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# Physicochemical and microbiological effects of geological biomethane storage in deep aquifers: introduction of $O_2$ as a cocontaminant<sup>†</sup>

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Biomethane is considered one of the most promising energy vectors to substitute fossil fuels during the global energy transition. Its production is steadily increasing, and high storage volumes are needed to cover seasonal needs. Existing underground gas storage (UGS) aquifers, which have been used for natural gas storage, are excellent candidates. Underground aguifers are known for being anoxic systems. However, dioxygen (O<sub>2</sub>) can be injected as an impurity with biomethane into these anoxic environments. O<sub>2</sub> limitations in the underground vary worldwide; however projects are conducted to optimize these limitations. It has been shown that O<sub>2</sub> presence can affect the aquifer's ecosystems and induce mineral reactions. Thus, a multidisciplinary study was conducted in which the in situ conditions were simulated in a high-pressure reactor. Water containing autochthonous microorganisms and reservoir rock were used as the aqueous and solid phases, respectively. Initially, the gas phase was composed of methane, 1% CO<sub>2</sub>, benzene and toluene under 60 bar and 36 °C conditions. Sulfate was depleted from the aqueous phase due to sulfate-reducing microorganismes. After 50 days, 100 ppm O<sub>2</sub> was injected into the gas phase. Sulfate reducers were inactivated; however, other taxonomic groups became dominant, such as members of the class Acidobacteriae and the families Desulfitobacteriaceae and Kineosporiaceae. Hydrocarbon biodegradation was demonstrated by a benzene decrease in the aqueous phase, which was barely affected by  $O_2$  injection. However microbial analyses suggested a shift in the ecosystem to adapt to this new 'low aerobic' conditions. The findings of this study can help for better understanding of any other process including O2 as an impurity in UGS such as CCS and CAES.

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#### **Environmental significance**

With the biomethane sector development, there is an increase of its injection in the natural gas network, including geological storages. Before its injection,  $O_2$  is added to eliminate sulfides. A co-injection of  $O_2$  is expected in geological storage (accepted limit of 100 ppm), such as deep aquifers, which host autochthonous microorganisms. This  $O_2$  threatens strictly anaerobic microorganisms and their biodegradation activities of monoaromatic hydrocarbons coming from the gas storage. Multidisciplinary studies mimicking deep aquifer conditions are essential to confirm or adapt the authorized limits of  $O_2$ . Our study showed no major modification of aquifer rocks (minerals and porosity) but a shift of the diversity of the community, eliminating the less resistant ones and creating a new equilibrium allowing benzene degradation.

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#### Introduction

Biomethane may provide a new, reliable and sustainable source of energy and an effective alternative to fossil fuels. After carbon dioxide ( $\rm CO_2$ ), hydrogen sulphide ( $\rm H_2S$ ) and water removal, biogas is essentially composed of biomethane. Biomethane is gaining interest and governments are encouraging methanization units. In France, for example, the aim is to increase the share of renewable gas in the gas network to up to 10% by 2030. Concurrently, Europe launched the REPowerEU project that aims to speed up the roll-out of renewable gases. One of the

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objectives was to attain a biomethane production of 35 billion cubic metres per year by 2030, while its production was at 3.6 billion cubic metres by the end of 2021.² To optimize biomethane use as a major energy source, gas storage facilities of sufficient volumes must be made available. The use of existing underground geological storage (UGS) sites that are known for their large capacities is important, especially to store large volumes to supply during periods of high gas demand, energy crises or even in the case of political issues. Moreover, the use of UGS seems rational since it is beneficial to use the existing infrastructure and network.

Among UGS solutions, deep aquifers are known for their capacity to store large volumes of gas under pressure under anoxic conditions and to be home of anaerobic microbial communities.<sup>3,4</sup> Sulfate-reducing microorganisms (SRMs) were the most frequently detected microorganisms in deep aquifers.<sup>4-12</sup> Sulfate quantities in UGS aquifers have been observed to decrease at multiple sites.<sup>4</sup> Nevertheless, an important product of SRMs is H<sub>2</sub>S, which can lead to UGS problems such as corrosion of wells and souring.<sup>13</sup> However, studies have also shown the capability of SRMs to degrade monoaromatic hydrocarbons found in natural gas storage mixes, especially benzene, toluene, ethylbenzene and xylene (BTEX).<sup>6,9,12,14,15</sup>

One of the main differences between natural gas and biomethane is the potential presence of oxygen (O<sub>2</sub>) in the latter.<sup>16</sup> In fact, O2 is used to remove the undesirable H2S compounds present in the biomethane product. The current European recommendations are that the O2 content must not exceed 100 ppm within the gas circulating in the network and possibly in UGS.  $^{17-19}$  In UGS,  $\mathrm{O}_2$  can also be present as an impurity during CO<sub>2</sub> capture and storage (CCS)13,18,20-22 and in compressed air for energy storage (CAES).23-25 O2 can also be intentionally injected for enhancing oil recovery26-28 and for the bioremediation of groundwater.<sup>29,30</sup> O<sub>2</sub> that is injected into deep aquifers not only affects microbial life but also interacts with mineral phases.<sup>25</sup> One example is mineral oxidation, which depends on the nature of the solid phase. 9,22,31-33 Recently, a study with 1% O2 was carried out since it corresponds to the maximum quantity found in biomethane. 9,16 The results indicated the deleterious effect of O2 on microbial life. Bioattenuation of monoaromatic hydrocarbons ceased.

Here, the experiment was performed to understand the effect of 100 ppm (0.01%)  $O_2$ , the maximum quantity authorized to circulate in the gas grid today. The conditions of an UGS aquifer were reproduced in a high-pressure (HP) reactor for 86 days and the results provided evidence for the potential impact of  $O_2$  introduction based on the current recommended limits. This multidisciplinary study permitted to understand the effect of  $O_2$  on the physicochemical parameters of the storage, on the microbial ecosystem developed underground and on the minerals found in the reservoir rock. The experimental work consisting of reproducing the aquifer used as UGS in a HP reactor with its different phases (gas, solid and liquid with autochthonous microorganisms) allowed observing the effects of 100 ppm  $O_2$  under the *in situ* conditions, the maximum concentration recommended in Europe. Indeed, this

concentration limit of  $O_2$  was established based on gas transport optimization, and storage conditions were not considered. Thus, studies are still needed to reconsider  $O_2$  limits in injected biomethane to protect UGS reservoirs. This work is the first to evaluate the effects of this  $O_2$  concentration limit on UGS in a deep aquifer.

