Electronic Properties and Dissociative Photoionization of Thiocyanates, Part III. The Effect of the Group’s Electronegativity in the Valence and Shallow-Core (Sulfur and Chlorine 2p) Regions of CCl₃SCN and CCl₂FSCN

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ABSTRACT: Both photoelectron spectroscopy (PES) data and PhotoElectron-PhotoIon-Coincidence (PEPICO) spectra obtained from a synchrotron facility have been used to examine the electronic structure and the dissociative ionization of halomethyl thiocyanates in the valence and shallow-core S 2p and Cl 2p regions. Two simple and closely related molecules, namely, CCl₃SCN and CCl₂FSCN, have been analyzed to assess the role of halogen substitution in the electronic properties of thiocyanates. The assignment of the He(I) photoelectron spectra has been achieved with the help of quantum chemical calculations at the outer-valence Green’s function (OVGF) level of approximation. The first ionization energies observed at 10.55 and 10.78 eV for CCl₃SCN and CCl₂FSCN, respectively, are assigned to ionization processes from the sulfur lone pair orbital [n(S)]. When these molecules are compared with CX₃SCN (X = H, Cl, F) species, a linear relationship between the vertical first ionization energy and electronegativity of CX₃ group is observed. Irradiation of CCl₃SCN and CCl₂FSCN with photons in the valence energy regions leads to the formation of CCl₂X⁺ and CClXSCN⁺ ions (X = Cl or F). Additionally, the achievement of the fragmentation patterns and the total ion yield spectra obtained from the PEPICO data in the S 2p and Cl 2p regions and several dissociation channels can be inferred for the core-excited species by using the triple coincidence PEPIPICO (PhotoElectron-PhotoIon-PhotoIon-Coincidence) spectra.

1. INTRODUCTION

The interest of haloalkyl thiocyanates is very much extended in the organic chemistry. They are used as key intermediates to produce important biocides. In particular, the species chloromethyl thiocyanate, CH₂ClSCN, shows a wide activity as fungicide, nematocide, and bactericide. The production of the fungicide 2-(thiocyanomethylthio)-benzothiazole, used in the leather industry, uses also this intermediate. It is known that CCl₃SCN displays biological activity as an agent to control insect pests.

Taking into account the singular importance of these kinds of molecules, few spectroscopic data have been reported. The infrared and Raman spectra of both CCl₃SCN and CCl₂FSCN molecules have been interpreted in terms of C₃ and C₁ molecular symmetry, respectively. Recently, the title molecules have been structurally studied in both gas and crystal phases by means of gas electron diffraction (GED) and low-temperature single-crystal X-ray diffraction (XRD). The experimental results concluded that the crystal structure of CCl₃FSCN contains solely the gauche-conformer (FC–SC dihedral angle near to 60°) while in the gas phase both gauche- and anti-conformations (FC–SC dihedral angle near to 180°) are present in equilibrium at room temperature, with the gauche-conformer being the most stable form. On the other hand, the crystal and gas phase conformations of the CCl₁SCN molecule show that the CCl₃ group results staggered with respect to the SCN group.

Electronic properties for the simple CH₃SCN derivative were previously studied by Hitchcock et al. More recently, our group has investigated the ionic dissociation of CH₃SCN and
CH₂ClSCN molecules induced by photon impact excitation at the S 2p and Cl 2p levels and the dissociation dynamics was determined by using synchrotron radiation and multicoincidence techniques. In both species, the HOMO corresponds to n₆(S) lone pair electrons and the fragmentation dynamics is dominated by the rupture of the C=S bond.

Herein, and as part of a general project aimed to elucidate the electronic properties of thiocyanates compounds by using synchrotron radiation in combination with photoelectron spectroscopy, we present a study of two closely related molecules, i.e., CCl₃SCN and CCl₂FSCN, in the 11.2 and 300.0 eV photon energy range. Outermost valence electrons as well as S 2p and Cl 2p shallow-core electrons were excited, and the positive ions formed after ionization were detected by means of coincidence techniques.

