Studies of Unoccupied Orbitals of BF₃ and BCl₃ by Electron Transmission Spectroscopy and Multiple Scattering Xα Calculations

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Abstract
Electron transmission spectroscopy (ETS) and bound-state and continuum multiple-scattering Xα (Ms-Xα) calculations are employed to characterize the unoccupied a₂', a₁' and e' orbitals of BF₃ and BCl₃. The a₂' orbital of BF₃, which produces a peak about 7 eV below threshold in the x-ray absorption spectrum (XAS), generates a scattering resonance at 3.5 eV in ETS. Similarly, the e' orbital that lies about 2 eV above threshold in XAS occurs about 13–16 eV above threshold in ETS. Dissociation of F⁻ from BF₃ due to electron attachment is attributed to a core-excited shape resonance involving an e' → a₂' excitation and electron capture into the a₂' orbital. In BCl₃ all the unoccupied orbitals lie at lower energy than in BF₃ and are closely spaced, making definitive spectral assignments difficult. Both Hartree-Fock (HF) and Ms-Xα methods apparently underestimate the stability of the unoccupied e' orbital of BCl₃. Vibronic coupling due to out-of-plane bending may significantly affect the spectral intensities. Feshbach resonances are observed for BCl₃ at energies close to those observed in the vacuum-UV absorption spectrum. The lower energies of the unoccupied orbitals of BCl₃ are consistent with their stronger bonding to nucleophiles.

Introduction
It is well known that the boron trihalides are strong Lewis acids [1]. However, for BF₃ the adiabatic electron affinity is negative [2], and it is known that the adiabatic electron affinity is considerably more positive than the vertical electron affinity due to large, stabilizing, out-of-plane geometric distortion in the anion [3]. Only orbitals with negative vertical electron affinities are observable by electron transmission spectroscopy (ETS). Although there has been no previous direct observation of electron capture into unoccupied orbitals of the boron halides, there are both experimental measurements and theoretical calculations that relate to the energies and other properties of these orbitals. For example, a resonance presumably associated with temporary anion formation has been observed in the electron impact vibrational excitation of BF₃ [4]. Furthermore, some information is available on UV and x-ray absorption [5,6] that involves electron excitation into the unoccupied orbitals. Theoretical studies using HF-based Stieltjes imaging [6] and equivalent ionic-core virtual orbital [6] methods, as well as multiple-scattering (MS) Xα techniques [7] have been employed in the interpretation of the x-ray absorption spectra. In the present work we have used
ETS to determine the energies of temporary negative-ion states in BF$_3$ and BCl$_3$ that arise from electron capture into low-lying unfilled molecular orbitals. In addition, we have undertaken MS-X$\alpha$ calculations, using the Slater transition-state concept [8] to relate the electron transmission, x-ray, and UV absorption results.

**Experimental Method**

The electron transmission experiment involves the measurement of the transparency of a gas to an electron beam as a function of energy. The transparency depends in an inverse fashion upon the electron-scattering cross section. Temporary negative-ion formation occurs with large cross section only over a narrow energy range. Treated as a scattering process, such a temporary ion appears as a phase change of $\pi$ in the scattering electron wave function and as a maximum in the electron-scattering cross section. We shall refer to this process, as well as the corresponding feature in the transmission-vs.-energy spectrum, as a "resonance."

The electron spectrometer consists of an electron source followed by an electron monochromator, a gas cell, and an electron collector [9]. In practice, the first derivative of the transmitted current as a function of energy is recorded, since the derivative is sensitive to the abrupt change in transmitted current associated with a resonance [10]. The energy associated with a resonance is known as an "attachment energy" (AE) and, with respect to the derivative spectrum, is defined as the point vertically midway between the minimum and maximum that characterize the resonance. For the present purposes an attachment energy may be identified with the negative of the corresponding vertical electron affinity [11]. The chief limitation is that ETS gives only the energy associated with unstable negative ions. That is, only negative electron affinities can be obtained with ETS. A time-of-flight mass spectrometer has recently been appended to the ETS apparatus to monitor anions from dissociative attachment. Thus, for example, we observe anions from the process $e^- + BX_3 \rightarrow BX_2 + X^-.$