#### Materials and methods

### Site description, formation water and reservoir rock characteristics

The UGS aquifer is situated in the Aquitaine sedimentary basin (582 m depth) in southwestern France.9 The selected monitoring well, denoted as Ab\_L\_1, is closest to the UGS.4 The formation water was obtained more than a year before the experiment due to COVID restrictions. The samples were stored in anaerobic bottles at 4 °C. The main anions comprising the formation water were chloride (0.35 mM) and sulfate (0.02 mM) (Table S1†). Two underground samplings were performed to recover 0.578 L and 0.556 L of water containing autochthonous microorganisms. Samplings were carried out from the site using two downhole samplers equipped with a PDS Sampler (Leutert Bottom Hole Positive Displacement Sampler<sup>33</sup>). Formation water was also sampled from the wellhead and was filter-sterilized (PES 47 mm membranes, 0.1 µm, Sartorius) under anoxic conditions and stored at 4 °C before use. A volume of 0.591 L of the filtered wellhead water was added to both volumes of the recovered formation water to yield the aqueous phase that was injected into the HP reactor. A volume of 0.1 L was taken from the final mixture for microbial diversity study (day 0), and the remaining (1.39 L) was injected into the reactor. On day 39 of the experiment, formation water supplemented with sulfate (0.60 mM) was added to maintain the system. The reservoir rock consisted of inframolassic sands. 4,9 Cuttings were collected from a drilling carried out near the studied site.

#### Experimental apparatus and analysis techniques

The entire experimental protocol has been schematically summarized in the form of a flowchart (Fig. S1†).

High-pressure (HP) reactor. The experimental apparatus consisted of a HP reactor as previously described. 9,10 Briefly, the reactor was made of Hastelloy C-276 to avoid corrosion problems. The autoclave was heated by using heating resistors with insulating coatings. To maintain the temperature of the apparatus, a double jacket was installed. The temperature was monitored by using thermocouples in both gas and aqueous phases. Both phases were mixed using a double disc stirrer with four vertical blades for the gas one and a Rushton turbine for the aqueous one, with a stirring speed of 20 rpm. The solid phase was in a basket, which was made of Teflon. Its bottom was composed of a Hastelloy C-276 disc with 10  $\mu$ m pores to prevent mineral particles from falling out.

**Physicochemical analyses.** Ion chromatography (Dionex Integrion HPIC; Thermo Fisher Scientific) was used to quantify fluoride, acetate, chloride, and sulfate ions. Water samples from the reactor were collected and immediately analysed for

the redox potential Eh (Mettler Toledo International Inc.). The gas phase was monitored throughout the experiment using inline gas chromatography with a microthermal conductivity detector (GC- $\mu$ TCD; Micro GC Fusion; Chemlys). Using this technique, CH<sub>4</sub>, CO<sub>2</sub>, O<sub>2</sub>, H<sub>2</sub> and N<sub>2</sub> were quantified in the gas phase. The aqueous and gaseous phase compositions were detected with a measurement uncertainty of 5%. More detailed specifications were presented previously. The gas quantities were calculated based on the total pressure measured by using a barometer and the gas phase composition was analysed by gas chromatography according to the real gas law corrected by the compressibility factor estimated by PhreeqC (2.4). The Eh of the aqueous phase collected from the reactor under strict anoxic conditions was monitored weekly at atmospheric pressure (Mettler-Toledo<sup>TM</sup> SevenCompact<sup>TM</sup>).

Benzene and toluene were quantified in the aqueous and gaseous phases throughout the experiment using a Thermo Fisher Scientific gas chromatograph coupled with a quadrupole mass spectrometer (GC-MS) (ISQ QD Single Quadrupole MS – Trace 1310), in agreement with the specifications detailed previously.<sup>9</sup>

X-ray tomography and mineralogical analyses. Three capillaries made of borosilicate with an internal diameter of 2 mm and a height of 3 cm were filled with the solid phase and placed in the Teflon basket. The top, middle and bottom of each capillary were scanned by X-ray tomography before and after the experiment. A comparison of both enabled the detection of morphological alterations in the solid phase and the presence of microorganisms in the water phase or on the surface of the solid grains. The initial scans were performed in a dry state. For the final scans performed at the end of the experiment, the capillaries were sealed with waterproof glue at room temperature in an anaerobic glove box. A Zeiss Xradia Versa 510 X-ray tomograph was utilized for the scans, and a voxel size of 2.5 microns and a square field of view of 2.5  $\times$  2.5 mm² were employed.  $^{9,10}$ 

The remaining solid phase was analysed by X-ray diffraction (XRD) before and after the experiment to detect any variations in the mineral phase. Samples were collected from the basket at four different depths at room temperature in an anaerobic glove box. The samples were then dried with  $N_2$  gas flux to limit mineral oxidation, manually ground and sieved (<100  $\mu m$ ) into a homogeneous powder. The solid powders were then mounted on holders and directly analysed by XRD. The analyses were performed using a Bruker D2 Phaser powder diffractometer equipped with a Cu K $\alpha$  radiation source. The XRD patterns were recorded over a 5° to 90° 20 range with a 0.02° step and a 0.5 s counting time per step. DIFFRAC.EVA software was used to identify the mineral phases.

Aliquots of the same samples collected at four different depths from the basket were submitted for a petrographic study using scanning electron microscopy coupled to energy dispersive spectroscopy (SEM-EDS). Solid pieces were directly mounted on PIN stubs and coated with carbon. Observations and mineral identification were performed with a JEOL JSM 7800F Prime SEM-FEG equipped with an Oxford Instruments AZtecEnergy EDS SDD X-Max 80 mm<sup>2</sup> detector at the Castaing Center in Toulouse, France.