2. EXPERIMENTAL SECTION

As already described earlier, we have used the facilities kindly offered at the Laboratório Nacional de Luz Síncrona (LNLS), Campinas, São Paulo, Brazil, to use synchrotron radiation.  The toroidal grating monochromator TGM, which is operative in the range between 11.2–300 eV, offers linearly polarized light interacting with the sample inside a high-vacuum chamber kept with a pressure of ca. 10⁻⁶ mbar. The fwhm of the light beam was determined to be 1 mm. The pressure of the sample was maintained below 5×10⁻⁶ mbar during the measurements. A resolution power better than 400 can be achieved in the TGM beamline under these conditions. The well-known S 2p→6a₁g and S 2p→2t₂g signals of SF₆ were used to calibrate the energies.

The use of a gas-phase harmonic filter in the 11.2–21.5 eV region allowed the provision of high-purity vacuum-ultraviolet photons due to the removal of contamination by high-order harmonics. The intensity of the emergent beam was recorded with a light-sensitive diode. The ions produced when the synchrotron light interacts with the sample were detected using a time-of-flight (TOF) mass spectrometer of the Wiley–McLaren type for both PhotoElectron-PhotoIon-Coincidence (PEPICO) and PhotoElectron-PhotoIon-PhotoIon-Coincidence (PEPIPICO) experiments. The characteristics and performance of this electron–ion coincidence TOF spectrometer have been already reported.

The average kinetic energy release (KER) values of the fragments have been deduced from the coincidence spectra by assuming both an isotropic distribution of the fragments, that they are perfectly space focused and that the electric field applied in the extraction region is uniform. Thus, the peak width contains the information to determine the energy release in the fragmentation processes. When the ideal conditions are not fully obtained an increase of the peak width is observed. A peak width of 0.05 eV was obtained for the Ar⁺ ion during the measurements of the argon mass spectrum under very similar experimental conditions. Thus, this value constitutes a reliable estimation for the instrumental resolution since this broadening in Ar⁺ can be originated from thermal energy and instrumental broadening.

The experimental conditions to determine the HeI PE spectra of CCl₃SCN and CCl₂FSCN have been already reported. A resolution of about 30 meV can be deduced as indicated by the Ar⁺(5P₁/₂) photoelectron band. The calibration of the vertical ionization energies was made by addition of small portions of argon and iodomethane to the sample.

OVGF calculations using the cc-pVTZ basis set and MP2/cc-pVTZ optimized geometries of CCl₃SCN and CCl₂FSCN have been computed using the Gaussian 03 program suit. The samples of CCl₃SCN and CCl₂FSCN were obtained from the following reactions.

Chlorine monoxide AgCN → CCl₃SCN + AgCl

CCl₂FSCN:

CCl₂SCI + HF → CCl₂FSCI + HCl

The liquid samples were purified by several trap-to-trap vacuum distillations. The purity of the compounds was measured by FTIR spectroscopy.

3. RESULTS AND DISCUSSION

3.1. Photoelectron Spectra. The HeI photoelectron (PE) spectra of CCl₃SCN and CCl₂FSCN are shown in Figure 1. The experimental and computed (OVGF/cc-pVTZ) vertical ionization energies can be shown in Table 1.

The molecular conformation of a molecule results decisive to derive its ionization energies. The CCl₃SCN molecule shows only one stable form in the gas phase, the gauche conformer. CCl₂FSCN presents both gauche- and anti-conformations in equilibrium at ambient temperature, with the gauche-rotamer being the most stable. Thus, ionization values have been computed for both conformers, as shown in Table 1. Similar ionization values were computed for both forms being the same the characters deduced for the molecular orbitals. The characters of the highest occupied molecular orbitals for CCl₃SCN and for the most stable rotamer of CCl₂FSCN are shown in Figure 2. In the electronic ground state, the CCl₃SCN species belongs to the C₃ point group of

