Effective field theory for the bound states and scattering of a heavy charged particle and a neutral atom

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We show the system of a heavy charged particle and a neutral atom can be described by a low-energy effective field theory where the attractive $1/r^4$ induced dipole potential determines the long-distance, low-energy wave functions. The $1/r^4$ interaction is renormalized by a contact interaction at leading order. Derivative corrections to that contact interaction give rise to higher-order terms. We show that this "induced-dipole EFT" (ID-EFT) reproduces the π^+ -hydrogen phase shifts of a more microscopic potential, the Temkin-Lamkin potential, over a wide range of energies. Already at leading order it also describes the highest-lying excited bound states of the pionic-hydrogen ion. Lower-lying bound states receive substantial corrections at next-to-leading order, with the size of the correction proportional to their distance from the scattering threshold. Our next-to-leading order calculation shows that the three highest-lying bound states of the Temkin-Lamkin potential are well described in ID-EFT.

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I. INTRODUCTION

When a charged particle interacts with a neutral atom in an *S* state at distances significantly larger than the Bohr radius, it experiences an attractive $1/r^4$ potential with a strength given by the atom's polarizability α [1]. The theory of singular potentials [2] therefore governs this situation, producing a particular pattern of bound states and scattering of the charged particle from the atom as a function of energy. Low-energy properties are given by $1/r^4$ dynamics, just as the $1/r^6$ potential determines the phase shifts and bound states of atom-atom systems for wave numbers of order one over the van der Waals length scale. The analog of the van der Waals length scale is $\beta_4 \equiv \sqrt{\mu\alpha/4\pi\epsilon_0 m_e a_B}$, where μ/m_e is the reduced mass of the atom-particle system in units of the electron mass and a_B is the Bohr radius.

Traditionally most attention has been devoted to electron scattering [3], in particular the simplest case of the hydrogen ground state, whose polarizability is $\alpha_{\rm H} = 9a_B^3/2$ [1,4]. Of particular current interest is the scattering of a heavier particle such as a negative muon (μ^-), a positive pion (π^+), or a proton. These systems have richer spectra than the electron case does, and rearrangement channels open up when the projectile is negatively charged, or positively charged and heavier than the proton.

Negatively charged heavy particles can also be captured in states with high orbital quantum number and cascade down

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to lower states, where they provide sensitive probes of nuclear properties [5]. The quantum mechanics of their higher ionic levels is determined by the atomic polarizability and so it can also be explored with positively charged heavy particles.

Such ions have $\beta_4 \gg a_B$. This scale separation enables an effective field theory (EFT) treatment of this problem. EFT is a general tool that uses the separation of scales within a system, or class of systems, to make systematically improvable predictions for observables within a well-defined energy window. (For an introduction to EFT, see Ref. [6], while Ref. [7] constitutes a pedagogical guide to implementing EFT principles in a Schrödinger equation setting.) EFTs can be used in all areas of physics to account systematically for small effects which are frequently simply neglected. Nuclear and atomic systems are particularly interesting because they require the most important, leading-order (LO) interactions to be treated nonperturbatively and generate bound states and resonances. They also frequently involve long-range interactions, such as the Coulomb potential. Nonsingular long-range interactions do not impose significant modifications to the case of purely short-range interactions (see, for example, Ref. [7]). Singular long-range interactions, on the other hand, dramatically affect the theory at small distances. EFTs for singular potentials have been studied extensively [8–16] for their relevance in atomic and nuclear physics [17]. They update and systematize the work of Case [18] and others on singular potentials in quantum mechanics [2]. The $1/r^3$ potential is especially relevant as it is a leading piece of the interaction between two nucleons in chiral EFT [17, 19-24] in the limit that the pion mass is taken to zero [25,26].

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The impact on observables of the $1/r^6$ potential in a van der Waals EFT has been discussed recently [27]. Van der Waals EFT is similar to quantum defect theory, which has been applied extensively to predict bound states and scattering in the situation that a $1/r^6$ potential determines the long-distance wave function of an atom-atom system [28,29]. Similar calculations have been done for the attractive $1/r^4$ potential [30].

In contrast to the goals of EFT, where an accurate description of low-energy physics is primary, the semiclassical Wentzel-Kramers-Brillouin (WKB) approximation is relevant in energy regimes where potential variations are slow in comparison to the wavelength of the wave function. The WKB predictions for attractive $1/r^n$ potentials were derived in Ref. [31]. At low energies, and certainly at threshold, the WKB approximation does not work well. Near threshold, the effective-range expansion (ERE) developed for finite-range potentials can be modified to account for the long-range tail; doing so generates additional terms that are nonanalytic in energy. For $1/r^4$, these terms can be expanded [32,33] as a series in powers of the wave number in units of β_4 , $k\beta_4$, times powers of β_4/a_0 , where a_0 is the scattering length. The series starts with a linear correction to $k \cot \delta$.

Instead of using either the ERE or a WKB approach, here we develop an EFT, induced-dipole EFT (ID-EFT), that does not employ an expansion in $k\beta_4$, but instead expands observables in powers of the ratio a_B/β_4 and the wave number in atomic units, ka_B [34]. At LO in this expansion the effect of the finite size of the atom on observables is captured in a smeared delta function. The Schrödinger equation is then solved for a potential consisting of that delta function and the $1/r^4$ induced-dipole potential. While the strength of the $1/r^4$ potential is fixed to be β_4^2 , the strength of the delta-function piece depends on short-distance details of the atom-charged particle interaction and must be fit to one datum. Accuracy is improved in a next-to-leading-order (NLO) calculation, which introduces an additional piece of the short-distance potential, with an additional coupling constant, that must be fit to an additional datum. The same procedure is repeated at higher orders

In this problem the standard ERE is limited to wave numbers k for which $k\beta_4 \lesssim 1$. In contrast, ID-EFT can handle wave numbers comparable to $1/\beta_4$. ID-EFT can be regarded as a systematic implementation of a modified ERE [35] that includes the long-range effects of the $1/r^4$ potential on the relevant matrix elements. It can therefore be derived by expanding the short-range potential in local operators, and considering matrix elements of that operator on the basis of distorted waves [20,36–38]. The standard ERE in powers of k described above then results if the induced-dipole potential is treated in perturbation theory as a higher-order effect.

