The harmonic hyperspherical basis for identical particles without permutational symmetry

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Abstract

The hyperspherical harmonic basis is used to describe bound states in an $A$–body system. The approach presented here is based on the representation of the potential energy in terms of hyperspherical harmonic functions. Using this representation, the matrix elements between the basis elements are simple, and the potential energy is presented in a compact form, well suited for numerical implementation. The basis is neither symmetrized nor antisymmetrized, as required in the case of identical particles; however, after the diagonalization of the Hamiltonian matrix, the eigenvectors reflect the symmetries present in it, and the identification of the physical states is possible, as it will be shown in specific cases. We have in mind applications to atomic, molecular, and nuclear few-body systems in which symmetry breaking terms are present in the Hamiltonian; their inclusion is straightforward in the present method. As an example we solve the case of three and four particles interacting through a short-range central interaction and Coulomb potential.

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I. INTRODUCTION

The Harmonic Hyperspherical (HH) method provides a systematic way to construct an expansion basis for a system of $A$ particles. The $N$ Jacobi vectors ($N = A - 1$) are transformed to the hyperradius $\rho$ plus $3N - 1$ hyperangular coordinates which are used to define the HH functions. These functions are the eigenfunctions of the hyperangular part of the Laplacian operator for a given number of particles (see Ref. [1] and references therein).

Applications of the HH method to describe bound states of $A = 3, 4$ nuclei are well documented in the literature (for a recent review see Ref. [2]). In these applications the HH basis elements, extended to spin and isospin degrees of freedom, have been combined in order to construct antisymmetric basis functions. In fact, the HH functions, as normally defined, do not have well defined properties under particle permutation; this results from the selection of a particular ordering of the particles in the definition of the Jacobi coordinates and, as a consequence, of the hyperangular coordinates used to define the HH functions. Changing the ordering of the particles, it is possible to define a new set of Jacobi coordinates and, accordingly, HH functions depending on the hyperangular variables obtained from this new set. To be noticed that the HH functions defined using a particular choice of the Jacobi coordinates form a complete basis.

The HH functions defined in one set of Jacobi coordinates can be transformed to HH functions defined in another set. In this transformation (permutation) the grand angular quantum number $K$, which identifies a subset of HH functions, is conserved. For finite values of $K$, the dimension $N_K$ of this subset is finite, and therefore a finite number of HH functions, having all the same value of $K$, are necessary to describe a HH function having the same value of $K$ but defined in a different Jacobi set. The coefficients of the transformation can be collected in a matrix having the dimension $N_K$ for each number of particles. For $A = 3$ these matrix elements are the Raynal-Revai coefficients [3]. For $A > 3$ the coefficients cannot be given in a close form, and a few methods have been devised for their calculations [4, 5, 6, 7]. The knowledge of these coefficients allows for the construction of basis elements with well defined permutational symmetry. In fact, each subset defined by $K$ is invariant under particle permutation, as a consequence, the constructions of basis elements with that property is performed as linear combinations of HH functions having the same value of $K$. Different schemes to construct hyperspherical functions with an arbitrary
permutational symmetry are given in Refs. [6, 8, 9, 10, 11]. Recently, a procedure for constructing HH functions in terms of a single particle basis has been proposed in Ref. [12].

In problems in which the $A$-body system is composed by identical particles the wave function of the system has to be completely symmetric or antisymmetric in the case of bosons or fermions, respectively. Considering a Hilbert space extended to spin and isospin degrees of freedom, the construction of HH functions having well defined permutational properties allows for a reduction of the large degeneracy of the basis. In general the completely symmetric or antisymmetric basis functions are a small part of the total Hilbert space. However, the difficulties of constructing HH functions with well defined permutational symmetries increase with $A$ and $K$; therefore, the preliminary step of constructing basis functions with well defined permutational symmetry for $A$ particles could be sometimes very difficult to carry out.

In the present paper we investigate a different strategy. We intend to perform the description of a $A$-body system using the HH basis defined on one set of Jacobi coordinates, the reference set, and not having a well defined behaviour under particle permutation. We will loose the advantage of using a reduced part of the total Hilbert space; however, we will gain in simplicity in the calculation of the matrix elements. By including all HH basis elements up to a certain grand angular momentum $K$, the diagonalization of the Hamiltonian matrix will produce eigenvectors reflecting its symmetries. If the Hamiltonian commutes with the group of permutations of $A$ objects, $S_A$, in the case of non degenerated eigenvalues, the eigenvectors will have a well defined permutation symmetry, and can be organized in accordance with the irreducible representations of $S_A$. Therefore, identifying those eigenvectors with the desired symmetry, the corresponding energies can be considered variational estimates. In particular, it will be possible to identify a subset of eigenvectors and eigenvalues corresponding exactly to those that would be obtained performing the preliminary symmetrization of the states. The disadvantage of this method results in the large dimension of the matrices to be diagonalized. However, at present, different techniques are available to treat (at least partially) this problem.

For a system interacting through a two-body potential $V(i, j)$, the potential energy operator results in a sum over pairs. Its matrix elements can be reduced to one term, let us say $V(1, 2)$, times the number of pairs when symmetric or antisymmetric state functions are considered. When HH functions without well defined permutation behaviour are used, the
calculation of the potential energy operator cannot be reduced to the computation of one term. So we have to face the problem of computing the matrix elements of a general term $V(i, j)$ between HH functions defined in the reference set of Jacobi coordinates in which the distance $r_{ij}$ between particles $(i, j)$ has not a simple form.

The calculation of $V(i, j)$ in the reference set of Jacobi coordinates is performed in two steps: (i) first, we use a property of the HH basis which allows to expand a general function of the coordinates $(i, j)$ in terms of a subset of the basis called the potential basis (PB) [1]; (ii) then, as for the case of a generic HH basis element, each PB element is transformed to the HH basis defined in the reference set of Jacobi coordinates. In the case of the PB, the transformation coefficients are known analytically for each value of $K$ and for a general number of particles $A$. In this way, each term $V(i, j)$ of the potential energy can be first expanded in the PB, and then transformed to HH functions defined in the reference set. So, after this procedure, the potential energy will be expressed in terms of HH functions. As we will see, the computation of the matrix elements of the potential energy is now very simple since it results in a combination of integrals of three HH functions. A compact form suitable for a numeric treatment of the problem will be given.

The derivation and implementation of the final expression for the potential energy in the calculation of bound states is the main subject of the present paper. As a simple application, a system of three and four nucleons interacting through a central potential will be analyzed. Different symmetries will appear considering or not the Coulomb interaction between two protons. To be noticed that when antisymmetrized basis functions are used to describe three or four nucleons, the presence of the Coulomb interaction implies that states with total isospin $T = 1/2, 3/2$ (for $A=3$) and $T = 0, 1, 2$ (for $A = 4$) have to be considered. The extension of the Hilbert space to include these terms increases the dimension of the problem resulting comparable to that one in which the antisymmetrization of the basis is not performed. Finally we would like to stress that the present paper is the first step in a program devoted to applications of the HH basis to systems with $A > 4$ interacting through realistic potentials.

