XUV-laser excitation of molecular fluorine

P.F. Levelt, K.S.E. Eikema, S. Stolte, W. Hogervorst and W. Ubachs

Laser Centre, Vrije Universiteit, De Boelelaan 1081, 1081 HV Amsterdam, The Netherlands

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Excited states of the F₂ molecule are investigated in high resolution by means of 1 XUV + 1 UV resonance-enhanced photo-ionization. Perturbations in the C'Σ⁺ state and the spectroscopy of the H'Π, Rydberg state are studied in detail. It is shown that F₂ molecules may be state-selectively detected with a sensitivity of 5 x 10⁷ particles per cm³.

1. Introduction

Although fluorine is the simplest halogen molecule its spectrum is not as extensively studied as that of chlorine, bromine and, in particular, iodine. While I₂ has important band systems in the visible, the electronic structure of F₂ is such that absorption on transitions to bound states only occurs at wavelengths shorter than 102 nm. Relatively weak continuous absorption was observed in the near-UV at around 300 nm [1]. Emission in the visible range was recorded as early as 1924 [2], but a determination of term energies of the electronic levels involved had to await data on absorption. Investigations of the discrete absorption spectrum started in 1959 [3]. The most detailed and accurate study of the extreme ultraviolet (XUV) absorption spectrum was reported by Colbourn et al. [4]. In the wavelength range between 95 and 102 nm they observed several vibrational progressions, the most intense being denoted as the C'Σ⁺ state. Vibrational energy spacings indicate a large internuclear equilibrium distance for these states with almost purely ionic binding at large distances. Because of small Franck-Condon overlap factors for transitions to low vibrational levels of the C'Σ⁺ state, this vibrational sequence could not be followed to the bottom of the potential well and vibrational quantum numbers could not be given. In the wavelength range 92-95 nm Colbourn et al. located in addition a Rydberg state denoted as H'Πₙ with a characteristic small equilibrium bond distance. Photo-ionization studies have been performed by Berkowitz et al. [5], who established values for the ionization potential, the dissociation limit of the F₂⁺ ion and the limit for ion-pair formation. A theoretical study on the electronic structure of F₂ and F₂⁺ was performed by Cartwright and Hay [6].

Laser excitation of molecular fluorine has been applied by Bischof and co-workers [7,8], employing various resonance-enhanced multi-photon ionization (REMPI) techniques. States of gerade symmetry, F'Πₗ and f'Πₗ were investigated by 2+1 REMPI, while states of ungerade symmetry, the H'Πₙ and h'Πₙ states, were probed with 3+1 REMPI. In the present study narrow-band XUV-laser radiation is applied in a 1 XUV + 1 UV excitation process in F₂. In contrast to the 3+1 scheme, not only Rydberg states but also valence states, most notably the C'Σ⁺ state, are probed. Limited wavelength regions, covering the H'Πₙ, n=0, 1 and 2 states and the C'Σ⁺ vibrational progression near 96 nm, are studied spectroscopically with improved resolution. Moreover it is shown that the 1+1 REMPI scheme allows for a high detection sensitivity: F₂ concentrations as low as 5 x 10⁷ molecules per cm³ can be detected.

2. Experimental

The XUV-laser spectrometer has been described in some detail in a previous report [9,10]. In short the radiation of a Nd: YAG pumped dye laser system
(10 Hz repetition rate) is frequency-doubled and thereupon tripled in a pulsed gas jet. Xenon is used as the non-linear medium in the present study. For the harmonic conversion process lenses with focal lengths of $f = 15$ cm or $f = 35$ cm are used. The spatially and temporally overlapping XUV and UV beams perpendicularly intersect a skimmed supersonic molecular beam produced with a second pulsed valve. The intersection zone is at a constant distance from the frequency-tripling zone, so that XUV and UV power densities depend on the choice of the lens. To prevent clogging of the nozzle and corrosion of the equipment the fluorine gas is diluted with He with a ratio 10% F2:90% He. A low-density molecular beam from an expansion with a 1 bar backing pressure is used. F2 spectra are thus recorded from a molecular beam with a partial pressure of $5 \times 10^{-6}$ Torr in the interaction zone. Adjustment of the nozzle-skimmer distance and the time delay between the opening of the valve and the laser pulse allows for measurements in a relatively hot molecular beam (i.e. 250 K), which is favorable for a spectroscopic study including states with high J quantum numbers.