Figure 1. HeI photoelectron spectra of CCl₃SCN (a) and CCl₂FSCN (b).
The Journal of Physical Chemistry A

### Table 1. Experimental and Computed Ionization Energies (OVGF/cc-pVTZ, in eV) and MO Characters for CCl₃SCN and CCl₂FSCN Molecules

<table>
<thead>
<tr>
<th></th>
<th>CCl₃SCN</th>
<th>CCl₂FSCN</th>
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<tbody>
<tr>
<td></td>
<td>exptl</td>
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</table>

![Character of the main occupied valence orbitals of the CCl₃SCN and CCl₂FSCN molecules](image)

**Figure 2.** Characters of the main occupied valence orbitals of the CCl₃SCN and CCl₂FSCN molecules.

3.2. Correlation between Group Electronegativity and Ionization Energy. When the photoelectron spectra of the title molecules are compared with related species, such as CH₃SCN and CH₂ClSCN, a dependency of the vertical ionization energy with the electronegativity of the −CX₃ (X = H, Cl, F) group is observed. The concept of electronegativity is well established in chemistry. It refers to the capacity of an atom (or functional group) to attract electrons to itself. Linus Pauling defined this concept as “the power of an atom in a molecule to attract electrons to itself.” Then, the extension of this concept to groups of atoms gave rise to group electronegativity, in which a functional group can be treated as a pseudatom. Hardness represents another related concept. It has been defined to be half the derivative of electronegativity with respect to charge. Hardness is a property inversely related to polarizability. A charge distribution in a molecule (the electronegativity of its component atoms) and its polarizability (the hardness of the component atoms) govern the inner-shell ionization energies. Therefore, the correlation between the core-ionization energy of a central atom and the electronegativity sum of the substituents attached to the atom can be expected. Thomas et al. have shown that both electronegativity and hardness are very much related with the ionization energies in a previous report over halomethanes.

The family of the molecules studied in this work, the halomethyl thiocyanates, are also good candidates for this determination. Whereas in this series the molecules present basically similar structures and bonding, the electronegativities and polarizabilities of the −CX₃ (X = H, Cl, F) group, however, vary over a wide range depending on the kind and number of halogen atoms which are present in the molecule. These changes in the electronegativity of group attached to the sulfur atom affect directly to the firsts ionization energy of these compounds, since it is associated with the ionization process starting at the π lone pair orbital localized on the S atom.

Figure 3 shows the experimental and theoretical vertical ionization energies of CX₃SCN species as a function of the electronegative of the CX₃ group, with X = H, F, and Cl. Taking the methyl derivative, i.e. CH₃SCN, as a reference, it is observe that the substitution of the hydrogen by fluorine and chlorine atoms cause an increment of the electronegative of this...
group in agreement with the electron-withdrawing property of halogens, leading to higher first ionization energy values.

A molecular orbital diagram for a series of CCl$_3$SCN molecules is depicted in Figure S1 (Supporting Information). The formal change of the hydrogen atom by more electronegative groups is related to higher energy potential values for the valence electrons. Therefore, the CF$_3$SCN species presents a higher stabilization of both the $n$$_\pi$(S) and $\pi$(CN) orbitals.

### 3.3. Photoionization in the Valence Region

The title compounds were irradiated with synchrotron monochromatic light in the energy region extended between 11.2 and 21.0 eV. Simultaneously, the electrons and cationic fragments produced upon photoionization and eventually subsequent fragmentation events were collected in coincidence, and PEPICO spectra were taken.

The coincidence spectra for both CCl$_3$SCN and CCl$_2$FSCN irradiated in this energy range are shown in the Figure 4. In Table 2, the branching ratios for ion production are also listed. When photons with 11.2 eV are used (the lower energy delivered by the TGM line when experiments were performed) very simple spectra are obtained. This value of energy is higher than the first ionization potential of these molecules (10.55 and 10.78 eV for CCl$_3$SCN and CCl$_2$FSCN, respectively), being possible to observe in these spectra ionization processes starting from the HOMO, corresponding to $n$$_\pi$(S) lone pair electrons formally located on the sulfur atoms.

