

Article

Low-Energy Elastic Electron Scattering from Helium Atoms

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Abstract: We reinvestigate a key process in electron-atom collision physics, the elastic scattering of electrons from helium atoms. Specifically, results from a special-purpose relativistic polarized-orbital method, which is designed to treat elastic scattering only, are compared with those from a very extensive, fully ab initio, general-purpose B-spline R-matrix (close-coupling) code.

Keywords: electron scattering; helium; cross section; polarized orbital; B-spline R-matrix; close-coupling

1. Introduction

One of the most-frequently investigated collision problems in atomic physics has been the elastic scattering of electrons from helium atoms. Although often considered a “solved problem”, especially when it comes to modelling applications in plasmas or astrophysics, attempts to increase the accuracy of the predictions continue to be reported. In fact, many special-purpose computational approaches have been developed solely to achieve this goal. The most recent example of such an attempt is the paper by Wan et al. [1] who state in their abstract: “Our calculated phaseshifts improve the best published results by at least one order of magnitude.”

While the method employed by Wan et al. is indeed capable of determining highly accurate phaseshifts, there are certain aspects of their approach, and others like it, which ultimately require sacrificing the rigour of the underlying physics to achieve these impressive gains in numerical accuracy. We note, for example, that Equation (1) of Wan et al. [1] contains a model potential with an approximate form of a *local* dipole-polarization potential. Both the cut-off radius and the actual form of the cut-off function, while known to be working well, are not based on first principles. Furthermore, the parameters chosen by Wan et al. yield a highly accurate nonrelativistic total energy of the ground state for an infinite nuclear mass, but this is not the actual energy of the system. As pointed out by those authors themselves, going from an infinite nuclear mass to the proper reduced mass will change the results, and so will relativistic and even quantum electrodynamic effects. In addition to such inaccuracies based on the omission of some known (or even unknown) physics in a model, the numerical accuracy of any computational method is limited as well.

Oleg Zatsarinny, to whom this Special Issue is dedicated, was well aware of both the physical and numerical limitations of calculations for structure and particularly collision problems. He would first define the physics (e.g., non-relativistic, semi-relativistic, or full-relativistic) of his approach and then try to solve the resulting equations as best he could with the computational resources available. He would compare with experiment, but use experimental data only reluctantly to further “improve” his ab initio calculations. For example, he might adjust the theoretical energies from his structure calculations to the values recommended by his colleagues and friends at the National Institute of Standards and Technology (NIST) [2], but only if that seemed necessary to allow for a direct comparison with experimental data that would have a smaller uncertainty in their



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energy resolution than the deviations between his theoretical threshold values and the NIST recommendations.

For many practical collision experiments, such high accuracy is not easy to achieve. Furthermore, experimental data for cross sections often have relatively large uncertainties due to the difficulties associated with their *absolute* calibration. The e-He collision system remains very important, since results for this problem are often utilized as a reference for both the energy scale and to normalize *relative* data for electron collisions with more complex targets via a relative-flow technique [3].

The purpose of the present contribution to this Special Issue is to investigate how predictions from a very *general* approach developed by Oleg to treat elastic scattering, excitation, and ionization not only from the ground state but also from and between excited states of atomic and ionic targets (including helium), compare to those from models that were designed to treat just one. Elastic e-He (a closed-shell target) scattering below the first excitation threshold, specifically below the famous $((1s2s^2)^2)$ resonance at 19.367 (1) eV [4], is such a process. There are numerous special-purpose methods around, some of which are mentioned by Wan et al. [1]. Here we present results from another such special-purpose method, the relativistic polarized orbital (RPO) approach described in the next section.

The B-spline R-matrix (BSR) approach developed by Oleg is a general-purpose approach. It is, in principle, applicable to any atom or ion, since it treats all $N + 1$ electrons explicitly, i.e., N in the target plus the projectile. There are, of course, practical limitations due to the number of configurations and channels that can be handled. An overview of the method and many applications to structure, photoionization, and electron collision problems can be found in [5]. In 2006, Oleg published the code [6] and continuously worked on further extensions and improvements. The latest version of his programs (including a relativistic D(irac)BSR version) are freely available from Oleg's GitHub repository [7]. Furthermore, the BSR version is being preserved through the Atomic, Molecular, and Optical Sciences (AMOS) Gateway [8], where the source code as well as many sample input files and run scripts for test cases are made available to all interested users.

