Van der Waals five-body size-energy universality

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A universal relationship between scaled size and scaled energy is explored in five-body self-bound quantum systems. The ground-state binding energy and structure properties are obtained by means of the diffusion Monte Carlo method. We use pure estimators to eliminate any residual bias in the estimation of the cluster size. Strengthening the inter-particle interaction, we extend the exploration from the halo region to classical systems. Universal scaled size-scaled energy line, which does not depend on the short-range potential details and binding strength, is found for homogeneous pentamers with interaction potentials decaying at long range predominantly as $r^{-6}$. For mixed pentamers, we discuss under which conditions the universal line can approximately describe the size-energy ratio. Our data is compatible with generalized Tjon lines, which assume a linear dependence between the binding energy of the pentamers and the one of tetramers, when both are divided by the trimer energies.

Universality in few-body systems connects physical systems at vastly different energy and length scales. It manifests as the independence of system’s characteristics upon the shape of the interaction potential and the length scale. The most famous universal phenomenon is the Efimov’s prediction of the geometric series of three-body bound state levels which occur when two-body state has zero energy, i.e., in the unitary limit. Although the first Efimov candidates were expected in nuclear physics, the first signature came from an ultracold gas of cesium atoms. This was possible due to the ability to control interactions between atoms by a magnetic field, thanks to the presence of Feshbach resonances. The Efimov effect was very soon observed in other cold atom systems, including those with non-identical particles and $(N > 3)$-body systems, in which a variety of universal bound states linked to the Efimov trimer was found. A further unexpected van der Waals universality appeared for three atoms interacting with potential $-C_0 r^{-6}$ in the ultracold regime, near Feshbach resonances. The ground-state trimer dissociation scattering length $a_{3}$, which acts as a three-body parameter, appeared universally proportional to the van der Waals length $\lambda_{vdW}$. Wang et al. explained the emergence of an effective repulsive three-body barrier, which prevents the three particles from getting close together, thus preventing configurations with small hyperradii, $\rho > 2\lambda_{vdW}$. In the limit of zero-range interactions and large scattering lengths, there are evidences for scales beyond three-body and the consequent necessity of a four-body scale when particles interact through an attractive contact or soft core pairwise potential.

The Efimov effect was also observed by the Coulomb explosion imaging in the experimentally elusive atomic trimer $^3\text{He}_3$, which is weakly-bound under natural conditions. Clusters which are even more weakly bound than $^3\text{He}_3$ present also different types of universality. Importantly, they are examples of quantum halo states, i.e., systems which prefer to be in classically forbidden regions of the space. Their large spatial extent makes the details of their interparticle interactions less important, leading to universal properties. The search for a universal relation between size and energy, in quantum halo states, began in nuclear physics and was later continued in atomic systems. The precise knowledge of the interparticle interactions in atomic clusters made it possible to determine universal ground-state size-energy ratios in weakly-bound dimers, trimers, and tetrathers. The progress in Coulomb explosion imaging enabled measurements of the distribution functions in dimers, trimers, and tetrathers of Argon and Neon, as well as weakly-bound Helium trimers and dimer $^4\text{He}_2$. A thorough analysis for a large set of pure and mixed weakly-bound atomic dimers, trimers, and tetrathers showed that universal size-energy scaling extends even below the halo area, in the so called quasi-halo region. Four-body systems with large size were found in Helium and Helium-alkali tetrathers but also in pentamers. It is therefore interesting to explore the existence of a universal relation between energy and size of five-body clusters, in a wide range extending from weakly-bound quantum halo systems to classical ones.

