1 2 3 4	The chemistry of hyperalkaline springs in serpentinizing environments: 1. the composition of free gases in New Caledonia compared to other springs worldwide.
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22 23	Key Points:
24	• Hyperalkaline spring waters are anoxic. An elevated high O2 content in free gases results from air
25	contamination.
26	• The relationship between H_2 and CH_4 contents of free gases is consistent with the conversion of fully
27	oxidized carbon (CO ₂).
28	• The H ₂ -CH ₄ relation is consistent with a Sabatier reaction in the gas or a methane production by
29	hydrogenotrophic methanogens in the water.
30	

32 Abstract

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34 Serpentinization is a natural process that transforms ferromagnesian minerals such as olivine into 35 serpentine and that produces waters at very high pH and gases enriched in methane (CH₄) and 36 hydrogen (H₂). We report the composition of gases venting at two springs (Bain des Japonais and 37 Rivière des Kaoris) of the serpentinizing environment of the Prony Bay (New Caledonia) collected 38 eight times between 2011 and 2014, along with in situ measurements (temperature, pH, oxydo-39 reduction potential, dissolved oxygen content) of on-land alkaline springs of the Southern New 40 Caledonia ophiolite. Venting gases are mainly composed of H₂, CH₄ and N₂ and their composition has slightly varied during the 4-year field survey. An elevated oxygen (O₂) content in a high-pH water 41 42 sample is due to air uptake during surface flow. O2-corrected gas compositions along with those published for gas data obtained at similar serpentinizing environments (Italy, Turkey, Philippines, 43 44 Oman) show that the H₂ and CH₄ concentrations display a linear correlation with a slope close to the value corresponding to the CH₄ production from carbon dioxide rather from a less oxidized carbon 45 such as carbon monoxide. Although these data are consistent with the stoichiometry of the Sabatier 46 47 reaction, as such in the gas phase, it is also possible that microbial hydrogenotrophic methanogenesis 48 takes place in the aqueous phase followed by degassing. A diagram is proposed that outlines the partitioning of H₂ and CH₄ between the gaseous and aqueous phases and the need to consider a two-49 50 phase flow in the hydrology of these hyperalkaline environments.

51

53 Plain Language Summary

54 Serpentinization is a natural process that transforms ferromagnesian minerals such as olivine into 55 serpentine. This process is accompanied by the formation of high-pH water, hydrogen and methane. This forms a unique ecological niche for microorganisms adapted to extreme pH and/or involved in 56 57 the hydrogen and/or methane cycles. There is an ongoing debate on the mechanisms at play and on the contribution of abiotic versus biogenic reactions. An analysis of the composition of free gases 58 collected in hyperalkaline springs of New Caledonia combined with literature values for similar sites 59 60 worldwide shows that the formation of methane follows the stoichiometry of a Sabatier-type reaction 61 involving fully oxidized carbon (CO₂) and not carbon monoxide (CO) as required by a Fischer-Tropsch-Type process. Although it could be concluded that methane production occurs in the gas 62 63 phase, and as such would be abiotic, it does not rule out the possibility that the relationship between gaseous hydrogen and methane is due to methane production and hydrogen consumption by 64 microorganisms in the aqueous phase followed by degassing. 65

67 1 Introduction

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Natural environments with waters at pH above 11 have received considerable attention in the 69 70 last decades due to the wide range of scientific questions that they relate to : abiotic formation of 71 hydrocarbons (e.g. [Etiope and Sherwood Lollar, 2013; Price et al., 2017; Sissmann et al., 2019], 72 origin of life and deep subsurface biosphere (e.g., [Menez et al., 2012; Schrenk et al., 2013], alteration of the oceanic crust (e.g. [Debret et al., 2019; Fryer et al., 2020; Ligi et al., 2013], natural formation 73 of hydrogen as an energy source (e.g. [Truche et al., 2020], geological storage of CO₂ [Kelemen and 74 75 Matter, 2008; Kelemen et al., 2011], the analogy with planetary surfaces like Mars or Europe [Holm et 76 al., 2015; Szponar et al., 2012], energy and matter fluxes between geochemical reservoirs [Plank and 77 Manning, 2019].

78 Sites of discharge of such very high pH waters are found on land in Oman [Chavagnac et al., 79 2013b; Miller et al., 2016; Paukert et al., 2012; Stanger, 1985], in Northern Italy; [Boschetti and 80 Toscani, 2008; Chavagnac et al., 2013b; Cipolli et al., 2004], in Cyprus [Neal and Shand, 2002], in the Philippines [Cardace et al., 2015], in California [Boschetti et al., 2017; Morrill et al., 2013], in 81 82 Ontario [Sader et al., 2007], in Newfoundland [Szponar et al., 2012], in Portugal [Marques et al., 83 2008], in Greece [D'Alessandro et al., 2018a], in Turkey [Meyer-Dombard et al., 2015; Yuce et al., 2014], in Jordan [Khoury et al., 1992], in New Caledonia [Monnin et al., 2014] and in Spain 84 [Giampouras et al., 2019]. Such hyperalkaline waters are also found underwater in the ocean, at the 85 Lost City Hydrothermal Field on the Atlantis massif off the Mid-Atlantic ridge [Kelley et al., 2001; 86 87 Kelley et al., 2005] and in the mud volcanoes of the Marianna forearc [Fryer et al., 1989; Fryer et al., 88 2018; Mottl et al., 2004]. The Maqarin site in Jordan is a peculiar case where the natural combustion 89 of bituminous marls and the alteration of the rocks created by a phenomenon that has been called 90 pyrometamorphism lead to the formation of extremely alkaline water, with pH value up to 12.5 91 [Alexander et al., 1992; Khoury et al., 1992]. Apart from the Magarin site, these hyperalkaline waters 92 originate from the serpentinization of ultramafic rocks that can be found on land in ophiolites (e.g. 93 Oman, Liguria, etc.), in an ultramafic complex created by continental collision such as Cabeço de Vide 94 in Portugal [Marques et al., 2008; Marques et al., 2018], at the seafloor where plate tectonics has 95 exposed ultramafic rocks to alteration, such as at the Lost City site (Northern Atlantic), or at the 96 decollement zone of the subducting plate in the case of the Marianna forearc mud volcanoes [*Mottl et* 97 *al.*, 2004]. The discharge of these waters is focused and occurs at springs where they lead to the 98 formation of carbonate concretions, in the form of tuffs in Oman [*Chavagnac et al.*, 2013a] or 99 submarine pinnacles at Lost City [*Kelley et al.*, 2005] and in the Prony Bay in New Caledonia [*Launay* 100 *and Fontes*, 1985; *Monnin et al.*, 2014; *Pelletier et al.*, 2006].

