Scattering of heavy charged particles on hydrogen atoms

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Abstract. The low energy scattering of heavy positively charged particles on hydrogen atoms (H) are investigated by solving the Faddeev equations in configuration space. A resonant value of the pH scattering length, $a = 750 \pm 5$ a.u., in the pp antisymmetric state was found. This large value indicates the existence of a first excited state with a binding energy $B = 1.14 \times 10^{-9}$ a.u. below the H ground state. Several resonances for non zero angular momenta states are predicted.

1 Introduction

The scattering of heavy charged particles ($X^+$) on atoms at kinetic energies smaller than the inelastic thresholds is dramatically influenced by the presence of atomic electrons. Their virtual excitations in presence of the incoming charged particle result into long range attractive forces which dominate the low energy scattering. These states were proposed as a possible source of metastability in the $\bar{p}$He system [1] and were found to play a determinant role in the low energy $\bar{p}$H annihilation [2]. A full solution of these problems is however made extremely difficult by the presence of annihilation channels and only approximate solutions were achieved. We present in what follows a rigorous solutions for the simplest problem of scattering on hydrogen atoms.

A simple two-body approximation is first discussed in Section 2. Section 3 is devoted to the solution of the 3-body ($X^+e^-p^+$) problem, obtained by solving the Faddeev equations in configuration space. We treat with special care the pH case, for it exhibits the more interesting properties and constitutes moreover a realistic experimental challenge. Some final remarks about the present calculations and plans for future work are given in the conclusion. We use all along the paper electronic atomic units ($m_e = e^2 = \hbar = 1$).

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2 Two-body approach

To get a first qualitative insight into the underlying physics, it is interesting to consider a simple 2-body $X^+H$ problem. The H atom is supposed to be point-like and to interact with the incoming particle via a central potential

$$V(r) = \frac{1}{2} \frac{\alpha(r)}{r^4}$$

(1)

$\alpha(r)$ tends to the H dipole polarizability ($\alpha_d = \frac{9}{2}$) for large values of $r$ and regularizes the $\frac{1}{r^4}$ singularity at $r=0$. Its precise form is given in [3].

We have displayed in Fig. 1 the binding energies of the $\pi^+H$ bound states with interaction (1). One can see a large number of states with angular momentum values up to $L=7$. Some of them are very close to the dissociation threshold. The elastic $\mu^+H$ cross section in the momentum range $k \in [0.48, 0.64]$ has been plotted in Fig. 2 with the most relevant partial wave contributions shown separately. The $L=5,6,7$ states display a clear resonant behaviour also visible in the total cross section. A direct calculation of their position and width – using the complex rotation method [4] – provides the values $E_5=(7.9-1.4i)10^{-4}$, $E_6=(1.1-0.25i)10^{-3}$, $E_7=1.0 \times 10^{-3}-1.3i \times 10^{-7}$ respectively [5]. These resonances concern all high angular momentum states, involving large centrifugal barriers and relatively large energies. It’s worth noticing however that they can be found even for $L=1$ and at extremely small energy values. In the $\pi^+H$ case one has e.g. $E_1=(4.9-1.4i) \times 10^{-7}$. The examples shown in Figs. 1 and 2 illustrate well the kind of physics governed by the polarization forces. An extensive work as a function of the projectile mass has been done [6] showing a very rich spectrum of bound and resonant states, with a complexity increasing as a function of the projectile mass.

To what extend the results of this simple approach are reliable? Answering this question was the main motivation of this work. A definite answer will only appear by letting the electron dynamics in H play its full role, that is by
considering the \((X^+p^+e^-)\) three body problem. And this is the aim of the next section.

### 3 Three-body calculations

The 3-body \((X^+p^+e^-)\) calculations are performed using Faddeev equations in configuration space. Three different sets of Jacobi coordinates are involved, defined by

\[
x_\alpha = -\left[\frac{2m_\beta m_\gamma}{m_\beta + m_\gamma}\right]^{1/2} (r_\beta - r_\gamma),
\]

\[
y_\alpha = -\left[\frac{2m_\alpha (m_\beta + m_\gamma)}{m_\alpha + m_\beta + m_\gamma}\right]^{1/2} \left(\frac{r_\alpha - r_\beta m_\beta + m_\gamma r_\gamma}{m_\beta + m_\gamma}\right)
\]

where \((\alpha\beta\gamma)\) denote cyclic permutations of \((123)\), and \(m_\alpha\) the particle masses.

