Stable bright solitons in two-component Bose-Einstein condensates

A.I. Yakimenko\textsuperscript{1,2}, K.O. Shchebetovska\textsuperscript{1}, S.I. Vilchinskii\textsuperscript{1}, M. Weyrauch\textsuperscript{3}

\textsuperscript{1} Department of Physics, Taras Shevchenko National University, Kiev 03022, Ukraine
\textsuperscript{2} Institute for Nuclear Research, Kiev 03680, Ukraine
\textsuperscript{3} Physikalisch-Technische Bundesanstalt, Bundesallee 100, D-38116 Braunschweig, Germany

Two-dimensional (2D) fundamental soliton-soliton pairs are investigated in binary mixtures of Bose-Einstein condensates with attractive interactions between atoms of the same type. Both attractive and repulsive interactions between atoms of different type are considered. The general properties of the stationary states are investigated variationally and numerically, and the stability regions of the soliton-soliton pairs are determined.

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I. INTRODUCTION

Binary mixtures of Bose-Einstein condensates (BECs) are interacting quantum systems of macroscopic scale which exhibit rich physics not accessible in a single-component degenerate quantum gas. They open up intriguing possibilities for a number of important physical applications, including quantum simulation \[1\], quantum interferometry \[2\], and precision measurements \[3\], \[4\]. Experimentally, multi-component BECs are generated as mixtures of atoms in different hyperfine states or by simultaneously trapping different atomic species. The key difference between multi-component and single-component BECs is the inter-component interaction. It is significant that the strength and even the sign of the interactions depend on the component interactions. General properties of the steady states of such systems are investigated by means of a variational analysis and numerical simulations. The conditions for the existence and stability of matter-wave vector solitons are revealed.

II. MODEL

We consider a binary mixture of BECs at zero temperature described in mean-field approximation by two coupled Gross-Pitaevskii equations (GPEs) \((j \in \{1, 2\})\)

\[ i\hbar \frac{\partial \Psi_j}{\partial t} = \left[ \hat{H}_j + g_{jj}|\Psi_j|^2 + g_{j,3-j}|\Psi_{3-j}|^2 \right] \Psi_j, \quad (1) \]

with \(\hat{H}_j = -\frac{\hbar^2}{2M_j} \nabla^2 + V_j(r)\), and \(M_j\) the mass of an atom of type \(j\) loaded into the axially-symmetric harmonic external trapping potential \(V_j(r) = M_j\omega_z^2 (x^2 + y^2)/2 + M_j\omega_z^2 z^2/2\). Interactions between atoms of the same type (\(j\)-\(j\) interactions) are characterized by the diagonal coupling coefficients \(g_{jj} = 4\pi\hbar^2 a_{jj}/M_j\), where \(a_{jj}\) are the s-wave scattering lengths for binary collisions between these atoms. Inter-component interactions are controlled by the off-diagonal coupling terms \(g_{12} = g_{21} = 2\pi\hbar^2 a_{12}/M_*\) with the reduced mass \(M_* = M_1 M_2/(M_1 + M_2)\). Note, that our simple mean field approach provides a reasonable approximation for the evolutionary scenarios investigated here, however, a mean-field approach of course fails at the final stage of the collapse when the atomic density increases catastrophically.

We assume that the longitudinal trapping frequency \(\omega_z\) is much larger than the transversal trapping frequency \(\omega_\perp\), and that the nonlinear interactions \(\sim g_{jj}\) are weak with respect to the confinement strength of the potential in the longitudinal direction. In this case the BEC is “disk-shaped” and we may assume that the longitudinal motion of condensates is frozen in, \(\Psi_j(r,t) = \Psi_j(x,y,t)\Gamma_j(z,t)\), where \(\Gamma_j(z,t) = (l_{zj}/\sqrt{\pi})^{-1/2} \exp(-\frac{1}{2}\omega_z t - \frac{1}{2}z^2/l_{zj}^2)\) is the ground state.

