Nitrogen-Broadening Parameters for Atmospheric Spectra Modelling of the $v_3$ Band of SF$_6$

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Abstract: The infrared absorption of the $v_3$ band region of SF$_6$, at temperatures spanning the 130 to 297 K range, has been reexamined using improved instrumentation with one goal: to estimate the broadening of parameters by nitrogen gas. These parameters are compared to previous literature predictions and an extended set of IR cross-sections is proposed and compared to other existing datasets.

Keywords: SF$_6$; IR absorption; nitrogen-broadening; cross-sections

1. Introduction

SF$_6$, as a synthetic gas with several technological applications, is now present in the atmosphere and, although at relatively small concentration, it has been steadily growing [1,2]. With a long lifetime and very strong absorptions in mid-infrared atmospheric transmission windows, it is an important greenhouse gas to be monitored and one of the species to be controlled mentioned in the Kyoto protocol, due to its exceptionally high global warming power (23,900 times that of CO$_2$). Additionally, as it is mostly produced in the northern hemisphere, it is used as a probe to investigate the stratospheric circulation processes and test the currently accepted models; see for instance [3] and references therein. The infrared spectrum of gaseous SF$_6$ has been extensively studied at low resolution (see Ref. [4] and references therein) and all its IR-active [5–7], Raman active [8–10] as well as its IR and Raman-silent [11] fundamentals have been observed at high to very high resolution and analyzed in detail [12]. In particular, the region of the very strongly absorbing $v_3$ mode, which is easiest to detect in situ, is used in various atmospheric spectrometric monitoring experiments [13,14]. Until recently, there existed no line list providing a complete description of the $v_3$ region, as it is cluttered by many hot bands that contribute significantly to the absorption (only about 30% of the population in the ground state near 300 K). The attempts to complete this linelist by adding linelists sequentially provided by the analysis of different transitions involved in the series of $(v_3 + (n + 1)\nu_6) - n\nu_6$ hot bands [15,16] still fails in the absence of a correct parameter set describing the main hot bands of the $(v_3 + (n + 1)\nu_6) - n\nu_6$ series. Instead, several authors have proposed empirical cross-sections [17,18] to help quantification in the remote sensing atmospheric spectra, providing spectra at given pressures and temperatures.

However, this useful approach reaches its limits for the correct retrieval of the molecular column in low-pressure stratospheric conditions as spectra cannot be recalculated in the exact physical conditions. Another, innovative approach has been only recently...
implemented by the Reims and Tomsk groups, which calculated from first principles potential energy and dipole moment multi-dimensional surfaces and used them to compute synthetic spectra, potentially including transitions originating from all populated vibrational levels [19]. This is extremely promising, yet also very challenging in computational power for such large molecular systems with many vibrational degrees of freedom. In [19], Nikitin et al. have thus operated empirical corrections to compensate for the lack of accuracy of the pure ab initio calculations. It thus seems interesting to provide high-quality, high-resolution reference spectra, as well as the most complete line list possible based on effective Hamiltonians, to serve as reference points for some of the most important transitions. Additionally, even supposing that these theoretical efforts will eventually reach perfection, a good knowledge of the pressure-broadening coefficients and their temperature dependence will still be necessary for the comparison with high-altitude limb atmospheric spectra, where SF$_6$ is clearly observable [13,14]. To our knowledge, two studies have been devoted to the assessment of the pressure-broadening coefficients and their temperature dependence. First, the computational study of Tejwani and Fox [20] has calculated the broadening coefficients of SF$_6$ by several gases (He, Ar, O$_2$, N$_2$, HF and SF$_6$ itself) at 100, 200, and 300 K within the Anderson–Tsao–Curnutte model. For N$_2$, they found a very weak $J$-dependence of the pressure-broadening and calculated values for the $\gamma_0$ coefficient and $n$ temperature law exponent of 0.05 cm$^{-1}$/atm on average and 0.623, respectively. More recently, Gamache and coworkers [21] have calculated N$_2$-induced pressure-broadening coefficients using the more advanced, complex Robert–Bonamy (CRB) formalism [22] at 200, 250, 296, and 350 K. They also compared their predictions with experimental measurements at 295 K and N$_2$ pressures ranging from 200 to 1000 mbar. Their calculations also predict a very weak $J$-dependence of the broadening coefficient. Their estimation of the $\gamma_0$ coefficient and $n$ temperature law exponent are 0.078 cm$^{-1}$/atm and 1.05 on average, respectively. In the course of previous studies in our laboratory [23–25], we found evidence that the N$_2$ pressure-broadening coefficient or its temperature dependence departed noticeably from these predictions and this motivated this new investigation using extended experimental capabilities.