The paper is organized as follows, section II is devoted to a brief description of the HH basis. In sections III the expression for the potential energy in terms of HH states are given. In section IV the results for the examples proposed are shown. Section V includes a brief discussion of the results and the perspectives of the present work.
II. THE HARMONIC HYPERSPHERICAL BASIS

In this section we present a brief overview of the properties of the HH basis following Ref.[1]. We start with the following definition of the Jacobi coordinates for an $A$ body system:

$$x_{N-j+1} = \sqrt{\frac{2m_{j+1}M_j}{(m_{j+1} + M_j)m_j}} (r_{j+1} - X_j), \quad j = 1, \ldots, N,$$

(1)

where $m$ is a reference mass, $N = A - 1$ and we have defined

$$M_j = \sum_{i=1}^{j} m_i, \quad X_j = \frac{1}{M_j} \sum_{i=1}^{j} m_i r_i.$$

(2)

Let us note that if all the masses are equal, $m_i = m$, Eq. (1) simplifies to

$$x_{N-j+1} = \sqrt{\frac{2j}{j+1}} (r_{j+1} - X_j), \quad j = 1, \ldots, N.$$

(3)

For a given set of Jacobi coordinates $x_1, \ldots, x_N$, we can introduce the hyperradius $\rho$

$$\rho = \left( \sum_{i=1}^{N} x_i^2 \right)^{1/2} = \left( 2 \sum_{i=1}^{A} (r_i - X)^2 \right)^{1/2} = \left( \frac{2}{A} \sum_{j>i}^{A} (r_j - r_i)^2 \right)^{1/2},$$

(4)

and the hyperangular coordinates $\Omega_N$

$$\Omega_N = (\hat{x}_1, \ldots, \hat{x}_N, \phi_2, \ldots, \phi_N),$$

(5)

with the hyperangles $\phi_i$ defined via

$$\cos \phi_i = \frac{x_i}{\sqrt{x_1^2 + \cdots + x_i^2}}, \quad i = 2, \ldots, N.$$

(6)

The radial components of the Jacobi coordinates can be expressed in terms of the hyperspherical coordinates

$$x_N = \rho \cos \phi_N$$

$$x_{N-1} = \rho \sin \phi_N \cos \phi_{N-1}$$

$$\vdots$$

$$x_i = \rho \sin \phi_N \cdots \sin \phi_i \cos \phi_i$$

$$\vdots$$

$$x_1 = \rho \sin \phi_N \cdots \sin \phi_3 \sin \phi_2.$$
Using the above hyperspherical angles \( \Omega_N \), the surface element becomes

\[
d\Omega_N = \sin \theta_1 d\theta_1 d\varphi_1 \prod_{j=2}^{N} \sin \theta_j d\theta_j d\varphi_j (\cos \phi_j)^2 (\sin \phi_j)^{3j-4} d\phi_j ,
\]

and the Laplacian operator

\[
\Delta = \sum_{i=1}^{N} \nabla_{x_i}^2 = \left( \frac{\partial^2}{\partial \rho^2} + \frac{3N-1}{\rho} \frac{\partial}{\partial \rho} + \frac{\Lambda_N^2(\Omega_N)}{\rho^2} \right),
\]

where the \( \Lambda_N^2(\Omega_N) \) is the generalization of the angular momentum and is called grand angular operator.

The HH functions \( \mathcal{Y}_{[K]}(\Omega_N) \) are the eigenvectors of the grand angular momentum operator

\[
\left( \Lambda_N^2(\Omega_N) + K(K+3N-2) \right) \mathcal{Y}_{[K]}(\Omega_N) = 0.
\]

They can be expressed in terms of the usual harmonic functions \( Y_{lm}(\hat{x}) \) and of the Jacobi polynomials \( P_{\nu}^{a,b}(z) \). In fact, the explicit expression for the HH functions is

\[
\mathcal{Y}_{[K]}(\Omega_N) = \prod_{j=1}^{N} Y_{l_jm_j}(\hat{x}_j) \prod_{j=2}^{N} (j)P_{K_j}^{l_j,K_j-1}(\phi_j),
\]

where \([K]\) stands for the set of quantum numbers \( l_1, \ldots, l_N, m_1, \ldots, m_N, n_2, \ldots, n_N \), the hyperspherical polynomial is

\[
(j)P_{K_j}^{l_j,K_j-1}(\phi_j) = \mathcal{N}_{n_j}^{l_j,K_j}(\cos \phi_j)^{l_j}(\sin \phi_j)^{K_j-1} P_{n_j}^{\nu_j-1,l_j+1/2}(\cos 2\phi_j),
\]

where the \( K_j \) quantum numbers are defined as

\[
K_j = \sum_{i=1}^{j} (l_i + 2n_i), \quad n_1 = 0, \quad K \equiv K_N,
\]

and the normalization factor

\[
\mathcal{N}_{n_j}^{l_j,K_j} = \sqrt{\frac{2\nu_j \Gamma(\nu_j - n_j) n_j!}{\Gamma(\nu_j - n_j - l_j - 1/2) \Gamma(\nu_j + l_j + 3/2)}},
\]

with \( \nu_j = K_j + 3j/2 - 1 \). The quantum number \( K \) is also known as grand angular momentum.

The HH functions are normalized

\[
\int d\Omega_N \left( \mathcal{Y}_{[K]}(\Omega_N) \right)^* \mathcal{Y}_{[K']}(\Omega_N) = \delta_{[K],[K']},
\]
moreover, the HH basis is complete

\[ \sum_{[K]} \left( \mathcal{Y}_{[K]}(\Omega_N) \right)^* \mathcal{Y}_{[K]}(\Omega'_N) = \delta^{3N-1}(\Omega'_N - \Omega_N). \]  

With the above definitions, the HH functions do not have well defined total orbital angular momentum \( L \) and \( z \) projection \( M \). It is possible to construct HH functions having well defined values of \( LM \) by coupling the functions \( Y_{l_j m_j}(\hat{x}_j) \). This can be achieved using different coupling schemes. Accordingly we can define the following HH function

\[ \mathcal{Y}_{[K]}^{LM}(\Omega_N) = \left[ Y_{l_1}(\hat{x}_1) \otimes \ldots \otimes Y_{l_N}(\hat{x}_N) \right]_{LM} \prod_{j=2}^{N} \mathcal{P}_{K_j}^{l_j, K_j-1}(\phi_j), \]  

having well defined values of \( LM \), although the particular coupling scheme is not indicated. The set of quantum numbers \([K]\) now includes the values of \( LM \) plus \( N - 2 \) intermediate \( l \)-values instead of the \( N \) magnetic numbers \( m_j \). When necessary, the explicit coupling scheme of the above HH function will be given.