In the present work we perform a detailed theoretical analysis of nonlinear matter-wave structures in binary mixtures of atoms with attractive intra-component interactions and attractive as well as repulsive inter-component interactions. General properties of the steady states of such systems are investigated by means of a variational analysis and numerical simulations. The conditions for the existence and stability of matter-wave vector solitons are revealed.
wave function of the longitudinal harmonic trapping potential, \( l_{2j} = \sqrt{\hbar/(M_j\omega_2)} \). After integrating out the longitudinal coordinates, the GP equations take the effective 2D form
\[
\frac{i}{\hbar} \frac{\partial \Psi_j}{\partial t} + \left( \hat{h}_j + |\Psi_j|^2 + b_{j,3-j}|\Psi_{3-j}|^2 \right) \Psi_j = 0,
\]
where \( \hat{h}_1 = \Delta_1 - \mu^2 \), \( \hat{h}_2 = \kappa\Delta_2 - \frac{\kappa^2}{\hbar^2}, \kappa = M_1/M_2 \), \( r = \sqrt{x^2 + y^2} \), and \( \Delta_2 = \partial^2/\partial x^2 + \partial^2/\partial y^2 \) is the 2D Laplacian. Here we have introduced the dimensionless variables \((x, y) \rightarrow (x, y)/l_{11}, t \rightarrow t/\tau, \) and \( \Psi_j \rightarrow \Psi_j/\sqrt{C_j}, \) where \( l_{11} = \hbar/(M_1\omega_1), \tau = 2/\omega_2, \) \( C_j = \hbar\omega_2 \sqrt{\pi(l_{11}^2 + l_{22}^2)}/(2|g_{j2}|) \). Dimensionless coupling parameters are defined by \( b_{12} = -\frac{g_{21}}{|g_{22}|} \sqrt{\frac{2}{1 + \kappa}} \) and \( b_{21} = -\frac{g_{21}}{|g_{22}|} \sqrt{\frac{2}{1 + \kappa}} \). For simplicity, we consider here only the symmetric case \( M_1 = M_2 \) and \( g_{11} = g_{22}, \) so that \( \kappa = 1 \) and \( b_{12} = b_{21} = \sigma \). Furthermore, we specifically consider only cases where the diagonal part of the interaction matrix \( g_{ij} \) is attractive \((g_{11} < 0, g_{22} < 0)\) while the off-diagonal terms \( g_{12} = g_{21} \) may be attractive \((g_{12} < 0, \sigma > 0)\) or repulsive \((g_{12} > 0, \sigma < 0)\).

The coupled differential equations (2) have the following integrals of motion:

(i) the number of particles in each component
\[
N_j = \int |\tilde{\Psi}_j|^2 d^2r, \tag{3}
\]
(ii) the total energy
\[
E = E_1 + E_2 - \sigma \int |\tilde{\Psi}_1|^2|\tilde{\Psi}_2|^2 d^2r, \tag{4}
\]
with
\[
E_j = \int \left\{ |\nabla \tilde{\Psi}_j|^2 + r^2|\tilde{\Psi}_j|^2 - \frac{1}{2}|\tilde{\Psi}_j|^4 \right\} d^2r, \tag{5}
\]
(iii) momentum, and (iv) angular momentum.

III. STATIONARY SOLUTIONS

We consider now stationary vector soliton solutions of the coupled GPEs (2). We are looking for a ground state of the form
\[
\tilde{\Psi}_j(r, t) = \psi_j(r)e^{-i\mu_j t}, \tag{6}
\]
where \( r = \sqrt{x^2 + y^2} \). Each solution is characterized by chemical potentials \( \mu_j \). The real functions \( \psi_j(r) \) satisfy the coupled equations
\[
\mu_j \psi_j + \psi_j'' + \frac{1}{r} \psi_j' - r^2 \psi_j + [\psi_j^2 + \sigma \psi_{3-j}^2] \psi_j = 0 \tag{7}
\]
and the boundary conditions \( \psi_j'(0) = 0 \) and \( \psi_j(\infty) = 0 \). At fixed strength of the inter-component interaction \( \sigma \) we then obtain a two-parameter family of vector soliton solutions (with the chemical potentials \( \mu_1 \) and \( \mu_2 \) as parameters).

Obviously, vector solitons with the same chemical potential \( \mu_1 = \mu_2 = \mu \) and the same radial profiles \( \psi_j(r) \) are described by a single GPE with a harmonic potential and an effective interaction strength \( \tilde{\sigma} = 1 + \sigma \). It is known (see e.g. Ref. [21]) that 2D solitonic solutions of this equation exist for \( \mu < 2 \) if \( \tilde{\sigma} > 0 \) and for \( \mu > 2 \) if \( \tilde{\sigma} < 0 \). The value \( \tilde{\mu} = 2 \) coincides with the eigenvalue of the ground state of a linear Schrödinger equation with harmonic oscillator potential, i.e. nonlinear terms vanish and the number of atoms tends to zero when \( \tilde{\mu} \rightarrow 2 \). If \( \tilde{\sigma} < 0 \), the number \( N \) of atoms grows rapidly for \( \tilde{\mu} > 2 \); if \( \tilde{\sigma} > 0 \), \( N \) saturates at a critical value \( N_{\text{cr}}/\tilde{\sigma} \) where \( \tilde{\mu} \rightarrow -\infty \). Here, \( N_{\text{cr}} = 11.68 \) is the number of atoms in the fundamental Townes soliton, which is the localized solution of a single GPE without external trap.