As a specific example, we consider here a heavy charged particle for which we can expect several bound states within the regime of validity of the theory. Since an LO description of this system in which the potential consists solely of a delta function can support at most one bound state, an EFT description of multiple bound states relies on the (singular) interaction being included as a leading effect. To be definite, we study the bound states and phase shifts of the π^+ -hydrogen ion. The spectrum is rich but scattering is not afflicted by open rearrangement channels. We use the pion-atom scattering

PHYSICAL REVIEW A 108, 062817 (2023)

The additional short-range parameter is fitted to the shallowest bound-state energy. Other energy levels in the system, as well as the scattering phase shifts, are then predicted by ID-EFT, up to corrections to each observable that have a fractional size $\sim k^4 a_B^4$, where k is the characteristic wave number of the scattering or bound state. An important aspect of our calculation is that by carrying it out for several different choices of the delta-function smearing we can assess which of our observable predictions are independent of the details of this short-distance piece of the potential.

Several studies, some of which treated $\pi^+ H$ as a twobody system and some of which treated it as a three-body system, have been conducted previously [39-41]. In the twobody treatment the pion-atom interaction was taken to be an analytical, parameter-free potential-the Temkin-Lamkin polarization potential [3,42,43]—that superposes some shortrange effects onto the $1/r^4$ tail. (The Temkin-Lamkin potential is also useful in complementing approximate solutions of the three-body system [44].) We take the Temkin-Lamkin potential's results for the $\pi^+ H$ system's bound-state energies and S-wave scattering phase shifts as data that allow us to assess the efficacy of ID-EFT for this system. We use those results as a laboratory to demonstrate the ability of ID-EFT to capture the low-energy portion of the rich spectrum and multifaceted phase-shift behavior that results from the induced-dipole interaction. We are particularly interested in the fact that in many other applications of singular potentials-for example, in nuclear physics-one contends with a single bound state, while the Temkin-Lamkin potential produces seven bound states in the π^+H system. It is therefore interesting to see how certain details of the EFT renormalization and calculation play out in this more complex situation. Results from the Temkin-Lamkin potential are clean and allow sharp conclusions about the convergence of the EFT and its breakdown scale, conclusions that were much fuzzier in other contexts where a LO singular interaction was considered—for example, Ref. [45].

Nevertheless, the Temkin-Lamkin treatment of π^+H is an approximation, and there are significant corrections to that approximation in a full three-body treatment [39]. Having developed the basic ideas of ID-EFT in this work, we intend to return to this problem in subsequent papers. There we will instead use data from three-body treatments of the π^+H system as input to our EFT.

The remainder of our paper is structured as follows. Our theoretical formulation is given in Sec. II, with details of its implementation relegated to Appendixes A–C. The bound-state and scattering results at LO and NLO are presented and discussed in Sec. III, while details of extrapolations to small cutoffs are given in Appendix D. Appendix E describes the evaluation of WKB phase shifts to which we compare our results. Conclusions and future prospects are discussed in Sec. IV.

II. THEORY

A. Leading order

The LO interaction in ID-EDT takes the coordinate-space form

$$V(r) = -\frac{C_4}{r^4} \ \rho(r; R) + g_{\rm LO}(R) \ \chi(r; R), \tag{1}$$

which, for context, is input to the radial Schrödinger equation at energy E, TL

$$\left[-\frac{\hbar^2}{2\mu}\frac{d^2}{dr^2} + V(r)\right]u(r) = Eu(r),$$
(2)

where μ is the reduced mass of the charged particle-atom system. The regulator functions $\rho(r; R)$ and $\chi(r; R)$ act, respectively, to overcome the $1/r^4$ potential and mimic the delta function at short distances. Both interactions are regulated at the radius *R* related to the short-distance physics that we account for at LO through a contact interaction of strength $g_{\text{LO}}(R)$. The precise forms of $\rho(r; R)$ and $\chi(r; R)$ are not important, only that

$$\lim_{r \to 0} \rho(r; R) / r^4 = 0,$$
(3)

$$\lim_{R \to 0} \chi(r; R) \propto \delta(r). \tag{4}$$

Here, we take

$$\chi(r;R) = e^{-(r/R)^4},$$
(5)

$$\rho(r;R) = [1 - e^{-(r/R)^2}]^4.$$
(6)

Once $g_{\text{LO}}(R)$ is determined from one low-energy datum, LO is renormalized [8], namely, other low-energy observables converge as 1/R increases beyond the breakdown scale of the theory, $\Lambda_b \sim 1/a_B$.

The short-range interactions of ID-EFT capture the lowenergy effects of physics at distances comparable to the atom's size. While the asymptotic form of the potential is $1/r^4$, as the charged particle approaches its impact on the atom's distortion can no longer be accounted for solely by the polarizability [46]. Here, as an illustration of the method, we tune $g_{LO}(R)$ at each value of R to match the scattering length, a_0 , obtained with the Temkin-Lamkin (TL) potential [3,42,43],

$$V_{\rm TL}(r) = -\frac{e^2}{8\pi\epsilon_0} \frac{\alpha(r)}{4\pi\epsilon_0} \frac{1}{r^4},\tag{7}$$

where *e* is the electron charge, ϵ_0 is the vacuum permittivity, and

$$\frac{\alpha(r)}{4\pi\epsilon_0} = \frac{9}{2}a_B^3 \left\{ 1 - e^{-2r/a_B} \left[1 + 2r/a_B + 2(r/a_B)^2 + \frac{4}{3}(r/a_B)^3 + \frac{2}{3}(r/a_B)^4 + \frac{4}{27}(r/a_B)^5 \right] \right\}.$$
 (8)