XUV-laser excitation spectra are recorded by means of 1 XUV + 1 UV photo-ionization. After resonant excitation of F2 into a particular state by XUV photons the molecules are subsequently ionized by UV photons, which are present in abundance. The ions produced are extracted from the interaction region with a static electric field of about 50 V/cm. In a field-free time-of-flight tube mass separation takes place before detection of the charged particles with an electron multiplier. Ions at selected masses are detected with a boxcar integrator. This wavelength and mass-selective spectroscopic method enables the recording of contamination-free spectra of F2, although absorption of N2 and O2 rest gas molecules occurs simultaneously. In addition to F + parent ions also F+ ions are observed with a relative signal intensity of about 5%-20% (see below).

Excitation spectra are frequency-calibrated by simultaneous recording of an I2 absorption spectrum at the fundamental wavelength of the dye laser [11]. Spectral line positions in the XUV domain are calibrated with an absolute accuracy of 0.1 cm$^{-1}$ by computerized interpolation. The observed spectral linewidths of 0.3 cm$^{-1}$ are solely determined by the bandwidth of the XUV radiation. The fluorine molecules are irradiated with XUV intensities estimated of several nJ/pulse, and with UV intensities up to 30 mJ/pulse (in a 5 ns pulse). Pulse-to-pulse fluctuations in the signal intensities of 30% are observed. Spectra are recorded by averaging over 10 shots per wavelength setting.

3. Measurements and interpretation

3.1. Spectroscopy

In fig. 1 an overview of an XUV-laser excitation spectrum of fluorine in the wavelength interval 94.58–97.15 nm, recorded by detection of F2+ ions, is shown. The range corresponds to the typical sextupled output of a pulsed dye laser, in this case with a mixture of rhodamine 590 and rhodamine 610, while scanning over its gain profile. The spectrum consists of a vibrational progression of red-shifted bands to the C 2Σu+ valence state in the notation of Colbourn et al. [4]. The vibrational numbering is not yet established and is therefore given with an arbitrary offset n. The transition to the C 2Σu+, ν = n + 26 state is strongest and individual spectral lines are recorded with a signal-to-noise ratio of up to 1000. The decreasing signal intensity for transitions to C 2Σu+, ν < n + 26 states in the range 96.25–97.15 reflects the fact that this part is recorded in the low-output wing of the dye gain profile. The lower signal intensities of the C 2Σu+, ν > n + 26 bands at the short-wavelength side, between 94.58 and 96.0 nm, correspond to a real decrease and cannot be ascribed to low dye-laser output.

At the short-wavelength side of the recorded spectrum, at $\lambda = 94.65$ nm, a blue-shifted band is observed. It shows the typical signature of a Rydberg state with an equilibrium internuclear distance smaller than the ground state. Transitions to the three lowest vibrational levels of this Rydberg state, denoted as H 1Πu, are recorded in more detail and line positions determined. A spectrum of the H 1Πu – X 1Σg+ (1, 0) band is presented in fig. 2. The R branch in the spectra of these three bands is clearly resolved for all J states. Spectroscopic constants for the H 1Πu ν = 0–2 states are deduced from least-squares fits using energy expressions for the excited state,
Fig. 1. Overview F2 spectrum in the wavelength range 94.58–97.15 nm. Part of the vibrational progression of transitions to the C⁺Σ⁺⁺ valence state as well as the H⁺Π⁺ v=0 state is shown.