### A. Potential Basis

If we have a function which depends only on the difference of two particle positions, \( f(r_1 - r_2) \), we can use a subset of the HH’s to expand that function, called the potential basis (PB) \([\text{I}]\). Let’s introduce the Jacobi coordinates such that \( \mathbf{x}_N = r_1 - r_2 \); then the PB subset is defined by the following condition \( (\Omega_{12} \equiv (\hat{x}_N, \phi_N) \) and \( \Omega_N = (\Omega_{N-1}, \hat{x}_N, \phi_N) ) \)

\[ \Lambda_{N-1}^2(\Omega_{N-1}) \mathcal{P}_{2n+l}^{l_m}(\Omega_{12}) = 0, \]  

where \((n,l,m) \equiv (n_N,l_N,m_N)\), and by

\[ \Lambda_{N}^N(\Omega_N) \mathcal{P}_{2n+l}^{l_m}(\Omega_{12}) = -K(K + 3N - 2)\mathcal{P}_{2n+l}^{l_m}(\Omega_{12}), \]  

with \( K = l + 2n \). Thus, the PB is a subset of the HH’s which depends only on \((\hat{x}_N, \phi_N)\) variables, and which is specified by only three quantum numbers \( n, l, m \), instead of the \( 3N - 1 \). The PB basis element has well defined angular momentum \( l \) and projection \( m \). The expression of the PB elements is:

\[ \mathcal{P}_{2n+l}^{l_m}(\Omega_{12}) = Y_{l_m}(\hat{x}_N)(\cos \phi_N)^l P_n^{3(N-1)/2-1, l+1/2}(\cos 2\phi_N)Y_{[0]}(D - 3), \]
where \((D = 3N)\)

\[
Y_{[0]}(D - 3) = \left[ \frac{\Gamma[(D - 3)/2]}{2\pi^{(D-3)/2}} \right]^{1/2}
\]

is the normalization verifying

\[
\int [Y_{[0]}(D - 3)]^2 d\Omega_{N-1} = 1 .
\]

The surface element is conveniently written as

\[
d\Omega_N = d\Omega_{N-1} d\Omega_{12} = d\Omega_{N-1} d\hat{x}N d\phi_N (\cos \phi_N)^2 (\sin \phi_N)^{3N-4} .
\]

We can extend the definition of the PB elements to depend on the coordinates of a general pair \((i, j)\) as \(P^{l,m}_{2n+1}(\Omega_{ij})\). The coordinates \(\Omega_{ij} \equiv (\hat{x}_N, \phi_N)\) are now defined by a different ordering of the particles entering in the Jacobi coordinates such that \(\mathbf{x}_N = \mathbf{r}_i - \mathbf{r}_j\).

One important property of the PB elements is the following. When a PB element is defined in the space spanned by the coordinates \(\Omega_{ij}\), its expression in terms of HH functions defined in the reference set \(\Omega_N\) corresponding to the ordering of the particles \(1, 2, \ldots, N\) is known and results

\[
P^{l,m}_{2n+1}(\Omega_{ij}) = \sum_{[K']=2n+l} (N)C^{n,l}_{[K']}(\varphi^{ij}) Y^{lm}_{[K']}(\Omega_N),
\]

where the coefficients \((N)C^{n,l}_{[K']}(\varphi^{ij})\) are given by the following relation

\[
(N)C^{n,l}_{[K']}(\varphi^{ij}) = \left( (N)P^{0,0}_{2n+1}(0) \frac{\Gamma(3(N - 1)/2)}{2\pi^{3(N-1)/2}} \right)^{-1} \int d\hat{x}Y^*_{lm}(\hat{x}) Y^{lm}_{[K']}(\Omega^{ij}_{z}) .
\]

The angles \(\varphi^{ij} = \{\varphi^{ij}_N, \ldots, \varphi^{ij}_2\}\) defined from the following kinematic rotation vector

\[
\mathbf{z}(\varphi^{ij}) = \mathbf{x}_N \cos \varphi^{ij}_N + \mathbf{x}_{N-1} \sin \varphi^{ij}_N \cos \varphi^{ij}_{N-1} + \cdots + \mathbf{x}_1 \sin \varphi^{ij}_N \sin \varphi^{ij}_{N-1} \cdots \sin \varphi^{ij}_2
\]

are chosen to verify \(\mathbf{z}(\varphi^{ij}) = \mathbf{r}_j - \mathbf{r}_i\). The hyperangles \(\Omega^{ij}_z\) are defined as \(\Omega^{ij}_z \equiv \{\hat{x}, \ldots, \hat{x}, \varphi^{ij}_N, \ldots, \varphi^{ij}_2\}\), with \(\hat{x}\) repeated \(N\)-times. The particular form of the HH function \(Y^{lm}_{[K']}(\Omega^{ij}_z)\) produces the coefficients of Eq.(25) to be independent of \(m\). In Eq.(21) the sum over all quantum numbers \([K']\) is limited by the condition \(2n + l = K'\), showing that a PB basis element depending on \(\Omega_{ij}\) can be given as a linear combination of HH basis elements having the same value of grand angular quantum number but depending on \(\Omega_N\). A generic function \(f(\mathbf{r}_i - \mathbf{r}_j)\) can be expanded in terms of the PB as

\[
f(\mathbf{r}_i - \mathbf{r}_j) = \sum_{nlm} f_{nlm}(\rho) P^{l,m}_{2n+1}(\Omega_{ij}) ,
\]
with
\[ f_{nlm}(\rho) = \int d\Omega_{ij} f(\mathbf{r}_i - \mathbf{r}_j) \int d\Omega_{N-1} [\mathcal{P}_{2n+l}^{lm}(\Omega_{ij})]^* \]
\[ = \frac{1}{Y_0(D - 3)} \int d\Omega_{ij} Y_{lm}^* (\cos \phi_N)^l P_n^{3(N-1)/2-1,l+1/2}(\cos 2\phi_N) f(\mathbf{r}_i - \mathbf{r}_j). \]

The functions \( f_{nlm}(\rho) \) are the hyperradial multipoles. Introducing the transformation of Eq. (24) in the above expressions it is possible to write a general function \( f(\mathbf{r}_i - \mathbf{r}_j) \) in terms of HH functions given in the reference set. We will use this property for the potential energy of an \( A \)-body system.