The TL potential is one of several semiphenomenological polarization potentials [47] that account for various effects associated with the interaction of the charged particle with the full charge distribution of the atom. However, at distances $r \gg a_B$ only the $1/r^4$ piece of the potential survives. Matching to Eq. (1), we determine the length-scale associated with this piece of the potential as $\beta_4^2 \equiv 2\mu C_4/\hbar^2$, where

$$C_4 = \frac{e^2}{8\pi\epsilon_0} \lim_{r \to \infty} \frac{\alpha(r)}{4\pi\epsilon_0} = \frac{1}{2} \left(\frac{e}{4\pi\epsilon_0}\right)^2 \alpha_{\rm H} \tag{9}$$

represents the strength of the $1/r^4$ potential and $\alpha_{\rm H} = 9a_B^3/2$ is the polarizability of the hydrogen atom [1,4]. Details at short distances are not important in this application, and other

potentials that curb the growth of $1/r^4$ would do as well. The TL potential is a rich example as it supports many bound states when the charged particle is heavy.

 β_4 and the scattering length a_0 are the two physical scales that are inputs to ID-EFT at leading order. In atomic units (a.u.), where $\hbar = e = m_e = 4\pi\epsilon_0 = 1$, lengths are given in Bohr radii, $a_B = 4\pi \epsilon_0 \hbar^2 / m_e e^2$; energies are given in Hartrees, $E_h = \hbar^2 / m_e a_B^2$; and $\alpha_{\rm H} = 9/2$. For the pion-hydrogen system, $\beta_4 = 32.7$ a.u. and the TL potential gives $a_0 = -65$ a.u. Details of the determination of a_0 in the presence of a $1/r^4$ tail are given in Appendix A. In addition, since this is a local theory, there are, in principle, infinitely many values of g_{LO} that yield the desired scattering length a_0 . Each such value produces a different number of bound states [8]. Here, we choose the branch of the implicit function $g_{LO}(a_0)$ that corresponds to 14 S-wave bound states, as described in Appendix B. There are only seven states allowed by the TL potential, but in order to study the renormalization of continuum and bound-state observables above the approximate breakdown scale of the theory, we choose a "lower" branch such that the repulsion of the LO counterterm is not numerically prohibitive.

With β_4 and a_0 fixed we can predict all binding energies at LO. We also compute the phase shifts

$$\delta^{(LO)} = \tan^{-1} \left(\frac{t^{(LO)}}{1 + it^{(LO)}} \right)$$
 (10)

from

$$t^{(\rm LO)} = V_{\rm LO} + V_{\rm LO} G_0 t^{(\rm LO)}, \tag{11}$$

in a short-hand notation where an integral over the momentum in the two-body propagator G_0 is implicit.

We note that the LO potential (1) contains no direct information on Λ_b . As is typical in EFTs, quantitative information on the breakdown scale enters the calculation only in the context of higher-order corrections.

B. Next-to-leading order

At next-to-leading-order (NLO) the interaction is modified to

$$V(r) = -\frac{C_4}{r^4} \rho(r; R) + [g_{\rm LO}(R) + Eg_{\rm NLO}(R)]\chi(r; R), \quad (12)$$

where we have chosen an energy-dependent NLO contribution to the short-distance potential. As is typical, a momentumdependent NLO term was first tried. It was found to cause unnecessary numerical difficulty. The energy-dependent scheme is less singular and therefore more tractable [37,48]. The new parameter $g_{\rm NLO}(R)$ is obtained from a second lowenergy datum, and other observables converge as R decreases as long as perturbation theory is employed. Corrections to the long-range polarization potential from the quadrupole polarizability and nonadiabatic contributions are $\propto 1/r^6$ [47]. They should be included at this order if a phenomenological analysis were to be performed. Here, we do not consider these corrections as we are interested only in demonstrating the ability of ID-EFT to reproduce the low-energy effects of a given underlying potential (chosen to be the parameter-free TL). There is no difficulty of principle in including them along the lines of Ref. [14], where a $1/r^4$ correction to an attractive $1/r^2$

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potential was considered. The additional long-range potential would give rise to perturbative corrections at large distances as well as a different running of $g_{\text{NLO}}(R)$ with R.

For bound states NLO corrections to the LO calculation are computed using standard first-order perturbation theory, i.e.,

$$B_{\rm NLO}^{(n)} = B_{\rm LO}^{(n)} \left(1 + g_{\rm NLO}(R) \left\langle \psi_{\rm LO}^{(n)} \right| \chi \left| \psi_{\rm LO}^{(n)} \right\rangle \right).$$
(13)

The value of $g_{\rm NLO}(R)$ is determined here by demanding that the binding energy of the shallowest *S*-wave state, $B_{\rm NLO}^{(6)}$, is fixed to the Temkin-Lamkin result for the pion-hydrogen system, 1.2×10^{-4} a.u.

Phase shifts are similarly computed in first-order perturbation theory according to the distorted-wave Born approximation (DWBA), described in detail for example in Ref. [49], where the scattering amplitude at on-shell momentum $k = \sqrt{2\mu E}$ is

$$t^{(\text{NLO})} = t^{(\text{LO})} - \frac{2\mu}{k} \langle \phi^{(\text{LO},-)} | V_{\text{NLO}} | \phi^{(\text{LO})} \rangle,$$
 (14)

with "-" denoting an outgoing wave. At NLO, we compute the phase shift perturbatively according to

$$\delta^{(\text{NLO})} = \delta^{(\text{LO})} - \frac{2\mu}{k} \langle \phi^{(\text{LO},-)} | V_{\text{NLO}} | \phi^{(\text{LO})} \rangle e^{-2i\delta^{(\text{LO})}}.$$
 (15)

Alternative but equivalent ways to calculate the NLO scattering amplitude are discussed in Appendix C.