Using these results for one-component solitons as a guide, one may expect qualitatively different properties for vector solitons with strong repulsion \((\sigma < 1)\) or weak repulsion \(0 < \sigma < 1\) or weak attraction \((0 < \sigma < 1)\) and strong attraction \((\sigma > 1)\).

A. Attractive inter-component interactions \((\sigma > 0)\)

The coupled differential equations (3) were solved numerically by a relaxation method similar to that described in Ref. [21]. Furthermore, we performed a variational analysis of the fundamental vector solitons. The variational results not only provide an appropriate initial condition for our numerical relaxation procedure but also they show a good agreement with our numerical results as is seen e.g. from Figs. 1(b)-(d) and (f)-(h).

If all interactions between particles are attractive, then both solitonic components are basically bell-shaped and a simple trial function of the form
\[
\psi_j(r) = \sqrt{\frac{N_j}{\pi a_j^2}} e^{-\frac{r^2}{a_j^2}} \tag{8}
\]
is expected to be a good approximation. Here \( a_j \) is the effective width of component \( j \). The trial function is normalized to the number of particles: \( \langle \psi_j|\psi_j \rangle = N_j \). Substituting the trial function (7) into Eq. (4), we obtain the energy
\[
E = E_1 + E_2 - \sigma E_{12} \tag{9}
\]
with
\[
E_j = N_j \left( \frac{1}{a_j^2} + a_j^2 - \frac{N_j}{4\pi a_j^2} \right), \quad E_{12} = \frac{N_1N_2}{\pi(a_1^2 + a_2^2)}.
\]
A soliton solution corresponds to the stationary point of the total energy at the fixed number of particles in each component: \( \frac{\partial E}{\partial a_1} N_{1,2} = 0, \frac{\partial E}{\partial a_2} N_{1,2} = 0 \).
Chemical potential \( \mu \) with respect to the line \( N_j \) as functions of the difference \( \sigma \). We present the number of particles for each component (black solid line for numerical results, black dashed-dotted line for variational results) and the chemical potential \( \mu \) in (a) and (e) correspond to the profiles given in (b)-(d) and (f)-(g), respectively.

To utilize this symmetry, \( N_j \leftrightarrow N_k \) the solutions are symmetric, \( \sigma = 1 \), so that in the plane of \( \mu_j \) and \( \mu_k \) vanishes. For decreasing total chemical potential \( \mu_j \), \( N_{j\max} \) increases towards \( N_{cr} = 11.68 \) for numerical solutions and \( N_{cr} = 4\pi \) for the variational trial functions.

The remarkably different behavior of the functions \( N_j(\mu_j) \) for weak or strong inter-component interactions is evident from a comparison of Figs. (d) and (h). In the first case \( 0 < \sigma < 1 \) the number of atoms saturates at \( N_{j\max} \leq N_{cr} \). In the second case \( \sigma > 1 \) the number of atoms decays rapidly from \( N_{j\max} \leq N_{cr} \) at one boundary to zero at the other boundary of the existence domain. It is easy to verify that for the case \( \sigma > 1 \) the sum of the number of particles in each component is always less than the critical value \( N_{cr} \). For weak inter-component interactions \( 0 < \sigma < 1 \) we may find \( N_1 + N_2 > N_{cr} \).

In summary, if inter-component as well as intra-component interactions are attractive, then both solitonic components are localized at the bottom of the potential trap, and the number of atoms in each component is subcritical. These facts change dramatically for repulsive inter-component interactions, which we discuss next.

B. Repulsive inter-component interactions \( \sigma < 0 \)

To gain insight into the properties of the fundamental vector solitons with inter-component repulsion we first present the results of a variational analysis. They provide a rather complete overview of the steady states which correspond to stationary points of the total energy at a fixed number of atoms in each component.

