Series of broad resonances in atomic three-body systems

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Abstract

We re-examine the series of resonances found earlier in atomic three-body systems by solving the Faddeev-Merkuriev integral equations. These resonances are rather broad and line-up at each threshold with gradually increasing gaps, the same way for all thresholds and irrespective of the spatial symmetry. We relate these resonances to the Gailitis mechanism, which is a consequence of the polarization potential.
I. INTRODUCTION

A couple of years ago we observed an accumulation of resonances above the thresholds [1, 2]. Other independent calculations have not confirmed these findings, only a few narrow resonances have been independently calculated. This is understandable, since calculation of broad resonances, especially in a multi-body system, is very complicated. The wave function of narrow resonances behaves very much like a bound-state wave function, so they can be calculated by some slight modification of bound state techniques. The most common method is the complex rotation of coordinates. This technique renders the resonance state wave function to a square integrable one, and thus standard techniques like variational expansion of the wave function become applicable. These methods, however, run into technical difficulties for broad resonances. Here, to uncover the resonances, they need a large rotation angle and the rotated continuum becomes more and more scattered making the identification of resonances increasingly difficult.

Our method is different. We start with the Faddeev integral equations with the modification proposed by Merkuriev [3] and solve them by using the Coulomb-Sturmian potential separable expansion method [4]. Since the investigations of Refs. [1, 2] we improved the technique in Ref. [5] making it more amenable to calculate broad resonances. Therefore in Sec. II we outline our technique of solving the Faddeev-Merkuriev integral equations for resonant states while in Sec. III we show our results for $e - Ps$ resonances. Finally we summarize our findings and provide an explanation for the formation of the series of broad resonances.

II. CALCULATION OF COULOMB THREE-BODY RESONANCES

A. Faddeev-Merkuriev integral equations

The Hamiltonian of an atomic three-body system is given by

$$H = H^0 + v_1^C + v_2^C + v_3^C,$$

(1)

where $H^0$ is the three-body kinetic energy operator and $v_\alpha^C$ denotes the Coulomb interaction of each subsystem $\alpha = 1, 2, 3$. We use the usual configuration-space Jacobi coordinates $x_\alpha$ and $y_\alpha$, where $x_\alpha$ is the distance between the pair $(\beta, \gamma)$ and $y_\alpha$ is the distance between the center of mass of the pair $(\beta, \gamma)$ and the particle $\alpha$. The potential $v_\alpha^C$, the interaction of the pair $(\beta, \gamma)$, appears as $v_\alpha^C(x_\alpha)$. In an atomic three-body system two particles always have the same sign of charge.
Without loss of generality, we can assume that they are particles 1 and 2, and thus $v_C^{(2)}$ is a repulsive Coulomb potential.

The wave function of a three-particle system is very complicated. It exhibits different asymptotic behaviors reflecting the possible asymptotic fragmentations. In the Faddeev approach we split the wave function into components such that each component describes only one kind of asymptotic fragmentation [3]. The components satisfy a set of coupled equations, the Faddeev equations.

The Hamiltonian (1) is defined in the three-body Hilbert space. Therefore, the two-body potential operators are formally embedded in the three-body Hilbert space,

$$v^{(2)}_\alpha = v^{(2)}_\alpha(x_\alpha)\mathbf{1}_{y_\alpha},$$

(2)

where $\mathbf{1}_{y_\alpha}$ is a unit operator in the two-body Hilbert space associated with the $y_\alpha$ coordinate.

The Coulomb potential is a long range potential as it modifies the motion even at asymptotic distances. On the other hand, it also possesses some features of a short-range potential as it correlates the particles strongly and supports two-body bound states. These two properties are contradictory and require different treatments. In Merkuriev’s approach the three-body configuration space is divided into different asymptotic regions [6]. The two-body asymptotic region $\Omega_\alpha$ is defined as a part of the three-body configuration space where the conditions

$$(|x_\alpha|/x_0)^\nu < |y_\alpha|/y_0,$$

(3)

with parameters $x_0 > 0$, $y_0 > 0$ and $\nu > 2$ are satisfied. It has been shown that in $\Omega_\alpha$ the short-range character of the Coulomb potential prevails, while in the complementary region the long-range character of the Coulomb potential becomes dominant.

