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Benchmark calculations of electron impact electronic excitation of the hydrogen molecule

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Abstract
We present benchmark integrated and differential cross-sections for electron collisions with H₂ using two different theoretical approaches, namely, the R-matrix and molecular convergent close-coupling. This is similar to comparative studies conducted on electron–atom collisions for H, He and Mg. Electron impact excitation to the b³Σ⁺ᵤ, a³Σ⁺_g, B¹Σ⁺_u, c³Π_u, E F¹Σ⁺_g, C¹Π_u, e³Σ⁺_u, h³Σ⁺_g, B¹'iΣ⁺_u and d³Π_u excited electronic states are considered. Calculations are presented in both the fixed nuclei and adiabatic nuclei approximations, where the latter is shown only for the b³Σ⁺_u state. Good agreement is found for all transitions presented. Where available, we compare with existing experimental and recommended data.

Keywords: electron molecule collisions, electronic excitation, R-matrix, convergent close coupling, differential cross sections

Supplementary material for this article is available online

(Some figures may appear in colour only in the online journal)

1. Introduction
Molecular hydrogen is one of the simplest, most abundant molecules in the Universe. Understanding of how it interacts with its surroundings is of vital importance for a large variety of physical systems, both naturally occurring and man-made e.g. fusion plasmas, planetary atmospheres and interstellar medium. In these environments, H₂ molecules are subject to frequent collisions with low to high-energy electrons.

The equations that govern electron–molecule collisions are well understood; however, accurate and reliable cross-sections for the different processes that can occur are few and far between. Several recommended cross-section datasets for H₂ have been assembled and published (Pitchford et al 2017, Tawara et al 1990, Yoon et al 2008), and yet, in their most recent review, Anzai et al (2012) note that benchmark cross-sections are still not available for a variety of cases. Thus far the vast majority of recommended H₂ data are based on experimental results. However, due to practical reasons these data...
cannot always be obtained via experiment. For example, the required target may be unstable (short-lived), or hazardous, or both e.g. T₂.

Furthermore, it is often difficult to obtain complete sets of data that contain all the cross-sections of interest across the required energy ranges. Therefore we must often rely on theory to provide this information. In addition, if cross-sections are required from an initial state other than the ground state then theory is presently the only realistic option.

In this work we use molecular convergent close-coupling (MCCC) theory and R-matrix theory to produce a set of high-accuracy, benchmark cross-sections for electron impact electronic excitation. This is similar in spirit to the convergence of the continuum. Use of BTO greatly extends the range of the possible R-matrix radius. Here we use a BTO-only continuum, a large molecular R-matrix radius of Rₐ = 1.40 Å and a triply-augmented target basis set especially designed for Rydberg-like orbitals.

### 2.1. Target model

The R-matrix method relies on a balanced description of the target and scattering wavefunctions, N and N + 1 respectively (Tennyson 1996b). Where N is the number of electrons in the target. Molecular hydrogen is a two electron system. Therefore we have aimed to use the most comprehensive models available in each case. Full-CI is the hallmark of accuracy in electronic structure methods and it provides an exact solution to the Schrödinger equation within a given finite-sized one-electron basis set. This method is used with an augmented Dunning basis set, especially designed to describe Rydberg-type excitations in molecules, x-aug-cc-pVXZ (Dunning 1989, Woon and Dunning 1994). x-aug signifies that the basis set is doubly, triply, quadruply-augmented where x = d, t, q, etc. Triply augmented means that three additional, even-tempered basis functions are added for each angular symmetry available in the original cc-pVXZ set. Traditional Dunning basis sets, cc-pVXZ, are correlation consistent and hence provide a systematic way of approaching the complete basis set limit as the number X of zeta functions is increased. Preliminary work found that moving from a singly augmented basis set to a triply augmented basis set had a more profound effect on the target description than increasing the number of zeta functions i.e. pVXZ for X = D, T, Q, etc. For the R-matrix calculations, presented in this work, we found that t-aug-cc-pVTZ was the optimal choice (tAVTZ thereafter).

As mentioned previously, we are using Full-CI and the tAVTZ basis set. Therefore, our target model (in D₂h symmetry) can be expressed as (19, 9, 9, 4, 19, 9, 4).² Using

### Table 1. Absolute target energies, E (a.u.), at the equilibrium bond length Rₑ = 1.40 Å compared to accurate electronic structure calculations.