We identify \(1 \equiv X^+\), \(2 \equiv p^+\), \(3 \equiv e^-\). The standard Faddeev equations read

\[
(E - H_0 - V_\alpha)\Psi_\alpha = V_\alpha \sum_{\alpha \neq \beta} \Psi_\beta,
\]

where \(H_0\) is the 3-particle free Hamiltonian and \(V_\alpha\) the 2-body Coulomb potential for the interacting \((\beta\gamma)\) pair

\[
V_\alpha(x_\alpha) = \frac{e_\beta e_\gamma}{|r_\beta - r_\gamma|} e_j = \pm 1
\]

For projectile masses \(m_X < m_p\) we restrict ourselves to scattering energies below the first rearrangement threshold \(X^+ \rightarrow p^+ + (X^+ + e^-)\). In that case, amplitudes \(\Psi_2\) and \(\Psi_3\) have no asymptotics.

Equations \((4)\) provide satisfactory solutions for bound states but are not suitable for scattering Coulomb problems. The reason is that their right hand side does not decrease fast enough to ensure the decoupling of Faddeev amplitudes in the asymptotic region and to allow unambiguous implementation of boundary conditions. In order to circumvent this problem, Merkuriev \([7]\) proposed to split the Coulomb potential \(V\) into two parts by means of some arbitrary cut-off function \(\chi\)

\[
V(x) = V^s(x,y) + V^l(x,y)
\]
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\[ V_s(x, y) = V(x) \chi(x, y) \]
\[ V_l(x, y) = V(x)[1 - \chi(x, y)] \]

and to keep in the right hand side of equation (4) only the short range \( V_s \) contribution. One is then left with a system of equivalent equations

\[ (E - H_0 - W_\alpha - V_s^e)\Psi_\alpha = V_s^e \sum_{\alpha \neq \beta} \Psi_\beta, \quad (6) \]

in which \( W_\alpha \) are some 3-body potential containing the long range parts:

\[ W_\alpha = V_l^\alpha + V_l^\beta + V_l^\gamma \quad (7) \]

This approach was found to be very efficient in calculating the e+Ps and e+H cross sections [8, 9].

Figure 4. 3-body zero energy X+H cross section as a function of the projectile mass \( m_X \)

Equations (8) were solved by expanding \( \Psi_i \) in the bipolar harmonics basis

\[ \Psi_i(x_i, y_i) = \sum_{\alpha_i} \varphi_{\alpha_i}(x_i, y_i) x_i y_i B^{LM}_{\alpha_i}(\hat{x}_i, \hat{y}_i) \quad \alpha_i \equiv \{l_i, i_i\} \quad (8) \]

and their components \( \varphi_{\alpha_i} \) in the basis of two-dimensional splines. In practical calculations we took the cutoff function:

\[ \chi(x, y) = 2 \left\{ 1 + \exp \left[ \left( \frac{x}{x_0} \right)^\nu \right] \right\}^{-1} \]
Final results are independent of the parameters \( x_0, y_0, \nu \) but an appropriate choice for their values makes the convergence of expansion \( 8 \) faster. The values \( x_0 = 2.0, y_0 \approx 2\sqrt{m_X}, \nu = 2.3 \) are suitable.

Binding energies for the lower 3-body \( \pi^+\text{H} \) and \( \mu^+\text{H} \) bound states are plotted in Fig. 6. They are compared to the results of the 2-body approach \( 1 \). One can see that, although there is a qualitative agreement, 2-body energies are systematically underestimated. These results can be used to improve the short range part of the 2-body potential.

Some interesting features of the 3-body Coulomb system can be learned from Fig. (4), where the zero energy \( X^+\text{H} \) cross section as a function of the projectile mass \( m_X \) is displayed. Each peak corresponds to the appearance of a new S-wave bound state. The critical mass values \( m_i \) at which they occur, would enable to generalize the ground state stability triangle \( 10 \) to higher excitations. A zoom in the region of physical interest, \( \mu^+ \) and \( \pi^+ \), is shown in Fig. 5. The calculated values are respectively \( a_{\mu \text{H}} = 69.1 \) and \( a_{\pi \text{H}} = 24.4 \). Some care has to be taken in extracting the scattering observables specially at zero energy, from the asymptotic solution at finite distance. The long range polarization force makes the convergence of the observables as a function of the \( X^+\text{H} \) distance very slow and requires an appropriate extrapolation procedure \( 8 \).

\[ \text{Figure 5. Scattering length as a function of } m_X, \text{ in the region of } \mu \text{ and } \pi. \]

\[ \text{Figure 6. Energies for lower } \pi \text{H and } \mu \text{H bound states compared to 2-body results } 1. \]

\( \mu \text{H} \) scattering deserves special comments. The 3-body wave function has to be antisymmetric with respect to the \( p \) exchange. This can be realized in two different ways following the proton spin coupling. For the case when the two protons spins are antiparallel (singlet) the spatial part of the wave function is symmetric, while for the parallel case (triplet) it is antisymmetric. In the 2-body approach, these two cases give rise to completely different potentials.

