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Raman lidar for hydrogen gas concentration monitoring and future radioactive waste management

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Abstract: A multi-channel Raman lidar has been developed, allowing for the first time simultaneous and high-resolution profiling of hydrogen gas and water vapor. The lidar measures vibrational Raman scattering in the UV (355 nm) domain. It works in a high-bandwidth photon counting regime using fast SiPM detectors and takes into account the spectral overlap between hydrogen and water vapor Raman spectra. Measurement of concentration profiles of H2 and H2O are demonstrated along a 5-meter-long open gas cell with 1-meter resolution at 85 meters. The instrument precision is investigated by numerical simulation to anticipate the potential performance at longer range. This lidar could find applications in the French project Cigéo for monitoring radioactive waste disposal cells.

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1. Introduction

Long-term management of radioactive wastes is an important scientific and society concern. In France, reversible deep geological disposal is under study as a solution for high-level and intermediate level long-lived radioactive wastes. Since the 2006 planning act, the French...
National Radioactive Waste Management Agency (Andra) is in charge to design and build an industrial disposal site. In this project called Cigéo, nuclear waste packages are expected to be buried 500 meters below the ground inside horizontal galleries of a few hundred meters long (disposal cells, into the dark). Some waste packages generate hydrogen gas ($\text{H}_2$) by radiolysis. The release rate, expected to be slow (1.6 mol/package/year), may occur during a long time. To ensure that $\text{H}_2$ concentration remains well below the lower explosive limit (4% in air), it is important to develop means for profiling $\text{H}_2$ concentration within disposal cells’ atmosphere. Performed at high spatial resolution, such monitoring could also highlight potential deterioration of waste packages by detecting unexpected local release rates of $\text{H}_2$. The lidar technique has the potential to perform non-intrusive measurement, which could, in radioactive environment, prove very useful compared to in situ sensors [1,2].

As a diatomic homonuclear molecule, $\text{H}_2$ has no dipole moment that could create a strong optical absorption in the infrared region. The absorption spectrum of $\text{H}_2$ is limited to vibrational bands of very weak electric quadrupole transitions. Absorption-based detection techniques for $\text{H}_2$ therefore require specific cavity-enhanced setups like Cavity Ring Down Spectroscopy [3], so open-path remote sensing techniques such as Differential Absorption Lidar (DIAL) cannot be used. Nevertheless, $\text{H}_2$ has a strong Raman cross section and Raman lidar is recognized as one of the most appropriate technique for remote $\text{H}_2$ detection [4]. $\text{H}_2$ measurement by Raman lidar has already been demonstrated in the past for pure $\text{H}_2$ point leaks at short range [5]. However, the Shikoku Research Institute system is designed to probe a limited close area and does not allow monitoring over hundreds of meters. Moreover, it disregards the potential impact of water vapor on $\text{H}_2$ measurements (discussed in Sec.2.1).

Detection is often challenging in Raman lidar due to intrinsic signal weakness. In the Cigéo scenario, low $\text{H}_2$ concentration (<1%) must be measured and this makes photon detection even more difficult. For this reason the mentioned overlap between $\text{H}_2$ and $\text{H}_2\text{O}$ spectra cannot be ignored, and simultaneous profiling of $\text{H}_2$ and $\text{H}_2\text{O}$ must be performed. Our goal is also to monitor the atmosphere of future disposal cells with a high spatial resolution (at least 3m gates) in order to distinguish successive waste packages. It is thus crucial to design a very efficient, sensitive, fast and robust detection stage.

In this paper, we first recall Raman lidar measurement basics, highlighting $\text{H}_2$ detection specificities. Then we detail the lidar design, and we report on first experiment results for concept demonstration on hydrogen gas and water vapor. Finally, we conclude on the expected performance at longer range for our system.