#### Microbial analyses

Nucleic acid extraction and RNA reverse transcription. Samples from the aqueous phase were collected throughout the experiment with the aim of coextracting nucleic acids (DNA and RNA). Aqueous samples were collected from the reactor and directly filtered using 47 mm PES membrane filters of 0.1 µm porosity (Sartorius Stedim). The filters containing the samples were then stored at -80 °C. Subsequently, the filters were crushed in liquid nitrogen, and the nucleic acids were collected using a Fast RNA Prosoil Direct kit (MP BIO). Then, DNA and RNA were separated using AllPrep RNA/DNA (Qiagen). The extracted DNA and RNA were quantified using a Quant-it™ dsDNA HS (Invitrogen) kit and a Quant-it™ RiboGreen (Invitrogen) kit, respectively. To measure the extracted DNA and RNA, a BioTEK SYNERGY HTX microplate reader was used. The reverse transcriptase M-MLV (InvitrogenTM) was used to achieve reverse transcription of the RNA.

Polymerase chain reaction and sequencing. To study global microbial diversity, the V4-V5 region of 16S rRNA genes and complementary 16S rDNA were amplified by PCR (2700 Thermal Cycler, Applied Biosystems). To reduce the deleterious effects of PCR inhibitors, 1 mg mL<sup>-1</sup> of BSA (Bovine Serum Albumin, NEB-B9200S) was added to the reagent mix (Taq PCR Core Kit, Roche). For sequencing, the PCR primer set (515F-928R34) contained the adapters GTGYCAGCMGCCGCGGTA (forwards) and CCCCGY-CAATTCMTTTRAGT (reverse). Sequencing was performed by using the GenoToul genomic platform (Toulouse, France) using Illumina MiSeg  $2 \times 250$  bp technology. The resulting sequencing data represented 511 618 reads. Raw sequences were archived in the public NCBI database (BioProject ID PRJNA930342). These data were analysed via the FROGS pipeline (GenoToul genomics platform in the Galaxy interface).35 Preprocessing resulted in 424 552 reads that were clustered into OTUs taxonomically classified with the Silva database (release 138.1) and into groups at the taxonomic family level to monitor the evolution of taxonomic diversity during the experiment.

Quantitative PCR. Genes and transcripts of the 16S rRNA, dsrB and mcrA genes were quantified to monitor the activity of the overall microbial community, SRM and methanogenic archaea separately at different times during the experiment in the HP reactor. Quantitative PCR was performed with a Takyon NO ROX SYBR 2X MasterMix blue dTTP kit (Eurogentec) according to the supplier's instructions and with a Bio-Rad CFX Connect. The primer pairs consisted of 515F-928R34 (400 nM), DSR 2060F-DSR 4R³6,37 (300 nM), and mlas-mcrArev³8,39 (400 nM). The number of copies of transcripts was calculated using a standard with serially 10-fold diluted pCR™ 2.1-TOPO plasmid (TOPO TA cloning kit, Invitrogen).

#### **Experimental protocol**

Before reservoir rock samples were introduced into the HP reactor, they were rinsed with isopropanol and demineralized water to eliminate potential hydrocarbons and drilling brine salts. The solid phase was then dried overnight at 90  $^{\circ}$ C. Once the solid phase was ready, capillaries were filled with solid samples and were analysed via X-ray tomography before the

experiment.9 Filled and scanned capillaries were placed in a basket containing the solid phase inside the reactor. When the experiment was performed, the capillaries were analysed again to compare with the initial solid state.

Once the reactor was closed with the basket inside, a sterilization step was carried out. A volume of 60 mL of demineralized water was injected, and a flush of N2 gas was applied to eliminate O2 in the reactor. The reactor was then heated to 100 °C for 24 h for sterilization under anoxic conditions. Afterwards, the reactor was cooled to 36 °C, which corresponded to the experimental temperature, and a light vacuum was created before the formation water was injected. The aqueous phase containing the autochthonous microorganisms was then injected into the reactor. Following the injection of the aqueous phase, the gaseous phase was added. The gas mixture initially consisted of CH<sub>4</sub> with 1% CO<sub>2</sub>, 3.57 ppm benzene and 7.95 ppm toluene to simulate natural gas. Initially, the piston was at a high level such that the solid phase was completely immersed in the formation water. This step was important so that the solid phase came into contact with the aqueous phase and the microbial community. After 9 days, the piston was lowered, and only 1 cm of the solid phase remained in contact with the aqueous phase and in contact with microorganisms (2 cm in the gaseous phase). The gas mixture was reinjected to adjust for the loss of pressure. After 50 days, O2 was injected into the system at a concentration of 100 ppm in the gas phase. The biotic experiment lasted 86 days. The basket containing the solid phase was carefully removed from the reactor and stored in a sealed anaerobic jar. An anaerobiosis generator and indicator pockets (Dutscher Ref. 0260001) were used in the jar. The basket was then transported to an anaerobic glove box. In the glove box, samples of the solid phase were collected for X-ray tomography and mineralogical analyses.

#### Thermodynamic modelling

To estimate the thermodynamic liquid-gas equilibrium, PhreeqC software was used to estimate the solubility of each gaseous compound in the system. To do so, a simplified abiotic system comprising the aqueous phase and the gas phase was modelled to quantify the solubility of the gas in the liquid phase at different stages of the experiment. The calculated values were then compared to the values measured with gas chromatography. The chosen database was 'Phreeqc.dat', which uses the extended Debye-Hückel law for the activity coefficient and the Redlich-type equation of state for the gas phase. Redox reactions were not taken into account due to their slow kinetics in such environments.40-42 The calculated solubility corresponded to a simplified abiotic system in which only the gas and liquid phases were present. Estimating the dissolved gas in the aqueous phase helped to evaluate the amount of gases consumed by chemical reactions and microbial metabolism.