<table>
<thead>
<tr>
<th>State</th>
<th>Ref</th>
<th>RM²</th>
<th>MCCC³</th>
</tr>
</thead>
<tbody>
<tr>
<td>X₁⁺Σ₂⁺</td>
<td>−1.174⁴</td>
<td>−1.173</td>
<td>−1.162</td>
</tr>
<tr>
<td>b₁⁺Σ₂⁺</td>
<td>−0.784⁵</td>
<td>−0.784</td>
<td>−0.770</td>
</tr>
<tr>
<td>a₁⁺Σ₂⁺</td>
<td>−0.713⁵</td>
<td>−0.713</td>
<td>−0.710</td>
</tr>
<tr>
<td>B₁⁺Σ₂⁺</td>
<td>−0.705⁵</td>
<td>−0.705</td>
<td>−0.697</td>
</tr>
<tr>
<td>c₁⁺Π₂₀</td>
<td>−0.706⁵</td>
<td>−0.706</td>
<td>−0.701</td>
</tr>
<tr>
<td>EF₁⁺Σ₂⁺</td>
<td>−0.691⁶</td>
<td>−0.691</td>
<td>−0.687</td>
</tr>
<tr>
<td>C₁⁺Π₂₀</td>
<td>−0.688⁶</td>
<td>−0.688</td>
<td>−0.683</td>
</tr>
<tr>
<td>e₁⁺Σ₂⁺</td>
<td>−0.643⁶</td>
<td>−0.643</td>
<td>−0.640</td>
</tr>
<tr>
<td>h₁⁺Σ₂⁺</td>
<td>−0.630⁶</td>
<td>−0.630</td>
<td>−0.628</td>
</tr>
<tr>
<td>d₁⁺Π₂₀</td>
<td>−0.628⁶</td>
<td>−0.628</td>
<td>−0.626</td>
</tr>
<tr>
<td>B₁⁺Σ₂⁺</td>
<td>−0.625⁶</td>
<td>−0.625</td>
<td>−0.625</td>
</tr>
</tbody>
</table>

²Kolos and Szalewicz (1986).
³Staszewska and Wolniewicz (1999).
⁴Staszewska and Wolniewicz (2002).
⁵Wolniewicz and Dressler (1994).
⁷This work.
2.1.2. Scattering model. In the R-matrix method the electronic density of the target must be contained within the R-matrix sphere, which is of radius \( r \). Due to the extremely diffuse nature of our basis set we used a radius of \( r = 100 \text{a}_0 \). Usually an R-matrix sphere of this size would be impossible, as the continuum basis set required to fill the space would suffer from severe linear dependence. However, as mentioned previously, the new UKRmol+ codes allow the use of BTOs which are numerically stable regardless of the size of the R-matrix sphere. We found that, for molecular hydrogen, using a BTO only continuum basis not only removed linear dependence issues but it also gave a better description of the continuum. Details of the continuum basis can be found in table 2.

To solve the scattering problem we are using a close-coupling expansion. This is necessary for describing exchange and polarisation effects in addition to modelling electronic excitation. To ensure balance between the \( N+1 \) and \( N \)-electron contributions in the close-coupling expansion we use a similar treatment in the \( N+1 \) electron system as we did for the target. We adopt two types of configuration state function (CSF) in the \( N+1 \) system. There are those where one electron occupies a continuum orbital, and those where all of the \( N+1 \) electrons occupy the target molecular orbitals. This amounts to:

\[
\text{(target)}^2\text{(continuum)}^1,
\]

\[
\text{(target)}^3,
\]

where target stands for the complete set of target molecular orbitals. Note that in the first configuration step it is necessary to couple the target electrons to the appropriate symmetry, in order to facilitate the identification of the correct target states (Tennyson 1996a). However, there are no such constraints on the \( \ell^2 \) configurations generated in the second step. Using this model we retain all the target states below 30 eV vertical excitation energy, which is 98 states. This generates an average of 65 000 CSFs per molecular symmetry.

Up to now differential cross-sections (DCS) obtained from R-matrix calculations were generated using the program POLYDCS (Sanna and Gianturco 1998) which includes rotational excitation of the molecule but is limited to electronically elastic transitions. Therefore we have developed a new program for the calculation of DCS which includes only orientational averaging of the molecule but is applicable to electronically inelastic transitions and optionally employs the standard top-up procedure based on the first Born approximation for inelastic dipolar scattering. For details see appendix A.