**Computational Method**

The continuum MS-X$\alpha$ method has been previously described [12, 13]. We have applied this method using transition state potentials to the series (C, Si)X$_4$, X = H, F, Cl [14]. In the present work we use D$_{3h}$ symmetry for BF$_3$ and BCl$_3$ with experimental bond distances of 1.307 Å and 1.742 Å, respectively [15]. Atomic sphere radii were chosen according to the prescription of Norman [16] with approximately 20% overlap [17]. Alpha values were obtained in the usual way by using the tabulations of Schwarz [18]. For the calculations of elastic electron-scattering cross-section, experimental dipole polarizabilities [19] were used, and the polarization potential was assumed to have the functional form $-\alpha/r^4$. All calculations were carried out in a spin-restricted formalism. Maximum azimuthal quantum numbers were 4 on the outer sphere and on the central atom, and 2 on the halogens.

**Results**

The ground state MS-X$\alpha$ orbital energy diagram of BCl$_3$ is shown in Figure 1. The 1a$_1'$ and 1e' orbitals are predominantly Cl3s in character, the 2a$_1'$ and 2e' are B2s-Cl3p...
and B3p-Cl3p σ bonding, the orbitals from 1a'' to the HOMO 1a'_1 are predominantly Cl3p nonbonding, the 2a'_1 LUMO is B2p-Cl3p π*, and the 3a'_1 and 4e' are B2p-Cl3p σ* orbitals. The BF₃ eigenvalue spectrum is similar, except that all the empty a'_1 orbitals are higher in energy and more diffuse than for BCl₃. We shall employ the same orbital numbering scheme for BF₃. Based on this simple orbital model, we would expect that transient negative ions could arise by capture of electrons into the 2a'_2, 3a'_1, or 4e' orbitals, although the stability of the 2a'_2 in BCl₃ is great enough that anions formed by its occupation could be quite close to threshold. In Figure 2 we present the derivative ETS spectrum for BF₃ and BCl₃. Figure 3 is a correlation diagram showing the attachment energies for the two molecules and giving the most probable orbital assignments. Continuum ms-Xα elastic scattering cross sections for BF₃ and BCl₃ are shown in Figure 4. For both molecules the potential used in the continuum ms-Xα calculation was obtained self-consistently with 0.5 electrons in the 2a'_2 orbital. Previous experience suggests that cross sections near threshold in the totally symmetric channel are exaggerated by the method [20] so that the threshold behavior shown in Figure 4 is certainly suspect.

The electron scattering between 0.1 and 0.8 eV in BF₃ is due to vibrational excitation of neutral BF₃. An expanded spectrum of this region is shown in Figure 5. This process has been observed previously for CF₄ [21]. The spacing between the vibrational features alternates between two values of approximately 0.08 eV and 0.10 eV. Four modes of vibrational excitation are possible for neutral BF₃ [22]. The vibrational energies of the different modes are ΔE(ν₁) = 0.110 eV, ΔE(ν₂) = 0.0085 eV, ΔE(ν₃) = 0.180 eV, and ΔE(ν₄) = 0.060 eV. The spacing of the vibrational features in Figure 5 thus indicates that ν₁ (the asymmetric stretch mode) and ν₂ (the umbrella mode) are excited. The first feature near 0.18 eV corresponds to ν₃ = 1 and ν₂ = 0.
Figure 2. Derivative electron transmission spectra of BF$_3$ and BCl$_3$. The spike at 0 eV in each spectrum is simply the derivative of the abrupt turning on of current at threshold. The spike in the BF$_3$ spectrum is characteristic of the resolution of the experiment. The broadening of the spike for BCl$_3$ implies an attachment process at threshold. For BF$_3$, the insert shows an expanded version of the region from 10–16 eV, showing a weak but reproducible resonance.

