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Vibrational shifts of absorption bands of linear molecules diluted in high-density rare gases: Measurements and modeling for CO₂-Rg and OCS-Rg

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Abstract

Increase of buffer gas pressure causes redistribution of intensity in the IR absorption bands and, indicating the presence of vibrational perturbation, changes the first spectral moment value responsible for the band-origin position. Band-origin shift coefficients for the ν_3 , $2\nu_3$ and ν_2 bands of OCS and the ν_3 band of CO₂ diluted in high-density Ar, Kr and Xe are measured and calculated by two methods: directly with available in the literature vibrationally dependent potential-energy surfaces for both initial and final vibrational states and assuming that vibrational dependence arises from the interaction of the permanent /transient dipole moment of the absorbing molecule and the dipole moment induced on the perturber. A comparative analysis of results is given, showing a general underestimation of calculations with respect to measurements.

Keywords: Infrared absorption spectra, Band-origin shift, Spectral moments, Vibrational dependence, DID model, Carbon dioxide, Carbonyl sulfide, Rare gases

1. Introduction

Though molecular spectra contain information on the nature of intermolecular interactions and features of molecular structure, their interpretation is associated with numerous difficulties [1] still to be resolved. For instance, spectral band shapes change significantly with increasing pressure and a set of characteristics is required for description of the band under various experimental conditions. In particular, the spectral moments M_n account for numerous band properties and thus can be used for practical purposes, though even a complete set of spectral moments does not characterize a unique band. For examining the interaction mechanisms that result in specific-band properties, particularly suitable candidates are benchmark systems composed of linear molecules, such as CO₂ and OCS, perturbed by rare gases.

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These two molecules are present in planetary atmospheres (94.9% and 96.5% of CO₂ for Mars [2] and Venus [3], respectively, and trace-gas level [4, 5] of OCS), comets [6, 7], and interstellar space [8, 9]. Precise knowledge of their spectral band shapes and band origins is **therefore** essential for reliable radiative transfer simulations and remote sensing of **atmospheric environments**. Line positions, intensities, widths and shifts have been closely considered in numerous works. For OCS-Ar e.g., line widths in the ν_1 [10] and hot [11] bands, line widths and shifts in the $2\nu_3$ band [12] have been measured. For CO₂-Ar, line shifts in the 00⁰3-00⁰0 band [13], line shifts, speed-dependence parameters and line mixing coefficients in the 30013-00001 and 30012-00001 bands [14] have been reported. In contrast, studies of band-origin shifts remain very scarce up to now due to two major reasons.

From the experimental standpoint, determination of the band-origin shift (defined by M_1 — see Sec. 3.2) requires accurate intensity measurements up to very far spectral wings, which is a tedious and time-consuming task **because of the numerous neighboring bands to be removed**. That is why, traditionally, the band-origin shift is rather estimated from the position of the intensity maximum, as it was done in the work of Ozanne *et al.* [15] who investigated the ν_3 and $3\nu_3$ bands of CO₂-Ar at pressures up to 1000 bar. The band shifts were determined as a deviation of the measured intensity maximum from its position computed by the Energy-Corrected Sudden model. The shifting coefficients were found to be $\gamma(\nu_3) = (-5.4 \pm 1.0) \times 10^{-3} \text{ cm}^{-1} \text{ amagat}^{-1}$ and $\gamma(3\nu_3) = (-17.0 \pm 1.0) \times 10^{-3} \text{ cm}^{-1} \text{ amagat}^{-1}$.

From the theoretical viewpoint, accurate band-origin shift calculations require vibrationally dependent potential energy surfaces (PESs) for both initial (ground) and final (excited) states, that are currently available only for a limited amount of systems. If the vibrational dependence of **the** final-state PES is unknown, mechanisms responsible for inducing the band-origin shift must be examined (e.g., Ozanne and coauthors [15] suggested that the polarizability matrix elements of a linear molecule are vibrationally dependent).

In the present work we perform accurate high-pressure measurements of CO₂-Rg (up to 100 atm) and OCS-Rg (up to 300 atm) band shapes and rigorously compute the band-origin positions from the first moment definition (see Sec. 4.1). Moreover, we suggest that, in the absence of vibrationally-dependent PES, the interaction between the dipole moment induced on the atom and the dipole of the molecule (DID interaction) can also result in a band-origin shift and test this hypothesis by comparing the results of the DID model and those obtained with vibrationally-dependent PES for both initial and final states. In Section 2 we briefly describe the experimental conditions. Theoretical background is summarized in Section 3. Details of modeling for CO₂ and OCS are given in Sec. 3.2. The bulk of our experimental and theoretical results is discussed and compared with that of Ref. [15] in the final section.