2.2. Molecular convergent close-coupling

The MCCC method is a momentum-space formulation of the close-coupling theory. The target spectrum is represented by a set of (pseudo)states generated by diagonalising the target electronic Hamiltonian in a basis of Sturmian (Laguerre) functions. For a suitable choice of basis the resulting states provide a sufficiently accurate representation of the low-lying discrete spectrum and a discretisation of the continuous spectrum, which allows the effects of coupling to ionisation channels to be modelled. Expanding the total scattering wave function in terms of the target pseudostates and performing a

<table>
<thead>
<tr>
<th>Table 2. Continuum basis parameters for the continuum basis.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Property</td>
</tr>
<tr>
<td>Number of B-splines (per ( l ))</td>
</tr>
<tr>
<td>B-spline order</td>
</tr>
<tr>
<td>( l_{\text{max}} )</td>
</tr>
</tbody>
</table>

Figure 1. Potential energy curves for the ground state and the lower-lying excited states, relative to the ground state equilibrium energy. Data from Kolos and Szalewicz (1986), Staszewska and Wolniewicz (1999), Staszewska and Wolniewicz (2002), Wolniewicz and Dressler (1994) and Wolniewicz and Stasewska (2003).
partial-wave expansion of the projectile wave function leads to a set of linear integral equations for the partial-wave T-matrix elements, which are solved using standard techniques. The strength of the MCCC method is the ability to perform calculations with very large close-coupling expansions, allowing for the explicit demonstration of convergence in the scattering quantities of interest with respect to the number of target states included in the calculations and the size of the projectile partial-wave expansion.

The MCCC method has been implemented for electron and positron scattering on diatomic molecules in both spherical and spheroidal coordinates. The spherical implementation is simpler and provides an adequate description of the molecular structure at the mean internuclear separation of the H₂ ground state. We have utilised the spherical MCCC method for detailed convergence studies and the calculation of elastic, excitation, ionisation, and grand-total cross-sections over a wide range of incident energies. Spheroidal coordinates are a more natural system for describing the electronic structure at larger R, where the target wave functions become more diffuse. We have utilised the spheroidal MCCC method to calculate vibrationally-resolved cross-sections for excitation of a number of low-lying states of H₂, including scattering on all bound vibrational levels of the ground electronic state (Scarlett et al 2020a, submitted). This has allowed detailed studies to be performed for dissociation of H₂ in the ground and vibrationally-excited states (Scarlett et al 2019a, Tapley et al 2018a, Tapley et al 2018b, Scarlett et al 2018), and vibrational excitation of the X1Σ⁺ ground state via electronic excitation and radiative decay (Scarlett et al 2020c). For clarity of presentation, in the present paper we present only the spherical MCCC results. For details of the spheroidal MCCC method and comparisons of the spherical and spheroidal MCCC cross sections see Scarlett et al (2020b).

2.2.1. Target model. The MCCC target structure is obtained using a CI calculation. The basis for the CI expansion consists of products (nℓ, n’ℓ’) of one-electron Laguerre-based orbitals. To reduce the number of two-electron states generated, we allow one of the target electrons to occupy any one-electron orbital, while the other is restricted to the 1s, 2s, and 2p orbitals. The largest target basis used was N = 8 ω functions for ℓ ≤ 3, which generates a total of 491 states. To improve the accuracy of the X1Σ⁺ ground and b1Σ⁺ states, where the multicentre effects are strongest, we replace the 1s Laguerre function with an accurate H₂ 1sσg state obtained via diagonalisation of the H₂ Hamiltonian in a basis with N = 60 – ℓ functions for ℓ ≤ 8.

2.2.2. Scattering models. Fixed-nuclei (FN) MCCC calculations were performed at R = 1.448 a₀ using a number of scattering models, ranging from 9–491 states included in the
close-coupling expansion. This allowed for a detailed investigation of convergence and the effects of including various reaction channels (see Zammit et al. (2017b) for details). The MCCC results presented here were obtained from the 491-state model, which yielded convergent cross-sections for each of the transitions of interest. With regards to the partial-wave expansion of the projectile wave function, we have included angular momenta up to $L_{\text{max}} = 8$, and all total angular momentum projections up to $M_{\text{max}} = L_{\text{max}}$. To account for the contributions from higher partial waves we utilise an analytical Born subtraction (ABS) technique, which is equivalent to replacing the $L > L_{\text{max}}$ cross-sections with their respective partial-wave Born cross-sections. We have found that the partial-wave expansion with $L_{\text{max}} = 8$ produces convergent integrated cross sections (ICS) for all transitions considered here when used in conjunction with the ABS technique. For dipole-allowed transitions, the partial-wave convergence of the DCS can be considerably slower than it is for the ICS. The method we have adopted to resolve this issue is discussed in Zammit et al. (2017b). For the $X^1\Sigma_g^+ \rightarrow b^3\Sigma_u^+$ transition, adiabatic-nuclei calculations have been performed at low incident energies using a model consisting of 12 target states which yields convergent cross-sections for the $b^3\Sigma_u^+$ state below approximately 15 eV. These calculations are described in Scarlett et al. (2017).

