

CLUSTER CALCULATION FOR ${}^9_{\Lambda}\text{Be}$ HYPERNUCLEUS IN FADDEEV
APPROACH

VLADIMIR SUSLOV^{a,b}, IGOR FILIKHIN^{a,b} and BRANISLAV VLAHOVIĆ^{b,1}

^a*Department of Mathematical and Computational Physics,
Sankt-Petersburg State University, 198504, Petrodvorets, Russia*

^b*Department of Physics, North Carolina Central University, Durham, NC 27707, USA*

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Configuration-space Faddeev calculations are performed for cluster model $\alpha\alpha\Lambda$ of the ${}^9_{\Lambda}\text{Be}$ hypernucleus using various $\alpha\Lambda$ potentials. For the $\alpha\alpha$ interaction, the nuclear component is only taken into account (phenomenological Ali-Bodmer potential). The binding energy of the ${}^9_{\Lambda}\text{Be}$ hypernuclei is calculated for two different potential models. In the first model, the s-wave $\alpha\Lambda$ potential acting in all partial waves in the $\alpha\Lambda$ subsystem is used. In the second model, a recent more realistic $\alpha\Lambda$ potential having the s- and p-partial components is employed. The core effect of nuclear $\alpha\alpha$ potential is also studied.

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1. Introduction

Configuration-space Faddeev calculations are performed to study the ${}^9_{\Lambda}\text{Be}$ hypernucleus considered as two α clusters and a Λ hyperon. We estimated higher partial-wave contribution to the binding energy of the ${}^9_{\Lambda}\text{Be}$ ground state ($\frac{1}{2}^+$) and particularly contribution coming from higher partial waves of the $\alpha\Lambda$ interaction. The Coulomb force between the α 's is not taken into account. The phenomenological Ali-Bodmer potential [1] is used for the description of the $\alpha\alpha$ interaction. For the $\alpha\Lambda$ potential, both form and parameters are uncertain, because $\alpha\Lambda$ interaction data are limited by the experimental value of binding energy of the ${}^5_{\Lambda}\text{He}$ hypernucleus, which is considered as the bound s-wave state of the $\alpha\Lambda$ system. To calculate the binding energy of the $\alpha\alpha\Lambda$ system, an s-wave potential model was

¹Corresponding author, e-mail address: branko@jlab.org

used in Ref. [2]. In this model, one and the same s-wave potential acts in all partial waves of decomposition of the $\alpha\Lambda$ interaction. It leads to unrealistic large p -wave contribution to the binding energy. In a recent work [3], a new $\alpha\Lambda$ potential having the s- and p-wave components was proposed. This potential was constructed on the basis of the NSC97 model of ΛN interaction [4]. In the present work, we use both $\alpha\Lambda$ potential models and compare p-wave contributions to the binding energy. We also study the core effect of nuclear $\alpha\alpha$ potential.

2. Theory

The bound state of the $\alpha\alpha\Lambda$ system is calculated by solving the differential Faddeev equations [5], which have the following general form:

$$\{H_0 + V_\alpha(|\bar{x}_\alpha|) - E\}\Psi_\alpha(\bar{x}_\alpha, \bar{y}_\alpha) = -V_\alpha(|\bar{x}_\alpha|) \sum_{\beta \neq \alpha} \Psi_\beta(\bar{x}_\beta, \bar{y}_\beta),$$

where V_α is the short-range pair interaction in the channel α , $H_0 = -\Delta_{\bar{x}_\alpha} - \Delta_{\bar{y}_\alpha}$ is the kinetic energy operator, E is the total energy and the wave function of the three-body system Ψ is given as a sum over the three Faddeev components, $\Psi = \sum_{\alpha=1}^3 \Psi_\alpha$.

