Precise comparison of the Gaussian expansion method and the Gamow shell model

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We perform a detailed comparison of results of the Gamow shell model and the Gaussian expansion method supplemented by the complex scaling method for the same translationally invariant cluster-orbital shell model Hamiltonian. As a benchmark test, we calculate the ground state 0^+ and the first excited state 2^+ of mirror nuclei 6 He and 6 Be in the model space consisting of two valence nucleons in a *p* shell outside of a 4 He core. We find a good overall agreement of results obtained in these two different approaches, also for many-body resonances.

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I. INTRODUCTION

In recent years, the playground of nuclear physics has extended towards neutron and proton drip lines [1-3]. A huge amount of new experimental data on nuclei far from the valley of stability has been provided by new rare-isotope facilities. The knowledge of these nuclei has largely improved also due to the progress in theoretical methods and computing power which allows to calculate light nuclei in *ab initio* framework taking into account the proximity of the scattering continuum. The description of various manifestations of the continuum coupling requires the generalization of existing many-body methods and calls for theories which unify structure and reactions.

Realistic studies of the coupling to continuum in the manybody framework can be made in the open quantum system extension of the shell model (SM), the so-called continuum shell model (CSM) [4,5]. A recent realization of the CSM is the complex-energy CSM based on the Berggren ensemble [6], the Gamow shell model (GSM), which finds a mathematical setting in the rigged Hilbert space [7]. This model is a natural generalization of the standard SM for the description of configuration mixing in weakly bound states and resonances. The Berggren completeness relation can be derived from the Newton completeness relation [8] for the set of real-energy eigenstates by deforming the real momentum axis to include resonant poles which are located in the fourth quadrant of the complex k plane. Thus the Berggren completeness relation, which replaces the real-energy scattering states by the resonance contribution and a background of complex-energy continuum states, puts the resonance part of the spectrum on the same footing as the bound and scattering spectra. As the benefit of the explicit inclusion of the nonresonant continuum and resonant poles, the contribution of the unbound states to the one- and two-body matrix elements can be discussed. The Berggren ensemble has found the application in the GSM [9], time-dependent Green's function approach [10], the no-core GSM [11], the coupled cluster approach [12], the density matrix renormalization group (DMRG) approach [13], and in the coupled-channel GSM [14,15] to study various nuclear structure and reaction problems.

Another approach is the complex scaling (CS) method [16], which has been used to solve many-body resonances in many fields including atomic physics, molecular physics [17,18], and nuclear physics [19,20]. In the CS method, asymptotically divergent resonant states are described within \mathcal{L}^2 -integrable functions through the rotation of space coordinates and their conjugate momenta in the complex plane. As basis functions, the Gaussian expansion method (GEM) [21] has been extensively employed for the cluster-orbital shell model (COSM) [22] and coupled rearrangement channel model such as the TV model [23]. The CS-COSM has successfully been applied to a description of resonant states observed above the many-body decay threshold in *p*-shell nuclei (A = 5-8) using a ⁴He+XN model, where X = 1-4 and N = p, n [24,25]. The CS-TV model for the core+2N systems has been shown to reproduce the observed Coulomb breakup cross sections for three-body continuum energy states [26,27].

The purpose of these studies is to perform a detailed comparison of the GSM and the GEM+CS results for ⁶He and ⁶Be using the same COSM coordinates for valence nucleons [22] and the same Hamiltonian. In COSM, all coordinates are taken with respect to the core center-of-mass (c.m.), so that the translational invariance is strictly preserved. COSM combined with the CS method has been employed in numerous

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FIG. 1. Coordinate system of the COSM approach.

studies of weakly bound states and resonances in light nuclei [20,23,28–34]. COSM coordinates have been also used in the GSM [35] to investigate isospin mixing in mirror nuclei [36] and charge radii in halo nuclei [37].

The paper is organized as follows. In Sec. II we present our COSM Hamiltonian and the model space. In Sec. III, the two theoretical approaches, namely the GEM+CS (Sec. III A) and the GSM (Sec. III B), are briefly introduced. GEM+CS and GSM results for ⁶He and ⁶Be are presented and discussed in Sec. IV. Finally, Sec. V gives the main conclusions of these studies.