Minima of the total energy at a given number of atoms can be attained for two different types of the solutions. The first type we call “bell-shaped” since the density distribution of both components looks Gaussian-like. To stabilize such a “miscible” configuration for negative \( \sigma \),
the particle number in each component must increase with respect to the previously discussed case $\sigma > 0$ in order to compensate for the additional inter-component repulsion by increased intra-component attraction. Solutions of the second type will be referred to as “phase-separated” or “immiscible”. These solutions minimize the “component-overlapping” part of the total energy [the last term in Eq. (1)] by substantially changing the shape of the density distributions. One component is pushed outwards and forms a ring-like shell, while the other component is noticeably compressed - it has a higher peak density and narrower width than its non-interacting counterpart. The “immiscible” states are typical for BECs with both repulsive intra- and inter-component interactions 22,27. The notable feature of BECs with attractive intra-component but repulsive inter-component interactions is the possible coexistence of “miscible” and “immiscible” states for the same chemical potentials.

The simplest normalized trial function $\psi_j(r)$ supporting spatially separated components is given by the Ansatz

$$\psi_j(r) = A_j \left(1 + \delta_j \frac{r^2}{a_j^2}\right) e^{-\frac{r^2}{2a_j^2}}, \quad (9)$$

FIG. 2: (Color online) Variational results for $\sigma = -0.5$ [(a)-(c)] and $\sigma = -2.0$ [(d)-(f)]. Shown are the number of atoms $N_j$ (red dashed curve for solutions with $\delta = 0$, red curves with open circles for solution with $\delta > 0$) and $N_2$ (solid black curves for solutions with $\delta = 0$ and black curves with filled dots for solutions with $\delta > 0$) as the functions of the chemical potential difference $\mu_+ = \mu_1 - \mu_2$ at different values of the total chemical potential $\mu_0 = \mu_1 + \mu_2$. Green dotted line: $N = N_{at} = 4\pi$.

where $a_j$ is the effective width of component $j$ of the soliton-soliton pair, and $A_j = \sqrt{N_j/(\pi \alpha_j^2(1 + 2\delta_j(1 + \delta_j)))}$. The parameter $\delta_j \geq 0$ introduces deviations from the Gaussian-like shape of the soliton. If $\delta_j > 1/2$, the density distribution has a local minimum at the bottom of the external potential trap. Evidently only one of the two BEC components is forced outwards by repulsive inter-component interactions. Consequently, $\delta_2$ may be set to zero without loss of generality and we need to determine only three variational parameters: $\alpha_1$, $\alpha_2$ and $\delta = \delta_1$. Variational results for the number of atoms as functions of the chemical potential difference $\mu_0 = \mu_1 - \mu_2$ are shown in Fig. 2.

Let us first discuss the properties of the “bell-shaped” solutions with $\delta = 0$. These solutions are presented in Fig. 2 by the red dashed curves for $N_1$ and by the solid black line for $N_2$. As it should, solutions with the same chemical potential and same number of atoms only exist for $\mu_+ < 4$ if $-1 < \sigma < 0$ and for $\mu_+ > 4$ if $\sigma < -1$. For $-1 < \sigma < 0$, the existence domain of vector solitons is bounded in the $(\mu_1, \mu_2)$ plane like in the case $\sigma > 0$.

However, if inter-component repulsion dominates over intra-component attraction ($\sigma < -1$) “bell-shaped” solutions exist for any chemical potential. It is remarkable that even at $\mu_0 = 0$ the solitonic components differ essentially from each other for $\mu_+ < 4$ [see Fig. 2 (f)], but for $\mu_+ > 4$ there appears an additional branch of solutions that share the same profile at $\mu_0 = 0$ [see Fig. 2 (d),(e)]. The gap between these two branches gradually disappears when $\mu_+$ increases [compare Figs. 2 (e) and (d)]. For relatively weak inter-component interactions ($-1 < \sigma < 0$) and decreasing $\mu_+$ the absolute value of the derivatives $|\partial N_j/\partial \mu_+|$ grows rapidly in the vicinity of the intersection point. Furthermore, it is interesting, that in the profile for $N_j(\mu_-)$ a fold appears for $N_j(0) > 16\pi/(3 + \sigma)$ provided that $-1 < \sigma < -1/3$. In this case two additional crossing points are observed at $\mu_1 = \mu_2$ [see Fig. 2 (c)].

The properties of the second type of solutions (“phase-separated” states with $\delta > 0$), predicted by the variational method, are illustrated in Fig. 2 for $N_1(\mu_-)$ by red lines with open circles, and for $N_2(\mu_-)$ by black curves.