Fig. 2. Spectrum of the H⁺Π⁺ –X⁺Σ⁺⁺ (1,0) transition. The peaks denoted with an asterisk indicate perturbations in the R-branch sequence.

\[
E_{\Pi}(J) = \nu_0 + BJ(J+1) - DJ^2(J+1)^2 + q_nJ(J+1), \quad (1)
\]

\[
E_{\Pi}(J) = \nu_0 + BJ(J+1) - DJ^2(J+1)^2. \quad (2)
\]

Here \( \nu_0 \) is the position of the bandhead, \( B \) the rotational constant, \( D \) the centrifugal distortion parameter and \( J \) the rotational quantum number. The subscripts \( e \) and \( f \) refer to the common convention of electronic symmetry of the two non-degenerate \( \Pi(J) \) levels, probed in the Q branch, and the P and R branch respectively. The constants derived from a least-squares minimization routine are listed in table 1. A significant value for the \( A \)-doubling constant \( q_n \) could only be determined for the H⁺Π⁺ v=1 state.

In the analysis the level energies of the X⁺Σ⁺⁺ v=0 ground state were calculated using the accurate rotational constants of Edwards et al. [12], obtained from Raman spectroscopy: \( B_0=0.88331 \text{ cm}^{-1} \) and \( D_0=3.48 \times 10^{-6} \text{ cm}^{-1} \). The observed line positions coincide with the calculated values within the error.
Table 1
Spectroscopic constants for the H 'IIu=0-2 states of F2. All values are in cm⁻¹

<table>
<thead>
<tr>
<th>H 'IIu=0</th>
<th>H 'IIu=1</th>
<th>H 'IIu=2</th>
</tr>
</thead>
<tbody>
<tr>
<td>ν₀</td>
<td>105606.41(2)</td>
<td>106674.82(2)</td>
</tr>
<tr>
<td>B</td>
<td>1.0087(5)</td>
<td>0.9976(4)</td>
</tr>
<tr>
<td>q</td>
<td>-0.0041(5)</td>
<td>-0.0041(5)</td>
</tr>
</tbody>
</table>

margin of 0.1 cm⁻¹, except for the R(20) and R(22) lines in the (1, 0) band that split into two components, probably as a result of a perturbation with a yet unknown state.

Because the transitions to the C 'Σ⁺, v= n+25 and v= n+26 states are strongest in intensity, they might provide a suitable and efficient scheme for state-selective probing of F₂ molecules. For this reason detailed measurements on the P and R branches of two bands were performed in order to clarify the spectroscopy. In fig. 3 recordings of the C 'Σ⁺ – X 'Σ⁺ (n+26, 0) band are presented, for F₂⁺ as well as F⁺ detection. From the analysis of Colbourn et al. [4] it was established that the C 'Σ⁺, v= n+26 state is perturbed by a vibrational level of another 'Σ⁺ state, denoted D 'Σ⁺, v= m+31. Because of the high spectroscopic precision of the present work (absolute accuracy of 0.1 cm⁻¹) their deperturbation procedure is repeated and extended. In the analysis rotational levels of the D 'Σ⁺, v= m+31 and v= m+32 are included and excited state energies are calculated by diagonalizing the energy matrix.
for each value of the rotational quantum number J. It is assumed that the perturbation parameters $H_{CD}$ are independent of J, as is expected to be the case for homogeneous interactions between electronic states of the same symmetry. Energies of the deperturbed levels are given in the common notation as in expression (2). P(J) and R(J) transitions for the three bands are calculated from the eigenvalues of the matrix. After assignment of the spectral lines the spectroscopic constants and interaction parameters are obtained from a least-squares minimization routine.

We could assign the R(0)-R(27) and P(1)-P(26) lines of the C-X (n+26, 0) band, the R(15)-R(17) and P(14)-P(19) lines of the D-X (m+31, 0) band and the R(19) and P(21)-P(22) lines of the D-X (m+32, 0) band. All lines are included in the fit.