While the discharge zones (springs) are easily recognized in the field, the recharge of these 101 102 hydrological systems is poorly or not at all characterized. The geographic extension of a given 103 hydrothermal system, the characterization of the inflowing water, its residence time and the role of the 104 composition of the geological formation (its lithology, its physical and thermal structure) are general 105 questions on any hydrological system. They are rarely documented for hyperalkaline hydrothermal 106 systems [Christofi et al., 2020; Dewandel et al., 2005; Lods et al., 2020; Marques et al., 2018]. A 107 classification of high-pH springs based on the location of the output (marine or continental) and on the 108 origin of the circulating fluid based on the salinity of the spring waters has been proposed [Monnin et 109 al., 2014]. Thus, a first category is defined by the fresh high-pH spring waters found in continental environments like in Oman, Liguria, Portugal, Philippines, Greece, and Turkey. The second main 110 category contains hyperalkaline waters with salinities similar to that of seawater discharging at springs 111 112 located in the marine environment, the Lost City site being the archetype of this category [Kelley et 113 al., 2001; Kelley et al., 2005; Seyfried Jr et al., 2015]. This is also the case of the Marianna mud 114 volcanoes, where the high-pH waters have seawater-like salinities, but a very complex geological 115 history linked to their origin in the dehydration of the subducting Pacific plate in the Marianna 116 subduction zone [Hulme et al., 2010; Mottl et al., 2004]. In nearly every case, the inflowing water 117 (recharge) of these hyperalkaline hydrothermal systems is continental (meteoric, runoff) water (low 118 salinity) for systems fully located on continents and seawater for fully marine environments.

119 Two hyperalkaline hydrothermal systems do not fit in this classification. The Aqua de Ney 120 spring is located inland about 200 km from the Northern California Coast. Its water (pH \approx 12) has a 121 salinity similar to that of seawater and is a case of a high-pH saline water discharging in a continental 122 environment [*Blank et al.*, 2017; *Feth et al.*, 1961]. It has been proposed that it originates from the dehydration of the Pacific plate subducting below the American continent [*Boschetti et al.*, 2017]. The
other system that does not fit in the classification is in Prony Bay (southern lagoon of New Caledonia)
where alkaline springs located in the lagoon discharge fresh waters into the marine environment
[*Monnin et al.*, 2014].

127 Globally, these hyperalkaline waters contain elevated concentrations of methane and hydrogen 128 [Etiope and Whiticar, 2019]. Gas venting is commonly observed but not at every location, while 129 dissolved gases have been analyzed in numerous hyperalkaline springs (see the review of [Etiope and 130 Whiticar, 2019]. Gas bubbles have been sampled in Oman [Boulart et al., 2013; Vacquand et al., 2018], in the Voltri ophiolite in Liguria (Northern Italy) [Boschetti et al., 2013; Boulart et al., 2013; 131 132 Etiope and Whiticar, 2019], at several locations in the Philippines [Abrajano et al., 1988; Vacquand et al., 2018] and in Turkey [D'Alessandro et al., 2018b; Vacquand et al., 2018] and in New Caledonia 133 [Vacquand et al., 2018]. The gases are mainly composed of dihydrogen (H₂), dinitrogen (N₂) and 134 135 methane (CH₄), with trace amounts of higher alkanes, carbon dioxide and monoxide and helium. 136 Oxygen is not always measured or reported. It can be found in noticeable concentrations, for example 137 up to 15% for one of the samples of the Kisecik spring in the Kizildag ophiolite in Turkey [D'Alessandro et al., 2018b] or the Zambales springs in the Philippines [Abrajano et al., 1988]). 138

The Prony Bay hydrothermal field (PBHF), that has been studied during the HYDROPRONY 139 140 cruise in 2011 [Pelletier et al., 2011], comprises several underwater springs in the Prony Bay, with the 141 Aiguille de Prony being the most prominent, and two costal main sites at the northernmost part of the 142 Prony Bay, the Bain des Japonais, a spring that is uncovered at low tide and the Bain des Kaoris that is 143 located above the water level [Monnin et al., 2014]. A number of other springs and seepages of high pH waters are known inland from the coast (Grande Terre) many 10s of kilometers north from Prony 144 145 Bay ([Maurizot et al., 2020]; Fig. 1). We here present in situ measurements of pH, oxidation-reduction 146 potential (ORP), dissolved oxygen and temperature of these on-land spring waters, along with the compositions of free gases collected at the Bain des Japonais and at the Kaoris springs during a survey 147 carried out between 2011 and 2014. 148

149 The oxic/anoxic characteristic of the high pH waters has been determined from in situ 150 measurements of the oxygen content directly at the springs, but also from the gas chromatography

151	analysis of the free gases. We address the question of air contamination of the samples and propose a
152	way to correct it. These new data are then compared to sites of alkaline springs in other countries
153	where gas venting is observed. This allows to address the question of the stoichiometry of methane
154	production and hydrogen consumption and therefore the nature of the carbon pool in low temperature
155	serpentinizing environments.
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157 158	2 Location and geological setting of the springs
159 160	2.1 The springs of the Prony Bay
161 162	The Prony Bay springs have been previously described ([Monnin et al., 2014] and references
163	therein). The Kaoris spring is located above the sea level at the northeastern end of the Bay along the
164	Kaoris River. The Bain des Japonais, which is submerged at high tide, is located on the Baie du
165	Carénage, at the northwestern end of the larger Prony bay [Monnin et al., 2014]. These two sites, 2.2
166	km apart, are easily accessed using a shallow-draft boat. They have been visited several times between
167	2011 and 2014 (Table S2).
168	
169 170	2.2 On-land springs
171	The spring north of the Yaté Lake (called Nord Yaté) has been visited only once on August 8,
172	2013, while the other on-land springs were sampled during the 2014 fall (Nov. 17 to Dec. 15) (Fig. 1).
173	



175 Figure 1 – Location of the high pH springs in southern New Caledonia (after [Maurizot et al.,
176 2020]).

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The spring "La Coulée" is located in the outskirts of Nouméa in the valley of the river "La Coulée" and is easily accessed. The main water outlet is about two meters above the river bed in a shallow (about 10-cm deep) pool (Fig. 2). The pools below are fed by water overflowing from the upper pool. Very intermittent gas bubbling has been observed in the upper pool, but no sample could be taken because of the very slow discharge (a bubble every 10 mn or so).

