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# *In situ* continuous monitoring of dissolved gases (N<sub>2</sub>, O<sub>2</sub>, CO<sub>2</sub>, H<sub>2</sub>) prior to H<sub>2</sub> injection in an aquifer (Catenoy, France) by on-site Raman and infrared spectroscopies: instrumental assessment and geochemical baseline establishment†

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The establishment of a baseline of gases from an aquifer appears to be an essential prerequisite for monitoring and securing underground storage operations such as the storage of carbon dioxide (carbon capture and storage: CCS), methane or hydrogen. This study describes an innovative metrological technique dedicated to the *in situ* and continuous quantification of dissolved gases (CO<sub>2</sub>, O<sub>2</sub>, N<sub>2</sub>, CH<sub>4</sub> and H<sub>2</sub>) in a shallow aquifer, on the site of Catenoy (Paris Basin) with a water table at a depth of 13 m. Monitoring was carried out from May 7, 2019 to November 19, 2019, before the simulation of H<sub>2</sub> injection. Gases as vapors were collected from the aquifer through a nine-meter long, half-permeable polymer membrane positioned below a packer in a 25-meter deep well. Collected gases were analyzed simultaneously at the surface by fiber Raman (CO<sub>2</sub>, O<sub>2</sub>, N<sub>2</sub>, CH<sub>4</sub> and H<sub>2</sub>) and infrared sensors (CO<sub>2</sub>). Gas concentrations were determined from Raman and infrared data, and then converted into dissolved concentrations using Henry's law. The dissolved gas concentrations were about constant over the 6 months period with average values of 31–40 mg L<sup>-1</sup> (CO<sub>2</sub>), 8 mg L<sup>-1</sup> (O<sub>2</sub>), 17 mg L<sup>-1</sup> (N<sub>2</sub>), and 0 mg L<sup>-1</sup> (H<sub>2</sub>, CH<sub>4</sub>) indicating a very low variability in the aquifer. This is believed to allow for rapid detection of any possible abnormal concentration variation, in particular linked to an accidental arrival of gases such as hydrogen. Such an online gas measurement system can be deployed as is on any site type of underground storage without any need for adaptation.

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## Introduction

Fossil fuels (oil, gas and coal) have been the primary sources of energy for over a century. However, the combustion of these carbon-bearing materials produces greenhouse gases (GHG, *e.g.* CO<sub>2</sub>, CH<sub>4</sub>, N<sub>2</sub>O), which cause environmental issues when emitted into the atmosphere. Over the past several decades,

surging amounts of these GHGs have been emitted, leading to global concerns about the Earth's environment.<sup>1</sup> For instance, GHG emission is considered the first cause of global warming that has many adverse impacts on our society as warned by the Intergovernmental Panel on Climate Change.<sup>2</sup> The replacement of fossil fuels with renewable carbon-free energies is considered the most critical solution to achieve balance between the environment, economy and public health.<sup>3–5</sup> Among promising renewable energy sources (including solar, hydro, wind power, biomass, and geothermal energy), molecular hydrogen (H<sub>2</sub>) is the most frequently mentioned. It can be produced from these sources and used as an energy vector.<sup>6</sup> Currently, 60 million tons of H<sub>2</sub> are produced from fossil resources ("grey H<sub>2</sub>") each year in the United States and Europe.<sup>7,8</sup> H<sub>2</sub> production from fossil resources can be coupled with carbon capture and storage ("blue H<sub>2</sub>"). However, H<sub>2</sub> production from these sources is not carbon neutral. By further developing this energy as part of the energy transition, global needs may increase rapidly in the coming decades with production of H<sub>2</sub> in carbon neutrality as by electrolysis of renewable energies ("green H<sub>2</sub>").<sup>7–10</sup> Currently, the International Energy Agency (IEA) brings together twenty

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† Electronic supplementary information (ESI) available: Appendix 1: schematic representation of the continuous gas monitoring device in the Catenoy site (according to Derrien and Navelot, 2019). Appendix 2: completion module installation in PZ2 ter in the Catenoy site with the packer and the gas circulation lines (Solexperts' photography, according to Derrien and Navelot, 2019). Appendix 3: mathematical development for the calculation of the peak area and the quantitative calculation of the gas concentration. Appendix 4: determination of the optical characteristics: signal-to-noise ratio (SNR) and the instrumental derivative factor (*F*). See DOI: 10.1039/d1ay01063h



countries to avoid the increase in CO<sub>2</sub> emissions (300 Gt estimated up to 2050)<sup>6</sup> and thus to study pure and applied research subjects related to the H<sub>2</sub> development until 2050.<sup>6,10</sup> The part of renewable energy in energy consumption, involving H<sub>2</sub>, was in 2019 expected to increase by 51% in India, 20.3% in Europe, and 16% in China by 2023.<sup>11</sup> The Hydrogen Council was created in 2017 in Davos (Switzerland) to join forces with the IEA in order to encourage the main actors for the development of this sector.<sup>12–14</sup> This Council currently estimates that hydrogen will supply 18% of global energy demand.<sup>13</sup> In practical operations at the international, European, and local levels, the massive use of H<sub>2</sub> is strongly dependent on the capacity to develop H<sub>2</sub> safety and sustainable underground storage<sup>15–19</sup> unlike conventional storage methods such as compression, liquefaction, physisorption, metallic hydrides or complex hydrides.<sup>10,20–22</sup>

For all types of gas underground storage (deep aquifers, saline caverns, mined caverns and depleted fields),<sup>17,23,24</sup> effective monitoring systems should be set up in the surrounding geological media from deep (–1000 m) to surface for timely detection of potential H<sub>2</sub> leakage, thus enabling a quick alert and response to adverse, sometimes disastrous, impacts (*e.g.* for the studies dealing with saline caverns where the permeability of the rocks is very low.<sup>15,16,19,23–27</sup> In any storage operations, project integration inside the territory requires securing the site before (measuring the gas emission level of the site before injection which is equal to baseline definition), during and after operations (survey monitoring). These operations are in direct link with geophysical and geochemical monitoring methods adapted to the studied site.<sup>28–39</sup> In the last 30 years, numerous studies have been devoted to metrological developments for gas monitoring mainly in the context of underground disposal of nuclear wastes<sup>40</sup> and greenhouse gas underground storage including CCS (Carbon Capture and Storage) applications.<sup>33,36,41–45</sup> Most of these developments concern the temporal evolution of gas concentrations (mainly CO<sub>2</sub> and CH<sub>4</sub>) in the hydrosphere, soil and atmosphere.<sup>46–51</sup> The successive developments have aimed to optimize all measurement systems, to simplify experimental protocols, and to establish gas calibration curves for accurate prediction and simulation of gas dissolution, diffusion and transfer in the surrounding geological medias.<sup>52–61</sup>

The most commonly adopted analytical setups for the purpose of *in situ*, long-term and continuous gas monitoring comprise three modules: specific completion, gas circulation tubing and sensor(s) compartment, installed in a dedicated borehole from the shallow surface down to –600 m deep.<sup>32,50,53,56,57,60,62</sup> Raman and infrared spectrometers have proven to be suitable sensors: these techniques have been constantly developed from the laboratory measurement scale to the continuous field measurement scale for soil and underground gas monitoring.<sup>32,50,60</sup> The two techniques are complementary (at similar cost) as Raman spectroscopy can detect a lot of gases (*e.g.* N<sub>2</sub>, O<sub>2</sub>, CO<sub>2</sub>, CH<sub>4</sub> and H<sub>2</sub>) but with a low sensitivity while only CH<sub>4</sub> and CO<sub>2</sub> can be detected by infrared spectroscopy but with higher sensitivity.<sup>63</sup> However, if some studies referred to *in situ*, discontinuous sampling monitoring of dissolved gases in freshwater aquifers such as in Montana, USA

and in Oise, France,<sup>64–66</sup> very few referred to *in situ* continuous gas monitoring in this environment. The most recent study refers to the work of Petit *et al.* (2021)<sup>67</sup> which consists of the injection of dissolved CO<sub>2</sub> into a carbonate freshwater aquifer at 21 m deep in a pilot site in Gironde, France. The dissolved CO<sub>2</sub> concentrations were monitored five days before the injection by a probe immersed in several wells.