2. Measurements

IR absorption spectra of CO₂-Ar, CO₂-Kr, CO₂-Xe, OCS-Ar and OCS-Kr gas mixtures at 298 K were registered with Bruker IFS-28 and Bruker IFS 125/HR spectrometers in the region 400–4500 cm⁻¹ with a spectral resolution 0.8 cm⁻¹. Active-gas concentrations were at trace-gas level, and the buffer-gas pressures varied from 15 to 100 atm for CO₂-Rg and from 25 to 300 atm for OCS-Rg. In function of the studied spectral range and the mixture

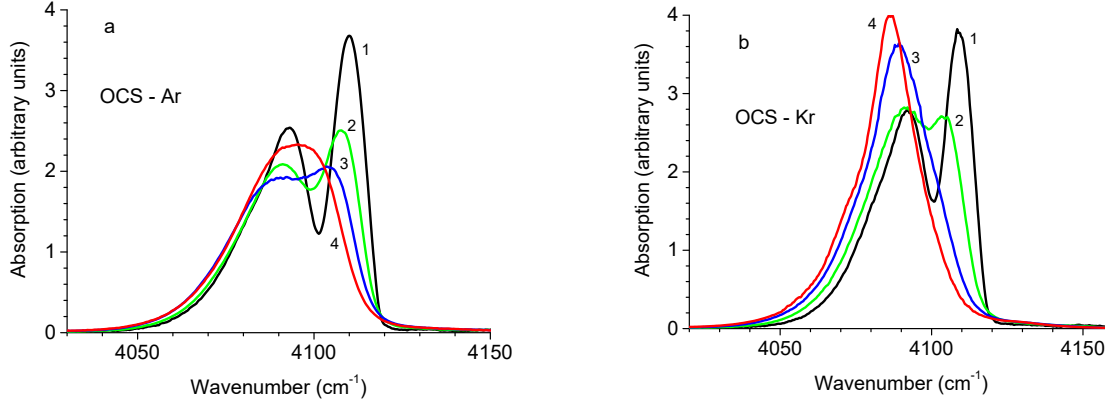


Figure 1: $2\nu_3$ OCS-Ar (a) and OCS-Kr (b) band shapes at buffer-gas pressures of 25 (1), 100 (2), 200 (3) and 300 (4) atm (active gas present at trace level).

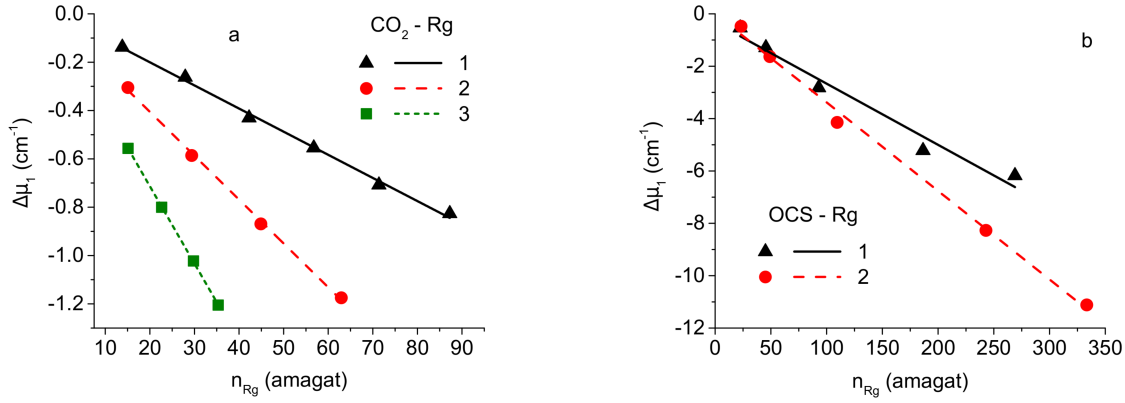


Figure 2: Deviations of the first moment $\Delta\mu_1 = \mu_1 - \mu_1^0$ with increasing gas density and corresponding linear fits for the ν_3 CO_2 (a) and $2\nu_3$ OCS (b) bands: perturbation by Ar (1), Kr (2) and Xe (3).

pressure, two cells were used: from 400 to 1000 cm^{-1} and pressures up to 100 atm a 13.5 cm cell with thallium bromoiodide (KRS-5) windows and from 1000 to 4500 cm^{-1} and pressures up to 300 atm a 7.4 cm cell with fluorite (CaF_2) windows. The mixtures were composed by pumping the compounds into a cell. The purity was controlled spectroscopically (no impurity-gas bands were detected above the noise level).

