Five-flavor pentaquarks and other light- and heavy-flavor symmetry partners of the LHCb hidden-charm pentaquarks

Fang-Zheng Peng,1 Ming-Zhu Liu,1,2 Ya-Wen Pan,1 Mario Sánchez Sánchez,3 and Manuel Pavon Valderrama1

1School of Physics, Beihang University, Beijing 100191, China
2School of Space and Environment, Beihang University, Beijing 100191, China
3Centre d'Études Nucléaires, CNRS/IN2P3, Université de Bordeaux, 33175 Gradignan, France

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The discovery of three pentaquark peaks — the \(P_c(4312)\), \(P_c(4440)\) and \(P_c(4457)\) — by the LHCb collaboration has a series of interesting consequences for hadron spectroscopy. If these hidden-charm objects are indeed hadronic molecules, as suspected, they will be constrained by heavy-flavor and SU(3)-flavor symmetries. The combination of these two symmetries will imply the existence of a series of five-flavor pentaquarks with quark content \(b\bar{c}sdu\) and \(b\bar{c}sd\), that is, pentaquarks that contain each of the five quark flavors that hadronize. In addition, from SU(3)-flavor symmetry alone we expect the existence of light-flavor partners of the three \(P_c\) pentaquarks with strangeness \(S = -1\) and \(S = -2\). The resulting structure for the molecular pentaquarks is analogous to the light-baryon octet — we can label the pentaquarks as \(P^{D^{+}}_{Q'Q''}, P^{D^0}_{Q'Q''}, P^{D^-}_{Q'Q''}, P^{D^0}_{Q'Q''}\) depending on their heavy- and light-quark content (with \(N, \Lambda, \Sigma, \Xi\) the member of the light-baryon octet to which the heavy-quark structure resembles and \(Q', \bar{Q}\) the heavy quark-antiquark pair). In total we predict 45 new pentaquarks from heavy- and light-flavor symmetries alone, which extend up to 109 undiscovered states if we also consider heavy-quark spin symmetry. If an isouquet (\(I = 3/2\)) hidden-charm pentaquark is ever observed, this will in turn imply a second multiplet structure resembling the light-baryon decuplet: \(P^{\Delta}_{Q'Q''}, P^{\Delta^+}_{Q'Q''}, P^{\Delta^0}_{Q'Q''}, P^{\Delta^-}_{Q'Q''}\).

1. INTRODUCTION

The discovery by the LHCb collaboration of three hidden-charm pentaquarks \([1]\) — the \(P_c(4312)\), \(P_c(4440)\) and \(P_c(4457)\) — extends the previous observation of the \(P_c(4450)\) peak in 2015 [2]. Their masses and widths (in MeV) are

\[
m_{P_{c1}} = 4311.9 \pm 0.7^{+0.8}_{-0.6}, \quad \Gamma_{P_{c1}} = 9.8 \pm 2.7^{+3.7}_{-4.5}, \quad (1)
m_{P_{c2}} = 4440.3 \pm 1.3^{+1.4}_{-1.7}, \quad \Gamma_{P_{c2}} = 20.6 \pm 4.9^{+8.7}_{-10.1}, \quad (2)
m_{P_{c3}} = 4457.3 \pm 0.6^{+0.4}_{-1.7}, \quad \Gamma_{P_{c3}} = 6.4 \pm 2.0^{+5.7}_{-1.9}, \quad (3)
\]

where from now on we will use the notation \(P_{c1}, P_{c2}\) and \(P_{c3}\) for these three pentaquarks. The \(P_{c1}\) is 8.9 MeV below the \(\bar{D}\Sigma\) threshold, while the \(P_{c2}\) and \(P_{c3}\) are 21.8 and 4.8 MeV below the \(\bar{D}'\Sigma\) threshold, respectively (where we have considered these thresholds in the isospin-symmetric limit). This, together with the existence of hidden-charm pentaquark predictions in the molecular picture before their experimental observation \([3, 9]\), suggests a molecular interpretation of these pentaquarks, i.e. that they are bound states of a charmed antimeson and a charmed baryon \([10–18]\), though this is not the only explanation that has been considered by theoreticians \([14, 18]\).

Heavy-hadron molecules are highly symmetrical: their light- and heavy-quark content implies that they are constrained both by SU(3)-flavor symmetry \([19, 20]\) and heavy-quark symmetry \([21, 22]\). Heavy-quark symmetry has in turn different manifestations, namely heavy-quark spin symmetry (HQSS), heavy-flavor symmetry (HFS) and heavy-antiquark-diquark symmetry (HADS) \([23]\), which altogether provide deep insights into the molecular spectrum \([24, 25]\). The application of HQSS to the particular case of the LHCb pentaquarks implies that the \(P_{c1}, P_{c2}\) and \(P_{c3}\) actually belong to a multiplet composed of seven members \([12, 14, 33, 34]\), four of which have not been observed yet. Before knowing that the \(P_c(4450)\) peak contained two peaks, HQSS was already used to predict a \(J^P = 5/2^-\) \(\bar{D}\Sigma^*\) molecular pentaquark and other partner states \([6, 35–38]\). In the past HFS and HADS have been applied to heavy meson-antimeson molecules to explain spectroscopic relations among known molecular states \([30]\) or to deduce the existence of new states \([31]\). In this manuscript we will explore what are the consequences of SU(3)-flavor symmetry and HFS if the hidden-charm pentaquarks are indeed molecular.

2. SYMMETRIES

First, we will consider the constraints that HFS and SU(3)-flavor symmetry impose on the potential between a heavy antimeson and a heavy baryon. HFS refers to the fact that the structure of a heavy-light hadron (i.e. the “brown muck” around the heavy quark) is independent of the flavor of the heavy quark. As applied to heavy-hadron molecules, HFS implies that the potential among heavy hadrons is independent of the flavor of the heavy quarks inside the heavy hadrons. The clearest example of this symmetry in molecular states are the \(Z_s\)’s and \(Z'_s\)’s resonances \([30]\), which are repeated in the charm and bottom sectors and are conjectured to be \(D^{(*)}\bar{D}^*\) and \(B^{(*)}\bar{B}^*\) bound states, respectively. If applied to the molecular pentaquarks, from HFS we expect the potentials in the \(\bar{D}\Sigma, \bar{D}\Sigma_b, B\Sigma\) and \(B\Sigma_b\) two-body systems to be identical (plus similar relations for the \(\bar{D}\Sigma^*, \bar{D}'\Sigma\) and \(D^*\Sigma\) family of molecules). For simplicity we will often use the generic notation \(P\) and \(P'\) for the \(J^P = 0^-\), \(1^-\) heavy mesons and \(\Sigma_Q\) and \(\Sigma_Q'\) for the \(J^P = 1/2^+\) and \(3/2^+\) heavy baryons, irrespective of whether they are their charm or bottom versions. In addition we will use the notation \(P'_s\) for the heavy mesons...
TABLE I. The SU(3)-flavor structure of the potential for heavy meson-baryon molecules, where the heavy meson belongs to a SU(3)-flavor triplet and the heavy baryon to a sextet. The heavy meson-baryon potential can be decomposed into an octet and decuplet component, from which the octet piece corresponds to the potential for the hidden-charm molecular candidates. As a consequence other molecular pentaquarks belonging to the octet representation are also expected to bind. In addition to the SU(3)-flavor decomposition, the S-wave potential can be further decomposed into its light-quark structure, which is not explicitly shown here.