Close to the unitary regime, Tjon predicted a linear relation between the binding energy of the $\sigma$ particle and the triton, which was shown to approximately hold for different nuclear models. It was argued that, in the universal regime, a four-body parameter is not needed for determining the energy of the four-body cluster. The
so-called Tjon lines were later investigated in atomic systems close to unitarity\textsuperscript{32–36}. Hanna and Blume\textsuperscript{33} did not find that the energies of \(E_{N+1}\) and \(E_N\) clusters are well described by linear relations. However, they and others\textsuperscript{34–36} showed an approximate validity of generalized Tjon lines connecting linearly the relative energies, \(E_{N+1}/E_{N-1}\) and \(E_N/E_{N-1}\). There are some differences between the predictions of the generalized Tjon lines slope in previous studies, that occur most likely due to analysis of different ranges around the universality limit\textsuperscript{37}. Additionally, Yan and Blume\textsuperscript{39} showed that, at unitarity, the energies of few-body systems are not fully independent of the shape of the two-body short-range potentials. However, they found that in the case of van der Waals two-body interactions the binding energies at unitarity are approximately given solely in terms of the van der Waals length. It has not been reported how the relationship between \(E_{N+1}/E_{N-1}\) and \(E_N/E_{N-1}\) changes when moving away from the unitary limit, in direction of even more weakly bound states or when approaching the classical limit, or how well realistic atomic clusters approach the results obtained by model Lennard-Jones systems in these limits. Such findings are relevant for a better understanding of the limits of universality in Lennard-Jones systems.

In the present work, we study the energies and sizes of five-body Lennard-Jones clusters with the goal of determining the extension of their universality, from strongly to extremely weakly bound systems, that can be regarded as quantum halo states. Besides model systems, we study a range of realistic clusters containing up to three different atomic species. We rely on the use of quantum Monte Carlo simulations which provide exact results, within some statistical errorbars. We also compare the obtained five-body energies with the energies of four and three-body Lennard Jones systems in order to test the accuracy of generalized Tjon lines.

The rest of the paper is organized as follows. Section Methods describes the quantum Monte Carlo methods used in our work and introduces the energy and size scaling. Section Results discusses first five-body size-energy universality and then the obtained Tjon lines. The main conclusions of our work are summarized in Sect. Conclusions.

Methods

The ground-state properties, energy \(E\) and mean square of inter-particle separations \((\langle r^2 \rangle)\), were obtained by solving the Schrödinger equation

\[
-\frac{\hbar^2}{2m} \frac{\partial^2 \Psi(R,\tau)}{\partial \tau^2} = \left( H - E_\tau \right) \Psi(R,\tau),
\]

written in imaginary-time \(\tau = it/\hbar\), for the Hamiltonian \(H\). The reference energy \(E_\tau\) is introduced for numerical convenience. The positions of particles in five-body systems are stored in the so-called walker \(R = (r_1, r_2, r_3, r_4, r_5)\). The Schrödinger equation is solved stochastically utilizing the second-order diffusion Monte Carlo (DMC) method\textsuperscript{38} which, within statistical errorbars, leads to the calculation of the exact binding energy \(B = -E\), when the time-step \(\Delta \tau \to 0\), the imaginary time \(\tau \to \infty\), and the number of walkers \(\to \infty\).

As usual, importance sampling is introduced in DMC\textsuperscript{38} to reduce the variance by multiplying the ground-state wave function by a trial wave function optimized using the variational Monte Carlo (VMC) method. Estimators which do not commute with the Hamiltonian, e.g. \((\langle r^2 \rangle)\), can be biased by the mixed distributions produced by the use of importance sampling. In order to completely remove any bias from the trial wave function, we do not use the extrapolation approximation \((\langle r^2 \rangle)_{\text{VMC}} \approx 2(\langle r^2 \rangle)_{\text{DMC}} - (\langle r^2 \rangle)_{\text{VMC}}\) but implement much more sophisticated pure estimators\textsuperscript{39} to get unbiased estimations. Masses and trial wave-functions were taken from our previous works\textsuperscript{21,22,27,29,40}. The use of pure estimators proved to be successful in Helium clusters\textsuperscript{41}, where theoretical predictions on distribution functions reproduced accurately experimental results\textsuperscript{18,24} drawn from Coulomb explosion imaging.