### III. THE POTENTIAL ENERGY IN TERMS OF HH FUNCTIONS

A local two-body interaction can be put in the form
\[ V(i, j) = \sum_l \left[ A_l(i, j) \otimes Y_l(i_r) \right] \bar{V}_l(i_{r}) \tag{29} \]
where we use the compact notation
\[ \left[ A_l(i, j) \otimes Y_l(i_r) \right]_{LM} = \sum_{m_1 m_2} (l_1 m_1 l_2 m_2 |LM) A_{l_1 m_1}(i, j) Y_{l_2 m_2}(\hat{r}_{ij}). \tag{30} \]

\( A_{lm}(i, j) \) is an operator independent of the coordinates \( \mathbf{r}_{ij} \), and the coupling with the spherical harmonics to zero in Eq. (29) shows that the potential is a scalar in total space. We can use the PB elements to expand each \( l \)-term of the expansion
\[ V(i, j) = \sum_{ln} \left[ A_l(i, j) \otimes \mathcal{P}_{2n+l}^l(\Omega_{ij}) \right]_0 V_n^l(\rho), \tag{31} \]
where the functions \( V_n^l(\rho) \) are obtained from the following integral in the hyperangular space
\[ V_n^l(\rho) = \int d\Omega_{ij} V_l(i_{\mathbf{r}}) Y_{lm}(\hat{r}_{ij}) \int d\Omega_{N-1} [\mathcal{P}_{2n+l}^{lm}(\Omega_{ij})]^* \]
\[ = \frac{1}{Y_0(D - 3)} \int d\phi_N (\cos \phi_N)^{2+l}(\sin \phi_N)^{3N-4} P_n^{3(N-1)/2-1,l+1/2}(\cos 2\phi_N) V_l(i_{\mathbf{r}}). \tag{32} \]

The complete potential energy is
\[ \sum_{i < j} V(i, j) = \sum_{i < j} \sum_{ln} \left[ A_l(i, j) \otimes \mathcal{P}_{2n+l}^l(\Omega_{ij}) \right]_0 V_n^l(\rho). \tag{33} \]

It would be convenient to have the potential energy expressed in the coordinates defined by \( \Omega \) (in the following we drop the suffix \( N \) for the reference set). To this end we transform
the PB elements obtaining
\[
\sum_{i<j} V(i, j) = \sum_{ln} V_n^l(\rho) \sum_{[K'=2n+l]} \sum_{i<j} \langle N \rangle C_{[K'](\phi^{ij})}^{m,l} [A_l(i, j) \otimes Y_{[K'](\Omega)}]_0
\]
\[
= \sum_{ln} V_n^l(\rho) G_n^l(\Omega),
\]
where we have defined
\[
G_n^l(\Omega) = \sum_{[K'=2n+l]} \left( \sum_{i<j} \langle N \rangle C_{[K'](\phi^{ij})}^{m,l} \right) [A_l(i, j) \otimes Y_{[K'](\Omega)}]_0.
\]

The final form of Eq. (34) gives a general expression for the potential energy in terms of the HH basis elements. In the case of central potentials \( l = 0 \) and \( A_{lm} = 1 \), and the above expressions reduce to (omitting the indices \( l = 0, m = 0 \))
\[
\sum_{i<j} V(i, j) = \sum_{i<j} \sum_{n} \left( \sum_{[K'=2n]} \langle N \rangle C_{[K'](\phi^{ij})}^{m,l} \right) V_n(\rho)
\]
\[
= \sum_{n} V_n(\rho) \sum_{[K'=2n]} \left( \sum_{i<j} \langle N \rangle C_{[K'](\phi^{ij})}^{m,l} \right) Y_{[K'](\Omega)}
\]
\[
= \sum_{n} V_n(\rho) G_n(\Omega),
\]
with
\[
G_n(\Omega) = \sum_{[K'=2n]} \left( \sum_{i<j} \langle N \rangle C_{[K'](\phi^{ij})}^{m,l} \right) Y_{[K'](\Omega)}
\]
\[
= \sum_{i<j} P_{2n}(\Omega_{ij}).
\]

The matrix elements of the potential energy between two different HH basis elements result
\[
\langle Y_{[K_1]}^{l_1 m_1} | \sum_{i<j} V(i, j) | Y_{[K_2]}^{l_2 m_2} \rangle_{\Omega} = \sum_{n,l} V_n(\rho) \langle Y_{[K_1]}^{l_1 m_1} | G_n(\Omega) | Y_{[K_2]}^{l_2 m_2} \rangle_{\Omega}. \tag{38}
\]

The above expression represents an integral in the hyperangular space, and shows the tensor product form between the hyperradius and the hyperangular coordinates which is typical using the HH basis. The matrix elements of the operators \( G_n^l(\Omega) \) are independent of the potential
\[
\langle Y_{[K_1]}^{l_1 m_1} | G_n^l(\Omega) | Y_{[K_2]}^{l_2 m_2} \rangle_{\Omega} = \sum_{[K'=2n+l]} \sum_{i<j} \langle N \rangle C_{[K'](\phi^{ij})}^{m,l}
\]
\[
\times \sum_{m} \frac{(-1)^{l-m}}{\sqrt{2l+1}} A_{l-m}(i, j) \int d\Omega \left[ Y_{[K_1]}^{l_1 m_1}(\Omega) \right]^* Y_{[K_2]}^{l_2 m_2}(\Omega) \tag{39}
\]
Each $G_n^l$ is a combination of HH functions with grand orbital momentum $K' = 2n + l$, therefore its matrix elements follow a triangular relation. In fact, given $K_1$ and $K_2$, the values of $n, l$ to be considered in the sum of Eq.(38) are limited by the relation $|K_1 - K_2| \leq 2n + l \leq K_1 + K_2$. A triangular relation is also verified by the orbital angular momenta: $|L_1 - L_2| \leq l \leq L_1 + L_2$. Furthermore, the matrix elements of $G_n^l$ includes the computation of integrals of three HH basis elements:

$$\int d\Omega \left[ Y_{[K_i]}^{L_1M_1}(\Omega) \right]^* \sum_{i<j} \mathcal{P}_{2n}(\Omega_{ij}) Y_{[K_j]}^{L_2M_2}(\Omega) Y_{[K_2]}^{L_2M_2}(\Omega).$$

(40)

These integrals factorize in products of one-dimensional integrals consisting of either three hyperspherical polynomials or three spherical harmonics that can be obtained analytically or very efficiently using quadratures.

As shown in Eq.(35), each function $G_n^l(\Omega)$ is symmetric in the particle indices, therefore its corresponding eigenvectors will have well defined symmetry under particle permutations. For example, when $A_{lm}(i,j) = 1$, $K_1 = K_2 = K$, $l = 0$, implying $(L_1, M_2) = (L_2, M_2) = (L, M)$, and $2n = 2K$, the following elements

$$\langle Y_{[K]}^{LM} | G_K(\Omega) | Y_{[K]}^{LM} \rangle_{\Omega} = \langle Y_{[K]}^{LM} | \sum_{i<j} \mathcal{P}_{2n}(\Omega_{ij}) Y_{[K]}^{LM} \rangle_{\Omega},$$

(41)

form a matrix by varying all the quantum numbers in $[K]$ with fixed values of $K$ and $L, M$. The dimension of the matrix is given by all HH functions with grand angular quantum number $K$ coupled to $L, M$. Its eigenvectors, which are combinations of this family of HH functions, will have well defined permutational symmetry. This reflects the fact that each $K$-subset is invariant under particle permutations. Therefore, the diagonalization of the above matrix is a way to construct basis elements with well defined permutational symmetry using HH functions with fix values of $K$ and $L$.

