Ab initio no-core Gamow shell model calculations with realistic interactions

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No-core Gamow shell model (NCGSM) is applied to study selected well-bound and unbound states of helium isotopes. This model is formulated on the complex energy plane and, by using a complete Berggren ensemble, treats bound, resonant, and scattering states on equal footing. We use the density matrix renormalization group method to solve the many-body Schrödinger equation. To test the validity of our approach, we benchmarked the NCGSM results against Faddeev and Faddeev-Yakubovsky exact calculations for ³H and ⁴He nuclei. We also performed *ab initio* NCGSM calculations for the unstable nucleus ⁵He and determined the ground-state energy and decay width, starting from a realistic N³LO chiral interaction.

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I. INTRODUCTION

In the last decade our knowledge of nuclei far from the valley of stability has radically improved. This improvement is a byproduct of advances in both experiment and theory. New experimental facilities that have already been built (RIBF at RIKEN) or are being constructed (SPIRAL2 at GANIL, FAIR, FRIB at MSU) will give us a better insight of areas in the nuclear chart that have never been explored, pushing even farther the limits of nuclear existence. A few decades ago, the nuclear chart consisted of approximately 1000 isotopes, whereas in 2011 this number has been expanded to approximately 3000 species, and an estimated number of nuclei that can exist in nature or can be synthesized in the laboratory is approximately 7000 [1]. The increase in computing power has made it possible to calculate the properties of nuclei in an *ab initio* manner, using realistic interactions, which reproduce the nucleon-nucleon scattering data. For few-body systems ($A \leq 4$) methods such as Faddeev [2] and Faddeev-Yakubovsky (FY) [3] provide an exact solution to the many-body problem. Methods such as the Green's function Monte Carlo (GFMC) [4], the hyperspherical harmonics (HH) [5,6], the no-core shell model (NCSM) [7,8], the coupled-cluster approach (CC) [9], and more recently, the in-medium similarity renormalization group method (IM-SRG) [10,11] and Dyson self-consistent Green's function (Dyson-SCGF) method [12] have been applied successfully for the *ab initio* description of light and medium mass nuclei.

Nuclei with a large isospin that can be found in regions far away from the valley of stability have attracted a great deal of interest. They belong to the category of open quantum systems (OQSs) [13], which in the case of the nucleus are interconnected via the decay and reaction channels. They are very fragile objects with small separation energies and very large spatial dimensions. The proximity of the continuum affects their bulk properties (matter and charge distributions) and their spectra. Phenomena such as the anomalous behavior of elastic cross sections and the associated overlap integral near threshold states in multichannel coupling (Wigner cusps) [14,15], the isospin and mirror symmetry-breaking threshold effects [16,17], the resonance trapping [18–20] and superradiance phenomenon [21,22], the appearance of cluster correlations in the vicinity of the respective cluster emission threshold [23], the modification of spectral fluctuations [24–26], and deviations from Porter-Thomas resonance width distribution [20,27,28], are all unique manifestations of the continuum coupling.

For their theoretical explanation it was necessary to generalize existing many-body methods, and create theories which unify structure and reactions. Examples of these attempts are the shell model embedded in the continuum (SMEC) [29-31] and the Gamow shell model (GSM) [32-35]. The SMEC is a recent realization of the real-energy continuum shell model [36,37], which uses the Feshbach projection technique [38] in order to take into account the coupling to the scattering continuum. The GSM is a generalization of the harmonicoscillator-based shell model in the complex energy plane by using the Berggren ensemble [39], which treats resonant and nonresonant states on equal footing. Ab initio calculations that can describe bound and unbound states of nuclei, include the NCSM coupled with the resonating group method [40-42], the CC approach generalized in the complex-energy plane using the Berggren basis [43–45], and the GFMC [46,47].

In this work we introduce the no-core Gamow shell model (NCGSM) as an alternative for calculations of weakly bound and unbound states of light nuclei using realistic interactions and allowing all the nucleons to be active. The paper is organized in the following manner: in Secs. II and III, we describe the basic ingredients of our method, such as the many-body Hamiltonian, the single-particle basis we employ, the way the two-body matrix elements are calculated within the Berggren basis, and we discuss the translational invariance of our Hamiltonian. In Sec. IV we describe the density matrix renormalization group (DMRG) method, which is an efficient

tool for a diagonalization of large complex-symmetric GSM matrices. In Sec. V we present our calculations for the ³H, ⁴He, and ⁵He nuclei and, finally, in Sec. VI we discuss the conclusions and the future perspectives.

II. HAMILTONIAN

Our goal is to solve the A-body Schrödinger equation

$$H|\Psi\rangle = E|\Psi\rangle,\tag{1}$$

where H is the intrinsic Hamiltonian

$$H = \frac{1}{A} \sum_{i < j}^{A} \frac{(\vec{p}_i - \vec{p}_j)^2}{2m} + \sum_{i < j}^{A} V_{ij}^{NN}, \qquad (2)$$

and V^{NN} a realistic NN interaction. For (2) the following identities are useful:

$$\vec{P} = \sum_{i=1}^{A} \vec{p_i},\tag{3}$$

where \vec{P} is the center of mass (CoM) momentum, and

$$\sum_{i=1}^{A} \vec{p}_{i}^{2} = \frac{1}{A} \left[\vec{P}^{2} + \sum_{i < j} (\vec{p}_{i} - \vec{p}_{j})^{2} \right], \quad (4)$$

resulting in:

$$\sum_{i=1}^{A} \frac{\vec{p}_i^2}{2m} - \frac{\vec{P}^2}{2mA} = \frac{1}{2mA} \sum_{i < j} (\vec{p}_i - \vec{p}_j)^2.$$
(5)

In the NCGSM there is no restriction on the type of the NN interaction, contrary to the GFMC approach for example, where difficulties arise for nonlocal potentials. One can use a local interaction, such as the Argonne v_{18} potential [48], or a nonlocal interaction, such as the CD-Bonn 2000 [49] and various chiral interactions. There is also the possibility to use renormalized versions of the aforementioned forces, by applying techniques such as $V_{\text{low-}k}$ [50], the similarity renormalization group (SRG) approach [51,52], or the G matrix [53,54]. In this work we employ the phenomenological Argonne v_{18} potential and the chiral N³LO interaction [55], which is consistent with the symmetries of the QCD Lagrangian. Both potentials were renormalized via the V_{low-k} method with a sharp momentum cutoff $\Lambda = 1.9 \text{ fm}^{-1}$ to decouple high from low momentum degrees of freedom and, henceforth, improving the convergence of nuclear structure calculations [56]. Moreover, specific interactions will be used to compare NCGSM with other approaches.

III. BERGGREN BASIS

In previous applications of the GSM, where a tightly bound core was assumed (⁴He or ¹⁶O), the single-particle (s.p.) basis was usually generated by solving the one-body Schrödinger equation with a Woods-Saxon (WS) potential, parameterized to reproduce the core plus nucleon spectrum. In the case of the NCGSM, the s.p. basis will be generated by the realistic two-body interaction itself by solving the integrodifferential Schrödinger equation, which contains both local and nonlocal parts [57,58]. This numerical method is known as the Gamow Hartree-Fock (GHF) method since it generates a microscopic basis that includes resonant and nonresonant states. The GHF method can be applied not only in spherical cases (closed shells) but also in deformed cases (nonclosed shells) [57]. The one-body self-consistent potential $U_{HF}(r)$ is then used to solve the one-body Schrödinger equation:

$$u_k''(r) = \left[\frac{\ell(\ell+1)}{r^2} + \frac{2m}{\hbar^2}U_{HF}(r) + V^c(Z_c, r) - k^2\right]u_k(r),$$
(6)

where $V^{c}(Z_{c}, r)$ is the one-body Coulomb potential:

$$V^{c}(Z_{c},r) = \frac{C_{c}Z_{c}\operatorname{erf}(\alpha r)}{r}$$
(7)

and C_c is the Coulomb constant, Z_c the proton number and α is a constant, which is given by $\alpha = 3\sqrt{\pi}/4R_0$. The reason we choose an error function to approximate the Coulomb field and not, for example, the field that is produced by a uniformly charged sphere at R_0 , lies in the fact that the latter is nonanalytic at R_0 . The value of R_0 is chosen in a way that the Coulomb potential of Eq. (7) and the potential of a uniformly charged sphere are equal at the origin. The wave number k is defined as $k = \sqrt{2mE}/\hbar$ and is, in general, complex. Equation (6) is solved with the requirement that at large distances the wave function will behave as a linear combination of Hankel or Coulomb functions for neutrons and protons, respectively:

$$u_k(r) \sim C^+ H_{\ell,\eta}^{(+)}(kr) + C^- H_{\ell,\eta}^{(-)}(kr), \tag{8}$$

where η is the Sommerfeld parameter. The C^+ and C^- coefficients are determined by the normalization of the radial functions u(r) to a Dirac's δ distribution:

$$\int_0^\infty u_k(r)u_{k'}(r)\,dr = \delta(k-k') \tag{9}$$

The solutions of (6) that satisfy pure outgoing boundary conditions $[C^- = 0$ in Eq. (8)] correspond to the poles of the *S* matrix and they are represented as dots in the complex *k* plane of Fig. 1. While normalization of bound states does not pose any difficulty, one should pay more attention on the normalization of resonances. The latter diverge exponentially for large distances and the regularization method that is used for the calculation of their norm is the external complex scaling [34]. The method facilitates from the fact of using complex radii for the integration of resonant wave functions:

$$\int_0^\infty u_k^2(r) \, dr = \int_0^R u_k^2(r) \, dr + \int_R^\infty u_k (R + x e^{i\theta})^2 e^{i\theta} \, dx,$$
(10)

with *R* chosen sufficiently large so as to match the condition (8) for $C^- = 0$, while θ is the angle of external rotation, which satisfies the condition that $u_k(R + xe^{i\theta}) = 0$ for $x \to \infty$.

It was shown by Berggren [39] that for a given partial wave (ℓ, j) the scattering states, which are distributed along



FIG. 1. (Color online) An illustration of the Berggren singleparticle (s.p.) basis used in the NCGSM, showing the position of resonant (bound states and resonances) states in the complex k plane. The nonresonant continuum states lie along the complex contour L_+ .

the L_+ contour and the resonant solutions (bound states and/or resonances) of (6) form a complete set:

$$\sum_{n} |u_{n}\rangle \langle \widetilde{u_{n}}| + \int_{L_{+}} |u_{k}\rangle \langle \widetilde{u_{k}}| dk = 1.$$
 (11)

The tilde symbol means that the complex conjugation arising in the dual space affects only the angular part and leaves the radial part of the wave function unchanged. The completeness reassures us that any function that lies between the contour and real k axis and exhibits outgoing wave asymptotic (e^{ikr}), can be expanded using (11). In practice the integral in Eq. (11) is discretized by means of an appropriate quadrature rule (the Gauss-Legendre quadrature in our case) and we end up with a discretized completeness relation:

$$\sum_{i=1}^{N} \omega_n |u_n\rangle \langle \widetilde{u_n}| \simeq 1, \qquad (12)$$

where $\omega_n = 1$ both for resonant states and the Gauss-Legendre weight for nonresonant states along the discretized contour. The approximate equality in Eq. (12) arises from the finite discretization of the contour. In addition, the discretized contour does not extend to infinity and we use a maximum cutoff (k_{max}) at around 4 fm⁻¹. We have checked that results do not depend on the values of the k_{max} for an adequately large number of discretization points.

In Fig. 2 we show the radial behavior of a few Berggren basis states that were generated from the N³LO interaction with a $V_{\text{low-}k}$ cutoff $\Lambda = 1.9 \text{ fm}^{-1}$. The states that are plotted refer to neutron states. The $0p_{3/2}$ resonant state has a complex energy and lies in the fourth quadrant of the complex k plane (see also Fig. 1). It is a solution of Eq. (6) with outgoing wave boundary conditions at large distances. We, indeed, observe a localization of the state in the region of the attractive nuclear potential ($r \leq 2$ fm) and an outgoing wave behavior



FIG. 2. (Color online) Selected squared radial basis functions in the NCGSM. Resonant (bound, resonances) and nonresonant (scattering) states are plotted. The basis generating potential is produced by the $V_{\text{low-}k}$ N³LO interaction for the ⁵He nucleus. The notation *nlj* is explained in the text.

for distances beyond the range of the nuclear potential. This is the basic characteristic of a metastable s.p. state.

In order to satisfy Berggren's completeness, the scattering contour L_+ has to be complex. It is then understood that the $10p_{3/2}$ is a point along the complex contour. It corresponds to a state, which is given as a linear combination of Hankel functions, as it is seen in Eq. (8). Here we are using the notation $n\ell j$, where n is the radial quantum number and is identified as the tenth Gauss-Legendre discretization point on the L_+ contour; ℓ is the s.p. angular momentum of the state ($\ell = 1$ in this example), and j is the s.p. total angular momentum. The $0s_{1/2}$ resonant state is bound and lies on the imaginary momentum axis with a real and negative energy. At large distances, the state is decaying as an exponential. Also shown are the $10s_{1/2}$ and $10p_{1/2}$ scattering states, which both lie on the real momentum axis, since there is no complex resonant state associated with these partial waves. Similar to the $10p_{3/2}$ complex state, they correspond to the temth discretization point of the Gauss-Legendre rule, but the contour lies along the real momentum axis. At this point we would like to highlight that the Berggren basis not only imposes the correct quantum mechanical asymptotic behavior of the s.p. states¹ but also includes the continuum states in a rigorous manner, promoting it to an ideal and realistic basis for the description of metastable and weakly bound states.

The completeness relation (12) is similar to a usual discrete completeness relation, such as the harmonic oscillator (HO) one, and results in an eigenvalue problem. The many-body basis states are Slater determinants $|SD_n\rangle = |u_1, \ldots, u_A\rangle$, where u_k is a resonant (bound state or resonance) or nonresonant (scattering) state. In this basis, the Hamiltonian matrix is complex symmetric and upon diagonalization, many-body

¹This can be immediately checked by the fact that a bound resonant state (e^{ikr}) behaves as $e^{-\kappa r}$ for $r \to \infty$, since $k = i\kappa$.

correlations and coupling to the continuum are taken into account simultaneously. As a direct consequence of Eq. (12), the many-body states also satisfy the completeness relation:

$$\sum_{i=1}^{N} |\widetilde{SD}_n\rangle \langle SD_n| \simeq 1.$$
(13)

The squares of the linear expansion coefficients and not by their absolute values, satisfy the relation:

$$\sum_{i=1}^{N} c_n^2 = 1.$$
(14)

Furthermore, the completeness relation (13) can be used for the calculation of two-body matrix elements (TBMEs) between Berggren basis states.

A. TBMEs of realistic interactions in the Berggren basis

Nuclear part. Matrix elements of realistic interactions are defined in a relative and CoM system of coordinates. In order to work in a basis of Slater determinants, a transformation from the relative and CoM to the laboratory system is necessary. When working in the HO s.p. basis, this is possible through the Brody-Moshinsky brackets [59]. For a different basis such as the Berggren basis, one has to perform a multiple decomposition of the realistic NN interaction and calculate two-dimensional radial integrals. Matrix elements between scattering states need to be regularized by means of the complex scaling (CS) technique [34], which unfortunately does not work for just any type of integrals and can cause numerical instabilities. The problem is alleviated by expanding the NN interaction in a truncated HO basis [60]:

$$V_{\rm NN} = \sum_{\alpha\beta\gamma\delta}^{N_{\rm max}} |\alpha\beta\rangle\langle\alpha\beta|V_{\rm NN}|\gamma\delta\rangle\langle\gamma\delta|.$$
(15)

The matrix elements of the NN interaction in the Berggren ensemble become then:

$$\langle \widetilde{ab} | V_{\rm NN} | cd \rangle = \sum_{\alpha\beta\gamma\delta}^{N_{\rm max}} \langle \widetilde{ab} | \alpha\beta\rangle \langle \alpha\beta | V_{\rm NN} | \gamma\delta\rangle \langle \gamma\delta | cd \rangle.$$
(16)

We end up calculating overlaps between HO and Berggren states $\langle \alpha\beta | ab \rangle$ and $\langle \gamma\delta | cd \rangle$, where the Latin letters denote Berggren states and Greek letters HO states. Due to the Gaussian falloff of the HO states, no complex scaling is needed for the calculation of these integrals. On the other hand, matrix elements of the NN interaction in the HO basis $\langle \alpha\beta | V_{NN} | \gamma\delta \rangle$ can be conveniently calculated using the Brody-Moshinsky brackets [59].