The aim of this article is to present a pilot study to test the applicability and robustness of continuous and *in situ* dissolved gas monitoring by Raman and infrared spectroscopies. The monitoring device was tested under real conditions in a shallow aquifer in the Paris basin (Catenoy, France) over a period of more than 6 months to establish a real gas reference before any type of anthropogenic disturbance such as the injection of H<sub>2</sub> to simulate a leak from an underground storage. For this, a completion module comprising a very long semi-permeable membrane allowing the extraction of dissolved gases from the water table was placed in a dedicated borehole. The gases collected were analyzed simultaneously by a fiber Raman sensor (CO<sub>2</sub>, O<sub>2</sub>, N<sub>2</sub>, CH<sub>4</sub>, H<sub>2</sub>O and H<sub>2</sub>) and a Fourier transform infrared spectrometer (FTIR) (CO<sub>2</sub>). The quantitative determination of dissolved gases is explained as well as the evaluation of the technical capacity between Raman and infrared spectroscopy. Knowing such a dissolved gas baseline is a preamble to follow-up to a future dissolved H<sub>2</sub> injection experiment. Moreover, the transposition of this monitoring protocol for other underground storage facilities is also discussed.

## Equipment and methods

### The injection pilot site (Ineris, Catenoy, France)

The establishment of a geochemical baseline for major gases and H<sub>2</sub> was carried out in a shallow aquifer located in the experimental site in Catenoy within the Paris Basin (Hauts-de-France area, Oise), 20 km north of the Ineris applied research center. This site has been instrumented since 2012, and has been used for several similar injection experiments with helium (He) and carbon dioxide (CO<sub>2</sub>) in the context of former research projects, CIPRES and CO<sub>2</sub>QUEST.<sup>65,66,68</sup>

The aquifer is unconfined within a chalky reservoir layer of Senonian (Cretaceous) comprising 95% of calcite, and stretches over a total thickness of around 220 m. It is overlain by 6 to 7 m thick Bracheux sands (Tertiary), which are composed of clay sands, fine flint sands, silty sands, sandy silts and silts. The water table of the aquifer is located at an average depth of 13 m, below which water flows in the WSW-ENE direction at around 3 to 10 meters per day<sup>66</sup> (Fig. 1). The knowledge of the geochemical baseline is important before any gas injection,<sup>66</sup> so the physico-chemical, chemical and hydrological parameters of the aquifer were acquired between October 27, 2018 and November 6, 2019 using mainly in the field two multiparametric probes (“HI 9828” of Hanna instruments and “Quanta” of Hydrolab Corporation).<sup>69</sup> Major hydrological and chemical parameters are summarized below (Fig. 1). The aquifer groundwater is always close to neutrality (pH of 7.3 ± 0.3), oxygenated (dissolved O<sub>2</sub> concentration of 5.4 ± 1.7 mg L<sup>-1</sup>), and oxidizing (positive oxidation–reduction potential (ORP):



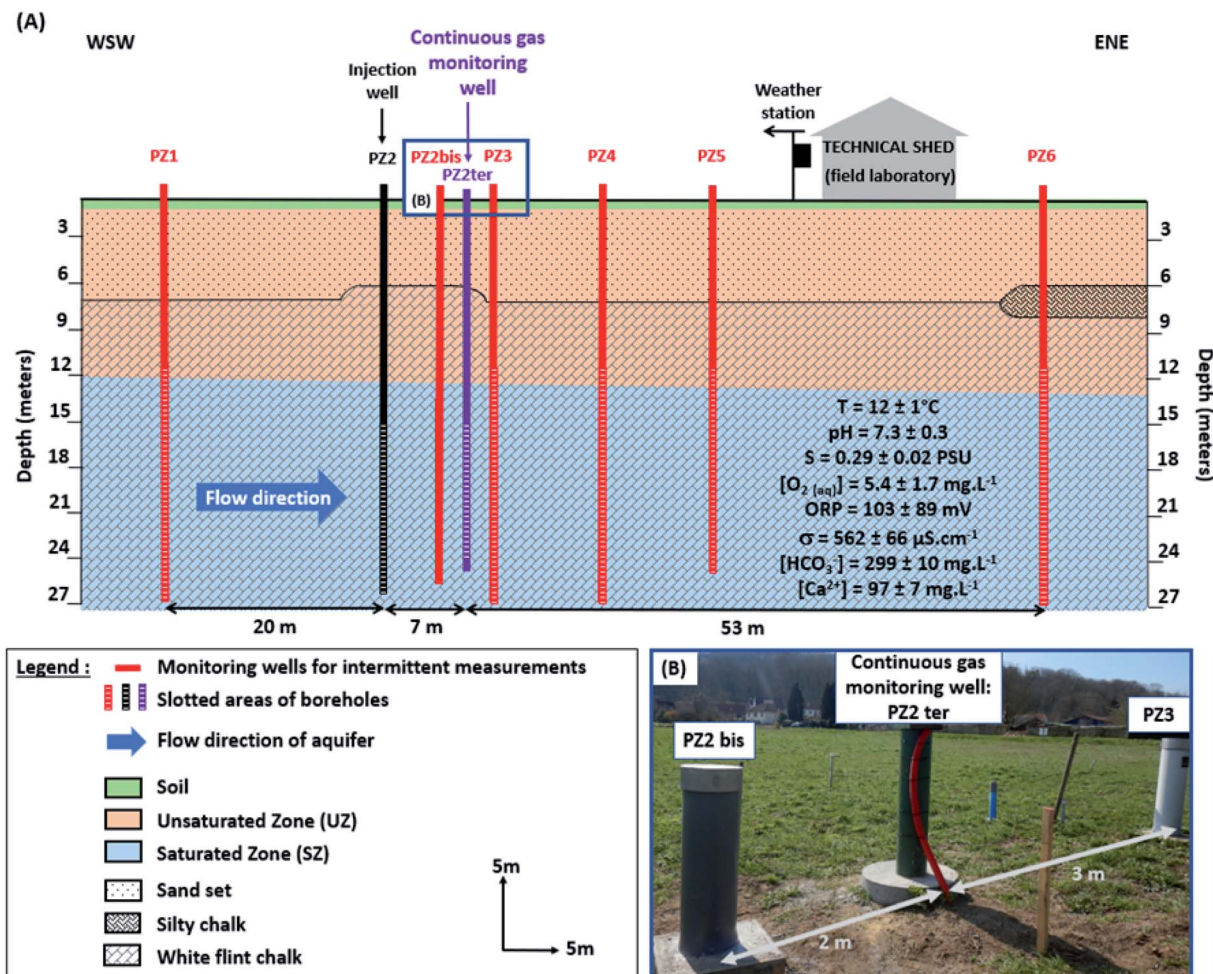


Fig. 1 (A) Schematic vertical section of the Catenoy experimental injection site (geological formations, equipment, injection and monitoring wells) (according to Gombert *et al.* 2014; Lafortune *et al.* 2020<sup>66,69</sup>). For better visibility, the dimensions of the monitoring wells, the weather station and the technical shed are not to scale, and the UZ monitoring wells (boreholes only reaching the UZ) are not represented here. They will be described in a future study. (B) Photograph of the location of the continuous gas monitoring well (PZ2 ter) in relation to two other monitoring wells for discontinuous measurements.

103 ± 89 mV). The aquifer is moderately mineralized (electrical conductivity ( $\sigma$ ) of 562 ± 66  $\mu\text{S cm}^{-1}$ ), with an average temperature ( $T$ ) of 12 ± 1 °C and a salinity ( $S$ ) of 0.29 ± 0.02 PSU. This water exhibits a dominant bicarbonate–calcium facies ( $\text{HCO}_3^-$  concentration of 299 ± 10  $\text{mg L}^{-1}$ ) with a slight alteration of sulfate and nitrate ions ( $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$  concentration of 28 ± 3 and 33 ± 3  $\text{mg L}^{-1}$  respectively). Ammonium ions are also present in water but at very low concentrations (0.10 ± 0.13  $\text{mg L}^{-1}$ ).<sup>69</sup> The concentration of metallic trace elements (Fe and Mn) is around 0.99  $\text{mg L}^{-1}$  (Fe) and 0.11  $\text{mg L}^{-1}$  (Mn).<sup>69</sup>

The experimental site is equipped with one injection well (PZ2 in Fig. 1A) and seven monitoring wells (PZ1, PZ2 bis, PZ2 ter, PZ3, PZ4, PZ5 and PZ6 in Fig. 1A). All these wells are arranged linearly and parallel to the flow direction over 80 m and reach the saturated zone (SZ) until 27 m deep. The upper limit of the strainers varies from 11 to 15 m in depth depending on the piezometers<sup>66,70</sup> (Fig. 1). The monitoring wells represented in red in Fig. 1A are designed for intermittent sampling of physico-chemical and gas measurements.<sup>66,69,71</sup> In contrast,

PZ2 ter (represented in purple in Fig. 1A and B), which is located 7 m downstream from the injection well (PZ2), is designed for *in situ* and continuous gas measurements. Moreover, the depth in each well was checked manually again in 2019 in order to verify the good operation (absence of mud plugs inside) of the piezometers several years since their last use using a depth probe.

