

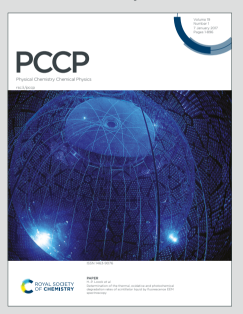




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Spectroscopic characterisation of radical polyinterhalogen molecules†

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Spectroscopic characterisations of the radical polyinterhalogen molecules IF2 and I2F are reported using anion photoelectron spectroscopy. The corresponding parent anions, IF_2^- and I_2F^- , are common products formed in hard Ar-CF₃I plasmas and are relevant in the semiconductor manufacture industry. The I_2F^- species, which is present as the [I-I-F] isomer, is a "non-classical" polyinterhalogen.

The propensity for halide anions to combine and form polyhalogen anions, such as I₃, has been known for around 200 years. 1 However, examples and spectroscopic characterisation of polyinterhalogen molecules, particularly open-shell species, have proven to be elusive.² To date, most spectroscopic characterisations of polyinterhalogen anions are as crystallised solids or have been performed in matrix isolation with a counter cation. 2-4 There are no known vibrationally-resolved spectroscopic determinations of radical (open-shell) polyinterhalogen molecules. On the other hand, closed-shell polyinterhalogen molecules such as IF₃, IF₅, IF₇ and I₂F₄ are relatively stable compounds. Polyinterhalogen anions and their corresponding radical neutrals are appealing targets for gas-phase spectroscopy due to being textbook examples for valence-shell electron pair repulsion (VSEPR) theory and the Rundle-Pimentel scheme for hypervalency. 5,6 They are also desirable systems to benchmark high-level quantum chemical calculations due to challenges associated with describing heavy atoms. 7,8

electronic structure theory study on the IF₂, I₂F⁻, IF₂ and I₂F species. There have been several reports of synthesis, crystallography and spectroscopic characterisation of IF₂ embedded in a solid state matrix, 9,10 and computational studies on ${\rm IF}_2^-$ and ${\rm I}_2{\rm F}^$ and related species, 7,10-14 however, a recent review by Riedel and co-workers² noted "Thus far, ... [I₂F] have not been detected experimentally." While this statement might be true for condensed phases, I₂F⁻ appears to be a common product formed in hard Ar-CF₃I plasmas.

There are only a handful of examples of gas-phase spectroscopic studies on polyhalogen species, mostly targeting I₃. A series of pioneering investigations by Neumark and co-workers 15-18 applied anion photoelectron spectroscopy with different laser sources to characterise vibronic properties of the radical neutral I₃, and photodissociation dynamics of the parent anion. Several other studies have focussed on photodissociation dynamics. $^{19-22}$ There have been two investigations on I₂Br⁻ probing the dissociation dynamics, with one of these studies providing a single-colour photoelectron spectrum at 267 nm. ^{23,24} In the present work, we have coupled a jet-cooled plasma discharge source with time-offlight mass spectrometry and anion photoelectron spectroscopy. This strategy allows for generation and mass selection of IF₂ and I₂F⁻, and determination of spectroscopic properties for the radical neutral IF2 and I2F species.

Intriguingly, radical polyinterhalogen molecules may have a significant bearing in industrial plasma processes. In particular, in the present study we produced the IF₂ and I₂F species through plasmisation of an Ar-CF3I mixture; this mixture and process is relevant in the semiconductor manufacture industry. In this industry, dry-etching plasma processes utilise the reaction between silicon wafers, which are the substrate for semiconductor and microprocessors, and fluorine atom and CF3 radicals to etch the substrate at specific locations on the wafer. ²⁵ Conventionally, these radical species are produced in a plasma consisting of an intert carrier gas seeded with CF₄ or C₃F₈. In recent years, there has been growing interest in the use of CF₃I as an alternative to CF₄ because the use of CF₄ is restricted in many countries by the Montreal Protocol due to the molecule's high global warming potential (GWP). ^{26,27} For example, even though CF₃I is infrared active, CF₃I has an atmospheric residence life of ≈ 1 day

This paper reports a combined photoelectron spectroscopy and

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[†] Electronic Supplementary Information (ESI) available: Experimental methods, Theoretical methods, Calculated vertical excitation energies, Further Franck-Condon simulations. See DOI: 00.0000/00000000.