The method of handling matrix elements in the Berggren basis using the projection of continuum onto the HO states should not be confused with basis expansion methods suitable for the description of closed quantum systems. Only the short-range part of the nuclear interaction is expanded in the HO basis. The kinetic energy operator is calculated in the Berggren basis, so the calculations of weakly bound and unbound systems are possible. With this formulation it is also clear that there is no restriction on the type of NN interaction one can use in the NCGSM. What we need is just the nuclear matrix elements in the HO basis.

Coulomb interaction treatment. For nuclear systems with two or more protons the two-body Coulomb interaction is included in the Hamiltonian (2). The method we adopt for the treatment of the long-range Coulomb interaction was first used in the description of isospin breaking due the continuum coupling [17] and it was also recently applied to calculate reaction observables [61,62] (see also Ref. [63] for a detailed numerical analysis). The basic idea of the method is to add and subtract the one-body Coulomb potential (7) (with Z = 2, e.g., in case of ³He or ⁵He) from the two-body Coulomb interaction: $V_c(1, 2) = V^c(1) + [V_c(1, 2) - V^c(1)]$. Then the second term in the parentheses has a short-range character and the HO expansion method of (15) can be applied. Matrix elements of the Coulomb interaction can be calculated using the Brody-Moshinsky brackets without the need to perform an external CS calculation.

We would like to mention here that this method of treating the two-body Coulomb interaction would be of particular interest when one has to deal with many-body proton resonances, such as in the ⁶Be nucleus. In this case, calculating the Coulomb interaction potential by simply expanding it in a HO basis [see Eq. (15)], would be a rather poor approximation.

Center of mass (CoM) motion in the NCGSM. By adopting a s.p. basis upon which we build many-body basis states, we effectively localize the nucleus in space and, hence, we break the translational invariance of the Hamiltonian. Moreover, we would need 3A - 3 coordinates to describe it, but the nuclear wave function we construct, depends on 3A coordinates, where A is the total number of particles. These redundant degrees of freedom are responsible for the CoM spuriosity that appears in many-body methods. On the other hand, plane-wave s.p. states are eigenstates of the momentum operator and, hence, preserve the translational invariance, but unfortunately cannot be used to describe a localized system. The alternatives are: (i) Solving the many-body problem using relative coordinates (e.g. Jacobi coordinates), which reassures the translational invariance of the system, with the price of unfeasible antisymmetrization of states for A > 8, and (ii) using the unique analytical properties of the HO s.p. basis in a full $N\hbar\omega$ space, in which the total wave function is factorized into $|\psi_{rel}\rangle \otimes |\psi_{CoM}\rangle$, limiting though the application to well-bound systems only. In the case of the NCGSM the latter factorization is not guaranteed and it has to be demonstrated numerically. Since our Hamiltonian (2) is intrinsic, we are expecting that in an infinite space there would be no spuriosity. However, because we are working in a finite space, it is necessary to check numerically this condition.

Assuming that the factorization into a CoM and a relative wave function is valid and also the CoM wave function has a Gaussian shape, we calculate the expectation value of the CoM operator [64]:

$$H_{\rm CoM} = \frac{1}{2mA}\vec{P}_{cm}^2 + \frac{mA\omega^2}{2}\vec{R}_{cm}^2 - \frac{3}{2}\hbar\omega, \qquad (17)$$

where $\hbar\omega$ is the parameter that characterizes the Gaussian wave function. The matrix elements of (17) are calculated with the HO expansion method of (15) and the analytical formulas for

their expression are found in Ref. [65]. Following the assertion of Ref. [66], if $\langle H_{cm} \rangle \sim 0$ then the factorization is valid.

IV. RESOLUTION OF THE MANY-BODY SCHRÖDINGER EQUATION WITH THE DMRG METHOD

The Schrödinger equation (1) is solved within a many-body basis constructed from the discrete set of single-particle states $|u_i\rangle$ (12). The discretization of the integral along the contour L_+ in Eq. (11) should be precise enough so that the discretized completeness relation (12) is fulfilled. In other words, the number of discretized shells should be increased until Eq. (12) holds. As a consequence, the dimension of the many-body model space will increase dramatically with the number of nucleons and number of shells. Efficient numerical methods allowing the diagonalization of large Hermitian as well as complex-symmetric matrices are then required to solve the NCGSM problem.

In this paper, we have used one of these methods, namely, the DMRG method [67,68], which has been generalized in the context of the GSM in Refs. [69,70]. The GSM/DMRG approach has been applied previously to study several weakly-bound/unbound nuclei described as few-valence-nucleon systems interacting via schematic two-body forces above an inert core. In this paper, all nucleons are considered active and realistic two-body interactions are used but nevertheless, the application of the DMRG method is similar. In the following, we recall the main ideas of the DMRG in the multishell GSM problem [70].

The purpose of the DMRG method is to allow the calculation of the many-body poles of the scattering matrix of the NCGSM Hamiltonian \hat{H} by performing efficient truncations of the many-body model space. As the contribution of the nonresonant continuum to the structure of many-body bound/resonant eigenstates of \hat{H} is usually smaller than the contribution from the bound/resonant orbits, the following separation is performed: the many-body states constructed from the s.p. poles form a subspace H (the so-called reference subspace), and the remaining states containing contributions from nonresonant shells form a complement subspace P. The set \mathcal{E} of many-body basis states (13) can then be written as

$$\mathcal{E} = H \otimes P. \tag{18}$$

The DMRG technique is then used to perform truncations in \mathcal{E} by keeping only selected optimized states in P in the sense of a criteria based on the density matrix in P (see below for a more explicit explanation).

Let us assume that we want to calculate an eigenstate $|\Psi\rangle$ of \hat{H} for a nucleus coupled to the total angular momentum J and parity π . The number of proton(s) and neutron(s) are respectively n_{π} and n_{ν} . One begins by constructing all states $|k\rangle_H$ forming the subspace H. The set of those states is denoted as $\{k_H\}$. The many-body configurations in H can be classified in different families $\{n; j_H^{\pi}\}$ according to their number of nucleons n, total angular momentum j_H , and parity π . States with a number of protons (neutrons) larger than n_{π} (n_{ν}) are not considered since they do not contribute to the many-body states in the composition of subspaces H and P.



FIG. 3. Schematic illustration of the NCGSM/DMRG procedure during the *s*th step of the warmup phase. States $\{k_H\}$ from *H*, previously optimized states α_H , and states $\{(lj)_s\}$ constructed by occupying the *s*th shell with *n* particles are coupled to generate the new set of states $\{k_H \otimes i_P\}^J = \{k_H \otimes \{\alpha_P \otimes (lj)_s^n\}\}^J$.

The matrix elements in H of the suboperators of the NCGSM Hamiltonian \hat{H} expressed in the second quantization form, are calculated and stored:

$$\{O\} = \{a^{\dagger}, (a^{\dagger}\widetilde{a})^{K}, (a^{\dagger}a^{\dagger})^{K}, ((a^{\dagger}a^{\dagger})^{K}\widetilde{a})^{L}, (a^{\dagger}a^{\dagger})^{K}(\widetilde{a}\widetilde{a})^{K}\},$$
(19)

where a^{\dagger} and \tilde{a} are the nucleon creation and annihilation operators in shells forming the subspace *H*. The NCGSM Hamiltonian is then diagonalized in the pole space *H* to provide the zeroth-order approximation $|\Psi\rangle^{(0)}$ to $|\Psi\rangle$.

In the following stage, the subspace P is built, step by step, by adding scattering shells one by one during the so-called warmup phase. At each step, many-body states constructed within the new added shell are coupled to optimized many-body states constructed during previous steps, i.e., constructed within previously added shells. Moreover, the matrix elements (19) of the suboperators acting among the optimized states have been stored during previous steps.