When two particles of the three-body system are identical as α particles in ${}^9_{\Lambda}\text{Be}$ ($\alpha\alpha\Lambda$), the coupled set of Faddeev equations reduces to two equations:

$$(H_0 + V_{\alpha\alpha} - E)U = -V_{\alpha\alpha}(W + P_{12}W), \quad (H_0 + V_{\alpha\Lambda} - E)W = -V_{\alpha\Lambda}(U + P_{12}W), \quad (1)$$

where P_{12} is the permutation operator for the α particles (particles 1,2), $V_{\alpha\alpha}$ and $V_{\alpha\Lambda}$ are the nuclear potentials of $\alpha\alpha$ and $\alpha\Lambda$ interactions, respectively, U is the Faddeev component corresponding to the rearrangement channel $(\alpha\alpha) - \Lambda$ and W corresponds to the rearrangement channel $(\alpha\Lambda) - \alpha$. The total wave function is expressed by the components U and W : $\Psi = U + (1 + P_{12})W$. The total orbital angular momentum is given by $\mathbf{L} = \boldsymbol{\ell}_{\alpha\alpha} + \boldsymbol{\lambda}_{(\alpha\alpha)-\Lambda} = \boldsymbol{\ell}_{\alpha\Lambda} + \boldsymbol{\lambda}_{(\alpha\Lambda)-\alpha}$, corresponding to two available channels of rearrangements.

The ground state of the $\alpha\alpha\Lambda$ system, having zero total angular momentum L , is considered as the $(\frac{1}{2})^+$ state of the ${}^9_{\Lambda}\text{Be}$ hypernucleus. Possible combinations for relative momenta $\ell_{\alpha\alpha}$ and $\lambda_{(\alpha\alpha)-\Lambda}$ are (0,0), (2,2), (4,4), . . . , and for $\ell_{\alpha\Lambda}$, $\lambda_{(\alpha\Lambda)-\alpha}$ are (0,0), (1,1), (2,2), (3,3), (4,4), Thus the pair of quantum numbers $\ell_{\alpha\alpha}$ and $\ell_{\alpha\Lambda}$ describe completely the angular states of subsystems.

To describe interactions in the $\alpha\alpha\Lambda$ system, local pairwise potentials are used. The nuclear $\alpha\alpha$ interaction is given by version ‘‘a’’ of the phenomenological Ali-Bodmer potential [1] having s-, d- and g-wave components which was modified in Ref. [6]. This potential has the following form, $V_{\alpha\alpha}(r) = \sum_{l=0,2,4} V_{\alpha\alpha}^l(r)P_l$, where P_l is a projector onto the state of the $\alpha\alpha$ pair with the orbital momentum l and the functions $V_{\alpha\alpha}^l(x)$ have the form of one or two ranges Gaussians

$$V_{\alpha\alpha}^l(r) = V_{\text{rep}}^l \exp(-r/\beta_{\text{rep}}^l)^2 - V_{\text{att}}^l \exp(-r/\beta_{\text{att}}^l)^2. \quad (2)$$

The parameters of partial components $V_{\alpha\alpha}^l(r)$ are given in the Table 1. The s-wave component $V_{\alpha\alpha}^0(r)$ has strong repulsive core which simulates Pauli blocking effect for the α 's at short distances.

TABLE 1. Parameters of the $\alpha\alpha$ and $\alpha\Lambda$ potentials for different pair angular momenta l .

System	Potential	l	V_{rep}^l (MeV)	β_{rep}^l (fm)	V_{att}^l (MeV)	β_{att}^l (fm)
$\alpha\alpha$	Ali-Bodmer [1]	0	125.0	1.53	30.18	2.85
		2	20.0	1.53	30.18	2.85
		4	–	–	30.0	0.35
$\alpha\Lambda$	TH [7]	0	–	–	60.17	1.2729
	Gibson [8]		–	–	43.48	1.5764
	Isle(DA) [11]		450.4	1.25	404.9	1.41
	MSA [3]		91.0	1.3	95.0	1.7
		1	33.4	1.3	39.4	1.7

For the $\alpha\Lambda$ interaction, a number of various potentials is used. These potentials have different shape and parameters, but reproduce well the experimental value of the binding energy of the ${}^5_{\Lambda}\text{He}$ hypernucleus which is considered as an s-wave bound state of the $\alpha\Lambda$ system. The Tang-Herndon (TH) potential [7] and Gibson I (Gibson) potential [8, 9] have the form of one-range Gaussian. The Maeda-Schmid potential (MS) [10] is a sum of two Woods-Saxon type functions

$$V_{\alpha\Lambda}(r) = V_{\text{rep}}(\exp((r - r_{\text{rep}})/a_{\text{rep}}) + 1)^{-1} - V_{\text{att}}(\exp((r - r_{\text{att}})/a_{\text{att}}) + 1)^{-1},$$

