The interaction between H-dibaryons


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Abstract

We investigate the interaction between H-dibaryons employing a quark cluster model with a one-gluon-exchange potential and an effective meson exchange potential (EMEP). A deeply-bound state of two H-dibaryons due to the medium range attraction of the EMEP is obtained. The bound H–H system has a size of about 0.8 ∼ 0.9 fm because of the short-range repulsion generated by the color-magnetic interaction and the Pauli principle.

1 Introduction

The H-dibaryon is a doubly strange six-quark state with spin and isospin 0. Since its first prediction in 1977 [1], it has been the subject of many theoretical and experimental studies as a possible candidate of a strongly bound exotic state. The main question is whether its mass is lighter than the ΛΛ threshold, because such a light H-dibaryon is stable against strong decay. However, theoretical estimates of its mass have varied widely ranging from deeply bound to unbound, depending on the model or theory used for the calculations [2]. Experimentally, there is no conclusive result yet, although several candidates for an H-dibaryon decay have been reported [5]. Many efforts to search for the H-dibaryon are under way [6] and planned. A double-hypernucleus
event with a ΛΛ binding energy less than about 30 MeV, has recently been found in
an emulsion experiment at KEK [7]. This excludes the possibility of a deeply-bound
H-dibaryon.

The main dynamical factor responsible for a deeply-bound H-dibaryon is the color-
magnetic interaction (CMI) in the one-gluon-exchange potential (OGEP). The CMI
plays an important role in explaining the low-lying hadron spectra and the short-range
part of the baryon–baryon interaction. In the H-dibaryon, the CMI is attractive. On
the other hand, it acts as a repulsive force for the nucleon-nucleon interaction and also
for almost all nucleon-hyperon and hyperon-hyperon interactions. As is shown later,
the CMI also generates a repulsive force between two H-dibaryons. Another important
factor determining the quark dynamics is the Pauli exclusion principle. It is known
that the Pauli principle gives repulsion for some channels of the nucleon–hyperon and
hyperon–hyperon systems. Furthermore, because of the color exchange character of
the OGEP, the CMI is only effective when the quark exchange interaction between
different baryons is taken into account.

Quark exchanges between clusters can be correctly treated in the quark cluster
model through the antisymmetrization of the wave function. Therefore, we employ the
quark cluster model to study the interaction between H-dibaryons in this paper.

Assuming that the CMI plays a key role in determining the properties of the H–H
interaction, Tamagaki suggested the possibility that a new form of hadronic matter,
called H-matter, appears at densities several times higher than normal nuclear density
[8]. In this pioneering study, a simple H–H interaction model consisting of a hard-core
potential plus a square well attractive potential outside the core was used, because no
microscopic calculation of the H–H interaction was available at that time.

In the present study, we extract more quantitative information on the H–H inter-
action by employing the quark cluster model used in Ref.[3]. In this model, quarks are
confined by an inter-quark potential and subject to the OGEP [9]. The medium range
attraction is taken into account by an effective meson exchange potential (EMEP) be-
tween H-dibaryons. Here we take only flavour singlet scalar meson exchange between
the H-dibaryons into account. In the next section, we briefly explain the model used
in the calculation and give the necessary matrix elements of the spin-flavor-color op-
erators. The results of the calculation and the discussion of the properties of the H–H
interaction are given in section 3. Section 4 is devoted to a summary.
2 Model

In the present quark cluster model, the H-dibaryon wave function $\phi(\xi)$ is assumed to be a $(0s)^6$ configuration divided by the $0s$ wave function of the center-of-mass of the H-dibaryon with the size parameter $\sqrt{1/6b}$ for the orbital part, $\varphi(\xi)$, where $\xi$ is the internal coordinate.

$$\phi(\xi) = \varphi(\xi)S([222]_{SF})C([222]_C).$$  \hspace{1cm} (1)