2. Measurement principle

2.1 Vibrational-rotational Raman scattering of $\text{H}_2$, $\text{H}_2\text{O}$, $\text{O}_2$ and $\text{N}_2$

Scattering of light in the atmosphere returns various signals dependent on scattering sources. Aerosols yield strong elastic scattering, whereas air’s molecules yield elastic Cabannes scattering, pure rotational Raman (RR) scattering, and vibrational–rotational Raman (VRR) scattering [6]. Measurement of the Stokes VRR scattering is chosen for the present Raman lidar system. In that case, scattered wavelengths are red-shifted with respect to the excitation wavelength by the quantity $\Delta \tilde{\nu} = \tilde{\nu}_s - \tilde{\nu}_i = \Delta E / hc$ characteristic of the molecule ($\tilde{\nu}_i$ and $\tilde{\nu}_s$ are wavenumbers (in cm$^{-1}$) of excitation incident light and Raman scattered light respectively, $\Delta E$ is the energy difference between incident and scattered photons, $h$ is the Planck’s constant and $c$ the velocity of light). For diatomic homonuclear molecules like $\text{H}_2$, $\text{O}_2$ and $\text{N}_2$, VRR transitions consist of a central Q branch with a wavenumber shift $|\Delta \tilde{\nu}| = \tilde{\nu}_m$ surrounded on both sides by O and S branches. For the latter, the wavenumber shift is (adapted from [7]):

$$|\Delta \tilde{\nu}| = \tilde{\nu}_m + 2B(J+3) - D[(2J+3)(2J+5)]$$ with $J = 0,1,2,... \quad (1)$

$$|\Delta \tilde{\nu}| = \tilde{\nu}_m - 2B(2J+1) + D[(2J-1)(2J+1)]$$ with $J = 2,3,4,... \quad (2)$
where \( \nu_v \) is the specific vibrational wavenumber of the molecule, \( B_1 \) and \( D_1 \) are respectively the specific rotational and stretching constant of the vibrational level, and \( J \) is the rotational quantum number corresponding to each rotational level.

The measured intensity of these VRR transitions for an observation at 180° is proportional to the backscatter coefficient (in m\(^{-1}\cdot\text{sr}^{-1}\)). This coefficient is given by \( \beta_\nu = N \cdot (d \sigma / d \Omega) \), where the molecule number density \( N \) (in m\(^{-3}\)) is multiplied to the differential Raman scattering cross section (in m\(^2\cdot\text{sr}^{-1}\)) for scattering at 180°. Figure 1 shows the computed backscattering spectra for \( \text{O}_2, \text{N}_2, \text{H}_2\text{O} \), and \( \text{H}_2 \) for a case study gas mixture. The calculation results from [6,7] with parameters data taken from [6,8–10]. For \( \text{H}_2 \) concentration measurement with a 355 nm laser excitation, the Q branch scattering at 416.1 nm is used along with the two first S-lines. The reference signal is given by \( \text{N}_2 \) scattering centered at 386.7 nm. The important distance between transitions lines of \( \text{H}_2 \) compared to other molecules is explained by the high value of \( B_1 \) for \( \text{H}_2 \) (due to a lower reduced mass value and a shorter internuclear distance).

A spectral overlap between \( \text{H}_2 \) and \( \text{H}_2\text{O} \) Raman spectra is visible on Fig. 1. This overlap could induce a bias on the hydrogen concentration calculation. Therefore a third channel is needed in the Raman lidar system to also monitor water vapor concentration and correct the bias. This third channel is centered on the Q branch of water vapor at 407.5 nm. The few \( \text{H}_2 \) O-lines inside the pass band of the water vapor filter are not sufficiently intense (compared to the sum of all VRR \( \text{H}_2\text{O} \) lines) to create a notable bias on the water vapor channel.

2.2 Mixing ratio retrieval

In photon-counting mode, the number of photons measured per meter \( N_i(z) \) from the Raman scattering of a gas species \( i \) at wavelength \( \lambda_{R,i} \) from distance \( z \) is given by the lidar equation:

\[
N_i(z) = K_i \beta_{i_\nu}(z) \gamma(z) T_{\lambda_L}(z) T_{\lambda_{R,i}}(z)
\]

where \( K_i \) comprises all range-independent system parameters, \( \gamma(z) \) is the overlap function between the transmitted laser beam and the receiver field of view, \( T_{\lambda_L} \) and \( T_{\lambda_{R,i}} \) are atmospheric transmissions at the laser wavelength \( \lambda_L \) and at Raman wavelength. For high signal intensities the receiver response is no longer linear. The true number of photons \( N_i \) is then given by \( N_i = N_m / (1 - N_i t) \) (non-paralyzable behaviour) where \( N_m \) is the measured number of photons and \( t \) the measured pulse pair time resolution, or dead time.

