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7 **Manuscript Title:** Estimating Microbial Hydrogen Consumption in Hydrogen Storage in  
8 Porous Media as a Basis for Site Selection

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29 Estimating Microbial Hydrogen Consumption in  
30 Hydrogen Storage in Porous Media as a Basis for  
31 Site Selection

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## 45 **ABSTRACT**

46 Subsurface storage of hydrogen, e.g. in depleted gas or oil fields (DOGF), is suggested  
47 as a means to overcome imbalances between supply and demand in the renewable energy  
48 sector. However, hydrogen is an electron donor for subsurface microbial processes,  
49 which may have important implications for hydrogen recovery, gas injectivity and  
50 corrosion. Here, we review the controls on the three major hydrogen consuming  
51 processes in the subsurface, methanogenesis, homoacetogenesis, and sulfate reduction,  
52 as a basis to develop a hydrogen storage site selection tool. Testing our tool on 42 DOGF  
53 showed that seven of the fields may be considered sterile with respect to hydrogen-  
54 consuming microbiota due to either temperatures  $>122$  °C or salinities  $>5$  M NaCl. Only  
55 three fields can sustain all of the hydrogen consuming processes, due to either  
56 temperature, salinity or pressure constraints in the remaining fields. We calculated a  
57 potential microbial growth in the order of  $1-17 \times 10^7$  cells  $\text{ml}^{-1}$  for these fields. The  
58 associated hydrogen consumption is negligible to small ( $<0.01-3.2$  % of the stored  
59 hydrogen). Our results can help inform decisions about where hydrogen will be stored in  
60 the future.

61 *Keywords:* Hydrogen, underground storage, microbial hydrogen consumption,  
62 homoacetogens, methanogens, sulfate reducers

63

## 64 **Highlights**

- 65 • Review of the most important hydrogen-utilizing microorganisms in the  
66 underground.
- 67 • Elucidation of the growth criteria for 480 strains of the mayor hydrogen-  
68 utilizers.

- 69 • Development of a site selection tool for sterile hydrogen storage.
- 70 • Evaluation of the site selection tool on 42 depleted oil and gas fields
- 71 (DOGF).
- 72 • Calculation of the microbial growth and hydrogen consumption in DOGF.

73

## 74 **Abbreviations and units**

75 SSR Sulfur species reduction

76 SSRM Sulfur species reducing microorganisms

77 DOGF Depleted oil and gas fields

78 EPS Extracellular polymeric substances

79 M Molarity (mol L<sup>-1</sup>)

80 MPa Megapascal

81

## 82 **1. Introduction**

83 Zero-carbon energy generation from renewable sources can help mitigate carbon emissions and  
84 abate climate change [1-3]. One of the most significant challenges for renewable energy is the  
85 imbalance between supply and demand [3, 4]. The generation of hydrogen (H<sub>2</sub>) via electrolysis  
86 of water during periods of renewable energy oversupply and subsequent H<sub>2</sub> storage is one way  
87 of overcoming this imbalance, as H<sub>2</sub> can be recovered and used for electricity generation during  
88 periods of renewable energy shortage [1, 5]. Subsurface storage of H<sub>2</sub> in salt caverns, depleted  
89 gas or oil fields or saline aquifers is being considered as an alternative to expensive purpose-  
90 built storage containers [6]. However, the artificial elevation of the H<sub>2</sub> concentration in the  
91 subsurface may stimulate the growth of H<sub>2</sub>-oxidizing (hydrogenotrophic) bacteria and archaea,  
92 here collectively referred to as microorganisms, with possible adverse implications for gas

93 injectivity and withdrawal via permeability reduction, H<sub>2</sub> volume loss and corrosion of metal  
94 infrastructure [4, 7]. Understanding the controls on microbial H<sub>2</sub> metabolism is therefore highly  
95 important.

96 Much of the subsurface is characterized by combinations of elevated temperature [7], high salt  
97 concentrations and high pressure [3], reduced void space [8], limited nutrient availability [9]  
98 and typically highly reducing conditions [9-11]. The evidence for microbial life at depth is  
99 plentiful (e.g. [12-16]). Most microorganisms in nature grow in biofilms attached to surfaces  
100 (communities of aggregated microbial cells embedded in a secreted matrix of extracellular  
101 polymeric substances (EPS)) [17, 18]. Even small amounts of biofilm can reduce pore throat  
102 sizes and increase the flow-path tortuosity, resulting in dramatic decreases in permeability [19].  
103 It has been postulated that biofilms may not form in the nutrient-limited underground when the  
104 groundwater flow is low [19]. However, subsurface biofilms are commonly encountered during  
105 geoenergy activities such as fracturing, hydrocarbon recovery or in geothermal plants [20-25].  
106 Biofilm formation may actually be enhanced under the harsh subsurface conditions as the EPS  
107 layer acts as a protective clothing which ensures the normal reproduction and metabolism of  
108 microorganisms [18].

109 Hydrogen plays a central role in the energy metabolism of subsurface life [9]. Yet, a  
110 quantitative assessment of the consumption of H<sub>2</sub> by deep microbial communities in the context  
111 of the global H<sub>2</sub> cycle is lacking [26]. In underground gas storage sites and oil reservoirs the  
112 most abundant H<sub>2</sub>-oxidizers are hydrogenotrophic sulfate reducers, that couple H<sub>2</sub>-oxidation to  
113 sulfate reduction to produce hydrogen sulfide (H<sub>2</sub>S); hydrogenotrophic methanogens that  
114 reduce carbon dioxide (CO<sub>2</sub>) to methane (CH<sub>4</sub>) by oxidizing H<sub>2</sub>; and homoacetogens that  
115 couple H<sub>2</sub> oxidation to carbon dioxide (CO<sub>2</sub>) reduction producing acetate [7, 24, 27]. These

116 three groups of microorganisms are, amongst others, implicated in causing subsurface  
117 corrosion [7, 27, 28].

