Non-Equilibrium Dynamics and Superfluid Ring Excitations in Binary Bose-Einstein Condensates

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We revisit a classic study [D. S. Hall et al., Phys. Rev. Lett. 81, 1539 (1998)] of interpenetrating Bose-Einstein condensates in the hyperfine states $|F = 1, m_f = -1⟩\equiv |1⟩$ and $|F = 2, m_f = +1⟩\equiv |2⟩$ of $^{87}$Rb and observe striking new non-equilibrium component separation dynamics in the form of oscillating ring-like structures. The process of component separation is not significantly damped, a finding that also contrasts sharply with earlier experimental work, allowing a clean first look at a collective excitation of a binary superfluid. We further demonstrate extraordinary quantitative agreement between theoretical and experimental results using a multi-component mean-field model with key additional features: the inclusion of atomic losses and the careful characterization of trap potentials (at the level of a fraction of a percent).

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ping potential for species $i$, each with the same number of atoms $N$, resulting in a series of post-expansion top- and side-view images depicting the evolution of each component of the condensate.

After the transfer, the system can be described theoretically by a pair of coupled nonlinear partial differential equations of the Gross-Pitaevskii (GP) type \cite{14, 15, 16, 17, 18}:

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\begin{align*}
\hbar \frac{\partial \psi_1}{\partial t} &= \left[ -\frac{\hbar^2 \nabla^2}{2m} + V_1 + g_{11} |\psi_1|^2 + g_{12} |\psi_2|^2 - i \Gamma_1 \right] \psi_1, \\
\hbar \frac{\partial \psi_2}{\partial t} &= \left[ -\frac{\hbar^2 \nabla^2}{2m} + V_2 + g_{22} |\psi_2|^2 + g_{12} |\psi_1|^2 - i \Gamma_2 \right] \psi_2,
\end{align*}
$$

where $g_{ij} = 4\pi \hbar^2 a_{ij}/m$ are the coupling constants for elastic interactions between atoms in states $|i\rangle$ and $|j\rangle$ with scattering lengths $a_{ij}$, and $V_i$ represents the trapping potential for species $|i\rangle$. The key features that distinguish the present model from earlier ones are (i) the different trap frequencies for different species, and (ii) the explicit inclusion and accurate quantitative characterization of the loss rates $\Gamma_i$. Here, $\Gamma_1 = \frac{1}{2}(\gamma_{1111}|\psi_1|^4 + \gamma_{12}|\psi_2|^2)$ and $\Gamma_2 = \frac{1}{2}(\gamma_{12}|\psi_1|^2 + \gamma_{22}|\psi_2|^2)$ represent losses for species $|1\rangle$ and $|2\rangle$, respectively. Physically, three-body recombination losses dominate in a pure $|1\rangle$ BEC (decay constant $\gamma_{111} = 5.4(11) \times 10^{-30}$ cm$^6$/s \cite{22}), but two-body inelastic collisions dominate in mixed and pure $|2\rangle$ BECs (decay constants $\gamma_{12} = 0.780(19) \times 10^{-13}$ cm$^3$/s and $\gamma_{22} = 1.194(19) \times 10^{-13}$ cm$^3$/s, values we report here for the first time, as obtained from loss data with mixed and pure condensates, respectively). Other loss processes are of limited experimental significance in this particular system and thus are not included in the GP equations.

The binary condensate begins in a highly non-equilibrium state as a result of the transfer of atoms and small differences in the inter- and intra-species scattering lengths. One would then naturally expect the system to evolve in the direction of its stationary ground state, which consists of a ball of $|2\rangle$ atoms surrounded by a shell of $|1\rangle$ atoms \cite{23}. Such evolution was observed in Ref. \cite{2}, in which the system apparently arrived at its ground state without exhibiting any oscillatory behavior. In our experiment the motion is not rapidly damped \cite{24}, in marked contrast to the earlier results \cite{23}. A typical time sequence is shown in Fig. 1 which we present alongside the results of the corresponding numerical simulations \cite{26} with and without the key model additions. The superb agreement of the improved model (Num. A) with the present experiment emphasizes one of our main results: the motional damping observed in Ref. \cite{2} is not intrinsic to the component separation process.

The shortcomings of the unimproved model (Num. B) are clearly visible in Fig. 4 as it exhibits ring structures when the experiment does not (e.g., at 60 ms) and does not exhibit rings when the experiment does (e.g., at 120 ms). We find the modified timing of these features in the improved model is due primarily to the different trap frequencies experienced by states $|1\rangle$ and $|2\rangle$. Both of the key model enhancements also contribute to more subtle changes in the coetaneous atomic density distributions.

The ring patterns in state $|1\rangle$ at $t = 30$ and 120 ms, as well as that of state $|2\rangle$ at $t = 70$ ms, constitute the most striking manifestation of the dynamical evolution and phase separation of the hyperfine state components of our system. Taking one period to be the time between the two ring-like patterns appearing at 30 ms and 120 ms, we determine an oscillation frequency $\nu \approx 11$ Hz — much lower than the radial trap frequency, $\nu_r \approx 31$ Hz. This behavior is quite different from that of single-species BECs where excitation frequencies are always greater than the trap frequency \cite{15, 22}, and further emphasizes the essentially two-component nature of the exci-
FIG. 2: (color) Top panels: 3D renderings of the density distributions in a binary BEC computed from the model at 30 and 80 ms. Each component is depicted by a contour slice at about half of its corresponding maximal density. Red and green surfaces correspond to components $|1\rangle$ and $|2\rangle$ respectively. The bottom (side) projection corresponds to the $z$-($x$-) integrated density for the $|1\rangle$ component in our model. Bottom panels: radial profiles of the density distributions at 30 and 80 ms. Symbols (squares for $|1\rangle$ and circles for $|2\rangle$) correspond to the experimental data, which are averaged over the azimuthal angle. Solid lines depict the model results. The middle curve depicts half of the total density, which is nearly preserved during the time evolution [2].