#### Continuous gas measurements by Raman and infrared spectroscopies in a PZ2 ter well

A new 220 mm diameter borehole (PZ2 ter) was rotary drilled to a depth of 25 m. This piezometer was then equipped with a plastic filter between 15 and 24 m deep with a diameter of 80 mm (int.) to 90 mm (ext.). The annulus was filled up with quartz sand, intercalated with two layers of clay balls. The top of the borehole was sealed with cement and bentonite (Appendix 1 in the ESI section†). After the drilling operations, the piezometer was cleaned up by airlift, before installing the completion system.



**In situ and continuous gas monitoring system.** The PZ2 ter well is dedicated to a combined Raman and infrared monitoring device for the continuous determination of dissolved gas concentrations in the saturated zone as shown in Fig. 1. The device consists of three modules: (i) a completion module designed by the company Solexperts (Vandœuvre-lès-Nancy, France) (Fig. 2A and Appendix 2 in the ESI†) connected to (ii) an external circulation module (Fig. 2B) carrying gas which is itself attached to (iii) an optical sensor module consisting of Raman and infrared spectrometers (Fig. 2C). This setup resembles the system installed at the Lacq-Rousse CO<sub>2</sub> storage pilot site to continuously monitor gas concentrations in soils.<sup>32,50</sup> Some technical improvements are made at the Catenoy site to account for application in aquifers and greater depths (25 m vs. 3.4 m). The description of these three different modules is detailed below.

**Completion (module A).** The collection of the gases dissolved in the aquifer required the development of a specific completion module. The completion module consists of, from bottom to top, a nine-meter length collecting chamber, which is isolated by a one-meter length mono-packer system topped with PVC pipes (Fig. 2A). This collecting chamber comprises a semi-permeable polymer membrane to collect the gas dissolved in

the aquifer in vapor form while avoiding water circulation into the chamber. A thermocouple is placed at the bottom end of the chamber for measuring temperature (Fig. 2A). Moreover, Gombert *et al.* (2014) showed the existence of a vertical heterogeneity, which can induce a multichannel flow, governed by a fissure and matrix porosity.<sup>69</sup> Therefore, a long membrane technology (9 m) was favored, which should allow, in a first approach, to average the analytical bias brought by the vertical heterogeneity of the aquifer. The system is first installed with plastic rods in the borehole, and then the packer was inflated (Fig. 2A and Appendix 2 in the ESI†).

Inside this completion system, steel lines cross the entire length of the well (Appendix 2 in the ESI†) allowing gas to circulate in the second module (see the next section).

**Gas circulation module (module B).** A gas circulation module allows the gas to circulate from the completion module to the measuring devices located on the ground, through a closed loop circuit. The module is placed in a cabinet. This module is equipped with a specific gas circulation pump, a gas flow control valve, an atmospheric pressure sensor, a circulation pressure sensor and a temperature sensor. In addition, the gas flow module has a water trap to prevent any water in the lines that could interfere with gas flow. Installed in the

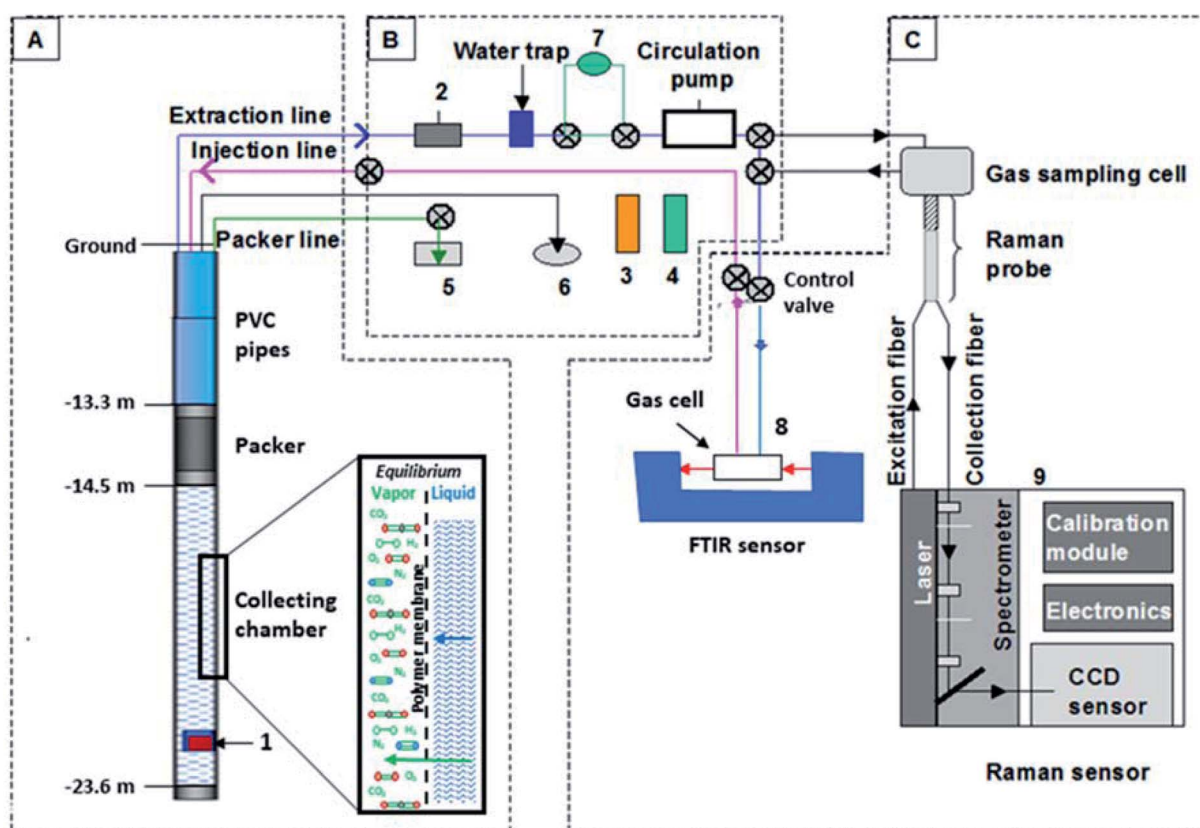


Fig. 2 Experimental device used in the Catenoy site for *in situ* and continuous gas measurements by FTIR and Raman spectroscopies (according to Taquet *et al.*, 2013; Derrien and Navelot, 2019<sup>32,79</sup>). (A) Completion module set up in the PZ2 ter well with a drawing of gas and liquid phase reactions at equilibrium through the membrane (1: temperature sensor); (B) gas circulation module (2: pressure sensor, 3: atmospheric pressure sensor, 4: atmospheric temperature sensor, 5: packer pressure sensor, 6: borehole temperature indicator and 7: fluxmeter); (C) optical sensor module (8: simple compartment FTIR spectrometer BRUKER ALPHA and 9: Kaiser Optical System RXn1 Raman spectrometer).



technical shed, all of this equipment is enclosed in a box sheltered from general humidity and small animals. All the stainless-steel lines from the well to the gas circulation module are buried (~1 m deep) to prevent from heating/cooling. An automatic data acquisition system collects the data, which are sent to a web visualization platform (Fig. 2B).

**Optical sensor module (module C).** The sensor module consists of a Raman spectrometer equipped with a Raman gas probe and an optical fiber and an infrared spectrometer connected to a specific gas cell placed therein (Fig. 2C). The gases are first analyzed by the Raman probe and then 5 seconds later (propagation time) by the infrared cell. The details of the measurements by Raman and infrared spectroscopies are given below.