In both cases, the single charged molecular ion is detected with low intensity and the PEPICO spectra measured are dominated by only two intense signals in all range of energy.

![Figure 4. PEPICO spectra of CCl$_3$SCN (left panel) and CCl$_2$FSCN (right panel) at selected energies into the valence regions.](image)

**Table 2. Branching Ratios (%) for Fragment Ions Extracted from PEPICO Spectra Taken at Photon Energies into the Valence Regions for CCl$_3$SCN and CCl$_2$FSCN Molecules**

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<td></td>
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<tr>
<td>58</td>
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<tr>
<td>124</td>
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<tr>
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*Contributions of the 35/37 natural isotopologues of Cl were summed together.

The main photodissociation channel is produced for the rupture of the sulfur–carbon single bond from CCl$_3$XSCN$^+$ (X = Cl or F) species (see Scheme 1), leading to the formation of CCI$_3^+$ (117 amu/q) and CCl$_3$F$^+$ (101 amu/q), respectively.

The second ionic contribution is the loss of one Cl atom from the single charged molecular ion (M$^+$ − 35), resulting in...
the formation of CClXSCN+ (X = Cl or F) species. These ions can be characterized as derived from the parent ion by rupture of the C−Cl bond, with the charge being retained in the molecular fragment (see Scheme 2). Naturally occurring isotopic contributions are plainly resolved, giving confidence to this assignment.

Scheme 2

When the samples are irradiated with photon energies higher than 14.0 eV, the observation of ionization fragmentation channels for the production of the ions XCS+ (X = Cl or F) and SCN+ from ionized states of CCl3SCN and CCl2FSCN becomes possible. In the case of CCl3SCN, the ionization channel process for the formation of CCl3+ is opened at a photon energy near 16.0 eV.

3.4. Total Ion Yield Spectra (TIY). When the parent and fragment ions are detected with different incident photon energy a complement to the information gained from absorption spectroscopy is reached.45 The TIY spectra of CCl3SCN and CCl2FSCN between 160.0 and 180.0 eV depicted in the Figure 5 were obtained in the region near the S 2p edge by recording the count rates of the total ions.

Figure 5. Total ion yield spectra for CCl3SCN (a) and CCl2FSCN (b) near the S 2p edge.

A group of well-defined features in the region below the S 2p ionization threshold ranged between 164.0 and 169.0 eV has been clearly observed in TIY spectra. These features should be originated from processes involving permitted transitions between the S 2p electron and an antibonding molecular orbital. Following the proposed assignment for S 2p transitions for both CH3SCN12 and CH2CISCN,14 in which the most relevant signals were attributed to states related with (S 2p, σ*SCN) and (S 2p, σ*CS) electronic configurations, the clear observed structures in the TIY spectra can be due to electronic transitions involving the spin–orbit split of the 2p sulfur excited species (2p1/2 and 2p3/2 levels) to unoccupied orbitals, mainly the LUMO (σ*Cl−C) and LUMO+1 (σ*Cl−S) antibonding orbitals, in accordance with the calculated characters for the first unoccupied molecular orbitals displayed in Figure S2 (Supporting Information).

TIY spectra measured for CCl3SCN and CCl2FSCN near the Cl 2p region are depicted in Figure 6. The Cl 2p threshold is positioned in the region of 210.0 eV, in accordance with the values previously reported for similar molecules containing chlorinated groups.13,14,46−52 Centered at about 200.0, 202.0, and 204.0 eV, three wide and not well resolved bands can be observed in this region. Two of these signals could be attributed to electronic transitions related the spin–orbital split of the 2p term in the 2p1/2 and 2p3/2 levels of the excited species toward the LUMO+2 (σ*Cl−C) vacant orbital. The prominence of transitions toward C−Cl antibonding (vacant) orbitals has been already noted for Cl 2p excited organo-chlorinated species.53,54