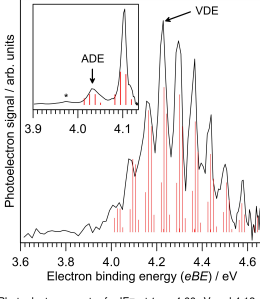


Fig. 1 Photoelectron spectra for IF_2^- at $hv=4.66\,\mathrm{eV}$ and $4.13\,\mathrm{eV}$ (inset) and Franck-Condon simulation (red sticks) of the $D_0 \leftarrow S_0$ detaching transition for the [F-I-F]⁻ isomer. Electron binding energy (eBE) and electron kinetic energy (eKE) are related by eBE=hv-eKE, where hv is the photon energy.

and corresponding GWP \approx 1. 28,29 In contrast, CF $_4$ has a residence life of \approx 50,000 years and corresponding GWP \approx 6,000. 29,30 Furthermore, although iodine atoms formed through UV photodissociation of CF $_3$ I are exceptionally destructive towards stratospheric ozone, 31 the short residence time of CF $_3$ I released at terrestrial altitudes means that these molecules do not reach the stratosphere and contribute to ozone destruction. Ultimately, modelling and tuning of the etching process allows maximisation of process efficiency and minimisation of unwanted by-products. Theoretical plasma models require a detailed qualitative and quantitative understanding of the rich ion-molecule and electron-molecule chemistry that occurs in dry-etching plasmas (particularly the chemistry associated with fluorine-containing radicals). $^{32-36}$

IF₂ & **IF**₂

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Photoelectron spectra for IF₂⁻ are shown in Fig. 1, revealing a clear vibrational progression with spacing $600\pm50\,\mathrm{cm}^{-1}$. The ADE, assigned as the lowest energy discernible detaching vibration in the hv=4.13 eV spectrum, is ADE= 4.03 ± 0.02 eV. The VDE, determined as the most intense vibration in the hv = 4.66 eVphotoelectron spectrum, is VDE=4.23±0.04 eV. These ADE and VDE values are most consistent with the calculated values (Table 1) for the [F-I-F] isomer shown in Fig. 2; there was no evidence for the [F-F-I] - isomer, which was calculated to lie 2.74 eV (264 kJ mol⁻¹) higher in energy. Calculated ADE and VDE values for [F-I-F] at the CCSD(T)//def2-TZVPD level of theory (Table 1) are both \approx 4% larger than the experimental values (similar to the result for I⁻ given in the ESI). Increasing the basis set to def2-QZVPD gave calculated values that were ≈6% larger than the experimental values (Table 1). The better agreement with the smaller basis set is presumably a fortuitous cancellation of errors with some fraction of this associated with heavy atom/relativistic

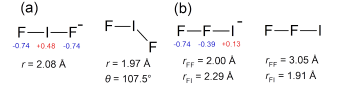


Fig. 2 Calculated equilibrium geometries at the CCSD(T)//def2-TZVPD level of theory for: (a) [F-I-F] $^-$ and [F-I-F], and (b) [F-F-I] $^-$ and [F-F-I]. NBO charges (red and blue font) are given for the anions. The lowest energy isomer, [F-I-F] $^-$ has D $_{\infty h}$ symmetry, and [F-I-F] has $C_{2\nu}$ symmetry.

effects.

A Franck-Condon simulation of the photoelectron spectrum for the [F-I-F] configuration is shown in Fig. 1 and has good agreement with the experimental spectrum. The simulation suggests that the vibrational progression is dominated by detaching transitions to the v_2 (asymmetric stretch) and v_3 (symmetric stretch) vibrations of the [F-I-F] radical neutral species; calculated frequencies suggest that these two vibrations are nearly degenerate - see Table 2. We can therefore assign the experimental value of v_2/v_3 at $600\pm50\,\mathrm{cm}^{-1}$ for the [F-I-F] species. The small feature denoted by * in the inset in Fig. 1 is probably a hot band from hot ions produced in the plasma (see ESI). The ADE assignment should be reliable based on alignment of the VDE with the Franck-Condon simulation. It is worth noting that a Franck-Condon simulation of the photoelectron spectrum for the [F-F-I] isomer (see vibrational frequencies for [F-F-I] in Table 2) is inconsistent with the experimental spectra, lending support to the above assign-

Calculated vertical excitation energies for the [F-I-F] $^-$ isomer are given in Table S1 in the ESI. These data show that although an excited state resonance is energetically accessible with a $hv=4.66\,\mathrm{eV}$ photon, oscillator strengths are zero and therefore the photoelectron spectra in Fig. 1 should be described by direct photodetachment to the ground electronic state of the neutral. Calculated vertical excitation energies for neutral [F-I-F], i.e. electronic excited states of [F-I-F] at the [F-I-F] $^-$ equilibrium geometry, suggested that only the ground electronic state is accessible using the photon energies in Fig. 1, consistent with a single vibrational progression.