Therefore we split the Coulomb potential in the three-body configuration space into short-range and long-range parts

$$v^{(C)}_\alpha = v^{(s)}_\alpha + v^{(l)}_\alpha.$$  

(4)

The splitting is carried out with the help of a splitting function $\zeta_\alpha$,

$$v^{(s)}_\alpha(x_\alpha, y_\alpha) = v^{(C)}_\alpha(x_\alpha)\zeta_\alpha(x_\alpha, y_\alpha),$$

(5)

$$v^{(l)}_\alpha(x_\alpha, y_\alpha) = v^{(C)}_\alpha(x_\alpha)\left[1 - \zeta_\alpha(x_\alpha, y_\alpha)\right].$$

(6)

The function $\zeta_\alpha$ vanishes asymptotically within the three-body sector, where $x_\alpha \sim y_\alpha \to \infty$, and approaches 1 in the two-body asymptotic region $\Omega_\alpha$, where $x_\alpha << y_\alpha \to \infty$. As a result, in the
three-body sector, the short-range potential $v_{\alpha}^{(s)}$ vanishes and long-range potential $v_{\alpha}^{(l)}$ approaches $v_{\alpha}^{C}$. In practice, the functional form

$$\zeta_{\alpha}(x_{\alpha}, y_{\alpha}) = 2/ \left\{ 1 + \exp \left[ \left( x_{\alpha} / x_0 \right)^{\nu} / \left( 1 + y_{\alpha} / y_0 \right) \right] \right\}$$  \hspace{1cm} (7)

is used. Typical shapes for $v^{(s)}$ and $v^{(l)}$ are shown in Figures [1] and [2]. In fact, these parameters were adopted in our $e - Ps$ calculations. We can see that $v^{(l)}$ is a valley which opens up as $y_{\alpha}$ goes to infinity and becoming shallower and shallower. Finally, in the $y_{\alpha} \to \infty$ limit there is no two-body bound state in $x_{\alpha}$. 

FIG. 1. Short range ($v^{(s)}$) and long range ($v^{(l)}$) parts of an attractive Coulomb potential. The parameters are $Z = -1$, $x_0 = 4$, $y_0 = 12$ and $\nu = 2.1$.

FIG. 2. Short range ($v^{(s)}$) and long range ($v^{(l)}$) parts of an attractive Coulomb potential. The parameters are $Z = -1$, $x_0 = 15$, $y_0 = 35$ and $\nu = 2.1$. 

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The Coulomb potential $v^C_3$ is repulsive. So, it does not support bound states and there are no two-body channels associated with this fragmentation. Consequently, the entire $v^C_3$ can be considered as a long-range potential. Then the long-range Hamiltonian is defined as

$$H^{(l)} = H^0 + v_1^{(l)} + v_2^{(l)} + v^C_3,$$

and the three-body Hamiltonian looks like an ordinary three-body Hamiltonian with only two short range interactions

$$H = H^{(l)} + v_1^{(s)} + v_2^{(s)}.$$  (9)

So, we split the wave function into two components only

$$|\Psi\rangle = |\psi_1\rangle + |\psi_2\rangle.$$  (10)

Then for components, we have the set of Faddeev equations,

$$(E - H^{(l)}_1)|\psi_1\rangle = v_1^{(s)}|\psi_2\rangle$$

$$(E - H^{(l)}_2)|\psi_2\rangle = v_2^{(s)}|\psi_1\rangle,$$

where

$$H^{(l)}_\alpha = H^{(l)} + v_\alpha^{(s)}.$$  (12)

By adding these two equations we recover the original Schrödinger equation. So, the Faddeev procedure is a clever way of solving the quantum mechanical Schrödinger equation. We can write these differential equations into an integral equation form

$$|\psi_1\rangle = G^{(l)}_1(E) v_1^{(s)} |\psi_2\rangle$$

$$|\psi_2\rangle = G^{(l)}_2(E) v_2^{(s)} |\psi_1\rangle,$$

where $G^{(l)}_\alpha(E) = (E - H^{(l)}_\alpha)^{-1}$. With Merkuriev’s procedure Faddeev’s aim is achieved for the Coulomb potential as well. Now each component describes only one kind of asymptotic fragmentation.