#### Results

Throughout the 86 days of the experiment, liquid and gas samples were collected and directly analysed by ion and gas chromatography, respectively. During the first 49 days, the system was under a pressure of 60 bar and a temperature of 36 °C, with a gaseous mixture composed of  $CH_4$  and 1%  $CO_2$  with traces of benzene and toluene. On day 50,  $O_2$  was injected with a concentration of approximately 100 ppm in the gas phase. The results obtained during this experiment were divided into two main stages: before  $O_2$  injection, when the simulated aquifer was under strict anoxic conditions similar to present-day conditions, and after  $O_2$  injection, simulating the introduction of  $O_2$  at a concentration equivalent to the maximum limit authorized today.

#### Evolution of the aqueous phase composition

The formation water sampled from the studied aquifer had very low salinity, approximately 0.8% of seawater salinity.  $^{9,43}$  Underground sampling of the formation water showed that it contained  $2.84\times 10^{-2}\pm 1.5\times 10^{-3}$  mmol of sulfate (ESI Table S1† and Fig. 1(a)). The sulfate quantity increased to  $3.77\times 10^{-1}\pm 1.9\times 10^{-2}$  mmol after aqueous phase injection into the reactor due to solid–liquid interactions and compounds release from the solid phase. From day 9 to day 39, a decrease in dissolved sulfate was observed. After day 39 and until day 44, the sulfate quantity increased to  $4.16\times 10^{-1}\pm 2.1\times 10^{-2}$  mmol after 0.1  $\mu$ m-filtered wellhead water was added with sulfate into the system before  $O_2$  injection. A decrease in sulfate quantity was then observed until day 51. Starting at this point, which corresponded to  $O_2$  injection (day 50), the sulfate quantity was stable until the end of the experiment.

The acetate quantity was  $1.16\pm5.0\times10^{-2}$  mmol at the beginning of the experiment (Fig. 1(b)). The concentration increased through the experiment to reach  $1.64\pm8\times10^{-2}$  mmol on day 44. After  $O_2$  injection, the acetate quantity reached a plateau around a value of  $1.60\pm8\times10^{-2}$  mmol and stayed stable until the end of the experiment.

#### Evolution of the gas phase composition

After the injection of the aqueous phase into the reactor, the gas mixture of CH<sub>4</sub> + 1% CO<sub>2</sub> was added at a pressure of 60.6  $\pm$  0.6 bar and a temperature of 36 °C to simulate the conditions in the targeted UGS. Using the PhreeqC model based on Peng-Robinson's equation, the compressibility factor of the gas mixture was calculated to be 0.9. Thus, by considering the compressibility factor,  $4.45 \pm 2.2 \times 10^{-1}$  mol of CH<sub>4</sub> was injected at the beginning of the experiment (Fig. 2). After 3 days, the CH<sub>4</sub> quantity decreased from the gas phase and reached 4.40  $\pm$  2.2 imes $10^{-1}$  mol. As shown in Fig. 2, the calculated values of CH<sub>4</sub> corresponded to the quantity assumed to remain in the gas phase after thermodynamic equilibrium, based on PhreeqC calculations. On day 9, the piston's position was readjusted in the reactor to keep 1 cm of the solid phase immersed in the aqueous phase. After the piston level was lowered, the gas pressure was readjusted, and one more injection of the gas mixture was carried out. The gas injection induced an increase in CH<sub>4</sub> estimated to be approximately 3.07  $\times$  10<sup>-1</sup>  $\pm$  1.53  $\times$  $10^{-2}$  mol. Until day 39, the quantity of CH<sub>4</sub> in the gas phase remained constant. After aqueous phase supplementation on

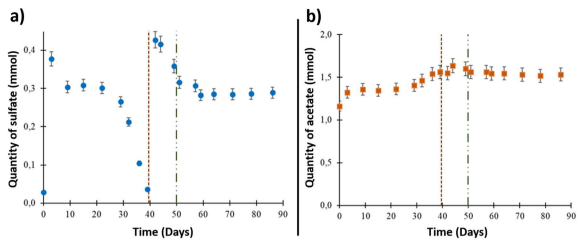


Fig. 1 Results of the aqueous phase analyses during the high-pressure experiment. The vertical dotted line on day 39 represents aqueous phase supplementation. The vertical line on day 50 represents oxygen injection. (a) Sulfate variation in the aqueous phase and (b) acetate variation in the aqueous phase.

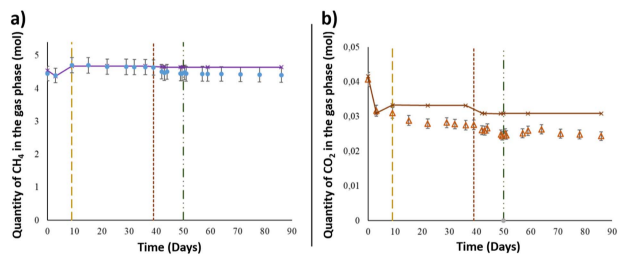


Fig. 2 Results of the gaseous phase analyses during the high-pressure experiment. The vertical dotted line on day 9 represents CH<sub>4</sub> + CO<sub>2</sub> reinjection. The vertical dotted line on day 39 represents aqueous phase supplementation. The vertical line on day 50 represents oxygen injection. (a) CH<sub>4</sub> variation in the gas phase. The purple dots and line represent the values calculated with PhreeqC. (b) CO<sub>2</sub> variation in the gas phase. The orange dots and line represent the values calculated with PhreeqC.

day 39, the CH<sub>4</sub> quantity in the gas phase decreased as it dissolved in the liquid phase. The quantity lost was  $1.29 \times 10^{-1} \pm$  $6.47 \times 10^{-3}$  mol. Following the aqueous phase addition and the gas-liquid thermodynamic equilibrium, the quantity of CH<sub>4</sub> in the gas phase remained stable until the end of the experiment. The estimated CH<sub>4</sub> given by the thermodynamic model fit well with the experimental data obtained during the study (Fig. 2).