118 A recent review addressed the many possible abiotic and biotic H<sub>2</sub>-producing and H<sub>2</sub>-  
119 consuming processes in the subsurface [7]. However, it lacked a quantitative assessment of the  
120 processes of microbial growth and H<sub>2</sub> consumption relevant for H<sub>2</sub> storage. Strobel et al. [29]  
121 summarized the concept and potential of underground methanation using experimental data  
122 from the Sun Storage project. These authors highlighted controls on the growth of methanogens  
123 and changes in gas composition due to methanogenesis, but did not quantify microbial growth.  
124 Many studies report changes in gas composition, biofilm growth and clogging near injection  
125 wells but hardly any studies report quantitative figures on microbial growth or on permeability  
126 changes [30].

127 To date it remains unclear how subsurface microorganisms might react to elevated H<sub>2</sub>  
128 concentrations [7] and hence whether microbial growth is a concern for H<sub>2</sub> storage. Even in  
129 natural, non-engineered subsurface environments, there is little information on microbial H<sub>2</sub>  
130 turnover rates [31] and the behavior and population kinetics of microorganisms are not fully  
131 understood [29]. The majority of the available data on microbial H<sub>2</sub> turnover rates come from  
132 batch cultures at optimal growth conditions where the kinetics [31], the pace of life [32, 33],  
133 the physiological states and the prominent organisms may differ widely from the subsurface  
134 environment [7, 32]. A further complication arises from the fact that many microorganisms in  
135 the deep subsurface are not culturable with modern enrichment techniques [12, 34].

136 In this work, we review the state-of-the-art understanding of the controls of temperature,  
137 salinity, pH, pressure and nutrients and water on microbial growth on H<sub>2</sub> in the subsurface, with  
138 emphasis on the three major H<sub>2</sub>-consuming processes methanogenesis, sulfate reduction and  
139 homoacetogenesis, to determine what reservoir conditions will be unfavorable to microbial

140 activity and as such more suitable sites for long term gas storage operations of 30 years or  
141 longer, such as the UK Rough gas storage site.

142 Physicochemical data from 42 depleted or close to depleted oil and gas fields (DOGF) of the  
143 British and Norwegian North Sea and the Irish Sea as well as five H<sub>2</sub> storage test sites provide  
144 the base for an evaluation of the number of sites where microbial growth of methanogens,  
145 sulfate reducers and homoacetogens can be expected. Using average nutrient contents of the  
146 microbial cells and site-specific dissolved ion concentrations, we calculate significant growth  
147 and a small H<sub>2</sub> consumption for growth-permitting DOGF.

## 148 **2. State of the art understanding**

### 149 **2.1 Likely microbial hydrogen oxidation in hydrogen storage systems**

150 Hydrogen oxidizing processes may be ranked according to the magnitude of their H<sub>2</sub> threshold  
151 and their standard free energy change ( $\Delta G^{0'}$ ), two useful metrics to compare the likelihood of  
152 reactions to take place and the order at which they proceed (Table 1). The H<sub>2</sub> threshold defines  
153 the concentration of H<sub>2</sub> below which it is no longer consumed. Given all other factors are at  
154 optimum, the microbial population with the lowest H<sub>2</sub> threshold value is expected to be the  
155 most successful population in competing for H<sub>2</sub> [35].

156 The  $\Delta G^{0'}$  marks the thermodynamic favorability of a reaction at ambient pressure and  
157 temperature, pH 7 and 1 M of all reactants. In oligotrophic (nutrient poor) high pressure and  
158 temperature environments, the order of the  $\Delta G^{0'}$  may be used to determine which reaction is  
159 more energetically favorable. As can be seen from Table 1, more negative  $\Delta G^{0'}$  values (more  
160 available free energy) are generally accompanied by lower H<sub>2</sub> thresholds. Not included in Table  
161 1 are the kinetics which describe the rate of the electron transfer in the redox reaction.

162 **Table 1.** Biotic H<sub>2</sub>-consuming processes ranked according to their free energy yield ( $\Delta G^0$ ) and measured H<sub>2</sub> threshold. Not included are

163 Vanadium, Cobalt, Technetium, Uranium and Selenium reduction, due their limited relevance for H<sub>2</sub> storage. NA= not available.

