Efimov-van-der-Waals universality for ultracold atoms with positive scattering lengths

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We study the universality of the three-body parameters for systems relevant for ultracold quantum gases with positive $s$-wave two-body scattering lengths. Our results account for finite-range effects and their universality is tested by changing the number of deeply bound diatomic states supported by our interaction model. We find that the physics controlling the values of the three-body parameters associated with the ground and excited Efimov states is constrained by a variational principle and can be strongly affected by $d$-wave interactions that prevent both trimer states from merging into the atom-dimer continuum. Our results enable comparisons to current experimental data and they suggest tests of universality for atomic systems with positive scattering lengths.

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I. INTRODUCTION

The recent theoretical and experimental progress in the exploration of ultracold quantum gases in the strongly interacting regime have largely established the relevance of the three-body Efimov physics [1–3] for the understanding of both dynamics and stability of such systems [4–12]. The control of interatomic interactions through magnetic-field dependent Feshbach resonances [13] allows for dramatic changes in the $s$-wave two-body scattering length, $a$, making it possible to tune systems to the vastly different collective (mean-field) regimes of attractive, $a < 0$, and repulsive, $a > 0$ interactions. In the regime of strong interactions, $|a|/r_{vdW} \gg 1$, where $r_{vdW}$ is the van der Waals length [13], the Efimov effect is manifested through the appearance of an infinite series of three-body states that can lead to scattering resonances and interference effects accessible to experiments [2–8]. Such dramatic few-body phenomena open up the possibility to explore new quantum regimes in ultracold gases. One of the striking signatures of the Efimov effect is the geometric scaling of the system for many trimer properties, which interrelates all the three-body observables via the geometric factor $e^{\pi/\alpha_0}$, where $\alpha_0 \approx 1.00624$ for identical bosons. As a result, if universal scaling holds, the determination of a single observable — the three-body parameter — would allow derivation of all properties of the system. However, since the early days of Efimov's original prediction it was largely accepted that this three-body parameter would be different for every system. Nevertheless, a few years ago, as experiments in ultracold gases evolved, it became clear that this concept needed reassessment.

The turnaround came from the experimental observations in $^{133}$Cs [14] showing that the three-body parameter $a_*$, associated with the value of $a < 0$ at which the first Efimov state merges with the three-body continuum, were the same (within a 15% margin) for different resonances in $^{133}$Cs. Moreover, if the results were recast in terms of $r_{vdW}$, the observations in every other available atomic species also led to similar results, $a_*/r_{vdW} \approx -10$ (see Ref. [3] for a summary of such experimental findings). Theoretical works then successfully confirmed and interpreted the universality of the $a_*$ parameter [15–21] and consolidated a new universal picture for Efimov physics in atomic systems dominated by van der Waals forces.

This paper assesses the universality of the three-body parameter in the yet unexplored regime of positive scattering lengths, $a > 0$. The available experimental data for Efimov features within this regime is relatively sparse and, consequently, does not clearly display the same degree of universality found for $a < 0$. Although not explicitly demonstrated here, our present theoretical study shows that universality for $a > 0$ persists and is rooted in the same suppression of the probability of finding particles at short distances previously found for $a < 0$ [13–17]. The observables we analyze are related to the value of $a$ at which an Efimov state intersects the atom-dimer threshold, $a_*$, thus causing a resonance in atom-dimer collisions [22, 23], and the value $a_+$ at which a minimum in three-body recombination occurs as a result of a destructive interference between the relevant collision pathways [22, 24, 25]. One important feature that can help to interpret our computed values for $a_*$ and $a_+$ associated with the ground Efimov state is the existence of a variational principle [27, 28] that constrains its energy to always lie below a certain value lower than the dimer energy, thus preventing the trimer to cross the atom-dimer threshold. This has a direct impact on both the lowest atom-dimer resonance and on interference phenomena, even when, as we show here, the conditions for the validity of that variational principle are not strictly satisfied. Moreover, our analysis indicates that the presence of strong $d$-wave interactions [29, 30], and/or possibly some other finite-range effects, also prevents the first excited Efimov state from merging with the dimer threshold, although it still produces a resonance feature in atom-dimer observables.
and a recombination minimum for small $a$.