This manuscript is organized as follows: in the first part, new data concerning the $\nu_3$ region of SF$_6$ recorded at improved resolution over a large temperature range and with varied nitrogen pressures are presented. In the second part, we show that the traditional line-by-line retrieval methods are not tractable in such a case and an effective, global approach is presented. From this, an average value of the N$_2$-broadening coefficient is determined, even and foremost for a temperature low enough so that the contribution of the $\nu_3$ fundamental can be clearly separated from that of the hot bands.

2. Experimental

Because of the very high IR intensity of the $\nu_3$ band, a short optical path length at low to room temperature was required. High-resolution spectra were recorded on the Bruker 125HR interferometer AILES Beamline at Synchrotron SOLEIL, using a globar source with appropriate iris aperture diameters, a KBr/Ge beamsplitter and a high-sensitivity home-made, 4 K-cooled HgCdTe detector with a cold 880–1125 cm$^{-1}$ bandpass filter, aperture and focusing optics [26]. Given the strong intensity of the transition, we used the newly developed cryogenic very short path cell regulated at temperature between 130 and 296 K. This setup has been fully described elsewhere [27]. Briefly, it consists of a short, 51 mm long double-walled stainless steel cell with diamond windows, placed in the spectrometer light path under high vacuum (4 × 10$^{-5}$ mbar). Cold gas from liquid-N$_2$ evaporation is flowed in the double jacket surrounding the cell to vary the gas temperature. Three temperature sensors, placed inside and outside the cell, are used for temperature measurement and regulation, as well as a resistive heater. Because of the non-linear nature of the feedback loop for the temperature regulation, we estimate a $+/−2$ K temperature uncertainty only for the 130 to 270 K temperature measurements, $+/−1$ K for the room-temperature measurements. A high-precision MKS capacitive thermally regulated gauge
(1000 mbar range, 0.1% precision) is connected to the cell filling tube to read out the pressure some 30 cm away from the gas cell. Because of the thermomolecular effect [28,29], this can introduce a potential bias in the pressure reading at low temperatures if the gas cell pressure is kept low (less than 1 mbar). To alleviate this problem and for other reasons detailed below, it was decided to keep the total gas pressure larger than 3 mbar. Given the large IR intensity of the $\nu_3$ fundamental, this meant preparing very dilute SF$_6$–N$_2$ gas mixtures in a large, well outgassed 3 l stainless steel chamber using 1 mb (for SF$_6$) and 1000 mb (for N$_2$) Pfeiffer capacitive gauges, and waiting several hours for complete equilibration. Next, precisely measurable pressures of the gas mixture were introduced in the cell. Six series of spectra, corresponding to six different temperatures (130, 169, 213, 230, 270 and 296 K) were recorded at varying SF$_6$/N$_2$ volume mixing ratios. Spectra were ratioed against an empty cell background, post-zero-filled, corrected for channeling effects and calibrated using well-known water and carbon dioxide rotational lines [30]. The experimental conditions are reported in Table 1.

