2222}$ for instance, the energy then can be written

$$\langle \mathcal{H} \rangle = \sum_{\beta=0}^{N} C_\beta^* \mathcal{H} C_\beta = |C_\beta|^2(\epsilon_{11}N + \frac{g}{2}\Gamma_{1111}(2\beta^2 - 2N\beta + N^2 - N))$$

(23)

Which is concave and has a minimum with respect to $\beta$ at $\beta = N/2$. Finally, since $\mathcal{H}$ is diagonal when the functions $\phi_j$ don’t overlap, the ground state is a single configuration $C_{\beta} = \delta_{\beta,N/2}$, or Fock state and the system is in a Mott Insulator phase.

Conversely, there is no analogue for this proof for the de-localized functions $\{\phi_g, \phi_u\}$, or their nonlinear generalizations. We can, however, examine how relevant the excited states, as defined by the promotion of an additional particle from $\phi_g$ to $\phi_u$ are in the large $N$ limit. Looking at the energy difference between all particles in $\phi_g$ (since the quantities in 9 depend on particle number, we denote the ground state factors of $\epsilon$ and $\Gamma$ by $\circ$) and
$N - 1$ particles in $\phi_g$ one particle in the first excited state (quantities denoted by +):

$$\Delta E = \langle \mathcal{H} \rangle_+ - \langle \mathcal{H} \rangle_0 = (\epsilon_{uu}^+ + (N - 1)\epsilon_{gg}^+) + \frac{g}{2}((N - 1)(4\Gamma_{gugu}^+) + (N - 1)(N - 2)\Gamma_{gggg}^+) - (N\epsilon_{gg}^0 + \frac{g}{2} N(N - 1)\Gamma_{gggg}^0)$$

Since we are simply looking for a scaling argument we can equate the geometric quantities which only differ by the exclusion of a single particle (for instance $\Gamma_{gugu}^+ = \Gamma_{gggg}^0$) in the limit of $N \gg 1$ giving.

$$\Delta E = \epsilon_{uu} - \epsilon_{gg} + \frac{g}{2}(4(N - 1)\Gamma_{gugu} + (2 - 2N)\Gamma_{gggg})$$

Roughly, one can estimate this by noting that $\Gamma_{gugu}$ and $\Gamma_{gggg} \sim 1$ and $\epsilon_{uu} - \epsilon_{gg} \sim \hbar \omega$ giving

$$\Delta E \sim \hbar \omega + gN$$

A complimentary derivation of the energy gap for a dilute Bose gas first published in [31] and strong indication that the energy crossings and population of excited states reported in computational attempts on the splitting problem which use a de-localized basis [23][30] may only be relevant at low particle numbers and those results may not generalize to large-$N$ systems. We further point out that using these methods to compute factors of $\Gamma$ and $\epsilon$ exact spectra for arbitrary trapping potentials and particle numbers can be calculated when the de-localized basis is used.

We make two remarks here about the presence of this energy gap in models which employ a de-localized basis. First, this calculation shows that the alternate definition of “fragmentation” which has appeared in the literature, namely, that a condensate is “fragmented” when both the $g$ and $u$ states are macroscopically occupied is not equivalent to the definition of a single unit-probability partition of the particles in between wells which is native to Bose-Hubbard type models. The former definition corresponds to a state which is, generically, of more energy than the ground state of that theory (all particles in $g$) and therefore must be reached by a non-adiabatic process. Conversely, we have shown that the Mott insulator ground state is a generic feature when Wannier basis methods are used to model repulsive condensates.

Secondly, in the context of dynamical investigations of the splitting process, the non-invertability of $\rho_{jk}$ (as discussed in Sec. III) means that in order for the equations of motion such as those used in [18] and [24] to be nonsingular, the initial state must include some
finite population of an energy level above this gap which is increasingly unphysical in the
large-$N$ limit. We note that if the structure of the energy levels seen in Fig. 2a) is general
and the energy gaps maintain approximately-equal spacing as the barrier raises, this may
explain the oscillatory behavior observed in 23, for instance.

As a graphical summary of the physical interpretation we propose for our theoretical
description of the splitting process, we include figure 3. In this figure, we again find the
ground state of the theory outlined in II and III for the 200$^{87}$Rb atom system described
above. This figure shows relevant physical data for a range of barrier heights. Clearly
visible in Fig 3a) are three distinct regimes, a coherent regime in which all $N$ particles
reside in a single state and GP theory should be applicable; a two-mode state, where there
are two populated eigenvalues and it is sensible to talk about relative populations of two
mode functions; and finally a regime in which there are two “fragmented” BECs which are
described by two independent complex order parameters.