II. BRIEF THEORETICAL BACKGROUND

Here we use the adiabatic hyperspherical representation which offers a simple and conceptually clear description of few-body systems while still accurately determining their properties [3]. Within this representation, after solving for the hyperangular internal motion—which includes all interparticle interactions—three-body observables can be obtained by solving the hyperradial Schrödinger equation [31]

$$\left[-\frac{\hbar^2}{2\mu}\frac{d^2}{dR^2} + W_\nu(R)\right] F_\nu(R) + \sum_{\nu'\neq\nu} W_{\nu'\nu}(R)F_{\nu'}(R) = EF_\nu(R). \tag{1}$$

where the hyprerradius $R$ describes the overall size of the system, $\mu = m/\sqrt{3}$ is the three-body reduced mass and $\nu$ is an index including all necessary quantum numbers to characterize each channel. Equation (1) describes the radial motion governed by the effective hyperspherical potentials $W_\nu$ and non-adiabatic couplings $W_{\nu'\nu}$, which determine all bound and scattering properties of the system. In the present study, each pair of particles interacts via a Lennard-Jones potential

$$v_{LJ}(r) = -\frac{C_6}{r^6} \left(1 - \frac{\lambda^6}{r^6}\right), \tag{2}$$

where $\lambda$ is adjusted to give the desired value of $a$ and $C_6$ is the usual dispersion coefficient. Note that our calculations use van der Waals units (with energy and length units of $E_{vdW} = \hbar^2/mr_{vdW}^3$) such that the specification of the value of $C_6$ is unnecessary. Our present study is centered around the first three poles of $a$, which occur at the values denoted $\lambda = \lambda_1^*, \lambda_2^*$, and $\lambda_3^*$. One important point to keep in mind is that near $\lambda_1^*$ there can exist only a single two-body s-wave state, whereas near $\lambda_2^*$ and $\lambda_3^*$ multiple deeply bound states exist (4 and 9, respectively), owing to the presence of higher partial wave dimers.

III. RESULTS

Figure 1 shows the energies of the lowest three Efimov states, $E_{3b}$, for values of $a$ near the three poles considered ($\lambda_1^*, \lambda_2^*$, and $\lambda_3^*$), offering a global view of the degree of the universality of our results. Near $\lambda_1^*$, Efimov states (black filled circles) are true bound states while near $\lambda_2^*$ and $\lambda_3^*$ (red and green open circles, respectively) Efimov states are resonant states with the corresponding widths indicated as the error bars. Approximate values for $a_-, a_+$, and $a_+$ are also indicated.

![Image](image_url)

**FIG. 1:** Energy of Efimov states calculated near the first three poles of $a$, $\lambda = \lambda_1^*, \lambda_2^*$ and $\lambda_3^*$, in our model potential in Eq. (2). Near $\lambda_1^*$, Efimov states (black filled circles) are true bound states while near the $\lambda_2^*$ and $\lambda_3^*$ (red and green open circles, respectively) Efimov states are resonant states with the corresponding widths indicated as the error bars. Approximate values for $a_-, a_+$, and $a_+$ are also indicated.

Closer inspection of Fig. 1 reveals that the first excited Efimov state also fails to intersect the atom-dimer threshold, as expected from the variational principle in Refs. [27, 28], which state that $E_{3b} < 3E_{2b}$. In principle, this variational constraint applies only to bound states, i.e., only for Efimov states near $\lambda_1^*$, however, our calculations for the energies of Efimov resonances near $\lambda_2^*$ and $\lambda_3^*$ also follow the same non-crossing rule. Evidently, this effect strongly modifies the expected universality predicted by zero-range models since it prevents an atom-dimer resonance and can also modify the minima in recombination associated with the ground Efimov state. Table I summarizes our computed values of the three-body parameters—see also Fig. 1 for their approximate location. The values for $a_-$ were previously determined in Ref. [15] (and in unpublished work from that study). The additional index on the $a_-, a_+$, and $a_+$ parameters indicates their Efimov family parentage. The physics involved and caveats on the determination of these three-body parameters are given below.