The resulting parameters are listed in table 2. One of the interactions is clearly visible in the spectrum of fig. 3 as a change of sequence in the R branch of the C-X (n+26, 0) band at R(14)-R(15). The anti-crossing with D, $v=m+31$ takes place between J=15 and 16, while D, $v=m+32$ anti-crosses between 24 and 25. The next anti-crossing, with D, $v=m+33$ is expected at J=29, but here the data are insufficient for a detailed analysis. All lines included in the routine fit with a resulting standard deviation of 0.12 cm$^{-1}$. However, the R(6) and P(8) lines of the C-X (n+26, 0) band are both shifted downward in energy by a significant 0.6 cm$^{-1}$. This is an indication of another perturbation of the C, $v=n+26$ state, albeit a much weaker one than the other two. As the C$^1 \Sigma^+_u$, $v=n+26, J=7$ state lies at 104125 cm$^{-1}$ the band origin of the perturbing state is expected near 104115 cm$^{-1}$ assuming a rotational constant of 0.16 cm$^{-1}$. From interpolation of the vibrational spacing in the D$^1 \Sigma^+_u$ states it follows that the D$^1 \Sigma^+_u$, $v=m+30$ state cannot cause this perturbation. Its energy is 70 cm$^{-1}$ too low. However, Colbourn et al. observed another vibrational progression, denoted as E$^1 \Sigma^+_u$. Extrapolation of the band origins upward from E$^1 \Sigma^+_u$, $v=p+17$ indicates that the perturber of the C$^1 \Sigma^+_u$, $v=n+26, J=7$ state may well be the E$^1 \Sigma^+_u$, $v=p+20, J=7$ state.

The C$^1 \Sigma^+_u$, $v=n+25$ state, and a single perturber, the D$^1 \Sigma^+_u$, $v=m+29$ state, are treated in a similar procedure and the resulting parameters are also listed in table 2. In this case the anti-crossing occurs between J=11 and 12. The bandhead of the D-X (m+29, 0) band is observable as weak lines near 103885 cm$^{-1}$ in the spectrum of fig. 3. The signal-to-noise ratio in these lines is about 10, a factor of 100 lower than in the C-X (n+26, 0) band.

We note that the intensities of transitions to the perturber states largely depend on the mixing of the excited state eigenfunctions. Where at the bandhead the intensity of the D-X (m+29, 0) lines is only 1% of the intensity of the C-X lines, the R(11) line of the D-X (m+31, 0) band has 60% of the intensity of the R(11) line of the C-X (n+25, 0) band. At $J=12$ the Cn+25 state has 61% C-state character, while the Dm+29 state has 39% C-state character. The intensities of transitions to all D$^1 \Sigma^+_u$, $v, J$ perturber states is quantitatively analyzed and it is found that they are also determined by the C-state character of the excited state.

### Table 2

Spectroscopic constants of F$_2$ for the C$^1 \Sigma^+_u$, $v=n+25$ and $v=n+26$ states and the perturbing D$^1 \Sigma^+_u$, $v=m+29$, $m+31$ and $m+32$ states as well as the homogeneous perturbation parameters. All values are in cm$^{-1}$.

| $\nu_0$ | 103833.16±0.05 | 104093.09±0.02 |
| $B$ | 0.5175±0.0004 | 0.6049±0.0002 |
| $D$ | (4.73±0.08)×10$^{-5}$ | (6.97±0.04)×10$^{-5}$ |

| $\nu_0$ | 103885.1±0.1 | 104196.1±0.1 | 104357.2±0.1 |
| $B$ | 0.1602±0.0002 | 0.1692±0.0004 | 0.1749±0.0003 |
| $D$ | (2±1)×10$^{-5}$ | (2.8±0.4)×10$^{-5}$ | (5.0±0.1)×10$^{-5}$ |

$$
\begin{pmatrix}
E_{Cn+26}(J) & H_{CD31} & H_{CD32} \\
H_{CD31} & E_{Dm+31}(J) & 0 \\
H_{CD32} & 0 & E_{Dm+32}(J)
\end{pmatrix},
$$

3.2. F$^+$ parent ion and F$^*$ fragment ion formation

Molecular fluorine excitation spectra have been measured by simultaneous recording of signals de-
ected at mass 38 and mass 19, corresponding to $F_2^+$ parent and $F^+$ fragment ions respectively. The observations on $F^+$ production are the following:

- $F^+$ signals are found at the same frequency positions as the resonances in the $F_2^+$ spectrum. Moreover it is verified that both signals only occur when XUV radiation is present in the setup, so that direct UV-induced multi-photon excitation is excluded.