Figure 1 shows examples of OCS band-shape transformation with increasing buffer-gas pressure. For both active molecules, the band origin was tracked by the first moment definition (numerical integration of Eq. (2) of Sec. 3.2 with the subsequent normalization). Examples of experimental band-origin shifts versus gas density and their linear fits can be seen in Fig. 2. The experimental band-origin shift coefficients are given in Tables 1 and 2.

Table 1: Spectral ν_3 band-origin shift coefficients γ (in $10^{-3} \text{ cm}^{-1} \text{ amagat}^{-1}$) measured and calculated by the direct (Eq. (5)) and DID (Eq. (6)) methods for CO_2 perturbed by various rare gases. The references indicate the PES used for computations.

	expt	direct	DID
$\text{CO}_2\text{-Ar}$	-9.6(2)	-2.06 [16]	-6.28 [16], -6.57 [17], -6.27 [18]
$\text{CO}_2\text{-Kr}$	-18.2(2)	-9.51 [19]	-8.96 [19]
$\text{CO}_2\text{-Xe}$	-32.0(9)	-10.20 [20]	-13.29 [20], -2.83 [21], -3.36 [22]

Table 2: Experimental and DID-computed spectral band-origin shift coefficients γ (in $10^{-3} \text{ cm}^{-1} \text{ amagat}^{-1}$) for various bands of OCS perturbed by Ar (PES [23]) and Kr (PES [24]).

	ν_3 band		$2\nu_3$ band		ν_2 band	
	expt	DID	expt	DID	expt	DID
OCS-Ar	-8(2)	-7.25	-23(2)	-14.50	-2(2)	-0.12
OCS-Kr	-19(2)	-11.32	-34(1)	-22.63	-5(2)	-0.18

3. Theory

3.1. Absorption coefficient and correlation function

An incident electromagnetic wave with angular frequency ω and intensity I_0 , having propagated over a path length l in a media containing n_a absorbers per unit volume, loses its intensity I according to the Beer-Lambert law:

$$I(\omega) = I_0(\omega)e^{-n_a K(\omega)l}.$$

The absorption coefficient $K(\omega)$ is expressed [25] as

$$K(\omega) = \frac{4\pi^2\omega}{3\hbar c} \left(1 - e^{-\frac{\hbar\omega}{k_B T}}\right) F(\omega),$$

where $\hbar$, c and k_B stand for the Plank constant, speed of the electromagnetic wave in vacuum and Boltzmann constant, respectively, T is the temperature in Kelvin and $F(\omega)$ is the so-called spectral function. $F(\omega)$ may be regarded as the Fourier transform of a time correlation function $C(t)$ [26]:

$$F(\omega) = \frac{1}{2\pi} \int_{-\infty}^{\infty} dt e^{-i\omega t} C(t),$$

which is related to the dipole-moment absorption as

$$C(t) = \langle \mathbf{d}^\dagger(0) \mathbf{d}(t) \rangle. \quad (1)$$

Here, the dagger denotes the Hermitian conjugation and $\mathbf{d} = |v_f\rangle\langle v_f| \mathbf{D} |v_i\rangle\langle v_i|$ with $\langle v_f| \mathbf{D} |v_i\rangle$ standing for the vibrational $f \leftarrow i$ transition moment of the active-molecule dipole $\mathbf{D}$. The function $C(t)$, when known, determines completely $F(\omega)$ and, consequently, the full absorption band. To characterize the band-origin position we employ the spectral moments approach.

3.2. Spectral moments

The spectral moment of order n is defined by

$$M_n = \int_{-\infty}^{\infty} F(\omega) \omega^n d\omega. \quad (2)$$

Due to the fact that M_0 represents simply the integral intensity of the spectral band, normalized spectral moments $\mu_n = M_n/M_0$ are frequently used. From the Fourier-transform relationship $C(t) = \int_{-\infty}^{\infty} d\omega e^{i\omega t} F(\omega)$ one easily gets

$$\left. \frac{d^n C(t)}{dt^n} \right|_{t=0} = (i)^n \int_{-\infty}^{\infty} F(\omega) \omega^n d\omega.$$