| Temperature (K) | Spectrum | $P$ SF$_6$ (mbar) | $P$ N$_2$ (mbar) | Iris dia. (mm) | Resolution (cm$^{-1}$) | #Scans |
|----------------|----------|------------------|-----------------|---------------|------------------------|--------|
| 130 (2)       | P130K 3.27 mb | 0.017            | 3.25           | 1.15          | 0.00102               | 120    |
| 130 (2)       | P130K 6.57 mb | 0.035            | 6.53           | 1.3           | 0.0015                | 58     |
| 130 (2)       | P130K 9.46 mb | 0.051            | 9.41           | 1.7           | 0.002                 | 60     |
| 130 (2)       | P130K 20 mb   | 0.11             | 19             | 2             | 0.003                 | 60     |
| 130 (2)       | P130K 30 mb   | 0.16             | 29.9           | 2             | 0.003                 | 60     |
| 169 (2)       | P170K 10 mb   | 0.065            | 10.06          | 1.3           | 0.0015                | 90     |
| 169 (2)       | P170K 50 mb   | 0.065            | 40.1           | 1.3           | 0.005                 | 50     |
| 169 (2)       | P170K 100 mb  | 0.065            | 90             | 1.3           | 0.005                 | 50     |
| 169 (2)       | P170K 200 mb  | 0.065            | 202            | 1.3           | 0.01                  | 50     |
| 169 (2)       | P170K 800 mb  | 0.065            | 803            | 1.3           | 0.02                  | 50     |
| 213 (2)       | P213K 10 mb   | 0.064            | 9.94           | 1.3           | 0.0015                | 94     |
| 213 (2)       | P213K 50 mb   | 0.064            | 49.9           | 1.3           | 0.005                 | 100    |
| 213 (2)       | P213K 100 mb  | 0.064            | 89.9           | 1.3           | 0.005                 | 100    |
| 213 (2)       | P213K 200 mb  | 0.064            | 199.9          | 1.7           | 0.01                  | 100    |
| 230 (2)       | P230K 10 mb   | 0.064            | 9.97           | 1.3           | 0.0015                | 120    |
| 230 (2)       | P230K 20 mb   | 0.064            | 20.2           | 1.7           | 0.002                 | 80     |
| 230 (2)       | P230K 50 mb   | 0.157            | 49.9           | 1.7           | 0.006                 | 60     |
| 230 (2)       | P230K 100 mb  | 0.157            | 100.1          | 1.7           | 0.006                 | 60     |
| 230 (2)       | P230K 200 mb  | 0.157            | 200.1          | 1.7           | 0.01                  | 60     |
| 270 (2)       | P270K 15 mb   | 0.064            | 9.97           | 1.3           | 0.0015                | 120    |
| 270 (2)       | P270K 50 mb   | 0.160            | 50.0           | 1.7           | 0.006                 | 80     |
| 270 (2)       | P270K 100 mb  | 0.320            | 100.0          | 1.7           | 0.006                 | 60     |
| 270 (2)       | P270K 200 mb  | 0.320            | 200.1          | 1.7           | 0.006                 | 60     |
| 270 (2)       | P270K 800 mb  | 0.320            | 800            | 1.7           | 0.02                  | 60     |
| 297 (1)       | P297K 20 mb   | 0.129            | 20.2           | 1.3           | 0.0012                | 480    |
| 297 (1)       | P297K 50 mb   | 0.165            | 50.4           | 1.3           | 0.006                 | 480    |
| 297 (1)       | P297K 100 mb  | 0.319            | 100.2          | 1.3           | 0.006                 | 100    |
| 297 (1)       | P297K 100 mb  | 0.319            | 100.2          | 1.3           | 0.006                 | 100    |
| 297 (1)       | P297K 200 mb  | 0.319            | 216.3          | 1.3           | 0.01                  | 100    |
| 297 (1)       | P297K 800 mb  | 0.320            | 802            | 1.3           | 0.02                  | 100    |

3. Methods

The very high line density due to the small rotational constant, the additional splitting due to centrifugal distortion, and the presence of numerous hot bands had previously prevented a detailed and complete, line-by-line analysis of the $\nu_3$ region (except for the line
positions of the cold band itself [6,12]). Thus, our strategy was first to work at a temperature low enough to restrict observation to lines of the fundamental band. We progressively reduced the gas temperature until the onset of condensation inside the cell was reached, i.e., about 130 K in our experimental conditions. This corresponds to a SF₆ pressure of the order of less than 0.02 mbar. Since the gas pressure is measured with a pressure gauge regulated at 313 K and placed some 30 cm away from the cold gas cell, the precise measurement of a small gas pressure within the cold cell will be affected by thermomolecular effects [26,27]. To circumvent this difficulty, we diluted the SF₆ gas in nitrogen to introduce in the cell a total gas pressure well above 2 mbars, where the thermomolecular corrections become negligible. By carefully choosing the starting nitrogen pressure and using a spectral resolution high enough, this also had the advantage of slightly broadening the lines to the point that the correction due to interferometer instrumental line shape (ILS) becomes small to negligible.