FIG. 3: Top panels: Experimental and numerical density profiles of a 50:50 mix of $|1\rangle$ and $|2\rangle$ atoms at 30 ms for different $N$ as indicated (in thousands of atoms). Different experimental scalings [21] are used to avoid saturating the images. Bottom panels: same as the preceding for different relative numbers of atoms as indicated by the $|1\rangle$:$|2\rangle$ ratio. The atom number is fixed at $N = 3.50(5) \times 10^5$, as in Fig. 1.

We used radial moments (such as $I_1 = \int r|\psi|^2 dV$) to examine the consistency of the best theoretical values of the scattering lengths [27] by comparing numerically computed and experimentally measured density profiles. Through a detailed analysis of the periodicity of $I_1(t)$, which represents the collective excitation leading to the recurrence of the ring waveform in Fig. 1, we are able to show that the value $a_{22} = 95.00a_0$ agrees well with the experimental data. By using the previously computed values $a_{11} = 100.40a_0$ and $a_{12} = 97.66a_0$ [27], we have obtained remarkable agreement between the experimental and the numerical results in all the cases considered up to 130 ms. At longer times, effects of the imperfect cylindrical symmetry of the trap become increasingly apparent in the relative position of the two condensates and the crispness of the recurring ring patterns.

Our model can be used to infer the complex spatial configurations arising from the nonlinear interactions between the two components. We show in Fig. 2 a 3D rendering of the spatial distribution of the densities for both species based upon the numerical results, which reveals interesting aspects of the interaction effects such as the location of the rings in the $z$-direction [28]. The modeled density distributions provide a richer view of the separation phenomena and are more subtle than might be suspected from an analysis of the experimental data, which consist only of integrated planar projections of the atomic density distributions.

We have also explored the dependence of the period and shape of the ring-like structures on the total initial number of atoms and the relative atom fraction in each of the two spin states. Our observations at $t = 30$ ms as we vary $N$ and the relative number are shown in Fig. 3. We find that although the precise density distribution at any particular time depends sensitively on both of these variables, the periodicity of the patterns and the quality of agreement between the experimentally obtained and the numerically computed profiles do not.

The observed ring patterns are reminiscent of radial excitations, and the further we drive the system out of equilibrium the more rings are formed. In the top panels of Fig. 3 the number of rings increases as the
mean-field (nonlinear) interaction energy of the system increases. Similarly, in the bottom panels of Fig. 3, the number of rings increases as more atoms are transferred from [1] to [2]. We understand this behavior by noting that the excess energy after transfer, as determined by differences in scattering lengths and trapping potentials, places the system far from its ground-state equilibrium and consequently induces a long oscillatory journey through the system’s configuration space. Stronger interactions, larger transfer ratios, and greater differences in the trapping potentials should therefore produce more dramatic dynamics. The experiment demonstrates that the first two conditions lead to the most pronounced ring-like structures; the last will be explored elsewhere.

In conclusion, we have observed for the first time a largely undamped two-component collective excitation in which each component forms striking ring-like patterns that recur on a timescale much longer than the radial period. Numerical solutions to a theoretical model based on a modified pair of coupled Gross-Pitaevskii equations, along with careful experimental characterization of loss rates and trapping potentials, accurately and quantitatively describe the dynamics of the system, and help to visualize aspects of the complicated component separation behavior that are difficult to observe directly. We anticipate the possibility of assuming the accuracy of the model and using experimental data to extract scattering length ratios from the periodic behavior of the density distributions, much as in Ref. [20]. This technique would be sensitive to the interspecies scattering length, of crucial importance to the question of the miscibility of two atomic species that depends on the value of the determinant $D = a_{11}a_{22} - a_{12}^2$. A further extension involves self-consistently including beyond mean-field effects, such as thermal excitations, in order to match the corresponding component separation experiments at finite temperature. These directions are subjects of future work.

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[21] For $|1\rangle$ state detection, a resonant laser pulse removes the $|2\rangle$ atoms; then a fraction (typically 12%) of the $|1\rangle$ atoms are transferred to the $|2\rangle$ state with a short two-photon pulse. The $|2\rangle$ atoms are subsequently imaged on the condensate. For $|1\rangle$ state detection this procedure is preceded by a population-inverting two-photon $\pi$-pulse. This new imaging method avoids difficulties associated with lensing and saturation.
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[24] Atomic losses become significant on a timescale $\gtrsim 100$ ms and should not affect the oscillatory behavior significantly at shorter times. Other possible damping mechanisms, such as Beliaev damping and Landau damping, may be inhibited by the low frequency of the excitation and by the low sample temperature, respectively.
[25] The origin of the highly damped behavior leading to the ground state “ball and shell” observed in Ref. [2] remains mysterious. We were unable to reproduce this damping at different trap strengths, at higher temperature, or with the introduction of additional weak trap asymmetries.
[26] We numerically integrate the coupled GP equations using a standard finite-difference in space, fourth-order in time scheme with the equations expressed in cylindrical coordinates. From the symmetry of the system the equations become effectively two-dimensional. We adaptively increase the number of mesh points to keep up with the
growing domain after releasing the condensate from the trap.

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[Publications] to view movies showing the interpenetrating time evolution between the two components over a span of 220 ms.