II. THE COSM HAMILTONIAN

In these studies, we employ the three-body model for ⁴He plus two-nucleon system in the COSM coordinates [22] (see Fig. 1). The Hamiltonian is written as follows:

$$\hat{H} = \sum_{i=1}^{2} \left(\hat{t}_i + \hat{V}_i^{(C)} \right) + \left(\hat{T}_{12} + \hat{v}_{12} + \hat{V}_{12}^{(C)} \right), \tag{1}$$

where \hat{t}_i and $\hat{V}_i^{(C)}$ are the kinetic and potential energy operators for the ⁴He core and an *i*th valence nucleon subsystem. In Eq. (1), the first parenthesis corresponds to the single-particle Hamiltonian for the *i*th valence nucleon, which is defined as

$$\hat{h}_i \equiv \hat{t}_i + \hat{V}_i^{(C)}, \quad (i = 1, 2).$$
 (2)

In the second parenthesis of Eq. (1), \hat{v}_{12} is the nucleon-nucleon interaction for valence particles, and

$$\hat{T}_{12} = -\frac{\hbar^2}{M^{(C)}} \nabla_1 \cdot \nabla_2 \tag{3}$$

is the recoil part which comes from the subtraction of thec.m. motion, due to the finite mass $M^{(C)}$ of the core nucleus. The last term of Eq. (1) is the three-body potential of ⁴He and two valence nucleons.

The interaction $\hat{V}_i^{(C)}$ between the core and the *i*th valence nucleon contains three terms:

$$\hat{V}_{i}^{(C)} = \hat{V}_{i}^{\alpha n} + \hat{V}_{i}^{\text{Coul}} + \lambda \,\hat{\Lambda}_{i}, \quad (i = 1, 2).$$
(4)

The nuclear interaction part $\hat{V}_i^{\alpha n}$ is the modified *KKNN* potential [23,38], which reproduces the α -*N* phase shifts in the low energy region. This potential contains a central and an

LS parts as

$$\hat{V}_{i}^{\alpha n}(r_{i}) = V_{0}^{\alpha n}(r_{i}) + 2V_{LS}^{\alpha n}(r_{i}) \boldsymbol{L} \cdot \boldsymbol{S}, \quad (i = 1, 2), \quad (5)$$

where r_i is the relative coordinate between ⁴He and the *i*th valence nucleon. The central part of Eq. (5) is written as

$$V_0^{\alpha n}(r_i) = \sum_{k=1}^5 \left[(-1)^{\ell_i} \right]_k V_k^0 \exp\left(-\rho_k^0 r_i^2\right), \tag{6}$$

where $[(-1)^{\ell_i}]_k$ is given by

$$[(-1)^{\ell_i}]_k = \begin{cases} 1 & (k=1,2)\\ (-1)^{\ell_i} & (k=3,4,5) \end{cases}.$$
 (7)

The LS part is

$$V_{LS}^{\alpha n}(r_i) = \sum_{m=1}^{3} f_m^{LS} V_m^{LS} \exp\left(-\rho_m^{LS} r_i^2\right),$$
(8)

where the factor f_m^{LS} is

$$f_m^{LS} = \begin{cases} 1 & (m=1) \\ 1 - 0.3 \times (-1)^{\ell_i} & (m=2,3) \end{cases}.$$
 (9)

Parameters of the modified *KKNN* potential [23,38] are shown in Table I.

For the Coulomb part \hat{V}_i^{Coul} in Eq. (4), we use a folded-type Coulomb interaction for the ⁴He+*p* subsystem:

$$\hat{V}_i^{\text{Coul}}(r_i) = \frac{2e^2}{r_i} \operatorname{Erf}(\alpha r_i),$$
(10)

where Erf(r) is the error function, and $\alpha = 0.828 \text{ fm}^{-1}$.

