Overlap of exact and Gross-Pitaevskii wavefunctions in Bose-Einstein condensates of dilute gases

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It has been proven theoretically for bosons with two-body repulsive interaction potentials in the dilute limit that the Gross-Pitaevskii equation provides the exact energy and density per particle as does the basic many-particle Schrödinger equation [Lieb and Seiringer, Phys. Rev. Lett. 88, 170409 (2002)]. Here, we investigate the overlap of the Gross-Pitaevskii and exact ground state wavefunctions. It is found that this overlap is always smaller than unity and may even vanish in spite of the fact that both wavefunctions provide the same energy and density per particle. Consequences are discussed.

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Since the experimental discovery of Bose-Einstein condensates (BECs) consisting of dilute atomic gases two decades ago [1–3], there has been vast interest in their properties [4–6]. In the respective theoretical studies, the Gross-Pitaevskii equation which is obtained by minimizing the Gross-Pitaevskii energy functional [7] has played a particularly leading role. The simplicity of this mean field (which we call GP) in the dilute limit.

We shall show that the LS theorem applies also for cases not covered by the available proof.

As a first step we introduce a many-body perturbation theory (MBPT) where the unperturbed Hamiltonian is the GP one. The situation is similar to the so called Möller-Plesset MBPT widely and successfully employed in electronic structure calculations which is based on the Hartree-Fock unperturbed Hamiltonian [12]. The GP Hamiltonian $h_{GP}(\mathbf{r}) = h + v$, where $h$ comprises the kinetic energy of a boson and its trap potential and $v = \Lambda |\varphi_{GP}(\mathbf{r})|^2$, possesses a complete set of eigenfunctions $\varphi_{i}(\mathbf{r})$ of which the one with the lowest eigenvalue $\mu_{GP}$ called the chemical potential is $\varphi_{GP}(\mathbf{r})$. The eigenvalue equation of the GP operator reads:

$$[h + v] \varphi_{i}(\mathbf{r}) = \mu_{i} \varphi_{i}(\mathbf{r}).$$

We note that the GP equation can also be derived from e-field methods [13], but being interested here in the wavefunction of the system at zero-temperature we rely here on the quantum mechanical origin of this equation. We may now rewrite the many-body Hamiltonian of $N$ interacting bosons $H = \sum_{j=1}^{N} h(\mathbf{r}_{j}) + \sum_{j>\ell} \lambda_{0} V(\mathbf{r}_{j} - \mathbf{r}_{\ell})$ where $V(\mathbf{r}_{j} - \mathbf{r}_{\ell})$ is the boson-boson interaction potential and $\lambda_{0}$ its strength to give:

$$H = H_{0} + \lambda_{0} W.$$  

Here, $H_{0} = \sum_{j=1}^{N} h_{GP}(\mathbf{r}_{j})$ and $\lambda_{0} W = \lambda_{0} V - v$ are now the unperturbed Hamiltonian and the residual interaction suitable for our MBPT.

The orthonormal eigenstates of $H_{0}$ can all be cast into the simple form

$$|q_{1}, q_{2}, \ldots, q_{m}\rangle = \left( a_{1}^{\dagger} q_{1} a_{2}^{\dagger} q_{2} \ldots a_{m}^{\dagger} q_{m} \right)_{\sqrt{q_{1} q_{2} \cdots q_{m}}} |0\rangle,$$  

where the $a_{i}^{\dagger}$ are the usual boson creation operators corresponding to the solutions $\varphi_{i}(\mathbf{r})$ in Eq. [1] |0⟩ is the boson vacuum, and the total number of bosons
identifying $a^+_i$ with $a^+_{GP}$, the N boson GP ground state is just \( (GP) = |N\rangle = |N\rangle^{-1/2}(a^+_{GP})^N|0\rangle \). Note that zero occupations $q_i = 0$ are not indicated in the eigenstates. It is easily seen that $H_0(|q_1, ..., q_m\rangle) = \sum q_i \mu_i |q_1, ..., q_m\rangle$, and, in particular, $H_0(|N\rangle) = N\mu_{GP}|N\rangle$.