Substitution of this equation in Eq. (2) gives

$$M_n = (-i)^n \left. \frac{d^n C(t)}{dt^n} \right|_{t=0}.$$

For our purposes the key interest is in the first spectral moment giving the band origin:

$$M_1 = -i \left. \frac{dC(t)}{dt} \right|_{t=0}.$$

With the definition (1) of $C(t)$ the time derivative can be re-written as

$$\left. \frac{dC(t)}{dt} \right|_{t=0} = \left\langle \mathbf{d}^\dagger(0) \frac{d}{dt} \mathbf{d}(t) \right\rangle \Big|_{t=0},$$

where

$$\frac{d}{dt} \mathbf{d}(t) = \frac{i}{\hbar} [H, \mathbf{d}(t)]$$

is determined by the commutator containing **the full** Hamiltonian H of the **interacting** molecule. The first spectral moment becomes

$$M_1 = \langle \mathbf{d}^\dagger(0) [H, \mathbf{d}(t)] \rangle / \hbar. \quad (3)$$

Moreover, H can be decomposed as a sum of the free-active-molecule Hamiltonian H_a and the absorber-perturber interaction term **(PES)** W :

$$H = H_a(\Omega, \mathbf{q}) + W(\mathbf{R}, \Omega, \mathbf{q}).$$

Here $\mathbf{R}$ is the intermolecular distance vector (between **the** centers of mass of the absorber and perturber), Ω specifies the active-molecule axis orientation and $\mathbf{q}$ is a set of vibrational coordinates. Thus, the rhs side of Eq. (3) reads

$$\langle \mathbf{d}^\dagger(0) [H, \mathbf{d}(t)] \rangle / \hbar = \langle \mathbf{d}^\dagger(0) [H_a, \mathbf{d}(t)] \rangle / \hbar + \langle \mathbf{d}^\dagger(0) [W, \mathbf{d}(t)] \rangle / \hbar.$$

The intermolecular potential W , in its turn, consists of the isotropic part $W^{iso}(R)$ and two compounds related to rotations and vibrations:

$$W(\mathbf{R}, \Omega, q) = W^{iso}(R) + W^{rot}(\mathbf{R}, \Omega) + W^{vib}(\mathbf{R}, \Omega, q).$$

The rotational perturbations W^{rot} and W^{iso} commute with the dipole moment $\mathbf{d}$ of a vibrational transition, thus the presence of solely rotational perturbation doesn't influence the value of the first spectral moment. For the vibrational transition $f \leftarrow i$ one gets

$$M_1 = M_0 \mu_1^0 + M_0 \langle W_{ff}^{vib} - W_{ii}^{vib} \rangle / \hbar \quad (4)$$

with the normalized first spectral moment in the absence of vibrational perturbation $\mu_1^0 = \langle v_f | H_a | v_f \rangle - \langle v_i | H_a | v_i \rangle$ and the double indices $ii(ff)$ denoting the vibrational diagonal matrix elements of $W^{vib}(\mathbf{R}, \Omega, q)$. Dividing Eq. (4) by M_0 one obtains

$$\mu_1 = \mu_1^0 + \gamma n_{Rg},$$

where n_{Rg} is the number density of the perturbing gas and the band-origin shift coefficient $\gamma = \langle W_{ff}^{vib} - W_{ii}^{vib} \rangle / (n_{Rg} \hbar)$ is introduced.

4. Band-origin shift computations

4.1. Direct computation of band shifts

Since the binary collision regime is confirmed by the linear dependence of measurements on n_{Rg} (see Fig. 2), the average required for the shifting coefficient calculation can be performed with the pair distribution function [27]:

$$\langle W_{ff}^{vib} - W_{ii}^{vib} \rangle = n_{Rg} \int d\mathbf{R} \int d\Omega e^{-\frac{W_{ii}}{k_B T}} (W_{ff}^{vib} - W_{ii}^{vib}), \quad (5)$$

where the first integral is a volume integral and the second one is the integral over the active molecule orientation; the diagonal matrix element W_{ii} corresponds to the full potential. Practical computations with Eq. (5) are determined by the availability of PES with vibrational dependence in the initial (typically, ground) and final (excited) states. Owing to relative simplicity of "linear molecule + atom" systems, their *ab initio* potential energy surfaces have been intensively studied in the literature. In the pioneering works, the absorbing molecule was treated as a rigid rotor [17, 18, 21, 22, 23, 24, 28], thus the vibrational dependence was totally neglected. However, accounting for antisymmetric stretching vibrations is essential for calculating band shifts [29] for both CO₂ and OCS. For direct computations we employed therefore only the vibrationally dependent potentials [16, 19, 20] constructed with noniterative inclusion of connected triples (CCSD(T)) and basis functions sets depending on the considered system.