Here, $S([222]_{SF})$ and $C([222]_C)$ are the spin-flavor and color parts, respectively, with their permutational symmetries in parentheses. The total wave function of the H–H system is written as

$$\Psi(\xi_1, \xi_2, r) = \mathcal{A}[\phi(\xi_1)\phi(\xi_2)\chi(r)],$$  \hspace{1cm} (2)

where the wave function of the relative motion between the H-dibaryons $\chi(r)$ is introduced and $r$ is the relative coordinate between the H-dibaryons. The wave function of the relative motion is further decomposed into components of the angular momentum of the relative motion, $L$, as

$$\chi(r) = \sum_{LM} \frac{\chi_L(r)}{r} Y_{LM}(\hat{r}).$$  \hspace{1cm} (3)

In this paper, only relative S-waves are considered.

The antisymmetrization operator $\mathcal{A}$ can be expressed as

$$\mathcal{A} \equiv 1 - 36 P_{67} + 225 P_{57} P_{68} - 400 P_{47} P_{58} P_{69},$$  \hspace{1cm} (4)

where $P_{ij}$ is the permutation operator. Because each H-dibaryon consists of 6 quarks, there are 0,1,2 and 3 quark exchanges between H-dibaryons which are shown in Fig. 1. Because we symmetrize the two H–dibaryons system under a permutation of the two H-dibaryons, 4,5 and 6 quark exchanges are taken into account by the 0,1,2 and 3 quark exchanges and the exchange of the clusters as a whole.

The Hamiltonian $H$ consists of the kinetic energy term, $K$, and the interaction term, $V$.

$$H = K + V,$$  \hspace{1cm} (5)

$$K = \sum_{i=1}^{12} \frac{p_i^2}{2m} - K_G,$$  \hspace{1cm} (6)

where $K_G$ is the center-of-mass energy of the twelve-quark system, and

$$V = \sum_{i<j} (V_{ij}^{\text{OGEp}} + V_{ij}^{\text{conf}}).$$  \hspace{1cm} (7)
Here, $V_{ij}^{\text{OGEP}}$ and $V_{ij}^{\text{conf}}$ are the one-gluon-exchange potential (OGEP) and the confinement potential, respectively:

$$V_{ij}^{\text{OGEP}} = \frac{\alpha_s}{4} \left[ \frac{1}{r_{ij}} - \frac{\pi}{m_i m_j} \delta(r_{ij}) \left( 1 + \frac{2}{3} \sigma_i \cdot \sigma_j \right) \right] \lambda_i \cdot \lambda_j, \quad (8)$$

$$V_{ij}^{\text{conf}} = -a_c r^2_{ij} \lambda_i \cdot \lambda_j. \quad (9)$$

The term containing $\sigma_i \cdot \sigma_j \lambda_i \cdot \lambda_j$ in eq.(8) is known as the color-magnetic interaction (CMI) term. It is important for the short-range behavior of the baryon-baryon interaction and also plays a crucial role in the short-range H–H interaction.

Flavor SU(3) symmetry breaking in the CMI is taken into account by using the parametrization:

$$\frac{\pi}{m_i m_j} \rightarrow \xi_{ij} \frac{\pi}{m_u^2},$$

where $\xi_{ij} = 1$ when both $i$ and $j$ are not an s-quark, $\xi_{ij} = \xi_1$ when either $i$ or $j$ is an s-quark, and $\xi_{ij} = \xi_2$ when both $i$ and $j$ are s-quarks. Note that we take $\xi_2$ as an independent parameter, instead of being $\xi_2^2$, which is straightforwardly expected from eq.(8). As in ref.[3], we use here the parameter set which reproduces the experimental values of the mass splittings among $\Lambda\Lambda$, $N\Xi$ and $\Sigma\Sigma$ correctly.

The other parameters entering OGEP are determined from the experimental masses of octet and decuplet baryons [3]. Their values are reproduced here in Table 1.