2.3. Adiabatic-nuclei approximation

So far we have discussed FN calculations. However, in reality, the molecular geometry is not fixed and experimentally samples from a range of initial and final states. This will have an impact on both the integrated and differential cross-sections. This behaviour is most notable near threshold (Stibbe and Tennyson 1998). At higher scattering energies, away from the threshold, the two approximations converge as nuclear motion effects become less significant. We use the Adiabatic-nuclei (AN) approach detailed in Lane (1980) which has been recently demonstrated by Scarlett et al. (2017) on molecular hydrogen. In this work we use the ground vibrational wavefunction to vibrationally average multiple FN calculations, carried out at a range of different nuclear geometries. Although, in general, this method can also be used to produce vibrationally resolved cross-sections.

3. Results

In this section, we present FN ICS and DCS for elastic and inelastic processes. For inelastic processes we consider the first ten electronic excited states. In the second section we use the adiabatic-nuclei approximation to introduce nuclear motion effects which are particularly important close to threshold. The FN R-matrix ICS and DCS data are provided as supplementary data, which can be found online at https://stacks.iop.org/JPB/53/145204/mmedia.

The scattering calculations that follow were carried out at the mean vibrational bond length, $R_0 = 1.448\, a_0$, to
Figure 7. ICS for the $X^1\Sigma^+_g \rightarrow c^1\Pi_u$ transition. Comparison of the UKRMol+ and MCCC calculations with the recommended data of Yoon et al (2008).

Figure 8. ICS for the $X^1\Sigma^+_g \rightarrow e^3\Sigma^+_u$ transition. Comparison of the UKRMol+ and MCCC calculations with the recommended data of Yoon et al (2008).

provides the best comparison to experiment, within the FN approximation. Table 3 lists the target states and the vertical excitation energies obtained for both methods. Similarly, compared to table 1, the R-matrix target energies are more accurate than the MCCC method, as they are lower in energy (note that both methods are variational). However, it should be noted that the absolute energy is of less significance for this work, and that the vertical excitation energies (relative to the ground state) are in good agreement.

3.1. Fixed-nuclei cross-sections

We present ICS and DCS for the first ten target states (see table 3): $X^1\Sigma^+_g$, $b^3\Sigma^+_u$, $a^3\Sigma^+_g$, $B^1\Sigma^+_g$, $c^3\Pi_u$, $E^1\Sigma^+_g$, $C^4\Pi_u$, $e^3\Sigma^+_u$, $h^3\Sigma^+_u$, $B^1\Sigma^+_g$ and $d^3\Pi_u$. Where available, recommended cross-sections and experimental results are plotted against the two theoretical calculations.

3.1.1. Elastic cross-sections. The elastic ICS (figure 2) demonstrates good agreement between MCCC and R-matrix theory. The calculated data lie within the error bars of the experiment conducted by Muse et al (2008). For the DCS (figure 3) at scattering angles exceeding $15^\circ$ the two theories essentially overlap. At energies greater than 15 eV the R-matrix calculations have a diminished forward peak and this is due to a lack of convergence of the partial wave expansion. Due to computational constraints $L_{\text{max}} = 6$ for the R-matrix calculations. This compares to $L_{\text{max}} = 8$ for the MCCC calculation, which also employs the ABS technique. Nevertheless, scattering angles close to $\theta = 0$ or $\theta = 180^\circ$ do not contribute as much to the ICS due to a sin $\theta$ term in the integrand. Therefore, despite the differences in the DCSs the resulting ICSs are similar.

The recommended data of Yoon et al (2008) for the ICS are noticeably lower than those obtained from the R-matrix and MCCC calculations (figure 2). Whilst they are within their specified margin of error ($\pm 20\%$) we believe that, due to the excellent agreement between both theories and experiment for the DCS (figure 3), the recommended data should be revised.