We are interested in universal relations, so it is not crucial for us to use the most realistic potential, but to calculate accurately the ground-state energy and size of a system, for a given potential and particle masses. Our potential function sums only pair interactions. We take the Lennard-Jones (LJ) \(12-6\) model \(V(r) = 4\epsilon[(\sigma/r)^{12} - (\sigma/r)^6]\), with adjustable depth \(\epsilon\) and zero-point distance \(\sigma\), for a systematic exploration of van der Waals pentamers. For real clusters, we use the following model potentials: JDW\textsuperscript{42} for spin-polarized hydrogen isotopes \(2^3\)H \(_2\), denoting \(^1\)H also as T; Silvera\textsuperscript{43} for hydrogen molecules \(^3\)H\(_2\); TW\textsuperscript{44}D, DWW\textsuperscript{45}, TY\textsuperscript{46}, MF\textsuperscript{47} and MFmod\textsuperscript{48} for He-\(\text{H}\); semi-empirical HFDB\textsuperscript{49} for helium isotopes \(^4\)He; KTTY\textsuperscript{50} for interaction of an alkali metal isotope and a helium isotope; and TT\textsuperscript{51} for noble gases Ne and Ar.

To be able to compare quantities differing in several orders of magnitude, we scale the energy and the size with a characteristic length and analyze dimensionless quantities. Similar to what was done in previous works\textsuperscript{5,21}, we measure the size of the system by the mean-square hyperradius with subscript \(r\),

\[
\rho^2_r = \frac{1}{Mm} \sum_{i<k}^N m_im_k(r^2_{ik}),
\]

with \(r_{ik} = |r_i - r_k|\) and \(M\) the total mass of the \(N\)-body system. The particle masses \(m_i\) are given in an arbitrary mass unit \(m\). The characteristic hyperradius \(\rho^2_r\) is defined by substituting in Eq. (2) the pair size \(r^2_{ik}\) by the square of the corresponding van der Waals length,

\[
r^2_{ik} = \left( \frac{\mu C_6}{\hbar^2} \right),
\]

where \(C_6\) is the dispersion coefficient and \(\mu = m_im_k/(m_i + m_k)\) the reduced mass of a given pair. Notice that there is a different definition in the literature\textsuperscript{8} for this length, \(\rho_{vdW} \approx 0.5R\). In previous research of four-body systems\textsuperscript{22}, we showed that the van der Waals length \(R\) is convenient for scaling weakly and strongly bound systems, and
it was also used in the context of universal relations. We scaled the size of the pentamers as $Y_\rho = \rho^2 r_\rho - 2 R$ and analyze in the next Section how it depends on the dimensionless scaled binding energy, $X_E = m B \rho^2 R^2 / h^2$.

### Results

First, we discuss homogeneous five-body quantum systems $A_5$, i.e., van der Waals clusters of five identical atoms or molecules $A$. In a previous study of four-body systems, no mass effect on scaling was noticed. Therefore, and for practical reasons, we first explored clusters of particles with equal mass $m_i = 4u$, multiple of the atomic mass constant $u$. As a pair-potential model, we chose LJ 12-6 $V(r) = \varepsilon \left( (r/r_m)^{12} - 2(r/r_m)^6 \right) = 4\varepsilon \left( (\sigma/r)^{12} - (\sigma/r)^6 \right)$, where $-\varepsilon$ is the minimum at inter-particle separation $r = r_m = \sqrt{2\sigma}$, and $r = \sigma$ is the zero-point of the potential. The dispersion coefficient in this case has the simple form $C_6 = -\varepsilon r_m^6 m$. We used a repulsive core $\sigma = 4$ Å and varied the potential depth $\varepsilon$. This allowed us to explore a wide range of binding strengths, from $0.08$ mK for weakly-interacting ($\varepsilon = 3.32$ K, $\sigma = 4$ Å) to $50.752$ K for strongly-interacting ($\varepsilon = 20$ K, $\sigma = 4$ Å) pentamers. Corresponding $\langle r^2 \rangle$ appear in reverse order, from $990$ Å$^2$ to $36$ Å$^2$. The scaled size $Y_\rho = \rho^2 r_\rho^2$ and energy $X_E = m B \rho^2 R^2 / h^2$ for these model systems are shown with points in panel (a), Fig. 1. They span many orders of magnitude so a logarithmic scale is used.

