Nonlocalized motion in two-dimensional container of $\alpha$ particles in $3^-$ and $4^-$ states of $^{12}$C

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The first $3^-$ and $4^-$ states of $^{12}$C are studied in the present container model, in which the shift parameter is introduced to break the parity symmetry for projecting out the negative-parity states. Taking the limit as the shift parameter approaches zero and by variational calculations for one-deformed size parameter, the local energy minima are obtained for the $3^-$ and $4^-$ states. It is found that the obtained single THSR (Tohsaki-Horiuchi-Schuck-Röpke) wave functions for $3^-$ and $4^-$ states are 96% and 92% equivalent to the corresponding GCM wave functions, respectively. The calculated intrinsic densities further show that these negative-parity states of three clusters, different with the traditional understanding of rigid triangle structure, are found to have nonlocalized clustering structure in the two-dimensional container picture.
The $^{12}$C is a specially interesting atomic nucleus due to its rich cluster structure which has been investigated for half a century by various models [1–9]. In particular, describing and understanding the essential correlations of $3\alpha$-cluster states is a fundamental problem in this nucleus. Recently, using the THSR wave function [10–11], the Hoyle state was proposed to have a three $\alpha$ condensate characters where the $3\alpha$ clusters have $70\%–80\%$ possibility of making relative $(0S)$ motion [12] in $^{12}$C.

Quite recently, inspired by the original THSR wave function, we proposed a container model [13–16] for the description of general cluster states in nuclear systems. Different from the traditional understanding, the clusters are making a nonlocalized motion in the container which is only confined by the size parameter instead of the inter-cluster separation-coordinates. One unique feature of this new model is that we can obtain with high-precision a single optimal wave functions for various cluster states. It enables us to discuss the correlations of clusters in a straightforward way based on the explicit relative wave functions. This was highlighted by the successful descriptions of many nuclear systems [16–19].

However, up to now, the important negative-parity cluster states in $^{12}$C still have not been studied in the container model due to the fact that the THSR wave function in its original form described only positive-parity states. Traditionally, in analogy with the usual belief of the typical rigid $\alpha+^{16}$O structure in $^{20}$Ne system [20], the structure of the $^{12}$C has also long thought to be associated with a definite triangular shape. One typical state is the $3^-$ state at 9.64 MeV, which could have threefold symmetry and three alpha particles are in an equilateral triangular arrangement. Recently, Bijker et al. [21–22] proposed a description of cluster states in nuclei in terms of representations of unitary algebras. The $^{12}$C was assumed to have $D_{3h}$, $3\alpha$ cluster structure. Based on the algebraic cluster model, the rotational band of an oblate equilateral triangular with the cluster states $0^+$, $2^+$, $3^-$, $4^\pm$, $5^-$ was predicted. The negative-parity states was considered as the important support for their rigid $3\alpha$ cluster picture in $^{12}$C. However, this model became also criticized of not respecting the Pauli principle [23] or the positive and negative states can be interpreted as two separate bands [24]. Quite recently, a nuclear fluorescence experiment through patterns of polarized $\gamma$ rays was proposed [25] to discriminate the geometric configurations for $\alpha$-cluster nuclei like $^{12}$C in a model-independent way. On the other hand, the THSR wave function has taught us that the positive-parity states, especially the Hoyle state, should not be seen as being in rigid geometrical $3\alpha$ arrangements but rather the $\alpha$’s are in a low-density gas state akin to a Bose condensate. Thus, it is highly desirable to study the negative-parity states in the container model and clarify the corresponding structure of $3\alpha$ clustering in a microscopic way using a generalized THSR wave function.

To describe the negative-parity states, we here introduce shift parameters to generate negative-parity states in the THSR wave function. This has successfully been used already for the negative-parity states of the two-cluster structure in $^{20}$Ne [13]. The simple THSR wave function for three $\alpha$’s with one size parameter $\beta$ can be constructed as,

$$\Phi(\beta, S_1, S_2) = \int d^3R_1 d^3R_2 \exp[-\frac{(R_1 - S_1)^2}{2\beta^2} - \frac{2(R_2 - S_2)^2}{3\beta^2}] \Phi^B(R_1, R_2)$$

$$\times \phi_{G,A}[\exp\left(-\frac{(\xi_1 - S_1)^2}{B^2}\right) - \frac{(\xi_2 - S_2)^2}{3/4 B^2}\phi(\alpha_1)\phi(\alpha_2)\phi(\alpha_3)]}, \quad (1)$$

