Resonance structure in low-energy electron scattering from OCS

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Abstract

Energy-dependent absolute angle-differential cross sections (θ = 135°) for elastic and vibrationally inelastic electron scattering from OCS molecules have been investigated at high resolution (13 meV). The elastic cross section, reported over the range E = 0.06–20 eV, exhibits a deep Ramsauer–Townsend minimum near 0.55 eV and the π∗(1.2 eV) and σ∗(3.8 eV) resonances known from earlier experimental and theoretical work. The excitation functions for vibrational excitation (VE) of the fundamental CS stretch (100), the CO stretch (001) as well as for the fundamental and harmonics of the bending (0n0) mode are reported from threshold up to E = 8 eV. VE is mediated—in part in a mode-selective way—by threshold peaks and the higher lying resonances. The selectivity in the threshold region can be rationalized in terms of symmetry arguments where the ²Σ virtual state excites preferentially the σ vibrational states with even quanta of bending vibration over the π vibrational states with odd quanta of bending vibration. The σCS∗ resonance is essentially inactive in VE of the CO stretch and the bending mode.

1. Introduction

The series of related triatomic linear molecules CS₂, OCS and CO₂ is of fundamental interest in connection with low-energy electron scattering. It allows for systematic studies of, e.g., the influence of the molecular polarizability whose average value increases from 19.6 au (CO₂) via 38.5 au (OCS) to 59.8 au (CS₂) [1]. OCS as the intermediate case is the only one to possess a dipole moment of 0.715 D [2]. The vertical electron affinity (VEA) at the equilibrium bending angle α = 180° strongly increases from CO₂ to CS₂; for the latter, calculations [3, 4] indicate a slightly positive value of the VEA. The calculated adiabatic electron affinities (AEA) range from −0.54 eV (CO₂, α ≈ 138°) via −0.01 eV (OCS, α ≈ 136°) to 0.46 eV (CS₂, α ≈ 143°) [4].

Both scattering theory (numerous studies, see for example [5, 6]) and measurements of total and backward cross sections at very low energies [7] indicate that electron scattering from CO₂ at low energies is dominated by a virtual state, which also causes large threshold peaks in the vibrational excitation (VE) cross sections (see, e.g., [8, 9]). Calculations of the potential energy surfaces in the region of geometries where the electron is bound (in the fixed-nuclei picture) [4, 10, 11] reveal that the anion potential surface bends down before crossing the potential of the neutral molecule. This downward-bent section of the potential surface, where an electron is weakly bound to a bent CO₂ by a combination of dipole and polarization forces, supports vibrational Feshbach resonances (VFRs) manifested as narrow structures in VE cross sections [12–14], strongly reminiscent of those found in the prototype VFR case of HF [15–17].

For CS₂ giant resonances were observed in the total and backward cross sections at very low energies [18]. The VE cross sections are very large in magnitude and exhibit narrow vibrational structure which may be attributed to vibrational levels of the bound anion state [19].

The potential surface for OCS− bends down when approaching the crossing with the neutral OCS surface in a way similar to CO₂− [4], indicating the possibility for similar vibrational resonances to occur at energies around 0.2 eV. Recent studies of the total scattering cross sections of OCS at energies from 0.07 to 0.3 eV, however, have shown no vibrational structure [20].

In this connection, it is of interest to mention recent observations of anion formation in low-energy electron interactions with small clusters (XY)₅ of XY = CO₂, OCS...
and CS₂. Rather sharp VFRs were observed for CO₂ clusters \((N \geq 4)\) and OCS clusters \((N \geq 1)\) while no such structure was observed for CS₂ clusters. These results were interpreted to reflect the effects of solvation on the molecular ion state immersed in the cluster environment \([4, 21]\).