If particles 1 and 2 are identical particles, the Faddeev components $|\psi_1\rangle$ and $|\psi_2\rangle$, in their own natural Jacobi coordinates, must have the same functional forms

$$\langle x_1 y_1 |\psi_1\rangle = \langle x_2 y_2 |\psi_2\rangle.$$  (14)

On the other hand, by interchanging particles 1 and 2, we have

$$\mathcal{P}_{12} |\psi_1\rangle = p |\psi_2\rangle,$$

(15)
where \( p = \pm 1 \), depending the total spin of the two identical particles. So, \( \psi_1 \) and \( \psi_2 \) are not independent, and to determine one of them we need only one equation

\[
|\psi_1\rangle = G_{1}^{(l)} v_{1}^{(s)} p P_{12} |\psi_1\rangle.
\]

As we can see, we can easily incorporate the identity of particles into the Faddeev formalism, and this even leads to a considerable simplification of the equations.

**B. Solution method**

In order that we can solve the Faddeev-Merkuriev integral equations we represent them in Coulomb–Sturmian (CS) basis. The CS functions are given by

\[
\langle r|nl;b\rangle = \sqrt{\frac{n!}{(n + 2l + 1)!}} \exp(-br)(2br)^{l+1}L_n^{(2l+1)}(2br),
\]

where \( L \) denotes the Laguerre polynomials, \( l \) is angular momentum, \( n \) is the radial quantum number and \( b \) is a parameter. With \( \langle r|nl;b\rangle \equiv \langle r|nl;b\rangle/r \), the orthogonality and completeness relations take the forms

\[
\langle nl';nl;b\rangle = \delta_{nn'}
\]

and

\[
1 = \lim_{N \to \infty} \sum_{n=0}^{N} |nl;b\rangle \langle nl;b| = \lim_{N \to \infty} \sum_{n=0}^{N} |nl;b\rangle \langle nl;b|.
\]

The three-body Hilbert space is a direct product of two-body Hilbert spaces, so, as a basis, we may take the angular-momentum-coupled direct product of the two-body bases,

\[
|n\nu\lambda;b\rangle_\alpha = |nl;b\rangle_\alpha \otimes |\nu\lambda;b\rangle_\alpha, \quad (n,\nu = 0, 1, 2, \ldots),
\]

where \( |nl;b\rangle_\alpha \) and \( |\nu\lambda;b\rangle_\alpha \) are associated with the coordinates \( x_\alpha \) and \( y_\alpha \), respectively. With this basis, the completeness relation takes the form (with angular momentum summation implicitly included)

\[
1 = \lim_{N \to \infty} \sum_{n,\nu=0}^{N} |n\nu\lambda;b\rangle_\alpha \langle n\nu\lambda;b| = \lim_{N \to \infty} 1^{N}_\alpha.
\]

We insert a unit operator into the Faddeev equations

\[
|\psi_1\rangle = \lim_{N \to \infty} G_{1}^{(l)}(E) 11^{N} v_{1}^{(s)} 12^{N} \langle \psi_2|,
\]

\[
|\psi_2\rangle = \lim_{N \to \infty} G_{2}^{(l)}(E) 12^{N} v_{2}^{(s)} 11^{N} \langle \psi_1|,
\]

\[
(22)
\]
and keep $N$ finite. This amounts to approximating $v^{(s)}_{\alpha}$ in the three-body Hilbert space by a separable form

$$v^{(s)}_{\alpha} = \lim_{N \to \infty} 1_N^{N} v^{(s)}_{\alpha} 1_N^{N} \approx 1_N^{N} v^{(s)}_{\alpha} 1_N^{N} \approx \sum_{n,\nu,n',\nu'} |\tilde{n}_\nu l \lambda, b \rangle_{\alpha} v^{(s)}_{\alpha \beta} \langle n' \nu' l' \lambda'; b |,$$

where $v^{(s)}_{\alpha \beta} = \langle n \nu l \lambda; b | v^{(s)}_{\alpha} | n' \nu' l' \lambda'; b \rangle$. In general, we can calculate these matrix elements numerically. The completeness of the CS basis guarantees the convergence of the expansion with increasing $N$ and angular momentum channels.