### Measurement protocol and data processing

**Continuous measurements.** Both infrared and Raman spectrometers worked continuously for six months (from early May to mid-November 2019) to establish the geochemical baseline covering potential seasonal variation and before any voluntary disturbance of the environment.

All Raman spectral data are acquired and recorded by iCRaman software. This software automatically performs data processing by Fourier transform and apodization with a resampling interval of  $1\text{ cm}^{-1}$ . The signals of  $\text{CO}_2$ ,  $\text{CH}_4$ ,  $\text{H}_2$ ,  $\text{O}_2$ , and  $\text{N}_2$  were collected from May 7, 2019 to November 19, 2019, gathering a total of 7966 spectra. In order to optimize the signal, the accumulation number (which corresponds to the number of spectra recorded and then averaged to provide a unique spectrum) and the acquisition duration were optimized during the first weeks of measurement. From May 7 to 23, the measurements were acquired in 4 accumulations of 1 minute, then from May 23 to May 29, they were recorded in 8 accumulations of 1 minute and from May 29 to November 19 in 16 accumulations of 30 seconds. The acquisitions were performed every 30 minutes (sampling frequency) over the full period (from May 7 to November 19, 2019). This whole period was divided into nine sub-periods because of logistical interruptions during measurements (shutdown/restart due to power cut, fuse change, resumption and modification of the acquisition parameters).

The infrared spectra were recorded, *via* OPUS software, from May 7 to November 19, 2019 gathering a total of 3270 spectra. The spectra were acquired every hour from May 7 to October 24, 2019 and then every 30 minutes from October 24 to November 19, 2019. An OPUS macro-command was established to automatically calculate the areas of the bands associated with gas molecules. The whole baseline period was divided into seven sub-periods because of logistical interruptions during measurements (shutdown/restart due to power cut and electronic bugs).

**Data processing.** The Raman and infrared techniques are complementary and well-adapted for quantifying gas concentrations as the peak intensities or band areas are proportional to concentrations. All polyatomic molecules are Raman active, though all gases of interest here ( $\text{CO}_2$ ,  $\text{CH}_4$ ,  $\text{H}_2$ ,  $\text{O}_2$ ,  $\text{N}_2$ ). In contrast, only the vibrations associated with a change of the

dipolar moment are active in infrared spectroscopy. Though  $\text{CO}_2$  and  $\text{CH}_4$  can be detected but not the homonuclear gases such as  $\text{N}_2$ ,  $\text{O}_2$ , and  $\text{H}_2$ .

**Raman spectroscopy measurements.** The Raman spectrometer is an RXn1 Analyzer (Kaiser Optical Systems, Inc.) equipped with a 532 nm Nd-YAG laser for excitation, an optical fiber and a probe head (AirHead™, Kaiser Optical Systems, Inc.). This probe head is composed of a gas cell connected to the gas circuit and an optical device to amplify and collect the Raman signal. The Raman cell is placed inside the gas circulation module box after the water trap to avoid liquid water inside the cell, which would damage the mirrors therein and perturb the measurements. Raman spectra are collected in the wavenumber range of  $100\text{--}4400\text{ cm}^{-1}$  with an instrumental spectral resolution of about  $5\text{ cm}^{-1}$ . For each recorded spectrum, the fundamental vibration bands of the gaseous molecules ( $\text{CO}_2$ ,  $\text{O}_2$ ,  $\text{N}_2$ ,  $\text{CH}_4$  and  $\text{H}_2$ ) are identified based on the literature.<sup>72,73</sup> A very weak peak of water vapor was sometimes observable in the Raman spectra coming out weakly from the background noise ( $\sim 3650\text{ cm}^{-1}$ ). Due to this situation, an assumption was made not to consider the water vapor in the gas phase composition calculated from the Raman data. As a consequence, probably all concentrations of the other gases are a little overestimated. Considering the noise level (due to low laser power), it has no significant effect on the results. The peak area calculation of each gas by iCRaman software is determined integrating the signal between two anchor points on each side of the peaks (total interval around  $20\text{ cm}^{-1}$ ), with a linear baseline drawn between the two anchor points. The integration interval is fixed from  $1376$  to  $1396\text{ cm}^{-1}$  for  $\text{CO}_2$ , from  $1544$  to  $1564\text{ cm}^{-1}$  for  $\text{O}_2$ , from  $2320$  to  $2340\text{ cm}^{-1}$  for  $\text{N}_2$ , from  $2907$  to  $2928\text{ cm}^{-1}$  for  $\text{CH}_4$  and from  $4150$  to  $4165\text{ cm}^{-1}$  for  $\text{H}_2$  (Fig. 3).

It is possible to determine the molar proportion of each gas in a mixture from the peak area and from the Raman relative scattering cross section (RRSCS).<sup>73–76</sup> This last parameter depends on the excitation light wavelength used in the Raman spectrometer and on the wavenumber of each molecule vibration mode.<sup>73,77–80</sup> In this study, the laser is at 532 nm. However, there are no references of the RRSCS of the studied gases at this wavelength. Thus, those approaching this wavelength (514.5 nm) are used to calculate gas concentrations from their respective Raman spectral lines (Table 1) considering that the wavelength correction is negligible between 514.5 nm and 532 nm.<sup>80</sup> As the peak areas of  $\text{CH}_4$  molecules are zero or very negligible over the whole period, they are excluded from the calculation of the gas molar proportion. As explained before, water vapor was also neglected, probably affecting the results by a few percent.

Mathematical development for the calculation of the peak area and the quantitative calculation of the gas concentration is detailed in Appendix 3 in the ESI.†

**Infrared spectroscopy measurements.** The infrared spectra are recorded using a portable infrared spectrometer ALPHA (Bruker Optik GmbH, Ettlingen, Germany). A gas cell with a 5 cm path length equipped on both sides with zinc selenide (ZnSe) windows is placed in the sampling module (Fig. 2C) and connected to the gas circulation module (Fig. 2B). The spectra are acquired with 10 scans in the mid-infrared range ( $4000\text{--}600\text{ cm}^{-1}$ ) with a spectral resolution of  $1\text{ cm}^{-1}$ .



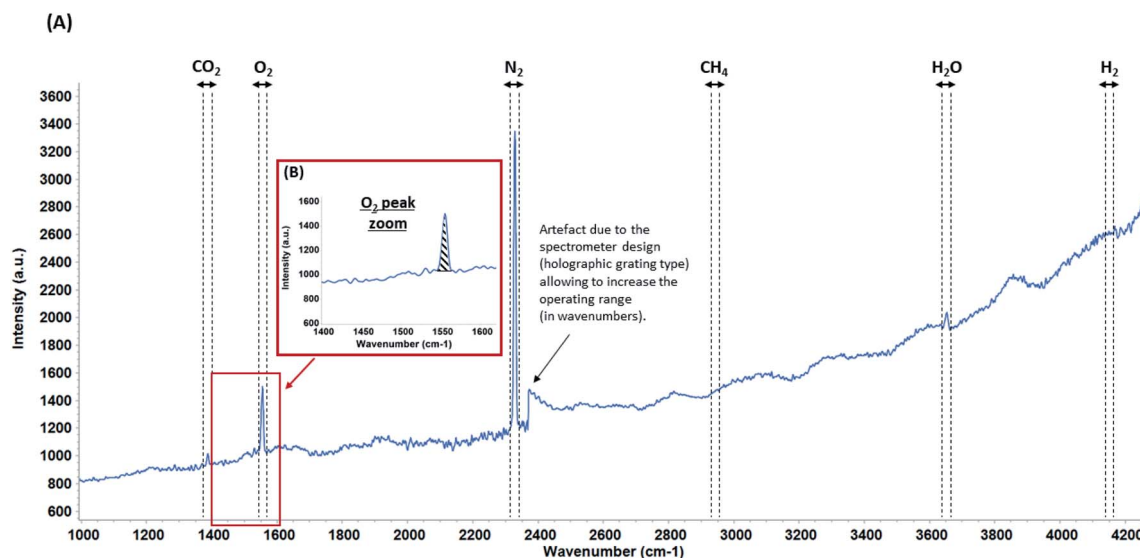


Fig. 3 Example of a Raman spectrum obtained during the baseline period. (A) Spectrum showing the spectral lines position of each dissolved molecule ( $\text{CO}_2$ ,  $\text{O}_2$ ,  $\text{N}_2$ ,  $\text{CH}_4$  and  $\text{H}_2$ ); (B) zoom on the  $\text{O}_2$  peak to show peak area integration; water vapor was sometimes detected  $\sim 3650\text{ cm}^{-1}$ .