To be more specific, let us assume that the *s*th step is reached. The method is illustrated in Fig. 3. The scattering shell $(lj)_s$ belonging to the discretized contour L^+ is added and within this shell, one constructs all possible many-body states $\{(lj)_s^{n_P}\}$. Matrix elements of suboperators (19) acting on $\{(lj)_s^{n_P}\}$ are also computed. One then couples previously optimized states denoted as $|\alpha\rangle_P$ to $\{(lj)_s^{n_P}\}$ to obtain the set of states $\{i_P\} = \{\alpha_P \otimes (lj)_s^{n_P}\}$. States in *H* are then coupled with the states $|i_P\rangle$ to construct the set $\{k_H \otimes i_P\}^J$ of states coupled to J^{π} that constitutes a basis in which the NCGSM Hamiltonian is diagonalized. The Hamiltonian matrix is constructed in this set with the Wigner-Eckart theorem and the matrix elements of the suboperators (19) acting on $\{k_H\}, \{\alpha_P\},$ and $\{(lj)_s^{n_P}\}$.

The target state $|\Psi\rangle$ is selected among the eigenstates of \hat{H} as the one having the largest overlap with the reference vector

 $|\Psi\rangle^{(0)}$. Based on the expansion

$$|\Psi\rangle = \sum_{k_H, i_P} c_{i_P(j_P)}^{k_H(j_H)} \{|k_H(j_H)\rangle \otimes |i_P(j_P)\rangle\}^J, \qquad (20)$$

by summing over the reference subspace H for a *fixed* value of j_P , one defines the reduced density matrix [71]:

$$\rho_{i_P i'_P}^P(j_P) \equiv \sum_{k_H} c_{i_P(j_P)}^{k_H(j_H)} c_{i'_P(j_P)}^{k_H(j_H)}.$$
(21)

Truncation in the subspace *P* is dictated by the density matrix. In the case of a Hermitian Hamiltonian, the eigenvalues of the density matrix are real and one can show that the truncation in *P* is optimal when one keeps the eigenstates of the density matrix with the largest eigenvalues [67]. More specifically, the error in the representation of $|\Psi\rangle$ (20) *after truncation*, is minimal in that case. In that sense, the eigenstates of the density matrix are optimal.

Within the metric defining the Berggren ensemble, the NCGSM density matrix is complex symmetric and its eigenvalues are, in general, complex. The trace of the density matrix being equal to one, the truncation is done by keeping eigenstates of the density matrix with the corresponding eigenvalue w_{α} such that the condition

$$\left|1 - \operatorname{Re}\left(\sum_{\alpha=1}^{N_{\rho}} w_{\alpha}\right)\right| < \epsilon \tag{22}$$

is satisfied. The quantity ϵ in Eq. (22) can be viewed as the truncation error of the reduced density matrix. The smaller ϵ , the larger number of eigenvectors must be kept. In particular, for $\epsilon = 0$, all eigenvectors with nonzero eigenvalues are retained.

One then keeps eigenstates of the density matrix according to Eq. (22). These are expressed as linear combination of the vectors $|i\rangle_P$ in P and all matrix elements of the suboperators in these optimized states are recalculated and stored. Note that at each step, we enforce that at least one state in each family $\{n; j_P^P\}$ is kept [70].

The warmup phase continues by having the *P* subspace grow by adding scattering shells one by one until the last shell is reached, providing a first guess for the wave function of the system in the whole ensemble of shells. At this point, all s.p. states have been considered, and all suboperators of the Hamiltonian \hat{H} acting on states saved after truncation in *P* have been computed and stored. The warmup phase ends and the so-called sweeping phase begins.

Starting from the last scattering shell $(lj)_{last}$, the procedure continues in the reverse direction (the sweep-down phase) using the previously stored information. At this stage, the truncations are done according to the density matrix, which is obtained by summing over states $|k_H\rangle$ of the reference subspace *H* and the states $|i_{prev}\rangle$ generated in the warmup phase. Scattering shells are added one at a time and at the last step of the sweep-down phase, the first scattering shell is reached. The procedure is then reversed and a sweep in the upward direction (the sweep-up phase) begins. Using the information previously stored, a first shell is added, then a second one, etc. The sweeping sequences continue until

TABLE I. Convergence of the NCGSM eigenvalues with respect to the $N_{\text{max}} = 2n_{\text{max}} + \ell_{\text{max}}$ of the HO expansion of the NN interaction. Values are in MeV. The length parameter of HO states is b = 1.5 fm. For this value of the *b* parameter, the CoM wave function is a Gaussian.

N _{max}	Energy
5	-5.321
7	-5.334
9	-5.336
11	-5.343
13	-5.343
15	-5.343
17	-5.346
19	-5.349
21	-5.352
23	-5.352
25	-5.352
27	-5.352
29	-5.352

convergence for the target eigenvalue is achieved. For more details, see Ref. [70].

V. APPLICATIONS OF THE NCGSM/DMRG

A. Test of convergence with respect to N_{max} of the two-body interaction in HO expansion

In Table I we check the energy convergence of the ³He nucleus with respect of the N_{max} parameter in the expansion of the two-body interaction. We choose ³He due to the existence of both nuclear and Coulomb parts in the NCGSM Hamiltonian. The interaction employed is the N³LO renormalized at a cutoff $\Lambda = 1.9$ fm⁻¹, and we used a s.p. Berggren basis consisting only of $s_{1/2}$, $p_{1/2}$, and $p_{3/2}$ orbitals, for both neutrons and protons. The $0s_{1/2}$ states are bound and the scattering contours lie on the real momentum axis. They are discretized with 20 points each and they extend up to 4 fm^{-1} . At this point, our purpose is not to describe realistically ³He but rather check the convergence of the many-body result with respect to a number of HO states of the expansion of the realistic interaction. In Table I we see that there is an overall small variation of the energy with increasing n_{max} (or N_{max}). The energy changes only by ~ 30 keV by changing N_{max} from $N_{\text{max}} = 5$ to $N_{\text{max}} = 29$. For $N_{\text{max}} = 21$ ($n_{\text{max}} = 10$), the energy shows the convergence and in all the following we will adopt $n_{\rm max} = 10$ for our calculations.

B. Convergence with respect to the angular momentum of the model space and with respect to the number of sweeps in the DMRG method

In this section, we are applying the NCGSM for ³H nucleus to test the DMRG method for solving the many-body Schrödinger equation in Berggren basis. ³H (and also ⁴He or ³He) is a well-bound system so results of the NCGSM/DMRG approach can be compared against other well-known



FIG. 4. (Color online) ³H results as a function of the (maximum) angular momentum, in the interval from $\ell = 0$ to $\ell = 4$.

bound-states methods. The numerical task in solving a three nucleon system is relatively easy so that one can check the convergence of NCGSM/DMRG results with respect to the maximal angular momentum of the model space and the truncation error of the reduced density matrix.

In Fig. 4, we compare NCGSM results obtained with the Argonne v_{18} against Faddeev calculations. We perform a calculation using $s_{1/2}$, $p_{3/2}$, $p_{1/2}$, $d_{3/2}$, $d_{5/2}$, $f_{7/2}$, $f_{5/2}$, $g_{9/2}$, $g_{7/2}$ partial waves for protons and neutrons. The basis generating potential is the GHF which gives the $0s_{1/2}$ proton and neutron states bound, with energies -10.417 MeV and -11.982 MeV along the imaginary momentum axis. For this reason, the scattering continua $i\{s_{1/2}\}, i\{p_{3/2}\}, i\{p_{1/2}\}$ are chosen along the real axis and each of them is discretized with 18 points. Here *i* denotes the number of discretization points, which ranges from 1 to 18 for the s wave and from 0 to 18 for the p waves. For the d waves we choose a discrete HO basis with b = 1.5 fm. In particular, we use five HO states for the $d_{3/2}$ and $d_{5/2}$ states, which from now on we denote as 5d. The physical argument behind this choice lies in the fact that for $\ell > 1$ the centrifugal barrier is large enough to confine the s.p. states and, hence, the HO basis becomes a realistic alternative. For fand g partial waves we used three HO states with b = 1.5 fm for both protons and neutrons. In total, the s.p. basis consisted of 154 partial waves. The result we obtain is

$$E_{\rm NCGSM} = -8.39 \,\,{\rm MeV},$$

whereas the Faddeev result [72] is

$$E_{\text{Fadeev}} = -8.47 \text{ MeV}.$$

In Fig. 4, we show also an exponential fit of the NCGSM results for different ℓ of the s.p. basis. The extrapolated result is

$$E_{\text{extrp}} = -8.449 \pm 0.087 \text{ MeV},$$

where the fit function is: $E = E_{\text{extrp}} + b \times e^{-c \times \ell}$. We see that inclusion of partial waves with angular momentum larger than $\ell = 4$, should have a very small contribution, of the order of

50 keV, to the many-body result. The ground state (g.s.) of ³H being well bound, the set of HO shells can also provide a basis in which the many-body Schrödinger equation can be solved. But nevertheless in our case, the $\ell = 0$, 1 shells are solutions of the HF potential, which admits two bound $s_{1/2}$ shells and a continuum set of shells. What we show in Fig. 4 serves as an illustration of the rate of convergence of the energy, for increasing the angular momentum of our basis states, and our aim is not to deduce any physical insight by using our Berggren basis on triton (and later on ⁴He). For three-particles systems we can perform an exact diagonalization using the Lanczos method and, therefore, we can test the precision of the NCGSM/DMRG approach. Since details of the DMRG algorithm can be found in Ref. [70], we only mention the basic ingredients of the DMRG calculation.