Closely inspection of Fig. 1 reveals that the first excited Efimov state also fails to intersect the dimer threshold. This is clearly shown in Fig. 2 for the binding energy of the Efimov states, $E_b = E_{2b} - E_{3b}$. Near $\lambda_1^*$ (black filled circles) the non-crossing of the first excited state is evident within the shaded region in Fig. 2. Near $\lambda_3^*$ (red open circles), the qualitative behavior is the same, however: As the energy of the Efimov state approaches the threshold its width increases to the point in which it exceeds the value of its binding energy—therefore, losing some its “bound” state characteristic—and eventually “dissolving” into the atom-dimer continuum (see shaded
TABLE I: Values for the three-body parameters $a_-, a_+$, and $a_+$ for the lowest two Efimov scattering features in recombination and atom-dimer collisions, near the lowest three poles in the scattering length. For $a_2$, we also show its dependence on the temperature by $\langle K_i \rangle$ (see text) at values of $k_B T/E_{vdW}$ (indicated in square brackets) listed in the last three columns below. In the bottom part of the table we list the universal ratios $\beta_{ij}^a$ [see Eq. (9)], resulting from the average value of the three-body parameters (see text for the comparison with the zero-range results).

| $(i, j)$ | $(a_-, i)/r_{vdW}$ | $(a_+, i)/r_{vdW}$ | $(a_+, 1)/r_{vdW}$ |
|---------|-----------------|-----------------|-----------------|
| $\lambda_1^+$ | $-9.60, -161$ | $3.41, 157$ | $1.41, 27.2$ | $28.0$ | $29.1$ | $32.1$ |
| $\lambda_2^+$ | $-9.74, -164$ | $3.26, 160$ | $1.41, 27.9$ | $28.7$ | $30.7$ | $34.8$ |
| $\lambda_3^+$ | $-9.96, -$ | $3.33, 160$ | $1.41, 28.0$ | $-$ | $-$ | $-$ |
| Avg. | $-9.77, -163$ | $3.33, 159$ | $1.41, 27.7$ | $28.4$ | $29.9$ | $33.5$ |
| $(i, j)$ | $\theta_{ij}^{0-}$ | $0.143$ | $0.195$ | $0.125$ | $0.170$ | $-$ |
| $\theta_{ij}^{0-}$ | $-$ | $-$ | $-$ | $-$ | $-$ |
| $\theta_{ij}^{0+}$ | $-$ | $-$ | $-$ | $-$ | $-$ |

region in Fig. 2. Passing this point, as $\alpha$ decreases further, the state recovers its bound character. Our physical interpretation of the non-crossing of the first excited Efimov state $\lambda_1^+$ is that it results from the existence of strong $d$-wave interactions near $a/r_{vdW} = 1$ [24, 30]. Within our theoretical model, since $s$- and $d$-wave interactions can not be separated, a more clear physical picture of the non-crossing of the first excited Efimov state still remains, leaving even the possibility of that being a generalization of the same variational principle [27, 28] which prevents the ground state to unbind. Figure 2 shows that only the second excited Efimov state displays the expected interaction with the atom-dimer threshold.

Evidently, the effects analyzed above have an important impact on the determination of the three-body parameter $a_+$. This is achieved here by directly calculating the corresponding atom-dimer scattering properties. Of particular importance for ultracold experiments is the atom-dimer scattering length $a_{vdW}$ and the atom-dimer loss rate $\beta$ [24]. Figure 3 shows our calculated values for these quantities. In Fig. 3 (a), around the shaded region (corresponding to the same shaded region in Fig. 2) $a_{vdW}$ is enhanced, however, remaining always positive and consistent with the failure of the first excited Efimov state in Fig. 2 to become unbound. (Note that in this regime $a_{vdW}$ for $\lambda_1^+$ and $\lambda_2^+$ displays a more complicated dependence on $\alpha$ due to the presence of strong couplings to nearby three-body channels.) For larger $\alpha$, $a_{vdW}$ is now enhanced and changes sign, implying that the second excited Efimov state intersects with the dimer energy (see Fig. 2). Note that here, $a_{vdW}$ for $\lambda_2^+$ and $\lambda_3^+$ does not actually diverge due to the presence of inelastic processes [35]. Figure 3 (b) shows a blow up of (a) near the second and first excited Efimov states, respectively.
shows the corresponding atom-dimer loss rates, which display the resonant behavior associated with the first and second excited Efimov states. Even though the first excited Efimov state does not become unbound, it approaches the atom-dimer threshold close enough to produce a clear enhancement in the atom-dimer loss rate. We define \( a_{+,1} \) and \( a_{+,2} \) as the value of \( a \) where \( \beta \) is maximum [see Fig. 3(b)], except for our calculations near the first pole, where no losses occur (\( \beta = 0 \)). In this case \( a_{+,1} \) and \( a_{+,2} \) were determined from the maximum value of \( a_{\text{vdW}} \) [see Fig. 3(a)]. Numerical values are listed in Table I. In order to contrast our numerical results with the universal predictions (based on two-body contact interaction models), we also display in Fig. 3 (dashed lines) the expected predictions (based on two-body contact interaction models) for Fig. 3(b), in order to better fit the data for \( \lambda^* \). Numerical values are listed in Table I. In principle, at finite temperatures one would also need to include higher partial-waves contributions to recombination. For identical bosons, however, the next leading contribution is for \( J = 2 \) and scale with the temperature and scattering length as \( T^2 a^8 \). Comparing those with the ones shown in the bottom part of Table I (calculated using the averaged values for \( a_{\text{vdW}} \)) is a balance between universal \( s \)- and \( d \)-wave physics. The inset of Fig. 3 shows the corresponding atom-dimer loss rates, which illustrate the temperature dependence of \( a_{+,1} \). Although the agreement is very good for large \( a \), near \( a_{+,1} \) not only finite range corrections become more important but also the fact that the first excited Efimov state fails to intersect with the atom-dimer threshold, imply strong deviations between universal zero-range theory and our results.