The spring "Rivière des Pirogues" is a little bit further away and can be accessed by dirt roads. The high pH water outlet is a seepage located within the bed of a tributary of the Pirogues River in fractured rocks above a large (about 30 m wide) water pool formed at the junction with the Pirogues river. Bubbling in this pool has been observed about two meters from the rocky bank, but again no sample could be taken. This spring does not build carbonate pools like the spring "La Coulée" (Fig. S1). 189 The spring of the Montagne des Sources (also named Rivière du Rocher) is located in a reserve. The access is totally restricted because this is where the city of Nouméa is getting its water supply 190 191 from (Fig. S1). A strenuous 5-hour hike in the river bed is required to get to the spring. The spring is a 30-m long spectacular suite of large pools located right above the river bed and below a forest (Figure 192 S1). These pools are about 1.5 m in diameter and about 1.5 m deep for the deepest one. Water flows at 193 a high rate through an outlet situated in the middle of the wall of the structure. The pools contain 194 195 varying amounts of branches and leaves from the trees above. Sampling and observations have been difficult and less extensive than anticipated because of the very bad weather and heavy rain on the day 196 197 of our visit (Dec. 3, 2014).

The spring on the Northern side of the Yaté Lake cannot be reached by land. Water samples were collected during a visit to the spring by helicopter on August 8, 2013, therefore with limited time on the site. Similarly to the Coulée spring, it is located right above the river bed (Fig. S1). The highpH water builds up small terraces that are common characteristics of these peculiar springs, as can be observed for example at La Coulée spring (Fig. 2), in Oman and Liguria [*Chavagnac et al.*, 2013a; *Chavagnac et al.*, 2013b] or in California at the Cedars spring [*Suzuki et al.*, 2013].

Although they have been included in previous studies of New Caledonia alkaline springs [*Cox et al.*, 1982; *Deville and Prinzhofer*, 2016; *Vacquand et al.*, 2018], the springs of the Canala area are somewhat different than those further South. The geological substratum is different and the pH (\approx 9) of its waters is lower than that at the other studied sites [*Cox et al.*, 1982; *Maurizot et al.*, 2020; *Quéméneur et al.*, 2021].

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3 Sampling procedures, in situ measurements and analytical methods 211

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3.1 Free gas sampling procedure and in situ measurements: pH, oxidation-reduction potential, temperature and dissolved oxygen

Gas samples were collected in 10-ml glass vials by the water displacement technique. It consists in filling the vial with spring water and immersing it upside down in the spring above the gas bubbles. The captured bubbles displace the water from the vial which is then closed with a rubber stopper when there is no water left (then sealed with an aluminum cap). It was noticed that an air bubble can be trapped in the rubber stopper, leading to air contamination. Even if great care is taken, air uptake during sampling is not totally unavoidable. For example, several gas compositions reported by [*Neal and Stanger*, 1983] and [*Sano et al.*, 1993] for Oman show that almost pure air has been sampled.

The samples were kept at 4°C in the laboratories in Nouméa (IRD) and in Marseille (MIO). They were kept at room temperature (i.e. plane cargo bays and lockers) while travelling between New Caledonia and metropolitan France (i.e. for about 30 hours).

225 Temperature, pH, oxidation-reduction potential (ORP) and the dissolved oxygen content were 226 measured directly in the springs with the appropriate probes using a wTW Multi 3420® 227 Multimeter that allows recording the measurements over a time period with a fixed step (Fig. S2). Temperature and pH were measured during all the campaigns using a SenTix 940® pH electrode in 228 229 which a thermocouple allows temperature to be measured. ORP (Oxidation-Reduction Potential in 230 reference to the Ag/AgCl standard) and dissolved oxygen were measured only in 2014 after the acquisition of the appropriate probes (SensoLyt ORP 900-P® for the redox potential and FDO 925® 231 232 for dissolved oxygen).

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234 3.2 Gas analyses

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The gas analyses (dissolved and gaseous) were carried out in three different laboratories: MIO (Mediterranean Institute of Oceanography) in Marseille (France), at the Laboratoire des Cycles Géochimiques et ressources (LCG) IFREMER Brest (France) and at the NASA Ames Research Center (Moffett Field, California).

At MIO (Marseille) the gas composition was determined using a Shimadzu GC 8A gas chromatograph equipped with a thermal conductivity detector (GC/TCD) and a concentric column CTR1 (Alltech, USA). Argon was used as carrier gas at a flow rate of 60 mL/min; temperature of the injector and the detector was fixed at 150°C. All analyses were run in duplicate.

At in the NASA Ames Research Center, The concentrations of free gases were analyzed by a Shimadzu GC 8A gas chromatograph equipped with a thermal conductivity detector (GC/TCD) and two columns (Alltech, USA). Operating conditions for H_2 , O_2 , CH_4 , and CO_2 analyses are as follows: the carrier gas is nitrogen at a pressure of 100 kPa; temperature of the column is fixed at 150°C and that of the injector and the detector is fixed at 200°C. N_2 was determined by difference based on previous estimates [*Monnin et al.*, 2014].

At IFREMER Brest, two analytical devices have been used: a gas chromatograph µGC R3000 250 from SRA equipped with three analytical modules to determine hydrogen, oxygen, nitrogen, methane 251 252 to hexane and carbon dioxide concentrations, and a Agilent 7890A gas chromatograph equipped with 253 a 32 m, 0.32mm Porapak Q column and a triple detection system (TCD-FID-mass detector) to analyze hydrocarbons at low concentrations. The TCD and FID are used for quantitative analysis while the 254 255 mass spectrometer is used for the purpose of verification of the compounds or identification of unknown compounds against the NIST05 library. The injection and calibration were performed as 256 257 described by [Donval et al., 2020].