The equation that determines the relative wave function $\chi(r)$ is called the Resonating Group Method (RGM) equation.

$$\int \{ H_{\text{RGM}}(r, r') - E N_{\text{RGM}}(r, r') \} \chi(r') dr' = 0, \quad (10)$$

where $N_{\text{RGM}}$ ($H_{\text{RGM}}$) is the normalization and Hamiltonian kernel. These kernels are calculated as matrix elements of 1 and $H$ using the wave function of eq.(2).

Details on how to evaluate the normalization and Hamiltonian kernels can be found, for example, in refs. [10, 11]. Here, we show the spin-flavor-color (SFC) parts of the matrix elements. Because the SFC parts of the H-dibaryon wave functions are totally antisymmetrized, it is sufficient to calculate the following matrix elements:

$$\langle H^{(\sigma c)} H^{(\sigma c)} | \xi_{ij} \lambda_i \cdot \lambda_j (\text{or} \xi_{ij} \lambda_i \cdot \lambda_j \sigma_i \cdot \sigma_j) | H^{(\sigma c)} H^{(\sigma c)} \rangle$$

$$\langle H^{(\sigma c)} H^{(\sigma c)} | \xi_{ij} \lambda_i \cdot \lambda_j (\text{or} \xi_{ij} \lambda_i \cdot \lambda_j \sigma_i \cdot \sigma_j) P_{67}^{(\sigma c)} | H^{(\sigma c)} H^{(\sigma c)} \rangle$$

$$\langle H^{(\sigma c)} H^{(\sigma c)} | \xi_{ij} \lambda_i \cdot \lambda_j (\text{or} \xi_{ij} \lambda_i \cdot \lambda_j \sigma_i \cdot \sigma_j) P_{57}^{(\sigma c)} P_{68}^{(\sigma c)} | H^{(\sigma c)} H^{(\sigma c)} \rangle$$
Here, $|H^{(\sigma \tau_c)}H^{(\sigma \tau_c)}\rangle$ is the SFC part of the shell model wave function in eq. (2). There are 66 pairs interacting via the two-body interaction, $V_{ij}$, and they are classified into ten topologically different types of graphs as illustrated in Fig. 2.

To obtain, for a given number of quarks exchanged, the total contribution of each graph, we must multiply the matrix element in eq. (11) by the number of ways of choosing the interacting pair without changing the structure of the graph and by the corresponding weight of the quark exchange operators in eq. (4). After multiplying these factors, the matrix elements in SFC space can be written as

$$A + B\xi_1 + C\xi_2.$$  \hspace{1cm} (12)

In Table 2, the values $A$, $B$ and $C$ for each type of graph and each number of quarks exchanged are shown together with the multiplying factors.

It is known that the OGEP is not sufficient to describe the medium-range part of the baryon–baryon interaction, and that an attractive contribution arising from the meson cloud is also necessary. An effective meson exchange potential (EMEP) is introduced here as a force between H-dibaryons in order to simulate this meson contribution, although there are several works which introduce the meson exchange interactions as a force between quarks[12, 18]. The EMEP has previously been used to obtain a realistic description of the baryon–baryon interaction [10]. We extend the EMEP to describe the flavor-independent scalar meson exchange interaction between H-dibaryons. There are more elaborate meson exchange potentials [12], where other scalar and pseudoscalar meson exchanges are included, in addition to the simple flavour-independent scalar meson exchange used here. We shall argue later that the neglect of other meson exchanges will not substantially affect the qualitative conclusions obtained in this paper.