III. RESULTS

In this section we present numerical results for the scattering of a charged pion on hydrogen, and for the bound states of this system. Solutions to Eq. (2) are computed with the Runge-Kutta method of order 8 with a relative tolerance of 10^{-8} and absolute tolerance of 10^{-12} . As R decreases, $g_{\rm LO}(R)$ becomes very large so it can provide the repulsion necessary to keep a_0 and the number of bound states fixed as more of the $1/r^4$ attraction is exposed. Although working with the 14-state branch alleviates the problem, we were unable to find accurate LO and NLO solutions once β_4/R became larger than 70. In order to calculate sufficiently precise phase shifts at LO with $\beta_4/R = 70$, g(R) was tuned to a relative accuracy of 10^{-12} . The total range of R values presented in this work is $\beta_4 =$ [20, 70] (or R = [1.59, 0.47] a.u.). As we are going to see, this range in β_4 extends high enough for many conclusions to be drawn about the scope of ID-EFT. At small R, a_0 must be calculated to high precision—approximately 0.01%. This limit was reached using the results of Appendix A. Convergence with respect to maximum r in the solution of Eq. (2) was demonstrated across a range of upper values up to 1000 a.u. The required precision was not reached until approximately 600 a.u.

A. Scattering

The efficacy of our proposed EFT is first tested in the continuum where we study the LO and NLO *S*-wave phase shifts. Figure 1 shows the ID-EFT predictions for the phase shifts, alongside the Temkin-Lamkin results, the ERE, and the WKB prediction of Ref. [31]. We find that the variation of the ID-EFT phase shifts for cutoffs near the breakdown scale is minimal: were we to draw bands of cutoff variation



FIG. 1. The *S*-wave phase shifts are shown as a function of the dimensionless quantity, $(k\beta_4)^2$. The EFT at LO (solid blue line) and NLO (green dashed line) at a cutoff $R = \beta_4/70$ are compared with the Temkin-Lamkin potential (orange dot-dashed line) [3,42,43] and a WKB approach [31] with *R* varied between $\beta_4/45$ and $\beta_4/70$ in the classically forbidden region (red band). ERE results are included in the inset. The purple dotted line represents the a_0 term. The black, densely dotted line includes the term linear in *k* as found in Refs. [32,33]. The LO prediction is included in the inset but overlapped by the TL and NLO lines.

for β_4/R above, say, 50, they would be barely visible on the scale of the figure. Therefore, our results are plotted at minimum R (or maximum $\beta_4/R \approx 70$). We find excellent agreement between ID-EFT and the Temkin-Lamkin phase shifts already at LO over a momentum range that extends well beyond $k\beta_4 \approx 1$, indicating that the breakdown scale of the theory is relatively high. In other words, the higher momentum range over which this agreement holds suggests that the curvature of the $1/r^4$ potential is a crucial piece of physics in the Temkin-Lamkin phase shifts. The agreement at lower momenta is due to two factors. First, the inclusion of the $1/r^4$ interaction allows us to capture physics at the $k \sim 1/\beta_4$ scale. Second, by fixing a_0 at LO, we demand agreement at threshold. For comparison, the inset of Fig. 1 shows also the first two terms in the ERE from the inverse scattering length a_0 and a correction linear in $k\beta_4$ [32,33]. While they approach the TL results for $k\beta_4 \ll 1$, ID-EFT captures the sign change of the phase shifts already at LO. At NLO, as expected, the agreement between ID-EFT and TL improves significantly at larger momenta.

Figure 2 offers a closer look into the errors in the LO and NLO predictions at minimum *R*. The most important features are the slopes of the LO and NLO lines—the rates at which the error grows—as $k\beta_4$ gets very large. To interpret this accurately, it is important to keep in mind that the LO and NLO predictions are both dependent on the cutoff, $\Lambda \sim 1/R$, and breakdown scale, Λ_b . Where *k* is greater than the typical momentum scales of the problem, but less than Λ_b , we expect the Λ dependence to dominate the errors, as our cutoff is not very large. This appears in Fig. 2 between $(k\beta_4)^2 \sim 10^3$ and 10^4 , where the slope of the NLO line is clearly greater than the slope of the LO line. This is of course by design, because



FIG. 2. Relative errors with respect to the Temkin-Lamkin *S*-wave phase shifts are shown for the LO (solid blue line) and NLO (dashed green line) predictions at maximum $\beta_4/R \approx 70$. The dot-dashed red line represents the WKB prediction [31]. The solid, grey, vertical line corresponds to the sixth-excited-state binding momentum. Both scales are logarithmic.

at NLO we have suppressed the Λ (or *R*) dependence. Finally, one can expect that these two lines cross above $(k\beta_4)^2 \sim 10^4$. This intersection indicates the breakdown scale of ID-EFT in this system,

$$\Lambda_b \sim 100/\beta_4 \approx \pi/a_B. \tag{16}$$

Thus, the phase-shift results indicate that ID-EFT holds in a region somewhat larger than naively expected.

Bearing this in mind also yields qualitative understanding of other features of Fig. 2. At low momenta, the NLO prediction is significantly closer to the Temkin-Lamkin result than the LO one is. However, the slopes of the LO and NLO deviations on the log-log plot of Fig. 2 are similar for $(k\beta_4)^2 \lesssim 10^3$ even though NLO is consistently almost an order of magnitude more accurate at these low values of k. (Note that $k\beta_4$ is still markedly larger than 1 there, so effective-range theory does not apply.) The similarity of the slopes is not coincidental. Forcing the NLO calculation to reproduce B_6 induces an error in the NLO calculation of order $(k\beta_4)^2 2\mu B_6/\Lambda_b^2$. For $k\beta_4 \lesssim 10^3$ this effect is larger than the $(k\beta_4)^4$ errors that dominate in the upper end of the EFT's validity range.