H <sub>2</sub> - oxidizing process	Reaction (number)	H <sub>2</sub> threshold (nM)	$\Delta G^0$ (KJ mol H <sub>2</sub> <sup>-1</sup> )	Typical ambient [H <sub>2</sub> ] (nmol L <sup>-1</sup> )	Relevance for H <sub>2</sub> storage
<b>Chromate reduction</b>	$\frac{1}{2}H_2 + \frac{1}{3}CrO_4^{2-} + \frac{5}{3}H^+ \rightarrow \frac{1}{3}Cr^{3+} + \frac{4}{3}H_2O$ (1)	<0.1 <sup>[36]</sup>	NA	NA	low
<b>Aerobic hydrogen oxidation (Knallgas)</b>	$H_2 + \frac{1}{2}O_2 \rightarrow H_2O$ (2)	0.051 <sup>[7]</sup>	-237 <sup>[7, 36]</sup>	NA	low
<b>Denitrification</b>	$H_2 + \frac{2}{5}H^+ + \frac{2}{5}NO_3^- \rightarrow \frac{1}{5}N_2 + \frac{6}{5}H_2O$ (3)	<0.05-0.5 <sup>[7]</sup>	-240.1 <sup>[7, 36]</sup> -224 <sup>[4, 37]</sup>	<0.05 <sup>[4, 35, 36]</sup>	low
<b>Halorespiration</b>	$H_2 + \text{halogenated compounds} \rightarrow \text{dehalogenated compounds} + HCl$ (4)	0.05-0.27 <sup>[36]</sup> <0.3 <sup>[38]</sup>	-230 to -187 <sup>[7]</sup>	NA	low
<b>Iron (III) reduction</b>	$H_2 + \text{ferric(oxy)hydroxides} \rightarrow \text{ferrous iron} + H_2O$ (5)	0.27-2 <sup>[7]</sup> <0.11-0.8 <sup>[36, 38]</sup>	-228.3 <sup>[7, 38]</sup> -182.5 <sup>[36]</sup> -114 <sup>[4]</sup>	0.2 <sup>[4, 35]</sup> 0.2-1 <sup>[36]</sup>	intermediate
<b>Manganese (IV) reduction</b>	$2H_2 + MnO_2 \rightarrow Mn(OH)_2 + 2H_2O$ (6)	<0.05 <sup>[35]</sup>	-163 <sup>[4, 35]</sup>	<0.05 <sup>[4, 35]</sup>	low
<b>Arsenate reduction</b>	$H_2 + HAsO_4^{2-} + 2H^+ \rightarrow H_3AsO_3 + H_2O$ (7)	0.03-0.09 <sup>[36]</sup>	-162.4 <sup>[36]</sup>	0.4-0.7 <sup>[36]</sup>	low
<b>Ammonification</b>	$4H_2 + 2H^+ + NO_3^- \rightarrow NH_4^+ + 3H_2O$ (8)	0.015- 0.025 <sup>[38, 39]</sup>	-150 <sup>[4, 38]</sup>	<0.05 <sup>[4, 35]</sup>	low
<b>Fumarate reduction</b>	$H_2 + \text{fumarate} \rightarrow \text{succinate}$ (9)	0.015 <sup>[38, 39]</sup>	-86.2 <sup>[38]</sup>	NA	low
<b>Hydrogenotrophic sulfate reduction</b>	$4H_2 + SO_4^{2-} + H^+ \rightarrow HS^- + 4H_2O$ (10)	1-15 <sup>[38, 39]</sup>	-38 <sup>[7, 38]</sup> -48 <sup>[36]</sup> -57 <sup>[4]</sup>	1-2 <sup>[4, 35]</sup>	high
<b>Hydrogenotrophic methanogenesis</b>	$4H_2 + CO_2 \rightarrow CH_4 + 2H_2O$ (11)	0.4-95 <sup>[38-40]</sup>	-34 <sup>[4, 38]</sup> -43.9 <sup>[36]</sup>	5-10 <sup>[4, 35]</sup> 7-13 <sup>[36]</sup>	high
<b>Sulfur reduction</b>	$H_2 + S \rightarrow HS^- + H^+$ (12)	2500 <sup>[7]</sup>	-33.1 <sup>[7]</sup>	NA	intermediate
<b>Homoacetogenesis</b>	$4H_2 + 2CO_2 \rightarrow CH_3COOH + 2H_2$ (13)	328-3640 <sup>[38, 39]</sup>	-26 <sup>[4, 38]</sup> -36.1 <sup>[36]</sup>	100- <sup>[4]</sup> , 117-150 <sup>[36]</sup>	high

164



165 Abiotically, most of the H<sub>2</sub>-oxidizing reactions are very slow but mediated by microbial  
166 enzymes the processes are catalyzed [37, 41].

167 The three main microbial processes with implications for H<sub>2</sub> storage, hydrogenotrophic sulfate  
168 reduction, hydrogenotrophic methanogenesis (for simplicity from now on just referred to as  
169 sulfate reduction, and methanogenesis unless otherwise specified) and homoacetogenesis,  
170 require the highest threshold [H<sub>2</sub>] and are among the processes with lowest  $\Delta G^0$  (Table 1).  
171 Nevertheless, e.g. sulfate reduction is instantaneous in most geologic settings [42] possibly due  
172 to fast kinetics [37] and/or a relatively high availability of sulfate.

173 Because sulfate reducers may use the same substrates as sulfur reducers (i.e. sulfide and  
174 thiosulfate [43, 44]), they are here collectively referred to as sulfur species reducing  
175 microorganisms (SSRM) performing sulfur species reduction (SSR). Direct respiration of  
176 sulfur is limited by its low solubility ( $1.6 \cdot 10^{-7}$  M) and hence requires cell attachment to the  
177 sulfur particle [45]. However, sulfur readily reacts with sulfide formed during the reduction of  
178 sulfate to form easily metabolizable polysulphides [45, 46].

179 Iron (III) reduction relies on the availability of iron oxides and iron-bearing minerals such as  
180 smectite and chlorite [47, 48], as well as the availability of organic carbon, since dissimilatory  
181 iron reducing bacteria (DIRB) are strict heterotrophs, i.e. synthesize cell carbon from organic  
182 compounds [49]. Iron oxides are abundant in many sediments and aquifers [47] but are  
183 typically not available in the carbon-rich oil fields because they have been reduced over  
184 millions of years and are not replenished [24]. Meanwhile, bacteria capable of reducing iron  
185 are frequently isolated from hydrocarbon-contaminated or oil-associated sites (reviewed in  
186 [50]). However, the mere observation of iron reduction by bacteria, which are given a DIRB  
187 enrichment medium in the laboratory, does not imply that these bacteria will reduce iron in  
188 nature. In addition, cell counts are often low to intermediate ( $10$ – $100$  cells ml<sup>-1</sup>) and may

189 include non-hydrogenotrophs (e.g. [51, 52]). In non-engineered environments rich in Fe oxides  
190 and organic carbon, IRB may have a great advantage over SSRM, methanogens and  
191 homoacetogens, due to a very high affinity for H<sub>2</sub> [47]. We evaluate this process as of  
192 intermediate relevance for H<sub>2</sub> storage in DOGF.

193 Many IRB and a few SSRM can also couple H<sub>2</sub> oxidation to reduction of a variety of other  
194 trace metal oxides, e.g. MnO<sub>4</sub><sup>2-</sup>/MnO<sub>2</sub>, CrO<sub>4</sub><sup>2-</sup>, Co, SeO<sub>4</sub><sup>2-</sup>, UO<sub>2</sub><sup>2</sup>, TcO<sub>4</sub><sup>-</sup>, AsO<sub>3</sub><sup>3-</sup>, and VO<sub>4</sub><sup>-</sup> [43,  
195 53, 54]. After Fe, the most abundant metal in sedimentary environments is Mn (~10 % of Fe  
196 abundance) [47, 53]. Due to the trace content of these compounds in the environment, their  
197 reduction has low relevance for H<sub>2</sub> storage.

198 Oxygen and nitrate are scarce in the subsurface [11, 24, 55, 56] and aerobic hydrogen oxidation,  
199 denitrification and ammonification hence only become significant when contamination of the  
200 aquifer occurs, e.g. by drilling fluid [57-59].