3.5. PEPICO Spectra. A number of PEPICO spectra have been registered during this study. The photon energies have been selected using the resonant values detected from the TIY spectra taken into consideration the regions corresponding to both S 2p and Cl 2p levels. It is also typical for this kind of study to have measurements of PEPICO spectra below and above the ionization value, generally 10 and 50 eV, respectively. The spectra are achieved by the arrival of only one ion during the period in which the window is open. This process can be due to either a single ionization or multiple ionization processes. In this case, only the lighter and faster ion can be detected. PEPICO spectra for CCl3SCN and CCl2FSCN are depicted in Figure 7, and the corresponding branching ratios for the main fragment ions are listed in Table 3.

In both compounds, the most abundant ion formed is Cl+ (14–20% approximately), reaching relative abundance near 25% at 199.9 eV of energy (Cl 2p edge). Other noticeable ions evidenced in the spectra with relative abundances between 4 and 13% are Cl(C=C)/ClC=F+ (m/z = 117/101), Cl(C=C) (m/z = 82), CICS+/FCS+ (m/z = 79/63), SCN+ (m/z = 58), ClC+/FC+ (m/z = 47/31), CS+ (m/z = 44), S+ (m/z = 32), CN+ (m/z = 26), N+ (m/z = 14), and C+ (m/z = 12). Few changes become apparent when the recorded PEPICO spectra obtained at different photon energies are compared. At higher energies by going from the S 2p to the Cl 2p region an expected atomization process results evident since an increase in the peak intensities corresponding to the lighter ions or fragments C+, N+, S+, Cl+, and SCN+, together with the concomitant decrease in the intensities of the heaviest ions CCl3FSCN+, CCl2+, CClF+, and CCl2+, can be observed.
Using the information deduced from the PEPICO peak contour, the kinetic energy release (KER) value can be inferred for the formed ions at the registered photon energies, as shown in Table 3. The Cl 2p region results are interesting since all ions present high KER values. Experimentally, this fact is associated with broad peaks as observed at the higher energies registered in the Figure 7 for both title compounds. This fact can be explained since the decay of core-shell-excited species usually promotes the formation of doubly charged parent ions in several excited states. Subsequently, they could then dissociate, releasing much of their internal energy as kinetic energy of the fragment ions (KER).

Species such as CClSCN⁺ (m/z = 140) and CClFSCN⁺ (m/z = 124) with relative low KER values can only be derived from the singly charged CCl3SCN and CCl2FSCN molecules, respectively, since these fragments, taken into account the basis of the PEPICO spectroscopy, must be the lightest in the PEPICO spectra. The loss of neutral fragments (chlorine atom) is similar to that proposed in the valence region (Scheme 2). As shown in Figure 8, panels b and d, the peak shapes observed for these ions are clearly symmetric.

\[ \text{CCl}_3\text{SCN}^{++} \rightarrow \text{CCl}_3\text{SCN}^+ + \text{Cl}^- \quad (\text{KER} = 0.17 - 0.22\text{eV}) \]

\[ \text{CCl}_2\text{FSCN}^{++} \rightarrow \text{CClFSCN}^+ + \text{Cl}^- \quad (\text{KER} = 0.19 - 0.21\text{eV}) \]

The same concept can be applied to the formation of CCl⁺ and CClF⁺ fragments (see Figure 8, panels a and c, respectively). The following simple mechanisms explain the experimental observation:

\[ \text{CCl}_3\text{SCN}^{++} \rightarrow \text{CCl}_3^+ + \text{SCN}^- \quad (\text{KER} = 0.18 - 0.68\text{eV}) \]

\[ \text{CCl}_2\text{FSCN}^{++} \rightarrow \text{CCl}_2\text{F}^+ + \text{SCN}^- \quad (\text{KER} = 0.19 - 0.53\text{eV}) \]