Along with methane, CO<sub>2</sub> was injected at the beginning of the experiment and was present in the gaseous mixture. Under a total pressure of 60.6  $\pm$  0.6 bar and a temperature of 36 °C, the quantity of  $CO_2$  initially injected into the system was 4.07  $\times$  $10^{-2} \pm 2.0 \times 10^{-3}$  mol (Fig. 2), representing 1% of the gas phase. After thermodynamic equilibrium was reached between the gaseous and aqueous phases, a  $CO_2$  decrease of  $8.97 \times 10^{-3}$  $\pm$  4.5  $\times$  10<sup>-4</sup> mol was observed from the initial quantity. Even with the reinjection of the gas mixture on day 9, the measured

CO2 quantity showed a decrease in the gas phase and was approximately  $3.10 \times 10^{-2} \pm 1.5 \times 10^{-3}$  mol. The CO<sub>2</sub> quantity presented a continuous decrease through the experimental period to reach  $2.47 \times 10^{-2} \pm 1.2 \times 10^{-3}$  mol on day 49, just before O<sub>2</sub> injection. Similar to CH<sub>4</sub>, the calculated values of CO<sub>2</sub> represented the corresponding gas-liquid thermodynamic equilibrium in a binary abiotic system. As shown in Fig. 2, a marked difference was observed between the PhreeqC modelling results and experimental measurements, suggesting that the amount of gaseous CO2 may be controlled by reactions other than aqueous phase dissolution, e.g., microbial activity.

As explained above, the first step in the experiment was the reproduction of the aquifer conditions currently used for natural gas storage. The second step consisted of identifying the effects of O2 presence in such ecosystems. The experimental conditions required an O2 content of approximately 100 ppm in

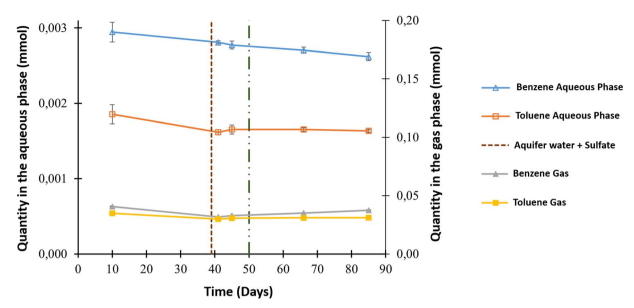


Fig. 3 Results of benzene and toluene variations in the gas and aqueous phases during the high-pressure experiment. The vertical dotted line on day 39 represents aqueous phase supplementation. The vertical line on day 50 represents oxygen injection. The blue and orange dots correspond to benzene and toluene values in the aqueous phase, respectively. The grey and yellow dots correspond to benzene and toluene values in the gas phase, respectively.

the gas phase, corresponding to the maximum quantity of  $O_2$  authorized in UGS storage in France today. Since these low concentrations were measured using small gas samples in  $\mu$ -GC, large uncertainties were generated in the measured values. The  $O_2$  quantity in the gas phase varied between 70 and 170 ppm. The system became microoxic, and the  $O_2$  content reached a plateau around this value. The solubility of  $O_2$  calculated by PhreeqC was lower than  $5.8 \times 10^2$  mM under these conditions. The redox potential increased in the formation water, confirming the presence of  $O_2$  in the system (Fig. 4(a)).

The aqueous and gas phases were sampled throughout the experiment for additional analyses, such as benzene and toluene quantification. The main source of both compounds was the gaseous mixture that contained initial concentrations of 3.57 ppm benzene and 7.95 ppm toluene. Fig. 3 shows the evolution in the quantity of these compounds in both phases. Before O2 injection, analyses of the aqueous phase showed a decrease in benzene from 2.95  $\times$  10  $^{-3}$   $\pm$  1.32  $\times$  10  $^{-4}$  to 2.77  $\times$  $10^{-3} \pm 5.31 \times 10^{-5}$  mmol and toluene from 1.86  $\times$   $10^{-3} \pm 1.26$ imes 10  $^{-4}$  mmol to 1.65 imes 10  $^{-3}$   $\pm$  0.60 imes 10  $^{-4}$  mmol. A benzene decrease was also observed in the gas phase, with a value of approximately 7.79  $\times$  10<sup>-3</sup>  $\pm$  7.18  $\times$  10<sup>-4</sup> mmol, but this was not the case for toluene. After the injection of O2 into the system, benzene only decreased in the aqueous phase to reach  $2.62 \times 10^{-3} \pm 5.39 \times 10^{-5}$  mmol; however, all the other values appeared to be constant.

#### **Evolution of the microbial community**

Members of the *Spirochaetaceae* family were detected throughout the experiment (Fig. 4(b)). Representing between 30 and 59% of the microbial community before the injection of  $O_2$ , their

abundance decreased to reach 16% at the end of the experiment. Methanogenic archaea (Methanobacteriaceae, Methanothermobacteriaceae, Methanosaetaceae and Methanosarcinaceae), which represented nearly 19% of the diversity at the beginning of the experiment, drastically decreased after the 36<sup>th</sup> day of incubation and O<sub>2</sub> injection, while the new equilibrium within the community favoured bacteria involved in the sulfur cycle. The Desulfurisporaceae family represented up to 39% of the community on day 36 and was most active (Fig. 4(c)). With the decrease in sulfate, the family Desulfurisporaceae (Desulfurispora genus) and its activity decreased from the 36<sup>th</sup> to the 42<sup>nd</sup> day. Supplementation with sulfate did not allow their regrowth before the injection of O<sub>2</sub> on the 50<sup>th</sup> day. On the other hand, the reduction of sulfates and the injection of O2 did not impact the growth of Desulfitibacteraceae, which were part of the dominant families. The members of this family identified in this study are all affiliated with the genus Desulfosporosinus. The injection of O<sub>2</sub> allowed a strong development of bacteria belonging to the class Acidobacteriae, which represented 64% of the microbial community at the end of the experiment. Acidobacteriae genes were very active and dominated the 16S rRNA gene transcripts (Fig. 4(c)). To a lesser extent, these new conditions also favoured the development of members of the Kineosporiaceae family (9% at the end of the experiment; affiliated with the genus Quadrisphaera). Although the low concentration of  $O_2$  in the gas phase made O2 quantification difficult, its presence led to an increase in the redox potential of the aqueous phase from approximately −160 mV before the injection of O<sub>2</sub> to +73 mV at the end of the experiment (Fig. 4(a)). In addition, this introduction of O<sub>2</sub> quickly stopped the decrease in sulfate in the 10 days that followed. The sulfate content was stable until the end of the experiment, clearly indicating the cessation of sulfate reduction, which was corroborated by the disappearance of the dsrB