**Table 1** Raman relative scattering cross section (RRSCS) of each gas (Schrötter and Klöckner, 1979; Burke, 2001; Garcia-Baonza *et al.* 2012<sup>73,78,80</sup>)

Gas	Peak position ( $\text{cm}^{-1}$ )	Cross-section (RRSCS) <sup>a</sup>
$\text{CO}_2$	1388	1.5
$\text{O}_2$	1555	1.2
$\text{N}_2$	2331	1
$\text{H}_2$	4155	2.3

<sup>a</sup> Determined for a laser at 514.5 nm.

The  $\text{CH}_4$  band expected around  $3015\text{ cm}^{-1}$  is not detected in the infrared spectra. The fundamental vibration band ( $\nu_3$ ) of the  $\text{CO}_2$  molecule (doublet between  $2400$  and  $2220\text{ cm}^{-1}$ ) is identified into the literature (ref. 81 and references therein). The band area calculation of  $\text{CO}_2$  by OPUS software is determined as a function of a wavenumber interval after baseline subtraction. Thus, the selected signal of  $\text{CO}_2$  is integrated between  $2400$  and  $2220\text{ cm}^{-1}$ . The  $\nu_3$  band is characterized by the presence of a doublet whose intensity of the second peak is always lower by a ratio of  $\frac{3}{4}$  compared to that of the first peak. Some spectra respect this characteristic (A to D in Fig. 4) despite interference due to the presence of vapor water (A, B, C in Fig. 4) or liquid water (D in Fig. 4). However, some spectra do not respect the  $\frac{3}{4}$  ratio (E in Fig. 4). These spectra are therefore removed from the dataset. Thereby, 2335 infrared spectra are kept for the period from May 7 to November 19, 2019 (71.4% of the initial dataset).

Furthermore, the band area data are also processed using "Chauvenet's criteria" (eqn (2) and (3) in Appendix 3 in the ESI†), while checking beforehand the data distribution follows a normal law. After the processing with Chauvenet's criteria (from 1 to 3 runs) of each time sub-period within the baseline period, 2325 spectra are selected (71.1% of the initial dataset).

It is also possible to quantify the  $\text{CO}_2$  absolute concentration from a previous mathematical calibration (polynomial type) of the band area data<sup>55</sup> extended by Adisaputro *et al.* (2021)<sup>82</sup> for  $\text{CO}_2$  measurements acquired with a resolution of  $1\text{ cm}^{-1}$ . It provides an absolute concentration of  $\text{CO}_2$  in ppmv.

In order to determine the dissolved  $\text{CO}_2$  concentration, the hypothesis of the equilibrium between the gas phase and the dissolved phase of the considered species is also put forward. Thus, a formula similar to eqn (4) in Appendix 3 in the ESI† is used. Then, a comparison between the Raman data and the infrared data is established to assess the quality of the measurements by integrating the optical parameters detailed in the following sections.

## Results and discussion

In the following figures (Fig. 5–8), the start of each numbered sub-period corresponds to logistical interruptions during measurements (shutdown/restart due to power cut, fuse change, resumption and modification of the acquisition parameters, gas cell cleaning) that can change a little the spectrometer response and induce some discontinuity in calculated concentrations especially for Raman. During the Raman sub-period #5 (from July 26 to August 22, 2019), the temperature in the technical shed increased sharply. Thus, the infrared device switched automatically to temperature safety mode while the Raman device continued to record measurements. It is possible that the Raman measurement bias during this period is linked to the heatwave: more than  $35\text{ °C}$  in the shade for several days.

### Raman results

The peak area evolution of  $\text{N}_2$ ,  $\text{O}_2$ ,  $\text{CO}_2$  and  $\text{H}_2$  is studied from 7633 Raman spectra (example of a spectrum in Fig. 3). In order to interpret the future variations in dissolved  $\text{H}_2$



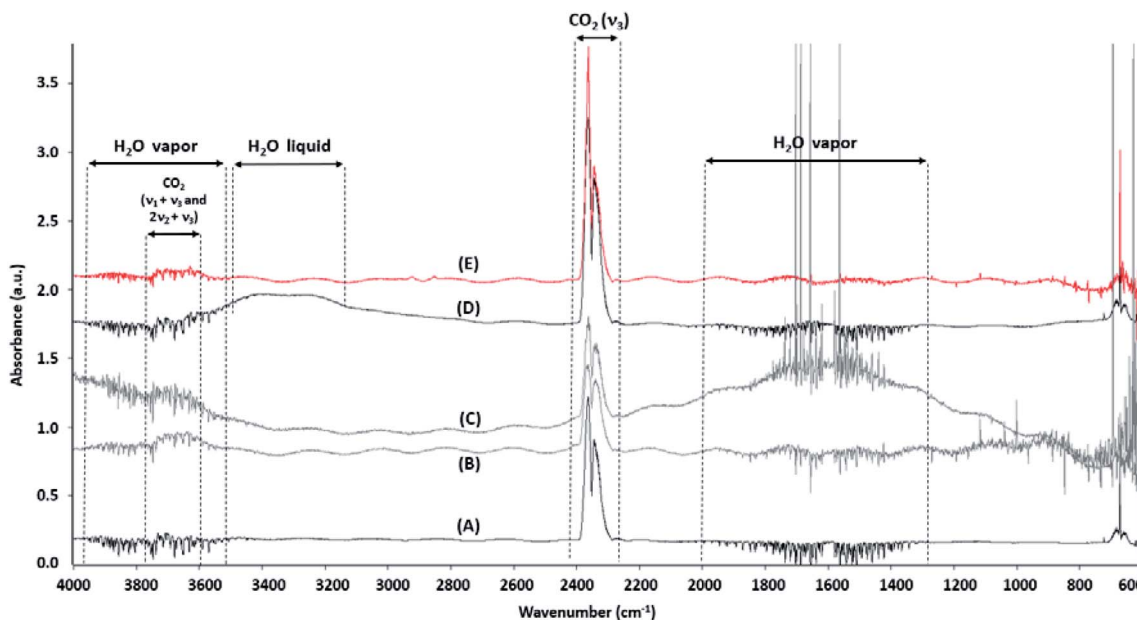


Fig. 4 Example of typical infrared spectra recorded in the continuous gas monitoring well on the Catenoy injection site. Four profiles were selected for the calculation of CO<sub>2</sub> concentrations (black and grey spectra: A–D) and one was deleted (red spectrum: E).

concentrations during the injection of this gas into the aquifer, H<sub>2</sub> concentrations are also monitored in the geochemical baseline period.

The dissolved gas concentrations are obtained with a water temperature of 12 °C and an average pressure in the membrane of 1.45 bar (relatively constant pressure during the baseline period) which conforms to the pressure of the gas system.

Since the signal for each gas is noised by several transient fluctuations throughout the duration of baseline monitoring, a concentration representation by a moving average is needed to attenuate signal fluctuations (Fig. 5–7). This parameter may be relevant for the future dissolved H<sub>2</sub> injection due to the high cross section of H<sub>2</sub>: a variation in signal intensity at time intervals with a magnitude of tens of minutes is expected there.