Figure 8 depicts also the PEPICO spectra for arriving times corresponding to the SCN⁺ (m/z = 58) ion for both molecules. The SCN⁺ species could be produced from both singly and doubly charged molecular ions. The analysis of the KER distribution is decisive in this case in order to determine the origin of this fragment. From the peak shapes observed in both ranges of energy (S 2p edge and Cl 2p edge), a clear double KER distribution can be observed. At higher photon energies, the formation of double charged ions is favored and broader peaks appear for this ion in the spectra. This behavior suggests that when the thiocyanates are excited at shallow-core S 2p and Cl 2p levels, SCN⁺ ions are originated by both single and double charged molecular ions, according to the equations:

\[ \text{CX}_3\text{SCN}^+ \rightarrow \text{CX}_3^- + \text{SCN}^+ \]

\[ \text{CX}_2\text{SCN}^{++} \rightarrow \text{CX}_2^- + \text{SCN}^+ \]

3.6. PEPICO Spectra. To further determine the ionic dissociation mechanism following the photon absorption, two-dimensional PEPICO spectra have been also recorded. In this method, correlation between one electron and the two
The evaluation of the experimental slopes for coincidence islands in the resonance and off-resonance S 2p photon energy region evidence that the observed differences in the dissociation mechanisms are less significant. The following discussion will be referred to slopes determined from the PEPIPICO spectrum taken at 166.8 and 199.9 eV photon energies for CCl3SCN and CCl2FSCN species, respectively. The contour plots for coincidence islands between selected pair of ions are shown in Figures S3 and S4 in the Supporting Information.

3.6.1. Dynamics of Fragmentation for the CCl3SCN2+ ion. Coincidence between Ions with m/z Values of 58 amu/q (SCN+) and 117 amu/q (CCl2+). This coincidence is observed in the PEPIPICO as a well-defined and intense island with the natural 35Cl/37Cl isotope distribution, as can be observed in Figure S3 (panel A) for the spectrum at 166.8 eV. A simple fragmentation mechanism involving the rupture of the C=S bond is deduced for the experimental coincidence island due to its characteristic parallelogram-like shape and the observed slope of −1.0. A similar behavior has been reported for the CH3SCN and CH2CISCN species.13,14

\[
\text{CCl}_3\text{SCN}^{2+} \rightarrow \text{CCl}_3^{+} + \text{SCN}^{+} \quad \alpha = -1.0
\]

Coincidence between Ions with m/z Values of 58 amu/q (SCN+) and 82 amu/q (CCl2+). The SCN+ and the CCl2+ ions coincide also with a slope of −1.0 (Figure S3B). The formation of both charged species and the neutral Cl atom might be produced by a three-body Coulombic explosion. However, a deferred charge separation mechanism can be postulated due to the observed broadening in the peak shape. This is a three-body reaction involving the loss of a Cl neutral fragment in the first step and the subsequent formation of the double charged intermediate CCl3SCN2+:  

\[
\text{CCl}_3\text{SCN}^{2+} \rightarrow \text{CCl}_3\text{SCN}^{2+} + \text{Cl} \quad \alpha = -1.0
\]

\[
\text{CCl}_2\text{SCN}^{2+} \rightarrow \text{CCl}_2^{+} + \text{SCN}^{+}
\]

Coincidence between Ions with m/z Values of 32 amu/q (S+) and 58 amu/q (SCN+). A probable mechanism for this coincidence could be a three-body secondary decay (SD) since the experimental slope for this island is −1.6 (calculated \(\alpha = -1.8\)). This graphic is shown in Figure S3C.

\[
\text{CCl}_3\text{SCN}^{2+} \rightarrow \text{CCl}_3^{+} + \text{SCN}^{+}
\]

\[
\text{SCN}^{+} \rightarrow \text{S}^{+} + \text{CN}
\]

Coincidence between Ions with m/z Values of 32 amu/q (S+) and 82 amu/q (CCl2+). A four-body secondary decay in competition (SDC) represents a reasonable mechanism for this coincidence:

\[
\text{CCl}_3\text{SCN}^{2+} \rightarrow \text{CCl}_3^{+} + \text{SCN}^{+}
\]

\[
\text{CCl}_2^{+} \rightarrow \text{CCl}_2^{+} + \text{Cl}
\]

\[
\text{SCN}^{+} \rightarrow \text{S}^{+} + \text{CN}
\]

When the kinetic energy release corresponding to the ejection of a neutral fragment is ignored, a slope of −1.2 can be calculated in the frame of a SDC mechanism. This value agrees with the determined experimentally of \(\alpha = -1.3\), as shown in Figure S3D.

