Methane monitoring by near infrared photoacoustic spectroscopy: The importance of relaxation phenomena

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Abstract. The importance of molecular relaxation in photoacoustic spectroscopy is discussed. The particular case of methane monitoring in dry oxygen using a 1.65-μm laser-based photoacoustic sensor is reported. The slow vibration-to-translation energy transfer occurring in this gas mixture results in a drastic reduction of the detection sensitivity. A quadratic response of the sensor to the methane concentration is also reported and explained by molecular relaxation effects.

1. INFLUENCE OF MOLECULAR RELAXATION IN PHOTOACoustics

Photoacoustic (PA) spectroscopy basically consists in exciting an absorbing gas sample with a modulated laser beam at proper wavelength and detecting the generated acoustic wave using a microphone. The acoustic wave results from molecular absorption of photons and subsequent deactivation of the excited vibrational state via inelastic collisions with the nearby molecules (collisional relaxation). When the excited vibrational energy is transferred into kinetic energy of the surrounding molecules, a periodic heating occurs in the sample and induces the pressure wave. In order to efficiently generate a PA signal, the typical timescale of the vibration-to-translation (V-T) energy transfer should be much shorter than the period of the laser modulation (generally in the millisecond range). This condition is most often fulfilled, as the collisional relaxation time of most molecules ranges from nanosecond to hundreds of microseconds at atmospheric pressure. However, the relaxation time of some particular molecules may be orders of magnitude longer. In particular, this is the case for a few diatomic molecules, such as N₂ or O₂ (see Table 1), and for collisions with some particular partners (N₂ and O₂, too). When the timescale of the V-T processes is comparable or larger than the modulation period, the PA signal is poorly excited.

Table 1. Examples of relaxation rates of some vibrational states with different collisional partners. Reactions labelled in bold correspond to V-T processes, the others to V-V processes.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Rate [s⁻¹ atm⁻¹]</th>
<th>Ref.</th>
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<th>Rate [s⁻¹ atm⁻¹]</th>
<th>Ref.</th>
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<tbody>
<tr>
<td>(R₁) CH₃(v₅) + CH₄ → CH₃(ν₅) + CH₄(v₅)</td>
<td>2.7·10⁸ [1]</td>
<td></td>
<td>(R₂) O₃(v) + O₂ → O₂ + O₂</td>
<td>6.3·10⁶ [3]</td>
<td></td>
</tr>
<tr>
<td>(R₂) CH₄(ν₅) + CH₄ → CH₄(ν₅) + CH₄</td>
<td>8·10⁹ [1]</td>
<td></td>
<td>(R₃) O₃(v) + N₂ → O₂ + N₂</td>
<td>4.10 [3]</td>
<td></td>
</tr>
<tr>
<td>(R₁) CH₃(ν₅) + O₂ → CH₃(ν₅) + O₃(v)</td>
<td>-3·10⁷</td>
<td></td>
<td>(R₄) O₃(v) + H₂O → O₂ + H₂O</td>
<td>1.1·10⁶ [3]</td>
<td></td>
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<tr>
<td>(R₅) CH₄(ν₅) + O₂ → CH₄(ν₅) + O₂</td>
<td>1.3·10⁷ [2]</td>
<td></td>
<td>(R₆) O₃(v) + CH₄ → O₂ + CH₃(v₅)</td>
<td>2·10⁷ [4]</td>
<td></td>
</tr>
<tr>
<td>(R₇) CH₃(ν₅) + N₂ → CH₃(ν₅) + N₂</td>
<td>8·10⁹ [1]</td>
<td></td>
<td>(R₈) O₃(v) + CH₄ → O₂ + CH₃</td>
<td>1·10⁷ [5]</td>
<td></td>
</tr>
<tr>
<td>(R₉) N₂(v) + N₂ → N₂ + N₂</td>
<td>1.0 [3]</td>
<td></td>
<td>(R₁₀) O₃(v) + He → O₂ + He</td>
<td>2.3·10⁴ [6]</td>
<td></td>
</tr>
</tbody>
</table>
2. EXPERIMENTS

A fibre-coupled PA sensor based on a 1.65-μm DFB laser and a resonant PA cell operated on its first longitudinal resonance around 1 kHz has been developed for CH₄ detection [7]. Methane has been measured in different buffer gases using this set-up, in order to emphasize the importance of molecular relaxation processes. Whereas a sub-ppm detection limit has been achieved for CH₄ in dry nitrogen, the performances are reduced by more than one order of magnitude when CH₄ is measured in pure oxygen, or even in dry N₂-O₂ mixtures containing more than 10% of O₂, as shown in Figure 1. Furthermore, the PA signal measured in O₂ does not present the usual linear dependence on the CH₄ concentration, but shows a quadratic variation, as illustrated in Figure 2. Finally, when a few percent of helium is added to O₂, the CH₄ PA signal retrieves the usual linear dependence on the gas concentration and the detection sensitivity becomes comparable to the value obtained for CH₄ in N₂. All these observed phenomena are related to molecular relaxation effects. They can be explained by the deactivation steps of the CH₄ excited state and the relaxation rates corresponding to collisions with the different types of surrounding molecules.