It is important to consider in the reference space H, not only resonant states, but also some nonresonant shells from the P space, in order to generate all possible many-body configurations in the warmup phase, which these shells could generate [70]. In addition to the $0s_{1/2}$ neutron-proton resonant states, the space H contained also the last $d_{5/2}$ and $d_{3/2}$ HO states. The choice of the shell to be included in the pole space H is arbitrary and the results do not depend on which shell is considered. The mixture of positive and negative parity states assures that we will not miss any couplings in the warmup phase. In the space P the shells are ordered as follows: $\{is_{1/2}, ip_{3/2}, ip_{1/2}, id_{5/2}, id_{3/2}, \ldots\}$, where *i* denotes the scattering shell starting from 0. In the case of the *d* states the index *i* (*i* = 0, 1, ..., 4) denotes the HO radial quantum number.

Usually in the DMRG applications, it is decided from the beginning how many states of the density matrix that have the largest eigenvalue will be kept. This number is then kept fixed throughout the whole iterative process. In this work we will use the truncation scheme defined in Eq. (22). The smaller the ϵ , the more vectors are kept. Usually with $\epsilon = 10^{-8}$ the exact result is reproduced. However, even for a larger ϵ (10⁻⁷ or 10⁻⁶) the agreement is very satisfactory, especially if the DMRG calculations are followed for more than two sweeps (see also Fig. 5). This method is called the dynamical block selection approach because the number of states N_{ρ} kept changes during the iterative process to satisfy the truncation limit (22). In Fig. 5, we show the iterative DMRG process that includes partial waves up to $\ell = 2$. The number of steps starts from zero, denoting the first shell in the sweep-down phase. As discussed earlier, each step corresponds to the addition of a new s.p. shell. We notice a periodic pattern of pronounced oscillations in energy, that appear in the middle of each sweep, with their amplitude continuously decreasing with the addition of more shells until the convergence is reached. Finally, the energy has converged in the end of the third sweep. The truncation error for this calculations was $\epsilon = 10^{-7}$ resulting in a maximum number of vectors kept $N_{\rho} \sim 85$ and a maximum dimension $D_{\rm max} \sim 1200$ of a matrix to be diagonalized. For a direct diagonalization in this basis, the maximum dimension would be 387 998 in m scheme and 96 883 in J scheme. The exact result of the diagonalization is

$$E_{\text{exact}} = -7.840 \text{ MeV},$$



FIG. 5. (Color online) Iterative process of the DMRG approach for $\epsilon = 10^{-7}$ and including only waves with $\ell = 2 (s - pd)$. Both Lanczos and DMRG diagonalizations of the Hamiltonian are shown.

and the result obtained by the DMRG is

$$E_{\epsilon=10^{-7}} = -7.832$$
 MeV.

In the following, using the V_{low-k} Argonne v_{18} potential, we will test the convergence of the DMRG method with respect to the truncation parameter ϵ , which controls how many vectors of the density matrix are kept at each step. The results are gathered in Figs. 6 and 7. Additionally, in Table II we show the number of vectors that are kept for different values of the parameter ϵ and the corresponding energy.

Each point in Figs. 6 and 7 corresponds to the value at the end of the fourth sweep of the DMRG process. The error bar reflects the extremum values of the energy in the last sweep (see also Fig. 8). We observe that with decreasing the value of ϵ



FIG. 6. (Color online) Convergence of ³H ground-state energy as a function of the truncation error ϵ of the density matrix (22). The s.p. basis consisted of all states up to $\ell = 4$. (a) corresponds to a fit with four points, (b) with five, (c) with six, and (d) with nine points. The fit function has the form: $E = E_{\text{extrp}} + b \times \epsilon^c$. For more details, see the description in the text.



FIG. 7. (Color online) Same as in Fig. 6 but omitting the lowprecision $\epsilon = 10^{-3}$ point.

the error bar also decreases and almost vanishes for precisions better than 10^{-5} , i.e., the DMRG result quickly converges to the exact result for ϵ smaller than 10^{-5} .

The purpose of this exercise is to test the extrapolation properties of the NCGSM/DMRG calculations. In general, one would like to perform calculations with the best precision possible, i.e., $\epsilon \sim 10^{-8}$, what is presently possible only in the lightest systems. Extending the NCGSM applications to somewhat heavier systems, one needs to choose a different strategy that makes the calculation feasible and, at the same time, gives a rather precise result. The ³H studies serve here as a testing ground for this investigation.

In Fig. 6 we show calculations with ϵ ranging from 10^{-3} up to 10^{-8} . Figure 6(a) consists of four points that correspond to several values of ϵ from 10^{-3} up to 10^{-4} . The result is almost 600 keV away from the exact one and the extrapolation is also of a poor quality, lying at ~380 keV away. The function we use for fitting/extrapolating numerical data has the form:

$$E = E_{\text{extrp}} + b \times \epsilon^{2}$$

Adding one more point, the situation is improved, but still the exact result is off the error bars of the extrapolation [Fig. 6(b)]. In Fig. 6(c) the exact result is in between the error bars of the

TABLE II. Vectors of the density matrix that are kept for different values of ϵ . The corresponding matrix dimensions and many-body energies for ³H are also shown.

ε	No. of vectors	dimension	energy (MeV)
10 ⁻³	5	43	-5.648
$5.0 imes 10^{-4}$	8	99	-6.878
2.5×10^{-4}	13	109	-7.396
10^{-4}	22	173	-7.821
5.0×10^{-5}	28	308	-8.042
10^{-5}	46	600	-8.287
10^{-6}	73	1075	-8.357
10^{-7}	96	1909	-8.381
10^{-8}	117	2575	-8.388



FIG. 8. (Color online) Energy variation with respect to N_{step} of the DMRG calculation for ³H and for several ϵ values. The iteration is during the fourth sweep.

extrapolation and considering all values up to 10^{-8} we fall onto the exact result.

The outcome is that carrying calculations with a relatively low precision, one can find the exact result within the error bars of the extrapolated value. In this specific example, the maximum dimension for $\epsilon = 10^{-5}$ [Fig. 6(c)] is $D_{\text{max}} = 600$ (see Table II), whereas the dimension in a direct diagonalization is 890 021 in *m* scheme and 123 835 in *J* scheme.

In Fig. 7 we make the same analysis but neglecting the energy point corresponding to $\epsilon = 10^{-3}$. This energy is a rather poor approximation of a final result, lying almost 3 MeV away. By extrapolating, we retrieve the exact result even in the case [Fig. 7(a)], where energies that correspond to ϵ up to 10^{-4} were used in the fit. For this value of ϵ , only $N_{opt} = 22$ vectors of the density matrix were kept, resulting in a maximum dimension of the matrix to be diagonalized $D_{max} = 173$, which is much smaller than the dimension in a full diagonalization. Notice that the extrapolated value in Fig. 7(a) is not accompanied by an error estimate. This is because we are fitting here three points with a function that is characterized by three parameters and by definition the χ^2 fit produces a zero error.

In addition, in Fig. 8 we show the behavior of the energy during the fourth sweep in the DMRG process. Due to the small number of vectors kept, calculations that correspond to truncation parameters 10^{-3} , 5.0×10^{-4} and to a lesser extent for 2.5×10^{-4} , show variations with respect to the number of shells added (N_{step}). The error bars in Figs. 6 and 7 were calculated by the extremum values of these variations. For ϵ values equal 10^{-4} and smaller, the energy is a flat curve with variations of the order of less than 1 keV.