We are now in the position to write down the relevant MBPT expansion. As can be found in text books 12 the exact eigenfunction $|\tilde{\Psi}\rangle$ in the intermediate normalization $\langle N|\tilde{\Psi}\rangle = 1$ can be expanded in orders of perturbation

$$|\tilde{\Psi}(n)\rangle = \left\{ \frac{\hat{Q}}{N\mu_{GP} - H_0}(\lambda_0 W - \Delta E) \right\}^n |N\rangle, \quad n > 0,$$

where $\hat{Q} = 1 - |N\rangle\langle N|$ is a projection operator which removes $|N\rangle$ from the terms $|\tilde{\Psi}(n)\rangle$, and $\Delta E = E_{exact} - N\mu_{GP}$ is the difference between the exact energy and that of the unperturbed Hamiltonian. This increment can also be expanded as

$$\Delta E = \sum_{n=1} E^{(n)}, \quad E^{(n)} = \langle N|\lambda_0 W|\tilde{\Psi}(n-1)\rangle.$$ 

Obviously, $E^{(0)} + E^{(1)}$, where $E^{(0)} = N\mu_{GP}$, is nothing but the total GP energy $E_{GP} = \langle N|H|N\rangle$ of the N boson system.

The normalized exact many-body ground state is, of course, given by $|\Psi_{exact}\rangle = |\tilde{\Psi}\rangle/\langle\tilde{\Psi}|\tilde{\Psi}\rangle^{1/2}$, and hence the overlap $S(N)$ between the GP and exact ground states simply takes on the form

$$S(N) = \langle GP|\Psi_{exact}\rangle = \langle \tilde{\Psi}|\tilde{\Psi}\rangle^{-1/2}$$

and because of the projector $\hat{Q}$, we see that

$$S(N) = (1 + \langle \Delta\Psi|\Delta\Psi\rangle)^{-1/2}, \quad \Delta\Psi = \sum_{n=1} |\tilde{\Psi}(n)\rangle.$$ 

Clearly, this overlap is smaller than 1.

Let us now evaluate $S(N)$ in the leading order of perturbation theory which should be valid for small values of the interaction parameter $\Lambda$. We will focus on the dilute limit $N \to \infty$ and $\Lambda$ kept fixed. To compute any term $|\tilde{\Psi}(n)\rangle$ one inserts in Eq. 1 the unity operator $1 = \sum |q_1, q_2, \ldots\rangle\langle q_1, q_2, \ldots|$. Being interested in the leading term $|\tilde{\Psi}(1)\rangle$, one immediately sees that only $|N - 1, 1\rangle$, $|N - 2, 1, 1\rangle$ and $|N - 2, 2\rangle$ unperturbed states with $i, j > 1$ contribute. Now, due to the choice $\lambda_0 W = \lambda_0 V - v$ it can be shown that the matrix elements $\langle N - 1, 1\rangle|\lambda_0 V|N\rangle$ cancel those of $-v$ in the residual interaction, and we are left only with the latter two kinds of states and their matrix elements of $V$ only. The general rules to evaluate matrix elements of operators in the basis of the Fock states 13 can be found in 14. The final result correct up to second order reads:

$$S(N) = [1 + \Lambda^2\alpha^2]^{-1/2} \quad \text{for} \quad N \to \infty.$$
appears:

$$\Psi_{\text{exact}}(Q_1, Q_2, \ldots, Q_N) = \left(\frac{\omega}{\pi}\right)^{D/4} \left(\frac{\delta_N}{\pi}\right)^{D(N-1)/4} e^{-\frac{g^2}{2}}$$

$$\times e^{-\frac{1}{2} \delta_N \sum_{k=1}^{N-1} Q_k^2},$$

where $D$ is the dimension of the problem (the $D=3$ result has been reported in [15]) and the relevant parameter $\delta_N^2 = \omega^2 + 2\lambda_0 N$ which for large $N$ becomes $\delta_N^2 = \omega^2 + 2\Lambda$. The GP wavefunction can also be expressed by the above coordinates and takes on the simple appearance [15]:

$$\Psi_{\text{GP}}(Q_1, Q_2, \ldots, Q_N) = \left(\frac{\delta_{N-1}}{\pi}\right)^{D/4} e^{-\frac{1}{2} \delta_{N-1} \sum_{k=1}^{N-1} Q_k^2}$$

We have computed explicitly the overlap of these two functions as a function of $N$, $\omega$ and $\lambda_0$. For the sake of brevity we present here the result for large $N$ with $\Lambda$ kept fixed:

$$S(N) = 2^{D/2} \left(1 + \frac{2\Lambda}{\omega^2}\right)^{D/8} \left(1 + \sqrt{1 + \frac{4\Lambda}{\omega^2}}\right)^{-D/2}$$  \hspace{1cm} (7a)$$