As discussed in the figure caption, the eigenvalues show how to interpret the state. When
there are $\{N, 0\}$ eigenvalues, only the final density (and phase, should one be doing dynam-
ics) has a clear physical interpretation through the Landau identification 18 and the $C_\alpha$
and $\phi_j$ are simply elements of a computational basis. On the shoulder of the best-fit $tanh$
functions, a two-mode model is appropriate and the $\phi_j$ become the mode functions. In
this case, the $C_\alpha$ also have a direct physical interpretation as $|C_\alpha|^2$ being the probability of
detecting $\alpha$ particles in $\phi_1$. Finally, when the system has two $N/2$ eigenvalues, only one $C_\alpha$
has non-zero probability and the system is correctly described as two independent condens-
sates. If this were at the end of a dynamical splitting process, one would say that the initial
condensate has been “fragmented” into two condensates.

We have shown that the basis we have selected to generate the theory of a splitting
BEC correctly displays a curve crossing in energies corresponding to the Superfluid to Mott
insulator transition and that this is a generic feature of the model. We have also shown three
distinct physical regimes of the splitting BEC and have carefully delineated what parts of
the computational basis have a clear physical interpretation in each of these regimes.
FIG. 3: A selection of the physical data which comprise our theory. In a) the system eigenvalues are shown as a function of barrier height, clearly spanning three distinct regimes. In b) the computational variables from zero-barrier state is shown, only the final density has direct physical meaning. In c) the data from the two-mode regime is shown, both the $\phi_j(r)$ and $C_\alpha$ have the interpretation of the spatial and Fock-space variables of a two-mode theory. In d) the data for a “fragmented” state is shown in which both $\phi_1$ and $\phi_2$ have the interpretation of independent complex order parameters and there is no relative particle number uncertainty.
V. ANALYTICS AND NUMERICS OF THE RELEVANCE OF A TWO-MODE MODEL

We use the formalism developed in section II to determine the ground state of a variable number of bosons in the trap discussed above. Our goal is to understand how various aspects of the splitting process scale with the total particle number \( N \) and what generic features can be observed. We observe that a type of universal behavior is associated with the splitting of a dilute Bose-condensed gas and we present a simple analytic model to understand this behavior. Finally, we discuss the ramifications of this universal feature of a splitting BEC.

A. Numerics

As in the preceding section, we use the complex-time form of the equations shown above and a relaxation algorithm in order to compute the ground state of a system of various values of \( N \) \(^{87}\)Rb atoms confined in an external potential characterized by trap frequencies \( \omega_x = 2\pi \times 44.7\ Hz, \omega_y = \omega_z = \omega_{\perp} = 2\pi \times 1.1kHz \) and a central Gaussian barrier, applied along the \( x \) direction (See A for details of how this system was modeled as an effective 1D system).

In accordance with discussion in II in order to discern in between a single-mode “GP-like” state and a system that truly requires a two-mode description, the quantity of interest is the density operator traced over the spatial degrees of freedom or the “Fock-space density operator.” (16). We plot the eigenvalues of (16) as a function of increasing barrier height for four decades of particles, \( N = \{20, 200, 2000, 20000\} \), in the trapping potential described above. The results of these calculations are shown in Fig 4.

In the context of the splitting of a BEC as a phase transition, this figure may be understood as the left-most region of each graph, characterized by eigenvalues of \( \Lambda_{+/-} = \{N, 0\} \) as being a “GP-like” state, namely, the Gross-Pitaevskii equation is sufficient to describe the zero-temperature physics of this state. At high barriers, when the state is characterized by \( \Lambda_{+/-} = N/2, N/2 \), the two generalized Wannier functions are separated enough that an arbitrary phase shift on one of them, will not affect the total system energy, satisfying the requirement that this process be a type of phase transition. The region in between these two asymptotes, or phases, is where it is appropriate to use a two-mode model to discuss
FIG. 4: Eigenvalues of the Fock-space density operator as a function of barrier height for four decades of $N$. In each case, each of the three regimes discussed in the text corresponds to a finite range of barrier heights.

Since the production of the first dilute alkali gas BEC, schemes have been proposed in which atom interferometry is performed on an initially phase-coherent two-mode BEC, see for instance [32] [33], therefore it is of fundamental interest to understand when a two-mode model is relevant. We devote the rest of this paper to understanding under what circumstances a single coherent, BEC can be brought into the two-mode regime.

First, in order to examine the splitting process in a system-independent manner we define the intensive quantity we call the entanglement of the system which is simply equal to the normalized difference in between the eigenvalues of (16): $entanglement = (\Lambda_+ - \Lambda_-)/N$.