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The data and the laboratories where the analyses have been made are reported in Table S2.

259 4 Results

4.1 In situ measurements: temperature, pH, ORP and dissolved oxygen

263 4.1.1 Temperature and pH264

The in-situ parameters (Table S1) were recorded during 30 mn at the springs of La Coulée and Pirogues. Dissolved oxygen and ORP required an equilibration time of a few minutes (Fig. S2). The water temperatures at all the on-land springs are between 26 and 32°C, similar to that for the Kaoris (32°C) but a little bit lower than those for the Bain des Japonais (38°). At La Coulée, the water temperature measured in the upper pool is 27°, lower than the air temperature which was 30° or even higher on the days of sampling.

pH is constant at Pirogues at a value of 10.88 but varies over 0.2 pH units at La Coulée between
10.81 and 11.05 (Table S1). There is a correlation between temperature and pH at La Coulée in the
upper pool (Fig. S2). The temperature variation can be due to a convection effect of the thin water

layer. Indeed, the discharging water temperature is between about 27°C while the air temperature on
the days of measurements were over 30°C.

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277 4.1.2 The dissolved oxygen content

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At La Coulée, the water discharges in the upper pool from where it flows down to the other pools. The oxygen content of the water in the upper pool is 0.4 mg/L, 7.56 mg/L in the lower pool and 8.24 mg/L in the river (Fig. 2). The Eh values are the same in all the pools. The water discharging at the upper pool becomes oxygenated by air uptake when it flows downhill. This air and thus the CO₂ uptake lead to a small decrease in pH of 0.4 unit. Thus, for a given spring, variable values of *in situ* parameters can be obtained when measured away for the outlet.

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Figure 2 – The high-pH spring of La Coulée. The water discharges in the upper pool (1) and flows
down toward the river through the upper (2) and lower (3) intermediate pools. It gets oxygenated on

the way, as indicated by the elevated O_2 content in the lower pool (4), at a value close to that of the river. n.m.: not measured.

The Pirogues and Coulée springs have similar oxygen content between 0.2 and 0.4 mg/L, which classifies them as anoxic [*Berner*, 1981]. At the Montagne des Sources, the oxygen content of the very large and deep pool is 3.0 mg/L while the water flowing at a fast rate from the outlet in the middle of the formation (Fig. S1) is quite oxygenated (with a O₂(aq) content of 6.9 mg/L; Table S1).

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296 4.1.3 The oxidation-reduction potential

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The ORP values are measured in situ in reference to the Ag/AgCl standard. They have been corrected for the potential difference between the Ag/AgCl and the hydrogen electrodes and thus termed Eh (Fig. S2). There are marked differences in the Eh values between the springs, between the days of measurement for the same spring and sometimes even within the time of recording (30 mn).

Eh at the spring Rivière des Pirogues has markedly varied between Dec. 2 and 12, 2014 (Fig 302 303 S1). There is also a small Eh maximum at Pirogues for the data of Dec. 2. Very sporadic bubbling (a few events per hour) have been observed at La Coulée and also at the Pirogues site. For this latter 304 spring bubbles were observed not in the seepage of the spring where the water flows on the rocks 305 306 without being retained in a pool, but on the side of the very large pool (about 20 m wide) built by the 307 river about 10 meters downstream from the spring (Fig. S1). It has been shown in many instances that 308 gas bubbles forming in these high pH springs contain elevated concentrations of hydrogen (along with 309 methane and nitrogen). This has been documented for the Prony bay [Deville and Prinzhofer, 2016; 310 Monnin et al., 2014]. We have observed at the Bain des Japonais in the Prony bay that gas bubbling 311 disrupts the ORP measurement when a gas bubble hits the probe. This may explain the Eh maximum 312 and the increase from -300 to -200 mv in Dec.2 at Pirogues (Fig. S1). This H₂-enriched gas venting 313 may also be the cause of the dispersion of the Eh values.

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315 4.1.4 Comparison with other high-pH springs worldwide

- 317 4.1.4.1 The dissolved oxygen content
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319 Unfortunately, the dissolved oxygen concentration is not reported for every spring or seepage 320 area documented in the literature. The dissolved oxygen concentration of the New Caledonia springs 321 and of other high-pH springs worldwide are displayed versus pH in Fig. 3. These measurements were 322 made using different methods: with an oxygen probe (this work; [Cardace et al., 2015; Etiope et al., 323 2016; Rempfert et al., 2017; Suzuki et al., 2013], on site spectrophotometric measurements based on 324 the indigo carmine reagent [Canovas et al., 2017] and gas chromatography analysis of the dissolved 325 gases using the headspace method [Etiope et al., 2016]. In order to give the order of magnitude of 326 oxygen saturation, we used the dissolved oxygen content of pure water at equilibrium with the 327 atmosphere (i.e. for an oxygen partial pressure of 0.2 bar) at 25°C (8.3 mg/L) (Fig. 3). An accurate 328 calculation would require taking into account the temperature and the composition of the aqueous 329 phase (the Setchenov correction to Henry's constant). For example, oxygen solubility in pure water at 40°C is 6.41 mg/L. The three values reported for river waters close to the alkaline springs in New 330 331 Caledonia and a non-alkaline spring in Ronda (Spain; [Etiope et al., 2016]) are close to values of 332 oxygen saturation in pure water (Fig. 3). The present data for New Caledonia, but also the data for 333 Oman [Canovas et al., 2017], for the Manleluag springs in the Philippines [Cardace et al., 2015] and 334 for the Ronda springs in Spain [Etiope et al., 2016] show that at almost constant pH, the oxygen 335 content of the water samples collected in the high-pH springs can vary from zero to values close to 336 saturation. The waters at The Cedars springs in California [Suzuki et al., 2013] and of the Argolida and 337 Othrys spring in Greece [D'Alessandro et al., 2018a; Etiope et al., 2013] are either totally anoxic or 338 have a very low oxygen content. Water samples collected in boreholes in Oman [Rempfert et al., 339 2017] or at the CROMO site (Coast Range Ophiolite Microbial Observatory) in Northern California [Crespo-Medina et al., 2014] are also oxygen free. The example of La Coulée spring show that 340 341 elevated dissolved oxygen concentrations measured away from the outlet result from atmospheric oxygen dissolution due to air uptake. On the other hand, this air contamination provides carbon 342 343 dioxide to the water that contributes to the formation of the carbonate concretions building the pools 344 [Chavagnac et al., 2013a; Leleu et al., 2016]. Similarly, [Meyer-Dombard et al., 2015] have shown how the carbon content of the waters in the stream created by the discharge of a high pH fluid atChimarea springs (Turkey) increases with the distance from the source because of air uptake.