To eliminate the spurious states in the relative motion between the 4 He-core and the valence nucleon in CS, we use a projection operator [40]:

$$\hat{\Lambda}_i = \lambda |FS\rangle \langle FS|, \tag{11}$$

where the forbidden state in the ⁴He+*N* case; $|FS\rangle = |0s_{1/2}\rangle$, is given by the harmonic oscillator function with the size parameter b = 1.4 fm.

In the GSM, the forbidden state is eliminated from the set of the single-particle states, ϕ_i , as

$$\phi_i \Rightarrow (1 - \hat{\Lambda}_i)\phi_i. \tag{12}$$

We can confirm that the core-particle potential (4) with the parameters given in Table I, reproduces experimental energies and widths of $3/2_1^-$ and $1/2_1^-$ resonances in the ⁵He(⁴He+n) and ⁵Li(⁴He+p) systems.

TABLE I. Parameters of the modified *KKNN* potential [23,38] used in this calculation.

| $\overline{k} =$ | 1 | 2 | 3 | 4 | 5 |
|-----------------------------------|-------|-------|------|-------|------|
| V_k^0 [MeV] | -96.3 | 77.0 | 34.0 | -85.0 | 51.0 |
| ρ_k^0 [fm ⁻²] | 0.36 | 0.90 | 0.20 | 0.53 | 2.50 |
| V_k^{LS} [MeV] | -8.4 | -10.0 | 10.0 | _ | _ |
| ρ_k^{LS} [fm ⁻²] | 0.52 | 0.396 | 2.20 | - | - |

TABLE II. Parameters of the Minnesota potential [39].

| k | 1 | 2 | 3 |
|-----------------------------------|---------|-----------|----------|
| V_k^{nn} [MeV] | 200 | -178 | -91.85 |
| ρ_k^{nn} [fm ⁻²] | 1.487 | 0.639 | 0.465 |
| $W_k^{(u)}$ | u/2 | u/4 | u/4 |
| $M_k^{(u)}$ | (2-u)/2 | (2-u)/4 | (2-u)/4 |
| $B_k^{(u)}$ | 0 | u/4 | -u/4 |
| $\hat{H_k^{(u)}}$ | 0 | (2 - u)/4 | -(2-u)/4 |

For the two-body interaction $\hat{v}_{12}(\mathbf{r}_{12})$ of valence nucleons, where $\mathbf{r}_{12} \equiv \mathbf{r}_1 - \mathbf{r}_2$, we use the Minnesota potential [39]:

$$\hat{v}_{12}(\mathbf{r}_{12}) = \sum_{k=1}^{3} V_{k}^{0} \left(W_{k}^{(u)} - M_{k}^{(u)} P^{\sigma} P^{\tau} + B_{k}^{(u)} P^{\sigma} - H_{k}^{(u)} P^{\tau} \right) \\ \times \exp\left(-\rho_{k} r_{12}^{2}\right).$$
(13)

Parameters of this interaction are summarized in Table II, and the exchange parameter is taken as u = 1.0. The Coulomb interaction between valence protons is taken as an ordinary 1/r-type functional form.

It was shown that the binding energy of ⁶He cannot be reproduced using the reliable one- and two-body potentials for core-particle and particle-particle parts, respectively [23,29]. The correct binding energy in a system ⁴He+N + N is recovered by using a simple two-body Gaussian interaction, mimicking a physical three-body effect in the system [29] as

$$\hat{V}_{12}^{(C)}(r_1, r_2) = V_{\alpha n n}^0 \exp\left(-\rho_{\alpha n n} \left(r_1^2 + r_2^2\right)\right)$$
(14)

with the parameters $V_{\alpha nn}^0 = -0.41$ MeV and $\rho_{\alpha nn} = 5.102 \times 10^{-3}$ fm⁻².

III. THE MODELS

In this section, we discuss two models for solving ${}^{4}\text{He}+2N$ (*N* is proton or neutron) systems with the COSM Hamiltonian. One is the GEM+CS approach, and another one is the GSM approach. The essential differences between the GEM+CS and GSM approaches are the choice of the basis functions and the treatment of continuum states.