3.6.2. Dynamics of Fragmentation for the CCl2FSCN2+ ion. Coincidence between Ions with m/z Values of 58 amu/q (SCN+) and 101 amu/q (CClF+). In coincidence with the halomethyl-thiocyanate family, the most abundant photo-fragmentation channels of core-excited CCl2FSCN is the rupture of C=S single bond from double charge molecular ion to form SCN+ and CCl2F+ species. This island of

| Table 3. Branching Ratios (%) for Fragment Ions Extracted from PEPIPICO Spectra Taken at Photon Energies around the S 2p and Cl 2p edges for CCl3SCN and CCl2FSCN Molecules |
|----------------|----------------|----------------|
|               | photon energy (eV) |               |
|               | 165.4 | 166.8 | 167.8 | 199.9 |
| m/z | ion | CCl3SCN | | | |
| 12 | C+ | 3.9 | 4.3/2.25 | 4.5 | 6.2/3.41 |
| 14 | N+ | 1.5 | 1.5/5.18 | 1.5 | 1.5/2.71 |
| 26 | CN+ | 3.5 | 3.7/2.15 | 3.6 | 4.1/3.47 |
| 32 | S+ | 8.8 | 9.2/1.37 | 9.8 | 11.9/2.54 |
| 35 | Cl+ | 17.2 | 19.6/2.61 | 20.4 | 25.6/3.50 |
| 44 | CS+ | 6.1 | 7.2/0.88 | 6.7 | 4.7/1.38 |
| 47 | CCl+ | 11.9 | 12.5/0.82 | 13.7 | 13.1/1.72 |
| 58 | SCN+ | 7.3 | 6.8/5.5 | 7.4 | 10.3/9.3 |
| 70 | C2SCN+ | 1.1 | 1.1/0.26 | 1.0 | 1.1/0.41 |
| 79 | C2ClS+ | 5.7 | 4.6/0.46 | 5.5 | 2.7/0.53 |
| 82 | C2Cl2+ | 11.7 | 11.5/1.04 | 9.2 | 9.8/1.73 |
| 117 | C3Cl+ | 12.7 | 10.3/0.18 | 9.5 | 5.8/0.68 |
| 140 | CCl3SCN+ | 7.1 | 5.9/0.17 | 6.0 | 2.5/0.22 |

“Kinetic energy release values (in eV) determined from the spectra at 166.8 and 199.9 eV are given in italics. Contributions of the 35/37 natural isotopologues of Cl were summed together. Ions with branching ratios lower than 1% are not listed. “Double KER distribution is observed (see text).”

lighter positive ions are observed, enabling the study of the dissociation of double ionized species. Thus, PEPIPICO spectra were registered as in the case of PEPIPICO at each of the resonant energies values on the S 2p and Cl 2p regions. Furthermore, the analysis of the PEPIPICO spectra is useful for identifying the dynamic of the fragmentation processes, and the occurrence of two-, three-, and four-body dissociation mechanisms can be determined.55,56

The evaluation of the experimental slopes for coincidence islands in the resonance and off-resonance S 2p photon energy region evidence that the observed differences in the dissociation mechanisms are less significant. The following discussion will
Coincidence is one of the most intense and can be observed with a parallelogram-like shape and a slope of $-1.0$ (see Supporting Information Figure S4A).

$$\text{CCl}_2\text{FSCN}^{+2} \rightarrow \text{CCl}_2\text{F}^+ + \text{SCN}^+ \quad \alpha = -1.0$$