201 Halogenated compounds are common in aquifers, and may arise from contamination or via  
202 natural processes in sediment [60, 61]. However, the concentrations of these compounds are  
203 extremely low: In aquifers of 167-1000 m depth, chloroflourocarbons reach maximum  
204 concentrations of ≤1.1 µg L<sup>-1</sup> [61] and for pristine aquifers 0.003-0.007 µg L<sup>-1</sup> of chlorinated  
205 hydrocarbons were measured [60]. We evaluate the relevance of this process to H<sub>2</sub> storage as  
206 negligible.

207 Literature on the importance of anaerobic fumarate respiration using H<sub>2</sub> is scarce. Fumarate  
208 may be used as an alternative electron acceptor by SSRM [43, 62] and homoacetogens [63-66].  
209 In the non-engineered subsurface, readily metabolizable organic matter, like fumarate, is rare  
210 [67]. Oil fields being rich in organic C compounds may contain more fumarate. Payler et al.  
211 [12] confirmed the presence of fumarate reductase, the key enzyme in fumarate reduction, in  
212 three out of five metagenomes from subsurface brines within sandstone. However, the

213 metagenomes belonged primarily to non-H<sub>2</sub> utilizing bacteria (*Halorubrum*) and fumarate  
214 concentrations were not reported. Acknowledging the lack of data in this field, we evaluate this  
215 process as being of low relevance for H<sub>2</sub> storage.

## 216 **2.2 Factors governing microbial growth**

217 Microbial growth and H<sub>2</sub> consumption rates vary with nutrient availability and environmental  
218 variables (e.g. [17, 68].) Each strain is adapted to an optimum set of nutrients and  
219 environmental conditions where potentially the greatest growth rates occur. Beyond the  
220 optimum conditions, organisms may grow but at reduced rate or they become dormant. In this  
221 section, we discuss the requirements for nutrients and water, and the overall impact of  
222 temperature, salinity, pH and pressure on the growth of the major microbial H<sub>2</sub>-oxidizers in  
223 DOGFs, in the ranges relevant to H<sub>2</sub> storage. The specific activity of microbial strains grown  
224 at optimum conditions varies as well (reviewed in [69]) but the elucidation of differences  
225 between strains is beyond the scope of this review.

### 226 **2.2.1 Nutrients**

227 Apart from water of sufficient thermodynamic activity (see Section 2.2.4), hydrogenotrophs  
228 require H<sub>2</sub> as a source of electrons (energy), an electron acceptor and a carbon source for cell  
229 division, together with a set of macro and trace elements as well as various organic nutrients  
230 [70]. Microorganisms can only access H<sub>2(aq)</sub> and hence the solubility of H<sub>2(g)</sub> is of direct  
231 relevance for all H<sub>2</sub>-consuming reactions. Given a gas phase of ~100 % H<sub>2</sub> in an H<sub>2</sub> storage  
232 system, the equilibrium solubility of H<sub>2</sub> exceeds the highest threshold value of an H<sub>2</sub>-  
233 consuming microorganism of 3.6 μM (Table 1) by ~3 orders of magnitude at ambient pressure  
234 and temperature and under static conditions (Fig. A.1a), with further increase at higher  
235 pressures (Fig. A.1b). While under non-static conditions hydrogenotrophs will consume part

236 of the H<sub>2</sub>, these figures suggest no limitation by the H<sub>2</sub> solubility on microbial growth under  
237 H<sub>2</sub> storage conditions.

238 Elemental requirements include the macro elements C, N, H, P, Ca, Mg, S and Fe (>95 % of  
239 the microbial cell dry weight), and the trace elements Co, Mn, Ni, Mo, Cu, Zn, W as well as  
240 Se for some metabolic groups [71, 72]. For optimum growth, many microorganisms  
241 additionally require different vitamins (e.g. lipoic acid, biotin, riboflavin, folic acid, thiamine,  
242 etc.), yeast extract, coenzyme M, aromatic acids and phospholipids or a combination of these  
243 (e.g. [8, 65, 73-75]).

244 Nutrients may be assimilated from the solution or directly from minerals (e.g., [76-79]), the  
245 latter being of particular importance in oligotrophic environments [77]. Carbon, sulfur,  
246 phosphorous and iron are amongst the key elements released by mineral weathering [77]. The  
247 extent to which subsurface microbial communities depend on mineral weathering is unknown  
248 [77]. For soils, Huang et al. [80] analyzed that >50 % of the 1100 microbial strains were  
249 capable of mineral weathering, as tested by their ability to mineralize biotite.

250 Microbial cell carbon may be assimilated from CO<sub>2</sub> alone (autotrophy) or from organic carbon  
251 compounds (heterotrophy) [81]. Methanogens and homoacetogens can grow autotrophically or  
252 heterotrophically, and several can grow mixotrophically (e.g. [66, 82, 83]). SSRM typically  
253 grow heterotrophically but some grow autotrophically or mixotrophically [84, 85]. Nitrogen  
254 may be assimilated from ammonia and nitrate or by nitrogen-fixation (diazotrophy).  
255 Diazotrophy is common amongst SSRM, methanogens and homoacetogens [86-89], though  
256 homoacetogens often inhabit ammonia-rich environments [88].

257 Little is known about the differences in the nutrient requirements on the level of functional  
258 groups and the variation in nutrient requirement within a functional group. SSRM have a higher  
259 requirement for iron ( $1.8 \cdot 10^{-6}$  M) than is usually observed for microorganisms [90] while

260 methanogens have a higher requirement for sulfur with optimal levels ranging from 0.03 to  
261 0.79 mM (reviewed in [91]).

262 Literature on when nutrients become limiting is also scarce. Sulfate reducing SSRM require a  
263 minimum sulfate and phosphorus concentrations of ~3 mM and  $\sim 3.2\text{-}320 \times 10^{-5}$  mM,  
264 respectively, for growth [92, 93]. Methanogens of the order *Methanosarcinae* require 29.6 mM  
265 Mg for optimum growth and growth ceases at 15.8 mM (reviewed in [91]). When grown under  
266 optimum conditions, the growth rate of autotrophs may be limited by the rate of transfer of H<sub>2</sub>  
267 and CO<sub>2</sub> from gas to liquid, as was shown for the methanogen *Methanobacterium*  
268 *thermoautotrophicum* [94] and for the sulfate reducers within *Desulfotomaculum sp* [95].