- Relative intensities in the $F^+$ spectrum follow that in the $F_2^+$ spectrum. In all cases the $F_2^+$ yield is largest. However, for two neighbouring lines near 103920 cm$^{-1}$ there is an anomalously high rate for $F^+$. These lines are indicated in fig. 3 and may be assigned as the R(24) and P(21) lines of the C-X ($n+26$, 0) band. The precise $F_2^+/F^+$ ratios depend on UV-laser input power, choice of the focusing lens for the generation of third-harmonic radiation and on the settings of the ion optics.

- The XUV radiation is generated in different focusing conditions, making use of $f=15$ cm or $f=35$ cm lenses. In case of the $f=15$ cm lens, UV power density in the $F_2$ excitation zone is a factor of 5 higher than in the case of $f=35$ cm, resulting in a $F^+$ yield a factor of 4 higher.

- For a typical line, the C-X ($n+26$, 0) R(3) line, the UV power dependence of the ion yields is carefully studied. The results are shown in fig. 4. From the slopes in a log–log plot it follows that the signal yield of $F_2^+$ ions has an $I_{\text{UV}}$ ($p=2.5 \pm 0.5$) power dependence and the $F^+$ yield an $I_{\text{UV}}^p$ ($p=4.5 \pm 0.5$) power dependence. The XUV intensity is found to depend on UV input according to $I_{\text{XUV}} \propto I_{\text{UV}}^{-3}$.

From these observations it may be concluded that all spectral features, including those in which $F^+$ ions are formed, are related to an initial XUV excitation of a neutral $F_2$ molecule. The production of $F_2^+$ ions is straightforward: after resonant excitation by incident XUV radiation excited $F_2$ molecules are subsequently ionized by a single UV photon. In the 1 XUV+1 UV process about 17.4 eV is absorbed, whereas the ionization limit of $F_2$ is at 15.686 eV [6]. As the $F^+$ yield depends more strongly on UV input power than the $F_2^+$ yield it follows that for the production of $F^+$ more than a single UV photon must be involved.

In order to facilitate an interpretation of the $F^+$ yield a schematic energy diagram of the $F_2-F_2^+-F^+$ system is presented in fig. 5, with the different possible product channels indicated. The bold arrows correspond to photon-induced excitations, either by XUV or by UV photons; the thin arrows indicate dissociative decay. An asterisk denotes an excited state, a double asterisk a doubly excited state.

Fig. 4. Dependence of $F_2^+$ and $F^+$ yields on intensity of the UV input. The full and dashed lines correspond to fits through the data points. The slopes indicated represent the UV power dependences.

Fig. 5. Schematic energy diagram (not on scale) for the interpretation of ion product channels. The bold arrows correspond to photon-induced excitations, either by XUV or by UV photons; the thin arrows indicate dissociative decay. An asterisk denotes an excited state, a double asterisk a doubly excited state.
ferent routes that may lead to fragment ion formation indicated. In view of the intensity ratios of XUV and UV laser beams (UV 10^6 more intense) it is unlikely that a second XUV photon will be absorbed.

The first possible channel is that of ion-pair formation after excitation of a doubly excited intermediate state. The energy threshold for ion-pair formation is 15.58 eV [5], which is reached by absorption of an XUV photon and a single UV photon. So the ion-pair channel requires the same number of UV photons as the parent-ion channel. The fact that F+ production more strongly depends on UV intensity is an argument against the ion-pair hypothesis.