$$\Phi^B(R_1, R_2) \propto \phi_{G,A}[\exp\left(-\frac{(\xi_1 - R_1)^2}{b^2}\right) - \frac{(\xi_2 - R_2)^2}{3/4 b^2}\phi(\alpha_1)\phi(\alpha_2)\phi(\alpha_3)]}, \quad (2)$$

where $B^2 = b^2 + 2\beta^2$. The $^{12}$C is assumed to have a $z$–axial symmetry shape, i.e., $\beta \equiv (\beta_x = \beta_y, \beta_z)$. $\xi_1 = X_2 - X_1$ and $\xi_2 = X_3 - (X_1 + X_2)/2$. The $X_1$, $X_2$, and $X_3$ are the center-of-mass coordinates of the three clusters of $^{12}$C. In the above equation, $\Phi^B(R_1, R_2)$ is the $3\alpha$ Brink wave function [26] with the corresponding generator coordinates $R_1$

![FIG. 1. Schematic diagram for the container picture in the description of negative-parity cluster structure of $^{12}$C.](image)
and \( R_2 \). The \( R_1 \) represents the specified distance parameter of two clusters and \( R_2 \) the distance between the center-of-mass of these two clusters and the third \( \alpha \) cluster. The shift parameters \( S_1 \) and \( S_2 \) are introduced to deal with the negative-parity states. Their coordinates, \( S_1 \equiv (S_{1x}, 0, 0) \) and \( S_2 \equiv (S_{2x}, S_{2y}, 0) \), can arrange arbitrary triangular shapes in the \( xy \)-plane as shown in Fig.1. Thus, applying angular momentum and parity projection techniques, we can obtain negative-parity components from the extended THSR wave function. It should be noted that the above extended THSR wave function can also be considered as a Hybrid-THSR-Brink wave function because there are shift and width parameters. As for the Hamiltonian, the Volkov No.2 \cite{27} (modified version) with Majorana parameter \( M = 0.59 \) and harmonic-oscillator size parameter \( \nu = 1/(2b^2) = 0.275 \text{ fm}^{-2} \) is used, which is the same as employed by Kamimura et al. for their 3\( \alpha \) RGM calculations \cite{3}.

In practical calculations for the negative-parity states of \(^{12}\text{C} \), one important problem is how to set the proper shift parameters \( S_1 \) and \( S_2 \) in the three-cluster system as shown in Fig.1. The treatment of negative-parity states in \(^{20}\text{Ne} \) told us that \cite{13} the introduced shift parameter only plays the role of breaking the parity symmetry of the THSR wave function in an intermediate step while the size parameter is the essential dynamical variable for the description of clustering motion. In analogy with the \(^{20}\text{Ne} \) case \cite{13}, we slightly separate the 3\( \alpha \) clusters to form an equilateral triangle shape on \( xy \)-plane, namely \( S_1 = (S, 0, 0) \), \( S_2 = (0, \sqrt{3}/2S, 0) \), and take the side-length very small, i.e., \( S=0.5 \text{ fm} \). Note that the shift parameter \( S_1 \) actually can be zero due to the 2\( \alpha \) positive-parity character and the shift orientation can also be taken three dimensional. Here this kind of numerical "infinitesimal shifted triangle" is our first simple try and later we will discuss our wave function with respect to the dependence on shift parameters.

Based on this simple THSR wave function, we performed variational calculations after angular-momentum and parity projections for the 3\(^- \) and 4\(^- \) states in the two parameter \( \beta_x = \beta_y \) and \( \beta_z \) space. Here we only consider the dominant \( K=3 \) component in the single projected THSR wave functions. The reason why we only consider the 3\(^- \) and 4\(^- \) states and not also the 1\(^+ \) and 5\(^- \) states has to do with the fact that the latter states apparently have a more complicated structure or resonance character than describable with the present ansatz of our wave function. It is planned to devote an extra study for those states by superposing THSR wave functions and also applying the resonance method in the future. Here we concentrate on the first two states which are well describable with our technique as it will be demonstrated below.