In an homogeneous pentamer $A_5$, there are 10 equal $A$-$A$ pairs of particles so we distinguish between clusters with zero or ten self-bound sub-dimers, represented with empty and full symbols, respectively. As one can see, all pentamers in Fig. 1 follow the same law, regardless of the interaction potential and number of sub-dimers. The empirical function, similar to the four-body universality, fitted well (thin blue line) the obtained data for the scaled energies below $10^6$. The parameters of the best fit are reported in the Table 1.

![Figure 1](image)

**Figure 1.** Scaled size-energy fit for various homogeneous quantum five-body systems $A_5$. Interactions are modeled by (a) LJ 12-6 pair potential and (b) potentials for realistic systems. Pentamers are classified according to the number of self-bound sub-dimers. For comparison, we report the classical approximation, with $Y_\rho$ given by Eq. (7).

| $Y_0$  | $X_0$  | $\xi$  | $k$  | $n$  |
|-------|-------|--------|------|------|
| $10^{-6}$ | 16.5348(64) | 8.86(38) | 0.8830(61) | 28.88(29) |

**Table 1.** Parameters of Eq. (4) that fit the DMC data in Fig. 1. Figures in parenthesis are the statistical errors.

In the limit $T \rightarrow 0$ and for strong interactions ($B \rightarrow \infty$), a system becomes classical and its structure is defined by its minimum potential energy. Two-, three- and four-body classical systems rest in equilateral geometrical arrangements with all inter-particle separations equal to the position of the pair-potential minimum $r = r_m$. Respectively, particles are located at the vertices of the line segment, triangle and tetrahedron, which are one-, two- and three-dimensional geometry objects. The structure of a five-body system is more complicated because it is not possible to form a geometrical structure in three-dimensional space where all vertices are equally separated. As an optimal structure in this case, we take a triangular dipyramid, i.e., a double tetrahedron with common base. Then, nine pairs span 9 edges of length $r_m$ and contribute to the binding energy by $\varepsilon$. The structural function, similar to the four-body universality,
remaining pair spans the only spatial diagonal whose length corresponds to double height of tetrahedron, $2H = \sqrt[3]{3/\rho_m}$, and thus contributes to the binding by an amount $\varepsilon \left( \frac{\rho_m^2}{2} \right)^{1/2} - 2 \left( \frac{\rho_m}{2} \right)^3$. If we take the mass of a particle as the mass unit, the hyperradius simplifies,

$$\rho_2^2 = \frac{me^2}{Nm^2} \left[ 9 \rho_m^2 + (2H)^2 \right] = \frac{7}{3} \rho_m^2.$$  \tag{5}

Scaling the size (5) with the characteristic hyperradius $\rho_2^2 = 2R^2$, as well as the binding energy,

$$X_E = \frac{mB\rho_2^2}{h^2} = \frac{2m\varepsilon}{h^2} \cdot \frac{2386215}{262144} \cdot R^2,$$  \tag{6}

leads to a straightforward relationship between scaled size and energy of classical systems,

$$Y^C_{\rho} \approx \left( 14.5/X_E \right)^{1/2}.$$  \tag{7}

This classical line is plotted in Fig. 1 as a thick dashed cyan line which, for scaled energies larger than $10^3$, smoothly continues the trend shown by quantum pentamers. All data of analyzed homogeneous five-body systems follow the same line. Thus, the universal law applies starting from purely quantum systems, defined by the relation (4), and then extends to classical systems, where for $X_E > 10^3$ it asymptotically takes a much simpler form (7). The universal quantum law starts differing from the simple classical estimation for scaled energies $X_E < 10^3$, when the contribution of the kinetic energy becomes significant, producing larger spatial structures than classical ones.