This analysis shows that ID-EFT is systematically improvable. The deviation from the underlying theory—the Temkin-Lamkin potential—is parametrically smaller at NLO than it is at LO. In contrast with the systematic improvement in ID-EFT, the WKB approximation [31] works as well as LO at high momenta but fails at low momenta. Details of the calculation of the WKB phase shifts are given in Appendix E and results are plotted in Fig. 1 as a band to indicate the variation with respect to *R* in the range of the classically forbidden region. It nearly overlaps with the LO curve, but differences are highlighted in the inset of Fig. 1: its assumptions are clearly not applicable at smaller values of *k*. Figure 2 reveals that the WKB approach describes the Temkin-Lamkin phase shift to better than 2% once $(k\beta_4)^2 \ge 10$ —and as long as ka_B



FIG. 3. Binding energies from both LO (solid blue lines) and NLO (dash-dotted green lines) are shown relative to the Temkin-Lamkin binding energies, $B_{TL}^{(n)}$. The results are plotted against the ratio of length scale associated with each state—defined in Eq. (17)—to the short-distance cutoff *R*. Vertically stacked pairs are shown successively for n = 3 (leftmost) through to n = 6 (rightmost). States with n < 3 are not shown as it is clear the EFT convergence breaks down for them.

remains small. The WKB curve crosses the Temkin-Lamkin curve at $(k\beta_4)^2 \approx 3 \times 10^2$, leading to the dip seen in Fig. 2, and for $(k\beta_4)^2 \gtrsim 10^3$ it nearly agrees with LO.

B. Bound states

Given the success of our description of phase shifts, we now turn to information from the bound-state spectrum, in order to obtain a parallel assessment of the ability of the EFT to capture the energy dependence below threshold for $|E| \ll \Lambda_b^2/2\mu$. As $a_0 \approx -2\beta_4$ is negative and not dramatically larger than β_4 , we do not expect that a very shallow bound state of size much greater than β_4 is present in this system, i.e., we anticipate that $B^{(6)}$ is not fine tuned.

The LO and NLO results for the binding energies $B^{(n)}$ of the four shallowest *S*-wave bound states are shown in Fig. 3 relative to the Temkin-Lamkin states. Each state is plotted against $\beta_{TL}^{(n)}/R$, where

$$\beta_{\rm TL}^{(n)} \equiv 1/\sqrt{2\mu B_{\rm TL}^{(n)}}$$
 (17)

is the characteristic size of the *n*th Temkin-Lamkin bound state with binding energy $B_{TL}^{(n)}$. Because the characteristic size decreases as we go down the spectrum, the lines do not cover the same horizontal span even though they are generated with the same *R* values. One can see the energies converge as *R* increases towards $70/\beta_4$, but much smaller values of *R* would be needed to see the deeper states "flatten out."

The shallowest, n = 6, state turns out to be two orders of magnitude deeper than the typical low-energy scale associated with β_4 in this system,

$$\epsilon_4 \equiv \frac{1}{2\mu\beta_4^2} \approx 1 \times 10^{-6} \,\text{a.u.} \tag{18}$$

Nevertheless, we obtain excellent agreement at LO with the Temkin-Lamkin result for this state, which is represented by the rightmost solid blue line in Fig. 3. This state is still of low energy compared to the energies of states in atomic hydrogen and corresponds to a length scale of approximately 4 a.u. The fact that this length scale is markedly smaller than a_0 demonstrates that our prediction is not a consequence of large-scattering-length universality. We predict $B^{(6)}$ so well using only β_4 and a_0 as inputs because we included the attractive $1/r^4$ potential at LO in our EFT and most of the state's wave function extends well outside hydrogen's electron cloud. At NLO, we renormalize $g_{\text{NLO}}(R)$ to the Temkin-Lamkin binding energy of this state, so agreement is by construction.

But introducing the NLO interaction reduces the disagreement between the other binding energies and the Temkin-Lamkin energies. The deeper states in Fig. 3 at smaller $\beta_{TL}^{(n)}/R$ values are an excellent visualization of how the theory scales with energy. $B^{(5)}$, the fifth excited state, second pair of lines from the right, is captured to within $\sim 10\%$ at LO and $\sim 1\%$ at NLO. Moving downward in the spectrum, where the pion's wave function has more overlap with the hydrogen atom, the NLO error for $B^{(4)}$ and $B^{(3)}$ grows systematically larger. This is a natural outcome in ID-EFT. We are fixing the scattering amplitude at threshold and continuing it to negative energies to find poles, including the analyticity properties implied by the $1/r^4$ potential in that continuation. In fact, by the time we reach $B^{(3)}$, NLO is not an improvement—both LO and NLO are off by 50%. This indicates that the series does not converge, which is why the ground and first two excited states are not shown in Fig. 3. As the continuation is made over a bigger energy range our prediction becomes less accurate.

In order to extrapolate our results to the $R \rightarrow 0$ limit, we assume that the effects associated with finite R can be accounted for via an expansion in R/β_4 . Therefore, we rely on the assumed convergence behavior of observables close to the renormalization point to extract asymptotic estimates. We expand

$$\mathcal{O}(R) = \mathcal{O}_{\infty} \left[1 + \sum_{m=1}^{\infty} c_m \left(\frac{R}{\beta_4} \right)^m \right], \tag{19}$$

with \mathcal{O}_{∞} being the asymptotic result for the LO or NLO energy of these bound states in ID-EFT and c_m the coefficients of the expansion. In the leading-order case the ID-EFT results for all seven binding energies are found by fitting the first coefficient of the expansion (19) and then reporting only \mathcal{O}_{∞} in Table I. At NLO the situation is more complicated: both the m = 1 and m = 4 term are needed to accurately fit the data. The rationale for this fit function is explained in Appendix D.

The NLO results show convergence to the TL results for the shallowest three states, but for the lowest three states NLO repulsion is so strong that they are no longer bound. (This further supports the use of the 14-state branch in our local regulator scheme.) This reordering of the states at NLO occurs already for the n = 3 state: its repulsive NLO correction is so large that it renders $B_{\text{NLO}}^{(3)}$ smaller than $B_{\text{NLO}}^{(4)}$. The matrix element in Eq. (13) is clearly no longer a perturbation for $n \leq 3$.