 I_2F^- & I_2F Photoelectron spectra for I_2F^- at three photon energies are shown in Fig. 3a, revealing two detachment bands. The lower bind-

	[F-I-F]-	[F-F-I]-	[I-I-F]-	[I-F-I] -
ADE_e	4.03 ± 0.02	_	$4.04{\pm}0.06$	_
VDE_e	4.23 ± 0.04	_	4.23 ± 0.04	
ADE_c	4.14 (4.15)	3.17	4.22 (4.28)	3.19 (3.38)
VDE_c	4.40 (4.50)	3.88	4.40 (4.55)	3.96 (4.08)

Table 1 Experimental (subscript *e*) and calculated (subscript *c*) electron detachment energies for IF_2^- and I_2F^- . Calculations are at the CCSD(T)/def2-TZVPD level of theory (CCSD(T)/def2-QZVPD in parentheses). Experimental uncertainties account for calibration uncertainity from I^- and pixel-to-energy conversion in image reconstruction.

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Mode	[F-I-F]-	[F-I-F]	[F-F-I]-	[F-F-I]
$\overline{v_1}$	193 ^a	96 ^a	137^{a}	18 ^a
v_2	193^{a}	547 ^c	137^{a}	18^{a}
v_3	401^{c}	548^{b}	230^{c}	38^b
v_4	437^{b}		247^{b}	614 ^c

Table 2 Calculated harmonic vibrational frequencies for IF $_2^-$ and IF $_2$ at the CCSD(T)//def2-TZVPD level of theory in units of cm $^{-1}$. a Bend. b Symmetric stretch. c Asymmetric stretch. [F-F-I] has C $_{\infty h}$ symmetry. Note the symmetric vs asymmetric stretch mode ordering.

ing energy band has ADE=4.04 \pm 0.06 eV and VDE=4.23 \pm 0.04 eV. There is no clear vibrational structure. The higher binding energy band has ADE2=4.60 \pm 0.04 eV and VDE2=4.68 \pm 0.04 eV, and vibrational spacing of 550 \pm 100 cm⁻¹.

The ADE and VDE parameters for the lower energy band are consistent with the calculated values for the [I-I-F] $^-$ isomer (Table 1), again with a small overestimation by theory. The two bands in Fig. 3a are thus assigned to detaching transitions to the ground and excited electronic states of the [I-I-F] radical neutral. Equilibrium geometries for the I_2F^- and I_2F isomers are shown in Fig. 4; the [I-I-F] $^-$ isomer is more stable by 1.35 eV (130 kJ mol $^{-1}$), consistent with it being the predominant isomer formed in the plasma. There was no evidence for the [I-F-I] $^-$

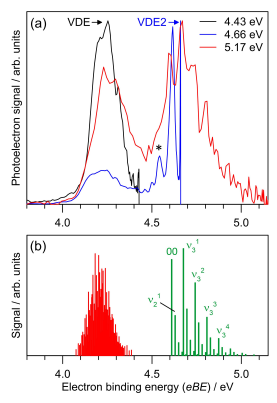


Fig. 3 (a) Photoelectron spectra for I_2F^- at hv=4.43, 4.66 and 5.17 eV. Electron binding energy (eBE) and electron kinetic energy (eKE) are related by eBE = hv - eKE, where hv is the photon energy. (b) Franck-Condon simulation of direct photodetachment to the first two neutral electronic states. The simulations suggest that the $D_0 \leftarrow S_0$ detaching transition is dominated by low frequency modes and combination bands. See ESI for Franck-Condon simulations at higher temperatures which assign the feature denoted by * to a hot band.

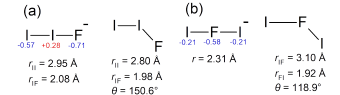


Fig. 4 Calculated equilibrium geometries at the CCSD(T)//def2-TZVPD level of theory for: (a) [I-I-F] $^-$ and [I-I-F], and (b) [I-F-I] $^-$ and [I-F-I]. NBO charges (red and blue font) are given for the anions. The lowest energy isomer, [I-I-F] $^-$ has $C_{\infty h}$ symmetry, and [I-I-F] has C_s symmetry.

isomer when photoelectron spectra were recorded at photon energies below the ADE for the [I-I-F]⁻ isomer. A Franck-Condon simulation of the photoelectron spectrum to give the ground electronic state of the neutral (red in Fig. 3b) shows no clear vibrational structure, consistent with experimental spectrum.

The second, higher binding energy photodetachment band in Fig. 3a was assigned to direct photodetachment to the A_1 state of the [I-I-F] species. Geometry optimisation of this state at the EOM-CCSD//def2-TZVPD level of theory produced a linear ($C_{\infty h}$ symmetry) structure with $r_{\rm II}$ =3.02 Å and $r_{\rm IF}$ =1.96 Å. Using this structure, ADE2 was calculated at 4.79 eV (0.57 eV difference to ADE), which is again \approx 4% larger than experiment. Calculated vibrational frequencies for the A_1 state state are v_1 =109 cm⁻¹ (bend), v_2 =157 cm⁻¹ (symmetric stretch) and v_3 =530 cm⁻¹ (asymmetric stretch). A Franck-Condon simulation of the photoelectron spectrum associated with formation of the A_1 state of the neutral is shown in green in Fig. 3b, and predicts that the vibrational structure is predominantly from the v_3 mode. The feature denoted by * in Fig. 3a at eBE=4.56 eV is assigned to hot band signal (see ESI).