3.1.2. Triplet states. The first excited electronic state is $b^3\Sigma^+_u$. For this state we have used a fine energy grid for both the MCCC and UKRMol+ calculations. This allows an accurate comparison of the two ICSs. In figure 4 prominent resonance structures are observed near 12 eV. Across the energy range considered the two calculations agree. The recommended cross-sections (Yoon 2008) are too large compared to the MCCC and UKRMol+ results above 12 eV. The newer experiment from Zawadzki et al (2018) is much closer to the two theories. The DCSs (figure 5) also agree closely with these experimental data. The R-matrix calculations are a little higher than the MCCC calculations for angles exceeding $135^\circ$ but, again, the effect on the ICS is insignificant.

For higher excited states i.e. those energetically above $b^3\Sigma^+_u$, MCCC results are presented on a coarser energy grid. Therefore, we can no longer compare the narrow resonant structures. The ICSs for states $a^3\Sigma^+_g$, $c^3\Pi_u$ and $e^3\Sigma^+_u$
Figure 10. DCS for the $X^1\Sigma^+_g \rightarrow c^3\Pi_u$ transition. Comparison of the UKRMol+ and MCCC calculations with the measurements of Wrkich et al (2002).

(figures 6–8, respectively) have good agreement between the MCCC and R-matrix theories.

The recommended data points are based on the EELS (electron energy loss spectroscopy) experiment of Wrkich et al (2002). The data points are sparse so it is hard to quantitatively compare against the two theory calculations. However, given agreement between the two theoretical calculations and more recent experiments, we believe that the recommended cross-sections should be revised for all of the triplet states considered so far.

The DCSs shed more light on the comparison. Figure 9 shows the $a^3\Sigma^+_g$ state. Agreement is best for 17.5 eV and 30 eV. The general shape is present at all three energies. That is, the cross-section dips around 60° and 120°. However, for intermediate angles the magnitude of the DCS is higher (especially for 20 eV) than the theoretical calculations. EELS experiments are hard to conduct for excited states of H$_2$ because the states overlap in the spectra and the individual components have to be deconvoluted. Based on the difficulty of these type of experiments for highly-excited states we suggest that the calculations are more reliable.

For the $c^3\Pi_u$ state (figure 10) the situation is similar to the $a^3\Sigma^+_g$ state. There is a slight downward slope towards higher scattering angles that is present in both the calculation and the experiment. However, the experimental DCS at 20 eV is approximately an order of magnitude higher.

For the $e^3\Sigma^+_u$ state there is no qualitative agreement between theory and experiment. At all three energies (shown in figure 11) we have large discrepancies for low angle scattering i.e. below 30°. This is not too surprising though as low and high angle scattering is difficult to measure due to the physical constraints of the experimental setup. Therefore, we suspect that the low angle cross-sections measured by Wrkich et al (2002), at 20 eV and 30 eV, are too high.

The ICSs for states $h^3\Sigma^+_g$ and $d^3\Pi_u$ (figures 12 and 13) show reasonable agreement between the two theories. However, the R-matrix calculation exhibits pronounced features around 16 eV and 19 eV. In the standard R-matrix approach used in this work, ionisation effects are not included. We include states above the ionisation threshold but we do not explicitly include pseudostates. To model ionisation pseudostates are required as implemented in the R-matrix with pseudostates (RMPS) method (Gorfinkiel and Tennyson 2005). As a result, the cross-section is overestimated above ionisation threshold. This behaviour was demonstrated
previously in MCCC calculations when only the bound states were used (Zammit et al. 2017b). In addition, weak transitions can also suffer from small oscillations, but the impact is reduced as the size of the close-coupling expansion increases. Therefore, the enhanced R-matrix cross-section is likely due to missing ionisation channels.

The R-matrix DCSs for state $h 3^2\Sigma^+_g$ (figure 14) show broad agreement with the MCCC data. There are no recommended data for either the $h 3^2\Sigma^+_g$ or $d 3^1\Pi_u$ states. For the $d 3^1\Pi_u$ state (figure 15) there are more significant differences between the two theories. As the target excitation increases, we typically expect less agreement between the two theories. Higher excited states tend to be less accurately described by the electronic structure calculations used in the R-matrix method.

3.1.3. Singlet states. Next we consider the singlet states. ICSs for three dipole-allowed states, $B^1\Sigma^+_u$, $C^1\Pi_u$ and $B^1\Sigma^+_g$, are shown in figures 16–18 respectively. All three ICSs show excellent agreement between MCCC and R-matrix theory. Furthermore, agreement with the recommended data, which is available for the $B^1\Sigma^+_u$ and $C^1\Pi_u$ states, is extremely good at the energies considered here. Contrary to the previous EELS experimental data, these recommended data were measured from the optical emission of the electron impact electronically excited B and C states (Liu et al. 1998).