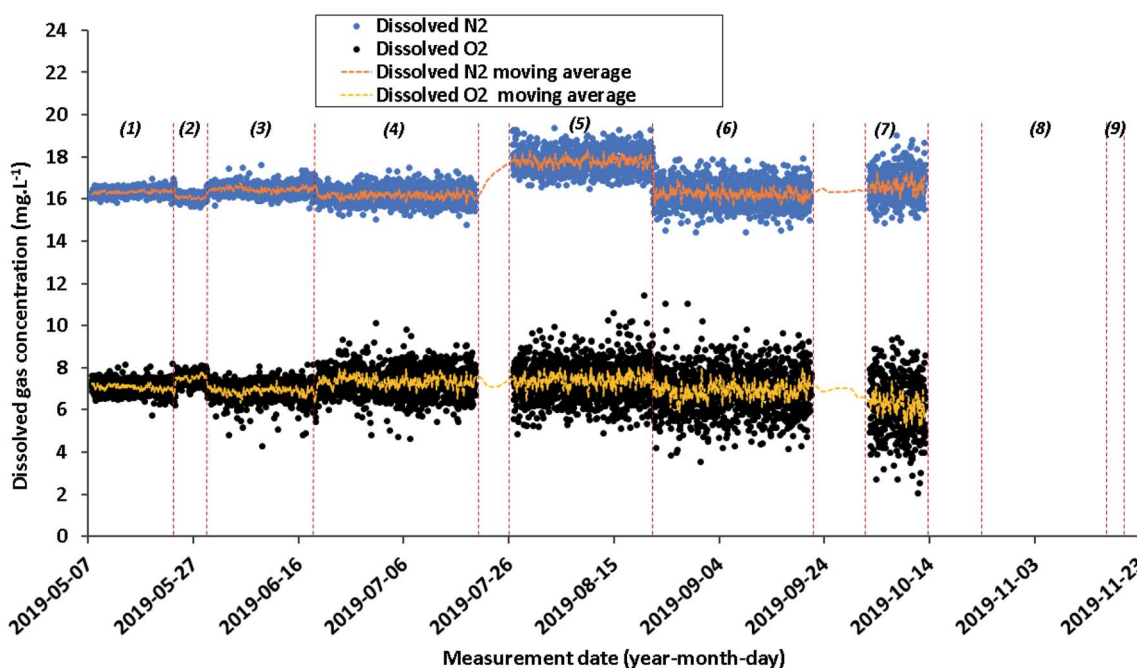


Fig. 5 Quantitative evolution of dissolved N<sub>2</sub> concentration (data values in blue and moving average in orange) and O<sub>2</sub> concentration (data values in black and moving average in yellow) from May 7, 2019 to November 19, 2019, calculated from Raman data.



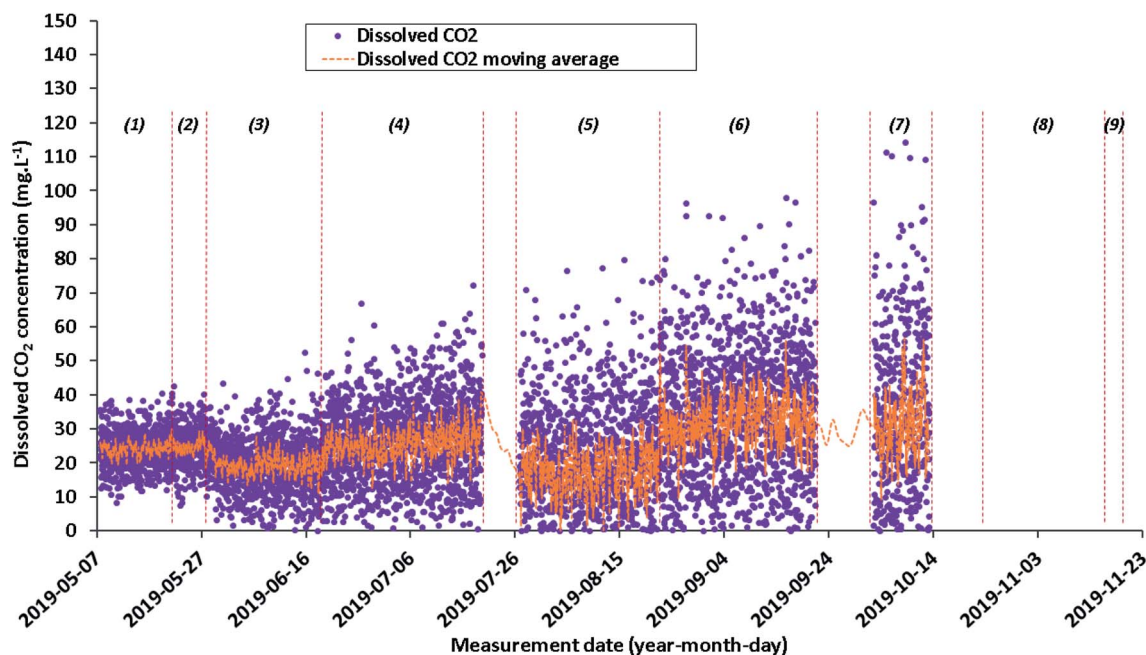


Fig. 6 Quantitative evolution of dissolved CO<sub>2</sub> concentrations (data values in purple and moving average in orange) from May 7, 2019 to November 19, 2019, calculated from Raman data.

Moving averages on various sets of data (from 5 to 25 data) are therefore tested. A moving average on a set of 10 individual data allows the general trend to be shown, which best represents individual data while attenuating signal fluctuations. Since the individual data are acquired every 30 minutes, each Raman moving average corresponds to a period of 300 minutes (*i.e.* 5 hours). Even if the measurements were carried out

continuously, the Raman signal cannot be accumulated *in situ* over 5 hours because of CCD detector saturation, background changes over time that do not average correctly or signal distortion owing to high background. The dissolved N<sub>2</sub>, O<sub>2</sub>, CO<sub>2</sub> and H<sub>2</sub> concentrations are plotted as daily individual data to better monitor how the situation evolves and as a moving average to show clearly the trend (Fig. 5–7).

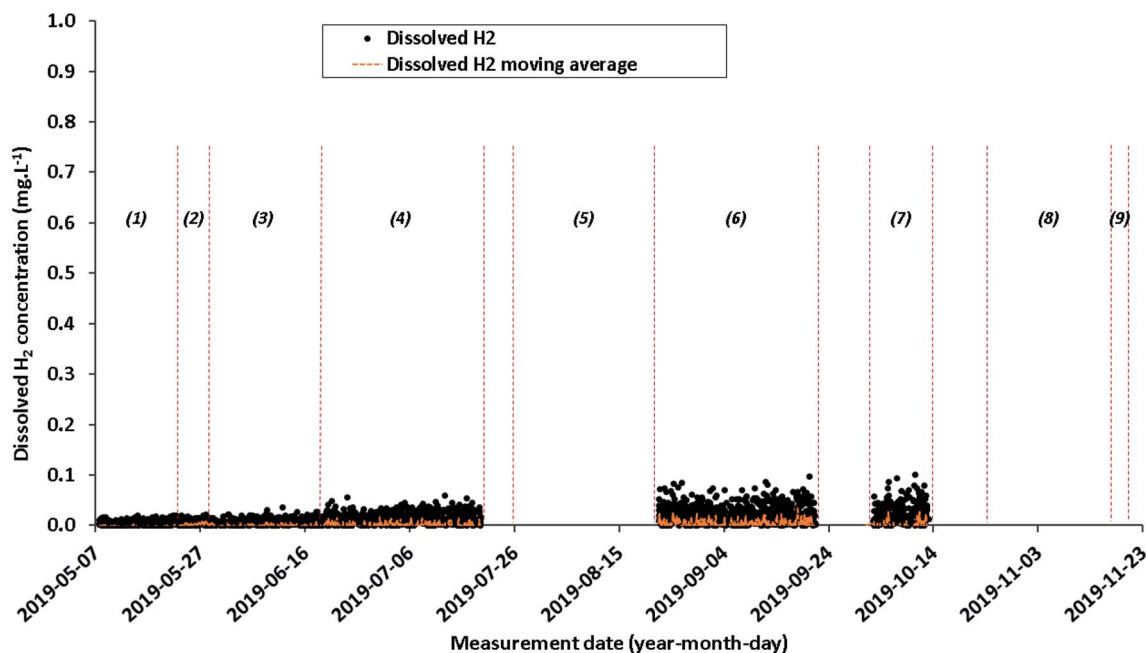


Fig. 7 Quantitative evolution of dissolved H<sub>2</sub> concentration (data values in black and moving average in orange) from May 7, 2019 to November 19, 2019 calculated from Raman data.