This paper is organized as follows. In Section 2, we summarize the most important aspects of the RPO and BSR methods. This is followed by our results in Section 3. The latter section is divided into subsections starting with some representative phaseshifts, followed by extraction of the scattering length, and finally the presentation of the angle-integrated elastic and momentum-transfer cross sections. In addition to comparing with the results of Wan et al. [1], we show representative data obtained by Nesbet [9] using his early benchmark variational approach for elastic scattering as well as extensive multi-channel close-coupling calculations, specifically the momentum-space convergence close-coupling (CCC) approach of Fursa and Bray [10] and the coordinate-space R-matrix with pseudo-states (RMPS) method used by Hudson et al. [11]. Finally, our findings and conclusions are summarized in Section 4.

2. Numerical Methods

We now briefly describe the numerical methods used to obtain the results presented in this paper. For more details, the reader should consult the references provided during the description. Unless indicated otherwise, atomic units (a.u.) are used throughout, i.e., energies (E) are given in Hartree (1 Hartree = 2 Ry \approx 13.6058 eV), linear momenta are obtained as $k = \sqrt{E [\text{Ry}]}$ since relativistic effects are very small, lengths are given in multiples of the Bohr radius $a_0 \approx 0.529 \cdot 10^{-10}$ m, and angular momenta in multiples of $\hbar = h/2\pi \approx 1.055 \times 10^{-34}$ kg \cdot m²/s.

2.1. Relativistic Polarized Orbital Approach

With this method, the various phaseshifts and cross sections were calculated using the relativistic optical potential method as originally described in [12]. This method is based upon an approximate solution of the Dirac-Fock scattering equations. It includes the static, polarization and exchange interactions as well as a complex optical potential

to simulate excitation and ionization processes. However, since we are only considering elastic scattering here, this optical potential is not required. The computer code has recently been used to describe electron and positron scattering from other two-electron systems, namely Be [13] and Mg [14].

The Dirac wavefunctions for the ground state of He were obtained in a single-configuration calculation using the multi-configuration Dirac-Fock code [15]. This simple procedure yields a total energy of -2.86181 a.u., i.e., practically identical to the single-configuration nonrelativistic Hartree-Fock result (-2.86168 a.u.) [16]. Using only one configuration misses a substantial amount of the correlation energy (see below).

Since we are dealing with just elastic scattering here, only three interactions need to be considered. The first two are given by the local potential $U(r) = U_s(r) + U_p(r)$, where $U_s(r)$ is the static potential and $U_p(r)$ is the (long-range) polarization potential. The third interaction is given by the (short-range) nonlocal exchange terms.

The static potential was determined in the usual manner from the ground-state atomic orbitals, while the polarization potential was generated by the polarized-orbital method [17]. It included the first five static multipole potentials plus the corresponding dynamic polarization potential. The precise form of this dynamic potential is discussed in McEachran and Stauffer [18,19] and references therein. Finally, the exchange interaction was given by anti-symmetrizing the total scattering wavefunction. The polarized-orbital method yielded dipole and quadrupole polarizabilities of $1.3222 a_0^3$ and $2.3362 a_0^5$, respectively. The dipole and quadrupole polarization potentials were then scaled to $1.3840 a_0^3$ [20] and $2.4459 a_0^5$ [21] to account for as much of the relevant physics as possible. Since there are no absorption processes, we refer to this method as the relativistic polarized-orbital method (RPO). Apart from the above scaling of the dipole and quadrupole polarization potentials, the RPO method is *ab initio*.