IV. APPLICATION TO SYSTEMS WITH $A = 3, 4$

In the description of bound states in an $A$-body system it is common to use basis elements having the required symmetry, symmetric states for bosons or antisymmetric states for fermions. In the present section we will analyze the use of the HH basis without the initial symmetrization or antisymmetrization of the basis. Although the basis elements have not the required symmetry, the eigenvectors of the Hamiltonian will have a well defined symmetry.
reflecting the symmetries appearing in the Hamiltonian. Therefore, among all eigenvectors and eigenvalues, the physical states have to be identified.

By taking opportune linear combinations of the HH basis elements, specific symmetries under particle permutation can be constructed for fixed values of $K$. Therefore two calculations, one in which all HH states up to a maximum value of $K$ are considered and the other in which states with a particular symmetry up to the same value of $K$ are considered, produce the same eigenvectors and eigenvalues. Of course, in the first calculation will appear eigenvectors and eigenvalues belonging to other symmetries not present in the second calculation. The simplification of avoiding the initial basis symmetrization is counterbalanced by the larger dimension of the Hamiltonian matrix.

Limiting the discussion to central potentials, Eq. (38) is well suited for a direct application of the HH basis. Let us introduced the following orthonormal basis element

$$\langle \rho \Omega | m [K] \rangle = \left( \frac{\beta^{(\alpha+1)/2}}{\sqrt{\alpha + m}} \right) L_m^{(\alpha)}(\beta \rho) e^{-\beta \rho/2} Y_{LM}^{[K]}(\Omega),$$

where $L_m^{(\alpha)}(\beta \rho)$ is a Laguerre polynomial with $\alpha = 3N - 1$ and $\beta$ a variational non-linear parameter. We will discuss the case $L = 0$ for $A = 3, 4$. The HH basis elements are

$$\mathcal{Y}_{[K]}(\Omega) = (2)^{P_{K1}^{l_1}K_1}(\phi) \left[ Y_{l_1}(\hat{x}_1) \otimes Y_{l_2}(\hat{x}_2) \right]_0$$

for $A = 3$, and

$$\mathcal{Y}_{[K]}(\Omega) = (2)^{P_{K2}^{l_2}K_1}(\phi_2) \left( 3 \right)^{P_{K3}^{l_3}K_2}(\phi_3) \left[ Y_{l_1}(\hat{x}_1) \otimes Y_{l_2}(\hat{x}_2) \right]_{l_3} \otimes Y_{l_3}(\hat{x}_3)$$

for $A = 4$. The corresponding matrix elements of the Hamiltonian are

$$\langle m' [K']|H|m [K] \rangle = \langle m' [K']|T + V|m [K] \rangle .$$

The matrix elements of the potential energy corresponding to the $\Omega$-space have been discussed in the previous section. Integrating also on $\rho$-space they result

$$\langle m' [K']|V|m [K] \rangle = \sum_n \langle m'|V_n(\rho)|m \rangle \langle m' [K']|\mathcal{G}_n(\Omega)|[K] \rangle \equiv \sum_n (V_{m'm}^n)(\mathcal{G}_{[K']}^[K])_n .$$
The matrix elements of the kinetic energy are the following

\[ T_{K'm';Km} = \langle m' [K'] | -\hbar^2/m \sum_{i=1}^{N} \nabla^2_{x_i} |m [K]\rangle \]

\[ = -\frac{\hbar^2}{m} \langle m' [K'] | \frac{\partial^2}{\partial \rho^2} + \frac{N-1}{\rho} \frac{\partial}{\partial \rho} + \frac{\Lambda_N^2(\Omega)}{\rho^2} | m [K]\rangle \]

\[ = -\frac{\hbar^2 \delta_{|K|,|K'|} T^{K}_{m'm}}{m} \]

\[ = -\frac{\hbar^2 \beta^2}{m} \delta_{|K|,|K'|}[T^{(1)}_{m'm} - K(K + 3N - 2)T^{(2)}_{m'm}], \] (47)

with

\[ T^{(1)}_{m'm} = \frac{1}{4} \delta_{m,m'} + \sqrt{\frac{m!}{(\alpha + m)!} \frac{m!}{(\alpha + m)!}} \times \int_{0}^{\infty} x^{\alpha} e^{-x} dx L^{(\alpha)}_{m'}(x) \left[ 1 - \frac{\alpha + 2m}{2x} - \frac{m}{x^2} L^{(\alpha)}_{m}(x) + \frac{m + \alpha}{x^2} L^{(\alpha)}_{m-1}(x)(1 - \delta_{m,0}) \right], \] (48)

and

\[ T^{(2)}_{m'm} = \sqrt{\frac{m!}{(\alpha + m)!} \frac{m!}{(\alpha + m)!}} \int_{0}^{\infty} x^{\alpha} e^{-x} dx L^{(\alpha)}_{m'}(x) \left( \frac{1}{x^2} \right) L^{(\alpha)}_{m}(x). \] (49)

Using the properties of the Laguerre polynomials, these integrals can be calculated analytically.

Therefore, the matrix elements of the Hamiltonian are sums of tensor products of two matrices, one calculated on \( \rho \)-space, depending on indices \( m, m' \), and one calculated on \( \Omega \)-space, depending on the indices \([K], [K']\)

\[ \langle m' [K'] | H | m [K]\rangle = -\frac{\hbar^2 \beta^2}{m} (T^{(1)}_{m'm} - K(K + 3N - 2)T^{(2)}_{m'm}) \delta_{|K|,|K'|} + \sum_{n}(V^{n}_{m'm})(G^{n}_{K'[K]}). \] (50)

If we introduce the diagonal matrix \( D \) such that \( \langle [K'] | D | [K]\rangle = \delta_{|K|,|K'|} K(K + 3N - 2) \), and the identity matrix \( I \) in \( K \)-space, we can rewrite the Hamiltonian schematically as

\[ H = -\frac{\hbar^2 \beta^2}{m} (I \otimes (1)T + D \otimes (2)T) + \sum_{n} G_{n} \otimes V_{n}, \] (51)

in which the tensor product character of the expression is explicitly given. A scheme to diagonalize such a matrix is given in the Appendix \( \Delta \).

In the following we give results for nucleon systems with \( A = 3, 4 \) using the Volkov potential

\[ V(r) = V_R e^{-r^2/R_1^2} + V_A e^{-r^2/R_2^2} \] (52)
with $V_R = 144.86$ MeV, $R_1 = 0.82$ fm, $V_A = -83.34$ MeV, and $R_2 = 1.6$ fm. The nucleons are considered to have the same mass chosen to be equal to the reference mass $m$ and corresponding to $h^2/m = 41.47$ MeV fm$^{-2}$. With this parametrization of the potential, the two-nucleon system has a binding energy of 0.54592 MeV.