Next, for each temperature, the nitrogen pressure was progressively increased to evaluate more precisely the nitrogen-broadening effect. As a first test, starting with the lowest temperature at which the spectrum is most simplified, the traditional approach of fitting individual lines in the spectrum using a program taking explicitly into account the ILS was attempted (WSpectra) [31]. The linelist present on the SHeCaSDa website [32,33] was used for positions and the measured gas temperature was used for the Gaussian component of the Voigt profile. Adjustment was thus made on the Lorentzian component.

Due to the severe line overlap, the analysis of only about 260 lines (corresponding to degenerate transitions) could be attempted.

The results are presented in Figure 1 with error bars estimated from the deviation of the fit. It confirms that this approach is too inaccurate to be useful in the case of such dense spectra, but the Lorentzian widths estimated here seem to be close to an average value of 0.125 ± 0.04 (1σ) and did not seem to indicate a notable dependence in J, confirming on that point the predictions of the theory [20,21]. This motivated us to attempt a more global and effective approach, based on the assumption that the J-dependence of the nitrogen-broadening coefficient and the pressure shifts can be neglected in a first approach. Starting with the coldest spectrum (at 130 K, where the hot bands are very weak), each individual spectrum was recalculated (using the XTDS, SPVIEW [34] and SGEN [35] software) with an effective broadening coefficient. It should be noticed that the XTDS software [34] whose results are reflected in the SHeCaSDa database [32] uses the effective dipole moment derivative for the ν₃ band that was derived by Person and Krohn a long time ago [36]. This value appears to be reliable, and it did not appear necessary to change it for the present simulations. The very small value of the SF₆ to N₂ mixing ratio (0.5% in most cases) justifies neglecting the self-broadening contribution. The program takes into account the correction due to the instrumental and the Doppler contribution.
Figure 1. Estimated Lorentzian widths of fitted lines in spectrum P130K 3.27 mb. The error bars are estimations from the fit.

4. Results

Figure 2 displays the comparison of experimental data and simulation obtained for a value of the broadening coefficient $\gamma(130 \, \text{K}) = 0.131 \, \text{cm}^{-1}/\text{atm}$. The insert presents a detail on the first J clusters on the R-branch side of the band, where lines of the first hot band ($\nu_3 + \nu_6 - \nu_6$, with a very weak Q-branch near 946.8 cm$^{-1}$) are the least noticeable (at 130 K, the $\nu_6 = 1$ level population is about 7%, while the ground state population represents about 90%).

Figure 2. Comparison of experimental spectrum (blue, P130K 3.27mb) and simulated (red) for the $\nu_3$ region of SF$_6$ the temperature is 130(2) K. The black trace is the difference between the upper traces, offset by $-0.05$ for clarity.

Please note that the residuals pictured on the black lower trace presents a root mean square deviation of 0.0065 over the whole band (of about 0.25 integrated intensity here), thus about four times that due to the measurement noise. Taking the strongest R cluster...
lines around 948.7 cm$^{-1}$, the rms value of the difference between observed spectrum and simulation represents 2.6% of the line intensity. Additionally, a departure of 0.02 cm$^{-1}$/atm produces a residual over the whole band that is four times higher. The uncertainty in the broadening coefficient estimate is thus determined, but is as a result of trial-and-error iteration. It is clear that the method does not allow the demonstration of fine differences in the broadening coefficients of lower and higher $J$ lines, but these seem small enough to be within the estimated error bar. Figure 3 presents a comparison for a gradually increasing pressure of nitrogen for the same, low temperature.