where $V_{\text{rep}} = 18.09$ MeV, $V_{\text{att}} = 35.98$ MeV, $r_{\text{rep}} = 0.88$ fm, $r_{\text{att}} = 1.72$ fm, $a_{\text{rep}} = 0.2353$ fm, $a_{\text{att}} = 0.3541$ fm. The Isle potential [11] has a form of two range Gaussian Eq.(2). In addition to these s-wave $\alpha\Lambda$ potentials, we use potential of Myint, Shimura and Akaishi (MSA) proposed in Ref. [3], having p-wave component and two rank Gaussian form. General formula for $\alpha\Lambda$ potential is written by

$$V_{\alpha\Lambda}(r) = \sum_{l=0,1,\dots} V_{\alpha\Lambda}^l(r)P_l, \quad (3)$$

where P_l is a projector onto the state of the $\alpha\Lambda$ system having the orbital momentum l , $V_{\alpha\Lambda}^l(r)$ are partial components of the potential. The parameters of the s- and p-wave components are listed in Table 1.

3. Results

The coupled configuration-space Faddeev equations, Eq. (1), for the $\alpha\alpha\Lambda$ system are solved numerically, applying the method given in Ref. [12]. We use two different potential models. In the first model, the s-wave $\alpha\Lambda$ potential acts in all partial

waves of the $\alpha\Lambda$ subsystem: $V_{\alpha\Lambda}^l(r) = V_{\alpha\Lambda}^0(r)$, $l = 1, 2, 3, 4$. In the second model, we use two partial components of the $\alpha\Lambda$ potential Eq. (3). These are the s- and p-components $V_{\alpha\Lambda}^l(r)$, $l = 0, 1$, where $V_{\alpha\Lambda}^0(r)$ can be any potential from potentials considered above, and $V_{\alpha\Lambda}^1(r)$ is the p-wave component of potential that is given in Ref. [3]. The results of the calculations are listed in Table 2. For the first model, they are in good agreement with those of a recent Faddeev calculation [2]. In our

TABLE 2. Calculated binding energy E_B (in MeV) of the $\alpha\alpha\Lambda$ system for the Ali-Bodmer $\alpha\alpha$ potential and different models of the $\alpha\Lambda$ interaction. The Coulomb potential between α 's is not taken into account. Here $l_{\alpha\Lambda}$ and $l_{\alpha\alpha}$ are relative angular momenta of subsystems, $l_{\max} = \max\{l_{\alpha\Lambda}, l_{\alpha\alpha}\}$.

Model	l_{\max}	$l_{\alpha\Lambda}$	$l_{\alpha\alpha}$	TH	MS	MSA	Isle(DA)
First	0	0	0	-6.844	-7.365	-8.392	-8.429
	1	0,1	0	-7.363	-8.278	-9.631	-9.768
	2	0,1,2	0,2	-7.786	-8.494	-9.758	-9.920
	3	0,1,2,3	0,2	-7.799	-8.503	-9.764	-9.935
	4	0,1,2,3,4	0,2,4	-7.813	-8.505	-9.765	-9.950
Cravo et al. [2]	-	0,1,2	0,2,4	-7.72	-8.49	-	-10.04
Second	4	0,1	0,2,4	-7.788	-8.050	-8.905	-8.948

calculations, all partial waves up to $l = 4$ were taken into account, but the first three waves ($l_{\alpha\Lambda} \leq 2$, $l_{\alpha\alpha} \leq 2$) give the main contribution. By comparing the first and the second model, one can conclude that the p-wave contribution to the $\alpha\alpha\Lambda$ binding energy is too large. The more realistic $\alpha\Lambda$ potential model of Myint et al. [3], which has the s- and p-wave components, reduces p-wave contribution of the $\alpha\Lambda$ interaction.

TABLE 3. Binding energy E_B (in MeV) of the $\alpha\alpha\Lambda$ system, rms distance between two α 's $r_{\alpha\alpha}$ (in fm), rms distance between the center of α 's pair mass and hyperon $r_{(\alpha\alpha)-\Lambda}$, the most probable distance between two α 's x_{\max} for different models of the $\alpha\Lambda$ interaction computed for $l_{\alpha\Lambda} = l_{\alpha\alpha} = 0$ (s-wave approximation). The binding energy $E_B(V_{\alpha\alpha} = 0)$ in the case $V_{\alpha\alpha} = 0$ and the interaction energy of α pair is $\Delta B_{\alpha\alpha}$.