The basis function $\Phi(r_1, r_2)$ in the COSM is defined with a product of the functions with respect to each coordinate from the core to a valence nucleon,

$$\Phi(\mathbf{r}_1, \mathbf{r}_2)_{JM} \equiv [\mathcal{A}\{\phi_{\alpha_1}(\mathbf{r}_1) \otimes \phi_{\alpha_2}(\mathbf{r}_2)\}]_{JM}.$$
 (15)

Here, α_i denotes the angular part of the *i*th particle $\{j_i, \ell_i\}$, and its *z* components are implicitly included. \mathcal{A} is the antisymmetrizer for particles 1 and 2.

The basis function for the *i*th valence nucleon is

$$\phi_{\alpha_i}(\mathbf{r}_i) = f(r_i) |j_i m_i\rangle. \tag{16}$$

The angular momentum part of the basis function is constructed by using the normal jj-coupling scheme as

$$|JM\rangle = |[j_1 \otimes j_2]_{JM}\rangle. \tag{17}$$

The above coupling procedure is the same both for the GEM and GSM.

A. The Gaussian expansion method with complex scaling

The radial part of the GEM wave function is not an eigenfunction of the single-particle Hamiltonian \hat{h}_i , but the Gaussian function with the width parameter *a* as

$$f_{n_i}(r_i) \equiv u_{\ell_1}^{n_i}(r_i)$$

= $N_i r_i^{\ell_i} \exp\left(-\frac{1}{2}a_{n_i}r_i^2\right), \quad (i = 1, 2), \qquad (18)$

where N_i is the normalization, and ℓ_i is angular momentum for the *i*th nucleon.

The width parameter $a_{n_i} = 1/b_{n_i}^2$ in the GEM basis functions is defined using the geometric progression as $b_{n_i} = b_0 \gamma^{n_i-1}$ [21]. Here, b_0 and γ are input parameters, and n_i is an integer. The model space of the system is spanned by basis functions from $n_i = 1$ to N_{max} . The *k*th eigenfunction $\psi_{k:\alpha_i}$ of the core+N system can be obtained by diagonalizing the single-particle Hamiltonian \hat{h}_i with the Gaussian basis functions,

$$\psi_{k:\,\alpha_i}(\mathbf{r}_i) = \sum_m^{N_{\text{max}}} c_m^{(k)} \phi_{\alpha_i}^{(m)}(\mathbf{r}_i).$$
(19)

Here, $\hat{h}_i \psi_{k:\alpha_i} = \epsilon_k \psi_{\alpha_i}$, and $c_m^{(k)}$ are determined by using the variational principle.

For solving the core+2N system, the basis function (15) is given by the product of basis functions in Eq. (18) for particles 1 and 2 as follows:

$$\Phi_{JM}^{(m)} = \mathcal{A} \Big\{ u_{\ell_1}^{(m)}(r_1) \cdot u_{\ell_2}^{(m)}(r_2) | JM \rangle^{(m)} \Big\}.$$
(20)

Here, the width parameters $a_i^{(m)}$ in $u_{\ell_i}^{(m)}$ are prepared independently for particles 1 and 2. *m* is the index of the one-body basis functions.

The calculation of two-body matrix elements (TBME), $\langle \Phi^{(m)} | \hat{O}_{12} | \Phi^{(n)} \rangle$ can be performed analytically. Even for different Gaussian width parameters, we can obtain the value of TBME without any approximations.