DCSs could not be determined in the emission experiments of Liu et al. (1998). However, Wrkich et al. (2002) produced a set of EELS DCS which have been plotted in figures 19 and 20. For the $B^3\Sigma^+_u$ state the agreement with experiment is good. At 30 eV however the R-matrix calculation displays oscillations that are not present in the MCCC calculation. This is due to a lack of convergence in the partial-wave expansion. Typically a Born correction would be applied to dipole allowed transitions. However, in the present work this has
not been possible. The Born top-up requires a sufficiently
converged cross-section, up to some intermediate number of
partial waves, $L_{\text{max}}$. For MCCC this is found to be $L_{\text{max}} = 25$, or more, depending on the scattering energy (Zammit et al 2017b). A similar approach was attempted for the R-
matrix calculation, although this was not tractable given
currently available software and computational power (see
appendix A).

Similarly, the oscillations observed for the $B^1\Sigma_u^+$ state at
higher energies are also observed in states $C^1\Pi_u$ and $B'^1\Sigma_u^+$
(figures 20 and 21). Furthermore, in all of the singlet state
DCSs, figures 19–22, the R-matrix calculation has a lower
forward peak. This is attributed, as in the elastic scattering
case, to a lack of convergence in the number of partial waves
used. Regardless, forward and backward scattering only make
a small contribution to the total ICS. Therefore the differences
caused by the oscillatory behaviour and lower forward peak are
lost upon integration. This highlights the importance of using
DCSs as a stringent test of theories. Two theories may produce
the same ICS but have different angular profiles.

In contrast to the dipole-allowed singlet states, the forbid-
den $EF^1\Sigma_g^+$ state DCS (figure 22) is not as sensitive to higher
partial-waves. Agreement between the two theories is good.
The agreement between theory and experiment is acceptable,
except for the scattering angles from 60° to 100° at 20 eV
where the experiment gives a larger cross-section, which could
be due to the analysis of the measured EELS.

Comparing the ICS (figure 23) between the two theories,
the R-matrix calculation is consistently above the MCCC data.
Again, this is likely due to the absence of ionisation chan-
nels in the R-matrix close-coupling expansion that leads to an
overestimated cross-section.

The recommended data are based on an emission experi-
ment carried out by Liu et al (2003). Whilst the $EF^1\Sigma_g^+$ state
is dipole-forbidden, the cross-section can be inferred using a
combination of theoretical and experimental considerations.
There is a difference in threshold for experiment, which occurs
near 15 eV as opposed to 13 eV for the FN MCCC and R-
matrix calculations. However the magnitude and qualitative
shape agree with theory.

### 3.2. Adiabatic-nuclei cross-sections

In this section we make use of the AN approximation described
previously. In figure 24 both FN (dot-dashed line) and AN
calculations (solid line) are shown side-by-side for electronic
excitation to the first excited state ($X^1\Sigma_g^+ \rightarrow b^3\Sigma_u^+$). For both
the MCCC (red) and UKRmol+ (black) calculations we can see
two main differences. The first is that resonant structures
are washed-out and the second is that the sharp turn-on near
the vertical excitation threshold (10 eV) is smoothed into a
ramp. This is due to the vibrational averaging over differ-
ent molecular geometries. The threshold for the $X^1\Sigma_g^+ \rightarrow b^3\Sigma_u^+$ transition is essentially the vertical excitation energy.
For some geometries this will be lower than 10 eV and for oth-
ers will be greater. The average is weighted by the square of
the ground vibrational wavefunction, which means the largest
correlations occur at the maximum of the wavefunction i.e.
about $R_0$. This is why the FN calculation at $R = R_0$ and the AN

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**Figure 17.** ICS for the $X^1\Sigma_g^+ \rightarrow C^1\Pi_u$ transition. Comparison of
the UKRmol+ and MCCC calculations with the recommended data

**Figure 18.** ICS for the $X^1\Sigma_g^+ \rightarrow B^1\Sigma_u^+$ transition. Comparison of
the UKRmol+ and MCCC calculations.

**Figure 19.** DCS for the $X^1\Sigma_g^+ \rightarrow B^1\Sigma_u^+$ transition. Comparison of
the UKRmol+ and MCCC calculations with the measurements of

**Figure 20.** DCS for the $X^1\Sigma_g^+ \rightarrow C^1\Pi_u$ transition. Comparison of
the UKRmol+ and MCCC calculations.