When mean particle separations become few times larger than van der Waals radius $R \sim \varepsilon^{1/4}$, while decreasing $\varepsilon$, the binding energy rapidly vanishes ($B \to 0$), but the size barely changes. In this case, particles are far away and pair potentials barely affect the probability outside the range of the van der Waals potential. That scenario is similar to the one of finite and contact interactions and thus, it is in agreement with previous theoretical findings\textsuperscript{10,31} that scaled size saturates in the weak binding limit. Weak binding, which does not support smaller clusters, holds pentamers through mediated interactions of additional particles.

If we change the short-range part, i.e., reduce the core size two times, $\sigma = 2 \AA$, we can see no effect in the scaling law thus confirming the universal ratio. Pentamers ($8\sigma_5$) for potential depths $\varepsilon = 8, 9, 14$ K, respectively, have ground-state binding energies $B = 554, 1318, 8768$ mK and sizes $(r^2) = 47.6, 34.0, 17.1 \AA^2$, which when scaled, $X_E = 3.36, 8.47, 70.3$, $Y^C_{\rho} = 2.59, 1.74, 0.70$, fit to the universal line.

In addition, we test the validity of the obtained law for the case of realistic homogeneous pentamers, whose interaction is formulated with elaborated potentials describing particles as induced fluctuating electric multipoles. Although their pair-potentials have different sort-range parts, they share the common feature that fall quickly with separation $r$ and so the London dispersion energy $-C_6 r^{-6}$ dominates at large $r$. The ground-state binding energy and size for the studied realistic systems ($T \downarrow_3$, $^4\text{He}_5$, ($^2\text{H}_2)_5$, Ne$_5$ and Ar$_5$ are reported in Table 2 and compared with the universal line in panel (b) of Fig. 1. They follow the universal line equally well, both in the regime of weak and strong binding.

**Second**, we test the validity of the obtained universal law for mixed realistic five-body systems consisting of up to three different components: spin-polarized H and He isotopes, an alkali atom, Ne, Ar, and hydrogen molecules H$_2$. Our results are summarized in Fig. 2.

In panel (a), different symbols are used to distinguish different species of particles that form the pentamer, while different filling of symbols is used to distinguish types of pentamers with regard to the number of self-bound sub-dimers. Among studied clusters, we distinguish pentamers that have 0 (empty symbols), 1 (quarter full), 3 (three quarter full), 5 (square with dash), 6 (two quarter full), 7 (two quarter empty), and 10 (full symbols) self-bound sub-dimers, as it is noted in the legend. The results with the realistic potential models are noted with additional black dots. Different symbols are spread everywhere and, at the first sight, it seems that no rule can be extracted regarding components, pentamer types or other characteristics. Only pentamers with all self-bound sub-dimers are always close to the A$_5$ line, while all other types can be on the line or above it. Only mixtures of Helium isotopes and an alkali atom sometimes go below the line. The latter can be understood in the following way. An alkali atom has a much larger electronic cloud repulsive core than Helium isotopes, so Helium isotopes tend to form a cluster on one side of an alkali atom\textsuperscript{28}. This feature limits the arrangements that the pentamer can

| Cluster | $B$ / K | $(r^2)$ / Å$^2$ | $X_E$ | $Y^C_{\rho}$ |
|---------|--------|----------------|------|-------------|
| $^3\text{T}_5$ | 0.399(9) | 158.5(9) | 2.63 | 2.99 |
| $^3\text{He}_5$ | 1.335(1) | 59.4(4) | 6.37 | 2.05 |
| $^3\text{H}_5$ | 44.34(2) | 27.8(4) | 218 | 0.470 |
| Ne$_5$ | 224.29(2) | 11.1(2) | 24970 | 0.082 |
| Ar$_5$ | 1110.1(3) | 14.67(2) | 1108500 | 0.024 |

Table 2. The ground-state binding energy $B$, mean square pair size $(r^2)$, scaled energy $X_E$, and scaled size $Y^C_{\rho}$ for five-body realistic clusters. Figures in parenthesis are errorbars.
exhibit, and its size is reduced. Other mixed systems separate from the line at different points; separation points also differ for the same type of pentamers.