In Figure 5 we show the entanglement plotted versus the height of the central barrier for a few decades of total particle number, showing the transition from a unit-entanglement “GP-like” state through a system correctly described by a two-mode model and finally to two independent condensates described by entanglement zero. We further roughly scale out the
FIG. 5: Scaled graphs of the entanglement which suggest a lack of non-analyticity in the splitting process as $N \to \infty$.

dependence of the barrier height for different particle numbers by plotting the entanglement versus $V_{\text{barrier}}(x = L/2)/\mu_N$, where $\mu_N$ is the chemical potential of each system, $N = \{20, 200, 2000, 20000\}$, at zero barrier.

While it is typical to expect some non-analyticity to emerge in the thermodynamic limit, Fig. 5 suggests, however, a universal feature associated with the process of transitioning from a one-mode state through a two-mode state in order to fragment a BEC. We conclude that this is, in fact, the case and present an analytic model to supplement this numeric data in the next subsection.

Since the peak barrier height, even when scaled to the chemical potential, is not independent of the system geometry, we show how the entanglement of the $N = 20,000$ particle system depends on the ratio of the self-energy to the hopping energy $\epsilon_{12}/\epsilon_{11}$. These quantities, typically denoted $J$ and $U$, are analogous to the on-site and hopping strengths of lattice Bose-Hubbard type models [34]. Figure 6 shows that, while a two-mode model is appropriate over several decades of the hopping rate before fragmentation, a single-mode GP-like state is a necessary and sufficient description until the ratio of on-site to hopping strength, $(\epsilon_{12}/\epsilon_{11})$ is well below $10^{-6}$. In addition to the entanglement, we plot the expected phase uncertainty of a two-mode model using the Heisenberg relation $\Delta N \Delta \Theta \approx 1$ [35] as a rough guide to when two-mode interferometry would fail for lack of a well-defined relative phase.
FIG. 6: Logarithmic scale graph of the entanglement as a function of (unitless) tunneling rate for \( N = 20,000 \).

**B. Analytic model of splitting regime**

In order to understand this behavior, we present an analysis of the basic mathematical quantity which dictates splitting process by assuming a simple analytical form for the distribution of the \( \{ C_\alpha \} \). Examining the Fock-space density operator \([15]\), for a symmetric geometry, the diagonal entries are known: \( \langle a_1^\dagger a_1 \rangle = \langle a_2^\dagger a_2 \rangle = N/2 \). Therefore the entanglement is determined solely by the off-diagonal entry \( \rho_{12} = \langle a_1^\dagger a_2 \rangle \) and its complex conjugate. Explicitly, this is quantity is given by

\[
\rho_{12} = \sum_{\alpha,\beta=0}^{N} \langle \alpha | C_\alpha^* (a_1^\dagger a_2) C_\beta | \beta \rangle = \sum_{\alpha=1}^{N} C_\alpha^* C_{\alpha-1} \sqrt{N - \alpha + 1} \sqrt{\alpha} \tag{26}
\]

When this quantity vanishes, the state is fragmented or two independent condensates and when it equals \( N/2 \) the system is at unit entanglement, and a description by the GP equation is appropriate. The sum in (26) is a weighted, unit-offset autocorrelation of the distribution \( \{ C_\alpha \} \) and, by assuming an physically-motivated analytic form of \( \{ C_\alpha \} \), we are able to get an insight into the process of fragmenting a BEC.

Assuming a continuous, Gaussian probability distribution of \( |C_\alpha|^2 = \frac{1}{\sqrt{2\pi}\sigma^2} e^{-\frac{(\alpha - N/2)^2}{2\sigma^2}} \), the limits discussed above are represented by means on the variance of the distribution,
FIG. 7: Graph of $\rho_{12}(2/N)$ vs. width of the Gaussian ansatz $C_\alpha$ distribution spanning four decades of $N$ showing the regimes of entanglement a) $N = 20$ $^{87}$Rb atoms, b) $N = 200$ $^{87}$Rb atoms, c) $N = 2000$ $^{87}$Rb atoms, d) $N = 20,000$ $^{87}$Rb atoms.

FIG. 8: Universal feature of $(2/N)\rho_{12} = \text{entanglement}$ as described by the Gaussian model discussed in the text.

i.e. $\sigma \approx \sqrt{N}$ for low-barrier ground states, well-described by GP theory and $\sigma \approx 1$ in the fragmented limit.