This approximation turns the homogeneous Faddeev–Merkuriev equation into a matrix equation for the component vector

$$\psi_1^{(l)} = G^{(l)}_1(E) \psi_2^{(s)} \psi_2$$

$$\psi_2^{(l)} = G^{(l)}_2(E) \psi_1^{(s)} \psi_1,$$

where

$$G^{(l)}_\alpha = \langle n \nu l \lambda; b | G^{(l)}_\alpha | n' \nu' l' \lambda'; b \rangle_{\alpha}.$$  

(25)

The Green’s operator $G^{(l)}_\alpha$ is too complicated for a direct evaluation. However, in the Faddeev-Merkuriev equation it generates only $\alpha$-type asymptotic configurations where particles $\beta$ and $\gamma$ form bound or scattering states. Therefore in this region of the three-body configuration space $G^{(l)}_\alpha$ can be linked to a simpler Green’s operator

$$G^{(l)}_\alpha(z) = \tilde{G}_\alpha(z) + \tilde{G}_\alpha(z) U_\alpha G^{(l)}_\alpha(z).$$

(26)

where $\tilde{G}_\alpha(z) = (z - \tilde{H}_\alpha)^{-1}$ and $U_\alpha = H^{(l)}_\alpha - \tilde{H}_\alpha$ with

$$\tilde{H}_\alpha = H^0 + v^C_\alpha + u^{(l)}_\alpha.$$  

(27)

Here $u^{(l)}_\alpha(y_\alpha) = Z_\alpha (Z_\beta + Z_\gamma) e^2 / y_\alpha$. This way $U_\alpha$ is of short range type, and can be approximated on the CS basis as before.

In our Jacobi coordinates, the three-particle kinetic energy can be written as a sum of two-particle free Hamiltonians

$$H^0 = h^0_{x_\alpha} + h^0_{y_\alpha}.$$  

(28)
Thus the Hamiltonian $\tilde{H}_\alpha$ of Eq. (27) appears as a sum of two two-body Hamiltonians acting on different coordinates

$$\tilde{H}_\alpha = h_{x\alpha} + h_{y\alpha},$$

where $h_{x\alpha} = h^0_{x\alpha} + v^C_{x\alpha} (x\alpha)$ and $h_{y\alpha} = h^0_{y\alpha} + u^{(l)}_{y\alpha} (y\alpha)$. So, $\tilde{G}_\alpha$ is a resolvent of the sum of two commuting Hamiltonians $h_{x\alpha}$ and $h_{y\alpha}$. Such resolvents can be expressed as a convolution integral of two-body Green’s operators

$$\tilde{G}_\alpha(z) = \frac{1}{2\pi i} \oint_C dz' (z - h_{y\alpha} - z')^{-1} (z' - h_{x\alpha})^{-1}$$

where $g_{x\alpha}(z) = (z - h_{y\alpha})^{-1}$ and $g_{y\alpha}(z) = (z - h_{x\alpha})^{-1}$. The contour $C$ should be taken in a counterclockwise direction around the singularities of $g_{x\alpha}$ such that $g_{y\alpha}$ is analytic on the domain encircled by $C$. So, to calculate the matrix elements $\tilde{G}_\alpha$, we need to calculate a contour integral of the two-body Green’s matrices $g_{y\alpha}$ and $g_{x\alpha}$. Those two-body Coulomb Green’s matrix elements can be calculated analytically for complex energies by continued fractions [7]. This is an exact representation of $g_{x\alpha}$ and $g_{y\alpha}$, consequently the thresholds are at the exact locations with the proper Coulomb degeneracy.

In this work we calculate the negative energy resonances of the $e^- Ps$ three-body system. We need to solve (16) such that in $x_1$ we have the $e^- - e^+$ pair. So, $g_{x_1}$ is a Coulomb Green’s operator with a branch-cut on the $[0, \infty)$ interval and accumulation of infinitely many bound states at zero energy. On the other hand $u^{(l)}_{y_1} (y_1)$ is absent and $g_{y_1}$ is a free Green’s operator with branch-cut singularity on the $[0, \infty)$ interval. The resonances are at $E = E_r - i\Gamma/2$. First, we need to formulate $\tilde{G}_1(E) = \tilde{G}_1(E_r + i\epsilon)$, with $\epsilon > 0$, then we need continue analytically to $E = E_r - i\Gamma/2$. For this purpose we take the contour of Fig. [3] With $\epsilon > 0$ the singularities of $g_{x_1}$ and $g_{y_1}$ are well separated and the contour encircles the spectrum of $g_{x_1}$ without touching the singularities of $g_{y_1}$. Then we change the contour analytically as shown in Fig. [4]. The contour encircles some low-lying singularities of $g_{x_1}$ resulting in its residue, while the other part of the contour is deformed to an integration along a straight line parallel to the imaginary axis. Now, we can take the $\epsilon \rightarrow -\Gamma/2$ transition. By doing so, the poles of $g_{x_1}$ submerge into the second Riemann sheet of $g_{y_1}$ but the contour stays away from the singularities of $g_{y_1}$ (Fig. [5]).