In this paper, we concentrate on elastic and vibrationally inelastic electron scattering by the molecule OCS at energies below \(E = 10\) eV. An overview of previous work up to the year 2000 has been given by Karwasz \(\text{et al}\) \([22]\), and we refer the reader to references in this review with regard to earlier work. The total cross section, studied before at energies above 0.2 eV, is dominated by the \(\pi^*\) resonance, peaking at about 1.2 eV; it is active in mediating VE \([23]\) and dissociative electron attachment (DEA) \([23–26]\). A Ramsauer–Townsend (RT) minimum has been predicted to occur in the \(\sigma\) elastic scattering channel near 0.7 eV in theoretical work \([27, 28]\).

We note that the energy dependence of the elastic differential cross section (DCS), reported by Sohn \(\text{et al}\) over the range 0.2–5 eV at 130° (see figure 1 in \([29]\)) exhibits a rather pronounced minimum at \(E = 0.6\) eV, but was not addressed as reflecting a RT minimum by these authors. VE was studied by Sohn \(\text{et al}\) \((E = 3.0–5\) eV\) \([29]\), Abouaf \(\text{et al}\) \((E = 0.15–2\) eV\) \([23]\) and Kawada \(\text{et al}\) \([30]\). Of most relevance to the present paper are the results of Abouaf \(\text{et al}\) obtained with a resolution of 0.025 eV. These authors reported excitation functions for the \((0 \, n \, 0; \, n = 4, 5, 6)\) bending modes at \(\theta = 90°\) and energies up to 2.5 eV which showed the \(\pi^*\) resonance at 1.2–1.3 eV and threshold peaks (notably for \(n = 4, 6\)); they interpreted the latter observation as indicating that a virtual state contributes to the scattering, as confirmed by the later theoretical work \([27, 31]\). It is of obvious interest, as noted in \([23]\), to also measure the VE functions for the lower \((0 \, n \, 0, \, n = 1–3)\) bending modes to check for the importance of threshold peaks.

In the present work, we investigated low-energy electron scattering from OCS molecules at improved energy resolution (13 meV) with two major aims: (i) search for structure and systematic behaviour in the elastic and vibrationally inelastic scattering channels; (ii) determination of accurate absolute cross sections for these two channels over a range of incident energies. The measurements were limited to one detection angle (135°), chosen such that the effects of the permanent dipole moment—which dominate at small scattering angles—are suppressed. Using previously published angular distributions, our absolute differential cross sections can be used to establish integral cross sections at selected energies. The paper is organized as follows. In section 2, we briefly describe the experimental setup and procedure. In section 3, we present and discuss the experimental results. We conclude with a short summary.

2. Experiment

Electrons emitted from a hot filament are energy-selected by a double hemispherical monochromator and focused onto an effusive beam target. A double hemispherical analyser allows for detection of elastically or inelastically scattered electrons. Further details of the experiment and procedures have been described elsewhere \([32–34]\). Absolute elastic cross sections were measured by comparison with the known helium cross sections using the relative flow method \([35]\). The pressure ratio between He and OCS was chosen as roughly 2:4 according to their molecular diameters, in order to get the same beam profile. The flow rates used were 0.106–0.136 mbar cm⁻³ s⁻¹ for OCS at pressures of 0.075–0.092 mbar and 0.892–1.294 mbar cm⁻³ s⁻¹ at 0.172–0.243 mbar for He. The measurement procedure compensated for small variations of pressure and beam current during the signal accumulation. The confidence limit (2 standard deviations) is about ±20% for the elastic cross sections and ±25% for the vibrationally inelastic cross sections. The incident electron resolution was about 13 meV with a beam current of about 40 pA, due to the low electron energies and deposits on the electron optics, which resulted in the frequent need for readjustment of the electron gun. The sample was introduced through a 0.25 mm diameter nozzle kept at about 30°C.