| Molecule | I | S | V | V_{eigen} |
|----------|---|---|---|----------|
| $P \Sigma_Q$ | 1/2 | 0 | $V^O$ | - |
| $P \Xi_Q$ | 1/2 | 0 | $V^D$ | - |
| $P \bar{\Xi}_Q$ | 0 | -1 | $V^O$ | - |
| $P \bar{\Omega}_Q - P \Sigma_Q$ | 1 | -1 | $\frac{1}{2} V^O + \frac{3}{2} V^D$ | - |
| $P \bar{\Omega}_Q - P \Xi_Q$ | 1/2 | -2 | $\frac{3}{4} (V^O - V^D)$ | - |
| $P \bar{\Omega}_Q - P \Xi_Q$ | 0 | -3 | $V^D$ | - |

with $S = 1$ and $\Xi_Q$, $\Xi_Q^*$ ($\Omega_Q$, $\Omega_Q^*$) for the heavy baryons with $S = 1$ ($S = -1$).

If we now consider SU(3)-flavor symmetry instead, it happens that the $\bar{P}$, $\bar{P}$ heavy antimesons and the $\Sigma_Q$, $\Xi_Q$ and $\Omega_Q$ heavy baryons belong to the 3 and 6 representation of the SU(3)-flavor group, respectively. Two-body heavy antimeson-baryon states can be decomposed into $3 \otimes 6 = 8 \oplus 10$, i.e. into the octet and decuplet representations, where the SU(3) Clebsch-Gordan coefficients can be consulted in Ref. [39]. This octet and decuplet decomposition is not dependent on the nature of the pentaquarks, but on their light-quark content, and it has indeed been previously pointed out for compact pentaquarks [40]. Within the molecular explanation, this decomposition specification implicitly that the heavy antimeson-baryon potential can be decomposed into a linear combination of an octet and decuplet contribution

$$V = \lambda^O V^O + \lambda^D V^D,$$

with $V^O$ and $V^D$ the octet and decuplet pieces and $\lambda^O$, $\lambda^D$ numerical coefficients. We show the full decomposition in Table I which happens to be surprisingly simple: for most heavy antimeson-baryon molecules, the potential is a pure octet or decuplet contribution. In turn, this is easily explained from the observation that the resulting pentaquarks have the same quantum numbers as the corresponding octet or decuplet light baryons. Even for the $P \Xi_Q - P \Sigma_Q$ and $P \bar{\Omega}_Q - P \Xi_Q$ molecules (where the dash indicates that these channels couple), for which the potential is a $2 \times 2$ matrix, when we look at the eigenvalues we recover

$$V = \{ V^O, V^D \},$$

depending on the linear combination of the two channels, with the octet eigenvalue corresponding to

$$|8\rangle = -\sqrt{\frac{1}{3}} |\bar{P} \Xi_Q^* (I = 1)\rangle + \sqrt{\frac{2}{3}} |\bar{P} \Sigma_Q\rangle,$$  

$$|8\rangle = -\sqrt{\frac{1}{3}} |\bar{P} \Omega_Q\rangle + \sqrt{\frac{2}{3}} |\bar{P} \Xi_Q\rangle,$$  

and the decuplet eigenvalue to

$$|10\rangle = \sqrt{\frac{2}{3}} |\bar{P} \Xi_Q^* (I = 1)\rangle + \sqrt{\frac{1}{3}} |\bar{P} \Xi_Q\rangle,$$  

$$|10\rangle = \sqrt{\frac{2}{3}} |\bar{P} \Omega_Q\rangle + \sqrt{\frac{1}{3}} |\bar{P} \Xi_Q\rangle.$$

These two molecular systems, $\bar{P} \Xi_Q^* - \bar{P} \Sigma_Q$ and $\bar{P} \Omega_Q - \bar{P} \Xi_Q$, will adopt the lowest-energy configuration, be it either the octet or decuplet one. In the absence of additional experimental information and knowing that the $P_{c1}$, $P_{c3}$ and $P_{c3}$-charm pentaquarks most probably belong to the octet, we naively expect the lowest-energy configuration to be the octet.

Owing to heavy-flavor symmetry, the potential is expected to be independent of the flavor of the heavy quarks. This implies in particular that the octet configurations

$$\bar{D} \Xi'_c (I = 0), \quad \bar{D} \Xi'_c (I = 1) - \bar{D} \Sigma_b,$$  

$$B \Xi'_c (I = 0), \quad B \Xi'_c (I = 1) - B \Sigma_c,$$

which contain the five quark flavors that hadronize, will display as much attraction as the hidden-charm pentaquarks. Out of the four five-flavor configurations, the strange-isoscalar molecules $[\bar{D} \Xi'_c (0), B \Xi'_c (0)]$ are relatively easy to deal with (they are single-channel systems). For the strange-isovector molecules $[\bar{D} \Xi'_c (I = 1) - \bar{D} \Sigma_b, B \Xi'_c (I = 1) - B \Sigma_c]$ we have a two-channel problem where the thresholds are separated by about 20 MeV and 40 MeV for the isovector $b \bar{c} s q q$ and $b \bar{c} s q q$ pentaquark configurations, respectively. The question is whether this energy gap will prevent a predominantly octet molecular state to form or not. The answer depends on the comparison of the momentum scales of the binding mechanism and the coupled-channel dynamics. The typical momentum scale of the coupled channels in the previous cases is about 250 MeV and 350 MeV for the $b \bar{c} s u d$ and $b \bar{c} s u d$ pentaquarks, while the binding mechanism is expected to be short-ranged (e.g. vector-meson exchange), with a momentum scale of the order of $(0.5 - 1.0)$ GeV give or take. As a consequence, we expect the isovector five-flavor pentaquarks to bind (a conjecture which we confirm by means of concrete calculations in what follows).
3. EFFECTIVE FIELD THEORY DESCRIPTION

To explicitly check the effects of the previous symmetries, we will describe the pentaquarks as non-relativistic meson-baryon bound states interacting by means of a contact-range potential that is heavy- and SU(3)-flavor symmetric.

This choice is not arbitrary, but corresponds with the lowest or leading order (LO) effective field theory (EFT) description of the heavy antimeson and heavy baryon two-body system. EFTs exploit the existence of a separation of scales to formulate generic low energy descriptions of physical systems. The idea is to identify characteristic low and high energy scales $Q$ and $M$ such that $Q/M \ll 1$ and then express every physical quantity as a power series in terms of the ratio $Q/M$. The first term in this series is the LO, the second is the next-to-leading order (NLO), and so on.