Coincidence between ions with m/z values of 32 amu/q ($S^+$) and 101 amu/q ($\text{CCl}_2\text{F}^+$). In Figure S4B, the coincidence between these ions has been observed with the slope of $-1.4$. The occurrence of the three-body secondary decay (SD) mechanism, with a calculated slope of $-1.8$ can be the origin of this coincidence island.

$$\text{CCl}_2\text{FSCN}^{+2} \rightarrow \text{CCl}_2\text{F}^+ + \text{SCN}^+$$

$$\text{SCN}^+ \rightarrow \text{S}^+ + \text{CN}$$

Coincidence between ions with m/z values of 32 amu/q ($S^+$) and 66 amu/q ($\text{CCIF}^+$). A coincidence island with an experimental slope of $-1.2$, as shown in Figure S4C, indicates that the four-body secondary decay in competition (SDC) can be a probable mechanism for this coincidence since its calculated slope is $-1.4$, if the kinetic energy release corresponding to the ejection of the neutral fragments is neglected.

$$\text{CCl}_2\text{FSCN}^{+2} \rightarrow \text{CCl}_2\text{F}^+ + \text{SCN}^+$$

$$\text{SCN}^+ \rightarrow \text{S}^+ + \text{CN}$$

$$\text{CCl}_2\text{F}^+ \rightarrow \text{CCIF}^+ + \text{Cl}$$

4. CONCLUSIONS

An exhaustive study of electronic properties of the CCl$_3$SCN and CCl$_2$FSCN molecules in the gas phase following continuum valence and shallow-core (S 2p and Cl 2p) excitations has been performed using multicoincidence techniques based of time-of-flight mass spectrometry and synchrotron radiation in combination with photoelectron spectroscopy. PES spectra of both molecules are analyzed, and the valence electronic structure determined. HOMO corresponds to an $n_\pi(S)$ lone pair electrons, characterized like a well-defined band in the photoelectron spectra. A dependency of the vertical ionization energy with the electronegativity of the $-\text{CX}_3$ ($X = \text{H}, \text{Cl}, \text{F}$) group have been observed for the CX$_3$SCN species.

TIV spectra of the title molecules in the S 2p and Cl 2p levels were recorded. The spectra for these species are dominated by a group of well-defined signals between 164.0 and 169.0 eV of energy, and these resonant transitions are due to dipole allowed transitions that involve electronic transitions from an atomic 2p to an antibonding molecular orbital.

The ionic fragmentation in both valence and shallow-core regions were analyzed with the PEPICO and PEPITICO techniques. Dissociation mechanisms have been proposed in order to explain the ionic fragmentation decay for single- and double-charged species. In the valence region, very simple PEPICO spectra are obtained. The single charged molecular ion is observed with low intensity, whereas the main photodissociation channel is produced for the rupture of the sulfur–carbon single bond, leading to the formation of CCl$_2X^+$ ions ($X = \text{Cl}$ or F). Furthermore, the analysis of the PEPITICO spectra has been used to identify the dissociation mechanisms for doubly charged parent ions. Fragmentation processes leading to the formation of CX$_4^+$ and SCN$^+$ ions from CX$_3$SCN$^{+2}$ dominate the dissociation of halomethyl-thiocyanates excited at the S 2p levels. Also, the three-body secondary decay (SD) mechanism is one of the main photodissociation channels observed in both molecules, which leads to the formation of S$^+$ and CX$_3^+$ ions in the coincidence island. In both cases, the experimental slopes have been in agreement with the calculated ones.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpca.7b08395.

Correlation diagram of the ionization potential of CX$_3$SCN molecules; characters of the four lowest unoccupied orbitals CCl$_3$SCN and CCl$_2$FSCN; contour plot for the main coincidence island observed in the PEPITICO spectra of CCl$_3$SCN and CCl$_2$FSCN.
Dedicated to Prof. Dr. Rüdiger Mews on occasion of his 75 birthday.

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