![Figure 3](image-url)

**Figure 3.** Comparison of experimental (blue), simulated (red, $\gamma = 0.131$ cm$^{-1}$/atm) spectra for increasing pressures of nitrogen at 130 K: (a) 3.27 mb, (b) 6.57 mb, (c) 9.46 mb, (d) 20 mb. The black traces represent the residuals for each pressure, shifted by $-0.05$ for clarity.

As the gas temperature rises, the effective broadening parameter must be progressively readjusted. This is somewhat complicated by the hot bands, as hot band line intensities increase progressively. Nevertheless, Figure 4 shows that the simulations remain convincing as the residuals for lines of the fundamental band are well within the hot band lines.

![Figure 4](image-url)

**Figure 4.** Comparison of experimental (black) and simulated spectra for two temperatures with 10 mb pressure of nitrogen. The dotted lines represent the residuals for each temperature, shifted by $-0.05$ for clarity.
The uncertainty on the estimation of the broadening coefficient increases, however, steadily with the temperature, as the hot band lines gain in intensity. Figure 5 presents a spanning larger view of the simulation in the R-branch at the highest temperature explored in this study.

![Absorbance vs Wavenumber (cm⁻¹)](image)

**Figure 5.** Comparison of experimental (black) and simulated spectra (red) for 297 K with 20 mb pressure of nitrogen. The blue line represents the residuals, shifted by −0.05 a.u. for clarity.

The estimation of the uncertainty on the effective broadening coefficient takes into account the relative uncertainties on pressure and temperature and, more important, the uncertainty on the adjustment of the observed profile obtained by varying slightly the value of the broadening coefficient around the optimum value until the residuals are clearly above the intensity of the hot band lines. Table 2 presents the estimation for the values of the effective nitrogen-broadening parameter for each temperature along with the estimated uncertainties.

**Table 2.** Estimated average nitrogen-broadening parameter for the $\nu_3$ band of SF$_6$. Estimated uncertainties are presented in parenthesis in units of the last digit.

| Temperature (K) | $\gamma(T)$ |
|----------------|-------------|
| 130 (2)        | 0.131 (5)   |
| 169 (2)        | 0.105 (5)   |
| 213 (2)        | 0.090 (6)   |
| 230 (2)        | 0.085 (7)   |
| 270 (2)        | 0.074 (8)   |
| 297 (1)        | 0.065 (1)   |

The temperature dependency of the effective broadening coefficient can be modelled using the usual power law:

$$\gamma(T) = \gamma_0 \left(\frac{T_0}{T}\right)^n$$

(1)

where the dependency is contained in the exponent $n$ and $T_0 = 297$ K. Figure 6 presents the results along with a linear fit using this power law. Please note that a simple least-square fit results in a convergence towards values close to 0.068 and 0.81 for $\gamma_0$ and $n$, but, when the results are weighted by the uncertainties, somewhat different values are obtained, 0.069 (1) and 0.77 (2) for $\gamma_0$ and $n$. The uncertainties given here correspond to 1 $\sigma$ deviation from the
fit and should be regarded as a lower limit. Given the approximation used in the method, we think it is safer to assume a 3σ deviation and the values obtained are thus $\gamma_0 = 0.069 \ (3)$ and $n = 0.77 \ (6)$.

Figure 6. Variation with temperature of the estimated effective nitrogen-broadening coefficient. The red line corresponds to a weighted least-square fit.

Another way to estimate the range of parameters compatible with our results is to take the two straight lines passing through all the error bars with either the higher or lower slopes. This yields the same average values but with larger uncertainties ($\gamma_0 = 0.069 \ (5)$ and $n = 0.77 \ (15)$).

In a second step, we used the recorded data to extract a set of absorption cross-sections in temperature and pressure ranges relevant for atmospheric measurements and which may be useful to compare and complete cross-section datasets recently published [17]. These are summarized in Figures 7–11 and available in the Supplementary Materials, which can be downloaded on the journal website.