Finally, we have calculated the expectation value of the CoM operator [Eq. (17)]. In the largest model space that we used, the expectation value of the H_{cm} is approximately 7 keV. The $\hbar\omega$ energy in this case is 18.5 MeV (b = 1.5 fm). Following the assertion of Ref. [66], we conclude that in a sufficiently large model space the NCGSM wave function factorizes and the CoM wave function $|\psi_{CoM}\rangle$ is a Gaussian with $\hbar\omega = 18.5$ MeV.

The smooth convergence properties of the NCGSM/DMRG procedure both with the number of partial waves and the truncation error, as they were tested in ³H and ³He, will be used later to calculate somewhat heavier nuclei. One can perform several smaller scale NCGSM calculations, changing both the number of density matrix vectors and the number of partial waves, and extrapolate these data to retrieve the exact result within the error bars.

C. ⁴He nucleus

In this section we apply the NCGSM to ⁴He nucleus, using the DMRG as the diagonalization technique. We will compare the NCGSM/DMRG results with FY calculations, using the Argonne v_{18} interaction with a $V_{\text{low-}k}$ cutoff $\Lambda = 1.9 \text{ fm}^{-1}$. The energies of the $0s_{1/2}$ neutron and proton as they are calculated from the GHF process are -26.290 MeV and -24.453 MeV, respectively. As in the case of ³H, the s and p scattering continua are taken along the real axis and are discretized with eighteen points. The contours extent up to 4 fm^{-1} . For the remaining d, f, g partial waves, we assume HO basis functions with b = 1.5 fm. We take 5d, 3f, and 3g states for protons and neutrons. For a truncation error $\epsilon = 10^{-6}$, the maximum number of vectors, which are kept is $N_{\text{opt}} = 180$, resulting in a Hamiltonian matrix of dimension $D_{\text{max}} \sim 6000$. On the other hand, the dimension of the Hamiltonian matrix for a direct Lanczos diagonalization is computed to be 119 864 088 in m scheme and 6 230 512 in J scheme.

In Fig. 9 we show the iteration pattern for 4 He, which is similar to the one in 3 H case, but we observe that already in the middle of the third sweep the energy is converged. This can be attributed to the larger number of vectors of the density matrix that are kept. The converged energy is

$$E_{\rm NCGSM} = -29.15 \,\,{\rm MeV},$$

and the FY result [72] is

$$E_{\rm FY} = -29.19$$
 MeV.



FIG. 9. (Color online) Comparison of the NCGSM with the FY result with the $V_{\text{low-}k}$ Argonne v_{18} interaction.



FIG. 10. (Color online) Same as in Fig. 9 but with the $V_{\text{low-}k} \text{ N}^3\text{LO}$ interaction.

Results of ³H and ⁴He using the Argonne v_{18} are in a nice agreement with both FY and CC calculations with triples corrections [73].

We also performed calculations using the chiral N³LO interaction ($\Lambda = 1.9 \text{ fm}^{-1}$). The s.p. energies for the 0s_{1/2} neutron and proton poles are -24.333 MeV and -24.303 MeV, respectively. The first point in the number of steps corresponds to approximately fortieth shell in the sweep-down phase. The results are shown in Fig. 10, and the converged energy:

$E_{\rm NCGSM} = -27.48$ MeV.

The experimental total binding energies are found in the 2003 Atomic Mass Evaluation II (AME) [74]. The difference with the experimental binding energy is attributed to the missing 3N interactions, which are both bare and induced. As bare we denote the 3N forces that appear due to the neglect of the quark degrees of freedom. The chiral potential we use has a certain cutoff, beyond which the missing physics is integrated out and this results in the appearance of many-body forces [4,75–78]. As induced we denote the 3N forces that are related to the renormalization technique [51,52,56].

D. ⁵He nucleus

⁵He is a challenge for any many-body theory due to its unbound character. In particular, both the ground and first excited states are many-body resonances, which obey outgoing asymptotics. Because of these characteristics, the complex energy formulation of the NCGSM using the Berggren ensemble is suitable for its description. Indeed, in our formalism the resonance parameters (g.s energy with respect to ⁴He and the width) will be identified as the eigenvalues of the complexsymmetric Hamiltonian matrix. The position of the resonance will then be the real part of the energy, while the imaginary part is related to the width by the formula: $\Gamma = -2Im(E)$. An advantage of using Berggren's states is that the complex eigenvalues (*E*) are the ones, which correspond to states with purely outgoing-wave solutions. Contrary to the previous applications, where we used a real-energy basis, in the case of



FIG. 11. (Color online) Same as in Fig. 10 but for the real part of ⁵He g.s. energy.

⁵He we are employing a complex basis, which also includes the $0p_{3/2}$ resonant state. Overall we include the bound $0s_{1/2}$ neutron state with a s.p. energy of -23.290 MeV, the bound $0s_{1/2}$ proton state with s.p. energy -23.999 MeV and the $0p_{3/2}$ s.p. resonance with a real part of energy 1.193 MeV and a width 1267 keV. Its position in the complex-energy plane is: k = (0.277, -0.068) fm⁻¹. The s.p. basis for protons and neutrons is produced by a GHF calculation using the N³LO V_{low-k} interaction with $\Lambda = 1.9$ fm⁻¹. The $p_{3/2}$ contour is taken complex to satisfy the Berggren completeness relation (11), whereas the $s_{1/2}$ and $p_{1/2}$ contours may be chosen along the real-k axis. For states with $\ell > 1$, we assume the HO basis functions (5d, 3f, 3g) as we described in the previous cases.

In Fig. 11 we show the DMRG convergence pattern of the real part of the g.s. energy in ⁵He. The calculation in Fig. 11 is presented starting from the fortieth shell in the sweep-down phase. The converged energy:

$$\Re e \left(E_{\text{NCGSM}} \right) = -26.31 \text{ MeV},$$

lies at about 1 MeV above the experimental total binding energy [74]. The truncation error in this calculation is $\epsilon = 10^{-6}$ and the maximum number of vectors we kept is $N_{\rm opt} \sim 300$. The corresponding dimension of the matrix is $D_{\rm max} \sim 10^5$, whereas in the direct diagonalization one deals with a matrix of a dimension $\sim 3 \times 10^9$.

The imaginary part of the 5 He g.s. energy is shown in Fig. 12. The converged value is:

$$\Im m(E_{\text{NCGSM}}) = -0.2 \text{ MeV},$$

i.e., $\Gamma_{\text{NCGSM}} = 400$ keV. Having calculated the g.s. total binding energies of ⁴He and ⁵He, we obtain then the position of the resonance at an energy 1.17 MeV above the $\alpha + n$ threshold with a width $\Gamma = 400$ keV.

We will compare this result with the value extracted from experiment. Experimentally, resonance parameters can be extracted in two ways [79]. First, one could apply the conventional R-matrix approach on the real axis and use the Lane and Thomas prescription for the extraction of positions and widths of a state [80]. The second way is the extended



FIG. 12. (Color online) Same as in Fig. 11 but for the imaginary part of 5 He g.s. energy.

R-matrix approach [81, 82], which associates the resonance parameters with the complex poles of the S matrix. According to Refs. [79,83], this is the recommended prescription, and it is also the most appropriate for comparison to the NCGSM results, since our complex eigenvalues correspond, indeed, to the location of the poles of the S matrix, with the correct asymptotic behavior. The extended *R*-matrix approach gives the position of the $(3/2)^{-}$ state at an energy 798 keV above the α + n threshold and with a width Γ = 648 keV. The difference we observe between our results and the experimental results is attributed to the specific interaction that we employ and/or to missing many-body forces. We know already that the position of such unbound states, which have a non Breit-Wigner character (broad states) and are close to threshold, heavily depends on the specific characteristics of the NN (or NNN) interaction. This was shown for calculations of ⁵He nucleus, in the many-body frameworks of the NCSM merged with the resonating group method (RGM) [40,84,85], (see also Ref. [42] for a recent application of the NCSM/RGM with 3N forces) and also in the Green's function Monte Carlo approach [46], with a $V_{low-k}\Lambda = 2.1 \text{ fm}^{-1}$ Argonne v_{18} and a SRG $\lambda = 2.02 \text{ fm}^{-1} \text{ N}^3 \text{LO}$ for the NCSM/RGM and a bare Argonne v_{18} with Urbana/Illinois 3N forces in the GFMC case.