3. DISCUSSION

3.1 CH₄ relaxation in O₂

The CH₄ molecule has four vibrational modes: two bending vibrations ν₂ (asymmetric) and ν₄ (symmetric), and two stretching vibrations ν₁ (symmetric) and ν₃ (asymmetric). The first excited bending levels ν₂ and ν₄ are located at 1533 and 1311 cm⁻¹, respectively, and the stretching levels ν₁ and ν₃ at 2917 and 3019 cm⁻¹, respectively, approximately two times higher than the two former ones. Consequently, the vibrational energies of methane appear as clusters (called polyads) of states interacting together through Fermi or Coriolis resonance. Due to the strong interactions existing between the states in a given polyad, the spectroscopic analysis of any vibrational state needs to take into account simultaneously all the states of the polyad to which the level belongs [2]. This characteristic is important for the understanding of the collisional relaxation of methane. In particular, very fast energy transfers (in the nanosecond range) occur between the states of the same polyad (intermodal transfer) for both CH₂-CH₄ and CH₃-O₂ collisions. The energy transfer between two polyads essentially occurs via the exchange of one ν₄ vibrational quantum (see reactions R₁ and R₃ in Table 1). For CH₄-O₂ collisions, the de-excitation of the CH₄ molecule is accompanied by a resonant excitation of the first vibrational state of oxygen, O₂ (ν), due to its close matching with the level.
For weak CH$_4$ concentrations, the transfer to the lower polyads occurs only via the resonant process $R_3$. Therefore, the whole 1.65-µm laser energy initially absorbed in the CH$_4^*(2ν_3)$ state is eventually stored in the O$_2^*(ν)$ vibrational state. According to Table 1 (see reaction $R_9$), this state has a relaxation rate that is much smaller than the laser modulation frequency (1 kHz), so that no PA signal is coherently generated with the laser modulation. This explains the very small PA signal observed when measuring CH$_4$ in O$_2$. The situation is totally different for CH$_4$ in N$_2$, as the first vibrational state of N$_2$ has a larger energy than CH$_4^*(ν_4)$ level, so that no V-V coupling can occur.

### 3.2 Quadratic dependence of the PA signal to CH$_4$ concentration

Whereas the collisional V-T relaxation of O$_2^*(ν)$ is very slow for O$_2$-O$_2$ collisions, it is more than four orders of magnitude faster for O$_2$-CH$_4$ collisions (see reaction $R_{11}$ in Table 1). So even for small CH$_4$ concentrations, the relaxation time of O$_2^*(ν)$ state is reduced when increasing the CH$_4$ concentration.

The relaxation rate of O$_2^*(ν)$ state in a O$_2$-M mixture changes with the concentration $C_M$ of the collisional partner $M$ according to:

$$\tau_{O_2}^{-1} = (1 - C_M)\tau_{O_2,0}^{-1} + C_M\tau_{O_2,M}^{-1}.$$  

(1)

For small CH$_4$ concentrations, a linear variation of the relaxation rate is obtained ($y = ax+b$), as shown in Figure 3. This produces a linear increase of the PA signal, as the ratio of the relaxation time to the modulation period is reduced. This linear dependence of the PA signal only results from the relaxation process. In addition to it, the PA signal also presents the usual linear variation on the gas concentration ($y = ax$), due to the increasing amount of energy absorbed in the sample. Therefore, a double linear dependence of the PA signal to the CH$_4$ concentration occurs, which results in the quadratic variation experimentally observed ($y = ax^2+bx$).

### 3.3 Effect of the presence of helium in the buffer gas

As methane, helium is an efficient collisional partner for the vibrational relaxation of O$_2^*(ν)$ (see reaction $R_{12}$ in Table 1). In a O$_2$-He mixture, the O$_2^*(ν)$ relaxation rate changes with helium concentration according to Eq. (1). With the addition of 10% of helium, the O$_2^*(ν)$ relaxation time decreases from 20 ms to 0.4 ms, which is smaller than the laser modulation period. Consequently, the absorbed laser energy fully contributes to the generation of the PA signal and the usual behaviour is again observed (linear variation of the PA signal with the CH$_4$ concentration).

### 4. CONCLUSION

The importance of collisional relaxation processes in the PA signal generation has been demonstrated in the particular case of CH$_4$ detection in dry O$_2$. A resonant coupling between CH$_4^*(ν_4)$ and O$_2^*(ν)$ vibrational states associated to the small relaxation rate of the O$_2^*(ν)$ excited state lead to an important loss of sensitivity in comparison to the case of CH$_4$ in N$_2$. A totally unusual quadratic variation of the PA signal with the CH$_4$ concentration has also been observed and explained. Finally, a sensitive CH$_4$ detection in dry O$_2$ with a linear response has been demonstrated by the addition of a few percent of helium to the gas mixture, which accelerates the vibrational relaxation of the O$_2^*(ν)$ excited state.
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