Figure 3. Correlation diagram giving the attachment energies (AE) for BF$_3$ and BCl$_3$. Probable assignments for the orbitals responsible for the observed anion states are also given.
Figure 4. Continuum ms-Xα elastic electron-scattering cross sections for BF₃ and BCl₃:
total, –– a¹¹ symmetry, ···· a¹ symmetry, -- e symmetry.

Figure 5. Derivative electron transmission spectrum of BF₃ expanded to show vibrational
excitation near threshold.
The second feature then corresponds to \( \nu_3 = 1 \) and \( \nu_2 = 1 \), and the next to \( \nu_3 = 2 \) and \( \nu_2 = 0 \). This pattern continues along the spectrum.

The feature at 3.54 eV in the spectrum of BF\(_3\) corresponds fairly closely to the 3.8-eV peak observed by inelastic scattering [4], and the broad feature around 16.9 eV is not far from the highest-energy inelastic electron-scattering peak at 13.7 eV [4]. Although the 16-eV feature is broad and weak, it is reproducible over the course of months and has a typical resonance shape, which is clearly shown in the insert in Figure 2. By contrast, the weak features from 6–14 eV are not reproducible. The 3.8- and 13.7-eV inelastic scattering peaks have been previously assigned [4] to the 2a\(_g^\circ\) and 4e' orbitals by drawing a correspondence of relative electron scattering and x-ray absorption energies and by reference to previous MS-X\(\alpha\) calculations [7]. The present continuum MS-X\(\alpha\) cross-section calculations (Fig. 4) certainly support this interpretation for the case of the a\(_g^\circ\) resonance, although the calculated resonance energy is considerably lower than experiment. Calculated x-ray term energies and oscillator strengths calculated by using the method of Noodleman [23] are given for BF\(_3\) and BCl\(_3\) in Table I. The 2a\(_g^\circ\) orbitals are calculated to be too stable by 1.3 to 1.6 eV. The present results thus give less accurate energies than the Hartree–Fock approaches of Ref. 6 (which give term energy errors of only a few tenths of an electronvolt), although they are more accurate than previous MS-X\(\alpha\) results [7] that employed only the ground-state potential with the objective of obtaining a comprehensive picture of the near threshold continuum structure. If we correct the calculated a\(_g^\circ\) resonance energy of 1.9 eV in BF\(_3\) by the difference of 1.1 eV between calculated and experimental x-ray term energies, we project the a\(_g^\circ\) resonance to be about 3.0 eV, in reasonable agreement with experiment. Assuming that the 4e' XAS term energy and the e' ETS resonance energy differ by the same amount as for the 2a\(_g^\circ\) (\(\sim 10.8\) eV) leads to a projection of 13.1 eV for the e' resonance, in reasonable agreement with Ref. 4 and the present data. Our MS-X\(\alpha\) cross-section calculations also give a broad maximum at 13.8 eV with a magnitude of 16.0 Å [2]. Although a number of channels contribute to \(\sigma\) in this energy region, only the e' shows a maximum, and it contributes more than 50% of the total, so it is reasonable to call this an e' resonance.

Dissociative attachment to BF\(_3\) gives rise to only one peak in the mass spectrum. As previously observed [24], capture of electrons of about 11.3 eV energy leads to