We verify this assertion also in our NCGSM calculations. For that purpose, we have compared NCGSM results for two different renormalization scale parameters: $\Lambda = 1.9 \text{ fm}^{-1}$ and 2.1 fm⁻¹. To this end we have performed the NCGSM calculation using the Davidson (variant of Lanczos) method for the diagonalization of the many-body Hamiltonian [34] and truncating the space of configurations to up to 4p-4h excitations. In addition, we have used a smaller s.p. basis by taking only the neutron $p_{3/2}$ states as Berggren states, whereas remaining states up to $\ell = 4$ are the HO basis functions with a HO length b = 1.5 fm. The many-body energy and width of ⁴He/⁵He in this NCGSM/4*p*4*h* calculation are close to the exact NCGSM/DMRG results. Using the V_{low-k} N³LO force with $\Lambda = 1.9$ fm⁻¹, we obtain $E_{\text{NCGSM}}^{(4p4h)} = -27.386$ MeV for the total binding ⁴He, and $\Re(E_{\text{NCGSM}}^{(4p4h)}) = -25.825$ MeV for

TABLE III. NCGSM_{DMRG} result as compared to experimental position and width of the ⁵He g.s. Energies are with respect to the α + n threshold.

Energy (MeV)	Γ (MeV)
1.17	0.400
0.798	0.648
0.963	0.985
0.771	0.644
0.890	0.651
0.79	0.525
	Energy (MeV) 1.17 0.798 0.963 0.771 0.890 0.79

^aIn the data evaluation of [87] the half life $T_{1/2}$ was given to be 700 ys, where ys stands for yoctosecond and equals 10^{-24} s. Then the width was obtained by the relation: $\Gamma_{\rm cm} T_{1/2} \approx \hbar \ln 2$, and $\Gamma_{\rm cm}$ is the level total width.

⁵He with a width $\Gamma_{\text{NCGSM}}^{(4p4h)} = 370$ keV. The ⁵He g.s. is found at an energy 1.56 MeV above the $\alpha + n$ threshold, with a width $\Gamma = 370$ keV. In the case of $\Lambda = 2.1$ fm⁻¹ renormalization parameter the total binding energy is $E_{\text{NCGSM}}^{(4p4h)} = -26.060$ MeV for ⁴He and $\Re(E_{\text{NCGSM}}^{(4p4h)}) = -23.903$ MeV for ⁵He with a width $\Gamma = 591$ keV. Hence, for $\Lambda = 2.1$ fm⁻¹ the g.s. of ⁵He is found at a position 2.15 MeV above the $\alpha + n$ threshold and with a width $\Gamma = 591$ keV. As expected, we observe that the unbound state of ⁵He is very sensitive on the choice of the NN interaction.

It needs to be mentioned that, because of the broad nature of the ⁵He g.s. and the proximity to the threshold, results would also depend on the method used to extract such a state from the experimental data. We find that if one uses the conventional *R*-matrix approach on the real axis, the position of the g.s. is at 0.963 MeV above the α + n threshold, with a width Γ = 985 MeV and both values are different by about 200 keV as compared to the extended *R*-matrix estimations. The *R*-matrix estimation of Ref. [86] is similar to the extended *R*-matrix method, since the authors determined the resonance energy and width by finding the *S*-matrix pole location implied by the fit parameters. In Table III we gather the existing measurements and/or the extracted values from the data of the position and width of ⁵He g.s.

E. Asymptotic normalization coefficient of a complex-energy ground state of ⁵He

Asymptotic normalization coefficients (ANCs), as the name implies, define the overall normalization of the tail of the overlap function between a system with A and A \pm N nucleons. In our case, we are interested in the addition of a neutron to ⁴He, so A = 4 and N = +1. Because of their definition, they have attracted a lot of attention, since the properties of the wave-function tail can be useful to test many-body methods. Additionally, their importance is manifested in the borderline between nuclear physics and astrophysics, where for example their knowledge is necessary for the correct description of neutrinos reaction rates [89,90]. Contrary to spectroscopic factors (SFs), ANCs are observables, due to their invariance under short-range unitary transformations. This can be understood intuitively because of their connection to the tail of the overlap function. Indeed, for a truly unitary transformation (such as V_{low-k} or SRG), where generated many-body terms will not be neglected, what will change is the short-range part of the potential and the interior region of the wave function. The tails of both potential and wave function, however, will remain unchanged. Hence, the ANC is a quantity that does not feel the changes of the short-range physics (see Ref. [91], discussion in Ref. [92] about observables and nonobservables and also Ref. [56] for an overview of unitary transformations and their impact on nuclear structure operators). Experimentally, ANCs have been measured for s-shell and p-shell nuclei and for a detailed list of references we refer the reader to Ref. [93]. Recently, theoretical calculations of ANCs have been performed in the framework of the HH approach [94], the GSM and SMEC [95], the GFMC [47,93,96], and also Ref. [97].

One of the criteria that should be met in order to calculate in a meaningful manner the ANC, is the correct asymptotic behavior of wave function. In the NCGSM, using the Berggren ensemble as basis, this condition is met for either bound, weakly bound or unbound states. The other condition is related to the separation energy S_{1n} . Indeed, the tail of the overlap is very sensitive on the S_{1n} and small differences in the S_{1n} for different Hamiltonians can cause large differences on the ANCs. This can be especially complicated for *ab initio* calculations, where there are no adjustable parameters. Recently, a method was proposed to alleviate this problem in the GFMC approach, where calculations could be performed using experimental separation energies [47,93].

Our approach to calculate the ANC of the reaction ${}^{4}\text{He} + n \rightarrow {}^{5}\text{He}$ is based on the calculation of the overlap:

$$I_{\ell j}(r) = \frac{1}{\sqrt{2J_a + 1}} \sum_{\mathcal{B}} \left\langle \widetilde{\Psi_A^{J_a}} \right| |a_{\ell j}^{\dagger}(\mathcal{B})| \left| \Psi_{A-1}^{J_{A-1}} \right\rangle \langle r \ell j | u_{\mathcal{B}} \rangle, \quad (23)$$

where the sum runs over the complete set of basis states \mathcal{B} . In our case, $a_{\ell j}^{\dagger}(\mathcal{B})$ creates neutron in any $p_{3/2}$ s.p. state of the Berggren basis and $\langle r \ell j | u_{\mathcal{B}} \rangle$ is the $p_{3/2}$ radial wave function, so the radial overlap integral takes the form:

$$I_{\ell=1, j=3/2}(r) = \frac{1}{2} \sum_{n} \langle {}^{5}\widetilde{\mathrm{He}}_{3/2^{-}} || a_{n,\ell=1, j=3/2}^{\dagger} || {}^{4}\mathrm{He}_{0^{+}} \rangle u_{n,\ell=1, j=3/2}(r).$$
(24)

Regarding the details of the calculation, we used the Davidson method to diagonalize the Hamiltonian with a $V_{low-k}\Lambda = 1.9 \text{ fm}^{-1}$ potential and the model space was the same as the one at the end of Sec. V D. We also performed calculations at a cutoff $\Lambda = 2.1 \text{ fm}^{-1}$.