For molecular pentaquarks the required scale separation manifest itself as follows: the typical low energy scale $Q$ is of the order of $(100-200)$ MeV and can be identified with the pion mass or the binding momentum of the pentaquarks. At this scale the meson-baryon dynamics is well known and involves the exchanges of pions and other pseudoscalar mesons. The high energy scale $M$ is in the $(0.5-1.0)$ GeV range and can be identified with the rho meson mass or the momentum scale at which the internal structure of the hadrons becomes evident. This part of the interaction is less well-known and might very well involve non-molecular components of the pentaquark wave function. EFT parametrizes it as a series of contact-range operators.

Our LO description of the pentaquarks only involves the contact-range potential. This choice is justified (i) from a well-known EFT observation that indicates that the existence of shallow bound states (e.g. the deuteron or near-threshold states such as hadronic molecules) increases the importance of contact-range interactions at low energies and (ii) from concrete EFT calculations for the LHCb pentaquarks that suggest that pion exchanges are NLO and thus a perturbative correction to the LO results.

From the previous, the LO S-wave interaction binding the $P_{c1}$, $P_{c2}$ and $P_{c3}$ molecular pentaquarks will be given by the Lagrangian

$$L_{\text{contact}} = C_i^O \sum_{IS} \langle \bar{q} IS \bar{u} (\gamma_5) p^I_3 b^c \rangle \langle \bar{q} IS \bar{u} (\gamma_5) p^I_3 b^c \rangle,$$  \hspace{1cm} (12)

where $C_i^O$ is the (octet) coupling constant, $i = 1, 2, 3$ is the index with which we label the hidden-charm pentaquarks, $M_a$ is a triplet heavy meson with the quark content $(\bar{Q}_a q_a)$, where $q_a = u, d, s$ depending on the flavor index $a$, $b^c$ is a sextet heavy baryon with quark content $(\bar{Q}_a b^c q_b + \bar{Q}_b b^c q_a)$ (i.e. symmetric in the flavor indices), $\pi_{IS}^{abc}$ is a tensor in flavor space that projects the heavy antimeson-baryon system in an octet state with given isospin $I$ and strangeness $S$ (the exact form of this tensor can be deduced from Table D, and $P^I_3$ is a projector into the corresponding spin channel $J$ if there is more than one.\footnote{The form of this projector is trivial ($P_1 = 1$) for the $P_{c1}$ pentaquark, while $J = \frac{1}{2}$, while for the $P_{c2}$ and $P_{c3}$ it will be either $J = \frac{1}{2}$ or $\frac{3}{2}$, though we do not know which of these two pentaquarks corresponds to each of the two possible spin configurations. We are also assuming that the decuplet contact-range interaction is subleading, which is why it is not included in the Lagrangian above.}

The previous Lagrangian generates a simple contact-range potential of the type

$$\langle p' | V | p \rangle = C_i^O (\Lambda) f(p^I_3) f(p'^I_3),$$  \hspace{1cm} (13)

where we have regularized the potential, originally a Dirac delta in momentum space, with the Gaussian regulator $f(x) = e^{-x^2}$ and a cutoff $\Lambda$. For the cutoff we choose the range $\Lambda = (0.5 - 1.0)$ GeV, i.e. around the $\rho$ meson mass. With this potential we solve a coupled-channel Lippmann-Schwinger equation of the type

$$\phi_A(p) + \sum_b \int \frac{d^3 p}{(2\pi)^3} (k | V_{AB} | p) \phi_B(p) = 0,$$  \hspace{1cm} (14)

where $A, B$ are indices for the channels we are considering, $\phi_A$ the vertex function (i.e. the wave function $\Psi_A$ times the propagator, $\phi_A(p) = [M_A + p^2/(2\mu_A) - M_F] \Psi_A(p)$), $V_{AB}$ the potential between channels $A$ and $B$, $M_B$ the total mass of the heavy antimeson and baryon comprising channel $B$, $\mu_B$ their reduced mass and $M_F$ the mass of the molecular pentaquark we are predicting. We notice that the only configurations with more than one channel are the $(I, S) = (1, -1)$ and $(\frac{1}{2}, -2)$, see Table D. For illustrative purposes we consider the bound-state equation for a Gaussian regulator in the single-channel case, in which it reduces to

$$1 + C_i^O (\Lambda) \frac{\mu A}{4\pi} I_0(\gamma_A, \Lambda) = 0,$$  \hspace{1cm} (15)

with $\gamma_A = \sqrt{2\mu_A (M_F - M_B)}$ the wave number of the molecular pentaquark and where $I_0$ is given by

$$I_0(\gamma_A, \Lambda) = \sqrt{2\pi} \Lambda - 2 \gamma_A^2 / \Lambda^2 \pi \gamma_A \text{erfc} \left( \frac{\sqrt{2} \gamma_A}{\Lambda} \right),$$  \hspace{1cm} (16)

where erfc$(x)$ is the complementary error function.

If we determine the $C_i^O$ couplings from reproducing the masses of the $i = 1, 2, 3 \ P_{c1}$ pentaquark, for $\Lambda = 0.75$ GeV we obtain the couplings

$$C_1^O = -1.19 \ (-(2.17 - 0.80)) \ \text{fm}^2,$$  \hspace{1cm} (17)

$$C_2^O = -1.44 \ (-(2.88 - 0.93)) \ \text{fm}^2,$$  \hspace{1cm} (18)

$$C_3^O = -1.02 \ (-(1.80 - 0.71)) \ \text{fm}^2,$$  \hspace{1cm} (19)

$P_2$ and $P_3$ depend on the spin of the $P_{c2}$ and $P_{c3}$ pentaquarks, which is either $J = \frac{1}{2}$ or $\frac{3}{2}$, where the projector for the $|JM\rangle$ spin configuration in the $D^* \Sigma_c$ system takes the form $\langle \ell | m_1 | P_{c1} | m_2 \rangle = \langle \ell m_1 m_2 | J M \rangle$, i.e. it coincides with the Clebsch-Gordan coefficients coupling a $D^*$ meson and $\Sigma_c$ baryon with spin wave functions $|m_1\rangle$ and $|m_2\rangle$ to total spin $|J M\rangle$.\footnote{The form of this projector is trivial ($P_1 = 1$) for the $P_{c1}$ pentaquark, while $J = \frac{1}{2}$, while for the $P_{c2}$ and $P_{c3}$ it will be either $J = \frac{1}{2}$ or $\frac{3}{2}$, though we do not know which of these two pentaquarks corresponds to each of the two possible spin configurations. We are also assuming that the decuplet contact-range interaction is subleading, which is why it is not included in the Lagrangian above.}
where the values in parentheses correspond to varying the cutoff in the (0.5 – 1.0) GeV window. With these couplings, for \( \Lambda = 0.75 \text{ GeV} \) we predict the location of the \( \bar{c}b \) five-flavor pentaquarks to be

\[
\begin{align*}
& m(P_{\bar{c}b}^5) = 7783^{+5}_{-6}, \quad 7907 \pm 7, \quad 7930^{+2}_{-3} \text{ MeV}, \quad (20) \\
& m(P_{\bar{c}b}^{2*}) = 7765^{+6}_{-7}, \quad 7892^{+8}_{-9}, \quad 7914^{+5}_{-4} \text{ MeV}, \quad (21)
\end{align*}
\]

where the uncertainty comes from varying the cutoff (i.e. taking \( \Lambda = (0.5 – 1.0) \text{ GeV} \)), but does not include the SU(3) symmetry breaking effects, which we discuss later. For the \( \bar{c}b \) five-flavor pentaquarks we predict instead

\[
\begin{align*}
& m(P_{\bar{c}b}^5) = 7829^{+10}_{-9}, \quad 7858^{+12}_{-10}, \quad 7883^{+10}_{-12} \text{ MeV}, \quad (22) \\
& m(P_{\bar{c}b}^{2*}) = 7804^{+6}_{-7}, \quad 7835^{+8}_{-7}, \quad 7858^{+8}_{-7} \text{ MeV}. \quad (23)
\end{align*}
\]