The radial integral equations for the large and small components of the scattering wavefunctions, $F_0(x)$ and $G_0(x)$, can be expressed in matrix form as

$$\begin{pmatrix} F_0(x) \\ G_0(x) \end{pmatrix} = \begin{pmatrix} v_1(kx) \\ v_2(kx) \end{pmatrix} + \frac{1}{k} \int_0^x dr G(x,r) \left[U(r) \begin{pmatrix} F_0(r) \\ G_0(r) \end{pmatrix} - \begin{pmatrix} W_P(\kappa;r) \\ W_Q(\kappa;r) \end{pmatrix} \right]. \quad (1)$$

Here the functions $v_1(kr)$ and $v_2(kr)$ are the corresponding free-particle wavefunctions given in terms of Riccati-Bessel functions, while $W_P(\kappa;r)$ and $W_Q(\kappa;r)$ are the large and small components of the exchange terms. The parameter κ is the relativistic angular momentum of the incident electron. The energy E and the wavenumber k of the incident electron are related according to $E = k^2/(1 + \gamma)$. Here $\gamma = 1/\sqrt{1 - v^2/c^2}$ is the usual relativistic parameter, where v is the speed of the incident electron and $c = 1/\alpha \approx 137$ is the speed of light in vacuum and α is the fine-structure constant. In the cases considered here, $\gamma \approx 1$ and hence $E \approx k^2$. The Green's function $G(x,r)$ can be expressed in terms of Riccati-Bessel and Riccati-Neumann functions (see Equation (23) of [12] for details).

Equations (1) were solved in a noniterative manner, and the corresponding phaseshifts δ_ℓ^\pm were determined from the asymptotic form of the large component of the scattering wavefunctions. Here δ_ℓ^+ and δ_ℓ^- are the spin-up and spin-down phaseshifts respectively corresponding to $j = \ell \pm \frac{1}{2}$, where j and ℓ are the total and orbital angular momenta of the incident electron. From these phaseshifts, the elastic cross section is obtained as

$$\sigma_{\text{el}}(E) = \frac{4\pi}{k^2} \sum_{\ell=0}^{\infty} \left[(\ell + 1) \sin^2 \delta_\ell^+ + \ell \sin^2 \delta_\ell^- \right]. \quad (2)$$

Since relativistic effects in He are small, the spin-up and spin-down phaseshifts are very similar, with the difference increasing as the incident energy increases. The largest difference of 0.00002 occurred in the *P*-wave at the highest energy (19.59 eV) considered here, and hence Equation (2) essentially reduces to the well-known nonrelativistic form.

2.2. Nonrelativistic B-Spline R-Matrix with Pseudostates

The present BSR calculations are based on the work described by Zatsarinny and Bartschat [22]. Specifically, it is a nonrelativistic model with 498 states included in the close-coupling expansion. We will refer to this model as BSR-498 below. With the R -matrix radius chosen as 40, the lowest 11 bound states up to a principal quantum number $n = 3$ are fairly well-described physical states, while the remaining states squeezed into this box are so-called pseudostates that represent coupling to the high-lying Rydberg states (the next 16) as well as the ionization continuum (the remaining 471). The highest state included had an energy of ≈ 76 eV.

For elastic scattering in the energy region discussed in this paper, the BSR-498 model appears to be a vast overkill. On the other hand, it is interesting to see how far one can get with a fully ab initio all-purpose approach (in contrast to special-purpose models designed to handle elastic scattering alone), like BSR-498, without adjusting any parameter. This particular model yields a ground-state energy of -2.90357 , i.e., it includes more than 99.6% of the nonrelativistic correlation energy $E_{\text{corr}} = 2.90372 - 2.86181 = 0.04191$ obtained by subtracting the highly accurate correlated result [23] from the single-configuration Hartree-Fock prediction. The static dipole polarizability comes out as 1.355, which is within 2% of the result given in [20].

3. Results and Discussion

We now present and discuss our results for the phaseshift of the partial waves with orbital angular momentum $L = 0, 1, 2$ (in this particular case $L = \ell$, due to the 1S target state) and compare them with a few selected predictions from other calculations. Since the spin-up and spin-down results for the RPO model are very close (practically identical to the number of figures given), the results shown here are the average of the two. We then concentrate on the S -wave at very low energies to extract the scattering length. We finish this section with a presentation of the angle-integrated elastic and momentum-transfer cross sections.

3.1. Scattering Phase

Tables 1–3 give the phaseshifts for the S -, P -, and D -waves, as obtained with the current RPO and BSR-498 models and compared to the results of Nesbet [9], Wan et al. [1], and Hudson et al. [11]. The latter model was the first extensive application of the then newly developed R -matrix with pseudostates (RMPS) method [24] in the framework of the widely-used Belfast R -matrix code [25].