3. Results and discussion

3.1. Elastic scattering

Absolute elastic cross sections were measured at eight different energies, and are shown by full circles in figure 1 and listed in table 1. The elastic signal was then measured as a function of the incident electron energy, corrected for the variation of the instrumental response function, normalized to the discrete absolute measurement at 1.15 eV, and is shown by a continuous line in figure 1.
as given by Altshuler (see equation (22) in [38]). Also indicated in the figure are somewhat larger than the present experiment, similarly to Gianturco and Stoecklin. We also include the cross sections of Michelin et al involving the Schwinger variational iterative method, a separable representation of the exchange potential and added correlation–polarization effects via a density functional formulation. We also include the cross sections of Michelin et al [37] involving the Schwinger variational iterative method, combined with a distorted wave approximation; their values are somewhat larger than the present experiment, similarly to those of Gianturco and Stoecklin. Also indicated in the figure is the rotationally inelastic cross section for scattering from a fixed molecule with dipole moment, averaged over orientation, as given by Altshuler (see equation (22) in [38]).

The dramatic increase of the cross section towards low energies is attributed to a virtual state [27, 31], the deep minimum around 0.55 eV is attributed to the Ramsauer–Townsend effect [27, 28]. The fact that the measured cross section is much larger than predicted by the Born approximation below 200 meV is taken as a manifestation of the virtual state, the fact that it is lower above about 350 meV as a consequence of the Ramsauer–Townsend minimum.

The peak at 1.33 eV has been assigned to a $^2\Sigma$ resonance, that at 3.7 eV to overlapping $^2\Sigma$ and $^2\Delta$ resonances, with the former being dominant [29]. This assignment of the resonances is compatible with the symmetry decomposition of the calculated integral cross section of Bettega et al [31] and of Gianturco and Stoecklin [28]. The latter group also calculated resonances at higher energies, a $^2\Pi$ resonance around 10 eV, which could correspond to the weak peak at 10.8 eV in our data in figure 1.

3.2. Vibrational excitation

The fundamental vibrations of OCS are summarized in table 2. Absolute values of the cross sections for vibrational excitation were determined from the energy-loss spectrum shown in figure 3, recorded with a constant incident energy of 1.15 eV, near the centre of the $^2\Pi$ resonance. The spectrum was corrected for the variation of the analyser response function and the area under the elastic peak was normalized to the measured absolute value. Peak overlap was partially resolved by fitting the observed spectrum to Gaussian profiles. The results are given in table 3. The spectrum is characteristic for vibrational excitation by a $^2\Pi$ resonance, as has already been pointed out in earlier work [23, 29]. The VE cross sections are large, the sum under all vibrationally inelastic peaks (up to an energy loss of 1 eV) amounts to $1.7 \times 10^{-20}$ m$^2$ sr$^{-1}$, i.e.

<table>
<thead>
<tr>
<th>No</th>
<th>Type</th>
<th>Symm.</th>
<th>Energy (meV)</th>
<th>$D$ (au)</th>
</tr>
</thead>
<tbody>
<tr>
<td>v1</td>
<td>CS str</td>
<td>$\sigma^+$</td>
<td>106.5</td>
<td>0.0243</td>
</tr>
<tr>
<td>v2</td>
<td>Bend</td>
<td>$\pi$</td>
<td>64.5</td>
<td>0.0185</td>
</tr>
<tr>
<td>v3</td>
<td>CO str</td>
<td>$\sigma^+$</td>
<td>255.6</td>
<td>0.134</td>
</tr>
</tbody>
</table>

Figure 2. Elastic cross section at an angle of 135°, shown as a function of electron energy with absolute values indicated by filled circles. Theoretical data by Bettega et al [31] are indicated by open triangles, calculated cross sections by Gianturco and Stoecklin [28] are depicted as squares and results from Michelin et al [38] are shown as diamonds. A Born approximation calculation for scattering from a polar molecule [38] is indicated by the dashed line.

Figure 3. Spectrum with constant incident energy at an angle of 135°. The ordinate scale is linear.
Differential cross section for excitation of the CS stretch vibration at an angle of 135°.

Differential cross section for excitation of the CO stretch vibration at an angle of 135°. Note that it is degenerate with four quanta of the bending vibration.

Table 3. Differential VE cross sections \((10^{-20} \text{ m}^2 \text{ sr}^{-1})\) at \(E_i = 1.15 \text{ eV}\) and 135°.