The solution of the core+2N system can be obtained by diagonalizing the Hamiltonian

$$\hat{H}\Phi_{k;JM} = E_k \,\Phi_{k;JM},\tag{21}$$

and the corresponding eigenfunction is expressed as a linear combination of the basis functions,

$$\Phi_{k:JM} = \sum_{m}^{N_{\text{tot}}} C_m^{(k)} \Phi_{JM}^{(m)}.$$
 (22)

In order to treat the many-body resonant states, we apply the CS method. In this method, the coordinate and momentum are transformed using a rotation angle θ as

$$r \to r e^{i\theta} \quad (k \to k e^{-i\theta}).$$
 (23)

Resonance wave functions, which diverge in the asymptotic region, can be converged with this transformation for a suitable rotation angle. This essential feature is proven by the ABC-Theorem [16,41]. After this transformation, all continuum



FIG. 2. Complex-scaled eigenstates of the three-body Hamiltonian for the Borromean system. Solid circles are bound and resonance states, and open circles are continuum states.

states are aligned along the rotated axis. Furthermore, using the GEM, the continuum states are automatically discretized through the diagonalization of the Hamiltonian. A schematic figure of the bound states and resonances and discretized continuum states are shown for the Borromean system like ${}^{4}\text{He}+N+N$ in Fig. 2.

B. Gamow shell model approach

Another approach to study many-body resonances is the GSM approach [9,11–13]. This generalization of the nuclear SM treats single-particle bound, resonance, and continuum states on the same footing using a complete Berggren single-particle basis [6]:

$$\mathbf{1} = \sum_{i \in b, r} |\phi_i\rangle \langle \phi_i| + \oint_{\Gamma_k} dk |\phi(k)\rangle \langle \phi(k)|, \qquad (24)$$

where Γ_k is a deformed momentum contour. For each (ℓ, j) of the resonant single-particle state in the basis, the set $(\ell, j)_c$ of continuum states along the discretized contour in the *k* plane enclosing the resonant state(s) (ℓ, j) is included in the basis (see Fig. 3):

$$\mathbf{1} \simeq \sum_{i \in b, r} |\phi_i\rangle \langle \phi_i| + \sum_{\eta \in \text{cont}} |\phi(k_\eta)\rangle \langle \phi(k_\eta)|, \qquad (25)$$

where k_{η} are linear momenta discretized on the deformed contour with the parameters of a maximum k and a number of discretized points. Different shapes of (ℓ, j) contours are equivalent unless the number of resonant states contained in them changes. The complete many-body basis is then



FIG. 3. Deformed contour on the complex momentum plane (a), and discretized continuum states along the deformed contour (b). Solid circles are bound and resonant states, and open circles are discretized continuum states.

formed by all Slater determinants where nucleons occupy the single-particle states of a complete Berggren ensemble [9].

In the Berggren basis, the basis function of the core+2N system is

$$\Phi_{JM}^{(\nu)} = \mathcal{A}\{\left[\phi_1^{(\nu)} \otimes \phi_2^{(\nu)}\right]_{JM}\}.$$
 (26)

Here, $\phi_i^{(\nu)}$ are single-particle bound, resonance, and discretized-continuum states for particles 1 and 2.

In the GSM, the deformed contour for each (ℓ, j) is varied to obtain the best numerical precision of calculated eigenenergies and eigenvalues for a given discretization of the contour. Since the direct calculation of the TBME using the continuum and/or resonant single-particle states is numerically demanding, and even difficult to define from a theoretical point of view for some particular instances, one calculates the TBME using the harmonic oscillator (HO) expansion procedure [35]. For the TBME between GSM basis functions $\Phi_{\rm GSM}^{(i)}$ and $\Phi_{\rm GSM}^{(j)}$, one obtains

$$\langle \Phi_{\text{GSM}}^{(i)} | \hat{O}_{12} | \Phi_{\text{GSM}}^{(j)} \rangle$$

$$= \sum_{\alpha,\beta} \langle \Phi_{\text{GSM}}^{(i)} | \Phi_{\text{HO}}^{(\alpha)} \rangle \langle \Phi_{\text{HO}}^{(\alpha)} | \hat{O}_{12} | \Phi_{\text{HO}}^{(\beta)} \rangle \langle \Phi_{\text{HO}}^{(\beta)} | \Phi_{\text{GSM}}^{(j)} \rangle$$

$$= \sum_{\alpha,\beta} d_{i,\alpha}^* d_{j,\beta} \langle \Phi_{\text{HO}}^{(\alpha)} | \hat{O}_{12} | \Phi_{\text{HO}}^{(\beta)} \rangle,$$

$$(27)$$

where $\Phi_{\text{HO}}^{(\alpha)}$ are HO basis functions and $d_{i,\alpha}$ is the overlap between the GSM basis function $\Phi_{\text{GSM}}^{(i)}$ and the HO basis function:

$$d_{i,\alpha} \equiv \left\langle \Phi_{\rm HO}^{(\beta)} \middle| \Phi_{\rm GSM}^{(i)} \right\rangle. \tag{28}$$