Figure 7. Absorption cross-sections in the $\nu_3$ region of SF$_6$ at 169 K for various nitrogen pressures. The insert shows detail of the Q-Branch for the first three pressures.
Figure 8. Absorption cross-sections in the $\nu_3$ region of SF$_6$ at 213 K for various nitrogen pressures. The upper traces are offset by steps of $2 \times 10^{-16}$ for clarity.

Figure 9. Absorption cross-sections in the $\nu_3$ region of SF$_6$ at 230 K for various nitrogen pressures. Curves for 20 to 200 mb nitrogen pressures are offset for clarity.

Figure 10. Region of SF$_6$ at 270 K for various nitrogen pressures.
which are likely to exist, but seem small enough to be within the estimated error bar. The calculation of Tejwani and Fox [20] indicate a very small variation of the broadening parameter over a large range of J quantum number ($\gamma$) between 0.108 and 0.097 cm$^{-1}$/atm and $\gamma$ (300 K) between 0.054 and 0.051 cm$^{-1}$/atm). Their estimate of an average value of $\gamma_0$ is 0.051 cm$^{-1}$/atm and their estimate of the $n$ exponent is 0.623. The calculation by Gamache and coworkers [21] also predicted a small variation of $\gamma_0$ with rotational quantum numbers, roughly from 0.074 to 0.0798 cm$^{-1}$/atm and an exponent $n$ value around 1.05. [21] also presents a comparison of an experimental spectrum and a simulation based on the $\gamma$ (295 K) value derived from theory. Their value around 0.075 cm$^{-1}$/atm is close to our estimate ($\gamma_0 = 0.069(5)$ cm$^{-1}$/atm), yet slightly outside the error bar of our estimation. Since our estimate accounts quite well for the temperature dependence of the broadening coefficient, we think this value represents at present the best starting point for future simulation of this important spectral region, based on future complete line lists for this region. We want to stress again that the method used here does not allow the demonstration of fine differences in the broadening coefficients of lower and higher J lines, which are likely to exist, but seem small enough to be within the estimated error bar.

The absorption cross-sections presented here are in good general agreement with those presented in the recent publication of Harrison [18], with small differences only. It is important to note that the absorption cross-sections in [18] were normalized to the PNNL integrated absorption intensities of [37], while the present measurements provide an independent determination. The integrated band strength found here for 169, 213, 230, 270, and 297 K are 1.87 (8), 1.88 (8), 1.88 (8), 1.81 (8), and 1.81 (8) $\times 10^{-16}$ cm/molecule, thus 1.85 (8) $\times 10^{-16}$ cm/molecule on average. This is to be compared to the value of 1.96 (3) $\times 10^{-16}$ cm/molecule of [37]. The difference is small and at the limit of the cumulated error bars, but the small dispersion in our measurements gives us confidence that this difference is physically significative and that our data can slightly improve the precision of concentration retrievals. These data have been incorporated in the 2018 release of the technical note for MIPAS data processing [38].

Finally, the inspection of our data, even at moderate resolution with the effect of pressure-broadening, is interesting for the spectral analysis of the region, for which the most intense hot bands have not yet been analyzed. It is interesting to note that it can be clearly seen on the spectra at 130 and 169 K that the Q-branch of the first hot band ($v_3 + v_6 - v_6$)
has an asymmetric, two-maxima structure, presumably due to at least two IR-active subcomponents (946.70 and 946.85 cm\(^{-1}\)). The Q-branch of the second hot band (\(v_3 + v_5 - v_9\)) produces the broad shoulder with two maxima near 945.91 and 946.16 cm\(^{-1}\) [15], while the structure near 945.62 cm\(^{-1}\), growing next to temperature, is likely to correspond to the \((v_3 + v_4 - v_4)\) and \((v_3 + 2v_6 - 2v_6)\) hot bands. The next challenge will be to supply reliable linelists for all these bands. Combined with the present nitrogen-broadening coefficient, these will supply the means for efficient atmospheric modelling at any temperature and pressure.

**Supplementary Materials:** The Supplementary Materials are available online. The absorptions cross sections at 169, 213, 230, 273 and 297 K are available online.

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**Data Availability Statement:** The cross-sections data presented in this study can be downloaded from the Supplementary Materials.

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