The complete list of predictions (including not only cutoff but also SU(3)-flavor uncertainties) can be consulted in Table II.

The spectrum of Table II implies that each of the observed hidden-charm pentaquarks belongs to a light/heavy-flavor multiplet with 16 members. As three hidden-charm pentaquarks have been observed, this means a total of 48 states (of which 45 are so far unobserved). The experimental observation of these pentaquarks could be achieved by means of the SU(3)-flavor and HFS analogues of the \( J/\psi N \) decay channel that has been used in the discovery of the \( P_{c1}, P_{c2} \) and \( P_{c3} \). For instance, the five-flavor pentaquarks \( P_{\bar{c}b}^5 \) and \( P_{\bar{c}b}^{2*} \) could be detected by means of their \( B_{c}^* \Lambda \) and \( B_{c}^* \Sigma \) decays.

Even though for the moment we have not considered HQSS explicitly, it is easy to figure out its consequences: from HQSS we expect the hidden-charm pentaquarks to come in multiplets of up to seven members. Within the scope of contact-range EFT incorporating HQSS, the observation of the \( P_{c1}, P_{c2} \) and \( P_{c3} \) pentaquarks suggests that the aforementioned septet is probably complete, meaning that there are 4 unobserved states. This result is reproduced in most schemes that include HQSS, e.g. models with a compact core coupled to the molecular degrees of freedom, indicating that it depends on HQSS instead of the specific dynamics generating the pentaquarks. The bottom-line is that if we compound the HQSS multiplets with the SU(3)-flavor and HFS ones, the heavy molecular pentaquark family could contain a total of 112 states (3 observed, 109 to be discovered), as we will discuss later.

Among the results in Table II it is interesting to notice the strange-isoscalar \( P_{c}^5 \) partners of the three LHCb pentaquarks, which were predicted (together with the pentaquarks) nearly a decade ago. This prediction has been recently updated in Ref. [49], which uses a contact-range theory where the couplings are saturated by vector-meson exchange and the regularization is set as to reproduce the \( P_{c}(4312) \) pentaquark. The prediction of Ref. [49] for the mass of the \( \bar{D}E \) molecule is 4436.7 MeV, which happens to be pretty close to ours (check Table II).

4. UNCERTAINTIES

We are predicting the molecular pentaquarks within a contact-range EFT, which entails that they are amenable to systematic error estimations. A conventional way to estimate these theoretical errors is to vary the predictions within a sensible cutoff window (which is what we have done for the five-flavor pentaquarks in Eqs. [20-23]). Here the cutoff floats from 0.5 to 1 GeV, which can be either identified with the mass of the vector mesons or with the momenta at which the internal structure of the hadrons starts to be resolved. For the \( c\bar{c} \) family of pentaquarks this translates into a systematic error of less than 1 MeV, which explains why the predictions of other theoretical works [49-51] are basically identical to ours. Yet this uncertainty is calculated under the assumption that SU(3)-flavor symmetry is perfectly preserved, which is not the case. Violations of SU(3)-flavor symmetry relations are usually of the order of 20%, as estimated from the difference between the pion and kaon weak decay constants (\( f_{\pi} \approx 130 \text{ MeV} \) and \( f_{K} \approx 160 \text{ MeV} \)). From this, within the EFT we are using we can be easily take into account the SU(3)-flavor symmetry breaking effects by randomly varying the \( C^{0} \) couplings by 20% around their central values. For \( \Lambda = 0.75 \text{ GeV} \), this translates into an uncertainty of 2 – 15 MeV depending on the specific \( c\bar{c} \) pentaquark, where the largest uncertainties correspond to the states with the largest binding energies.

For the \( \bar{c}b, cb \) and \( bb \) molecular pentaquarks the situation is different owing to the considerably larger cutoff dependence (about 5, 10 and 20 – 30 MeV respectively), which we will discuss in the next paragraph. The SU(3)-flavor uncertainties in these cases will be 10 – 20 and 15 – 25 MeV for the \( \bar{c}b/cb \) and \( \bar{b}b \) cases, respectively. That is, while for the \( c\bar{c}, cb, \bar{c}b \) the uncertainties are dominated by flavor symmetry breaking effects, for the \( bb \) pentaquarks cutoff variation tends to be the largest source of uncertainty.

However, the application of SU(3)-flavor symmetry remains theoretical in the sense that we do not really have a clear molecular example from where we can determine how well this symmetry works at the quantitative level. Two qualitative examples are already known:

\[ \text{Footnote 5: For simplicity, we have not considered the errors stemming from the uncertainties in the pentaquark masses, see Eqs. (10), nor from the further dependence of these masses on the resonance profile, check for instance Ref. [45] in which the } P_{c}(4312) \text{ is found to be a virtual (instead of a bound) state.} \]
The mass of the corresponding heavy antimeson baryon, for which we take the isospin symmetric limit of the masses listed in the Review of Particle Physics (RPP) [68] and “Partner” represents which hidden-charm pentaquark (\(P_{i}, i = 1, 2, 3\)) is the partner of the predicted state.

The coupled-channel cases, the binding energy is calculated relative to the channel with the lowest mass. For the calculations we use a contact-range EFT, with the potential of Eq. (11) and a Gaussian regulator with a cutoff \(\Lambda = 0.75\) GeV. The error comes from two different sources, which are added in quadrature: (i) varying the cutoff in the \(\Lambda = (0.5 - 1.0)\) GeV range and (ii) assuming a 20% uncertainty in SU(3)-flavor symmetry as applied to the contact-range couplings (this second error only pertains pentaquarks with strangeness). In general the SU(3)-flavor uncertainty dominates in the \(c\bar{c}, c\bar{b}, b\bar{b}\) sectors, while for the \(b\bar{b}\) pentaquarks the bulk of the errors come from the cutoff variation (in agreement with theoretical expectations [47]).