The advantage of this procedure is that the TBMEs with the HO expansion can be stored for a fixed b_{HO} , and one only needs to calculate the overlaps $d_{i,\alpha}$, whatever the Berggren states are.

IV. RESULTS

For numerical calculations, we define the number of basis states. In the GEM+CS approach, the number of radial wave functions for each valence nucleon N_{max} is $N_{\text{max}} = 22$. The typical value of the Gaussian width parameters are $b_0 = 0.1$ fm and $\gamma = 1.3$. Hence, the maximum size of the width parameter becomes $b = b_0 \gamma^{N_{\text{max}}-1} = 0.1 \times 1.3^{21} \simeq 25$ fm. In the GSM, the continuum is discretized with 40 points for each partial wave and the maximum momentum is $k_{\text{max}} = 3.5$ fm⁻¹.

A. ⁶He in the ⁴He+2N model space

First, we show results for the ground state 0_1^+ and the first excited state 2_1^+ of ⁶He. The ground state of ⁶He is bound one with an energy E = 0.97 MeV from the ⁴He+2*n* threshold. Hence, we can take the rotation angle as $\theta = 0$ for the calculation of this state in GEM+CS approach.

We calculate energies of ⁶He by changing the maximum angular momentum for the coordinates r_1 and r_2 from $\ell_{max} = 1$ to 5, where ℓ_{max} is the maximum angular momentum in the basis function for the ⁴He+*n* subsystem. Parameters of the

TABLE III. Energies of the ground 0_1^+ and the first excited 2_1^+ states of ⁶He calculated using the GEM+CS and GSM approaches. All units except for the angular momentum are in MeV.

| | $\ell_{\rm max}$ | GEM+CS | GSM |
|----------------|------------------|----------------|----------------|
| | 1 | -0.117 | -0.116 |
| | 2 | -0.737 | -0.737 |
| $E(0_{1}^{+})$ | 3 | -0.870 | -0.870 |
| | 4 | -0.933 | -0.932 |
| | 5 | -0.978 | -0.977 |
| | $\ell_{\rm max}$ | GEM+CS | GSM |
| | 1 | 0.805 - i0.086 | 0.804 - i0.086 |
| | 2 | 0.675 - i0.038 | 0.669 - i0.041 |
| $E(2_{1}^{+})$ | 3 | 0.628 - i0.027 | 0.619 - i0.030 |
| | 4 | 0.605 - i0.023 | 0.595 - i0.026 |
| | 5 | 0.589 - i0.021 | 0.577 - i0.024 |

interaction are chosen to reproduce the binding energy of the ground state of ⁶He in a model space with $\ell_{max} = 5$.

The energies of the 0_1^+ state are shown in Table III for different values of ℓ_{max} . One can see that the calculation for $\ell_{max} = 1$, which includes the $s_{1/2}$, $p_{3/2}$, and $p_{1/2}$ orbits of the ⁴He+N system, is not enough to reproduce the binding energy of ⁶He. The inclusion of higher angular momenta ($\ell_{max} \ge$ 2) improves the calculated energy significantly. Nevertheless, even $\ell_{max} = 5$ is not enough to obtain the converged ground state energy since the T-type Jacobi configuration of valence neutrons is very important [23]. However, since the scope of this paper is to compare results of the GEM+CS approach and the GSM, we restrict the maximum angular momentum for the core+N system to $\ell_{max} = 5$ and determine the interaction parameters in this model space.