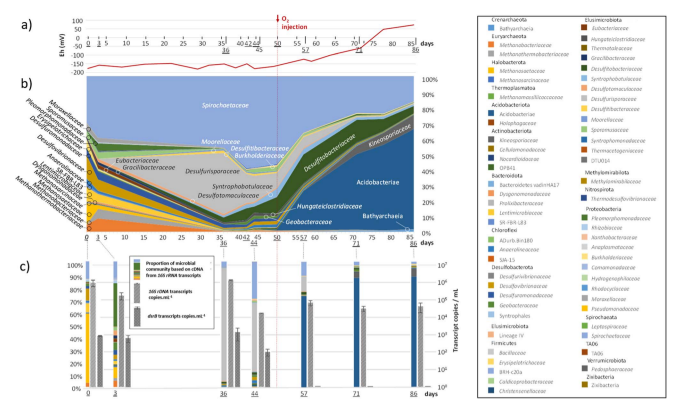


Fig. 4 Temporal monitoring of microbial community evolution during the experiment simulating gas (CH<sub>4</sub>, 1% CO<sub>2</sub>) storage in a deep aquifer under 60 bar and 36 °C conditions. The vertical dotted line separates samples collected before and after the injection of approximately 100 ppm O<sub>2</sub> on the 50th day of incubation. (a) Redox potential (mV) monitored throughout the experiment. (b) Tracking the taxonomic diversity of the microbial community at the family level based on the 16S rRNA gene. (c) Monitoring active taxonomic diversity of the microbial community based on 16S rRNA gene transcripts (relative quantification) and absolute RT-qPCR quantification of 16S rRNA gene transcripts (v4-v5) and dsrB (diversity of sulfate reducers) gene transcripts.

transcripts. The quantification results of the 16S rRNA gene transcripts (Fig. 4(c)) corresponded well to the observations made from DNA (Fig. 4(b)) and showed that although the microbial community evolved, it maintained its activity throughout the experiment (1.83  $\times$  10<sup>4</sup>  $\pm$  1.14  $\times$  10<sup>2</sup> to 1.50  $\times$  $10^6 \pm 7.26 \times 10^4$  16S rRNA transcript copies per mL). In addition, 16S rRNA gene concentration monitoring indicated a microorganism concentration of 3.93 imes 10<sup>5</sup>  $\pm$  1.31 imes 10<sup>4</sup> 16S rRNA gene copies per mL at the start of the experiment and  $1.51 \times 10^5 \pm$  $2.26 \times 10^4$  16S rRNA gene copies per mL at the end of the experiment (ESI Table S2†). Throughout the experiment, mcrA genes were detected in small quantities in the sampled aqueous phase. However, transcripts showed that those microorganisms were not active (Fig. 4(c)). At the end of the experiment, 96% of the microbial community was dominated by 6 microbial taxonomic groups: the class Acidobacteriae (64%), the family Spirochaetaceae (16%), the family Kineosporiaceae (9%), the family Desulfitobacteriaceae (5%), the class Bathyarchaeia (1%) and the family Syntrophobotulaceae (1%). Close OTUs of the Acidobacteriae class were related to environmental sequences (accession number: AF407714; 375 nt/99.7% identity and acc#: KP676769; 375 nt/99.7%) from Australia's Great Artesian Basin (52 °C) and Russian hot springs. OTUs affiliated with the Spirochaetaceae family were related to bacteria, in particular Rectinema cohabitans, from microbial communities degrading aromatic hydrocarbons (acc#: JF800779; AY214182; KP297860). OTUs related to the family Kineosporiaceae were affiliated with a bacterium described to have phosphate-solubilization activity44 (NII-1013). OTUs grouped in the Desulfitobacteriaceae family were affiliated with the genus Desulfosporosinus, with environmental sequences found in deep (acc#: KX822015, MF942638, JX470444, etc.) and acidic (acc#: DQ137901, GY127803, GQ342329, etc.) ecosystems. OTUs clustered in the class Bathyarchaeia were related to environmental sequences detected in consortia involved in the degradation and fermentation of organic matter (acc# CU916921; KJ424517; U81774). Finally, close OTUs of the Syntrophobotulaceae family were affiliated with environmental sequences of deep aquifers (acc#: LC198569).

#### Evolution of the solid phase from the beginning to end of the experiment

Analyses of the reservoir rock before and after the experiment were carried out to identify potential changes in mineral phases and porosity. X-ray diffraction patterns identified quartz as the dominant phase (81%), while calcite was also present (12%) at the beginning and at the end of the experiment (Fig. S2†). Clay minerals were detected as minor phases (kaolinite and illite < 2%) as well as muscovite (<4%). Iron sulfides such as marcasite were possibly present as trace phases. The results confirmed that the

reservoir rock was mainly composed of quartz, with a minor presence of calcite and clay minerals. Furthermore, samples analysed by scanning electron microscopy (SEM) showed the additional presence of feldspars. Ti-oxide minerals (likely rutile), as well as rare zircon and gypsum crystals, were initially contained in the reservoir rock. At the end of the experiment, no substantial changes were identified except the formation of microcrystals containing zinc oxides, which was observed by SEM-EDS.

Similarly, X-ray tomography was conducted on samples before and after the experiment. The analyses did not reveal any morphological alterations of the solid phase. The grey level of the aqueous phase did, however, exhibit slight variations by the end of the test. Such a difference can be a proxy for the presence of microorganisms. In fact, although microorganisms are principally composed of water, their density<sup>45</sup> (between 1 and 1.05) endows them with a slightly higher attenuation coefficient than that of the surrounding water. As a consequence, microorganisms appear slightly brighter in reconstructed images. To further explore this observation, the voxels constituting the water phase were separated into higher and lower attenuating classes, corresponding to brighter and darker pixels, respectively. The threshold between the two classes was set to the average grey level of all voxels constituting the water phase. The median value was also tested and yielded very similar results.