269 Carbon is unlikely to be limiting in the hydrocarbon-rich DOGF [56, 96, 97] but this is not a  
270 given in saline aquifers with no history of oil or gas. Sulfate is present in significant  
271 concentrations in most DOGF (Table 2) but H<sub>2</sub> injection can cause sulfate depletion due to  
272 accelerated growth of SSRM [98]. Nitrogen in the form of ammonia may be limiting in DOGF  
273 [51, 56, 90] but nitrate levels may be elevated [51], often due to contamination by drilling fluid  
274 [57-59].

### 275 **2.2.2 Temperature**

276 Temperatures of 22.5–80 °C or 20–100 °C have been suggested for H<sub>2</sub> storage based on a  
277 recommended depth range of 500- 2000 m for H<sub>2</sub> storage in DOGF and saline aquifers [99-  
278 101]. Microorganisms are classified according to their preferred growth temperature:  
279 psychrophiles grow optimally below 20 °C, psychotrophs grow optimally at or above 20 °C  
280 and may tolerate temperatures below 5 °C, mesophiles grow between 20 and 45 °C,  
281 thermophiles grow above 45-50 °C, and hyperthermophiles show optimal growth at  
282 temperatures of 80 °C or above [102, 103].

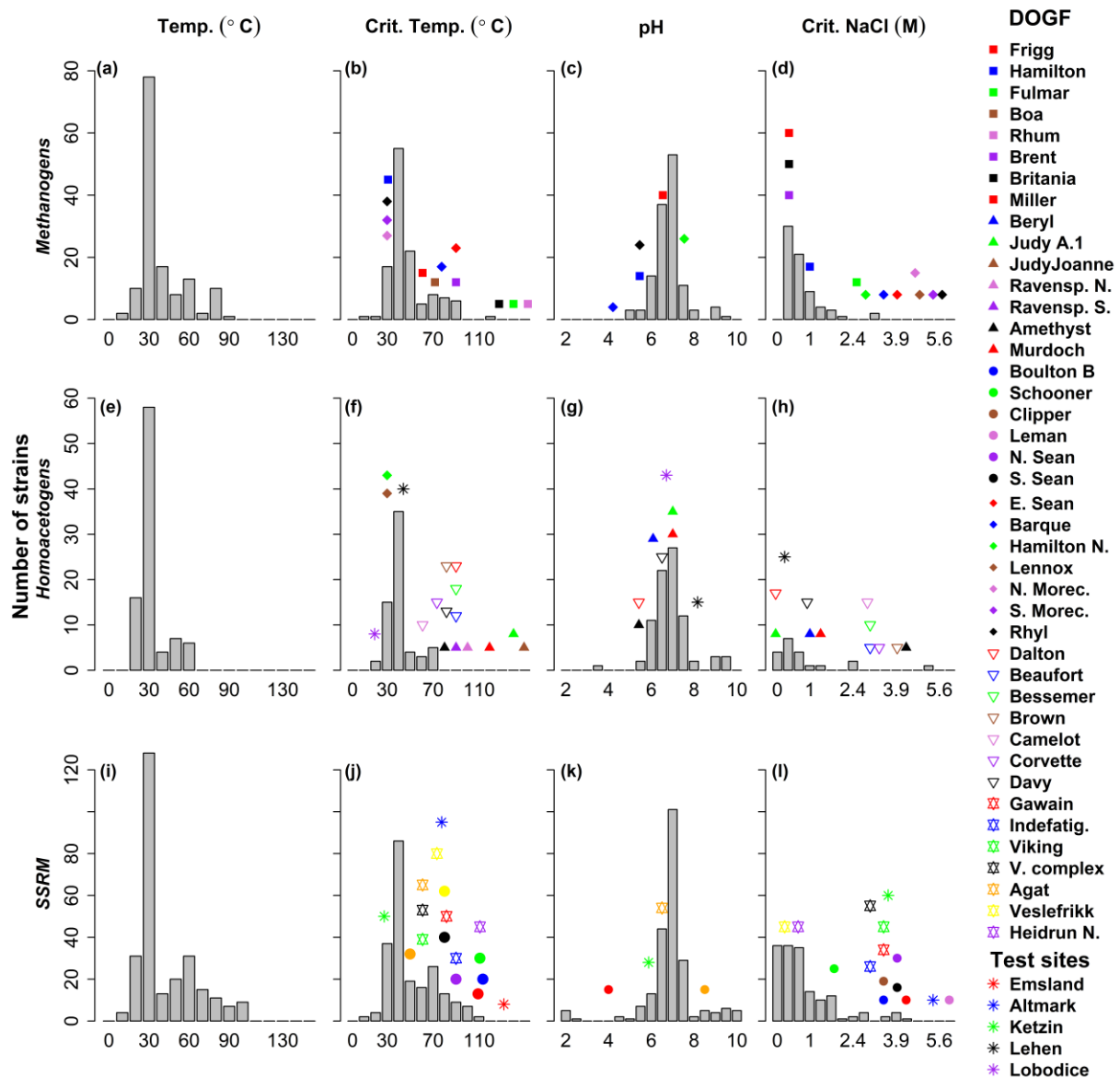
283 High temperatures alter the energetic properties (e.g., vibrational modes) of biomolecules in  
284 their aqueous solvent, change the substrate solubility or viscosity and the ionization of the  
285 aqueous medium [104]. Adverse effects of high temperature include DNA denaturing or  
286 damage, decreased protein stability, hydrolysis of ATP and ADP, amongst others [104, 105].  
287 The metabolic strategies of thermophiles are highly diverse. For a discussion, the reader is  
288 referred to [106].

289 Thermophiles and hyperthermophiles are challenged by increased reaction rates at elevated  
290 temperature which can imply that abiotic reaction rates are so fast that there is no benefit to the  
291 microorganism if it catalyzes the reaction [41]. High-temperature-adapted microorganisms are  
292 therefore thought to produce enzymes with faster reaction rates [107].