4.2. Computation via dipole-induced dipole interaction

In the cases where vibrationally dependent PES are unavailable and Eq. (5) can not be used, evaluation of γ can be made with a simple model suggested bellow. Collisions with a polar active molecule may alter the spherical symmetry of the atomic perturber and induce on it an instantaneous dipole moment which will interact with the permanent (or transient) dipole of the molecule. This interaction can be a cause of the band origin shift. Indeed, the dipole moment $\mathbf{D}^{ind}$ induced on an atom placed in the electric field $\mathbf{E}$ is proportional to the atom polarizability α_{Rg} :

$$\mathbf{D}^{ind} = \alpha_{Rg} \mathbf{E},$$

and its potential energy reads

$$W^{ind} = -\frac{1}{2} \frac{\alpha_{Rg}}{R^6} (3 \cos^2 \theta + 1) D^2$$

with the angle between $\mathbf{D}$ and $\mathbf{R}$ denoted by θ . For a transition from the ground state i to a state f Eq. (5) becomes

$$\gamma = \frac{\pi \alpha_{Rg}}{\hbar} (\langle v_f | D^2 | v_f \rangle - \langle v_i | D^2 | v_i \rangle) \int_0^\infty \frac{dR}{R^4} \int_{-\pi/2}^{\pi/2} e^{-\frac{W_{ii}}{kT}} (3 \cos^2 \theta + 1) \sin \theta d\theta. \quad (6)$$

In the harmonic approximation (D is proportional to q) **one gets** $\langle 1 | D^2 | 1 \rangle - \langle 0 | D^2 | 0 \rangle = \langle 2 | D | 1 \rangle^2 = 2 \langle 1 | D | 0 \rangle^2$ and $\langle 2 | D^2 | 2 \rangle - \langle 0 | D^2 | 0 \rangle = 4 \langle 1 | D | 0 \rangle^2$. For our calculations, the transition dipole moments $d = \langle 1 | D | 0 \rangle$ were taken as 0.374 D and 0.0473 D for the ν_3 and ν_2 **OCS** bands, [30] respectively, whereas $d = 0.3212$ D was used for the ν_3 band of CO_2 [31]. The polarizabilities α_{Rg} were 1.6404 [32], 2.4844 [33], 4.044 [34] Å³ for Ar, Kr and Xe, respectively. The results of computations are presented in Tables 1 and 2.

5. Discussion

The bulk of our experimental and theoretical results is collected in Tables 1 (for CO_2) and 2 (for OCS). From the experimental standpoint, the band-origin shift coefficients appear to grow with increasing polarizability of the perturbing gas. Moreover, for the OCS case, the order of magnitude for the band shift for the ν_2 band is significantly smaller in comparison with that for the ν_3 band, which confirms the proportionality of the band shift to the dipole matrix element associated with the transition. Our experimental γ -value for CO_2 -Ar ($(-8.9 \pm 0.2) \times 10^{-3} \text{ cm}^{-1} \text{ amagat}^{-1}$) has the same order of magnitude as the datum published by Ozanne *et al.* [15] ($\gamma(\nu_3) = (-5.4 \pm 1.0) \times 10^{-3} \text{ cm}^{-1} \text{ amagat}^{-1}$) and, likely their results for the ν_3 and $3\nu_3$ bands of CO_2 -Ar, we find roughly the factor two between our measurements for ν_3 and $2\nu_3$ bands of OCS-Ar and OCS-Kr.

Due to the absence of vibrationally dependent potentials for OCS-Rg, direct calculations with Eq. (5) were feasible solely for CO_2 -Rg cases, **and** comparison with the DID model (Eq. (5)) can be made for this absorbing molecule only (Table 1). **In the case where the**

same PES is used, for CO₂-Kr both methods give close results, but for CO₂-Ar and CO₂-Xe direct computation leads to a smaller absolute value of the band shift than the DID model. It can be noted also that the DID results for CO₂-Ar are very similar for various PES tested, while for CO₂-Xe choice of the potential plays a significant role.

To summarize, both computational methods may be reliably applied to CO₂/OCS mixtures with Ar, but with increase of the perturber polarisability the experiment gives significantly larger values. The nature of this discrepancy currently remains unclear and deserves further detailed consideration.

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