The error of these binding energies relative to the Temkin-Lamkin result is shown in Fig. 4. It ranges from 1% for the

TABLE I. Asymptotic $(R \rightarrow 0)$ results for LO and NLO binding energies compared to the Temkin-Lamkin spectrum. LO results are obtained from a fit to Eq. (19). Details about the NLO fit are given in Appendix D. Negative values indicate that the state is not bound.

n	$B_{\rm LO}$ (a.u.)	B _{NLO} (a.u.)	<i>B</i> _{TL} (a.u.)
6	1.19×10^{-04}	1.20×10^{-04}	1.20×10^{-04}
5	1.75×10^{-03}	1.56×10^{-03}	1.56×10^{-03}
4	8.59×10^{-03}	5.39×10^{-03}	6.12×10^{-03}
3	2.72×10^{-02}	4.91×10^{-03}	1.47×10^{-02}
2	6.84×10^{-02}	-3.02×10^{-02}	2.74×10^{-02}
1	1.49×10^{-01}	-1.84×10^{-01}	4.42×10^{-02}
0	2.98×10^{-01}	-6.95×10^{-01}	6.48×10^{-02}

sixth excited state, to 12% for the fifth excited state, to a factor of 4 for the ground state. It is notable that the LO error grows linearly with the energy of the bound state: the slope is \sim 1. The NLO interaction removes this error but leaves errors quadratic in energy, and indeed at NLO the slope is \sim 2.

Thus, we find evidence that the three shallowest states are within the regime of validity of ID-EFT, even though their binding energies vary by a factor of ~100. The breakdown binding energy ~ 10^{-2} a.u. inferred from where the LO and NLO trends intersect in Fig. 4 implies $\Lambda_b \sim 100/\beta_4$. This value is in good agreement with the determination from scattering, Eq. (16).

IV. CONCLUSIONS

We presented an effective field theory, ID-EFT, to describe the low-energy scattering of a heavy charged particle on a neutral atom, and the associated shallowest bound states. This EFT captures at leading order the physics of the long-range, but singular attractive, potential created by the atom's polarization. Renormalization requires at LO also a short-range interaction that is fixed by one datum. At next-to-leading



FIG. 4. The relative error between the Temkin-Lamkin binding energies B_{TL} and those obtained with ID-EFT. LO results are shown as blue circles. NLO results are shown in green—squares indicate bound states while x's indicate that the NLO correction pushes the state into the continuum.

order a second short-range interaction, determined by a second datum, systematically improves results for observables till momenta reach the breakdown scale.

We illustrated the workings of the EFT above and below threshold when the charged particle is a pion and the atom is hydrogen. We took as data results of the Temkin-Lamkin potential: the scattering length at LO and the shallowest binding energy at NLO. We compared the EFT outcomes with the exact results for other predictions using the same potential. Because, relative to the scale set by the long-range potential, the scattering length is not particularly large and the shallowest state is not particularly shallow, there is no fine tuning in this system and the EFT goes well beyond the effective range expansion.

We found a momentum breakdown scale somewhat larger (by a factor $\approx \pi$) than the inverse of the Bohr radius, $1/a_B$. For smaller momenta, phase shifts are well described at LO and the description improves systematically at NLO. The three shallowest bound states are also better reproduced at NLO than at LO. For larger momenta, the pion probes the inside of the atom; the atom can no longer be treated as a single unit. The four lowest-lying states of the pion-hydrogen ion have sizes somewhat smaller than a_B and are outside the regime of validity of the EFT.

Although here we used the Temkin-Lamkin potential as an example, ID-EFT offers a simple way to account for the long-range properties of this type of system without requiring detailed knowledge of the dynamics inside the atom. ID-EFT can be applied to other heavy charged particles and/or atoms and compared to data and/or other calculations where atomic structure is taken into account. For example, the protonhydrogen system has been studied with a three-body model and a uniquely shallow state predicted [39]. A comparison with our methods would be useful and the ability to quantify the uncertainties associated with the prediction of this shallow bound state could motivate experimental measurements. Regardless, we expect to find a similar convergence pattern, although details will depend on the values of leading-order parameters and the breakdown scale.

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APPENDIX A: ACCURATE CALCULATIONS OF THE SCATTERING LENGTH IN THE PRESENCE OF A $1/r^4$ POTENTIAL

The zero-energy solution to the reduced radial Schrödinger equation for a finite-range potential goes asymptotically like

$$u(r) \propto 1 - r/a_0. \tag{A1}$$

In general, the solution can be calculated numerically and the long-range portion of the wave function can be fit to a straight line such that the slope and intercept give an accurate and stable estimate of the scattering length. However, once a long-range potential is introduced, the tail of the interaction can make this extraction slow to converge.

In order to overcome this challenge, we derive the so-called "infrared corrections" perturbatively assuming that the $1/r^4$ potential is weak at large distances. The exact solution becomes a sum

$$u(r) = \sum_{i=0}^{\infty} u^{(i)}(r).$$
 (A2)

The first-order correction at zero energy is then

$$-\frac{d^2}{dr^2}u^{(1)}(r) = \frac{\beta_4^2}{r^4}u^{(0)}(r),$$
 (A3)

where $u^{(0)}(r)$ is taken to be Eq. (A1) up to an overall factor. After integration, we obtain

$$u^{(1)}(r) = \frac{\beta_4^2}{2a_0} \left(\frac{1}{r} - \frac{a_0}{3r^2}\right),\tag{A4}$$

and a better approximation for u(r) (again, up to an overall factor),

$$u(r) \approx 1 - \frac{r}{a_0} + \frac{\beta_4^2}{2a_0} \left(\frac{1}{r} - \frac{a_0}{3r^2}\right).$$
 (A5)

With this corrected form of the zero-energy solution, we are able to fit the coefficients of

$$u(r) = b_0 + b_1 r + b_{-1}/r + b_{-2}/r^2,$$
 (A6)

and reliably extract $a_0 = -b_0/b_1$ at much lower *r*. Additionally, we are able to compare the fit to the predicted coefficients of the 1/r and $1/r^2$ terms where

$$\frac{\beta_4^2}{2a_0} = \frac{b_{-1}}{b_0},\tag{A7}$$

$$\frac{b_{4}^2}{6} = \frac{b_{-2}}{b_0}.$$
 (A8)

We find excellent agreement between the fit results and the analytical predictions.