Calculated vertical excitation energies for the I_2F^- isomers are given in Table S1 in the ESI. For the $[I\text{-}I\text{-}F]^-$ isomer, the first few electronic states have very low oscillator strengths and the photoelectron spectra for photon energies <5 eV should be dominated by direct photodetachment. There is, however, a bright A_1 -symmetry resonance with a calculated vertical excitation energy in the 5.1–5.2 eV range (see ESI). Fortunately, the photoelectron spectra for both of these species are at lower binding energies and should not interfere with the spectra shown in Fig. 3. It is worth noting that the issue of bright excited state resonances in the detachment continuum is more serious for I_3^{-15-18} and larger polyinterhalogen anions due to an increased density of electronic states, hampering measurement of photoelectron spectra.

Polyinterhalogen molecules can be divided into two categories,

Mode	[I-I-F]-	[I-I-F]	[I-F-I]-	[I-F-I]
$\overline{v_1}$	124^{a}	102^{a}	99 ^a	20^{a}
v_2	124^{a}	158^{b}	99^{a}	72^b
v_3	136^{b}	502^{c}	120^b	606^{c}
v_4	402 ^c	_	229^{c}	_

Table 3 Calculated harmonic vibrational frequencies for I_2F^- and I_2F at the CCSD(T)//def2-TZVPD level of theory in units of cm⁻¹. ^aBend. ^bSymmetric stretch. ^cAsymmetric stretch.

classical and non-classical. The former is defined as containing an electropositive central atom surrounded by electronegative halogen atoms, e.g. [F-I-F] $^-$. In contrast, for a non-classical polyinterhalogen molecule, the central halide is more electronegative than the coordinating dihalogen or interhalogen molecules, e.g. $Cl(I_2)_4^-$ in which four I_2 molecules coordinate a Cl^- . 3,37 Interestingly, the [I-I-F] $^-$ species satisfies the non-classical criterion, possessing covalent-like bond lengths and a terminal F^- coordinated to an I_2 molecule (see NBO charges in Fig. 4). Similar arguments have been made about the [I-I-Cl] $^-$ species in the solid state. 3,38 Finally, it is worth noting that Mabbs $et\ al.\ ^{23}$ reported a 267 nm photoelectron spectrum of I_2Br^- , which presumably exists as the [I-I-Br] $^-$ isomer and also is an example of a non-classical polyinterhalogen, however, the photoelectron spectrum showed no vibrational structure.

Our experiments suggest that IF_2^- , $\mathrm{I}_2\mathrm{F}^-$ and most likely the corresponding neutrals are amongst the major products formed in a hard Ar-CF₃I plasma. In the semiconductor manufacture industry, radical neutrals including IF_2 and $\mathrm{I}_2\mathrm{F}$ formed in the etching plasma may collide and react with etchant substrates. In this context, the bond dissociation energy (BDE) to produce a fluorine atom is related to etching efficiency – a lower BDE gives enhanced etching capacity. For the [F-I-F] and [I-I-F] isomers, BDEs for fluorine atoms were calculated at 1.74 eV and 1.35 eV, respectively, using the CCSD(T)//def2-TZVPD level of theory. For comparison, the calculated BDE for a fluorine atom from the CF₃ radical is 3.51 eV, which is substantially larger than that for the [F-I-F] and [I-I-F] isomers and suggests that polyinterhalogen radicals formed in hard Ar-CF₃I plasmas are important etching species.

In summary, this paper has reported a combined photoelectron spectroscopy and electronic structure theory study on the IF₂, IF₂, IF₂ and I₂F species, determining the gas-phase structures and electron detachment parameters for the anions and vibrational structure of the neutrals. This determination provides the first vibrationally-resolved spectroscopic characterisation of isolated polyinterhalogen radicals. Whereas F_2 as the [F-I-F] isomer is a classical polyinterhalogen molecule, IF₂ as the [I-I-F] isomer is classified as a non-classical polyinterhalogen molecule. Theoretical modelling of these species has shown that the CCSD(T)//def2-TZVPD level of theory, on average, predicts ADE and VDE parameters to within \approx 4% of experiment, however, this level of theory suffers from some degree of Pauling-point agreement. The IF2 and I2F radicals may play an important role in plasma etching processes due to having low bond dissociation energies to give fluorine atoms.

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Conflicts of interest

There are no conflicts to declare.

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