293 Most cultivated hydrogenotrophic methanogens are mesophiles but known optimal growth  
294 temperatures for methanogens range from 15 to 98 °C (Fig. 1a). A considerable number of  
295 methanogens favor temperatures above 60 °C (Fig. 1a). The highest temperature that a  
296 methanogen was found to grow under is 122 °C (*Methanopyrus kandleri*) (Fig. 1b) [108].

297 Cultivated SSRM typically have optimum growth temperatures of 20-30 °C or 50-70 °C where  
298 sulfur reducing archaea have higher optimum growth temperatures than sulfur and sulfate  
299 reducing bacteria. The full range for optimum growth of SSRM spans 10-106 °C (Fig. 1i). The  
300 critical temperature for growth of cultivated SSRM is 113 °C (*Pyrolobus fumarii*) [109].

301 Homoacetogens typically have optimum growth temperatures between 20-30 °C (85 % of the  
302 here gathered cultivated strains; Fig. 1e). Thermophilic growth temperatures  $\geq 60$  °C have been  
303 reported for eight strains, only (e.g. *Moorella mulderi*, *Thermoanaerobacter kivui*,  
304 *Acetogenium kivui*) [110-112]. Corresponding upper limits for growth are 70-72 °C (Fig. 1f)  
305 [110-112].



306

307

**Figure. 1.** Distribution of optimum growth temperature, critical growth temperature, optimum pH values and critical salinity for 123-140 methanogens (a-d), 21-91 homoacetogens (e-h) and 151-255 sulfur species reducing microorganisms (SSRM) (i-l). Distributed between the graphs for the different groups of H<sub>2</sub>-oxidizers are the temperatures, pH values and salinities of 42 depleted oil and gas fields (DOGF) and five test sites for H<sub>2</sub> injection. Where ranges of a parameter were given (see Table A.4), the lower end value was plotted.

313

314

### 315 **2.2.3 Salinity**

316 The relevant salt concentration range for H<sub>2</sub> storage is 0-5 M NaCl [100], at which highly  
317 diverse prokaryote communities can be found [113]. Microorganisms are classified according  
318 to their salt tolerance: Non-halophilic microorganisms grow up to 0.2 M NaCl, slight halophile  
319 grow at 0.2–0.5 M NaCl, moderate halophile between 0.5–2.5 M NaCl, and extreme halophile  
320 that grow best in hypersaline media containing 2.5–5.2 M NaCl [113].

321 High salt concentrations exert osmotic stress [114], requiring any microorganism living at high  
322 salt concentrations to maintain its intracellular environment at least isosmotic with the  
323 environment [113]. Commonly, salt tolerance/requirement is enhanced at increased  
324 temperatures [113] but there are many examples of mesophilic halophiles.

325 Most hydrogenotrophic methanogens favor salt concentrations up to 0.77 M NaCl (the  
326 approximate salinity for seawater) but 16 known strains survive under more halophilic  
327 conditions. Two extremely halophilic mesophilic hydrogenotrophic methanogens, will tolerate  
328 salt concentrations of ~3.3- 3.4 M, *Methanocalculus halotolerans FRIT* [115] and  
329 *Methanocalculus natronophilus* [116] (Fig. 1d).

330 The large majority of SSRM grow optimally at low salinities between >0-0.4 M. However,  
331 fourteen SSRM (all mesophiles) have upper salinity limits for growth of  $\geq 1.7$  M NaCl (Fig.  
332 1h). *Desulfovibrio oxyclinae*, *Thiobacillus halophilus*, *Desulfohalobium utahense* and  
333 *Desulfohalobium retbaense*, have the highest upper salinity limits for growth of 4.0 to 4.2 M  
334 NaCl [117-120] (Fig. 1h).

335 The salt tolerance of homoacetogens is poorly investigated. The majority of homoacetogens  
336 have low optimum salinities of >0-0.4 M NaCl. However, a few strains, i.e. *Natroincola*  
337 *histidinovorans*, *Sporohalobacter lortii* and *Halanaerobium praevalens*, grow optimally at  
338 >1.4 M NaCl and will tolerate salinities up to 2.6-5.1 M (Fig. 1h) [121-123].



#### 339 **2.2.4 Brine complexity**

340 Natural brines contain dissolved ions whose interaction is extremely complex and may cause  
341 physicochemical stressors to brine habitability such as low water activity ( $a_w$ ), high ionic  
342 strength, chaotropy (ability to disrupt the network of H<sub>2</sub>-bonds between water molecules) or a  
343 combination of these [12]. Most bacteria grow well at an  $a_w$  around 0.98 (the approx.  $a_w$  for  
344 sea water) but relatively few species can grow at  $a_w$  of 0.96 or lower [124]. Halophilic  
345 microorganisms, including halophilic methanogens are one exception; several can grow at  $a_w$   
346 as low as 0.75 [125] in [124]; [126]. Steinle et al. [127] challenged these limits by detecting  
347 SSR in a nearly MgCl<sub>2</sub> saturated brine with  $a_w$  of ~0.4.

348 There are indications of a more important role of chaotropy over  $a_w$  in limiting microbial life  
349 [126]. Chaotropic agents include MgCl<sub>2</sub>, CaCl<sub>2</sub>, FeCl<sub>3</sub>, KI, LiBr, LiCl while examples of  
350 kosmotropic agents are NaCl, KCl, Na<sub>2</sub>SO<sub>4</sub>, MgSO<sub>4</sub>, K<sub>2</sub>SO<sub>4</sub>, FeSO<sub>4</sub> [128]. As such one may  
351 speculate that most subsurface brines due to their dominance of NaCl and richness in sulfate  
352 are kosmotropic and albeit also stress-inducing, more permissive of microbial growth [12, 126].  
353 Meanwhile, the interactions between chao- or kosmotropic agents,  $a_w$  and other  
354 physicochemical properties of brines may be very complex and hitherto not understood [12].  
355 The further elucidation of this topic is subject to more research and beyond the scope of this  
356 paper.