(i) The \(Z_{c} (3900)\) [58] and \(Z_{c} (3895)\) [59] (\(Z_{c}\) and \(Z_{c}^\prime\) from now on), which have been theorized to be \(I = 1\) \(D^{\prime}\bar{D}\) [31, 60, 61] and \(I = \frac{1}{2} D^{\prime}D - D, D^{\prime}\) \(62, 63\) molecules, respectively.

(ii) The \(P_{c1}(4459)\) pentaquark [52], which has been theorized to be an \(I = 0\) \(D\bar{Z}_{c}\) bound state [33, 50].

In the first case, the SU(3) decomposition of heavy meson-antimeson states is \(3 \otimes 3 = 1 \oplus 8\), i.e. a singlet and an octet representation, where the \(Z_{c}\) and \(Z_{c}^\prime\) both belong to the octet and thus their potential is expected to be the same [22, 62]. But it happens that the masses of the \(Z_{c}\) and \(Z_{c}^\prime\) resonances are above their corresponding meson-antimeson thresholds, which means that they are not necessarily bound states but more probably resonances (or even virtual states if we take into account that their Breit-Wigner masses might not correspond to their physical masses). This happens to be the case, they will require a different contact-range EFT description than the one we employ here for the pentaquarks (or the direct extraction of the couplings from the data instead of the masses, as done in Refs. [61, 62]), which renders it difficult to make direct comparisons between the \(Z_{c}\)’s and the \(P_{c1}\)’s.

In the second case, as pointed out previously, the \(\Xi_{c}\) charmed baryon is a flavor antitriplet and the \(D^{\prime}\bar{Z}_{c}\) system will essentially belong to a different and independent representation of SU(3). That is, the \(D^{\prime}\bar{Z}_{c}\) potential can be described with a new coupling constant \(D(\Lambda)\), i.e.

\[
\langle p' | V | p \rangle = D(\Lambda) f(\frac{P_{c1}}{\Lambda}) f(\frac{p'}{\Lambda}),
\]

the value of which is in principle unrelated to the \(C(\Lambda)\) couplings we have used to reproduce the three \(P_{c1}\) pentaquarks. However, phenomenological models based on vector-meson exchanges predict that \(D = C(\Lambda) [3, 4]\), i.e. the \(I = 0\) \(D\bar{Z}_{c}\) and \(I = \frac{1}{2}\) \(D\bar{Z}_{c}\) potentials are expected to be similar. Concrete
calculations with the same type of EFT, regulator and cutoff range we have used for the $P_{c1}, P_{c2}$ and $P_{c3}$ yield $D = 1.17 C_1^0$ when calibrating $D(\Lambda)$ to the $P_{c3}(4459)$ mass, showing a 17\% discrepancy from $D = C_1^0$. The more complete analysis of Ref. \[54\] (which includes a series of effects not considered here, like coupled channel dynamics or the double-peak solution considered in the experimental analysis of Ref. \[52\]) provides a compatible figure of $D = (0.90 - 1.11) C_1^0$, which deviates a merely 10\% away from the phenomenological relation $D = C_1^0$. The previous numbers are well within the 20\% SU(3) uncertainty estimated from the $f_\delta$ and $f_K$ difference. This is despite the fact that the $D = C_1^0$ relation is based on phenomenology, from which further uncertainties (beyond SU(3) symmetry breaking) should be expected.

Regarding HFS, as already pointed out, its application beyond the $c\bar{c}$ sector has a serious limitation in terms of model dependence within the contact-range EFT framework. The cutoff dependence of the predictions becomes larger as the reduced mass of the system is increased, from merely 1 MeV at most in the hidden-charm sector to a couple of tens of MeV in the hidden-bottom sector. This limitation was already pointed out in Ref. \[47\], where here we merely confirm the impossibility of making model independent predictions with HFS. Yet we notice that there is systematics in this model dependence, as increasing the cutoff $\Lambda$ invariably leans towards more binding. This is important, as it implies that the conclusion that the $c\bar{b}, b\bar{b}$ and $b\bar{c}$ molecular pentaquarks bind is indeed model independent, with the model dependence limited to how much they bind. In fact it can be shown that for two-body molecular systems where the potential respects HFS (i.e. the potential is independent of the heavy-quark mass), the binding energy $B_2$ increases monotonically with the reduced mass $\mu$, $\partial B_2/\partial \mu > 0$ (check Appendix \[A\] for further details). That is, though the specific masses of the $c\bar{b}, b\bar{b}$ and $b\bar{c}$ pentaquarks are model dependent to a certain extent, the fact that these systems bind is a model independent outcome of the calculations.

5. INCLUDING HEAVY-QUARK SPIN SYMMETRY

Previously we have made the simplifying assumption that the potentials binding the $P_{c1}, P_{c2}$ and $P_{c3}$ pentaquarks are unrelated. However, HQSS connects the potentials of these three configurations and allows for a common description of the $P_{c1}, P_{c2}$ and $P_{c3}$ molecules \[33,37\] (where here we will concentrate on the consequences of HQSS for the type of contact-range EFTs we are using). The disadvantage though is that we do not know which of the $P_{c2}$ and $P_{c3}$ pentaquarks corresponds to the $J = \frac{1}{2}$ and $\frac{3}{2}$ $D^* \Sigma_c$ configurations. As a consequence there are two possible set of predictions for the $P_{c}^{0,2,\Sigma^0_Q}$ family of molecules, depending on which spin identification we propose for the $P_{c2}$ and $P_{c3}$ pentaquarks.

HQSS indicates that the $|\bar{Q}q\rangle$ and $|Qqq\rangle$ family of heavy hadrons are related by means of rotations of the spin of the heavy quark. Indeed, we can group the ground and excited states of a heavy hadron in a single superfield, which for the S-wave heavy mesons and baryons are defined as

\[
H = \frac{1}{\sqrt{2}} \left[ P + \sigma \cdot P^\dagger \right],
\]

\[
\bar{S} = \frac{1}{\sqrt{3}} \left[ \sigma B_0 + \bar{P} B_b^\dagger \right],
\]

where for simplicity we are ignoring the SU(3)-flavor indices and with $P, P^\dagger$ the $J = 0, 1$ heavy mesons, $B_0, B_b^\dagger$ the $J = \frac{1}{2}, \frac{3}{2}$ heavy baryons and $\sigma$ the Pauli matrices. With the previous definitions, the lowest-order contact-range Lagrangian describing molecular pentaquarks reads \[38\]

\[
L_{\text{contact}} = C_a \text{Tr}[H^\dagger H] \bar{S}^\dagger \cdot \bar{S} + C_b \sum_{i=1}^{3} \text{Tr}[H^i \sigma_i H] \bar{S}^\dagger \cdot (J_i \bar{S}),
\]

where $J_i$ are the $i = 1, 2, 3$ spin-1 matrices. The terms proportional to the couplings $C_a$ and $C_b$ correspond to central and spin-spin contact-range interactions. Thus, the practical implication of the HQSS version of the contact-range Lagrangian is that the $C_1^0$ couplings we previously defined in Eq. \[12\] can be decomposed in central and spin-spin components:

\[
C_i^0 \rightarrow C_a^0 + \lambda_i C_b^0,
\]

where the explicit decomposition for the three known molecular pentaquark candidates is

\[
V_c(P_{c1} \bar{Q}q) = C_a^0, \quad V_c(P_{c2} \bar{Q}q) = C_a^0 - \frac{4}{3} C_b^0, \quad V_c(P_{c3} \bar{Q}q) = C_a^0 + \frac{2}{3} C_b^0,
\]

while for the four potentially unobserved configurations we will have

\[
V_c(P_{c1} \bar{Q}q) = C_a^0, \quad V_c(P_{c2} \bar{Q}q) = C_a^0 - \frac{5}{3} C_b^0, \quad V_c(P_{c3} \bar{Q}q) = C_a^0 + \frac{2}{3} C_b^0, \quad V_c(P_{c3} \bar{Q}q) = C_a^0 + \frac{2}{3} C_b^0.
\]