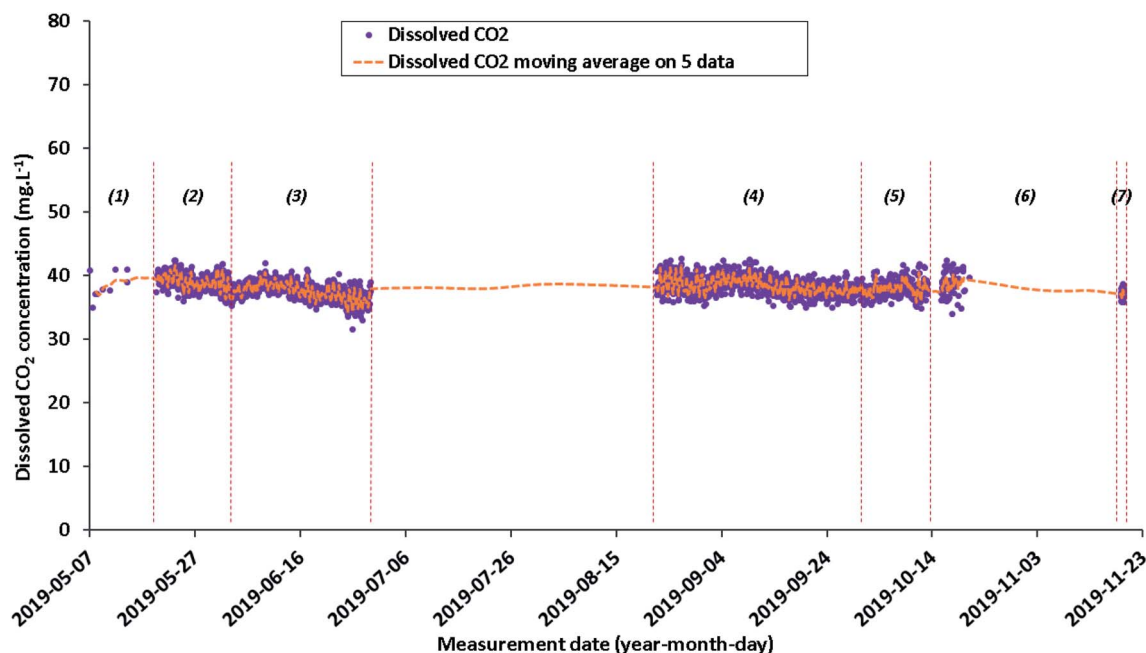


Fig. 8 Quantitative evolution of dissolved CO<sub>2</sub> concentration (data values in purple and moving average in orange) from May 7, 2019 to November 19, 2019, calculated from infrared data.

Although the Raman measurements were carried out from May 7, 2019 to November 19, 2019, a drift of the system is observed mainly due to a gradual decrease in laser power. This drift induces erroneous concentration values which prevent any physical interpretation. For this reason and in order not to confuse the interpretation, the data for periods #8 (October 24, 2019 to November 18, 2019) and #9 (November 18 to November 19, 2019) are not presented.

The concentrations of dissolved N<sub>2</sub> and O<sub>2</sub> are generally stable around an average of  $16.5 \pm 0.4$  mg L<sup>-1</sup> ( $\pm 1\sigma$ ) for N<sub>2</sub> and  $7.1 \pm 0.7$  mg L<sup>-1</sup> ( $\pm 1\sigma$ ) for O<sub>2</sub> with a data dispersion which widens from month to month. Small variations of dissolved N<sub>2</sub> concentrations were noted between the first seven time periods with average concentrations from  $16.1 \pm 0.1$  to  $17.8 \pm 0.5$  mg

L<sup>-1</sup> for N<sub>2</sub> and the O<sub>2</sub> concentrations were varied from  $6.2 \pm 1.3$  to  $7.5 \pm 0.3$  mg L<sup>-1</sup> (Fig. 5 and Table 2). As explained before, this fluctuation is mainly due to the decrease of the laser power. Under these conditions, it is hard to make a link with a fluctuation in the geochemical characteristics of the aquifer.

The evolution of the dissolved CO<sub>2</sub> concentration also varied over the nine time periods. During each period, the average concentration stabilizes around a different average value from that of the previous period: from 17.5 to 32.2 mg L<sup>-1</sup>. Moreover, the increase of the initial dispersion of concentrations over each of these 8 periods is due to a strong decrease of the laser power.

As shown in Fig. 7, the concentrations of dissolved H<sub>2</sub> are close to zero, at the level of the limit of detection (LOD) of the

Table 2 Average and standard deviation of dissolved N<sub>2</sub> and O<sub>2</sub> concentrations (mg L<sup>-1</sup>) determined from Raman data values for each of the nine time periods of the geochemical baseline

Period in 2019	Spectra number	Dissolved gases			
		N <sub>2</sub> (mg L <sup>-1</sup> )		O <sub>2</sub> (mg L <sup>-1</sup> )	
		Average	Standard deviation ( $\pm 1\sigma$ )	Average	Standard deviation ( $\pm 1\sigma$ )
May 7–23	735	16.3	0.2	7.1	0.3
May 23–29	271	16.1	0.1	7.5	0.3
May 29–June 19	968	16.5	0.2	6.9	0.4
June 19–July 19	1391	16.2	0.4	7.4	0.6
July 26–August 22	1223	17.8	0.5	7.4	0.9
August 22–September 21	1382	16.2	0.6	6.9	1.0
October 2–13	477	16.6	0.7	6.2	1.3
October 24–November 18	1167	—	—	—	—
November 18–19	19	—	—	—	—



**Table 3** Average and standard deviation of dissolved CO<sub>2</sub> concentrations (mg L<sup>-1</sup>) determined from Raman data values for each of the nine time periods of the geochemical baseline

Period in 2019	Spectra number	Dissolved gas	
		CO <sub>2</sub> (mg L <sup>-1</sup> )	
		Average	Standard deviation ( $\pm 1\sigma$ )
May 7–23	735	24.2	5.8
May 23–29	271	24.6	5.5
May 29–June 19	968	19.2	8.1
June 19–July 19	1391	25.7	12.8
July 26–August 22	1223	17.5	19.1
August 22–September 21	1382	32.2	21.0
October 2–13	477	31.2	27.5
October 24–November 18	1167	—	—
November 18–19	19	—	—

fiber Raman sensor during the entire baseline period. Note that during period #5, from July 26, 2019 to August 22, 2019, the dissolved H<sub>2</sub> values are not shown in Fig. 7.

### Infrared results

As with Raman data, infrared data are also represented in individual data and in moving averages (on five data) (Fig. 8). Since the individual data are acquired every 60 minutes, each infrared moving average data point corresponds to a period of 300 minutes (*i.e.* 5 hours).

During the entire baseline period, dissolved CO<sub>2</sub> concentrations are generally relatively stable with an average of 38.2  $\pm$  1.4 mg L<sup>-1</sup>. However, seven time periods can be defined. The dispersion of the infrared data is relatively low over time: from 0.7 to 2.0 mg L<sup>-1</sup> ( $\pm 1\sigma$ ) (Fig. 8 and Table 4). The lack of data between the end of June and the end of August (Fig. 8) is due to computer or spectrometer overheating because of higher summer temperatures. During the period from end-October to mid-November 2019, the cell was often filled in with liquid water, which resulted in incorrect measurements: these data are not shown in Fig. 8.

### Instrumental assessment between Raman and infrared data

**Characteristics of Raman and infrared data.** The dissolved N<sub>2</sub>, O<sub>2</sub> and H<sub>2</sub> concentrations obtained by Raman measurements remain relatively stable during the first three months and then seem to increase during the last two months of the monitoring period. At the same time, the dispersion of the measured concentration values increased (Fig. 5, Table 2, Fig. 6, Table 3 and Fig. 7). The dispersion of the Raman measurements can be expressed as follows over time: it varies from 1 to 4% of the average value of N<sub>2</sub> concentration, from 4 to 21% for O<sub>2</sub>, around 0% for H<sub>2</sub> and from 22 to 109% for CO<sub>2</sub> (according to Tables 2 and 3). In contrast, the infrared profile of dissolved CO<sub>2</sub> concentrations remains stable with a smaller data dispersion from 1.8 to 5.2% (Table 4). Thus, the concentrations of dissolved CO<sub>2</sub> obtained from Raman spectra have a dispersion factor between 12 and 21 times upper than that from infrared

spectroscopy. To explain this difference between the Raman and infrared signals, the determination of the optical characteristics (signal-to-noise ratio: SNR and the instrumental derivative factor: *F*) linked to the measurements of dissolved gas concentrations was quantified and appended (Appendix 4 in the ESI†).