<table>
<thead>
<tr>
<th>Final state</th>
<th>DCS (10^{-20} \text{ m}^2 \text{ sr}^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>(0 1 0)</td>
<td>0.38</td>
</tr>
<tr>
<td>(0 2 0)</td>
<td>0.11</td>
</tr>
<tr>
<td>(0 3 0)</td>
<td>0.076</td>
</tr>
<tr>
<td>(1 0 0)</td>
<td>0.11</td>
</tr>
<tr>
<td>(0 0 1)</td>
<td>0.26</td>
</tr>
</tbody>
</table>

1.8 times the elastic cross section, which is \(0.94 \times 10^{-20} \text{ m}^2 \text{ sr}^{-1}\) (including rotational transitions). Not only the fundamentals but also the overtones of all three vibrations, as well as many combination vibrations are excited, a consequence of the fact that the temporary occupation of the \(\pi^*\) orbital lengthens both the CS and the CO bonds and makes the OCS− bent.

The cross sections for exciting specific vibrations as a function of electron energy were then obtained by recording the respective excitation functions, correcting them for the instrumental response function and normalizing them to the absolute values given in table 3. The results are shown in figures 4–6. The agreement with the individual data points of Sohn et al. [29] is satisfactory. The contribution of direct scattering to vibrational excitation can be estimated using first Born approximation [41] and dipole transition matrix elements determined by infrared spectroscopy [40]. Though the results at high scattering angles will not be quantitative, they allow for a survey of the different contributions to threshold peaks [42], and are shown as dashed lines in figures 4–6.

The cross section for the excitation of the CS stretch mode \((1 \ 0 \ 0)\) in figure 4 shows a considerable threshold peak and two shape resonances, one around 1.3 eV and a second around 3.8 eV. As already discussed in connection with the elastic cross section, they can be assigned as \(2^\Pi_1\) and \(2^\Sigma_1\) [28, 31, 43]. The threshold peak must be primarily due to the virtual state since only a small fraction of it is explained by the Born approximation.

The CO stretch mode is nearly degenerate with four quanta of the bending mode (energy difference around 5 meV [44]), cannot be resolved in the present experiment, and figure 5 consequently shows the sum of the cross sections for exciting both the \((0 \ 0 \ 1)\) and the \((0 \ 4 \ 0)\) states. Both a threshold peak and the \(2^\Pi_1\) band are observed. The threshold peak is narrower than the Born prediction, but the Born prediction is not expected to be very reliable at large scattering angles. Interestingly the \(2^\Sigma_1\) resonance is practically absent, indicating that the temporarily occupied \(\pi^*\) orbital is localized on, and antibonding with respect to, the CS bond and has nearly no coefficients on the CO bond.

The excitation function for the bending mode (figure 6) does not exhibit a threshold peak and this aspect will be discussed further below. The \(2^\Pi_1\) resonance is prominent, consistent with its bent equilibrium geometry. The \(2^\Sigma_1\) resonance is absent, indicating a linear equilibrium geometry, consistent with its assignment.

Sohn et al. [29] and Abouaf et al. [23] already reported a preference for exciting even quanta of the bending vibration over odd quanta. We further study this effect by recording the cross sections for exciting the \((n \ n \ 0)\) states for \(n\) up to 8, as shown in figure 7. Some of the spectra are affected by band overlap, in particular the \((0 \ 8 \ 0)\) and the \((0 \ 0 \ 2)\) states as already mentioned above. Clear trends in the spectra emerge, despite this limitation:

(i) The relative intensity of the threshold peak over the \(2^\Pi_1\) resonant peak diminishes with increasing \(n\). This is consistent with the excitation process near threshold.
The difference in the height of the threshold peaks at an angle of $135^\circ$.

Figure 7. Excitation of different harmonics of the bending vibration at an angle of $135^\circ$.

having a greater autodetachment width than the $^2\Pi$ resonance.