Fragmented-like systems separated from the line share a common feature; all of them have at least one particle which appears less bound than the others and that is significantly separated from the others. In panel (b) of Fig. 2 we made a different analysis. In this case, the ratio of largest $\langle r^2_{\text{Max}} \rangle$ and smallest $\langle r^2_{\text{Min}} \rangle$ pairs are compared. One can notice that if all pairs are similarly bound, strongly or weakly, i.e., if the ratio is between 1 and 1.5, mixed systems (circles) follow the $A_5$ line. Triangles and diamonds are also close to the line. Thus, if the ratio is below 4, noticeable deviations from the $A_5$ line can happen only for very weakly bound quantum systems, i.e., in the area where scaled energies are less than $X_E < 1$, while in other areas only small deviations can occur. According to the position on the graph, each system can be recognized from the panel (b) of Fig. 1 and the panel (a) of Fig. 2. The larger the ratio is, the larger are the energies for which beginning of the start of separation can be expected. This happens because a very weakly bound component makes negligible contribution to the system energy, but significantly increases its size. In this case, a small displacement along $X_E$ axis results in a significant displacement along $Y_\rho$ axis and the separation occurs. Thus, separations which occur for large scaled energies diverge faster, feature which was also noticed in the case of tetramers.

To illustrate the separation from the universal line, some estimated quantities are extracted in Table 3 for the cluster $^4\text{He}_4T_5$ using different potential models for the $^4\text{He}–T_5$ interaction. This allowed modeling different strengths of the $^4\text{He}–T_5$ interaction, which is in neither case strong enough to support a dimer bound state. Thus, $^4\text{He}_4T_5$ has 6 self-bound sub-dimers. The results from Table 3 are shown by five two-quarter full cyan symbols that are furthest to the right and above the line in the panel (a) of Fig. 2, connected by a short dashed blue line to guide the eye. They deviate very fast from the universal line when the $^4\text{He}–T_5$ attraction decreases, starting from the symbol with black point on the line. They deviate very fast from the universal line when the $^4\text{He}–T_5$ attraction decreases, starting from the symbol with black point on the line. In the case of the strongest $^4\text{He}–T_5$ interaction model MFmod, which is the most realistic one, mean square $^4\text{He}–T_5$ pair size is already 1.58 times larger than $^4\text{He}–^4\text{He}$ and it is exactly on the universal line. Using less attractive $^4\text{He}–T_5$ models, respectively, MF, DF, TY, DW, TW, binding weakens up to 25%, almost reaching the pentamer threshold limit, i.e., the ground-state energy of $^4\text{He}_4 = 577.6(3)$ mK, while ratio of squared pair sizes doubles. Further weakening of the $^4\text{He}–T_5$
Among the studied realistic clusters, the largest ratios of mean square radii are in $^4$He$_3$(D ↓)$_2$, where $(r^2)_{-3} = 108, 140, 1900 \, \text{Å}^2$, respectively for pairs $^4$He $- ^3$He $- ^3$He $- ^3$He $- ^3$He and $D ↓ - D ↓$ and $D ↓ - D ↓$. Thus $(r^2)_{\text{Max}}/(r^2)_{\text{Min}} = 17.9$, while the binding energy is only 7 % larger than the energy of the trimer$^{41}$. $^4$He$_3$. On average He atoms are close to each other forming a sub-trimer $^4$He$_3$ which is surrounded with a halo cloud of far away $D ↓$ atoms. Their weak binding is barely mediated by $^4$He$_3$ so they very rarely find themselves on the same side of the $^4$He$_3$, contributing largely to the size of the cluster. Deviation from the line is obvious, $X_E = 0.58, Y_\rho = 18.5$. After substituting $D ↓$ with the heavier isotope $^1$T ↓, which has lower kinetic energy, the cluster becomes homogeneous-like with almost ten times lower ratio $(r^2)_{\text{Max}}/(r^2)_{\text{Min}} = 1.85$.