**Figure 21.** ICS for the $X^1\Sigma_g^+ \rightarrow B^1\Sigma_u^+$ transition.
calculation are broadly similar. Adiabatic effects have consequences for near-threshold electron impact dissociation of H$_2$ (Stibbe and Tennyson 1998).

The AN approximation requires FN calculations to be performed across a grid of different internuclear bond separations. For the R-matrix calculations, a grid size of \( \Delta R = 0.05 \) a.u. was used for \( 0.95 < R < 1.95 \) a.u. with a finer grid of \( \Delta R = 0.01 \) a.u. used in the region closer to the mean vibrational bond length, \( 1.35 < R < 1.55 \) a.u. Due to the large number of FN calculations required it was not possible to use the full model described previously. Therefore a smaller model was used which featured a singly augmented aug-cc-pVTZ basis set and an R-matrix radius \( a = 25 \) a.u. The smaller radius allowed the continuum representation to be simplified to 22 BTOs per angular momentum symmetry with \( L_{\text{max}} = 4 \) without sacrificing completeness. As before, all of the target states below 30 eV were included which led to a 59-state model. This model works well for the first excited state but due to the simplified target description it cannot represent higher-excited states.
In this paper we demonstrate good agreement with recent experimental data (Muse et al 2008, Zawadzki et al 2018), validated by two independent theories. The agreement with the recommended ICS data (Yoon et al 2008) and older experimental data (Wrkich et al 2002) is worse, predominantly for the triplet states but we believe this is due to the difficulties associated with the underlying experiments. That is, it is difficult for experiments to separate the overlapping contributions coming from different triplet excited states and therefore the error margin is larger for these types of experiment. Any other significant differences between the two theories and experiments are well understood.

This is the first time the CCC and R-matrix theories have been verified for a molecular target. This work presents one of the largest molecular R-matrix calculations to date. Many novel features have been exploited for the first time: a triply-augmented target basis set, a box size of 100 a.u. and the first B-spline only continuum for a molecular target. This shows that both MCCC and R-matrix methods can be used to perform large-scale, high-accuracy close-coupling calculations.

We have compared both fixed-nuclei and adiabatic-nuclei cross-sections obtained using the R-matrix and MCCC methods. For FN calculations, dipole-forbidden states generally show better agreement in the DCSs. Dipole-forbidden states do not require a born top-up and generally converge quicker for the same number of partial waves, compared to dipole allowed transitions (Zammit et al 2017b). For the dipole-allowed states the R-matrix calculations show oscillatory behaviour but this could be eliminated by using a higher cutoff in the number of partial-waves. However, this is currently not tractable given currently available hardware and software.

All of the ICSs show good agreement between the two theories with the exclusion of weak transitions that are more sensitive to the absence of ionisation channels in the R-matrix calculations, leading to slightly enhanced cross-sections. The AN ICS for the first excited state shows excellent agreement between the two theories and the recent experimental data.

There are several directions for future work. Firstly, it would be interesting to compare the effect of target model used in the MCCC calculations i.e. spherical versus spheroidal. Preliminary results for the EF $1\Sigma^+_g$ state suggest that the use of a spheroidal model could improve the agreement between both theories.

Secondly, in order to accurately describe ionisation effects in the R-matrix method we would need to employ the RMPS method. Whilst the RMPS method is implemented in UKRMol+ the calculations for this system are currently too expensive.

Additionally, for the R-matrix calculations presented in this work we have not been able to carry out systematic, quantitative analysis of the uncertainties. This is a common problem across the field for theoretical calculations (Chung et al 2016). For future work, we seek a tractable approach that is capable of providing uncertainties for our calculated data.

Finally, a general approach for handling Born top-ups, similar to the ABS method used in MCCC calculations would be desirable for the UKRMol+ calculations in order to reach convergence where larger numbers of partial-waves are required (as discussed in appendix A).

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Appendix A. Including higher partial waves
To include higher partial waves, specifically for dipole-allowed transitions, we require a top-up procedure. In R-matrix calculations this is done using an approach suggested by Norcross and Padial (1982). The MCCC uses an equivalent method described in (Zammit et al 2017b). For a DCS the top up procedure is given by

\begin{equation}
\frac{d\sigma}{d\Omega} = \left(\frac{d\sigma}{d\Omega}\right)_{\text{Born}} + \sum_{\lambda=0}^{\lambda_{\text{max}}} (A_{\lambda} - A_{\lambda}^{\text{Born}})P_\lambda(\cos \theta),
\end{equation}