Now, for the $P_{c1}$ pentaquark the identification of its particle and spin channel is trivial: $J = \frac{1}{2} \bar{D} \Sigma_c$. Meanwhile this is not the case for the $P_{c2}$ and $P_{c3}$ pentaquarks: both are expected to be $D^* \Sigma_c$ molecules, but what is not clear is which one is the spin $J = \frac{1}{2}$ and $\frac{3}{2}$ state, as their spins have not been experimentally determined yet. Thus there are two possibilities:

(i) that the $P_{c2}$ and $P_{c3}$ pentaquarks are $J = \frac{1}{2}$ and $\frac{3}{2}$ states, respectively, thus following the standard pattern of mass increasing with spin, which we will call scenario A, and

(ii) the opposite pattern, mass decreasing with spin, is scenario B.
These scenarios have been named following the convention found in Ref. [12]. Different theoretical works prefer scenario A [11, 64], scenario B [34, 48, 65, 66], do not find a strong preference [12, 44] or explore alternative possibilities [67, 68]. Scenario A has recently been explained as a consequence of the short-range interaction of the light-quarks within the heavy antimeson and heavy baryon composing the pentaquarks [69]. Scenario B appeared before the discovery of the pentaquark trio, for instance in Ref. [38], and has received explanations both in terms of pion [7] and vector meson exchanges [70].

Here a caveat is in place: the wave function is not an observable, and as a consequence there will always remain a degree of ambiguity on whether a particular state is composite or not (or how composite it is). In fact, the EFT framework usually does not rely on including new degrees of freedom at subleading orders in the wave function to improve predictions. Instead, it includes new contact-range operators acting on the degrees of freedom already present, which means that compact components often manifest as energy dependence.

Be it as it may, EFT can be used to derive a dimensional estimation of the compositeness (\(X_{\text{comp}}\), i.e. the probability of the meson-baryon component) of the pentaquarks

\[
X_{\text{comp}}(P_{QQ}) = 1 - O\left(\frac{Q}{M}\right) = 1 + O\left(\frac{Q}{M}\right)
\]

\[
\approx \frac{1}{1 + x_c \frac{\sqrt{2} \mu_B}{m_C}} + O\left(\frac{Q}{M}\right),
\]

where we have reordered the terms in order to obtain an expression that is suitable when \(Q/M\) is not small (i.e. when the binding energy is closer to the limit at which the EFT will fail, so we only have \(Q/M < 1\) but not \(Q/M \ll 1\)). In the second line we have particularized for the choice \(Q = \gamma_2 = \sqrt{2} \mu_B\) and \(M = m_C\), where \(x_c\) is a numerical constant of \(O(1)\) for which we will choose \(x_c = 1\). This yields a compositeness of around \(X_{\text{comp}}^\text{dim} = (0.85, 0.78, 0.88)\) for the \(P_{c1}, P_{c2}\) and \(P_{c3}\) pentaquarks in the \(c\bar{c}\) sector, \((0.76, 0.70, 0.79)\) and \((0.71, 0.67, 0.75)\) for \(b\bar{c}\) and \(b\bar{b}\) counterparts, respectively, while merely a value of \((0.60, 0.56, 0.63)\) for their \(b\bar{b}\) counterparts. As a comparison, for the deuteron (\(\gamma_2 = 45\) MeV) we will obtain a compositeness of 0.94, compatible with a pure molecular interpretation. Yet, we remind that these estimates are purely based on a comparison of scales and are not very precise. This is illustrated by the numerical factor \(x_c\) in Eq. (41), where by taking \(x_c = 1/2\) or \(x_c = 2\) instead of \(x_c = 1\) (all of which are \(O(1)\)), the compositeness will change by a factor of order \(Q/M\).

Actually, there is a rich literature dealing with ways of quantifying the compositeness of a state [71–73], which we can use to obtain a refined estimation of \(X_{\text{comp}}\). They began with the compositeness criterion proposed by Weinberg [71–73], which can be written as

\[
X_{\text{comp}}^W = \frac{1}{\sqrt{1 - 2r_0}}
\]

where \(a_0\) and \(r_0\) are the scattering length [8] and effective range and which showed in a model-independent way that the deuteron is probably composite. It actually returns \(X_{\text{comp}}^W > 1\) for the deuteron, which indicates we are using the previous formula beyond its domain of validity (\(r_0 < 0\) for obtaining \(X_{\text{comp}}^W < 1\) for a bound state, not to mention that there will be corrections coming from the range of the interaction, as already pointed in [73]), but this result is usually interpreted as molecular. The bottom-line though is that the Weinberg criterion relies heavily on the sign of the effective range of the purported components of the state: if positive (negative) the state will be predominantly composite (elementary). As a consequence the application of this criterion will lead to the conclusion that the pentaquarks we are dealing with here are mostly molecular. This however will be an artifact of the formalism we are using: our LO calculation automatically generates a positive effective range, which is a consequence of

\[\text{In our convention, for attractive potentials } a_0 < 0 \text{ in the absence of bound states and } a_0 > 0 \text{ when there is one bound state.}\]
the dynamics we are using. Besides, even though it is evident that the energy dependence of a compact core coupled to a two-hadron system is such that it will generate a negative effective range, a sufficiently short-ranged potential combined with a large binding energy implies a sizable superposition.
of the hadrons and, owing to their finite size, also a degree of non-compositeness. From this and other arguments, extensions of the Weinberg criterion have been proposed that apply to situations different from a bound state with negative effective range \([74–82]\).