For small $\Lambda$ one readily obtains

$$S(N) = (1 + \Lambda^2 \omega^2)^{-1/2} + \mathcal{O}(\Lambda^3) \hspace{1cm} (7b)$$

which demonstrates how the overlap decreases as the dimension of the trap increases and also when the “size” of the trap ($\sim 1/\omega$) increases. More importantly, we are now in the position to see what happens for large $\Lambda$, where perturbation theory, of course, does not apply. From (7a) one immediately gets:

$$S(N) = 2^{3D/8} \left(\frac{\Lambda}{\omega^2}\right)^{-D/8}$$  \hspace{1cm} (7c)$$

Obviously, the overlap between the GP and the exact ground state wavefunctions approaches zero as $\Lambda$ becomes large, and, interestingly, the faster the larger is the dimension of the problem. We stress that in the HIM model the energy and density (also density matrix) per particle in the dilute limit are exactly reproduced by the GP theory [15].

We would like to also study examples with short range interactions. In the absence of exactly solvable models we have to resort to a numerical solution of the full Schrödinger equation which is not an easy task for large boson numbers. We investigate a one-dimensional (1D) double-well trap potential and contact interaction $V(x - x') = \delta(x - x')$, a problem widely studied in the literature [24, 25]. Since the current proof of the LS theorem does not cover 1D, we also extend this example to 2D by choosing $V$ to be a normalized Gaussian, see [26, 32]. Two trap potentials are studied, see upper panel of Fig. 1. The trap potential is constructed by connecting two harmonic potentials $V_\pm(x) = \frac{1}{2}(x \pm x_0)^2$ with a cubic spline such that the resulting barrier is of a given height $V_0$. For the 2D example we add the harmonic trap $\frac{1}{2}y^2$.

The mass of the particles is chosen to be 1 as in the case of the HIM investigated above. We variationally solve the Schrödinger equation by using the multi-configurational time-dependent Hartree for bosons (MCTDHB) method [36, 37], finding the ground state by imaginary time propagation. The MCTDHB is a well established method in the literature [38]. In principle, it is a numerically exact method [39], but for large boson numbers it can only be solved approximately as the number of boson Fock states fiercely explodes. If MCTDHB is used with a single variational single-particle function, say $\varphi_g$, the working equations boil down to give exactly the GP equation, i.e., $\varphi_g = \varphi_{GP}$. If, on the other hand, we use two variational single-particle functions (called orbitals), say $\varphi_g$ and $\varphi_u$, then the many-body state obtained becomes a superposition of $N + 1$ Fock states and reads

$$|\Psi\rangle = \sum_{n=0}^{N} C_n |N - n, n\rangle,$$

where the first entry $N - n$ refers to the number of bosons residing in the gerade orbital $\varphi_g$ and $n$ residing in the second, ungerade, orbital $\varphi_u$. In MCTDHB, the orbitals and the coefficients $C_n$ are determined from the time-dependent variational principle [36, 37]. Numerically, we find that as the boson number $N$ grows, the gerade orbital $\varphi_g$ smoothly approaches the GP one $\varphi_{GP}$. This
finding is very useful, as the overlap \( S(N) \) can be simply computed from the first coefficient in (5): \( S(N) = C_0 \). The ungerade orbital is, however, found to be different from the ungerade solution of the GP equation (3).

We could solve the MCTDHB with two orbitals for up to \( N = 10^7 \) bosons. The results for the overlap are shown in the lower panel of Fig. 1 for three particle numbers and are similar for 1D and 2D. It is clearly seen that the overlap drops as the interaction parameter grows from \( \Lambda = 0 \) to \( \Lambda = 1 \) (in all calculations overlap drops as the interaction parameter grows from \( \omega = 1 \)). Although \( \Lambda \) is rather moderate, the overlap can fall below 0.5. We would like to stress that the results shown seem to saturate as \( N \) is increased: the curves for \( N = 10^5, 10^6 \) and \( 10^7 \) essentially fall on top of each other. In other words, the dilute limit is essentially achieved in this example. For the evolution of the overlap from few particles to 10 million particles for one value of \( \Lambda \), see supplemental material. One also sees that changes in the trap potential are reflected in the value of the overlap. The wider trap leads to smaller overlaps.