The second channel for F+ formation, the dissociation of the F2+ parent ion, requires absorption of an additional UV photon. Energetically 19.01 eV is needed for the process F2→F+ + F and this is provided by 1 XUV plus 2 UV photons. From the study of Cartwright and Hay [6] it follows that an F2+ ion in the X 2Π_g ground state may be vertically excited to the repulsive part of the first 2Π_u state by a 4 eV photon. So this channel can well explain the steeper UV dependence for F+ formation. The excitation to a repulsive state indicates a non-resonance behaviour in the ionic absorption as well. This explains why the relative intensities of the F+ yield closely follow those of the F2+ yield.

In the third channel after resonant absorption of an XUV photon the neutral molecules are further excited by absorption of a UV photon into a doubly excited state (F**) that may dissociate into a ground state neutral atom and an excited neutral atom, denoted by F*. If the dissociation energy of F2 is accounted for (1.59 eV [5]), via this process only atomic states of the (2s)^2(2p)^43s configuration may be populated. The excited atom may absorb further UV photons, first into a Rydberg state (F**) just below the atomic ionization threshold, followed by a second one producing F+. In the sequence of channel (3) in two of the steps coincidental UV wavelength-dependent excitation is supposed. Therefore this route might explain the coincidental anomalies in the F+ yield. However, the structure of the doubly excited molecular states (F2**) as well as the atomic Rydberg states (F**) is not known, so no definite conclusions may be drawn. It seems certain that channel (3) cannot provide an explanation for the major part of the F+ yield, that follows the intensity pattern of the F2+ yield.

As a fourth route atomic excitation by XUV photons might be considered. The UV intensity is such that 0.1% of F2 is dissociated during the laser pulse [1]. Ground state F atoms subsequently may be ionized in a 1 XUV+2 UV REMPI process analogous to the 3+2 REMPI excitations observed by Bischel and Jusinski [7]. In that case resonances are expected at wavelengths of 95.48, 95.19, 95.55 and 95.15 nm corresponding to transitions in the F atom [13]. However these atomic resonances are not observed.

It is concluded that channel (2), dissociative UV excitation of the F2+ parent ion, is the most probable route for the formation of F+, at least for the vast majority of features. The resonance features with anomalously high F+ yield may be interpreted as related to channel (3), where excited state F atoms are photo-ionized.

4. Conclusion and outlook

The method of 1 XUV+1 UV photoionization is shown to be a highly sensitive spectroscopic tool for the study of excited states of F2 with high resolution. As an example the perturbations in the spectra of two selected valence bands of the C 'Σ_u^+ state were treated in detail. Furthermore accurate spectroscopic constants could be determined for the H 'Π_u ν=0–2 states. In principle the present apparatus may be used for a study of the full wavelength range 90–102 nm.

The band intensities observed in the present study markedly differ from those obtained in 3+1 REMPI studies of Bischel et al. [7,8]. In those experiments, performed in a geometry with a focused UV-laser beam, the H 'Π_u and h 'Π_u Rydberg states were observed and no evidence of the C 'Σ_u^+ state was found. In agreement with observations of Colbourn et al. [4], it is found that the intensities of transitions to the D 'Σ_u^+ state are governed by intensity borrowing form the C 'Σ_u^+ state. While this was found to hold for whole bands at low vibrational levels, strong J-dependent intensity borrowing effects within a particular band are observed in the present work. No direct evidence of the E 'Σ_u^+ state is observed, other
than an indirect indication of a perturbation with the C' \Sigma_u^+ state.

The spectra of F2 have been recorded in a diluted, low-pressure molecular beam expansion with a relatively high rotational temperature. Assuming an F2 particle density of about $5 \times 10^{10}$ cm$^{-3}$ in the interaction zone, taking into account a distribution over rotational quantum states and a signal-to-noise ratio of 1000, a detection limit of $5 \times 10^7$ molecules per cm$^3$ or $5 \times 10^6$ molecules per quantum state per cm$^3$ is estimated. This implies that a 1 XUV + 1 UV photoionization scheme is a sensitive tool for the state-selective detection of F2 molecules. This method may prove fruitful in various applications, such as in gas-phase reaction dynamics and surface scattering experiments involving F2 (e.g. etching).

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References