For every pair $(0, n, 0)$ and $(0, n + 1, 0)$, left and right in figure 7, the threshold peak is higher on the right, for the excitation of even number of bending quanta.

The difference in the height of the threshold peaks between even and odd $n$ is largest for low $n$. The threshold peak is entirely absent for $(0 \ 1 \ 0)$, and completely dominates the spectrum for $(0 \ 2 \ 0)$. The difference is smaller but still very clear between $(0 \ 7 \ 0)$ and $(0 \ 8 \ 0)$.

Another way of visualizing the relative intensities of the threshold peaks in vibrational excitation is by means of an energy-loss spectrum recorded with a low constant residual energy, such as the spectrum recorded at $E_r = 0.05$ eV shown in figure 8. (A very similar spectrum, recorded at $90^\circ$ (for energies up to 2 eV with $E_r \approx 0$ eV), was presented by Abouaf et al [23].) The alternation of the intensities of the $(0 \ n \ 0)$ peaks, with even $n$ being consistently higher than odd $n$, is clearly visible. Note that the vertical scale is logarithmic and the enhancement of even $n$ over odd $n$ is generally more than a factor of 2.

Schulz and Wong [45] observed selectivity in vibrational excitation of a polyatomic molecule (benzene) and derived a selection rule to explain it. (A further discussion was given by Gallup [46, 47] and by Ben Arfa and Tronc [48].) They proposed that the vibrational state must have the same symmetry as the force field exciting it, that is, for our case of a $^2\Pi$ resonance, $\Pi \otimes \Pi = \Sigma (+\Delta)$. This selection rule would thus not allow the excitation of odd quanta of the bending vibration, having $\pi$ symmetry (neglecting angular momenta higher than 2 throughout). Schulz and Wong observed more vibrations than predicted by this selection rule, however. They were able to explain the additional vibrations by assuming that the electron may change angular momentum upon exciting a vibration, in particular that it can leave in an $s$-wave. In our case of a $^2\Pi$ resonance this adds vibrations of the $\Pi \otimes \Sigma = \Pi$ symmetry, i.e., the excitation of odd quanta also becomes allowed. The extended selection rule appears applicable to OCS because no clear preference for even quanta of bending is apparent in the $^2\Pi$ resonance region, both in figure 3 and in figure 7. (There is at least one case, however, where a $^2\Pi$ resonance leads to preferential excitation of even quanta of a bending vibration—that of diacetylene [49].) Abouaf et al [23] have shown both experimentally and theoretically that the angular distributions of the excitation of the odd and even quanta are quite different, however.

In the near-threshold region, dominated by the $^2\Sigma$ virtual state, the above rules predict that only $\sigma$ vibrational states will be excited, i.e., only even quanta of the bending vibration (and, of course, the CS and CO stretch vibrations). The selection rule thus explains the observations.

In view of these results it is interesting to pose the question of whether this selectivity is also manifest in the remaining two members of the present series, CO$_2$ and CS$_2$. The excitation of the $(0, 1, 0)$ state in CO$_2$ gives rise to the strongest vibrational peak in a spectrum recorded near threshold [8, 12], in an apparent violation of the symmetry rule, but it has been argued that the observed cross section (at $54^\circ$) can be entirely attributed to direct dipole excitation [8]. The excitation of the next odd number of quanta, $(0, 3, 0)$, is also (weakly) observed in CO$_2$, although the situation is complicated by the fact that it is mixed with the $(1, 1, 0)$ state by a Fermi resonance [12]. The problem has also been addressed by Sommerfeld et al [11]. It is thus difficult to decide whether the symmetry-induced intensity alternance of bending excitation occurs in CO$_2$. Vibrational states with odd quanta of bending vibration are readily excited in CS$_2$ near the threshold [19], but this is less surprising, because even the vertical electron affinity of CS$_2$ is positive and the $^2\Pi_u$ state probably dominates VE even at the threshold.