A recent proposal of a model-independent estimation of the compositeness of a state is the following \([80]\):

\[
\hat{X}_{\text{comp}} = \sqrt{\frac{1}{1 + 2\hat{\epsilon}}},
\]

which returns \(\hat{X}_{\text{comp}} < 1\), where the calculation of \(\hat{\epsilon}\) and \(r_0\) for our contact-range theory is explained in Appendix \(\text{[III]}\). This criterion would provide a compositeness of \(\hat{X}_{\text{comp}} = (0.73, 0.67, 0.76)\) for each of the three LHCb pentaquarks (i.e., \(P_{c1}, P_{c2}, P_{c3}\)), \((0.66, 0.62, 0.68)\) and \((0.63, 0.61, 0.65)\) for the \(\bar{c}b\) and \(c\bar{b}\) ones and \((0.59, 0.57, 0.60)\) for the hidden-bottom \(P_{b1}, P_{b2}\) and \(P_{b3}\) pentaquarks. However, the problem here is that we are using a LO EFT description with only one parameter (the binding energy), which means that the value of the effective range thus obtained is only a dimensional estimation within our EFT. For comparison the compositeness of the deuteron \((a_0 = 5.419\text{ fm}, r_0 = 1.753\text{ fm}\text{[83]})\) with this criterion will be 0.78, but in this case there is plenty of experimental information available about neutron-proton scattering, i.e. \(a_0\) and \(r_0\) are well-known.

Regardless of the specific criterion used to estimate compositeness (after all, the wave function is not an observable), it seems that in general the hidden-charm pentaquarks are less composite than the deuteron, and as we move into heavier flavor sectors their compositeness reduces further. This is in turn compatible with the observation that the EFT description is less convergent and has larger uncertainties for two-body systems with larger binding energies. Thus, as binding increases with the reduced mass, we expect compositeness to decrease accordingly.

| Molecule | \(I\) | \(S\) | \(B_P\) | \(M_P\) | \(J\) | Scenario |
|----------|------|-----|-------|-------|-----|---------|
| \(B^0\) | \(-\) | 0 | 27.0 \(^{+8.6}_{-7.3}\) | 7770.6 \(^{+7.3}_{-6.7}\) | \(-\) | \(A\) |
| \(B^-\) | \(-\) | 0 | 49 \(^{+10}_{-12}\) | 7794.3 \(^{-12}_{+10}\) | \(-\) | \(A\) |
| \(B^+\) | \(-\) | 0 | 35.6 \(^{+9.8}_{-8.5}\) | 7807.3 \(^{+9.8}_{-8.5}\) | \(-\) | \(A\) |
| \(B^0\) | \(-\) | 0 | 15.6 \(^{+7.0}_{-6.0}\) | 7827.3 \(^{+5.8}_{-5.5}\) | \(-\) | \(A\) |
| \(B^+\) | \(-\) | 0 | 12.8 \(^{+15.5}_{-12.6}\) | 7857.3 \(^{+15.5}_{-12.6}\) | \(-\) | \(A\) |

| Molecule | \(I\) | \(S\) | \(B_P\) | \(M_P\) | \(J\) | Scenario |
|----------|------|-----|-------|-------|-----|---------|
| \(B^0\) | \(-\) | 0 | 35.5 \(^{+9.8}_{-8.4}\) | 7762.9 \(^{+8.4}_{-8.0}\) | \(-\) | \(B\) |
| \(B^-\) | \(-\) | 0 | 15.6 \(^{+5.8}_{-7.0}\) | 7827.3 \(^{-7.0}_{+5.8}\) | \(-\) | \(B\) |
| \(B^+\) | \(-\) | 0 | 27.1 \(^{+7.3}_{-8.7}\) | 7815.7 \(^{+7.3}_{-8.7}\) | \(-\) | \(B\) |
| \(B^0\) | \(-\) | 0 | 49 \(^{+12}_{-10}\) | 7794.1 \(^{-10}_{+12}\) | \(-\) | \(B\) |

TABLE IV. Same as Table III but for the \(\bar{c}b\) and \(bb\) sectors.
7. FLAVOR SYMMETRY AND NON-MOLECULAR EXPLANATIONS

The present predictions have been done under the assumption that the hidden-charm pentaquarks are molecular. But, as a matter of fact, the light- and heavy-flavor symmetries we have used here are expected to apply to other light-heavy hadrons as well, independently of their nature (though the uncertainties stemming from the violations of these symmetries could be very different). For instance, the existence of this type of pentaquark multiplets has been predicted in the compact [40] and hadroquarkonium pictures [84, 85]. Theoretical explorations in the previous pictures have been mostly concentrated in the hidden-charm sector, where the mass splittings of the octet \( |m(P^\Lambda) - m(P^\Sigma)|\), \( |m(P^\Sigma) - m(P^\Xi)|\) and \( |m(P^\Xi) - m(P^\Omega)|\) are 141, 205 and 315 MeV for compacts pentaquarks [40] and 150, 217 and 327 MeV for hadrocharm [84]. These mass splittings happen to be larger than for molecular pentaquarks (125, 105 and 232 MeV) and might provide a way to distinguish their nature if they are observed. For the hidden-bottom sector there are indeed predictions of \( P^\Omega_b \) pentaquarks in the local hidden-gauge approach of Ref. [87] and in models considering a five-quark core and pion exchanges [86]. It is plausible that other theoretical models of \( Q^\prime \bar{Q} \) pentaquarks will lead to analogous predictions for their flavor partners, as these predictions are constrained by symmetry principles (instead of the details of the dynamics, which will matter for how the spectrum is organized in terms of quasi numbers, spin-spin splitting, etc.). Recent calculations of \( q\bar{q}Q^\prime \bar{Q} \) pentaquarks in the hadroquarkonium [85] and chiral quark models [88] provide further support for this conjecture.

8. SUMMARY

The observation of the LHCb hidden-charm pentaquarks in combination with SU(3)- and heavy-flavor symmetries leads to the prediction of a series of flavor partners. In particular, pentaquarks (molecular and non-molecular [40] alike) are expected to form a light-flavor octet reminiscent of the light-baryon octet and are also expected to appear in the \( \bar{c}b, \bar{c}b \) and \( bb \) sectors as well as in the original hidden-charm sector where they have been discovered. We denote these pentaquarks as \( P^N_Q, P^\Lambda_Q, P^\Sigma_Q, P^\Xi_Q, P^\Omega_Q \), with the superscript and subscript referring to their light- and heavy-quark structure, respectively (which we shorten to \( P^N_Q, P^\Lambda_Q, P^\Sigma_Q, P^\Xi_Q \) and \( P^\Omega_Q \) when the heavy flavors coincide \( Q = Q \), i.e. for hidden-flavor). For predicting their masses, we have made use of a contact-range description of the spectrum will be different than in the molecular case.

is worth noticing that the applicability of this description decreases with increasing binding energy, as this implies pentaquarks that are less composite, and with heavier reduced masses owing to the model-dependent nature of HFS [47]. This is reflected in the larger uncertainties, particularly in the hidden-bottom sector. Yet, it is sensible to expect these predictions to be more dependent on the general symmetry principles we have applied than on the details of the dynamics generating the pentaquarks, e.g. models with a compact five-quark core coupled to the meson-baryon degrees of freedom do reproduce the hidden-charm pentaquarks [34] and also predict the hidden-bottom ones [34], giving credence to the aforementioned conjecture. Thus it might be the case that the light- and heavy-flavor symmetry partners of the hidden-charm pentaquarks exist irrespective of the binding mechanism, though the details of the spectrum will be different than in the molecular case.