In Fig 7, we show how the value of $\langle a_1^\dagger a_2 \rangle$ depends on $\sigma$ for a few decades of total particle number $N$. In Fig 7a), the failure of $\rho_{12}$ to stay at the correct asymptotic value of $N/2$ is due to the tails of the Gaussians exceeding the physical boundaries of the integration, $\alpha \in \{0, N\}$ and is an artifact of using this approximation for small particles numbers.

Each of the graphs shows similar behavior as the system goes towards the fragmented limit. In order to see this in Fig 8, all four of the preceding graphs of $\rho_{12}$ has been scaled to unity and shown in the low-$\sigma$ limit.
Since a two-mode model is only an appropriate description of a double-well condensate when the entanglement is less than unity, this analysis indicates that a large particle number BEC, starting in ground state of a harmonic trap, must be very precisely controlled in order to be a true two-mode system. Specifically, $\sigma \approx 2$ has the physical meaning of a system which is well described by approximate Fock-states with an uncertainty in particle number of only a few particles since $2\sigma \approx \Delta N$. Condensates in splitting experiments performed to date, however, are typically on the order of $\sim 10^5$ particles, making it unlikely that such systems are in a regime where true two-mode physics is applicable for any significant amount of time.

We have identified a universal feature of the splitting process for identical bosons, based solely on the assumption of a unimodal distribution of particles around a mean value, which indicates that independent of the total particle number or trap geometry, two-mode physics is only applicable when the uncertainty, $\Delta N$ is on the order of a single particle. One of the primary implications of the $N$-independence of this feature is that reliably scaling up experiments such as the lattice SF-MI experiments to much larger than unit filling of each well would require extremely fine control over the splitting potential. Further, figure shows for an specific large-$N$ system that the tunneling rate corresponding to two-mode physics is very small.

VI. CONCLUSIONS

We have presented a tractable method to describe the splitting of a large number of indistinguishable bosons in an external trap based on using localized non-linear functions and a novel mapping criterion to the low-barrier state instead of the typical approach of using linear combinations of non-localized functions. This method captures all of the features of the splitting of a BEC as a phase transition from a one-mode theory whose dynamical law is the GP equation, through a correlated two-mode theory and finally to a fragmented condensate defined by two independent complex order parameters, giving that system a $U(1) \oplus U(1)$ symmetry. By exploring the region which connects the one-mode and fragmented limits numerically and with an analytic model which captures the relevant behavior of the splitting process in Fock space, we are able to conclude that the region in which a true correlated two-mode model is applicable corresponds to non-vanishing fluctuations around
a well-defined number state of no more than one or two particles - more than this and full
coherence is established between particles in the two wells and a single, de-localized, order
parameter $\Phi_{OP}$, whose time-evolution is dictated by the Gross-Pitaevskii equation, becomes
a sufficient description of the state.

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Appendix A: Effective parameters for 3D system

We discuss a method to self-consistently include trap and mean-field physics from the
condensate’s extent in the two transverse directions when we are interested in the spatial
profile (and ultimately dynamics) along only one principal splitting axis. This Appendix
deals with producing effective lower-dimension equations of motion for NLSE/GP like equa-
tions and multi-configurational systems of equations and is not concerned with systems in
which the transverse confinement is on the order of the healing length or scattering length,
at which point it has been shown \[37\] that a modified form of s-wave scattering takes place
and an effective $a_s$ needs to be used. One can verify that none of the systems under consid-
eration in this paper are in this limit, however, correct accounting of transverse degrees of
freedom have an important effect on the results given above.

For the sake of simplicity, we present this analysis by creating an effective 1D equation
for the $x$-direction from the full 3D GP equation, however it is exactly analogous to the
treatment of the coupled mode equations above \([15]\), which have all the same essential
mathematical features of the GP equation. In three dimensions, the GP equation may be
written:

$$
\mu \psi(x, y, z) = \left(V_{\text{ext}}(x, y, z) - \frac{\hbar^2}{2M} \nabla^2 + g(N-1)|\psi(x, y, z)|^2\right) \psi(x, y, z) \quad (A1)
$$

We start by assuming a separable solution of the GP equation $\psi(x, y, z) = \phi(x)\psi_\perp(y, z)$
with each factor independently normalized $\int dx |\phi(x)|^2 = \int dydz |\psi_\perp(y, z)|^2 = 1$, the time-
independent
In the commonly-discussed experiment where a barrier is raised along the \(x\)-direction and the transverse directions are harmonically trapped, this becomes, under our assumption of separability:

\[
E\phi(x)\psi_{\perp}(y, z) = \left( V_{\text{ext}}(x) + \frac{1}{2}\omega_y^2 y^2 + \frac{1}{2}\omega_z^2 z^2 - \frac{\hbar^2}{2M}\nabla^2 + g(N - 1)|\phi(x)|^2|\psi_{\perp}(y, z)|^2 \right) \phi(x)\psi_{\perp}(y, z) \tag{A2}
\]

Clearly, even under the assumption of a separable solution, the non-linear term does not allow the above equation to reduce to a set of two or three separable differential equations.