![Energy contour plot for the 3\(^- \) state in the two-parameter space \( \beta_x = \beta_y \) and \( \beta_z \) in the container model.](image)

Figure 2 shows the energy contour plot for the 3\(^- \) state after variational calculations with respect to the \( \beta \) parameters. The value for the side-length of the shift equilateral-triangle is taken 0.5 fm as mentioned. It can be seen that two local minimum points have appeared, which are connected by a long and narrow valley. The deeper local minimum point appears at \( \beta_x = \beta_y =1.5 \text{ fm} \) and \( \beta_z = 3.0 \text{ fm} \). The obtained corresponding energy for the 3\(^- \) state is...
about \( \approx -80.9 \) MeV, which is very close to the result of the previous RGM calculation \(-81.2\) MeV by Kamimura et al. [3]. The second local minimum point appears at \( \beta_x = \beta_y = 2.0 \) fm and \( \beta_z = 0.5 \) fm, whose energy is about \(-80.7\) MeV. The wave functions of these two local minimum points are quite similar and their squared overlap is about 98%. Figure 3 shows the contour plot for the \( 4^- \) state. We also can find two local energy minima and the deeper one appears at \( \beta_x = \beta_y = 1.9 \) fm and \( \beta_z = 0.2 \) fm, whose energy is \(-76.9\) MeV. Another local minimum point appears at \( \beta_x = \beta_y = 1.2 \) fm and \( \beta_z = 3.0 \) fm. The value of the squared overlap between these two obtained optimal wave functions with prolate and oblate shapes is also as high as 98%. It can be seen that, this kind of equivalence of projected prolate and oblate THSR wave functions is very similar to the \(^{20}\)Ne case. This actually has its reason in the nonlocalized character of the THSR wave function which has been discussed in the Ref. [14]. It also should be noted that the \( 3^- \) and \( 4^- \) states are shown to have quite similar oblate intrinsic shapes (\( \beta_x = \beta_y \approx 2.0 \) fm and \( \beta_z \approx 0.5 \) fm) and we will see later that this character is very useful for our analysis of the intrinsic shapes.

![Energy contour plot for the 4^- state in the two-parameter space \( \beta_x = \beta_y \) and \( \beta_z \) in the container model.](image)

**FIG. 3.** Energy contour plot for the \( 4^- \) state in the two-parameter space \( \beta_x = \beta_y \) and \( \beta_z \) in the container model.

| \( J^n \) | \( E^{\text{Brink}}_{\text{min}}(R_1, R_2) \) | \( E^{\text{THSR}}(\beta) \) | \( E^{\text{Brink}}_{\text{GCM}} \) | \( |\langle \Phi^{\text{Brink}}_{\text{GCM}} | \Phi^{\text{Brink}}_{\text{GCM}}(R_1, R_2) \rangle|^2 \) | \( |\langle \Phi^{\text{Brink}}_{\text{GCM}} | \Phi^{\text{THSR}}(\beta) \rangle|^2 \) |
|----------|------------------|------------------|------------------|------------------|------------------|
| \( 3^- \) | -78.4            | -80.9            | -81.6            | 0.78             | 0.96             |
| \( 4^- \) | -74.4            | -76.9            | -77.8            | 0.72             | 0.92             |

In Fig. 4 we compare the energy curves obtained from the THSR wave functions with those from the Brink cluster model. At first, we assume the equilateral triangle shape with the length of side \( S \) for 3\( \alpha \) clusters in the Brink model, which is just equivalent to \( \beta \to 0 \) for the extended THSR wave function in Eq. (1). After variational calculations, it can be seen that there are two distinct pockets around 3 fm for the \( 3^- \) and \( 4^- \) states. It seems that, from this microscopic cluster model, there is a support for the rigid geometrical structure due to the non-zero inter-cluster distance parameter \( S \). However, if we introduce the width variable of the relative wave function, i.e., the non-zero size parameter \( \beta \), the minimum energy points appear around \( S \approx 0 \). This means the introduced \( S \) parameter, like in the \(^{20}\)Ne(\( \alpha + ^{16}\)O) case [13], only plays an important role for treating the negative-parity states as a shift parameter in this three-cluster system. In particular, the energy curves from the THSR wave function are very flat within the range...