In calculating the three-body Coulomb Green’s matrix $\tilde{G}_\alpha$ the mathematical condition for the
integral in Eq. (30) is that the contour $C$ should encircle the spectrum of one of the two-body Green’s operators without incorporating the spectrum of the other. In Refs. [1,2] the contour was taken such a way that it encircled the singularities of $g_y$. However, for resonant-state energies, the bound-state poles of $g_x$ penetrate into the continuous spectrum of $g_y$. Then to meet the requirement for the contour $C$, the path around the spectrum of $g_y$ had to be taken in such a way that it descends down into the unphysical Riemann sheet. But, the integration on the unphysical sheet is rough, the Green’s matrix exhibits violent changes, and this is getting even worse for broader resonances as the contour dives deeper into the second sheet. A singularity is always very prominent, so this numerical inaccuracy does not eliminate the resonance poles and does not mask the whole phenomenon, but it makes the identification of individual resonances, especially the broad ones, less trustworthy. The contour adopted here avoids this pitfall. No integration goes on the unphysical sheet, the path of integration is far away from any singularities, so we get very reliable results with just a few integration points.

![Diagram](image.png)

**FIG. 3.** The analytic structure of $g_{y1}(E_r + i\varepsilon - z')g_{x1}(z')$ as a function of $z'$ with $\varepsilon > 0$. The operator $g_{x1}(z')$ has a branch-cut on the $[0, \infty)$ interval and accumulation of infinitely many bound states at zero energy, while $g_{y1}(E + i\varepsilon - z')$ has a branch-cut on the $(-\infty, E + i\varepsilon]$ interval. The contour $C$ encircles the spectrum of $g_{x1}$ and avoids the singularities of $g_{y1}$.

### III. RESULTS

We calculate the $L = 0$ resonances of the $e - Ps$ system. We used atomic units throughout. We have to select $l$ and $\lambda$ such that $l - \lambda = 0$. The two electrons are identical and the two spins can be coupled either to $S = 0$ or $S = 1$. The $S = 0$ state is antisymmetric with respect to the exchange of the spin coordinates, and consequently it should be symmetric with respect to exchange the
FIG. 4. The contour of Fig. 3 is deformed analytically such that it encircles the low-lying bound-state poles of \( g_{x_1} \) and the other part is taken along an imaginary direction.

FIG. 5. Analytic continuation to \( \varepsilon = -\Gamma/2 \). The low-lying poles of \( g_{x_1} \) submerge onto the second Riemann sheet of \( g_{y_1} \), and they are denoted by dotted contour.

electron spatial coordinates. So, if \( S = 0 \) we have \( p = 1 \) in Eq. (16), while if \( S = 1 \) we have \( p = -1 \).

We have two parameters to vary in the calculations. One is the scale parameter \( b \). We found a good stability in our results with \( b = 0.25 \). The other parameter is \( N \), the maximal radial quantum number in the expansion of the potentials in each angular momentum channel and in \( x \) and \( y \) coordinates. We can see that while the narrow resonances are very stable with increasing \( N \), the broad resonances are not so. This is understandable since the broad resonances are always hard to calculate. But this inaccuracy does not change the whole picture. Individual resonances may vary a little bit with increasing \( N \) and with changing the parameters \( x_0 \) and \( y_0 \), but the series of resonances originating from the threshold are there. We found \( N = 32 \) big enough for stable results.

Figures (6) and (7) show the resonances between various thresholds using three cut-off parame-
ters. False resonances may occur in the Faddeev-Merkuriev method. They are associated with the possible bound states of $H^{(l)}$. But, by taking $x_0$ about the same size as the two-body subsystem in $x$, and varying $(x_0, y_0)$, we can avoid them. We can see that the broad resonances line up along a straight line with increasing spacings. It seems that at higher thresholds we have more lines.

![Graph](image)

**FIG. 6.** $^1S$ and $^3S$ resonances of the $e - Ps$ system between the $n = 1$ and $n = 2$ thresholds. Black circles indicate data using cut-off parameters $x_0 = 4, y_0 = 12$, red dots indicate data using cut-off parameters $x_0 = 8, y_0 = 24$, and blue + symbols indicate data using cut-off parameters $x_0 = 8, y_0 = 12$. Thresholds indicated with vertical bars.