The inset of Fig. 4 shows the temperature dependence of \( K_3 \) obtained by calculating the thermally averaged recombination rate \( \langle K_3 \rangle \), which illustrates the temperature dependence of \( \langle a_{+,1} \rangle \) in the regime relevant for experiments — see also the values listed in Table I. In principle, at finite temperatures one would also need to include higher partial-waves contributions to recombination. For identical bosons, however, the next leading contribution is for \( J = 2 \) and scale with the temperature and scattering length as \( T^2 a^8 \). Comparing those with the ones shown in the bottom part of Table I (calculated using the averaged values for \( a_{\text{vdW}} \)) is a balance between universal \( s \)- and \( d \)-wave physics. The inset of Fig. 3 shows the corresponding atom-dimer loss rates, which illustrate the temperature dependence of \( a_{+,1} \).

Finally, we have also calculated the three-body recombination rate, \( K_3 \), in the lowest three-body angular momentum \( (J = 0) \) to determine the values of the three-body parameter \( a_{+,1} \). Figure 4 shows our results for \( K_3 \) in the zero-energy limit \( (E = 10^{-6} E_{\text{vdW}}) \) clearly displaying two minima, whose locations are identified as the values for \( a_{+,0} \) and \( a_{+,1} \) listed in Table I. Our numerical results obtained near \( \lambda^*_1 \) are compared with the analytical results in the absence of deeply bound dimers (dashed line). For large \( a \) our results agree well with the analytical ones while strong deviations can be observed for small \( a \). In particular, one can see that the predicted minimum in recombination near \( a/r_{\text{vdW}} = 1 \) is strongly affected by finite-range effects. We trace such effects to the presence of strong \( d \)-wave interactions. In fact, near \( a/r_{\text{vdW}} = 1 \) our results display an enhancement due to a universal three-body resonance with strong \( d \)-wave character. Therefore, our result for \( a_{+,0} \) is not consistent with ours but the corresponding results for \( a_{+,1} \) are not. A comparison with results originated from models which include finite-range corrections needs to be made carefully to ensure that the interaction parameters are the same. This, however, is beyond the
The calculations of Ref. 45, marked by LJ*, in Table II also list the value of $\kappa_0 = (mE^{(0)}_a/h^2)^{1/2}$ obtained from our calculations for $\lambda_0^*$ and the corresponding averaged result from Ref. 45. The agreement is generally good for all cases (the relative errors are indicated in Table II between square brackets), with the exception for the value of $a_{+-}$, most likely because the non-crossing of the first excited Efimov state is absent in the model of Ref. 45, clearly indicating a strong effect due to $d$-wave interactions. We note, however, that the agreement for the geometric factors $a_{+1}/a_{-0}$ and $a_{+1}/a_{+-0}$ are generally better than the absolute values of the three-body parameters. This indicates that the effect of the $d$-wave interactions in such parameters is mainly to introduce a shift:

$$a_x \rightarrow a_x e^{-\phi_d/\kappa_0},$$

or, equivalently, a change in the three-body phase: $s_0 \ln(a_{+1}/a_x) \rightarrow s_0 \ln(a_{+1}/a_x) + \phi_d$. Indeed, forcing our value of $a_{+1,0}$ to reproduce the one from Ref. 45, we obtain $\phi_d \approx -0.146$ and the resulting rescaled three-body parameters, marked by LJ* in Table II now agree much better, evidently, with the exception of $a_{+-1}$. The above rescaling process, therefore, can be seen as an attempt to subtract-off $d$-wave effects from our calculations, although a more rigorous study that can provide a more quantitative analysis of such effects still needs to be performed.