These data show that the high-pH waters discharging at the springs are oxygen-free, when they are collected directly at the outlet. This is in agreement with the fact that oxygen is the first oxidant consumed in the subsurface either through biological consumption or via the oxidation of ferric iron *[Leong and Shock*, 2020; *Neal and Stanger*, 1983]. It is then concluded that the hydrologic system discharging these high-pH waters is oxygen-free and that the venting gases should not contain any oxygen. Therefore, the compositions of the gases venting at the seeps must then be corrected for this air contamination.





Figure 3 – The dissolved oxygen concentration of the New Caledonia hyperalkaline springs and of other springs worldwide. The dashed line is the value of oxygen saturation in pure water at 25°C. 357 The symbols for New Caledonia are: C = La Coulée, P = Pirogues, M = Montagne des Sources, K =358

359 Kaoris.

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361 4.1.4.2 The oxidation-reduction potential

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The data set is very scattered with no clear relationship between Eh and pH (Fig. 4). The redox 363 364 conditions for the reduction of water are met for the Bain des Japonais and the Pirogues spring in New Caledonia (this work) and the Cedars springs in California. The ORP value for the Tahtakopru spring 365

of the Kilzidag ophiolite in Turkey [Yuce et al., 2014] and for a spring called Carénage in New 366 Caledonia, which corresponds to the Bain des Japonais [Deville and Prinzhofer, 2016] are below the 367 water reduction limit, -800 mV for a pH of 10.52 and -870 mV for a pH of 10.59, respectively. There 368 369 is an ambiguity in the Eh values reported in the literature as the reference potential (Ag/AgCl or H₂) is not always mentioned (e.g. [Paukert et al., 2012]). Some of the values reported in Fig. 4 may be the 370 371 values given by the ORP probe (i.e. in reference to the Ag/AgCl standard) and thus may be too high 372 by about 200 mV (Fig. S2). Such a correction would bring the redox potential of the Carenage and Tahtakopru springs close to the water reduction limit, i.e. corresponding to reaction (1): 373

$$H_2 O(l) + e^- = H_2(g) + OH^-(aq)$$
(1)

These data show that these high-pH waters display the most reducing conditions at the surface of the Earth. No correlation with the oxygen content of the waters was observed.



Figure 4 – The oxidation-reduction potential Eh versus pH. The dashed lines correspond to the water reduction reaction (eq. 2) at 25°C for the two mentioned hydrogen partial pressures. The symbols for New Caledonia are: B = Bain des Japonais, C = La Coulée, P = Pirogues, M = Montagne desSources, K = Kaoris. The Carenage spring is the name given to the Bain des Japonais by [Deville and

- 381 Prinzhofer, 2016]. The Tahtakopru spring is one of the springs of the Kilzilgag ophiolite (Turkey;
- 382 [Yuce et al., 2014]).

384 4.2 The composition of the gases venting at the Bain des Japonais and Kaoris springs 385 The gas samples were collected between October, 2011 and November, 2014 at eight different 386 387 dates in two springs of the Prony Bay (Table S2). At a given date, the concentration values are 388 scattered, between 5 and 40% for H₂ and between 5 and 25% for CH₄ (Fig. 5). H₂ values are similar 389 for the two springs, but CH₄ is lower at the Kaoris than at the Bain des Japonais. Some of the samples 390 contain elevated concentrations of oxygen that we have attributed to air contamination (Table S2). 391 Assuming that the nitrogen content of air is four times larger than its oxygen content and that air and 392 the sampled gases are perfect gases, a corrected nitrogen content (in percentage) in each gas sample 393 can be calculated using:

$$P_{N_2}(corr.) = P_{N_2}(raw) - 4 P_{O_2}(raw)$$
(2)

In eq. 2, "raw" designates the results of the chemical analysis (ionic chromatography). The hydrogen and methane concentrations are then recalculated using this corrected nitrogen concentration.

397 Data reveal a slightly increasing trend for H_2 and slightly decreasing for CH_4 during the four-398 year survey (Fig. 5). The scatter of the data is also due to the number of samples collected at a single 399 date (Fig. 6).





402 *Figure 5 – Uncorrected (filled dots) and O*₂*-corrected (crosses) hydrogen and methane contents*



404

405 When the number of samples collected at a given date allows it (e.g. Oct. 17, 2012; Table S2), 406 the data show a correlation between the methane and hydrogen contents (Fig. 6). This correlation also 407 appears on the entire data set, with different trends for the Kaoris and for the Bain des Japonais. Data 408 depicted in Figs. 5 and 6 also show that the hydrogen content of the gases can fluctuate during the time 409 of sampling on a given day, for example between 20 and 35% on Oct. 17, 2012. Because gas flow is 410 steady at these two springs, it takes less than half an hour to collect 5 samples. It also shows that the 411 conversion of hydrogen to methane (see below) is also very rapid, otherwise the methane concentration would not be correlated to that of hydrogen. In future studies, it will be necessary to 412 413 note, not only the date of sampling, but also the hour.

414



Figure 6 – The $CH_4(g)$ concentration of the Bain des Japonais and the Kaoris springs versus that of $H_2(g)$ for each date and for the whole data set. Open dots: Kaoris; filled dots: Bain des Japonais (BdJ); crosses: Vacquand et al. 2018. The dashed lines are fits to the data and the formulas are the expressions of the linear regressions. The plain lines represent the variation of the methane and hydrogen concentrations due to the reaction of fully oxidized carbon (such as CO_2) with hydrogen to produce methane, thus with a slope of -0.25 (see text and Fig. 7).