While full 3D stationary and time-dependent solutions of the GP equation are regularly found\(^2\)\(^3\)\(^8\), we want an effective 1D equation with an eye towards performing tractable dynamical calculations at large particle numbers. The solution we propose is to assume a form for the transverse solution \(\psi_{\perp}(y, z)\) which minimizes the energy of an appropriate energy functional given an initial solution of \(\phi(x)\), we then use a single value \(\bar{\psi}_{\perp}\), such as the peak density, in the non-linear term of the - now separable - complex-time version of the GP equation which generates solutions of \(\phi(x)\). The new solution of \(\phi(x)\) will give an updated energy functional for \(\psi_{\perp}(y, z)\) and the system of equations, eventually including the coupled mode and Fock-space equations for the two-mode theory presented in the body of this paper, can be iterated to convergence.

Taking the Gaussian ansatz for the transverse profile

\[
\psi_{\perp}(y, z) = \frac{1}{\sqrt{\pi}\sigma_y\sigma_z} e^{-\frac{y^2}{2\sigma_y^2}} e^{-\frac{z^2}{2\sigma_z^2}} \tag{A3}
\]

We left-multiply \(A2\) by \(\psi(x, y, z)\) and integrate over all space, utilizing the fact that the \(x\) and transverse factors of \(\psi\) are independently normalized in order to construct the full energy functional

\[
E = e^x + e^\perp + g(N - 1)\Gamma^x\Gamma^\perp \tag{A4}
\]

with the definitions

\[
e^x = \int dx\phi^*(x)(V_{\text{ext}}(x) - \frac{\hbar^2}{2M}\frac{\partial}{\partial x})\phi(x)
\]

\[
e^\perp = \int dx\phi(y, z)(\frac{1}{2}\omega_y^2 y^2 + \frac{1}{2}\omega_z^2 z^2 - \frac{\hbar^2}{2M}(\frac{\partial}{\partial y} + \frac{\partial}{\partial z}))\psi_{\perp}(y, z)
\]

and

\[
\Gamma^x = \int dy dz\phi^*(x)\phi(y, z)\phi(x)\phi(x),
\]

\[
\Gamma^\perp = \int dy dz\psi^*(y, z)\psi(y, z)\phi(y, z)\phi(y, z)
\]

The ansatz must minimize the energy \(E^\perp = e^\perp + g(N - 1)\Gamma^x\Gamma^\perp\) or, explicitly

\[
E^\perp = \frac{\hbar^2}{2M} \left( \frac{1}{2\sigma_y^2} + \frac{1}{2\sigma_z^2} \right) + \frac{1}{4}(\omega_y^2\sigma_y^2 + \omega_z^2\sigma_z^2) + g(N - 1)\frac{\Gamma^x}{2\pi\sigma_y\sigma_z} \tag{A5}
\]
Where $\phi(x)$ satisfies the effective 1D equation

$$
\mu \phi(x) = \left( V_{\text{ext}}(x) - \frac{\hbar^2}{2M} \frac{\partial}{\partial x} + g(N - 1)\overline{\psi}_{\perp}^2 |\phi(x)|^2 \right) \phi(x)
$$

(A6)

The two solutions can then be iterated to convergence through the parameters $\{\psi_{\perp}, \Gamma_x\}$, and provided that the dynamics in the transverse directions are negligible, yield effective 1D time-dependent equations.

Analogous to the case of the GP equation discussed above, the trap and particle data from the transverse directions can be accounted for in the two-mode theory discussed above. Summarizing the results from those calculations here $\epsilon_{jj} = \epsilon_{jj}^x + \epsilon_{jj}^\perp$, $\epsilon_{jk} = \epsilon_{jk}^x$ if $j \neq k$, $\Gamma_{jklm} = \Gamma_{jklm}^x \Gamma_{jklm}^\perp$. Further, in the nonlinear terms of the mode equations (A5), a prefactor of $\overline{\psi}_{\perp}^2$ is included in both the diagonal and off-diagonal interaction strength.

This procedure may be further generalized in a straight-forward manner to the case when dynamics in $m$ dimensions are of interest, while the condensate is constrained to $D > m$ dimensions.

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