One can conclude that the evolution over time of the optical characteristics of the Raman sensor is more important than for the infrared sensors. This leads to variations in the determination of the dissolved CO<sub>2</sub> concentration. Indeed, the dissolved CO<sub>2</sub> concentrations determined by Raman spectroscopy fluctuate between 17.5 and 32.2 mg L<sup>-1</sup> (Table 3) and those determined by infrared spectroscopy are around 37.2–38.4 mg L<sup>-1</sup> (Table 4). However, a ratio between the moving average concentrations of dissolved CO<sub>2</sub> measured from infrared and those measured from Raman was determined over the period from May 19 to June 19, 2019 (Fig. 9). This period is the one that presents the best quality of measurements over the entire period of the geochemical baseline. An average ratio of 1.6 was observed from May 19 to 29 (subperiod a''). This average ratio drops to 1.9 from May 29 to June 19, 2019 (sub-period b'') (Fig. 9). This confirms the progressive decrease about the

**Table 4** Average and standard deviation of dissolved CO<sub>2</sub> concentrations (mg L<sup>-1</sup>) determined from infrared data values for each of the seven time periods of the baseline period

Period in 2019	Spectra number	Dissolved gas	
		CO <sub>2</sub> (mg L <sup>-1</sup> )	
		Average	Standard deviation ( $\pm 1\sigma$ )
May 7–19	10	38.4	2.0
May 19–June 2	330	39.1	1.3
June 2–29	638	37.5	1.4
August 22–October 2	981	38.4	1.4
October 2–13	246	38.1	1.2
October 15–22	65	38.7	1.9
November 18–19	55	37.2	0.7



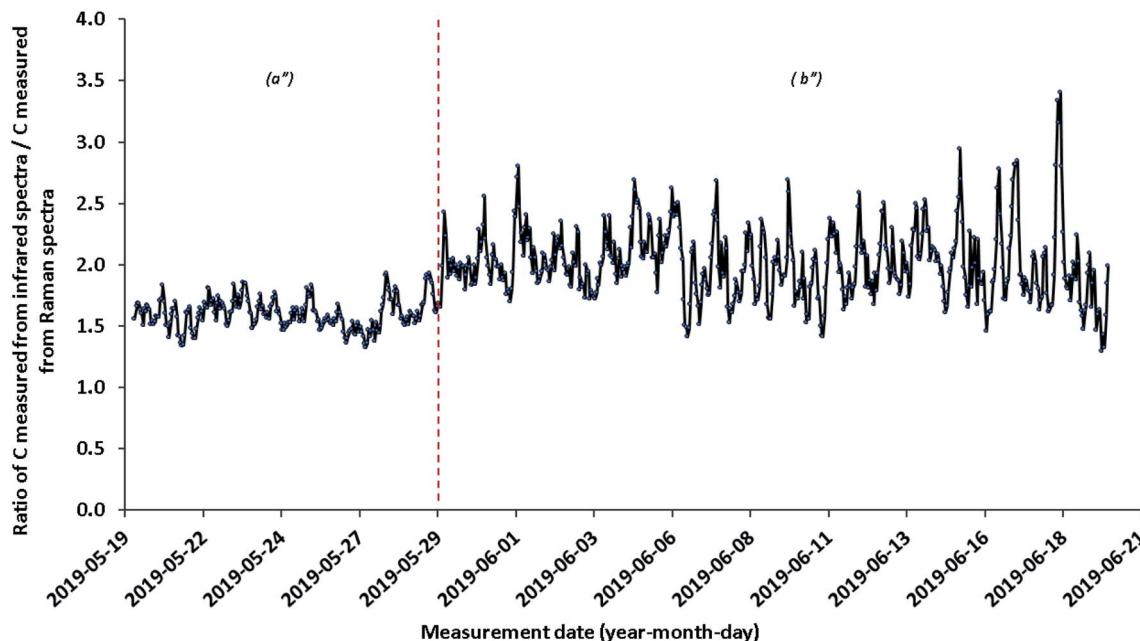


Fig. 9 Evolution of the ratio between the moving average concentrations (C) of dissolved CO<sub>2</sub> measured from infrared and those measured from Raman over the period from May 19 to June 19, 2019. The dotted red vertical line marks the two variations of this ratio (a'' and b'').

optical characteristic of the fiber Raman sensor (see Appendix 4†). Due to the greater stability and sensitivity of the infrared sensor, the infrared data for dissolved CO<sub>2</sub> can be used both for monitoring the deviation of the fiber Raman sensor and for the calibration of quantitative Raman measurements.

All these observations strongly argue in favor of a regular maintenance of the Raman spectrometer (spectrometer wavelength calibration, cleaning and purging the measuring cells, checking laser power) that is necessary every two months to obtain gas measurement of better quality with an almost zero derivative factor and better SNR values (see Appendix 4†). In order to ensure the relevance of the data measured by two different techniques, it is necessary to carry out a comparison for other gaseous species such as by regularly taking water samples on the same site (e.g. dissolved O<sub>2</sub> concentrations<sup>69</sup>).

**Comparison with previous geochemical data.** The monitoring methods for defining the geochemical baseline of this study are compared with the results in Lafortune *et al.* (2020)<sup>69</sup> by looking at the dissolved O<sub>2</sub> concentrations. This dissolved gas concentration was measured *in situ* within the freshwater aquifer of the Catenoy site during the same baseline period in these two studies. The dissolved O<sub>2</sub> concentrations are similar in these two studies with values measured of  $5.4 \pm 1.7 \text{ mg L}^{-1}$  in Lafortune *et al.* (2020) and  $7.5 \pm 1.4 \text{ mg L}^{-1}$  in this study. The difference is most probably due to the use of two very different methods (chemistry *vs.* spectroscopy, probe in a sample *vs.* calculation from the gas phase composition).

Moreover, using a mathematical simulation (Phreeqc software<sup>83</sup>) with the physico-chemical and ionic parameters of the Catenoy waters analyzed during the same period of the geochemical baseline,<sup>69</sup> an average dissolved CO<sub>2</sub> concentration of  $28 \text{ mg L}^{-1}$  is obtained. Moreover, discontinuous sampling in the aquifer under the same conditions in PZ1 and PZ3 exhibits

a concentration of CO<sub>2</sub>(aq) between 23.8 and 26.6 mg L<sup>-1</sup>. All these data are relevant to Raman and infrared *in situ* measurements in the dedicated well "PZ2 ter".

## Conclusion

Monitoring of dissolved gases was carried out under real conditions in a shallow water table in the Paris basin (Catenoy, France) over a period of more than 6 months. The measurement device involves 3 modules, a completion module placed in a dedicated well and equipped with a semi-permeable membrane with rapid gas transfer kinetics, a gas circulation module connected to a measurement module equipped with a fiber Raman sensor and an infrared sensor with a low optical path. It leads to the following main conclusions:

- The variability of dissolved gases can be continuously monitored in a saturated system such as an aquifer.
- The combination of Raman and infrared sensors makes it possible to quantify the concentrations of N<sub>2</sub>, O<sub>2</sub>, CO<sub>2</sub>, H<sub>2</sub> and CH<sub>4</sub> and to monitor their variability over more than one seasonal cycle.
- The Raman and infrared measuring cell must be placed in a temperature-controlled place. Under this condition, the signal-to-noise ratio remains stable for the infrared sensor during the period studied, requiring only an annual maintenance operation. For the fiber Raman sensor, the signal-to-noise ratio depends on the evolution of the laser power, which must be checked quarterly.
- Monitoring over a period of more than 6 months shows a very low variability of the dissolved gas concentrations in the aquifer, which will allow rapid detection of any possible drift, particularly linked to an accidental arrival of gases such as hydrogen.



Moreover, such online gas measurement systems can be deployed as it is on any type of site without the need for adaptation. For a new site and in order to validate the experiment protocol, it will be necessary to carry out a comparison by regularly analyzing water samples by conventional chemical methods. In order to improve the industrialization of these monitoring techniques in the context of underground gas storage, additional promising technologies are actually in development such as fiber optic network platforms<sup>84,85</sup> and/or sensors of the AlGaIn/GaN doped semiconductor type HEMT (High Electron Mobility Transistors).<sup>86</sup>

## Conflicts of interest

There are no conflicts to declare.

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