Third, we compare the present results with previous findings. The size-energy scaling laws for five-body systems (4) and (7) are compared in Fig. 3 with the four-body universal law$^{22}$. Both lines intersect at $(0.059, 11.1)$. In the classical limit, for the same scaled energy, strongly bound pentamers have larger scaled size than tetramer at the binding threshold. The quantitative comparison of the system size with our results is possible, because we used long-range decay which is characteristic for London dispersion forces between atoms and molecules that are electrically symmetric. Although they explored only the weak-binding regime, crossing of tetramer and pentamer curves is also just noticeable close to the end of their researched area.

Brunnian systems also show qualitatively the same behavior$^{22}$.

Having previously studied also the trimer$^{22}$ and tetramers$^{22}$, we are able to analyze their energies in comparison with the present pentamer results. Fig. 4 reports the generalized Tjon line, which describes the dependence of the energy ratios $E_5/E_3$ and $E_4/E_3$, where $E_N$ is the ground-state binding energy of the $N$-body system.

Rasmussen et al.$^{26}$ studied how the two lowest-lying weakly bound states of few bosons depend on the strength of two-body Gaussian interactions $V(r) = V_0 \exp(-r^2/b^2)$, where $b$ was chosen as the characteristic length scale. Their results are in qualitative agreement with ours. They also predicted that the pentamer has lower scaled size than the tetramer at the binding threshold. The quantitative comparison of the system size with our results is not possible, because we used long-range decay $-C_6 r^{-6}$, which is characteristic for London dispersion forces between atoms and molecules that are electrically symmetric. Although they explored only the weak-binding regime, crossing of tetramer and pentamer curves is also just noticeable close to the end of their researched area.

Brunnian systems also show qualitatively the same behavior$^{22}$.

Interaction would cause distancion of the T atom from the remaining tetramer, i.e., scaled size would diverge fast in logarithmic scale because scaled energy $X_E = 3.29$ is already close to the threshold limit $X_E = 2.74$, when the pentamer dissociates into $^4$He$_4$ and far away free T atom. The ground-state properties for all studied realistic systems are given in Supplementary Table S1 online.

Table 3. Van der Waals length $R$, mean square pair size $(r^2)$, ground-state energy $E$, scaled energy $X$ and scaled size $Y$, in the pentamer $^4$He$_4$T ↓ modeled with different pair potentials.

| Potential model | $R$ / Å | $(r^2)$ / Å$^2$ | $(r^2)_{\text{Max}}$ / Å$^2$ | $(r^2)_{\text{Min}}$ / Å$^2$ | $|E| / \text{mK}$ | $X$ | $Y$ |
|-----------------|---------|----------------|----------------|----------------|----------------|-----|-----|
| HFDB$^{44}$    | 5.38    | 6.69           | 65(1)          | 103(2)         | 1.58           | 885.7(7) | 4.74 | 2.27 |
| HFDB$^{45}$    | 5.38    | 6.10           | 65(1)          | 106(1)         | 1.63           | 866.9(8) | 4.30 | 2.48 |
| HFDB$^{46}$    | 3.58    | 5.75           | 66(1)          | 121(2)         | 1.83           | 791.4(9) | 3.75 | 2.79 |
| HFDB$^{47}$    | 3.58    | 6.10           | 67(1)          | 172(3)         | 2.57           | 682.8(9) | 3.39 | 3.22 |
| HFDB$^{48}$    | 3.58    | 6.10           | 67(2)          | 202(3)         | 3.01           | 662.8(9) | 3.29 | 3.55 |

Figure 3. Comparison of the universal size-energy scaling laws for four$^{22}$ and five body systems.
feature of a long-range behavior dominated by interparticle interactions were modeled by pair potentials with different short-range shape, but with the common tendency to the threshold limit, in the halo region, up to the limit of maximum binding of purely classical clusters. The approach of the trimer threshold (not shown in Fig. 4 to avoid loss of clarity), the estimated ratios $E_4/E_3 = 11.85$ and $E_5/E_3 = 34.63$ for a model system, with $a = 4\, \text{Å}$ and $\varepsilon = 4\, \text{K}$, when $E_3 = -8\, \text{mK}$, also verify the empirical relation.