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TABLE I. The calculated energies from the single optimal THSR wave functions in Eq. (1), the single optimal Brink wave functions in Eq. (2), and the Brink-GCM wave functions for the \( 3^- \) and \( 4^- \) states. The values of the squared overlap between the single optimal THSR/Brink wave functions and the Brink-GCM wave functions are also shown.
FIG. 4. Energy curves (calculated from the Eq. (1) and see the text.) of the 3− and 4− states of 12C within the THSR wave functions compared with those from the Brink wave functions. S < 2 fm. The obtained energies are almost degenerate without depending on the inter-cluster distance parameter S. Furthermore, it can be seen that the obtained energies in the container model for 3− and 4− states are 2.5 MeV deeper than those obtained from the Brink model. Moreover, we performed GCM calculations for 3− and 4− states by superposing hundreds of projected Brink wave functions with different inter-cluster generator coordinates and K numbers. The obtained GCM results are consistent with recent calculations from the real-time evolution method [28]. In Table I, we list the squared overlaps between the single THSR wave functions and the GCM wave functions, which are as high as 96% and 92% for the 3− and 4− states, respectively. On the other hand, the squared overlaps are only 78% and 72% in the case of the single optimal Brink wave functions, which were obtained by variational calculations from arbitrarily shaped triangle configurations. This indicates that the single specific triangle configuration cannot reproduce well these negative-parity states while these cluster states can be described very well in the container picture.

FIG. 5. Various density profiles from the THSR wave functions with different shift parameters before angular momentum and parity projections in the two-dimensional container picture. The size parameter is taken as βx = βy = 2.0 fm, βz = 0.5 fm. The squared overlaps between the corresponding projected THSR wave functions ΦJπ(S1x, S2x, S2y) and the Brink-GCM wave function are given. Figures share the same legend.

The obtained high accuracy of the THSR wave functions provides us with one clear nonlocalized cluster picture for understanding the intrinsic cluster structures of 12C. According to the obtained optimal THSR wave functions in Fig. 2 and Fig. 3, we take βx = βy = 2.0 fm and βz = 0.5 fm as the size parameters for the common intrinsic wave functions of 3− and 4− states in Eq. (1). Thus, the THSR wave function Φ(β, S1, S2) in Eq. (1) can be rewritten...
as $\Phi(S_{1x}, S_{2x}, S_{2y})$ with the tacit understanding that the size parameters are those from above. Various triangular shapes can then be constructed by taking the shift parameters $S_1$ and $S_2$ in the $xy-$plane. It can be seen that this adopted intrinsic THSR wave function has an oblate shape and it only slightly increases the widths of relative wave function in the $x y$ direction and almost makes no change in the $z$ direction compared with the Brink cluster model. Therefore, the actual situation can be considered as the three $\alpha$'s moving in a two-dimensional container. Figure 5 shows the densities of various intrinsic wave functions with different values of shift parameters. It should be noted that Figure 5(a) shows almost a positive-parity character of intrinsic density because of the "infinitesimal shifted triangle". As we know, by using the Brink-GCM wave functions, the energy levels, electric transitions and some other observables of $^{12}$C have been well reproduced [3][30]. Now, we found that these Brink-GCM wave functions are actually almost equivalent to our single THSR wave functions for the 3$^−$ and 4$^−$ states. This successful description of the 3$^−$ and 4$^−$ states can be considered as our first attempt to study the negative-parity states of 3$\alpha$ clusters in the new container model. The introduced shift parameter can not only handle the negative-parity problem, but also provides us with a useful way to understand the nonlocalized cluster motion. On the other hand, there remain other negative-parity states whose structure stays so far unexplained. In particular, as mentioned at the beginning, we found that the first 1$^−$ and 5$^−$ states cannot be well reproduced in our relatively simple picture of single THSR wave functions which indicates that these 1$^−$ and 5$^−$ states have some special characters compared with the 3$^−$ and 4$^−$ states. Quite recently, the 1$^−$ state at 10.8 MeV was reported [28] to have a strongly enhanced dipole transition strength from the ground state and could be regarded as an excitation mode of the Hoyle state. And the newly observed 5$^−$ state with the high-excitation energy 22.4 MeV [31] was actually difficult to be reproduced even with the GCM calculations. Therefore, in the next future step, it is expected that by including two $\alpha$ correlations or superposing enough configurations of the THSR-type wave function, these 1$^−$ and 5$^−$ states can be clarified in our model.

In summary, we successfully described the negative-parity 3$^−$ and 4$^−$ states in a simple container model. The obtained single optimal THSR wave functions for the 3$^−$ and 4$^−$ states have as high as 96% and 92% squared overlap with the corresponding GCM wave functions, respectively. From the densities of their intrinsic cluster wave functions, the nonlocalized cluster motion in the two-dimensional container was very clearly demonstrated.

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