where $A_{\lambda}$ are the coefficients from the original Born calculations, and $A_{\lambda}^{\text{Born}}$ are the coefficients from the top-up procedure, and $P_\lambda(\cos \theta)$ are the Legendre polynomials.
where the first term on the right-hand side is the DCS calculated for inelastic dipolar scattering in the first Born approximation and the second term includes the contribution of the lower partial waves $A_\lambda$ calculated with close-coupling and subtraction of the corresponding Born partial waves $A_\lambda^{\text{Born}}$. Only orientational averaging of the molecule is taken into account. This approach was used in previous R-matrix calculations for inelastic collisions, but only for ICSs (Baluja et al. 2000, Kaur et al. 2008, Mašín et al. 2012), which tend to converge quicker than DCSs. Recently, Zawadzki et al. (2020) employed the Born correction described above for the electronically inelastic DCS of CO. This method, however, requires a sufficiently high partial wave cutoff, $L_{\text{max}}$. At lower partial waves the analytic Born method is less accurate and tends to overestimate the cross-section, leading to unphysical negative cross-sections.

Born corrections have been successfully applied to DCSs for elastic collisions, see e.g. Zhang et al. (2009) and Mašín et al. (2012). However, these cross-sections are usually an order of magnitude larger than those for dipole-allowed inelastic transitions. Hence, they are less susceptible to the oscillatory behaviour seen in inelastic DCSs.

Figure A1 shows the DCS for the dipole-allowed $B^1\Sigma_u^+$ state. In solid black we have the original R-matrix calculation without the Born correction. If we apply the Born correction to the DCS we obtain the dotted line. At 17.5 eV, the Born corrected DCS displays unphysical behaviour around 150° where it becomes negative. The situation worsens for higher energies. This is due to an incomplete convergence of the partial-wave Born contribution $\sum \lambda A_\lambda^{\text{Born}} P_\lambda(\cos \theta)$.

To resolve this issue, the MCCC approach (Zammit et al. 2017b) has been to run a smaller-sized calculation but with a higher cutoff e.g. $L_{\text{max}} = 25$. The results of this calculation are then used to augment the T-matrices of the more expensive calculation. This allows the DCS contributions from higher partial waves to be calculated with the more accurate MCCC theory before including the additional contributions from the Born procedure.

A similar approach has been adopted in the R-matrix calculations, however $L_{\text{max}} = 25$ is currently not computationally feasible with the UKRmol+ codes. Calculations using a smaller model, but with $L_{\text{max}} = 10$, have been computed and these were used to augment the T-matrices of the accurate R-matrix calculation with $L_{\text{max}} = 6$. When augmenting the T-matrix elements, care must be taken to phase-match the two calculations. This can be achieved by comparing the transition dipole moments of the target states involved in each transition.

The result of augmenting the T-matrices and applying the Born correction is shown as the dashed line in figure A1. For the lowest scattering energy shown, 17.5 eV, the oscillatory behaviour is greatly reduced and the Born correction improves the quality of agreement between the MCCC and R-matrix calculations. At 20 eV Born-corrected DCS is improved but it still shows oscillatory behaviour that is characteristic of a lack of convergence. At 30 eV, even with the augmented T-matrix elements the DCS remains oscillatory when the Born correction is applied.

In theory, an approach similar to the MCCC method can be developed for the R-matrix calculations but there are two factors that currently inhibit further improvement. The first is that the target states from cheaper calculations need to be shifted to the more accurate values from the expensive calculation. For the R-matrix calculations, presented in this work, the energies were shifted in the outer-region. This is not ideal and instead we need to implement the energy shift in the $N + 1$ scattering calculation, similar to the approach used by Stibbe and Tennyson (1997). Secondly, the outer-region quickly dominates the computational resources required, both physical RAM and CPU-time, as a large number of channels are generated for higher partial waves. Furthermore a more sophisticated approach would need to be implemented in the outer-region to reduce the number of states included in the calculation.

As an alternative approach, we also attempted to top-up the DCS using a more basic method (not shown). We ran two cheaper calculations with small basis sets using $L_{\text{max}} = 6$ and $L_{\text{max}} = 10$. We took the difference between the two DCSs and used this to top-up the expensive calculation. This approach does help to capture the forward peak scattering but it was too susceptible to unphysical negative cross-sections when the differences between the cheap calculations became negative. This method behaved particularly poorly in regions where the cross-section was small.

In summary, we believe the MCCC approach to the Born top-up is the most sensible way forward, however there is still work to be done before it can be implemented in R-matrix calculations.
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