APPENDIX B: LOCAL BRANCHES

Several aspects of renormalization studies depend strongly on the ability to numerically approximate the limit $\beta_4/R \rightarrow \infty$. In local systems characterized by singular interactions, as *R* decreases and more of the singular well is exposed, the strength of the repulsive counterterm grows quickly. For these same local systems, there are an infinite number of "branches," each corresponding to a unique number of bound states, as noted in Ref. [8]. In order to achieve the practical $\beta_4/R \rightarrow \infty$ limit for the seven states mimicking the Temkin-Lamkin states, we chose to leverage this option and work on the n = 14 branch.

The advantages of this choice are highlighted in Fig. 5, where $g_{\text{LO}}^{(n=7)}$ begins to increase rapidly at $R \approx \beta_4/40$, but $g_{\text{LO}}^{(n=14)}$ does not reach the same value until $R \approx \beta_4/70$. Using this deeper branch, shallower states reach the asymptotic



FIG. 5. The running of the LO coupling for the n = 7 (solid purple) and n = 14 branches (dashed black).

regime—where Eq. (19) is valid—faster. On the n = 14 branch, the seven states of interest are now the shallowest states, and we are able to capture far more of the asymptotic behavior.

APPENDIX C: NLO IMPLEMENTATION COMPARISON

The NLO amplitude can be written in different forms, some of which we briefly compare here. In all cases, we define

$$t^{(\rm NLO)} \equiv t^{(\rm LO)} + \delta t, \qquad (\rm C1)$$

and the LO t matrix is given by Eq. (11).

In the first method, presented in Sec. II B,

$$\delta t = -\frac{2\mu}{k} \langle \phi^{(\text{LO},-)} | V_{\text{NLO}} | \phi^{(\text{LO})} \rangle.$$
 (C2)

From the relation between scattering wave function and the scattering amplitude, we can write instead

$$\delta t = (1 + t^{(\text{LO})}G_0)V_{\text{NLO}}(G_0 t^{(\text{LO})} + 1).$$
(C3)

This form is used, for example, in Ref. [50]. By substituting Eq. (11) in Eq. (C3),

$$\delta t = V_{\rm NLO} + V_{\rm NLO} G_0 t^{\rm (LO)} + V_{\rm LO} G_0 (1 + t^{\rm (LO)} G_0) V_{\rm NLO} (G_0 t^{\rm (LO)} + 1) = V_{\rm NLO} + V_{\rm NLO} G_0 t^{\rm (LO)} + V_{\rm LO} G_0 \, \delta t, \qquad (C4)$$

which is the form used in Refs. [51,52]. We have checked numerically that these three equivalent forms for δt indeed give the same result.

Since the calculation of δt respects unitarity only perturbatively, there are then different ways to express the relation between $t^{(\text{NLO})}$ and $\delta^{(\text{NLO})}$. Starting with the standard (unitary) relationship between the scattering amplitude and the phase shift,

$$t^{(\text{NLO})} = \frac{1}{2i} (e^{2i\delta^{(\text{NLO})}} - 1),$$
 (C5)



FIG. 6. (Upper panel) The real parts of the NLO phase shifts computed using Eq. (C9) (solid green line) and Eq. (C5) (dashed orange line). The TL results are shown as a dash-dotted red line. (Lower panel) Imaginary components of the NLO phase shifts (colors and styles are the same as in the upper panel). The (strictly real) difference between the TL results and the LO phase shifts is indicated with a dash-dotted red line.

we write $\delta^{(\text{NLO})} = \delta^{(\text{LO})} + \epsilon$ and expand in small ϵ , ignoring $\mathcal{O}(\epsilon^2)$ terms and higher. This yields

$$t^{(\text{NLO})} \approx \frac{1}{2i} \left[e^{2i\delta^{(\text{LO})}} (1+2i\epsilon) - 1 \right].$$
 (C6)

Simplifying, we get

$$t^{(\text{NLO})} = t^{(\text{LO})} + \epsilon e^{2i\delta^{(\text{LO})}},$$
 (C7)

where

$$t^{(\text{LO})} = \frac{1}{2i} \left(e^{2i\delta^{(\text{LO})}} - 1 \right)$$
(C8)

and the NLO correction to the scattering amplitude is related to the NLO piece of the phase shift by

$$\epsilon = e^{-2i\delta^{(LO)}}\delta t. \tag{C9}$$

This relation is strictly perturbative, in the sense that all quantities are computed to NLO accuracy and not further.

If instead one computes $\delta^{(\text{NLO})}$ from δt nonperturbatively, via Eq. (C5), thereby assuming that unitarity remains strictly valid at NLO, two key differences emerge. First, the real part of the phase shift is less accurate, as seen in the upper panel of Fig. 6. Second, an imaginary component accumulates

at higher momenta—shown explicitly in the lower panel of Fig. 6. Consistently computing the scattering amplitude and phase shifts perturbatively produces not only formally correct results but more accurate predictions.

Note that the imaginary component of the phase shift accumulated in such a calculation, although unphysical, does provide insight into the expected size of the next-to-next-toleading-order (NNLO) correction. The first term omitted in the expansion of $\exp(2i\epsilon)$ in Eq. (C6) is real and $\mathcal{O}(\epsilon^2)$. If we consider also the prefactors in Eq. (C6), we conclude that this is the piece of Eq. (C5) that generates the leading piece of the imaginary part of the nonperturbative phase shift, and hence $\Im(\delta^{(\text{NLO})})$ should also be $\mathcal{O}(\epsilon^2)$. In the lower panel of Fig. 6, we see in the context of the $\mathcal{O}(\epsilon)$ difference between the TL phase shifts and our LO phase shifts that this conclusion is well supported.