FIG. 3: (Color online) Variational parameter $\delta$ as a function of $\mu_- = \mu_1 - \mu_2$ for different values of the total chemical potential $\mu_0 = \mu_1 + \mu_2$. 

$\sigma = -0.5$

$\sigma = -2.0$

$\sigma = -0.5$

$\sigma = -2.0$
with filled dots. Recall that we show only solutions with \( \delta_1 = \delta \) and \( \delta_2 = 0 \) but the rest of solutions can be easily constructed using the corresponding symmetry. The parameter \( \delta \) describing the deviation of the radial profile from the Gaussian-like shape is given in Fig. 3. Here, \( \delta(\mu_-) \) is shown for the same values of \( \sigma \) and \( \mu_+ \) as in Fig. 2. As is seen from Fig. 3, the variational parameter \( \delta \) reaches zero at some value of \( \mu_- \), where the two types of solutions \( \delta > 0 \) and \( \delta = 0 \) merge. Indeed this is also observed in Fig. 2 for solutions with \( \mu_1 \approx \mu_2 \).

The above described features of phase-separated steady-states obtained using the variational approach are supported by numerical simulations of Eqs. (4). Note that only half of the phase-separated solutions are shown in Figs. (f)-(h) to avoid confusion. The traces in these figures can easily be completed using the replacements \( N_1 \to N_2 \) and \( \mu_- \to -\mu_- \). Unfortunately, our relaxation technique is not able to reproduce both types of solutions predicted by variational method for repulsive inter-component interactions since the “bell-shaped” solutions (with \( \delta_1 = \delta_2 = 0 \)) usually have higher energy than the phase-separated solutions at the same chemical potential. That is why our numerical procedure rather converges to the “phase-separated” solutions or to scalar solitons. Therefore, our analysis of the steady-states with \( \delta_1 = \delta_2 = 0 \) is limited to the variational method.

Typical radial profiles found numerically are shown in Fig. 4. Generally, the numerical results are found to be in good agreement with the variational predictions for the “phase-separated” solutions. One expects, however, differences between numerical and variational results for solutions with a very dense ring-shaped component. Indeed, a ring component with large amplitude could push away the atoms at the periphery of the core component. Consequently, an additional outer ring should be observed in the density distribution of the core component. Moreover, if the number of atoms increases even more, then more and more rings should appear in both solitonic components. Actually, an additional outer ring is seen for \( \psi_2(r) \) in Fig. 4(f), and such a modification

FIG. 4: (Color online) Examples of radial profiles \( \psi_1 \) (red dashed curves) and \( \psi_2 \) (black solid curves) for \( \sigma = -0.5 \) and \( \sigma = -2.0 \) found numerically.
of the core component explains the non-monotonic behavior of the diagrams $N_2(\mu_-)$ at negative $\mu_-$ in Figs. 5 (f)-(h).

IV. STABILITY ANALYSIS

A few general conclusions concerning stability follow from our variational analysis. In fact, it is easy to find a sufficient condition for stability against radially-symmetric collapse: if both solitonic components satisfy the condition $N_j < N_{cr}$, then the stationary solutions are expected to be stable since they correspond to the minimum of the total energy. As was shown in the previous section, for attractive inter-component interactions ($\sigma > 0$) the particle numbers in both solitonic components are always below the critical value and, therefore, the vector solitons should be stable in this case.

For $\sigma < 0$, the variational analysis predicts the existence of two types of stationary solutions: “phase-separated” states and colocated “bell-shaped” states with different stability properties. The solutions with $\delta_1 = \delta_2 = 0$ should be stable provided $N_1 < N_{cr}$ and $N_2 < N_{cr}$. At the same time, the phase-separated states are predicted to be stable against collapse even if the ring-component $j$ (with $\delta_j > 0$) is over-critical provided that the bell-shaped component (with $\delta_{3-j} = 0$) consists of an undercritical number of particles.

It is clear that a radially-symmetric variational analysis does not provide any information about the stability with respect to azimuthal perturbations. This question may be addressed by numerical simulations as discussed below. In contrast to the results of the variational analysis our numerical simulations predict the phase-separated states to be unstable against collapse even if only one component has a number of particles above the critical value $N_{cr}$. Note that numerical time evolution of the variational Ansatz for the steady-states with $\delta_1 = \delta_2 = 0$ confirmed their stability only in the region where these solutions practically merge with the second solitonic branch [see Figs. 4, 5 and 6].