Concentration measurement by Raman technique requires the measurement of a gas species taken as reference. Nitrogen was chosen here, since \( \text{N}_2 \) concentration in disposal cells is expected to remain very stable. This is not the case for oxygen, whose concentration is...
expected to decrease with time because of corrosion processes. Because of the spectral overlap previously discussed, water vapor in the atmosphere induces an overestimation of hydrogen mixing ratio. This bias is continuously corrected through a third channel dedicated to water vapor measurement. Therefore $H_2$ mixing ratio estimator is given by:

$$x_{H_2}(z) = C \frac{N_{H_2}^0(z) - k N_{H_2}^{\text{true}}(z)}{N_{H_2}^0(z)} \frac{T_{\text{atm}}^{\text{av}}(z)}{T_{\text{atm}}^{\text{av}}(z)}$$  \hspace{1cm} (4)$$

where $C$ is the calibration constant which depends on system characteristics and differential cross sections. $k$ is a correction factor determined through a calibration procedure, that consists in forcing to zero the residual signal measured on the $H_2$ channel in the absence of ‘true’ $H_2$ along the line of sight. The transmission ratio depends on aerosol distribution.

The 1σ error expected on $H_2$ mixing ratio estimation can be written as:

$$\sigma_{x_{H_2}} = \sqrt{\frac{V(N_{H_2}^0) + k^2 V(N_{H_2}^{\text{true}})}{(N_{H_2}^0 - k N_{H_2}^{\text{true}})^2} + \frac{V(N_{H_2}^0)}{N_{H_2}^0}}$$  \hspace{1cm} (5)$$

where $V(N_{H_2}^0)$ is the variance of $N_{H_2}^0$, for which the shot noise from signal and background radiation measured on each channel, along with the count correction are taken into account.

3. Simultaneous hydrogen and water vapor measurements

3.1 Raman lidar design

The emitter is a 355 nm laser source (Quantel – Ultra 100) that emits pulses of 30 mJ and 5.5 ns FWHM at 20 Hz. The Raman signal is collected via a Newtonian telescope of 152 mm diameter and 609 mm focal length. The received signal is coupled into a multimode fiber and directed towards the subsystem dedicated to spectrum analysis.

This multi-channel spectral analyzer, detailed in [11], is used to select VRR scattering spectral bands of interest seen in Sec.2.1. It uses bandpass optical filters ($\Delta \lambda$=15nm) placed in front of each detector, and a long-pass filter eliminating Rayleigh-Mie scattering at 355nm. There is a minimum of 100 dB attenuation at 355 nm on each channel (70dB is needed to obtain a Rayleigh scattering 100 times weaker than a 2%-H$_2$ Raman scattering). Dichroic mirrors are used for channel spatial separation. Silicon PhotoMultipliers detectors (SiPMs – SensL – 30035), working in photon-counting mode, have been chosen for the detection. This technology, still little used in lidar to date [12], suits well for this application [11]. It brings to the setup simplicity (no high voltage needed), high detectivity (photon counting), high speed (500 MHz count rate) and robustness (short dark state and undamaged by ambient light).

3.2 Experimental layout

Long-range measurements of $H_2$ and water vapor profiles have been performed for system concept demonstration. A 5-meter-long open gas cell, 10 cm diameter, has been developed to simulate $H_2$ release between waste packages. In order to redirect the laser beam in a more convenient area, a mirror has been placed at 50 m as shown in Fig. 2. The effective distance to the gas cell reached 85 m. Bottles containing either pure dry air or a mixture of $H_2$ at 2% in dry air were available for gas-release experiments (through the center of the tube).

For each measurement, in order to homogenize gas composition inside the tube, the latter was first filled with gas with the two openings blocked. Then obstructions were removed, leaving way for the laser beam, and Raman lidar acquisition started while maintaining the gas flow inside the tube. SiPM dark counts and background radiation contributions, measured between two lidar shots, are removed through signal processing. Mixing ratio calculations and displays were performed in real time by a computer. In these experiments, the laser energy
was reduced to 5 mJ to avoid signal saturation, and the telescope depth of focus was optimized at the center of the tube.