This potential has been used several times in the literature making its use very useful to compare different methods [11, 13, 14, 15]. The results will be obtained after a direct diagonalization of the Hamiltonian matrix of Eq. (50) including $m_{\text{max}} + 1$ Laguerre polynomials with a fix value of $\beta$, and all HH states corresponding to maximum value of the grand angular momentum $K_{\text{max}}$. The scale parameter $\beta$ can be used as a non-linear parameter to study the convergence in the index $m = 0, 1, \ldots, m_{\text{max}}$, with $m_{\text{max}}$ the maximum value considered. In the present analysis the convergence will be studied with respect to the index $K_{\text{max}}$, therefore, the number of Laguerre polynomials at each step, $m_{\text{max}} + 1$, will be sufficiently large to guarantee independence from $\beta$ of the physical eigenvalues and eigenvectors.

In Table I we show the different symmetries of the eigenvectors and the corresponding eigenvalues, for $A = 4$, in the particular case in which the Hamiltonian matrix has been diagonalized for $m_{\text{max}} = 0$, $\beta = 2$ fm$^{-1}$, and $K_{\text{max}} = 6$. In this case the total dimension of the matrix is 56 with 32 “even” elements, corresponding to even values of $l_3$, and 24 “odd” elements corresponding to odd values of $l_3$. In particular there are 6 totally symmetric states, irreducible representation [4] using the Yamaguchi symbol, 2 totally antisymmetric states, [1$^4$], 8 states belonging to the three-dimensional irreducible representation [3 1], 6 states belonging to the two-dimensional irreducible representation [2$^2$], and 4 states belonging to the three-dimensional irreducible representation [2 1$^2$]. The lowest eigenvalue of each irreducible representation is given in the table. In the last two columns of the table, the eigenvalues are reported considering separately even and odd basis elements. Symmetric states are formed exclusively by even-basis element whereas antisymmetric states are formed exclusively by odd-basis elements. The three mixed symmetries, one two-dimensional and the other two three-dimensional, show degenerate eigenvalues. In order to distinguish between the two three-dimensional mixed symmetries, we observe that the three degenerate eigenvalues divide differently in even and odd elements. The mixed symmetry [3 1] is twice degenerate when the expansion basis is restricted to even states whereas the mixed symmetry [2 1$^2$] is not. Therefore performing two different diagonalizations, one using a restricted basis con-
sidering only even states and one considering only odd states, all the symmetries can be identified.

Furthermore, we can see from Table I that a bound state appears in correspondence to a symmetric state. The fact that only one spatial symmetry is present in the bound state is a direct consequence of using a central potential. The final antisymmetrization of the state, as required in the case of four nucleons, is performed by multiplying the spatial symmetric wave function by the corresponding spin functions, singlet spin states $S_{12} = 0$ for the two protons labelled $(1, 2)$ and $S_{34} = 0$ for the two neutrons labelled $(3, 4)$. In the case of using the isospin formalism, the spatial symmetric state is multiplied by a four nucleon antisymmetric spin-isospin function having total spin $S = 0$ and total isospin $T = 0$.

In Tables II and III the convergence of the ground state binding energies for $A = 3, 4$ are given as a function of $K_{\text{max}}$, respectively. In the last column the point Coulomb interaction between the two protons, labelled as particles $(1, 2)$, has been considered. In the case without the Coulomb potential, spatial component of the ground state is completely symmetric. When the Coulomb potential is taken into account this component is symmetric with respect to particles $(1, 2)$. For $A = 4$, it is also symmetric with respect to the particles $(3, 4)$, the two neutrons. In this case it is convenient to introduce the $H$-type Jacobi coordinates (for a recent application see Ref. [16]):

\[
\begin{align*}
    x_3 &= r_2 - r_1 \\
    x_2 &= \frac{r_4 + r_3}{\sqrt{2}} - \frac{r_2 + r_1}{\sqrt{2}} \\
    x_1 &= r_4 - r_3
\end{align*}
\] 

(53)

and construct HH basis elements based on this type of coordinates. These HH functions are linear combinations of the HH function based on the $K$-type coordinates, introduced in Eq.(3) and used in the previous sections, at fixed values of the grand angular quantum number $K$. As example, in Table III the two different types of Jacobi coordinates have been considered. The dimension of the bases indicated corresponds to taking into account even basis elements which are the only ones entering in the construction of the bound states. As stated before, for the $K$-type Jacobi coordinates this means to take even values of $l_3$. For the $H$-type, both $l_1$ and $l_3$ are taken even. The dimension of the problem for obtaining the eigenvalue at $K_{\text{max}} = 30$ results to be 72 for $A = 3$, and 7872 (4056) for $A = 4$, using the $K$-type ($H$-type). The use of the $H$-type Jacobi coordinates reduces the dimension of the
problem by nearly a factor of two.

The calculations corresponding to the two different types of coordinates differ in the set of angles $\varphi^{ij}$ defined in Eq. (26) reflecting the different way of defining the interparticle distances in both cases. In the case in which the symmetric states are identified and constructed before diagonalization, the dimension is reduced to 27 for $A = 3$ and around 600 for $A = 4$. We observe a considerable reduction in the dimension of the eigenvalue problem for the symmetrized basis. However the computational cost of constructing HH states with specific permutational symmetry has to be compared to the simplicity in constructing the matrix elements of $G_n$ and in solving the system of Eq. (51). To be noticed that the results using the symmetrized HH basis of Ref. [8, 14] coincide with the results presented here for each value of $K_{max}$. For the sake of comparison, the results using the stochastic variational method (SVM)[13] are shown in the table.

When the Coulomb potential between protons is included the system can be treated as composed by two different species, the protons and the neutrons, having different interactions and slightly different masses. Using the complete HH basis this cause no extra difficulties since the following term can be added to the Hamiltonian

$$\sum_n (V_{m'm''}^{c,n}) (F_n^{[\kappa'][\kappa]}) ,$$

where $V_{m'm''}^{c,n}$ are the hyperradial matrix elements of the Coulomb potential multipoles and $F_n^{[\kappa'][\kappa]}$ is a matrix equivalent to $G_n^{[\kappa'][\kappa]}$ with the only difference that the sum over $(i, j)$ is limited to protons. This term has the tensor product form and therefore the Hamiltonian reads:

$$H = I \otimes (T^{(1)} + D \otimes (T^{(2)} + \sum_n G_n \otimes V_n + \sum_n F_n \otimes V_{n}^{c}.$$  