**IV. SUMMARY AND INTERPRETATION**

In this work we re-examined the broad resonances lining up around thresholds. We solved the Faddeev-Merkuriev integral equations by adopting a Coulomb-Sturmian based separable expansion approach on the potential in the three-body configuration space. This method approximates only the asymptotically irrelevant short range potentials. The asymptotically relevant parts are kept in the Green’s operator $\tilde{G}_\alpha$, and its CS matrix elements have been evaluated as a complex convolution integral of the two-body Green’s matrices. The adopted contour makes the calculation of $\tilde{G}_\alpha$ numerically exact, even for very broad resonances, and ensures that all the thresholds are at the right location. The only real approximation is that the basis in each coordinates are truncated to a finite $N$, but we found $N = 32$ big enough for very reliable results.

We found that some of those resonances lie along a straight line. From Fig. 6 we can see that for those resonances the ratio $\epsilon_m/\Delta\epsilon_m \approx 1.23 \pm 0.03$, where $\epsilon$ is the center of mass energy measured.
FIG. 7. $^1S$ and $^3S$ resonances of the $e^- Ps$ system between the $n = 2$ and $n = 3$ thresholds. Black circles indicate data using cut-off parameters $x_0 = 12, y_0 = 30$, red dots indicate data using cut-off parameters $x_0 = 15, y_0 = 35$, and blue + symbols indicate data using cut-off parameters $x_0 = 15, y_0 = 30$. Thresholds indicated with vertical bars.

form the thresholds. They must have a common origin.

It appears that these series of resonances have not been reproduced by other methods. Careful examination and comparison with previous resonance calculations revealed that long-lived resonances were calculated previously along with Feshbach resonances. They are called shape resonances in the atomic physics community. As early as 1962-1963 in the work of Gailitis and Damburg [8] their presence as T-matrix oscillations above thresholds were calculated.

Hu and Caballero [9] carried out a six open-channel high precision calculation of the $e^+ + H(n = 2)$ scattering system using the Faddeev-Merkuriev differential equations. The calculation involves no intermediate approximations nor truncation of any kind. The numerical method solved a half million coupled linear equations. All scattering properties were calculated in the range of energy just above the $Ps(n = 2)$ formation threshold. The results display singularities of the $K$-matrix $(\tan \delta)$ (the phase shift jumps by $\pi$), the cross section maxima and the six channel wave amplitudes. All display three resonances within a cutoff distance of $1000a_0$, where $a_0$ is the Bohr radius, in channels $p + Ps(n = 2, l = 0)$ and $p + Ps(n = 2, l = 1)$. These calculations revealed a previously unknown, but relatively simple formation mechanism for these kind of resonances.

For the $Ps(n = 2)$ target, the Coulomb degeneracy allows the incoming proton to induce a first order constant electric dipole moment $\mu_1$ on the target, known as the first order Stark effect. By analyzing the channel wave functions in resonance channels $p + Ps(n = 2, l = 0)$ and $p + Ps(n =$
Hu and Caballero [9] found that the resonant conditions arise when the center-of-mass energy of the proton satisfy the simple relations

$$\epsilon_m = m\mu_1/y_m^2, \quad \epsilon_m/\Delta\epsilon_m = \text{constant},$$

where $m = 1, 2, \ldots$ integer, and $y_m$ is the location (Jacobi coordinate) of the proton at the resonance.

The Stark effect is a universal phenomenon. Furthermore, if the target is not degenerate the second order Stark effect takes place. The induced electric dipole moment is $\mu_1 = \alpha/y^2$, where $\alpha$ is the second order polarizability. Then the resonant condition becomes

$$\epsilon_m = m\mu_1/y_m^2 = m\alpha/y_m^4, \quad m = 1, 2, \ldots.$$

These explain our observations that the resonances are lining up from the thresholds with increasing spacing. Also, in the energy region between $n = 2$ and $n = 3$ thresholds the induced electric dipole moment $\mu$ is different for $l = 0$ and $l = 1$, so we observe some splitting of the lines as well.

**Author Contributions:** This method of calculating three-body resonances has been designed by Z. Papp, the calculations were carried out by D. Diaz, and the interpretation has been provided by C.-Y. Hu.

**Conflict of Interest:** The author declare no conflict of interest.

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