Here we introduce an EMEP which is assumed to be flavour-independent. Since the EMEP is expressed as a function of the relative coordinate $r$ between the H-dibaryons, we here explain how we add the EMEP to the RGM equation in (10). First, we rewrite the RGM equation so that it has the same form as the usual Schrödinger equation, albeit with a non-local Hamiltonian. After introducing the renormalized H–H relative wave function, $\tilde{\chi}(r) = \int dr' N_{\text{RGM}}^{1/2}(r, r')\chi(r')$, \hspace{1cm} (13)
and the renormalized Hamiltonian,
\[ \tilde{H}(r, r') = \int dr'' dr''' N^{-1/2}_{RGM}(r, r'') [H_{RGM}(r'', r''') - E_{\text{int}} N_{RGM}(r'', r''')] N^{-1/2}_{RGM}(r'', r'), \]
(14)
the RGM equation is equivalent to the following non-local Schrödinger equation:
\[ \int dr' \tilde{H}(r, r') \tilde{\chi}(r') = E_{\text{cm}} \tilde{\chi}(r). \]
(15)
Here \( E_{\text{int}} \) is the internal energy, and \( E_{\text{cm}} \) is the energy of the relative motion in the center-of-mass system. Then we add the EMEP, \( V(r) \), to the renormalized Hamiltonian kernel \( \tilde{H}(r, r') \), i.e., \( \tilde{H}(r, r') \rightarrow \tilde{H}(r, r') + V(r) \delta(r - r') \). For the EMEP, we take a gaussian-type function for simplicity,
\[ V(r) = V_{0\text{HH}} \exp(-r^2/\alpha_{\text{HH}}^2). \]
(16)

The parameters of \( V(r) \), \( \alpha_{\text{HH}} \) and \( V_{0\text{HH}} \), are obtained as follows. In the same manner as in ref.[3], we employ the direct convolution of the EMEP determined from the scattering length and effective range for the \( ^1S_0 \) state of the nucleon-nucleon interaction, and obtain \( \alpha_{\text{HH}} = 1.00 \text{ fm} \). The parameter \( V_{0\text{HH}} \) is fixed so that the possible range of the H-dibaryon mass is consistent with the double hypernucleus event found at KEK [7], which is interpreted as \(^{10}_{\Lambda\Lambda}\text{Be} \) or \(^{13}_{\Lambda\Lambda}\text{B} \) with a \( \Lambda\Lambda \) binding energy of \( 8.5 \pm 0.7 \) or \( 27.6 \pm 0.7 \) MeV, respectively. The corresponding range obtained for \( V_{0\text{HH}} \) is \(-1096 (\Lambda\Lambda \text{ threshold}) > V_{0\text{HH}} > -1227 \text{ MeV} (^{13}_{\Lambda\Lambda}\text{B}) [-1136 \text{ MeV} (^{10}_{\Lambda\Lambda}\text{Be})] \). On the other hand, \( V_{0\text{HH}} = -1302 \text{ MeV} \) is obtained by the direct convolution of the EMEP determined from the nucleon–nucleon interaction, which gives a rather large binding energy of the H-dibaryon from the \( \Lambda\Lambda \) threshold, 43.4 MeV. The values obtained for \( V_{0\text{HH}} \) are summarized in Table 3.

3 Numerical results and properties of the H–H interaction

In this section, we show the numerical results obtained by solving the RGM equation including the EMEP with the parameters explained in the previous section.

\[ \mu_{\text{eff}} = \frac{2}{\alpha_{\text{HH}}} \approx 400 \text{ MeV}, \]
which is not very far from the empirical \( \sigma \)-meson mass.
In Fig.3, the equivalent local potentials of bound and scattering states and the adiabatic potential for $V_{0HH} = -1227$ MeV are shown in order to appreciate the qualitative features of the H–H interaction, first.

The equivalent local potential $V_{\text{EQLP}}(r)$ [10] is defined as a potential which satisfies the following radial Schrödinger equation:

$$-\frac{\hbar^2}{2\mu} \frac{d^2}{dr^2} \tilde{\chi}_0(r) + V_{\text{EQLP}}(r) \tilde{\chi}_0(r) = E_{\text{cm}} \tilde{\chi}_0(r),$$  \hspace{1cm} (17)

where $\mu$ is the reduced mass and $\tilde{\chi}_0(r)$ is the renormalized wave function obtained by solving the RGM equation with the non-local potential. The local potential obtained by this procedure is called the trivially equivalent local potential. We will discuss another way of defining the equivalent local potential later.