| Atom | $a_{+-1}/a_{+-0}$ | $a_{+1}/a_{+-0}$ | $a_+/a_{-0}$ | $a_+/a_{+1}$ |
|------|------------------|------------------|--------------|--------------|
| $^{133}$Cs | 0.08 0.13 4.2(0.1) | 0.13 4.2(0.1) | 54, 55 |
| $^{7}$Li | 0.03 0.24 6.5(0.3) | 0.24 6.5(0.3) | 55 |
| $^{39}$K | 0.02 0.09 13.0(0.6) | 0.09 13.0(0.6) | 48, 49 |
| $^{6}$Li | 0.32 0.05 6.0(0.1) | 0.05 6.0(0.1) | 56 |

As one can see from Table III, the values listed for $a_{+0}$ and $a_{+-1}$ are qualitatively consistent among themselves, with the exception of the data for $^{39}$K 53 — a new analysis presented in Ref. 58 suggests that this data might be subject of a new calibration. Although the values for $a_{+-1}$ in Table III are likely to suffer from thermal effects (the condition $|a| \ll a_c = \hbar/\sqrt{mkT}$ ensuring the absence of thermal effects is not strictly satisfied), our finite temperature calculations covering the range of temperatures relevant for the experiments (see Table I) indicate that thermal effects might lead to no more than a 10% variation from the zero temperature result. We also note that for $^{7}$Li and $^{39}$K the resonances are substantially less broad than the ones for $^{133}$Cs (see Ref. 13), thus opening up the possibility of finite-width effects as responsible for the deviations among the experimental data in Table III. In comparison to the values for $a_+$, the results for $a_{+-1}$ listed in Table III display a much stronger deviation among themselves. A more careful analysis, therefore, is necessary in order to understand some of the possible factors affecting such observations. For instance, the value for $a_{+-1}$ from Table III of $^{133}$Cs from Ref. 58, as well as the results for $^{7}$Li from Ref. 50, were obtained using a Feshbach resonance that is not well separated from another nearby resonance, possibly affecting the observed value for $a_{+-1}$. Most of the results marked in Table III by "i" present the largest variations compared with the total averaged result for $a_{+-1}$ ($\approx 6.63\text{r}_\text{vdW}$). They were, however, obtained based on the assumption that atom-dimer resonances can be observed in atomic samples by means of an avalanche mechanism 53. Although modifications on the description of such mechanism can lead to more reasonable results 50, 59, this hypothesis is currently considered questionable 55, 60, 61.

Therefore, accordingly to our analysis above, in order to properly compare the experimental data to theoretical predictions, we excluded the data from $^{39}$K 53 and those marked by "i" in Table III. From the remaining
experimental data, we determine an average value and corresponding relative error as listed in Table IV (the relative errors are indicated between square brackets). Using the zero-range (ZR) universal relations derived in Refs. 2,3 we determined the values for $a_{+0}$, $a_{++}$ and $a_{++}$, using the average value for $a_{-0}$ in Table II and list these in Table IV along with our corresponding averaged results (LJ) from Table II. As one can see, the zero-range results for $a_{+0}$ and $a_{++}$ perform better than our results when compared to the experimental data, while our result for $a_{++}$ outperforms the zero-range result. In fact, within the zero-range model the atom-dimer resonance associated to $a_{++}$ originates from an actual crossing between the first excited Efimov state while in our model it does not (see Fig. 2 and corresponding discussion in the text). We note, however, that our results for $a_{++}/a_{+0}$ better reproduces the value from the experimental data, indicating that a shift on the position of the three-body parameters for $a > 0$, in the same spirit than the one obtained from Eq. (4), can improve the comparisons. In fact, as shown in Table IV using the results (LJ°) listed in Table II, an overall improvement can be observed. If we now rescale our results (LJ) to reproduce the experimental value for $a_{+0}$, within a relative error of 0.10, our new rescaled results (LJ°) now display a much better overall comparison to the experimental data. Although there is no clear reason why such scaling should be allowed, the above analysis clearly indicates that our numerical results might suffer from finite-range effects, whether originated from the $s$- and $d$-wave mixing or the finite-width character produced by real interatomic interactions.