422

423 4.3 A comparison of the gas composition of New Caledonia alkaline springs with424 similar sites worldwide

425

426 Gas venting is commonly observed in hyperalkaline springs, but not in all of them. Whereas it is revealed by bubbles in spring waters, it can be detected as "dry" seeps in serpentinized peridotites 427 [Zgonnik et al., 2019]. The compositions of free gases venting in Oman, Italy (Liguria and Elba 428 429 Island), Turkey and the Philippines have been corrected for O2 contamination and recalculated for the 430 N₂-H₂-CH₄ system, as done for New Caledonia (Fig. 7; see plots for each location in Fig. S3). It has 431 already been shown that gases can be classified into a few types, such as H_2 -rich, CH_4 -rich or N_2 -rich 432 [Boulart et al., 2013; D'Alessandro et al., 2018b; Deville and Prinzhofer, 2016; Vacquand et al., 433 2018].

434 The present data point to three categories (Fig. 7):

435 1) Gases with little or no hydrogen (CH₄-N₂ gases). The methane content of these CH₄-N₂ gases varies within a large range, between 0 and 100%. They are found in the Voltri ophiolite in Liguria 436 (Northern Italy; [Boschetti et al., 2013; Boulart et al., 2013; Etiope and Whiticar, 2019], in the 437 Gulderen and Gokdere springs of the Kilzidag ophiolite in Turkey [D'Alessandro et al., 2018b], in the 438 submarine seeps offshore the Elba Island (Italy; [Sciarra et al., 2019], in the sample collected at the 439 submarine Aiguille de Prony in the Prony Bay in New Caledonia [Vacquand et al., 2018] and in the 440 Chimarea seeps (Turkey) where venting gases are almost pure methane with less than 10% of either 441 442 N₂ or H₂ [Etiope et al., 2011; Vacquand et al., 2018]. [Vacquand et al., 2018] report the very peculiar composition of a gas sample of the Aiguille de Prony (New Caledonia) collected in the vial containing 443

a degassing water, and as such not a free gas sample *per se*. It contains no hydrogen and only a very
small quantity (5%) of methane and as such is almost pure nitrogen. Several samples collected in
Oman by [*Neal and Stanger*, 1983] are merely air, but the gas sample pure nitrogen at the Huwayl
Qufays spring is pure nitrogen.

2) Gases with little or no nitrogen (CH₄-H₂ gases). They vent in the dry seeps of Los Fuegos Eternos and Nagsasa Bay in the Philippines [*Abrajano et al.*, 1988; *Vacquand et al.*, 2018], of the Kurtbagi area in the Kilzidag ophiolite (Turkey; [*D'Alessandro et al.*, 2018b; *Yuce et al.*, 2014] and also of the Chimarea seeps (Turkey; [*Etiope et al.*, 2011; *Vacquand et al.*, 2018]).

3) The compositions of gases collected in New Caledonia (this work; [Vacquand et al., 2018]), 452 453 in Oman [Boulart et al., 2013; Neal and Stanger, 1983; Sano et al., 1993; Vacquand et al., 2018], at 454 the Mangatarem site in the Zambales ophiolite (Philippines; [Vacquand et al., 2018]), and at the 455 Tahtakopru springs of the Kilzidag ophiolite (Turkey; [D'Alessandro et al., 2018b] can be grouped 456 into the N₂-H₂-CH₄ type. They build a trend from almost pure hydrogen, as found in Oman, to N₂-CH₄ 457 gases containing no hydrogen and no more than 20% of methane, as found in Liguria at Acquasanta 458 [Etiope and Whiticar, 2019], Branaga [Boschetti et al., 2013] and GOR35 (one of the springs of Lago Lavagnina; [Boulart et al., 2013]) and at Gulderen spring of the Kilzidag ophiolite [D'Alessandro et 459 al., 2018b]. Free gases for which hydrogen is dominant (>60%) have been collected in Oman by 460 461 [Sano et al., 1993] in 1982-1983 and by [Neal and Stanger, 1983] at dates that are not mentioned, but 462 likely at the same period as Sano et al., and by [Vacquand et al., 2018] between 2008 and 2012. While these data are in fair agreement at high hydrogen content, in spite of the scatter, they differ at 463 hydrogen concentrations below 60%. [Boulart et al., 2013] report Omani gas compositions with 464 465 hydrogen concentrations lower that those reported by [Vacquand et al., 2018], although the samples 466 have been collected mostly at the same locations. The three points with the lowest H₂ contents 467 (20<H2<40%) are nevertheless in agreement. Also, the data for New Caledonia reported by these two research groups [Boulart et al., 2013; Vacquand et al., 2018] are in full agreement, so that it is 468 469 unlikely that these differences for Oman are due to different sampling or analytical methods.

470



473 Figure 7 – A triangular plot of the O_2 -corrected free gas compositions in serpentinizing 474 environments. The black plain line is the line with a slope of -0.25 corresponding to the Sabatier 475 reaction involving CO_2 . The short dash line has a slope of -0.33 corresponding to a Fisher-Tropsch-476 type reaction involving carbon monoxide (see text). The long dash line represents the reaction of 477 elemental carbon with hydrogen (slope -0.5). 478

479 5 Discussion

481 The present analysis of the free gas composition is based on the observation that the endmember high-pH waters discharging at the springs are totally devoid of oxygen, meaning that oxygen is 482 483 consumed in the subsurface and that gases venting at the surface should also be oxygen-free. Small 484 amounts of oxygen would not drastically change the correction (eq. 2). Oxygen is very rapidly consumed when oxic waters flow into geological formations (see the examples of oxygen dynamics in 485 karstic environments; [Monnin et al., 2019; Young et al., 2018]). This lends support to the conclusion 486 487 that the hydrothermal systems responsible for the discharge of high-pH waters at the surface are 488 anoxic.