Recent estimates of $^4\text{He}_5$, binding energies, obtained within the framework of effective field theory at leading order and next-to-leading-order with a four-body force, that renormalizes the four-body system, respectively, $E_4/E_3 = 4.8(1), 4.35$ and $E_5/E_3 = 10.8(5), 11.3(3)$ deviate from the empirical relation, but also have large extrapolation errors. Our results with the HFDB potential are $E_4/E_3 = 4.335(6)$ and $E_5/E_3 = 10.02(2)$.

Our results show that the linear law is valid only for a limited range of ratios $E_4/E_3$. Increasing the attraction strength, and leaving the regime of weak binding, the slope collapses non-linearly and abruptly towards the classical boundary, where $E_4/E_3 = (6\varepsilon)/(3\varepsilon) = 2$ and $E_5/E_3 = 795405/262144 \approx 3.03$. Our realistic clusters $\text{Ne}_5$ (2.015, 3.07) and $\text{Ar}_5$ (2.004, 3.04) are very close to the classical ratio limit. This is to be expected, as the binding energy $1.110\, \text{kK}$ of $\text{Ar}_5$ is very close to the classical limit $1.306\, \text{kK}$.

**Discussion**

Five-body systems composed of one, two, and three different particles were explored by means of quantum Monte Carlo methods at $T = 0\, \text{K}$. Different strengths were analyzed, from very weak binding in quantum systems close to the threshold limit, in the halo region, up to the limit of maximum binding of purely classical clusters. The interparticle interactions were modeled by pair potentials with different short-range shape, but with the common feature of a long-range behavior dominated by $-C_6 r^{-6}$. This common characteristic enabled a simple choice of characteristic length for classical and quantum systems, the van der Waals length, which was used for defining the scaling energy and size.

The universal law, which relates scaled size and energy, has been found for homogeneous pentamers in their ground-state. For medium and weakly bound systems, it shows a non-linear non-logarithmic shape (4) valid for scaled energies $X_E < 10^6$, while after $X_E > 10^7$ it approaches its asymptotic simple linear shape (7) in log-log scale. The law is applicable if the pair potential asymptotically follows as $-C_6 r^{-6}$, while slower or faster decrease would produce a universal law with different log-log slope, as it can be deduced from the classical analysis. In the limit of the binding threshold, the scaled size of homogeneous pentamers monotonously converge to the finite value 15, below the tetramer size of $25^{[2]}$. Noticeably, this plateau is not present in the case of dimers and trimers which instead show a diverging size approaching the threshold for binding$^{[3,10,21]}$.

The universal size-energy line is also applicable to mixed systems which are homogeneous-like, i.e., if the distance between all the constituents is similar. If the mean square distance of the largest particle pair is few times larger than the shortest one, then mixed system could deviate above the line. In addition we find that if the cluster is spatially constrained, its size is reduced, so it can appear slightly below the line.

Finally, we analyzed the relationship of pentamer, tetramer, and trimer energies of homogeneous systems, confirming the range of approximate validity of generalized Tjon lines and demonstrating the convergence of Lennard-Jones systems to the classical limit.

**Data availability**

The data that support the findings of this study are available within the article and its Supplementary Information.
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Author contributions
P.S. calculated ground-state properties and fitted the results. L.V.M. compared data with the generalized Tjon lines. All authors discussed the results and reviewed the manuscript.

Competing interests
The authors declare no competing interests.

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