We find a good agreement between GEM+CS and GSM for a Borromean ⁶He nucleus. The ℓ_{max} dependence of the 0_1^+ and 2_1^+ energies is shown in Table III and Fig. 4.

The density of valence neutrons in the 0_1^+ state of ⁶He is plotted in Fig. 5. One can see that the GEM+CS and



FIG. 4. Convergence of the poles of the ground 0_1^+ and the first excited 2_1^+ states of ⁶He, which are calculated using the GEM+CS approach and the GSM for $1 \le \ell_{max} \le 5$. Open and solid circles denote GEM+CS and GSM results, respectively.



FIG. 5. (Color online) The density of valence neutrons in the COSM coordinate system for the ground 0_1^+ state of ⁶He. The normalization of the density distribution is 1.

GSM approaches give indistinguishable results for the density distributions in the 0^+_1 halo configuration of ⁶He.

Results for the 2_1^+ narrow resonance are shown in Table III and Fig. 4. The difference between GEM+CS and GSM results in this case is at most ~10 keV. The trajectories of the 2_1^+ state of the GEM+CS and GSM poles are shown in Fig. 4. Similarly, as for the 0_1^+ state, results of the GEM+CS and GSM approaches agree well.

B. ⁶Be in the ⁴He+2N model space

The ⁶Be nucleus, the mirror system of ⁶He, is unbound in the ground state. In this section, we shall compare results of the GEM+CS and GSM for the 0_1^+ and 2_1^+ states of ⁶Be described as a ⁴He+2*p* three-body system.

Calculated energies of the 0_1^+ and 2_1^+ states for different ℓ_{max} values are shown in Table IV. The difference of GEM+CS and GSM energies is less than ~ 10 keV for the 0_1^+ state and ~ 20 keV for the 2_1^+ state.

The trajectory of the 0_1^+ and 2_1^+ poles in the complex energy plane is shown in Fig. 6. Contrary to the 0_1^+ state, one may notice a slight difference between trajectories of 2_1^+ poles in the GEM+CS approach and in the GSM. This difference diminishes with increasing ℓ_{max} .

TABLE IV. Energies of the ground 0_1^+ and the first excited 2_1^+ states of ⁶Be calculated using the GEM+CS and GSM approaches. All units except for the angular momentum are in MeV.

| | ℓ_{max} | GEM+CS | GSM |
|----------------|------------------|----------------|----------------|
| | 1 | 1.932 - i0.152 | 1.926 - i0.146 |
| | 2 | 1.490 - i0.046 | 1.482 - i0.041 |
| $E(0_{1}^{+})$ | 3 | 1.380 - i0.036 | 1.374 - i0.030 |
| | 4 | 1.324 - i0.033 | 1.318 - i0.026 |
| | 5 | 1.285 - i0.031 | 1.279 - i0.024 |
| | $\ell_{\rm max}$ | GEM+CS | GSM |
| | 1 | 2.741 - i0.703 | 2.776 - i0.711 |
| | 2 | 2.614 - i0.559 | 2.610 - i0.596 |
| $E(2_{1}^{+})$ | 3 | 2.565 - i0.518 | 2.538 - i0.543 |
| | 4 | 2.537 - i0.500 | 2.512 - i0.518 |
| | 5 | 2.517 - i0.491 | 2.495 - i0.505 |
| | | | |



FIG. 6. Poles of the ground and first excited states of ⁶Be calculated using GEM+CS approach and GSM from $\ell_{max} = 1$ to 5. Open and solid circles correspond to GEM+CS and GSM results, respectively.

C. Discussion

In the comparison between the GEM+CS and GSM approaches, we obtain a good agreement for the bound state 0_1^+ in ⁶He, and narrow resonances; 2_1^+ in ⁶He and 0_1^+ in ⁶Be. A small difference appears only for the 2_1^+ broad resonance in ⁶Be. Below, we shall discuss a possible origin of such a small difference in the numerical results.