Potential	E_B	$r_{\alpha\alpha}$	$r_{(\alpha\alpha)-\Lambda}$	$E_B(V_{\alpha\alpha} = 0)$	x_{\max}	$\Delta B_{\alpha\alpha}$
TH	-6.844	3.75	2.65	-9.601	2.4	2.757
Gibson	-7.429	3.69	2.67	-8.934	2.5	1.510
MS	-7.364	3.68	2.72	-8.079	2.5	0.715
MSA	-8.392	3.65	2.80	-8.056	2.6	-0.335
Isle(DA)	-8.429	3.64	2.82	-7.839	2.8	-0.590

To explain substantial disagreements of the binding energies calculated with different $\alpha\Lambda$ potentials, we computed the interaction energy between the α particles for the bound $\alpha\alpha\Lambda$ system, which is defined by the formula

$$\Delta B_{\alpha\alpha} = E_B - E_B(V_{\alpha\alpha} = 0),$$

where the binding energy $E_B(V_{\alpha\alpha} = 0)$ of the $\alpha\alpha\Lambda$ system is calculated when $\alpha\alpha$ interaction is absent ($V_{\alpha\alpha} = 0$). Results of calculation, together with other physical characteristics, are presented in Table 3. In addition, we calculate the most probable configuration of the $\alpha\alpha\Lambda$ system. In Fig. 1, the distances between particles and their localization are shown. From the table and the figure, one can see that the $\alpha\Lambda$ potentials having a weak core (MS), or not having it at all (TH, Gibson), bring the α particles close together. However, at small distances, a repulsion starts between the α 's due to the presence of the strong repulsive core in the $\alpha\alpha$ potential, which decreases the $\alpha\alpha\Lambda$ binding energy.

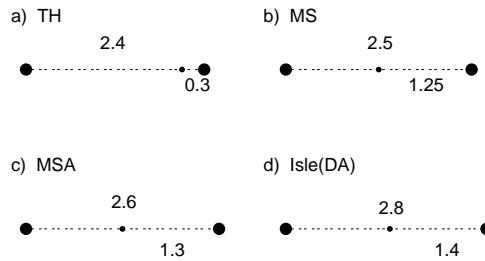


Fig. 1. Most probable configuration of the $\alpha\alpha\Lambda$ system for various $\alpha\Lambda$ potentials. The upper numbers are distances between the α particles (in fm) and the lower ones are distances between α particle and hyperon.

4. Conclusion

The ${}^9_{\Lambda}\text{Be}$ hypernucleus is considered in the framework of the $\alpha\alpha\Lambda$ cluster model. We have studied the contributions of higher partial waves of the $\alpha\Lambda$ interactions to the binding energy of the ground state. We found that the p-wave contribution of $\alpha\Lambda$ interaction is unnatural large when potential model is applied in which the s-wave of the potential acts in all partial waves of the $\alpha\Lambda$ subsystem. This contribution can be reduced by including a realistic p-wave component of the $\alpha\Lambda$ interaction. The differences between the $\alpha\alpha\Lambda$ binding energies calculated for various $\alpha\Lambda$ potentials can be explained by influence of a repulsive core of nuclear $\alpha\alpha$ interaction.

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NAKUPINSKI RAČUN ${}^9_{\Lambda}\text{Be}$ HIPERJEZGRE FADDEEVOM METODOM

Načinili smo račune Faddeevom metodom u konfiguracijskom prostoru za nakupinski model $\alpha\alpha\Lambda$ hiperjezgre ${}^9_{\Lambda}\text{Be}$ primjenom različitih potencijala. Za nuklearnu sastavnicu međudjelovanja $\alpha\alpha$ primijenili smo samo fenomenološki Ali-Bodmerov potencijal. Energiju vezanja hiperjezgre ${}^9_{\Lambda}\text{Be}$ smo računali dvama različitim potencijalnim modelima. U prvome se rabi s-valni potencijal $\alpha\Lambda$ koji djeluje na sve parcijalne valove u podsustavu $\alpha\Lambda$. U drugome, rabi se nov realističniji potencijal $\alpha\Lambda$ koji sadrži s- i p-parcijalne sastavnice. Proučavali smo također učinak sredice u potencijalu $\alpha\alpha$.