The singlet case has a broad attractive well which supports a great number of bound states. They have been calculated since the first days of Quantum Mechanics and they are presently known with a very high precision (see e.g. \( 11, 12 \) and reference therein). Our 3-body calculations cannot reach this kind of accuracy for bound states but are in good agreement for the lower excitations.
They provide furthermore the first result for the pH scattering length \( a_s = -29.3 \).
We notice that the zero energy scattering wave function shows 20 nodes in \( y_1 \)-direction, indicating the existence of 20 \( L=0 \) \( \sigma_g \) energy levels for \( H_2^+ \).

The triplet case – modeled by Landau [14] – is dominated by the Pauli repulsion between the two protons, overbalanced at \( r \sim 10 \) by the attractive polarization forces.

![Figure 7. Zero energy pH in the pp triplet state, compared to the results of Landau potential](image)

Our 3-body calculations give a scattering length of \( a_t = 750 \pm 5 \). The nodal structure of the Faddeev amplitudes indicates that such a big value is due to the existence of a first excited \( L=0 \) state with extremely small binding energy. By using the modified effective range theory [15] we are able to determine its binding energy, which turns to be \( B=(1.135\pm0.035)\times10^{-9} \) below the H ground state. To our knowledge, this is the weakest bond ever predicted, three times smaller than the \(^4\text{He} \) atomic dimer [13].

The existence of \( L=0 \) and \( L=1 \) ground states is well known and their binding energies have been very precisely calculated [11, 12]. These authors were however not able to conclude about the existence of a second S-wave bound state.

It is interesting to compare the three-body calculations with those provided by the simple Landau two-body potential. S-wave cross sections are plotted in Fig. [7]. At zero energy, both calculations differ by two orders of magnitude while at energies \( E \sim 10^{-6} \) they are already in quite a good agreement, despite the simplicity of the 2-body approach. This is due to the fact that the effective
interaction is highly repulsive at short distances, what prevents the incoming proton to penetrate inside the H atom and minimizes the effect of the 3-body dynamics.

![Figure 8. pH elastic cross section for L=3 in the pp spin triplet state](image1)

![Figure 9. pH elastic cross section for L=4 in the pp spin triplet state](image2)

The pH cross sections for higher partial waves have also been calculated. They exhibit some narrow resonances in several partial waves. Figs. 8 and 9 show the elastic cross section for the L=3,4 states. They are compared to the results of Landau potential. We can see that their agreement in this energy region is rather good. The 3-body resonances are a little bit shifted to the lower energy region and have smaller widths. The same effect is seen in the bound state calculations where the 2-body results are always slightly underbound. The position and width of these resonances were estimated to $E=(5.13-1.61i) \times 10^{-6}$ for L=3 and $E=(1.56-0.94i) \times 10^{-5}$ for L=4.

4 Conclusions

By solving the Faddeev equations in configuration space, we have calculated the bound and scattering states for the $(X^+ p^+ e^-)$ system with projectile masses $m_X$ larger than the electron. The long range polarization forces give rise to a rich spectrum of bound and resonant states with increasing complexity as $m_X$ increases.

Predictions for the physical cases $X^+ = \mu^+, \pi^+, K^+$ are obtained. We found in particular the scattering length values $a_{\pi^+ H}=24.4$ and $a_{\mu^+ H}=69.1$ a.u.

Of special interest is the pH system in the pp spin-triplet state. We predict a second S-wave bound state with binding energy $B=1.14 \times 10^{-9}$ a.u. below the H ground state. This constitutes the weakest bond ever predicted, even smaller than the $^4$He atomic dimer.

The existence of such a nearthreshold state dominates the low energy pH scattering and results into a scattering length of $a_t=750 \pm 5$ a.u. A low energy proton approaching an H atom will behave like colliding with a large nanoscopic object. Several resonances occurring in different partial waves, but visible in the total cross section, are also predicted. The experimental confirmation of these
results would be very interesting.

The calculations presented here are performed using a fully non relativistic dynamics with Coulomb pair-wise interactions. In this framework, they are parameter free with the only input of particles masses and charges. The perturbations induced by strong X\(^+\)-p interactions or higher electromagnetic corrections have not been included.

In view of the extreme sensibility of the pH results, it is necessary to quantify both relativistic and strong-interaction effects. A direct measurement of the pH cross section at very low energy seems unlikely. One can however access the low energy pH continuum in the final state of the H\(^+_2\) photodissociation cross section. Work is in progress in these two directions.

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