As expected, the overall agreement between the predictions from the various models is very good, with the largest deviations occurring in the S -wave. Especially in the low-energy region, this partial wave gives by far the largest contribution to the cross section, as will be further discussed below. The contribution from the D -wave is already very small. Interestingly, the predictions of all D -wave phaseshifts except for the one reported by Wan et al. [1] agree very well with the effective-range result of O'Malley et al. [26], which gives 0.0502 as the D -wave phaseshift for $k = 1.1$ when using the more accurate dipole polarizability, and 0.0490 when using the dipole polarizability included in the BSR-498 model. Assuming that the numbers in Table II of [1] are for finite nuclear mass, a nearly 10% difference from the effective-range formula due to this effect alone seems large. As noted already by Nesbet [9], partial waves with total orbital angular momenta $L \geq 3$ can be even better described by the simple formula of O'Malley et al. [26].

Table 1. *S*-wave phaseshift as a function of linear momentum *k* for infinite nuclear mass.

<i>k</i>	Nesbet [9]	RMPS [11]	Wan et al. [1]	Present RPO	Present BSR-498
0.1	−0.1282	−0.1260	−0.1281	−0.1262	−0.1286
0.2	−0.2655	−0.2607	−0.2650	−0.2614	−0.2560
0.3	−0.4021	−0.3979	−0.4030	−0.3978	−0.4043
0.4	−0.5388		−0.5380	−0.5319	−0.5398
0.5	−0.6684	−0.6617	−0.6678	−0.6609	−0.6699
0.6	−0.7930	−0.7848	−0.7907	−0.7835	−0.7932
0.7	−0.9067		−0.9062	−0.8990	−0.9088
0.8	−1.0155	−1.0085	−1.103	−1.0069	−1.0162
0.9	−1.1163	−1.1077	−1.112	−1.1071	−1.1151
1.0	−1.2056		−1.202	−1.1998	−1.2055
1.1	−1.2848	−1.2789	−1.291	−1.2854	−1.2863

Table 2. *P*-wave phaseshift as a function of linear momentum *k* for infinite nuclear mass.

<i>k</i>	Nesbet [9]	RMPS [11]	Wan et al. [1]	Present RPO	Present BSR-498
0.1	0.00308	0.0026	0.003029	0.00304	0.00305
0.2	0.01311	0.0129	0.01269	0.01264	0.01261
0.3	0.03063	0.0306	0.02964	0.02944	0.02945
0.4	0.05519		0.05390	0.05348	0.05352
0.5	0.08605	0.0880	0.08449	0.08380	0.08386
0.6	0.1209	0.1244	0.1209	0.11856	0.11865
0.7	0.1588		0.1588	0.15535	0.15556
0.8	0.1960	0.2000	0.1960	0.19175	0.19211
0.9	0.2305	0.2349	0.2305	0.22576	0.22657
1.0	0.2626		0.2626	0.25608	0.25772
1.1	0.2932	0.2934	0.2932	0.28211	0.28524

Table 3. *D*-wave phaseshift as a function of linear momentum *k* for infinite nuclear mass.

<i>k</i>	Nesbet [9]	Wan et al. [1]	Present RPO	Present BSR-498
0.1	0.0004	0.0004693	0.00042	0.00079
0.2	0.0017	0.001658	0.00165	0.00178
0.3	0.0037	0.003658	0.00367	0.00366
0.4	0.0066	0.006497	0.00650	0.00651
0.5	0.0104	0.01006	0.01017	0.01014
0.6	0.0149	0.01448	0.01471	0.01462
0.7	0.0203	0.01970	0.02010	0.02002
0.8	0.0265	0.02567	0.02630	0.02629
0.9	0.0335	0.0323	0.03318	0.03326
1.0	0.0414	0.0396	0.04060	0.04112
1.1	0.0501	0.0450	0.04837	0.04969

3.2. Scattering Length

Table 4 gives the *S*-wave predictions for very small *k*-values that allow the extraction of the scattering length. In the present work, we do this as follows. The RPO equations can be solved in a stable way for arbitrary energies, including $E = k = 0$, and hence the scattering length can be determined directly. For the BSR-498 calculations, the asymptotic coupled-channel equations are usually solved with the STGF [27] or FARM [28] packages. These codes have limitations and, consequently, parameters that need to be set regarding how close to a threshold (from both above and below) one can go before the results become numerically unstable. Consequently, we use a similar method as Wan et al. [1]. Specifically, we use their Equation (14) up to the term of order k^2 , calculate the value of the scattering length for each *k*, and then extrapolate to $k = 0$.