Having the rather involved and highly correlated wavefunction (S) at our disposal, we can compute more involved quantities which reflect the boson correlations. The coefficients \( C_n \) are shown for one calculation in 1D in the upper panel of Fig. 2. Although, their distribution is qualitatively extremely different from those dictated by GP (\( C_0 = 1 \), \( C_n = 0 \) for \( n > 0 \)) the energy and density (also density matrix) per particle in our example coincide numerically very well with the respective GP results (see also the supplemental material for 1D and 2D). This is a posteriori an interesting finding: a highly complex wavefunction and a one-term wavefunction give the same results.

Having a double-well trap, we calculate also the particle number fluctuation in one well, say the left well \( L \). This can be done by introducing the creation operators \( a_L^\dagger = (a_L^\dagger_1 + a_L^\dagger_2)/\sqrt{2} \) and \( a_L^\dagger = (a_L^\dagger_1 - a_L^\dagger_2)/\sqrt{2} \) corresponding to the orbitals \( \varphi_L \) and \( \varphi_R \) which define the left and right orbitals localized in the respective wells (see, e.g., [40]). The boson number fluctuation is as usual described by

\[
(\Delta n_L)^2 = \langle (a_L^\dagger a_L)^2 | \Psi \rangle - \langle (a_L^\dagger a_L | \Psi \rangle)^2.
\]

In GP theory (\( \Psi = \Psi_{GP} \)) the resulting number fluctuation is given by: \( (\Delta n_L)^2 = N/4 \). Obviously, the number of bosons in one well is just \( N/2 \).

Using the correlated wavefunction (S), the formal result reads

\[
(\Delta n_L)^2 = \frac{N}{4} + \frac{1}{2} \sum_{n=0}^{N} C_n^2 (N - n)n
\]

\[
+ \frac{1}{2} \sum_{n=2}^{N} C_n C_{n-2} \sqrt{(N - n + 1)(N - n + 2)n(n - 1)}.
\]

Our numerical results for 1D are depicted in the lower panel of Fig. 2. Surprisingly, the boson number fluctuations decrease dramatically with increasing interaction parameter \( \Lambda \). The finding rather reminds of a Mott insulator than of a superfluid [4, 32, 40], although the system is essentially condensed. The results are the more surprising if one notices that adding even a single boson outside of the \( N - 1 \) GP bosons enhances the number fluctuation by a factor of 3. Generally, \( (\Delta n_L)^2 = (2n + 1)N/4 \) if computed with \( |\Psi\rangle = |N - n, n\rangle \), for large \( N \). Obviously, the cross terms in (10) are those which are responsible for the substantial suppression of the boson number fluctuations.

In the GP limit in which \( N \to \infty \) and the interaction parameter \( \Lambda \) is kept fixed, the total energy as well as the density per boson are exactly reproduced by the GP theory. Nevertheless, we find that the overlap of the GP and exact many-body wavefunctions is always smaller than 1, and depending on the trap and \( \Lambda \), can be rather small, even vanishingly small. This in turn implies that the exact wavefunction describes substantial boson correlations, by definition not present in GP theory. Obviously, the energy and density per boson are mean field quantities in the GP limit. The situation is very different from that in fermion systems, e.g., in electronic systems like...
atoms and molecules, where the respective mean field theory is Hartree-Fock. Since two fermions cannot occupy the same one-particle state (orbital), they build up a shell structure and their total energy does not depend only on one interaction parameter and fermion correlations are reflected in the total energy.

Although we find it very interesting that in the dilute limit GP provides the energy and density per boson correctly even if the overlap of the GP wavefunction with the exact one can be essentially zero, one clearly does not catch the rich many-particle physics present in condensed boson systems by studying GP theory or by measuring energy and density. Indeed, this overlap behavior tells us that the underlying many-particle physics is rich. Other, boson-correlation susceptible quantities should be computed and measured. One example is the boson number fluctuation discussed here, but there are many other. We refer, for instance, to the recently proposed single-shot measurements which contain much information on the system beyond mean field and predictions of the effect of correlations in the GP limit on many-body vari- 

Finally, we would like to briefly remark on excited states and dynamics. In the ground state GP theory provides the lowest energy per particle in the dilute limit. In excited states other mean field (called best mean field) functionals can provide lower energy than GP even in the dilute limit. An example can be found in where the system is not condensed but exhibits a macroscopic fragmentation. Time-dependent GP theory is also often employed to compute the dynamics of a system. Even if one starts the process with a condensed state, it is clear that excited many-particle states will mix in as time proceeds and as boson correlation is expected to be more present in excited states, deviations from GP are expected to grow in time. Thus, investigating fragmentation and boson-correlation susceptible quantities in dynamical processes may show the deviations from mean field theory more clearly for large boson numbers.

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