3.3 Results on hydrogen and water vapor

Three successive experiments have been conducted on a winter night (T = 4°C, RH = 84%) and the calculated mixing ratios profiles for H₂ and H₂O are shown in Fig. 3. The first experiment (blue line) was made with the natural humid atmosphere (no gas released). The second experiment (green line) was made with pure dry air released inside the tube, and the third one (red line) with hydrogen gas at 2% in dry air. For water vapor profiles, a first-order calibration has been performed using an in situ hygrometer (Rotronic – HygroPalm HP23). H₂ profiles are given in arbitrary unit because no calibration sensor was available (currently under development). The water vapor cross talk is removed using a correction factor of \( k = 0.33 \) (determined 6 days before by a calibration procedure as explained in Sec. 2.2). All profiles are shown for 1 m resolution and 1 min accumulation time (1200 lidar shots). Drawn error bars show the expected 1σ standard deviation of H₂ mixing ratio given by Eq. (5) (a similar calculation is done for H₂O). As an example, in the 2% H₂ in dry air case study (red curves), the average numbers of photons/meter/shot for the measurement point at center of the tube (\( z = 84 \) m) are \( N_{tN} = 20.5 \), \( N_{HOtN} = 20.05 \) and \( N_{He} = 21.9 \).

The blue line shows no discontinuity inside and outside the tube, which means that the air composition without gas released is uniform along the line of sight. When the tube air is replaced with dry air a large decrease (95% extinction) of the water vapor mixing ratio inside the tube is observed as expected. Meanwhile there is no change on the H₂ mixing ratio. When the tube is filled with 2% H₂ in dry air, the same decrease is observed on the H₂O profile, but in the same time, a strong increase of the H₂ mixing ratio is observed. One can notice that the
H$_2$ signal is at least one order of magnitude above the level of signal fluctuation. Regarding the potential limit of detection, this result is therefore promising. For further works, the measurement range could be increased with additional mirrors, since the laser energy here was limited to 5 mJ, while 25 mJ (considering losses) are available. Range resolution could also be slightly degraded from 1 m to 3 m (size of a waste package) to reduce measurement fluctuations by spatial averaging if necessary. Finally the accumulation time (only 1 minute here) could be significantly increased (up to 1h) inasmuch as the experiment conditions and instrument behaviour remain stable enough along the integration time.

4. Expected LOD at long range

The expected limit of detection (LOD) of H$_2$, defined here as the expected 3$\sigma$ error (Eq. (5)) around a zero-H$_2$ mixing ratio, has been calculated in conditions comparable to long disposal cells. Its evolution at 500 m over accumulation time is shown on Fig. 4. All the available laser energy (25 mJ), a 3 m range resolution and a background noise similar to the previous experiment are assumed to be used for this calculation. Evolutions for two humidity conditions are drawn, humid (blue line, 0.7% of H$_2$O same as the real experiment) and dry (green) atmosphere. For 1 minute accumulation, the LOD of H$_2$ is about 0.2%, i.e. 10 times below the concentration used for concept demonstration, which looks consistent with our experimental observations. Assuming stable accumulation processes the 0.05% limit could be reached in 11 minutes in dry air and in 19 minutes in humid atmosphere.

![Fig. 4. Evolution over time of the LOD of H$_2$ at 500 m in humid (blue) and dry (green) atmosphere (laser energy of 25 mJ and a range resolution of 3m are assumed here).](image)

5. Conclusion and outlooks

Simultaneous high resolution profiles of hydrogen gas and water vapor have been performed along a 5-meter-long open gas cell at 85 meters with our multi-channel Raman lidar. This system could contribute to security, safety and reversible management of radioactive waste by monitoring future waste disposal cells if the Cigéo project comes into being.

Performance at longer range has been predicted by numerical simulation. A 0.05% LOD in dry air should be achieved for H$_2$ at 500 m in 11 minutes, insofar as the experiment conditions and instrument behaviour remain stable enough along the integration time.

In future works, the lidar calibration for quantitative hydrogen gas measurements will be performed using a local H$_2$ sensor inside the tube (currently under development by Andra). The lidar range will also be increased to several hundred meters, in order to demonstrate the measurement feasibility in long disposal cells galleries and confirm the expected detection limit of our system.

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