489 The composition of gas samples collected the same day at a given spring in New Caledonia is 490 variable (Fig. 6). Such a very rapid (hourly) variation of the hydrogen concentration of gases emitted 491 at the surface has been documented in the so-called "fairy circles" in Brazil [Myagkiy et al., 2020]. 492 The hydrogen concentration is roughly increasing (and the methane roughly decreasing) with time, for 493 the period of the survey (3 years). The four data sets for the Oman gas samples have been sampled at 494 dates that span from the early eighties ([Neal and Stanger, 1983; Sano et al., 1993] to the beginning of the years 2010 [Boulart et al., 2013; Vacquand et al., 2018]. All these data point to a time variation 495 of the free gas compositions, that for now cannot be related to seasonal or climatic change but 496 497 certainly reveal the dynamics of the hydrothermal system. Only long-term monitoring, with measurements at high frequency can provide further insight into this variation. 498

499 Serpentinization is the aqueous alteration of metallic minerals such as olivine to produce500 serpentine. It can be represented by:

$$3(Mg, Fe^{II})_{2}SiO_{4}(s) + 4H_{2}O(l) \rightleftharpoons Mg_{3}Si_{2}O_{5}(OH)_{4}(s) + Fe_{3}O_{4}(s) + H_{2}(aq)$$
(3)

501 This reaction produces hydrogen, which at first is dissolved (which is indicated by $H_2(aq)$ in Eq. 3). 502 When the hydrogen concentration reaches the saturation value, a gas phase is produced (degassing, 503 bubbling):

$$H_2(aq) \rightleftharpoons H_2(g) \tag{4}$$

Hydrogen can react with carbon-containing substances to produce methane. There are several ways to write this hydrogen consumption. All of them represent a well-defined mechanism. They also place strong constrains on the nature (for example carbon oxidation state) and the physical state (aqueous or gaseous) of the reactants. For example, hydrogen consumption to give methane through hydrogenotrophic methanogenesis (thus driven by methanogens), and therefore necessarily in the aqueous phase, can be represented by:

$$CO_2(aq) + 4H_2(aq) \rightleftharpoons CH_4(aq) + 2H_2O(l)$$
(5)

In eq. 5, carbon is totally oxidized, for example as in carbonic acid. Again the methane concentrationin the aqueous phase can reach the saturation value and methane will degas:

$$CH_4(aq) \rightleftharpoons CH_4(g)$$
 (6)

A methane production in the gas phase could be written in a similar way, which corresponds to theSabatier reaction:

$$CO_2(g) + 4H_2(g) \rightleftharpoons CH_4(g) + 2H_2O(l) \tag{7}$$

514 Strictly speaking, the Sabatier reaction is a catalyzed process at elevated temperatures (400-500°C) 515 converting carbon dioxide into methane (methanation). Both Eqs. 6 and 7 have the same 516 stoichiometry: four moles of hydrogen are consumed to produce one mole of methane. In a plot of the 517 methane versus the hydrogen content this would be represented by a line with a slope of -0.25.

518 The Fisher-Tropsch reaction is the conversion of carbon monoxide into alkanes:

$$nCO(g) + 2(2n+1)H_2(g) \rightleftharpoons C_nH_{2(n+1)}(g) + nH_2O(l)$$
 (7)

One of the criteria distinguishing the Sabatier reaction from the Fisher-Tropsch process is that the first
one leads to an almost pure methane (methanation) while the second one leads to a more complex
mixture of higher alkanes. Methane production following a Fisher-Tropsch scheme can be written as:

$$CO(g) + 3H_2(g) \rightleftharpoons CH_4(g) + H_2O(l) \tag{9}$$

In a plot of the methane versus the hydrogen content of the gas, eq. 9 would be represented by a linewith a slope of -0.33.

524 Elemental carbon has been considered as a possible source of carbon in methane production in
525 serpentinizing environments (e.g. [*Vacquand et al.*, 2018]):

$$C(s) + 2H_2(g) \rightleftharpoons CH_4(g) \tag{10}$$

Reaction 10 consumes two moles of hydrogen to produce one mole of methane, thus leading to a linewith a slope -0.5 in a methane versus hydrogen plot.

528 Serpentinization can be described as a process starting with the production of hydrogen (eq. 3) followed by hydrogen consumption, either in the gas phase (eq. 7) or in an aqueous solution (eq. 5) to 529 530 produce methane. The data reported in Fig 7 are more consistent with the stoichiometry of the 531 reactions involving fully oxidized carbon (eq. 5 and 7) than with those involving less oxidized carbon 532 (eq. 8) or reduced carbon (eq. 9). As such, in a strict sense, it should be termed Sabatier-type reaction instead of Fischer-Tropsch type (FFT). This distinction is more significant than what it seems as it 533 directly relates to the ongoing debate on the nature of the carbon source in low temperature 534 serpentinization. Increasingly complex reaction schemes and carbon substrates are now contemplated 535 536 and discussed (e.g. [Menez, 2020; Reeves and Fiebig, 2020]).

Yet this correlation between the hydrogen and methane contents of the H₂-CH₄-N₂ gases does 537 538 not distinguish a process taking place in the gas phase from a reaction in an aqueous solution followed by degassing. It also implies that nitrogen is produced simultaneously with hydrogen consumption and 539 540 methane formation. If this was not the case, for example if nitrogen was coming from deeper sources 541 such as sediments, crustal or mantle rocks below the ophiolites [Vacquand et al., 2018], it would be 542 found in all samples so that H₂-rich gases would be diluted by this deeply-sourced nitrogen and their 543 composition would move away from the pure H_2 apex and from the CH_4 - H_2 gas type (Fig. 7). It can be proposed that the chemical changes brought by serpentinization provide the necessary conditions for 544 545 the production of gaseous nitrogen from dissolved nitrate, i.e. denitrification. Such a process which is 546 mediated by bacteria at surface conditions, has not been studied so far in hyperalkaline environments.