Figure 8. Threshold excitation spectrum at an angle of $135^\circ$. 
The last question which this study addresses is that of the possible existence of sharp structures in the cross sections at low energies. As already mentioned in the introduction, such structures could be expected, based on several arguments:

(i) Sharp structures were observed in OCS clusters [4], less pronounced but otherwise similar to those found in CO₂ clusters.

(ii) Sharp structures were observed for isolated CO₂ molecules, in the excitation of the topmost members of Fermi polyads containing overtones (at least two quanta) of the symmetric stretch vibrations [12].

(iii) The structures in CO₂ are believed to be linked to a weak binding of an electron by a combination of dipole and polarization forces in the bend geometry, which causes the potential surface of the anion to bend down before crossing the potential of the neutral molecule [4, 10, 11]. This binding may be expected to be even stronger in OCS because of its larger polarizability and permanent dipole moment.

None of the curves in figures 4–7 exhibits structures near higher excited thresholds, however. Note that the third overtone of the CS stretch vibration, (300), overlaps with the fifth overtone of bending, (0 50), shown in figure 7, and any structure in the (300) cross section would be visible there. The second overtone of the CO stretch vibration, (0 02), is close to the (0 80) vibration, also shown in figure 7. No structures are thus observed, not even in the overtones of any of the three vibrational modes.

With this negative result in mind, we note that, according to the calculations [4], the potential surface of OCS− bends down, but less than that of CO₂, despite the larger polarizability of OCS. This result could explain the absence of structure in the cross sections. The fact that the structures in the OCS clusters are less pronounced than those in CO₂ clusters points in the same direction. Finally, we note that CO₂ and OCS differ in two other aspects:

(i) The structures in CO₂ appear only in the vibrations containing the symmetric stretch, they are absent in the antisymmetric stretch. This symmetry distinction is missing in OCS.

(ii) Both stretch and bending are important for the stabilization of the CO₂ − anion, and it is thus important that in the vibrational states for which structure appears in the cross sections, the stretch and bending motions are mixed, in phase, by strong Fermi resonances in CO₂ [13]. This aspect is missing in OCS where Fermi mixing is weaker.

4. Conclusions

We report absolute cross sections for elastic and vibrationally inelastic electron scattering from OCS molecules, as measured with high energy resolution (13 meV) at the selected angle $\theta = 135^\circ$ over the energy $0.06–20$ eV. In the range of overlap, satisfactory to good agreement is obtained with the previous experimental results of Sohn et al [29], Abouaf et al [23] and Tanaka and Hoshino [36] and the theoretical cross sections of Michelin et al [37], Bettega et al [27, 31] and Gianturco and Stoecklin [28].

More specifically, the present measurement of the elastic cross section confirms earlier measurements between 0.8 and 3 eV, but is larger between 3 and 5 eV and lower below 0.8 eV. Very good agreement is found with the static-exchange plus polarization approximation results of Bettega et al [31] between 3 and 8 eV and the calculation including correlation–polarization effects of Gianturco and Stoecklin [28] at energies of 1.5–20 eV. The present data are lower than the latter calculation below 1.5 eV, however.

The magnitudes of the present VE cross sections are in satisfactory agreement with the data of Sohn et al [29]. The 3.7 eV resonance excites selectively the CS stretch vibration, but not the CO stretch and the bending vibrations, confirming its assignment as $\sigma^*_{CS}$. The 1.33 eV resonance excites all three modes, consistent with a temporary $\pi^+$ occupation with a bent equilibrium geometry. A long progression of bending vibration with a strong preference for even quanta is excited near the threshold, consistent with a $^2\Sigma$ virtual state mechanism, which prefers $\sigma$ final vibrational states over $\pi$ states.

Despite thorough search, including overtone vibrations, no sharp resonant structures were found in the VE cross section near higher vibrational thresholds. This is somewhat surprising, because sharp VFRs were found in OCS clusters and because the potential energy surfaces of OCS−/OCS resemble those of CO₂−/CO₂ [4], and because sharp VFR were found in CO₂, both for isolated molecules [12] and in clusters [21].

Acknowledgments

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