The adiabatic potential $V_{\text{ad}}(r)$ is often used in order to get an idea in which regions of space repulsion and where attraction dominates. It is shown by the solid line in Fig.3. The adiabatic potential is the ratio of the diagonal element of the GCM (Generator Coordinate Method: see refs. [10, 11]) Hamiltonian kernel to the GCM normalization kernel.

$$V_{\text{ad}}(r) = \frac{H_{\text{GCM}}(r,r)}{N_{\text{GCM}}(r,r)} - H_{\text{GCM}}(r,r) N_{\text{GCM}}(r,r) \bigg|_{r \to \infty}. \hspace{1cm} (18)$$

Here the value at infinite H–H separation is subtracted, so that the $V_{\text{ad}}(r)$ goes to zero as $r \to \infty$. The GCM normalization and Hamiltonian kernels are the overlap and the expectation value of the Hamiltonian $H$ with a $(0s)^6$ configuration for each H-dibaryon, where the origins of the two H-dibaryons are separated by a distance $r$. Because the origin is not the center-of-mass of the H-dibaryon, the adiabatic potential represents a smeared H–H potential, where the center-of-mass of each H-dibaryon fluctuates. Although the local and the non-local parts of the potential are smeared out, the range of the fluctuations is sufficiently small to reflect the short-range repulsion due to the effect of the exclusion principle and the CMI.

We see from Fig.3 that the CMI leads to a strong short-range repulsion. The short range part of the EQLP, which is due to the CMI has an energy-dependent repulsion, i.e., the second bound and scattering states feel a stronger repulsion at short distances than the ground state. This energy-dependence is the typical one also seen in the nucleon–nucleon potential [13]. This behavior is shown to be partly due to the non-locality of the potential [14]. In the region below $r \lesssim 2$ fm, the contribution from the
OGEP has not yet died out and the EQLP still differs from the EMEP in the medium-range although the EMEP dominates there. Furthermore, the energy dependence seems to be reversed compared to the one at short distances. Namely, at short distances, the low energy state has a weaker repulsion than the high energy state, whereas at long range, the repulsion for the low energy state becomes stronger than the one for the high energy state. If we employ the usual procedure [13] to obtain the energy-dependent local potential from the non-local one, the ordering of the energy-dependence would be the same for short distances and long distances.

Our results for the energy-dependence of the H–H interaction can be understood in the following way. In the interacting region, the wave function \( \tilde{\chi}_0(r) \) for the attractive non-local potential is suppressed compared with the wave function \( \tilde{\chi}_{\text{EDL}}(r) \) determined from the energy dependent local (EDL) potential.

\[
\tilde{\chi}_0(r) = \sqrt{m^*(r)/m} \tilde{\chi}_{\text{EDL}}(r),
\]

where \( m^*(r) \) is the \( r \)-dependent effective mass [15]. This is known as the Perey-Buck effect [16]. In order to obtain the energy-dependent equivalent local potential, we should use \( \tilde{\chi}_{\text{EDL}}(r) \). In the actual calculation, however, we have employed \( \tilde{\chi}_0(r) \) instead of \( \tilde{\chi}_{\text{EDL}}(r) \). Then the kinetic energy given by the second derivative of the wave function \( \tilde{\chi}_0(r) \) differs from the one for \( \tilde{\chi}_{\text{EDL}}(r) \). This is the main reason for the different energy dependences at short and long distances.

The binding energy of the H–H system as a function of the parameter \( V_{0\text{HH}} \) is shown in Fig.4. The H–H system is a strongly self-binding system in the range, \(-1096 > V_{0\text{HH}} > -1227 \) MeV, and its binding energy in the ground state is 100–170 MeV. Even a second bound state, which is also an S-wave state, appears in the range \( V_{0\text{HH}} \approx -1200 \) MeV. This is also seen from the divergent behavior of the H–H scattering length \( a \) as a function of \( V_{0\text{HH}} \) shown in Fig.5, and from the change of the S-wave phase shifts with \( V_{0\text{HH}} \) shown in Fig. 6.