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Appendix A: Heavy-quark mass dependence of the binding energy

Here we consider the variation of the binding energy of a heavy hadron molecule with respect to the heavy-quark mass. If the potential between two heavy hadrons does not depend on the heavy-quark mass, it can be shown that the binding energy increases with the heavy-quark mass (in agreement with naive expectations).

At leading order in the \( 1/m_Q \) expansion, we can write the Schrödinger equation for a heavy hadron molecule as follows

\[
- \Psi^2 \Psi_Q(\hat{r}) + 2 \mu_Q V_Q(\hat{r}) \Psi_Q(\hat{r}) = -2 \mu_Q B_Q \Psi_Q(\hat{r}),
\]

(A1)

where the subindex \( Q \) indicates the dependence (explicit and implicit) on the heavy-quark mass, \( \Psi_Q \) is the wave function, \( \mu_Q \) the reduced mass of the molecule, \( V_Q \) the potential and \( B_Q \) the two-body binding energy. We can construct a Wronskian identity for the Schrödinger equation at two different heavy-quark masses as follows

\[
- \left( \Psi_Q^2 \Psi_Q - \Psi_Q \Psi_Q^2 \Psi_Q \right) + 2 \left( \mu_Q V_Q - \mu_Q^2 V_Q^2 \right) \Psi_Q \Psi_Q = -2 \left( \mu_Q B_Q - \mu_Q^2 B_Q^2 \right) \Psi_Q \Psi_Q^2.
\]

(A2)

where, again, \( \Psi_Q \) and \( \Psi_Q^2 \) represent the different quantities we are considering at \( m_Q \) and \( m_Q^\prime \), respectively. The Wronskian
identity can be integrated, leading to

\[ 2 \int d^3\bar{r} (\mu_Q V_Q - \mu_{Q'} V_{Q'}) \Psi_{Q'}(\bar{r}) \Psi_Q(\bar{r}) = -2(\mu_Q B_Q - \mu_{Q'} B_{Q'}) \int d^3\bar{r} \Psi_{Q'}(\bar{r}) \Psi_Q(\bar{r}) \]  

(A3)

where the kinetic term disappears because it is exactly differentiable and can be rewritten as a surface term, which vanishes if we consider bound state solutions. Now we will consider a small change in the heavy-quark mass, which we can symbolically indicate by

\[ Q' = Q + \delta Q. \]  

(A4)

We can deduce that

\[ \int d^3\bar{r} \Psi_{Q'}(\bar{r}) \Psi_Q(\bar{r}) = 1 + (\delta Q)^2 \]  

(A5)

which is a consequence of the normalization of the wave function (i.e. \( \langle \Psi_{Q'} | \Psi_{Q'} \rangle = \langle \Psi_Q | \Psi_Q \rangle = 1 \), which is why the \( \delta Q \) term vanishes). If we assume that the potential does not depend on the heavy-quark mass, i.e. \( V_Q = V_{Q'} \), we can use the previous result to prove that

\[ 2 \delta \mu_Q \langle V_Q \rangle = -2 \delta (\mu_Q B_Q), \]  

(A6)

which we can differentiate to obtain

\[ \langle V_Q \rangle = -B_Q - \mu_Q \frac{\partial B_Q}{\partial \mu_Q}. \]  

(A7)

If we take into account

\[ \langle T_Q \rangle + \langle V_Q \rangle = -B_Q, \]  

(A8)

where \( \langle T_Q \rangle \geq 0 \) is the kinetic energy of the heavy molecule, we can rewrite the binding energy dependence on the reduced mass as

\[ \langle T_Q \rangle = \mu_Q \frac{\partial B_Q}{\partial \mu_Q} \]  

(A9)

or, equivalently

\[ \frac{\partial B_Q}{\partial \mu_Q} \geq 0, \]  

(A10)

as a consequence of the fact that the kinetic energy is positive. That is, the system will become more bound the heavier the mesons (this is a model-independent result). What is difficult (and model-dependent) is to determine by what amount. Finally, we notice that including a heavy-quark mass dependence of the type \( V_Q = V_0 + \frac{m_Q}{M} V_1 + \ldots \) in the potential does only induce \( 1/m_Q \) corrections to the previous relation, which can be safely neglected in the heavy-quark mass limit.

**Appendix B: Calculation of the effective range expansion parameters**

The evaluation of the different compositeness conditions available in the literature usually require the effective range parameters as input. Here we briefly explain how to calculate them. We begin by writing down the relation between the effective range expansion and the on-shell T-matrix \( T_{os} \):

\[ -\frac{2\pi}{\mu} \text{Re} \left[ \frac{1}{T_{os}(k)} \right] = -\frac{1}{a_0} + \frac{1}{2} n_0 k^2 + \sum_{n=2}^{\infty} v_n k^{2n}, \]  

(B1)

where \( a_0 \) is the scattering length, \( n_0 \) the effective range, \( v_n \) the shape parameters, \( k \) the center-of-mass momentum and \( \mu \) refers to the reduced mass of the two-body system. For attractive potentials, the previous convention implies \( a_0 < 0 \) if there is no bound state (or an even number of bound states) and \( a_0 > 0 \) if there is an odd number of bound states. The on-shell T-matrix corresponds to the following matrix element of the full T-matrix

\[ T_{os}(k) = \langle k | T(k) | k \rangle, \]  

(B2)

where \( T \) obeys the Lippmann-Schwinger equation, which for scattering states takes the form

\[ T = V + V G_0(E + i\epsilon) T, \]  

(B3)

with \( G_0(E) = 1/(E - H_0) \) the resolvent operator and \( E = k^2/2\mu \) the center-of-mass energy of the system. If we consider a regularized contact-range of the type

\[ \langle p'|V_C|p \rangle = C(\Lambda) f(p'/\Lambda) f(p/\Lambda), \]  

(B4)

then the explicit solution of the Lippmann-Schwinger equation for the on-shell T-matrix reads

\[ \text{Re} \left[ \frac{1}{T_{os}(k)} \right] = \frac{1}{C(\Lambda)} - \frac{\mu}{\pi^2} \mathcal{P} \int_0^{\infty} \frac{p^2 dp f^2(p/\Lambda)}{k^2 + i\epsilon - p^2 f^2(p/\Lambda)}, \]  

(B5)

where \( \mathcal{P} \) denotes the principal value of the integral. By expanding in powers of the center-of-mass momentum, we arrive at

\[ \frac{1}{a_0} = \frac{2\pi}{\mu} \frac{1}{C(\Lambda)} + \frac{2}{\pi} \int_0^{\infty} dp f^2(p/\Lambda), \]  

(B6)

\[ n_0 = -\frac{4}{\pi} \int_0^{\infty} \frac{dp}{p^2} \left(f^2(p/\Lambda) - f^2(0)\right), \]  

(B7)

where we can appreciate that at LO in our contact-range theory \( n_0 \) merely provides a dimensional estimation of its size. If we particularize for our choices of regulator function, cutoff and couplings, we will obtain the values of \( a_0 \) and \( n_0 \) that we have used as input for Eq. (43).
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