Both GEM+CS and GSM approaches solve the non-Hermitian problem. In the GEM+CS approach, the wave function of a resonance becomes \mathcal{L}^2 integrable with the help of the complex rotation. As a result, the Hamiltonian becomes non-Hermitian. The standard procedure to find the optimum values of the parameters is to search for a stationary point of the eigenvalue with respect to the variational parameters. The variational parameters are the complex rotation angle θ and the parameter b_0 in a definition of the Gaussian width: $b_{n_i} = b_0 \gamma^{n_i - 1}$ [20] for the Gaussian basis functions. The optimization procedure is a simplified version of the generalized variational principle for complex eigenvalues [42]. The procedure works efficiently and gives very accurate solutions even for broad resonant states [20].

The GSM is formulated in the Berggren set, which includes bound single-particle states, single-particle resonances, and scattering states from the discretized contour for each considered (ℓ, j) . Consequently, the Hamiltonian matrix in this basis becomes complex-symmetric. The number of scattering states on each discretized contour (ℓ, j) and the momentum cutoff have to be chosen to assure the completeness of many-body calculations. Moreover, in the HO expansion procedure of calculating the TBMEs, the dependence on the oscillator length and the number of oscillator shells should be carefully examined.

Figure 7 presents a trajectory of the 2_1^+ narrow resonant pole of ⁶He calculated in the GEM+CS approach by changing the rotation angle θ , where the stationary point at the optimum value of the rotation angle is $\theta_{opt} = 13^\circ$, and the optimum point for the GSM calculation, which is obtained with the oscillator length and the number of oscillator shells are $b_{HO} = 2$ fm



FIG. 7. Poles of ⁶He (2⁺) with $\ell_{max} = 1$. For the GEM+CS, we change the rotation angle θ from 5° to 25° in steps of 1°.

and N = 41, respectively. In this case, the difference is only ~ 1 keV, and both methods give almost the equivalent result.

On the other hand, the 2_1^+ state of ⁶Be is a broad resonant pole due to the presence of the Coulomb force for all three particles. The optimum value of the rotation angle in the GEM+CS calculation is $\theta_{opt} = 17^\circ$, and the optimal HO oscillator length in GSM calculations is $b_{HO} = 3$ fm. The difference of complex GEM+CS and GSM eigenenergies becomes in the order of 10 keV. To improve the agreement for the eigenvalues obtained by the GEM+CS and GSM, it would be necessary to examine the optimization of the variational parameters more precisely. However, in the practical point of view, the difference is only less than 1 percent to the total energy.

The convergence can be tested by introducing an extrapolation procedure, e.g., the Richardson extrapolation [43]. We extrapolate the energy *E* as a function of $1/\ell_{\text{max}}$ to $1/\ell_{\text{max}} = 0$. The energies $E(1/\ell_{\text{max}})$ of the 2⁺ state of ⁶Be become 2.483 – *i*0.474 and 2.464 – *i*0.481 (MeV) for the GSM+CS and GSM, respectively. The difference becomes smaller than that of the $\ell_{\text{max}} = 5$ case. Hence, we can conclude that both methods provide a sufficient accuracy even for the calculation of the broad resonant states.

V. SUMMARY

The GSM and GEM+CS are two different theoretical approaches which allow to describe unbound resonant states. These two approaches differ in the choice of the basis functions and the numerical procedure to obtain the eigenvalues. To benchmark the GSM and GEM+CS approaches, we have performed a precise comparison for weakly bound and unbound states using the same Hamiltonian in the COSM coordinates preserving the translational invariance. For a weakly bound ground state of ⁶He, the GSM and GEM+CS give essentially identical results. For the three-body resonance states, the GEM+CS and GSM give very close results proving the reliability of both schemes of the calculation for unbound states. The slight difference between GSM and GEM+CS results for broad resonances may have different origins. The

HO expansion procedure in calculating the TBMEs in the GSM may lead to rounding errors, especially for broad many-body resonances. On the other hand, the stationarity condition in the GEM+CS approach could also be a source of small imprecision for broad resonances. Based on our results, we conclude that both approaches are essentially equivalent for all quantities studied. The other work for a comparison in the ⁶He system has been done and also shows a good agreement between two different approaches [44].

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