Fig.7 shows the equivalent hard-core radius \( a_h \), which is defined as

\[
a_h = -\frac{d\delta}{dk}, \tag{19}
\]

where \( \delta \) is the phase shift, and \( E_{\text{cm}} = \hbar^2 k^2/2\mu \). When \( k \) goes to zero, the value \( a_h \) coincides with the scattering length. For comparison, we show \( a_h \) for the case without
EMEP. For $k \gtrsim 2 \text{ fm}^{-1}$, $a_h$ is determined mainly from the repulsive core, and is of the order of $1 \text{ fm}$.

The wave function of the lowest bound state, $\tilde{\chi}_0(r)/r$, is shown for $V_{\text{HH}} = -1227 \text{ MeV}$ in Fig.8. Because of the strong repulsive core, the peak of the wave function is around $0.7 \text{ fm}$ in spite of the large binding energy of the H–H system. The wave function is clearly localized around the minimum of the EQLP. In order to see the effect of the renormalization, the unrenormalized wave function, $\chi_0(r)/r$, which does not contain the factor $N_{\text{RGM}}^{1/2}$, is also shown. The differences between the renormalized and unrenormalized wave functions can be understood as follows. The square root of the normalization kernel is expanded in terms of the harmonic oscillator wave functions, $u_{nLM}$, with the size parameter $\sqrt{2/3b}$ as

$$N_{\text{RGM}}^{1/2}(r, r') = \sum_{nLM} \sqrt{\mu_N} u_{nLM}^*(r) u_{nLM}(r').$$

(20)

Here, $\mu_N$ is the eigenvalue of the normalization kernel,

$$\mu_N = 1 - 2 \left( \frac{2}{3} \right)^N + \frac{49}{16} \left( \frac{1}{3} \right)^N - \frac{41}{32} \delta_{N0},$$

(21)

where $N = 2n + l$. The angular momentum $l$ is zero for the S-wave interaction. If $\mu_N = 1$ for all $N$ in the expansion of eq.(20), the left-hand side becomes $\delta$-function, $\delta(r - r')$. The deviation of $\mu_N$ from 1 is due to the quark exchanges between the H-dibaryons. The eigenvalues $\mu_N$ are $\frac{25}{32}$ for $n = 0$ and $\frac{65}{144}$ for $n = 1$, respectively. The large contributions from $n = 0$ and $n = 1$ cause the enhancement of the amplitude around the origin and the suppression at intermediate range by the renormalization.

These results are obtained by employing the simple flavor-singlet EMEP. Therefore, further examinations are required especially concerning the validity of the flavor independent EMEP. Here, we wish to make a brief comment on the use of the EMEP which is taken to simulate a flavor singlet scalar meson exchange. Recently, more elaborate nucleon-hyperon interactions based on the SU(6) quark model [12] have been suggested. In ref. [12], flavor-octet scalar meson exchanges are also included, in addition to the flavor-singlet scalar meson. Although we do not take into account the octet mesons in the present calculation, we qualitatively discuss their effect on the H–H system. The potential due to the flavor-octet scalar-meson exchange has the following structure:

$$\mathcal{V}(r) = \mathbf{f}_1 \cdot \mathbf{f}_2 V(r),$$
where $f_1$ and $f_2$ are flavor SU(3) operators for baryons 1 and 2, respectively. This form is based on the SU(6) quark model. In general, there are two types of meson-baryon couplings, which are called F and D. It is known that the F type coupling, which gives the above flavor-dependence for the baryon–baryon interaction, is dominant for scalar meson exchanges [12]. The expectation value of the flavor operator for the H-dibaryon is given as follows.