Fig. S3† depicts three orthogonal cuts through one of the obtained datasets. The images were colour-coded, where red was the solid grains, blue was the water phase and grey was the gas phase. It was immediately clear that the water phase mainly remained at the bottom of the sample, while the gas phase was mainly found in the upper portion of the capillary. The interface between the two phases was not flat, as it resulted from the complex interplay between capillary forces and gravity. In the water phase, the presence of small green areas was noted. This corresponded to the region where higher attenuation was found, which was explained by a higher-than-average concentration of microorganisms. As observed, microorganisms were concentrated in areas close to the grains, and the centres of the pores remained vacant.

#### Discussion

Similar to natural gas, massive amounts of biomethane are planned to be stored in UGS such as salt cavities and deep aquifers. <sup>9,46</sup> In this paper, the influence of a 100 ppm O<sub>2</sub> injection into deep aquifers was studied, particularly its impact on the microbial life. In fact, many studies have confirmed and presented a direct effect of O<sub>2</sub> on autochthonous microbial communities. <sup>9,13,26</sup> The same study site with the code name Ab\_L\_1 was used to study the impact of 1% O<sub>2</sub> on biomethane storage. <sup>47</sup> The aqueous phase had very low salinity, less than 0.8% of marine water salinity. During the experiment, the system was maintained under anoxic conditions for 49 days. O<sub>2</sub> was injected on day 50, and the system was further studied for 36 days.

#### Evolution of the system before O2 injection

During the first phase of the experiment, the reactor represented an UGS of natural gas at 60 bar and 36 °C. For 49 days, the liquid,

gas and solid phases and microorganisms interacted and a decrease in benzene and toluene was observed in the liquid phase (Fig. 3). During the first days, the sulfate content slightly increased in the aqueous phase. Initially, the measured sulfate in the aqueous phase was very low at approximately  $2.85 \times 10^{-2} \pm$  $1.42 \times 10^{-3}$  mmol. Once in contact with the solid phase, this quantity increased and reached  $3.37 \times 10^{-1} \pm 1.89 \times 10^{-2}$  mmol. Based on the reservoir rock analyses, the increase in the sulfate content may be explained by the dissolution of gypsum (CaSO<sub>4</sub>) as identified by SEM-EDS or barium sulfate (BaSO<sub>4</sub>) as detected by XRD (Fig. S3†) or from residual drilling brines contained in the pores (some might have remained even though the rock was washed with demineralized water). The same results and sulfate increase were obtained in previous experiments using aquifer reservoir rock. 9,10 After the increase, a consumption of sulfate was observed in the system until complete depletion on day 39. This observation reoccurred after sulfate supplementation on day 39, and the sulfate content decreased again under anoxic conditions (Fig. 1(a)). Furthermore, previous studies on the Ab\_L\_1 aquifer identified SRM activity and provided evidence for the presence of the Desulfovibrionaceae family.4,47

The presence of a necromass in the microbial community coinjected with the formation water in the HP reactor could explain the growth of SRM such as those belonging to the Desulfovibrionaceae family and fermentative microorganisms such as Spirochaetaceae, which lead to the production of organic acids such as acetate, CO2 and H2. These compounds could benefit the growth of methanogenic archaea and lead to the potential disappearance of sulfate (Fig. 4(b)). In fact, the Methanobacteriaceae and Methanothermobacteriaceae families include microorganisms described to be hydrogenotrophic, while the Methanosaetaceae and Methanosarcinaceae families include acetotrophs and methylotrophs.48 Fermentation continued during the first 20 days of incubation and potentially produced H2.49 Therefore, autotrophic metabolisms explain why the observed gaseous phase value of CO2 was lower than the theoretical value predicted from CO<sub>2</sub> dissolution into the aqueous phase (Fig. 2). The thermodynamic calculations using the PhreeqC model based on an abiotic binary system suggested that other phenomena contributed to the experimentally observed disappearance of CO<sub>2</sub>. Some microorganisms identified in this study at the start of the experiment, e.g., Methanobacteriaceae and Methanothermobacteriaceae, and a few Desulfovibrionaceae, which have lithoautotrophic metabolisms that employ H2 (from fermentation) as an energy and electron source and CO2 as an electron acceptor and carbon source for biomass production, could have played such a role in CO2 removal.

Among the SRMs, the physicochemical conditions and the presence of rock seemed to have favoured the growth and activity of the genus *Desulfurispora*, which are known to incompletely oxidize organic substrates to acetate. The quantity of the latter increased in the liquid phase before the injection of O<sub>2</sub> (Fig. 1(b), 4(b and c)). Since fermenters such as *Spirochaetaceae* and SRMs such as *Desulfurisporaceae* were not competitors for electron acceptors, the high relative decrease in the activity of *Spirochaetaceae* observed on the 36<sup>th</sup> day could be due to a decrease in the concentrations of accessible organic

substrates in the reactor. From the 36<sup>th</sup> day, the decrease in the quantity of sulfate (<1 mmol) and the probable limitation in organic substrates seemed to favour the members of the *Desulfitobacteriaceae* family to the detriment of the *Desulfurisporaceae* family.