547 The data for the Kaoris spring (and to a lesser extend for the Bain des Japonais) (this work)
548 show a CH₄ content lower than the stoichiometry of the Sabatier reaction (Fig. 7; Fig S3). This can be

549 attributed to a consumption of methane, but it is difficult to consider such a consumption in the gas 550 phase (apart from burning). Methanotrophs and methanogens have already been identified in several 551 alkaline springs such as Prony Bay (in both Kaoris and Bain des Japonais springs) [Frouin et al., 552 2018; Mei et al., 2016; Postec et al., 2015; Quemeneur et al., 2014], the Voltri ophiolite (Liguria, Italy) [Brazelton et al., 2017; Quéméneur et al., 2015], the Santa Elena ophiolite (Costa Rica) 553 554 [Crespo-Medina et al., 2017], the Oman ophiolite ([Kraus et al., 2020; Rempfert et al., 2017]; Kraus et 555 al., 2020) and the Cedars springs (California) [Suzuki et al., 2013]. Methane consumption or production by microorganisms take place in the aqueous phase. Again transfer of gases from the gas 556 557 phase to the aqueous phase (and conversely) plays a role in the methane and hydrogen budget of the 558 system. It would be erroneous to conclude that a Sabatier-type reaction in the gas phase necessarily implies an overall abiotic formation of methane. 559

560 The origin of methane (abiotic, biogenic, thermogenic, magmatic) can be inferred from a variety of criteria, each of them with its advantages and pitfalls (e.g. [Etiope and Whiticar, 2019; Reeves and 561 562 Fiebig, 2020]. Gases venting in fractures (dry) seeps and in springs were suggested to belong to the abiotic methane domain in a δ^2 H versus δ^{13} C diagram [Vacquand et al., 2018]. Data for Oman, 563 Philippines, Liguria, Turkey and Elba Island indeed plot within the of abiotic field empirically defined 564 from isotopic data for methane venting in serpentinizing environments [Milkov and Etiope, 2018] (Fig. 565 8). This field has likely been delimited in part using the same data as reported in Fig. 8. Unfortunately 566 567 the study of Milkov and Etiope does not provide a simple way of tracking down the data used to 568 delimit the abiotic methane domain. This field can be subdivided into the dry seeps and the springs 569 [Vacquand et al., 2018]. Note that the Gokdere spring of the Kildizag ophiolite (Turkey) lies in the "dry seeps" domain. 570



Figure 8 – The δ^2 The hydrogen-methane relationship can also be seen as a picture of different stages of the evolution of the hyperalkaline hydrothermal systems in different regions, as a result of the local hydrological conditions, differences in lithology thus leading to different residence times of the waters and the gases in the subsurface. It can be speculated that the H₂-rich gases in Oman would then be the first step, after which methane starts forming in increasingly large amounts up to the point where all the hydrogen is consumed to reach the end point of the CH₄-N₂ gases.

It is also very likely that the flow path and the rate of flow of gases and waters are different. Etiope and Whiticar (2019) have shown that the carbon in the methane is older than 50 000 years as it is ¹⁴C free whereas the age of the waters in hyperalkaline springs determined from ¹⁴C in aqueous carbonate is only a few thousand years. As such this infers in an apparent decoupling of the sources of the waters and gases sampled at the surface. Serpentinization is an aqueous alteration process, so that hydrogen is first produced in an aqueous (liquid) environment. A gas phase will form only when the 584 hydrogen concentration exceeds saturation. The formation of methane may take place in this gas 585 phase, as proposed by Etiope and Whiticar (2019). Methane can then dissolve into the water and be 586 used as a substrate by methanotrophs.

587 Figure 9 presents a diagram that summarizes the observations and conclusions of this work. It 588 highlights the relationship between the gas and aqueous phases, but cannot explain the reason why gases in dry seeps do not contain nitrogen. The $\delta^2 H$ and $\delta^{13}C$ isotope data (Fig. 8) point to isotopic 589 590 δ^2 H-CH₄(g) signatures different for dry seeps than those for springs. This may indicate different mechanisms at play in the generation of methane. Serpentinization is the first step of the water-rock 591 592 interactions during which hydrogen is generated by the reduction of water and the oxidation of metals. 593 Because it is an aqueous process, hydrogen is forming within the aqueous phase, and a gas phase will form only when its concentration exceeds saturation. Methane can then be generated within this gas 594 595 phase by a Sabatier reaction. Methane will dissolve into the aqueous solution where it can support the development of methanotrophs. The methane budget in the gas will be due to its production by the 596 Sabatier process and its consumption by its dissolution in the water. In the aqueous phase, the methane 597 598 budget will result from its uptake from the gas phase, its consumption by methanotrophs, but also its 599 formation by potential methanogens.

600 Clumped isotope data for CH_4 and an analysis of the dissolved gas data will be reported in 601 separate publications.

602



Figure 9 – A diagram summarizing the hydrogen, methane, and nitrogen budgets in the serpentinizing environment. The first step is hydrogen production by the serpentinization reaction. The hydrogen concentration increases in the water up to the point where saturation is exceeded, and a gas phase forms. In Step 2, methane is produced by a secondary process with the stoichiometry of the Sabatier reaction. Nitrogen is also produced during this step (see text). The gases are reactive in both the gas and the aqueous phases and their concentrations are linked by a solubility relationship. In the last step (3) gases and water flow toward the surface to discharge at dry seeps and springs.

- 611
- 612 6 Conclusions
- 613

The dissolved oxygen concentrations measured *in situ* in the New Caledonia on-land springs along with literature data on alkaline springs worldwide show that high-pH waters discharging at the springs are anoxic. Therefore gases venting at the spring should also not contain any oxygen. It is then necessary to correct gas compositions for atmospheric oxygen for atmospheric oxygen contamination.

The compositions of gases collected at the Kaoris and Bain des Japonais springs in New
 Caledonia over a four-year period (2011–2014) show daily variations and some trends during
 the duration of the survey. Free gases collected in Oman by four different research groups
 over a period of about 20 years also display varying compositions. It can be cautiously
 concluded that there is a time variation of composition of free gases venting in these
 serpentinizing environments, at the hour scale as well as at the seasonal or yearly scale.

The hydrogen and methane contents of free gases collected in New Caledonia along with
 those of Oman, the Mangatarem (Philippines) spring and the Tahtakopru (Turkey) springs
 define a trend of H₂-CH₄-N₂ gases going from almost pure hydrogen to N₂-CH₄ gases (with
 CH₄ around 25%). The slope of this trend is consistent with the stoichiometry of methane
 production from an oxidized carbon source such as carbon dioxide or carbonic acid. It also
 implies that nitrogen is produced simultaneously with methane.

• The stoichiometry of the hydrogen-methane reaction is consistent with a Sabatier reaction in the gas phase or a methane production by hydrogenotrophic methanogens in the aqueous 633 phase. The two mechanisms can take place at the same time and can be coupled through the 634 transfer of gases from the water to the gas phase and conversely. This last point can be 635 constrained by the study of the concentration of dissolved gases to investigate the conditions 636 at which the water degasses.

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651 (https://hal.archives-ouvertes.fr/).

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