Table 4. S-wave phaseshift used to extract the scattering length for infinite nuclear mass.

k	Nesbet [9]	Wan et al. [1]	Present RPO	Present BSR-498
0.01			−0.01181	−0.01224
0.02			−0.02386	−0.02430
0.03			−0.03612	−0.03679
0.04	−0.04926	−0.04919	−0.04858	−0.04949
0.05	−0.06207	−0.06219	−0.06119	−0.06236
0.06	−0.07504	−0.07516	−0.07396	−0.07539
0.07	−0.08815	−0.08824	−0.08686	−0.08855
0.08			−0.09987	−0.10183

The result is shown in Figure 1. The BSR-498 model suggests a scattering length of about 1.187 with a numerical uncertainty of less than 0.002, provided we extract it from the results generated by FARM. The largest uncertainty is most likely coming from the point at $k = 0.01$, where the extracted results start to deviate from a straight line. That such deviations will occur is more clearly seen in the STGF results. At large values of $k \geq 0.07$ they agree well with those from FARM, but start to deviate from the latter and the almost straight line when k is smaller than about 0.07. This issue could likely be addressed by changing the default numerical parameters, but our goal is to illustrate that care is required when using these general codes in such relatively extreme situations.

Looking at the FARM and RPO results, we see that the purely numerical uncertainty is much smaller (by at least an order of magnitude) than the differences between the actual results from the two models. The reasons for these differences, therefore, lie in the *physics* included in the two models.

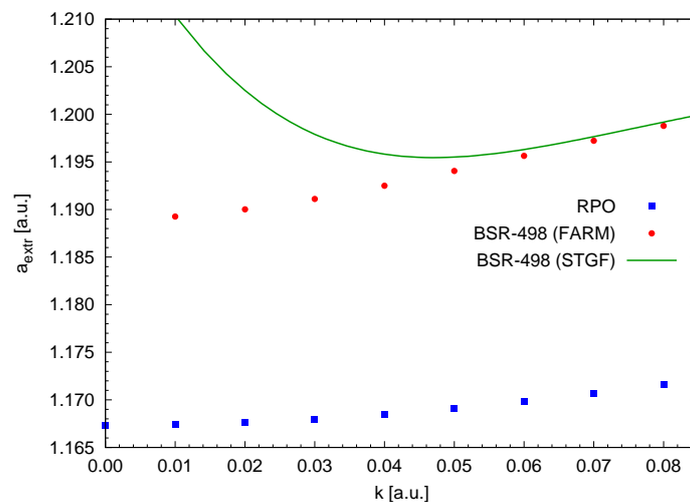


Figure 1. Extraction of the scattering length. Due to the limitations of the asymptotic FARM and STGF codes (see text for details), the smallest k -value in the BSR-498 calculation is 0.01, while the RPO code allows the calculation at zero collision energy.

3.3. Cross Sections

We now use these phaseshifts to calculate both the angle-integrated total elastic and momentum-transfer cross sections. Looking at Figure 2, we see that the RPO results are systematically lower than those obtained with BSR-498 and CCC [10], and also by Wan et al. [1]. This can be traced back to the *S*-wave, which accounts for more than 90% of the total cross section at the maximum around 1 eV and also (albeit in combination with the *P*-wave) essentially determines the momentum-transfer cross section. Even at the highest energy of 18 eV shown here, the *D*-wave only contributes about 1% of the total cross section, with the remaining 99% coming from the *S*- and *P*-waves at an approximate ratio of 3:1. For all practical purposes, all other partial waves are negligible in the energy

regime considered here. This is an example where authors have put significant effort into optimizing tiny corrections [29] rather than concentrating on the physics of the partial waves that matter most.