#### Evolution of the system after O2 injection

As part of the experiment, 100 ppm O<sub>2</sub> was injected into the gas phase on the 50<sup>th</sup> day of incubation. It is important to specify that the rock material of the targeted UGS site consisted mainly of quartz, which poorly reacts with oxygen. Only a limited amount of O2 could have been trapped a priori following mineral oxidation. Therefore, the effects were mainly observed in the liquid phase and the microbial community. Injection had several effects: (i) an increase in the redox potential to +73 mV; (ii) cessation of sulfate reduction and stabilization at 2.82  $\times$  $10^{-1} \pm 1.41 \times 10^{-2}$  mmol until the end of the experiment (40 days); (iii) cessation of the appearance of acetate and stabilization of its concentration at 1.56  $\pm$  7.80  $\times$  10<sup>-2</sup> mmol; (iv) cessation in the disappearance of CO<sub>2</sub> from the gaseous phase. These four effects were explained by the community modifications induced by the introduction of O2. SRMs that are under strictly anoxic conditions suffered from oxygen toxicity. Under conditions simulating the aquifer (60 bar and 36 °C), the O2 concentration was estimated by calculations to be approximately  $5.8 \times 10^{-2}$  mM in the aqueous phase. Two days after the injection of O2, Desulfurisporaceae represented only a small portion of the active microorganisms (Fig. 4(c)). The apparent antagonism between the disappearance of sulfate-reducing activity demonstrated by the absence of dsrB transcripts and the identification of numerous SRMs affiliated with the Desulfitobacteriaceae family (i.e., Desulfosporosinus sp.) may explain the versatility of their metabolism and their ability to ferment or use other terminal electron acceptors.51 These microorganisms have been identified in groundwater contaminated with BTEX and are regularly shown to be key players in the biodegradation of these monoaromatic hydrocarbons. 52-56 Interestingly, it seems that the cessation of sulfate reduction following O2 injection did not slow the degradation of benzene in our experiment, which could be explained by the microbes' ability to ferment this compound.57 The taxonomic group that was mainly favoured by the new conditions included bacteria affiliated with the class Acidobacteriae, which account for more than 80% of the microbial activity (Fig. 4(c)). The development of these taxonomic groups (Desulfosporosinus and Acidobacteriae) is not surprising since they are associated with acidic pH conditions.51,58 Although the pH could not be monitored in this experiment, the work of Sin and her collaborators<sup>25</sup> in another storage aquifer showed that the injection of O<sub>2</sub> and the presence of CO<sub>2</sub> led to an acidification of the formation water due to the dissolution of certain minerals such as pyrite. Slight acidification has previously been shown to be conducive to the biodegradation of BTEX.5,10 Finally, the new microoxic conditions also allowed the development of members of the Kineosporiaceae family (Quadrisphaera spp.), which are described as aerobic microorganisms<sup>59</sup> but can be assumed to

be able to ferment to maintain themselves in deep anoxic aquifers since they are unable to sporulate.

At the end of the experiment, the solid phase demonstrated no substantial changes except for the formation of microcrystals containing oxides (ESI Fig. S2 $\dagger$ ). X-ray tomography scans confirmed the absence of significant alterations (ESI Fig. S3 $\dagger$ ). Therefore, even though ionic compounds were released into the aqueous phase, the nature of the reservoir rock, which was mainly composed of quartz, was not strongly impacted by the O<sub>2</sub> injection. Finally, X-ray data provided a strong indication that microorganisms were not distributed uniformly throughout the aqueous phase but rather clustered in close proximity to grain boundaries.

#### Conclusions

Biomethane storage in underground aquifers is a mandatory step to ensure the energy transition and to substitute natural gas. During our experimental study, a deep aquifer situated in France was reproduced in a high-pressure reactor under a pressure of 60 bar and a temperature of 36 °C. As designed, the reactor cannot simulate a continuous flow of water in the rock, as in the aquifer. However, a small amount of water is displaced as the piston moves to rebalance the pressure following the various withdrawals during the experiment. Finally, we believe that we are very close to the reality, since the movement of the formation water is very slight (of the order of a meter or a few meters per year). To summarize the effects observed after the injection of 100 ppm of  $O_2$  in the biomethane mixture:

- The cessation of bacterial sulfate reduction and a change in the microbial community were detected. O<sub>2</sub>'s effect on the microbial community was not limited to sulfate reducers only, but greater changes occurred.
- Even though the ecosystem varied, the biodegradation of benzene was maintained.
- Conversely, no remarkable changes were observed in the solid phase.
- ullet The same on the gas phase initially composed of CH<sub>4</sub> with 1% CO<sub>2</sub>, and decrease of the latter one was detected. CO<sub>2</sub> consumption was related to microbial activity as their source of carbon. Thus once the amount of O<sub>2</sub> was injected and the microbial activity became very slow, CO<sub>2</sub> consumption stopped.

Shifts in the microbial community due to  $O_2$  injection can have larger impacts on aquifer's evolution and stability. In fact, existing and active microorganisms in the aquifers today are responsible to degrade hydrocarbons present *in situ*. Studying the effect of  $O_2$  as an impurity during biomethane storage has an added value to fix the  $O_2$  limit in biomethane. Here, injecting 100 ppm  $O_2$  into a UGS in a deep aquifer does not appear to affect its operation. These results compose developed knowledge about  $O_2$  biological geological and chemical impact on the underground. The effects of  $O_2$  on UGS must be correlated with  $O_2$  effects on the gas network facilities used today. Findings can be used in any other process including  $O_2$  as an impurity such as CCS and CAES. Further studies applied on aquifer storage are needed to dissect the multiple results of our findings in order to optimize  $O_2$  injection in the underground and possibly increase

the regulatory limits for O<sub>2</sub> to enhance the development of the biomethane sector in Europe.

#### **Abbreviations**

BTEX: Benzene, toluene, ethylbenzene, xylene

CAES: Compressed air energy storage

CCS: Carbon capture and storage

CH<sub>4</sub>: Methane
CO<sub>2</sub>: Carbon dioxide
H<sub>2</sub> Hydrogen

 $\begin{array}{lll} \text{H}_2\text{S} & \text{Hydrogen sulfide} \\ \text{HP} & \text{High pressure} \\ \text{N}_2 & \text{Nitrogen} \\ \text{O}_2 & \text{Dioxygen} \end{array}$ 

SRM Sulfate reducing microorganisms

UGS Underground gas storage

#### **Author contributions**

FC, MRP, PC and ARP co-conceived the study. PH, JM, FC, and PC carried out the high pressure and simulation experiments. MG, MRP and ARP applied the microbiological approaches. PS, M-PI, PM and GH used the imaging and mineralogical characterization. ML and ILH analyzed the hydrocarbons. All authors contributed to the interpretation of results and paper writing.

#### Conflicts of interest

PCh, GC and AP are employed by two French companies specialized in geological natural gas storage, which are Teréga and Storengy, respectively.

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