<table>
<thead>
<tr>
<th>(2a_g^\circ)</th>
<th>(3\alpha_1)</th>
<th>(4e')</th>
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<tbody>
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<td>8.4 (.105)</td>
<td>5.1 (forbidden)</td>
<td>-1.8 (.081)</td>
</tr>
<tr>
<td>7.3 (.082)</td>
<td>4.6 (.004)</td>
<td>-2.3 (.195)</td>
</tr>
<tr>
<td>9.0 (.100)</td>
<td>6.0 (forbidden)</td>
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</tr>
<tr>
<td>7.4 (.022)</td>
<td>3.7 (.0036)</td>
<td>2.9 (.043)</td>
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Ref. 6.
the production of F\textsuperscript{−}. This peak corresponds to the second peak observed by Tronc et al. [4] in the vibrational excitation spectrum of BF\textsubscript{3}. This ion peak and the structure in the vibrational excitation spectrum are not well understood because no corresponding feature appears in the electron transmission or photoabsorption spectrum. Possibly this structure is the result of a weak, repulsive resonance or core-excited shape resonance. The lowest energy resonance of this type is expected to correspond to the configuration (1e\textsuperscript{−})\textsuperscript{−}(2a\textsubscript{2g})\textsuperscript{2} arising from a dipole allowed excitation 1e\textsuperscript{−} → 2a\textsubscript{2g} and the capture of an electron into 2a\textsubscript{2g}. Our MS-Xα calculations give 12.3 eV for the resonance and 11.6 eV for the 1e\textsuperscript{−} → 2a\textsubscript{2g} excitation energy, leading to classification of the resonance as a core-excited shape resonance. Unfortunately, the only data on the vacuum-UV absorption spectrum of BF\textsubscript{3} indicate that there are no singlet absorption bands below 10.5 eV [25], although a triplet absorption may occur at about 8 eV [26]. The 11.3-eV feature is thus energetically consistent with a core-excited shape resonance of the type suggested, but we presently have no way to calculate the intensity of such a feature and no direct experimental correspondence with the UV spectrum.

The ETS spectrum of BCl\textsubscript{3} is considerably more difficult to assign. Although the x-ray term energies of the 2a\textsubscript{2g} orbitals of BF\textsubscript{3} and BCl\textsubscript{3} are very similar, direct calculation of the vertical AE's using the transition-state approach gives values of 1.6 eV for BF\textsubscript{3} and about 0.0 eV for BCl\textsubscript{3}. This substantial difference arises from the larger size of BCl\textsubscript{3}, which reduces the magnitude of the 2a\textsubscript{2g} orbital self-repulsion. If we assume that the underestimation of the 2a\textsubscript{2g} resonance energy in BCl\textsubscript{3} is similar to that in BF\textsubscript{3}, we would project an 2a\textsubscript{2g} resonance at about 1.9 eV, but the lowest energy ETS peak occurs at 2.9 eV. The calculated cross sections give only a very weak a\textsubscript{2g} maximum at low energy. The calculations do, however, indicate maxima in the a\textsubscript{1g} and e\textsuperscript{′} channels. The a\textsubscript{1g} maximum is somewhat unexpected since no such feature occurs in BF\textsubscript{3}, but a transition-state calculation of the BCl\textsubscript{3} 3a\textsubscript{1g} vertical AE indeed gives a value of 1.2 eV and a localization of about 15% of the electron density within the B atomic sphere. The e\textsuperscript{′} maximum near 4.0 eV is at lower energy than would be expected from the calculated XAS energies in Table I. Although the total cross section is smoothly decreasing from 5.0 to 9.0 eV, above 9.0 eV there is a very slow increase in cross section, leading to a flat maximum at 10.8 eV, with the main contribution being from the e\textsuperscript{′} channel. The complexity of the calculated cross section and the lack of quantitative correspondence with experiment make interpretation difficult.

Similar difficulties appear in the interpretation of the x-ray [6] and UV [25] absorption spectra of BCl\textsubscript{3}. Although both MS-Xα (Table I) and HF methods give reasonable Bi\textsubscript{s} XAS term energies and intensities for BF\textsubscript{3}, both methods have problems with the apparent 4e\textsuperscript{′} feature in the spectrum of BCl\textsubscript{3}, giving term energies and relative oscillator strengths that are too small. These discrepancies may arise either from a poor description of the 4e\textsuperscript{′} orbital or from vibronically allowed intensity to the 3a\textsubscript{1g} orbital, which is quite localized in BCl\textsubscript{3} as we have indicated. It has been argued that vibrationally excited BX\textsubscript{3} molecules will be nonplanar with C\textsubscript{3v} symmetry, so that the Bi\textsubscript{s}(a\textsubscript{1g}) → 3a\textsubscript{1g} transition is allowed [6]. Since the frequency of the out-of-plane a\textsubscript{2g} vibration of BCl\textsubscript{3} is one-third smaller than that of BF\textsubscript{3} [27], we expect such vibronic coupling effects to be larger in BCl\textsubscript{3}.
In the same way the MS-Xα calculations give intensities that are too small for the features attributed to 4e' final states in the vacuum-UV spectrum. The spectrum of Planckaert et al. [25] shows maxima at 7.2 and 9.0 eV and a shoulder at 8.3–8.6 eV. A sharp rise in absorption also occurs around 10.5 eV, suggesting another maximum beyond the limit of the study. MS-Xα calculations give two intense transitions: 1e'' → 2a'' at 6.3 eV, with \( f = 0.195 \), and 2a' → 2a'' at 11.1 eV, \( f = 0.154 \). All the intermediate energy transitions allowed in D₃h symmetry that involve 3a' and 4e' final states have oscillator strengths below 0.01, although their energies fall in the right range. It is conceivable that vibronic effects may also contribute to the intensity in this spectral region.