#### 357 **2.2.5 pH**

358 The brine pH may affect the growth of microorganisms via 1) a direct effect on the growth  
359 metabolism, and 2) an effect on the redox reaction. With respect to the former, most  
360 methanogens, homoacetogens and SSRM are adapted to a pH of 6.5-7.5 (Fig. 1c, g, k). Most  
361 methanogens and SSRM cannot grow outside the pH range 4–9.5 [29, 129, 130] (Fig. 1c, k).  
362 Ten known methanogens can endure a critical pH-value of 10 (e.g. the *Methanosalsum*

363 *natronophilum* and *zhilinae*, and the *Methanocalculus natronophilus* and *alkaliphilus* [131].  
364 At the other end of the spectrum, ten known methanogens can endure acidic conditions of pH  
365 4, e.g. the *Methanoregula boonei*, the *Methanothermococcus okinawensis*, the *Methanosarcina*  
366 *spelaei* and the *Methanocaldococcus bathoardescens* [132-135].

367 Eighteen known SSRM are adapted to highly alkaline environment >pH 10, e.g. the  
368 *Desulfonatronovibrio hydrogenovorans*, the *Desulforispira natronophila* and the  
369 *Desulfovibrio vietnamensis* [136-138]. Thirteen known SSRM grow down to a pH of 4. Nine  
370 known SSRM, all of them sulfur reducers, grow down to a pH of 1, e.g. the *Thiobacillus caldus*,  
371 the *Sulfolobus acidocaldarius*, the *Acidianus infernus* and *brierleyi*, and the *Stygiolobus*  
372 *azoricus* [139-142].

373 Seven known homoacetogenic strains have high critical pH values up to 10.0-10.7, e.g.  
374 *Clostridium ultunense*, *Natronella acetigena*, *Fuchsiella alkaliacetigena* and *ferrireducens*,  
375 *Natronoincola histidinovorans*, *Peptostreptococcus productus B-52* and *Moorella sp HUC22-*  
376 *1* [121, 122, 143-146]. The *Clostridium drakai*, *ljundahlia*, *scatologenes*, *coccoides* and  
377 *termoautrophicum* are the most acidophilic known strains; they can tolerate pH as low as 3.6-  
378 4.5 [147-151].

### 379 **2.2.6 Pressure**

380 Pressure ranges for H<sub>2</sub> storage of 5-20 MPa [99] or 1-50 MPa [100] have been reported. Life  
381 at high pressure requires homeostatic changes [103]. The high pressures encountered in pore  
382 spaces in the crust are generally less inhibitory to microbial cellular activity than the high  
383 temperatures, partly because of the relatively high osmotic pressure of cytoplasm [102], in  
384 particular in thermophiles and hyperthermophiles [41]. DNA synthesis and protein synthesis  
385 are among the most pressure-sensitive cellular processes [103, 152]. Protection against pressure  
386 includes biofilm [18] or spore formation [103].

387 At 30-50 MPa, the growth of various mesophilic, atmospheric-pressure-adapted  
388 microorganisms is inhibited [152] whereas pressure effects are generally favorable for the  
389 growth of hyperthermophiles; above 100 °C, elevated pressures are required to maintain a  
390 liquid environment [105]. Microorganisms that grow optimally at 10 MPa or above are obligate  
391 and facultative piezophiles, where the former do not tolerate ambient pressure and the latter do  
392 [103]. A recent publication listed all identified piezophiles and grouped them according to their  
393 growth temperature optimum [103]. The list of species is rather short (and as we find  
394 incomplete despite being published in 2020), possibly due to the fact that, to date, it has not  
395 been possible to isolate genes associated with piezophily, so the effects of pressure on any  
396 particular organism can only be determined empirically [103]. Empirical efforts however, do  
397 not commonly include pressure tolerance in the description of the environmental growth  
398 criteria of a microorganism. In addition, most mesophiles and thermophiles from habitats with  
399 pressures of <50 MPa will grow in enrichment cultures incubated at atmospheric pressure [34].  
400 The large majority of identified cultivated piezophiles are psychrophiles (27 strains) [103], the  
401 relevance of which is low to our study. Only four mesophilic strains were reported, three of  
402 them hydrogenotrophic sulfate reducers (the *Desulfovibrio profundus*, *piezophilus*, and  
403 *hydrothermalis*), growing optimally at 10-40 MPa [103]. Eight thermophiles were identified,  
404 including one hydrogenotrophic methanogen, *Methanococcus thermolithrophicus*, growing  
405 optimally at 50 MPa. The hyperthermophilic group hosts the hydrogenotrophic *Methanopyrus*  
406 *kandleri* and *Methanocaldococcus jannaschii* growing optimally at 20 to 75 MPa, respectively.  
407 Examples of hydrogenotrophic piezophiles that are not included in [103] are the mesophilic  
408 SSRM *Parococcus pantrophus* and *Pseudodesulfovibrio indicus* which growth optimally at 30  
409 and 10 MPa, respectively [153, 154], and the thermophilic SSRM *Piezobacter thermophiles*  
410 and *Archaeoglobus fulgidus TF2* which grow optimally at 30 and 42 MPa, respectively [75,  
411 155].

412 A temperature dependence of the pressure response was reported for the SSRM *Desulfovibrio*  
413 *indonesiensis* which has similar growth rates at high and ambient pressure 45 °C but reduces  
414 its growth rate at 20 °C and 30 MPa relative to at 0.1 MPa [156]. Elevated pressure may  
415 increase the maximum growth temperature by 2-12 °C relative to lower pressure (0.1-3 MPa)  
416 [105, 108, 157].

### 417 **2.2.7 Inhibitors**

418 Exposure to hydrogen sulfide, H<sub>2</sub>S, and its bisulfide ion, HS<sup>-</sup>, causes damage to microbial  
419 proteins and coenzymes [91, 158]. It remains unclear whether H<sub>2</sub>S or HS<sup>-</sup> is responsible for  
420 the toxicity effect but there is general consensus that H<sub>2</sub>S can penetrate the microbial cell  
421 membrane more easily than HS<sup>-</sup> [158]. Hydrogen sulfide dissociates with a pK<sub>1</sub> of 6.99 at 10  
422 MPa and 25 °C to form >99 % HS<sup>-</sup> at pH 8.5 [159].