In the above equation protons and neutrons are assumed to interact with the same short range potential. For realistic potentials this is not the case and this assumption can be relaxed dividing the potential energy in three parts, one for the interaction between protons, one for the interaction between neutrons and one for the interaction between protons and neutrons. To be noticed that using the complete HH basis the dimension of the problem does not change by distinguishing protons and neutrons or not. However, as is a common procedure, it is possible to treat the system as composed by identical particles using the isospin formalism. The Coulomb potential breaks the isospin symmetry and the use of antisymmetric states requires the inclusion of different isospin components in the wave function. The
A = 3 bound state will have isospin $T = 1/2, 3/2$ components whereas the $A = 4$ bound state will have $T = 0, 1, 2$ components. After including all these components the two procedures, one using the complete HH basis and the other using antisymmetrized states, will produce the same eigenvalues. An example for this case is given in Table IV in which the results for $A = 3, 4$ using antisymmetric basis states, including the different isospin components, are shown. For $A = 3$, the $T = 1/2$ component is by far the most important one; however, the exact result is obtained after including both components, $T = 1/2$ and $3/2$. To be noticed that the dimension of the basis using antisymmetrized HH states with isospin components $T = 1/2, 3/2$ is the same of the complete HH states using even basis elements. Therefore in this case the preliminary antisymmetrization of the basis is not convenient. For $A = 4$ the $T = 0$ component is by far the most important; however, the exact result is obtained after including the three isospin components $T = 0, 1, 2$. In this case the dimension of the basis using even HH states up to $K_{\text{max}}$ is greater than that using antisymmetrized basis elements, since in the symmetrization with respect the two neutrons is not included automatically in the even HH states and has to be constructed by the diagonalization procedure. However the difference in the dimension of the two cases is considerably reduced with respect to the case in which the Coulomb potential was not included.

The equivalence of the last columns of Tables II and III with columns third and sixth of Table IV illustrates the simplicity of treating symmetry breaking terms using the HH basis without permutational symmetry.

V. CONCLUSIONS

In this work we have presented a direct use of the HH basis in the description of a $A$-body system. The basis has neither been symmetrized nor antisymmetrized as required by a system of identical particles. However, the eigenvectors of the Hamiltonian have well defined permutation symmetry. Among all the eigenvectors, the physical ones can be identified. The benefit of the direct use of the HH basis is based on a particular simple form used to represent the potential energy. Each term of the two-body potential $V(i, j)$ has been expanded in the potential basis and then expressed in terms of the HH basis, defined in the reference set, by using the corresponding transformation coefficients. These coefficients are known for each value of $K$ and for a general number of particles $A$. Once the potential has been expressed
in terms of the HH basis, it results in a sum of tensor product terms originated from the separation of the hyperradial and the hyperangular coordinates inherent to the method. Moreover, the kinetic energy can be put in a tensor product form too. Therefore, the matrix representation of the Hamiltonian is expressed as a sum of tensor product matrices, and this particular form can be diagonalized very efficiently using the technique given in the Appendix. As a test case we have studied three and four nucleons interacting through a central potential, the Volkov potential, used many times in the literature. We have shown how the symmetries are present in the spectrum and can be identified. The symmetric and antisymmetric states appear as singlets, whereas the mixed symmetries appear as multiplets. We have identified all symmetries by dividing the spectrum in even and odd components. In the studied cases, only one bound state appears for \( A = 3 \) and \( 4 \) corresponding to a symmetric state. To be noticed that if the potential depends on the spin- isospin degrees of freedom, the Hamiltonian will still present the tensor product form in the hyperradial, hyperangular, spin and isospin spaces.

For \( A = 4 \) we have solved the problem using two different types of Jacobi coordinates, namely the \( K \)-type corresponding to a 3+1 configuration, and the \( H \)-type corresponding to a 2+2 configuration. The calculations using one or the other set differ in the values of the angles \( \varphi^{ij} \), which can be considered as input parameters. Therefore the method gives a systematic way of introducing the different types of Jacobi coordinates. The convenience of selecting one specific type is related to its capability to produce basis states having partially the required symmetry with a reduction of the total dimension of the problem. In the cases presented here, the \( A = 4 \) bound state is constructed using basis elements based on the \( K \)-type Jacobi coordinates with even values of \( l_3 \) or based on the \( H \)-type with both \( l_1 \) and \( l_3 \) restricted to even values. The latter resulting in a basis with a dimension smaller by a factor of two.

A further benefit of using the complete HH basis is obtained when symmetry breaking terms are included in the Hamiltonian. The complete basis will generate eigenvectors having specific permutation symmetries reflecting the symmetries present in the Hamiltonian. The complexity of the numerical problem does not increase when these terms are present. This is not the case when symmetrized or antisymmetrized basis are used. For example, in the case of a nuclear system, the presence of charge symmetry breaking terms requires the extension of the basis to include all the isospin components. As a specific example, here we have
analyzed the case of the Coulomb interaction between protons. The results using the HH basis without well defined permutation symmetry have been compared to the case in which antisymmetrized HH basis have been used. In the latter case the different isospin components entering in the wave function have to be included, resulting in spatial components having more than one symmetry. Accordingly the dimension of the basis increases. To this respect, the numerical effort to reduce the Hilbert space to subspaces with specific permutation symmetry is discussed in Ref. [8].

When one spatial symmetry is required, as for example a completely symmetric spatial state, the convenience of constructing symmetric HH state is obvious. When several spatial symmetries are present in the wave function, as in the case of an \(A\)-nucleus wave function, the convenience of constructing HH states with different spatial symmetries has to be compared to the capability of solving a large eigenvalue problem, for example that one given in Eq. (51).

In Refs. [14, 17, 18] the HH basis, used to describe three- and four-nucleon bound states, is antisymmetrized in the following way. The total wave function is expanded in angular-spin-isospin channels and, for each channel, it is written as a sum of Faddeev-like amplitudes, each of them antisymmetric in the pair \((i, j)\). In this way the total wave function results antisymmetric. Then, each \((i, j)\)-amplitudes is expanded in the HH basis defined from Jacobi vectors corresponding to the different ordering of the particles. As a consequence, the amplitudes for the different channels are not orthogonal, resulting in a non-orthogonal basis. For large values of \(K\) the non orthogonality of the basis could causes numerical instabilities.

In particular, for \(A = 4\), this problem is overcome performing an orthonormalization of the basis using the Gram-Schmidt technique with quadruple precision in the numerical treatment of the process. Therefore, the extension to \(A > 4\) systems appears to be difficult. On the other hand, the direct use of the HH basis without antisymmetrization circumvents this problem. Therefore, the method presented here has to be considered a first step in a program devoted to the application of the HH basis to systems with \(A > 4\). Further works along this line are the extension of the method to treat realistic interactions and the numerical implementation of the Hamiltonian of Eq. (51) to systems with \(A = 5, 6\). The extension of the method to treat three-nucleon interaction terms is also possible. In fact, the transformation of the spatial part of a three-nucleon interaction \(W(i, j, k)\) in terms of HH functions constructed in the reference set can be performed using the algorithm developed for the ”triplet basis” in Ref. [5].
APPENDIX A: EFFICIENT MATRIX-VECTOR PRODUCT FOR TENSOR-PRODUCT MATRICES

The algorithm we used to diagonalize the Hamiltonian is an iterative one, namely Lanczos’s \[19\]. These kind of algorithms are useful whenever an efficient matrix-vector product can be used, as in the case of sparse matrices; in the specific calculation, we have the product between a tensor-product matrix \( M = A_1 \otimes A_2 \), and a vector \( \mathbf{v} \)

\[
\mathbf{w} = M \cdot \mathbf{v} = (A_1 \otimes A_2) \cdot \mathbf{v}, \quad (A1)
\]

with \( A_1 \) a \( n \times n \) matrix, \( A_2 \) a \( m \times m \) matrix, and \( \mathbf{v} \) a \((n \cdot m)\)-dimensional vector.