Our analysis of the UV spectrum of BCl₃ does suggest an interpretation of the features observed in the ETS at 7.6 and 9.05 eV. The closeness of these energies to the UV absorption energies of 7.2 and 9.0 eV suggests that they are core-excited shape resonances. MS-Xα calculations give 6.4 eV for the energy of the \( (1e'')^{-1}(2a'') \) state, but this energy may be underestimated just as the x-ray term energy of the 2a'' orbital is underestimated.

It should be noted that the present MS-Xα calculations give values for orbital ionization potentials accurate to within about 1 eV for the 2a''-1a' orbitals, similar to that obtained from previous studies [28]. The errors in the description of the 2a''-4e' orbitals are apparently somewhat larger, perhaps due to their larger concentration of electron density in the interatomic region. In addition, ETS resonance energies are very small, so that modest absolute errors are large percentage errors.

**Conclusion**

The electron transmission spectrum of BF₃ shows resonances corresponding to the 2a'' and 4e' orbitals previously studied by XAS. In addition, a peak in the F⁻ dissociative attachment spectrum corresponds to a core-excited shape resonance probably involving a \( (1e'')^{-1}(2a'') \) configuration. For BCl₃, lower resonance energies are obtained for the 2a'', 3a', and 4e' orbitals, and the ETS shows two low-energy resonances and evidence for an electron capture process at threshold. Both HF and MS-Xα methods appear to underestimate the stability and degree of localization of the 4e' orbital of BCl₃, and this underestimation appears in both the x-ray and UV regions of the absorption spectra.

Previous MS-Xα studies have shown a strong dependence of photoionization intensity and electron-scattering cross section upon detailed geometry from some molecules, e.g., CO₂ [29] and CF₄ [14], although comparison with HF results for CO₂ [30] suggests that the MS-Xα distance effects may be exaggerated. The small energy difference of the resonant channels in BCl₃ and its low vibrational frequencies suggest the need for further studies at different geometries. We must also systematically explore the use of different transition-state potentials, i.e., those generated by the half-occupation of orbitals other than the 2a''.

The increase in BX₃-donor interaction from BF₃ to BCl₃, seen e.g., in the B—N bond lengths and stretching force constants of X₃B—NCCH₃ adducts [31], may be partially related to the greater stability of the 2a'' orbital in BCl₃. Within the context
of the perturbation theory of chemical reactivity [32] a stabilization of the 2a_1^2 orbital should increase the strength of its stabilizing interaction with nucleophiles. It also appears that for BCl_3 the 3a_1' orbital and perhaps the 4e' are also low enough in energy to accept electron density from a nucleophile. If the 4e' is active in this respect, one would expect that its relative heat-of-adduct formation would be larger for π-donor adducts, such as CO, that have the proper symmetry to interact with the 4e'. Studies on the BH_3—CO and BF_3—CO donor acceptor complexes using Morokuma's energy decomposition analysis [33] indicate a larger charge transfer component to the bonding in BF_3—CO, due to a lower energy for the empty a_2^" orbital. A similar analysis of BCl_3—CO would be interesting, particularly if a separation of the charge transfer contributions to the 2a_1^2, 3a_1', and 4e' orbitals could be made.

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Bibliography


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