Given the small differences in the theoretical predictions, it is no simple task to suggest which one of those may be most accurate. Looking at Figure 1 of Wan et al. [1], at low energies almost all the theoretical results fall within the margin of error of the experimentally obtained cross section, making selection by way of best agreement with experiment difficult. Somewhat more promising is the consensus expressed in Figure 1 of the review by Alves et al. [30]. Note that this consensus was *not* reached by including CCC results (they were accidentally omitted) or BSR-498 predictions. Instead, it was based on data from swarm measurements that are used extensively by the plasma community. The inset of this figure shows the momentum-transfer cross section peaking near 2 eV with a height between 6.9 and $7.0 \times 10^{-16} \text{ cm}^2$. This confidence range includes the BSR-498 predictions, while the peak height of the RPO results ($\approx 6.8 \times 10^{-16} \text{ cm}^2$) appears to be slightly too low.

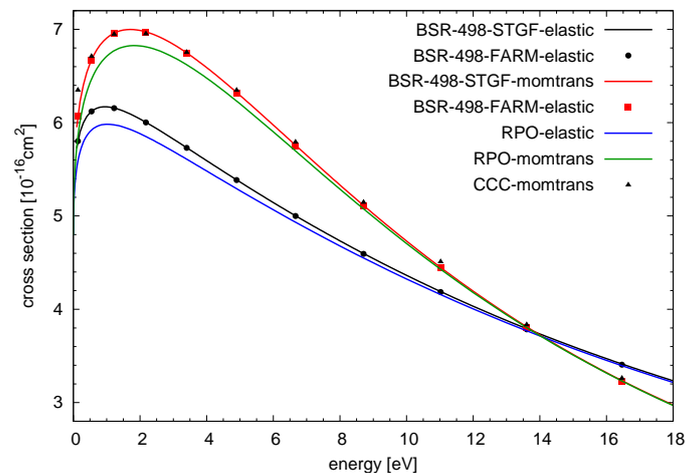


Figure 2. Angle-integrated elastic and momentum-transfer (momtrans) cross section for e-He scattering. The lines show the present results obtained with RPO and the STGF asymptotic code for the BSR-498 model. The symbols show the same results for the FARM asymptotic package and the CCC results of Fursa and Bray [10] taken from the LXCat database [31].

4. Conclusions

In this contribution, we compared results for elastic electron scattering from helium atoms obtained in a number of special-purpose (elastic scattering only) and general-purpose (multi-channel close-coupling) approaches. These included our current RPO and BSR-498 results. As expected, relativistic effects in this collision system are for all practical purposes negligible compared to other important physics, such as using a finite nuclear mass [1] and channel coupling. We obtain excellent agreement with other sophisticated calculations, even if *no adjustments* at all (such as using a more accurate value for the dipole polarizability) are made in the BSR-498 calculation. At collision energies below ≈ 2 eV, the differences between the RPO and BSR-498 results are as large as 3.5%, almost entirely due to the differences in the *S*-wave phaseshifts in this energy region. Furthermore, we have shown that the much simpler RPO method, which attempts to accurately represent the polarization interaction, can compete favorably with much more sophisticated methods (BSR and CCC) in the elastic scattering region. While one may argue which theoretical predictions are more accurate, we note that the BSR-498 results agree somewhat better with the consensus reached by the plasma community. Overall, it seems appropriate to assign a remaining uncertainty of a few percent to any of these predictions.

Finally, if the desired accuracy is indeed even higher than what has been achieved so far, it seems clear that efforts should be directed to solidifying the *S*-wave and *P*-wave

results in this purely elastic energy region, rather than worrying about small corrections to less significant contributions. This approach, of pursuing the fullest description of the physics of a problem rather than simply aiming for the greatest accuracy, was the true *modus operandi* of Oleg Zatsarinny. We believe that he would be very proud of what could be achieved with his approach to the e-He collision problem.

Author Contributions: The project was conceived R.P.M. who also carried out the RPO calculations. The BSR-calculations were performed by K.R.H. and K.B. Most of the manuscript was prepared by K.B. All authors have read and agreed to the published version of the manuscript.

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Data Availability Statement: The data shown in the manuscript are available in tabular form from the authors upon request.

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Abbreviations

The following abbreviations are used in this manuscript:

RPO Relativistic Polarized Orbital
BSR B-Spline R-matrix

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