The product is done in three steps: (i) first, the vector \( \mathbf{v} \) is reshaped in a \( m \times n \) matrix \( V \); (ii) then, the following matrix products are performed

\[
W = (A_1 \cdot (A_2 \cdot V)^T)^T; \quad (A2)
\]

(iii) finally, the matrix \( W \) is reshaped into the \((n \cdot m)\)-dimensional vector \( \mathbf{w} \), which is the result of the multiplication. The above algorithm is easily generalized to tensor-products of \( k \)-matrices \[20\].

[1] M. Fabre de la Ripelle, Ann. Phys.(NY) 147, 281 (1983).
[2] A. Kievsky, S. Rosati, M. Viviani, L.E. Marcucci, and L. Girlanda, J. Phys. G: Nucl. Part. Phys. 35, 063101 (2008).
[3] J. Raynal and J. Revai, Nuovo Cimento A 68, 612 (1970).
[4] R. Krivec and V.B. Mandelzweig, Phys. Rev. A 42, 3779 (1990).
[5] M. Viviani, Few-Body Syst. 25, 177 (1998).
[6] A. Novoselsky and J. Katriel, Phys. Rev A 49, 833 (1994).
[7] V.D. Efros, Few-Body Syst. 19, 167 (1995).
[8] A. Novoselsky and N. Barnea, Phys Rev A 51, 2777 (1995).
[9] N. Barnea and A. Novoselsky, Ann. of Phys. 256, 192 (1997).
[10] N. Barnea and A. Novoselsky, Phys. Rev. A 57, 48 (1998).
[11] N. Barnea, Phys. Rev. A 59, 1135 (1999).
[12] N. K. Timofeyuk, Phys. Rev. C 78, 054314 (2008).
[13] K. Varga and Y. Suzuki, Phys. Rev. C 52, 2885 (1995).
[14] M. Viviani, A. Kievsky, and S. Rosati, Phys. Rev. C 71, 024006 (2005).
[15] N. K. Timofeyuk, Phys. Rev. C 65, 064306 (2002).
[16] J. Vijande, E. Weissman, A. Valcarce and N. Barnea, Phys. Rev. D 76, 094027 (2007).
[17] A. Kievsky, M. Viviani, and S. Rosati, Nucl. Phys. A 551, 241 (1993).
[18] A. Kievsky, L.E. Marcucci, S. Rosati, and M. Viviani, Few-Body Syst. 22, 1 (1997).
[19] The IETL Project, http://www.comp-phys.org/software/ietl/.
[20] Paul E. Buis, and Wayne R. Dyksen, ACM Trans. Math. Softw. 22, 18 (1996).
TABLE I: Lowest Volkov-energy eigenvalues of each irreducible representations of $S_4$ for the $N = 4$ case, with $m_{\text{max}} = 0$, $K_{\text{max}} = 6$, and $\beta = 2 \text{fm}^{-1}$. The multiplets are further identified as being symmetric or anti-symmetric under permutation of particles 1-2.

| Irreps | Eigen's (MeV) | Sym(1-2) | AntiSym(1-2) |
|--------|---------------|----------|--------------|
| $[4]$  | -25.794       | -25.794  |              |
| $[2^2]$ | 27.680       | 27.680   | 27.680       |
| $[31]$ | 28.430       | 28.430   | 28.430       |
| $[21^2]$ | 102.85      | 102.85   | 102.85       |
| $[1^4]$ | 199.56       |          | 199.56       |
TABLE II: Results for the Volkov’s potential, as a function of $K_{\text{max}}$ using 30 Laguerre’s polynomials, and $\beta = 3 \text{ fm}^{-1}$ for the three-body case. In the last column the results including the Coulomb potential are given.

| $K_{\text{max}}$ | $N_{\text{HH}}$ | $E$ (MeV) | $E$ (MeV) |
|------------------|------------------|------------|------------|
| 0                | 1                | 7.7075     | 6.9926     |
| 10               | 12               | 8.4157     | 7.7083     |
| 20               | 36               | 8.4623     | 7.7566     |
| 30               | 72               | 8.4647     | 7.7693     |
| 40               | 121              | 8.4649     | 7.7694     |
| SVM[13]          |                  |            |            |

| SVM[13]          | 30               | 8.46       |            |

TABLE III: $A = 4$ results for the Volkov’s potential, using 25 Laguerre’s polynomials, and $\beta = 2 \text{ fm}^{-1}$. Two different types of Jacobi coordinates have been used. In the last two columns the results without and with Coulomb potential, using independently $K$- or $H$-type Jacobi coordinates, are given, respectively. At fixed value of $K_{\text{max}}$ the results, using either one or the other type of coordinates, coincide.

| $K_{\text{max}}$ | $N_{\text{HH}}(K\text{-type})$ | $N_{\text{HH}}(H\text{-type})$ | $E$ (MeV) | $E$ (MeV) |
|------------------|---------------------------------|---------------------------------|-----------|-----------|
| 0                | 1                               | 1                               | 28.580    | 27.748    |
| 10               | 136                             | 78                              | 30.278    | 29.456    |
| 20               | 1547                            | 819                             | 30.416    | 29.596    |
| 30               | 7872                            | 4056                            | 30.420    | 29.599    |
| SVM[13]          | 50                              |                                 | 30.42     |            |
TABLE IV: Contributions to the bound state energies, for $A = 3, 4$, of the different isospin components using antisymmetrized HH functions

| $K_{\text{max}}$ | $T = 1/2$ | $T = 1/2, 3/2$ | $K_{\text{max}}$ | $T = 0$ | $T = 0, 1, 2$ |
|------------------|-----------|----------------|------------------|---------|----------------|
| 0                | 6.9926    | 6.9926         | 0                | 27.748  | 27.748         |
| 10               | 7.7072    | 7.7083         | 10               | 29.453  | 29.456         |
| 20               | 7.7555    | 7.7566         | 20               | 29.594  | 29.596         |
| 30               | 7.7582    | 7.7593         | 30               | 29.596  | 29.599         |
| 40               | 7.7583    | 7.7594         |                  |         |                |