**TABLE IV: Comparison between the values for the three-body parameters from different theories and the average experimental data, marked by Exp (see text). Our average results (LJ and LJ°) are those from Table II and the results marked by LJ°E are those obtained from LJ° by forcing the result for $a_{+0}$ to agree with the experimental data within 10% (see text). The corresponding relative errors are indicated between square brackets.**

| Parameter | Exp | ZR | LJ | LJ° | LJ°E |
|-----------|-----|----|----|-----|------|
| $a_{+0}/\text{vdW}$ | 2.50[0.10] | 2.05[0.22] | 1.41[0.77] | 1.63[0.53] | 2.27[0.10] |
| $a_{+1}/\text{vdW}$ | 39.3[0.12] | 46.5[0.16] | 27.0[0.42] | 32.0[0.23] | 44.7[0.12] |
| $a_{+1}/a_{+0}$ | 15.7[0.22] | 22.7[0.31] | 19.7[0.20] | 19.7[0.20] | 19.7[0.20] |
| $a_{++}/\text{vdW}$ | 4.78[0.20] | 10.4[0.54] | 3.33[0.43] | 3.85[0.24] | 5.37[0.11] |

Evidently, there is much to be understood on the effects that realistic interactions can impose in the determination of the three-body parameters. In more realistic systems the short-range multichannel nature of the interactions can produce, for instance, a different mixing of $s$- and $d$-wave components than the single channel model does. One can expect $d$-wave interactions to be more important when the system possesses a small background scattering length, i.e., of the order of $r_{vdW}$, since in this case the entrance channel physics, obeying the universality of the van der Waals interactions 29, can include a weakly bound $d$-wave state. Finite-width effects can lead to values of the effective range different than the one produced in our model, also determined by the universal van der Waals physics 62. Such effects, although not entirely understood yet, can also lead to substantial deviations of the three-body parameters 18. In fact, the model developed in Ref. 63, which incorporates some of the multichannel physics of the problem, shows a much better agreement between theory and experiment 55, including for the $a < 0$ geometric scaling $a_{-1}/a_{-0} \approx 21.0$ from Ref. 61, indicating that both $s$- and $d$-wave mixing and finite-width effects might be at the heart of deviations of the three-body parameters for $a > 0$ here obtained, as well as the deviations among the currently available experimental data (Table III). A fundamental difference between the physics for $a < 0$ (where a more robust universal picture was found 12,21) —see Refs. 3,47 for a summary of such experimental findings—and for $a > 0$ is that corrections for the energy of the weakly bound $s$-wave dimer, whether originated from mixing of $s$- and $d$-wave interactions or finite-width effects, should already lead to modifications on the $a > 0$ three-body parameters. For $a_+$, the atom-dimer channel controls the interference effects in recombination via the exit channel while it represents the initial collision channel responsible for the resonant effects determining $a_+$. In fact, under this perspective, a simple criteria can be established to determine whether $s$- and $d$-wave mixing and finite-width effects are important: if the degree of deviation between the binding energy obtained from multichannel interactions and the one obtained from single channel models are substantially different, such effects are likely to be important.

**IV. SUMMARY**

In conclusion, our present study establishes the universal values for the three-body parameters $a_+$ and $a_+$, both relevant for ultracold quantum gases with positive scattering lengths. One of the most interesting results that has emerged from this study is the fact that the first excited Efimov resonance fails to intersect the dimer threshold, which is a surprising difference from the zero-range universal theories that always predict such an intersection. Our interpretation, that this failure of the resonance to intersect the threshold derives from important $d$-wave interactions, is consistent with findings from another recent study of this $a > 0$ region 63, which uses a nonlocal potential model having no $d$-wave physics, and which does show such an intersection. The robustness of the present prediction thus hinges critically on whether the $d$-wave two-body physics is tightly constrained in the way predicted by van der Waals physics in single channel potential models 29,57. Whether it is reasonable to expect that in the case of broad two-body Fano-Feshbach
resonances, this linkage of two-body s-wave and d-wave resonance positions is satisfied, remains an open question deserving further investigation. However, especially in the case of narrow two-body resonances, s-wave and d-wave resonances are likely to be largely uncorrelated which presumably invalidates the present predictions in the vicinity of $a/r_{vdw} \approx 1$. Nevertheless, the qualitative agreement between our results and the currently available experimental data partially confirms the notion of universality of Efimov physics for ultracold atoms. However, more experimental data and more sophisticated theoretical models incorporating the multichannel nature of the atomic interactions might be necessary in order to quantitatively address present discrepancies.

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