Retaining $\Re(\delta^{(\text{NLO})})$ from Eq. (C5), and simply ignoring $\Im(\delta^{(\text{NLO})})$ accounts for some effects beyond NLO. But this is an incomplete NNLO calculation. The line this produces in Fig. 2 has approximately the same slope as the NLO line, which is based on Eq. (C9). However, as $\Re(\delta^{(\text{NLO})})$ calculated nonperturbatively gives worse results (compared to the TL phase shifts) than the consistent perturbative calculation, the line this produces in Fig. 2 is displaced up, towards the LO curve. It then intersects the LO curve at a smaller momentum, leading to an underestimate of the breakdown scale. This is just a new example of a well-known phenomenon: including a partial subset of small corrections does not necessarily improve the result. For another example of the same type, see Fig. 5 of Ref. [53], and for a more dramatic example, with far-reaching consequences, Ref. [54].

APPENDIX D: R DEPENDENCE OF THE NLO BINDING ENERGIES

Extracting asymptotic binding energies at LO is relatively straightforward, but it turns out to be markedly more complicated at NLO. First, we observe that the NLO binding energy formula, Eq. (13), contains two different *R*-dependent terms. The first is $B_{\text{LO}}(R)$. The second includes a matrix-element ratio when the NLO renormalization condition

$$g_{\rm NLO}(R) = \left(\frac{B_{\rm TL}^{(6)}}{B_{\rm LO}^{(6)}} - 1\right) \frac{1}{\left\langle \psi_{\rm LO}^{(6)} | \chi | \psi_{\rm LO}^{(6)} \right\rangle} \tag{D1}$$

is inserted. $B_{LO}^{(n)}(R)$ is well described with a straight line, i.e., without including any of the m > 1 terms in Eq. (19). The matrix-element ratio is more complicated: we expect it to be even in *R* because of analyticity in the regulator parameter. But whether the nonlinear piece of the NLO binding energy is proportional to R^2 (simplest dependence) or R^4 (which is the leading *R* dependence of the regulator) is not immediately apparent. Consequently, we have tried several approaches to fit the NLO binding energies and record them here for posterity.

We require two features of our NLO binding energy analysis. First, we expect that the NLO renormalization procedure ought to reduce the linear *R* dependence for states where the binding momentum is not far from the additional renormalization point, $\gamma^{(6)}$: for these states the coefficient of the linear-in-*R* term will be smaller than those found when fitting



FIG. 7. Results of a fit to Eq. (19) where only m = 1 and m = 4 terms are included. (Top panel) Binding energy as a function of the cutoff distance *R*. Both quantities are in atomic units. Data are indicated as blue circles. The fit is represented with a solid green line. The asymptotic value of the binding energy is indicated on the *y* axis as a green square. Shaded (purple online) regions highlight the range of *R* over which the fit was conducted. (Middle panel) Data and fit results for the linear term of our fit are shown with the asymptotic binding energy and the best-fit quartic *R* dependence subtracted. (Bottom panel) The residuals of our $R + R^4$ fit. Note that the *y* scale in the bottom panel is expanded roughly 100 times compared to the top panel. Note also that the difference in *x*-axis range gets progressively smaller as one moves from the top to middle to bottom panel.

the LO binding energies. Second, we expect that the EFT fails *systematically*. The NLO corrections ought to increase with a positive power of the binding momentum.

Figure 7 displays the steps we took to understand the R dependence of the binding energy of the fourth excited state

in our NLO calculation. The upper panel of Fig. 7 shows the *R* dependence of this quantity is not exclusively linear. A successful fit to Eq. (19) is shown together with the data. While the ability of the fit to describe the data is clear (and highlighted in the bottom panel), it seems odd at first glance that data that is decreasing as *R* decreases leads to an asymptotic value that is larger than any value in the data set. The minimum, and subsequent change in derivative, of $B_{\rm NLO}^{(4)}$ with respect to *R* is due to the linear term in the fit assuming a dominant role, a role that it does not have for the *R* values at which the fit is performed.

The middle panel of Fig. 7 shows that we can confidently extract the linear coefficient in the region where we have data on the bound state energy. In this panel we have subtracted the asymptotic value and quartic term from the data, leaving only the linear dependence. A linear fit is clearly an excellent description of the residuals: the linear dependence that dominates at small R is indeed present in the data even where the quartic term is more significant.

The bottom panel of Fig. 7 then shows the residuals once the linear-in-*R* term in our fit is subtracted. The residuals show no systematic trend with *R* and are of order 10^{-7} in atomic units. Any remaining *R* dependence will therefore not affect the extrapolation at the level of accuracy we are quoting in this paper.

To be thorough, we also fitted our data using $R + R^2$ and $R + R^2 + R^4$ forms. While these approaches did align with the data in the region 0.467 < R < 0.54, they also produced coefficients of the linear-in-R term that were larger than the LO value of the same coefficient for states with $n \ge 4$, i.e.,

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states where including NLO corrections and renormalizing to B_6 should have decreased the linear dependence on R. These fits also yielded asymptotic values of the binding energies that had no systematic trend with n, and showed clear signs of overfitting. We defer the question of why there is no R^2 term present in the function $B_{\rm NLO}(R)$ to future work, only commenting here that we do not believe this behavior will prevail for all regulators.

APPENDIX E: DETAILS ABOUT THE WKB CALCULATION

Reference [31] derives the phase shift at momentum k to be

$$\delta_0(k) = \Phi + \frac{\pi}{4} - I_n \left[\beta_n^{2/(n-2)}k\right]^{2/(n-2)}$$
(E1)

for singular potentials of the form $1/r^n$ where n > 2. In this expression I_n is a straightforward integral while the calculation of

$$\Phi = \int_{r_0}^{\infty} dr \sqrt{-2\mu V(r)}$$
(E2)

is a little more delicate. Here, r_0 is the classical turning point where V(r) = 0. It is the point where the potential crosses zero from below as the short-distance repulsion takes over at shorter distances. This point was found with the FORTRAN library MINIPACK for each value of *R* tested in this manuscript. In the relevant range of *R*, $r_0 < R$. The integral was then computed numerically with another FORTRAN library, QUADPACK. The variation of Φ is $\mathcal{O}(10^{-2})$ in radians.

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