The stability of the stationary solutions with respect to azimuthal symmetry-breaking perturbations was investigated by a linear stability analysis. We write the order parameter of the two-component BEC in the form $\Psi_j(r, t) = \varphi_j(r) e^{\mu_j t}$ with a small perturbation $\varphi_j(r, t) = u_j(r) e^{i\omega t + iL\varphi} + v_j^*(r) e^{-i\omega t - iL\varphi}$ and insert it into Eq. 6. We linearized the resulting equations with respect to $\varphi$ and solved the resulting eigenvalue problem for $\omega$. Solutions exhibiting an imaginary part would indicate instability.

We find that $\gamma_L = \text{Im}(\omega)$ are identically zero for all $L$ if the inter-component interaction is attractive. If it is repulsive the azimuthal modes $L = 1$ and $L = 2$ may be unstable. A typical example for a maximum growth rate $\gamma_L$ as a function of the chemical potential $\mu_1$ at fixed $\mu_2$ is shown in Fig. 7. Note that above some threshold the growth rates vanish. It is important to realize that the mode $L = 1$ has a much wider instability region than $L = 2$, and that the growth rates of the mode $L = 2$ are smaller than those of the mode $L = 1$. Therefore, the $L = 1$ modes are most dangerously influencing the stability of repulsively interacting solitons.

The predictions of the variational analysis and the results of the linear stability analysis have been tested by numerical experiments of the time evolution of the perturbed vector solitons. For the numerical solution of the time-dependent GPEs a standard split-step Fourier transform method (see, e.g., Ref. [28]) has been used. While vector solitons with attractive inter-component interactions are confirmed to be stable over the full existence domain, repulsive inter-component interactions lead to various instabilities. Typical examples for unstable dynamics are given in Fig. 7.

We found that, if at least one solitonic component contains an overcritical number of atoms, then the stronger component finally collapses. However, we do not see a collapse, if both solitonic components are under-critical. Obviously, the potential trap prevents the unlimited expansion of the BEC. Furthermore, the final collapse is not possible, if the number of atoms in both components is subcritical, so as the result of symmetry-breaking instability we see a contraction and relative motion of the two solitonic components. Oscillations between such a quasi-collapse and an asymmetric state is shown in Fig. 7 (a) for times $t = 3.5$ and $t = 4.0$. Here, both solitonic components are under-critical, and the growth rate for the $L = 1$ mode is non-vanishing.

The growth of the $L = 2$ mode leads to the decay of the shell component of the soliton into two filaments. At the beginning, two humps and two holes appear in the ring, and the inner component rapidly leaks through these hollows into a ring-shaped “well”. Repulsive inter-
component interactions between the ring-shaped component and the leaking inner component leads to an increase of the hollows and finally the ring breaks up into two separate parts as seen in Fig. 7(b). Note that the dynamics of the unstable trapped two-component BEC is similar to the evolution of vector solitons reported in Ref. [19] within a similar model but without external trapping potential. The most outstanding feature of trapped matter-wave vector solitons is the fact that they can be completely stabilized even for strong repulsive inter-component interactions. We show the stability region in green color in the \((\mu_1, \mu_2)\) plane for weak \((\sigma = -0.5)\) and strong \((\sigma = -2.0)\) inter-component repulsive interactions in Figs. (a) and (e).

V. SUMMARY AND CONCLUSIONS

Fundamental 2D soliton-soliton pairs are investigated in two-component BECs with attractive intra-component interactions. General properties of vector solitons and their stability are studied variationally and numerically for both attractive and repulsive inter-component interactions. We found different types of soliton-soliton pairs including phase-separated pairs where one component is pushed outwards and forms a ring-like shell and the other component is compressed due to repulsive inter-component interactions. It turns out that for some values of the chemical potentials phase-separated steady-states coexist with collocated states characterized by bell-shaped density distributions in both components.

We performed a linear stability analysis of small azimuthal perturbations and checked these results by an extensive series of numerical simulations. For attractive inter-component interactions matter-wave bright vector solitons are demonstrated to be stable throughout the existence domain. For BEC components which repel each other various unstable evolution scenarios including collapse and azimuthal symmetry-breaking instabilities are observed. The instabilities, as a rule, lead either to separation of the condensed phases and then a collapse of the stronger over-critical \((N > N_{cr)}\) component or a periodic relative motion of the under-critical \((N < N_{cr})\) solitonic components backwards and forwards near the bottom of trapping potential. Nevertheless, there are conditions where complete stabilization of vector solitons is observed even in the case of repulsive inter-component interactions.

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