$$<H| f_1 \cdot f_2 |H> = \frac{1}{2}\{(f_1 + f_2)^2 - f_1^2 - f_2^2\} = -12.$$  

Here, we have used that $<(f_1 + f_2)^2 >= 0$ for the H-dibaryon and that the H-dibaryon consists of flavor-octet baryons for which $<f_i^2 >$ is 12. Thus, the contributions from the flavor-octet scalar-meson exchanges in the H-dibaryon are repulsive because the radial part $V(r)$ is always attractive for scalar meson exchanges. As a result, if we include the flavor-octet mesons and fix the strength of the EMEP from the binding energy of the H-dibaryon, the strength of the flavor-singlet part becomes more attractive than in the present calculation without the flavor-octet mesons. For the H–H system, the flavor-octet meson exchanges do not contribute because the H-dibaryon is a flavor SU(3) singlet. Therefore, if the flavor-octet mesons were included, the H–H system would be more strongly bound.

For the nucleon-nucleon system, octet scalar meson exchange is attractive in the $^1S_0$ channel. We expect, therefore, that the strength of the flavor-singlet part of the EMEP derived from the nucleon–nucleon potential is decreased compared to the case without octet mesons. On the other hand, as discussed before, the strength of the flavor-singlet part of the EMEP as derived from the binding energy of the H-dibaryon is increased when the octet mesons are taken into account. Thus, the strengths derived from the nucleon–nucleon scattering data on one hand, and from the H-dibaryon binding energy on the other hand, would become similar when we include the octet scalar meson exchanges. Recall that without the octet scalar mesons, the EMEP derived from nucleon-nucleon scattering was more attractive ($-1302$ MeV) than the EMEP derived from the limits on the binding energy of the H-particle ($-1227$ MeV $\sim -1096$ MeV).

Finally, we comment on the effect of pseudoscalar meson exchange. The pseudoscalar meson exchanges do not contribute to the H–H system. The contribution from pseudoscalar meson exchanges to the binding energy of the H-dibaryon is known
to be about 5 MeV [17, 18]. Therefore, even if we take into account the pseudoscalar meson exchanges, there is no drastic change for the binding energy of the H–H system. Thus, the conclusion that the H–H system has a strong medium-range attraction leading to a bound state remains unchanged.

A bound state of two H-dibaryons is also obtained in the Skyrme model[19]. The binding energy of the “tetralambda” state is $E_B = 15 \sim 20$ MeV, which is smaller than the value obtained in our calculation.

Finally, we comment on the implications of our results for the occurrence of H-matter. Tamagaki [8] discussed the possibility of a phase transition from neutron matter to H-matter at a density which is $6 \sim 9$ times greater than the normal nuclear density $\rho_0$. In Tamagaki’s calculation, the H-dibaryons interact via a hard core potential and an attractive square well potential outside the core. In their work, the depth of the attractive potential was assumed to be so weak that it can be treated as a perturbation. The depth of the square well potential has been determined as the value where the scattering length becomes zero for the first time, when the strength of the attractive potential is gradually increased, i.e. as the limiting strength that still gives a positive scattering length. The pertinent strength parameter corresponding to our EMEP can be obtained from Fig. 5. It is found to be $V_{\text{HH}} = -638$ MeV. Thus the attraction used in Ref. [8] is much weaker than the one used in the present calculation. If the attractive H–H potential is indeed as strong as the EMEP employed in the present calculation, the critical transition density beyond which H-matter formation is energetically favorable may be appreciably lower. However, a quantitative estimate is not simple because of the inapplicability of perturbation theory under a strong attractive potential as in the present calculation. A recalculation of the critical transition density for H-matter formation is beyond the scope of the present work (see however ref.[20]).

4 Summary

We have investigated the interaction between two H-dibaryons in the quark cluster model with a one-gluon-exchange and quadratic confinement potential between constituent quarks. In addition, a phenomenological attractive meson exchange potential between H-dibaryons is used. The parameters employed for the calculation are de-
determined so as to reproduce the octet and decuplet baryon ground state masses and from the requirement that the H-dibaryon mass is consistent with the observed binding energy of two Λ's in a double hypernucleus. These parameters were also used in our previous work on the N–H interaction [3, 4]. In our calculation, the effective scalar meson exchange potential is assumed to be a flavor SU(3) singlet. We have presented a qualitative discussion on the effects of octet scalar mesons and pseudoscalar mesons. We conclude that the following properties of the H–H system remain unchanged even if we include the octet scalar and pseudoscalar mesons.