The radial overlap integral for ${}^{4}\text{He}_{0^{+}} + n \rightarrow {}^{5}\text{He}_{3/2^{-}}$ is shown in Fig. 13 for $\Lambda = 1.9 \text{ fm}^{-1}$. By fitting real and imaginary parts of the radial overlap integral in the asymptotic region with a Hankel function, one can extract the ANC, which in this case equals 0.197 fm^{-1/2}, whereas for $\Lambda = 2.1 \text{ fm}^{-1}$ equals 0.255 fm^{-1/2}. Real and imaginary parts of the radial overlap integrals for $\Lambda = 1.9 \text{ fm}^{-1}$ and 2.1 fm⁻¹ are compared in Fig. 14. We immediately see that the



FIG. 13. (Color online) The radial overlap integral for ${}^{4}\text{He}_{0^{+}} + n \rightarrow {}^{5}\text{He}_{3/2^{-}}$. The asymptotic region is fitted by the Hankel function. See the text for more details.

two different Hamiltonians we employed produced different separation energies, the overlaps have different asymptotic behavior and the ANCs are different. Thus, ANC calculations are meaningful only in the case that the separation energy is under control, which is not true in our case. Of course, the reason for this behavior is that the scheme we used to alter the short-range physics is not a unitary transformation, and additionally, we neglected a bare 3N interaction before renormalization. Unitarity was violated by the neglect of many-body forces, which yield different separation energies and ANCs. Nevertheless, calculations of ANCs can be used as a consistency test for the many-body method proposed, as we will see in the following discussion. Indeed, the width and ANC are related via the relation (see Refs. [47,95] and references cited therein)

$$C = \sqrt{\frac{\Gamma\mu}{\hbar^2 \Re \mathbf{e}(k)}},\tag{25}$$



FIG. 14. (Color online) Real and imaginary parts of radial overlap integrals for two different V_{low-k} cutoff scales Λ .

which is mainly valid for narrow resonances, even though in our case we are describing a relatively broad state. This formula expresses actually the outgoing flux probability for the decaying (outgoing-wave) ⁵He g.s. In this relation, Cdenotes the ANC, μ is the effective mass (1.25 in our case) and k is the linear momentum associated with the one-neutron separation energy of ⁵He. We then find a width $\Gamma_{ANC} =$ 311 keV and $\Gamma_{ANC} = 570$ for $\Lambda = 1.9$ fm⁻¹ and 2.1 fm⁻¹ respectively, which is consistent with the width obtained in NCGSM/DMRG (or NCGSM/4p4h). As we said, relation (25) is mainly valid for narrow resonances, which is known for calculations on the real axis [98] and it was also proved in Ref. [95] for an unbound state in the complex plane. The approximation is that the real part of the linear momentum $k = \sqrt{\frac{2mS_{1n}}{\hbar^2}}$ is considered. However, for a complex state as in the case of 5 He, we have:

$$k \sim \sqrt{S_{1n} \left(1 - i \frac{\Gamma}{2S_{1n}} \right)},\tag{26}$$

where S_{1n} stands for the one-neutron separation energy. The condition then, for a real number linear momentum, is that of

$$\frac{\Gamma}{2S_{1n}} \to 0. \tag{27}$$

We see that the notion of a narrow resonance, does not imply $\Gamma \rightarrow 0$, but it is actually the value the width Γ has, with respect to the separation energy, that has to be small. In our calculations, this quantity is in the range of about 10% to 15% for the two renormalization parameters we considered and this error is compatible with the comparison between the width from the NCGSM diagonalization and its extraction from the ANC formula (25).

1. $p_{3/2}$ spectroscopic factor in the ground state of ⁵He

Spectroscopic factors are solely a theoretical invention and should not be interpreted as a measurable quantity. The spectroscopic factor is given by the real part of the norm S^2 of the radial overlap integral:

$$S^{2} = \sum_{\mathcal{B}} \left\langle \widetilde{\Psi_{A}^{J_{A}}} \middle| |a_{\ell j}^{+}(\mathcal{B})| \middle| \Psi_{A-1}^{J_{A-1}} \right\rangle^{2}.$$
(28)

The overlap is always a representation-dependent quantity, since it involves the interior of the wave function, which will change accordingly depending on the scheme the practitioner will use. As scheme or representation we mean here the renormalization of the short-range physics, namely the NN potential. Indeed, one has the freedom to change the off-shell behavior of the potential in any possible manner, while maintaining the NN phase shifts (phase shift equivalent potentials). Each method of changing the off-shell behavior is then considered as a different representation or scheme. Nevertheless, SFs can provide information on shell occupancies and they can be a measure of correlations, i.e., how much from the s.p. picture the nucleus deviates. Such overlaps have been calculated by several methods, either in ab initio or more phenomenological approaches [15,93-96,99-103] (see also discussion on ANCs).

TABLE IV. NCGSM_(4*p*4*h*) results for the *S* dependence on the V_{low-k} cutoff Λ with respect to the S_{1n} .

$\Lambda \ { m fm}^{-1}$	S_{1n} (MeV)	S
2.1	-2.15	0.812
1.9	-1.56	0.787
1.5	-1.38	0.774

Using NCGSM/4*p*4*h* solutions for ⁴He/⁵He, one finds that the spectroscopic amplitude of ⁵He corresponding to the channel [⁴He(g.s.) $\otimes p_{3/2}$]^{3/2-} is 0.787 ($S^2 = 0.62$) for $\Lambda = 1.9 \text{ fm}^{-1}$ and 0.812 ($S^2 = 0.66$) for $\Lambda = 2.1 \text{ fm}^{-1}$. At this point we perform a calculation at a cutoff scale $\Lambda =$ 1.5 fm⁻¹. The total binding for ⁴He is -28.670 MeV and for ⁵He -27.285 MeV. We gather our numbers for the three different cutoffs on Table IV.

The SF in the case of $\Lambda = 1.5 \text{ fm}^{-1}$ is also, slightly, reduced. Overall, the SF is reduced when the separation energy approaches the threshold. This behavior confirms the findings of GSM calculations on the anomalous behavior of SFs, for close to threshold states [15]. Such a quenching of SFs was found in coupled-cluster calculations [103] and also in Ref. [102].

VI. CONCLUSIONS AND FUTURE PERSPECTIVES

In this work, we applied the Berggren s.p. ensemble to perform *ab initio* NCGSM/DMRG calculations for selected light nuclei, both well bound and unbound. We used a translational invariant Hamiltonian and benchmarked our results against Faddeev and Faddeev-Yakubovsky calculations for ³H and ⁴He, respectively. We also investigated the extrapolation properties of the NCGSM/DMRG iterative procedure with the number of partial waves and the truncation error. We found that even if a relatively small number of vectors of the density matrix are kept, the NCGSM results can be extrapolated with high accuracy to the exact result. This methodology will be followed for heavier systems, where the matrix dimensions, even with the DMRG algorithm cannot be handled at present.

The NCGSM is a natural choice to calculate unbound nuclei, such as ⁵He. For the description of ⁵He, we employed a complex-energy Berggren basis consisting of bound $0s_{1/2}$ proton/neutron s.p. states, the $0p_{3/2}$ neutron s.p. resonance, and the associated real and complex nonresonant continua. We successfully reproduced the unbound character of this system from first principles using the N³LO chiral potential as the NN interaction. For $\Lambda = 1.9$ fm⁻¹, the calculated neutron separation energy and the neutron emission width of the ⁵He g.s. are in a reasonable agreement with the experimental data. Still the NCGSM binding energies of ⁴He and ⁵He for this interaction are less by about 1.5 MeV than their experimental values.

Truncating the NCGSM configuration space up to 4p-4h excitations, we were also able to calculate the radial overlap integral, the spectroscopic factor and the ANC for ${}^{4}\text{He}_{0^{+}} + n \rightarrow {}^{5}\text{He}_{3/2^{-}}$. The one-neutron emission width of ${}^{5}\text{He}$ associated with this ANC was found to be in agreement with the width obtained by the many-body diagonalization, which is a nice consistency test for our method.

Quantities that are related to the tail of the wave function, such as the ANC and the width, can be sensitive to modifications of the short-range part of the interaction, if the latter is not done in a consistent (i.e., preserving the unitarity) manner. Within the NCGSM we can probe the impact of missing many-body terms, which arise from the renormalization of the short-range physics, or the impact of neglecting many-body terms from the generic nuclear Hamiltonian, on quantities that are characteristic of unbound systems (i.e., widths) and/or quantities such as the ANCs, which are relevant in regions outside the nuclear attraction where the correct asymptotic behavior is important.

The correct asymptotic behavior of the system and the coupling to the continuum plays an important role in the reaction theory [41]. The GSM has been recently generalized for the study of reactions within a coupled channel (CC) GSM framework [61]. In this respect, the NCGSM can provide the realistic wave function for a target nucleus, which will include both many-body correlations and coupling to the continuum.

This work serves as a proof of principle of the application of the Berggren's basis in a NCSM framework. In the near future, we plan to apply the NCGSM supplemented with the DMRG iterative procedure to calculate excited states of ⁴He and ⁵He, heavier weakly bound systems, such as ⁶He, and very exotic systems in the hydrogen isotopic chain. Understanding the role of three-nucleon forces and continuum coupling in light nuclei at the limits of nuclear stability will be important challenges for future NCGSM studies.

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