The main properties of the interaction between H-dibaryons can be characterized as a short-range repulsion due to Pauli blocking and the color-magnetic interaction, and a medium-range attraction due to flavor singlet scalar meson exchange. As a result of the present RGM calculation, we have obtained a strongly-bound state of two H-dibaryons with a separation about 0.8 ∼ 0.9 fm due to the strong repulsive core.

The present results suggest that the critical density at which the formation of H-matter becomes energetically favorable may be lower than the value $\rho_{\text{crit}} = 6\rho_0$ found in Ref. [8]. This would have interesting astrophysical consequences. In the framework of the Walecka model, it has recently been shown [20] that with the present H–H interaction, H-matter is unstable against compression. Thus, if the central density of a massive neutron star exceeds the critical density for H-matter formation, the energetically favorable compression of H-matter provides a possible scenario for the conversion of a neutron star into a strange quark star.

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Figure 1: Quark exchange diagrams between two H–dibaryons

Figure 2: Ten types of two-body interaction with quark exchange
Figure 3: The equivalent local potential (EQLP) for bound states and a scattering state of the H–H system, and the adiabatic potential for $V_{0HH} = -1227$ MeV. The solid curve denotes the adiabatic potential, the dashed curve denotes the EQLP of the lowest bound state, the chain curve denotes the EQLP of the second bound state, the double-dotted chain curve denotes the EQLP of the scattering state with center-of-mass energy $E_{cm} = 50$ MeV.
Figure 4: The binding energy of the H–H system as a function of the strength parameter of the effective meson exchange potential $V_{0HH}$. Both bound states are S-wave states. The vertical dotted lines show the range of $V_{0HH}$ from $-1227$ to $-1096$ MeV.

Figure 5: The H–H scattering length $a$ as a function of the strength parameter of the effective meson exchange potential $V_{0HH}$. $a = -1.26, -4.91, 6.12$ and $3.51$ fm for $V_{0HH} = -1096, -1136, -1227$ and $-1302$ MeV, respectively. The vertical dotted lines show the range of $V_{0HH}$ from $-1227$ to $-1096$ MeV.
Figure 6: The HH S-wave phase shift $\delta$ as a function of the relative wave number $k$ for several values of the strength parameter of the effective meson exchange potential $V_{0HH}$.

Figure 7: The equivalent hard core radius as a function of the relative wave number $k$ for several values of the strength parameter of the effective meson exchange potential $V_{0HH}$. 
Figure 8: The wave function of the lowest S-wave bound state for $V_{0\text{HH}} = -1227$ MeV. The solid curve denotes the wave function without the renormalization $\chi_0(r)/r$ and the dashed curve denotes the renormalized wave function $\bar{\chi}_0(r)/r$. The root-mean-square radii of the H–H system are 0.92, 0.89, 0.84 and 0.81 fm for $V_{0\text{HH}} = -1096, -1136, -1227$ and $-1302$ MeV, respectively.
Table 1: Quark model parameters

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<th>$b$ (fm)</th>
<th>$m$ (MeV/c$^2$)</th>
<th>$\alpha_a$</th>
<th>$a_c$ (MeV/fm$^2$)</th>
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<th>$\xi_2$</th>
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Table 2: The matrix elements of spin-flavor-color parts

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<th>interaction type (factor)</th>
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<th>$\lambda_i \cdot \lambda_j$</th>
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Table 3: Strength parameter of the effective meson exchange potential

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<th>$V_{0HH}$ (MeV)</th>
<th>$2M_\Lambda - M_H$ (MeV)</th>
<th>[\Lambda\Lambda] threshold</th>
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<td>$^{13}\Lambda\Lambda$ B</td>
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