423 Growth of SSRM and methanogens is adversely affected at concentrations of H<sub>2</sub>S >3.8-4.0  
424 mM [160-162]. At 5.0-6.3 mM H<sub>2</sub>S growth is completely inhibited for SSRM [160, 162],  
425 without however stopping all metabolic activity [160]. For methanogens and homoacetogens  
426 3.8-7.5 mM H<sub>2</sub>S and total sulfide concentrations of 3.3 mM, respectively, stop the growth [158,  
427 162]. In systems with circumneutral pH and ferric ion concentrations above 1 mM, the  
428 concentrations of H<sub>2</sub>S are predicted to be kept below toxic levels due to its precipitation in  
429 makinawite [46].

430 Carbon dioxide pressure above 1 bar can be toxic for microorganisms as shown for the SSRM  
431 *Desulfotomaculum geothermicum* and the methanogen *Methanothermococcus*  
432 *thermolithotrophicus* [163]. For many anaerobes like methanogens and homoacetogens,  
433 oxygen is toxic too [64, 102].

434 Nitrate inhibits homoacetogenesis [164], and ammonium [165] and sulfate inhibit  
435 methanogenesis (reviewed in [166]), with minimum inhibitory concentrations varying

436 depending on the environment [162, 165]. For instance, sulfate concentrations as low as  $2 \times 10^{-4}$   
437 M were shown to inhibit methanogenesis for 10 hours in lake sediments, possibly by  
438 competition with SSRM for available  $H_2$  and C-substrate [161] (see section 2.3.9). Under  $H_2$   
439 storage conditions however, sulfate is likely not to affect methanogenesis, because sulfate  
440 inhibition was shown to be reversed by addition of  $H_2$  [161]. For a discussion of an inhibitory  
441 effect of  $H_2$ , see section 2.5.

### 442 ***2.2.8 Summary of environmental growth constraints***

443 Acknowledging the lack of data for the pressure sensitivity of many microorganisms [103],  
444 and considering a general abundance of nutrients in DOGF (Table 2), we evaluate temperature  
445 and salinity as the most crucial environmental factors constraining the growth of  
446 homoacetogens, methanogens and SSRM in DOGF. Pressures encountered in the crust are  
447 documented to have less effect than temperature on microbial cellular activity, particularly in  
448 thermophiles and hyperthermophiles [41, 102]. The pH does not pose a similar constraint to  
449 the growth of homoacetogens, methanogens and SSRM because the pH ranges for growth  
450 typically span two to three pH units (not shown) and for most species they comprise the typical  
451 aquifer pH values of 6-7 [167] (Table A.4). Brine complexity and inhibitors were not included  
452 in this analysis due a lack of information on the brine composition of DOGF beyond a limited  
453 set of dissolved ions.

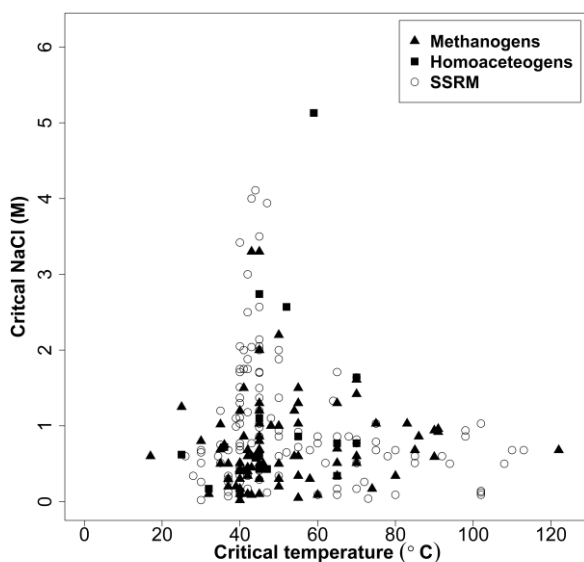
454 Figure 2 shows the critical temperature versus critical salinity for 269 cultivated strains and  
455 reveals that salt tolerances up to 1-1.7 M are widely distributed over the entire temperature  
456 range while salt tolerances  $>1.7$  M are mainly found at a critical temperature tolerances of 40-  
457 50 °C. Hence, from the point of view of minimizing microbial impacts on  $H_2$  storage, sites with  
458 temperatures  $>50$  °C and salinities  $>1.7$  M are preferred.

459 **Table 2.** Reservoir conditions for selected depleted, or soon to be depleted oil and gas fields. Except where otherwise indicated, the data are  
460 from [168]. <sup>®</sup>= reference [167]. The salinity was calculated from the chloride concentration and the concentrations of dissolved N<sub>2</sub> was  
461 estimated from the mol percentage in the gas phase, neglecting any effect of salinity. NA= not analyzed. See Table A.4 for extended data.

462

Field name	Area (Km <sup>2</sup> )	P (MPa)	Temp (°C)	Salinity (M)	pH	HCO <sub>3</sub> <sup>-</sup> (mM)	N <sub>2</sub> (mM)	SO <sub>4</sub> <sup>2-</sup> (mM)	K <sup>+</sup> (mM)	Ca <sup>+2</sup> (mM)	Mg <sup>+2</sup> (mM)	P (mM)	Na <sup>+</sup> (mM)	Cl <sup>-</sup> (mM)	Fe <sup>+2</sup> (mM)	Organic acids (mM)
<b>Frigg</b>	100	19.5	61	0.07-0.53	6.5-7.4	16.3	0.4	NA	26.3-31.2	0.4-2.0	1.9-7.1	NA	75.2-534.8	58.7-490.3	0.04-0.27	NA
<b>Hamilton</b>	15	9.6	30	1.59-4.18	5.8	4.8	2.1	0.6-7.4	8.4-29.7	72.8-720.0	19.5-37.6	0.012-0.028	1354.8-2210.9	1453.3-3700.7	4.03	NA
<b>Barque</b>	36	26.0	79	4.83	4.7	0.3	0.8	3.5	42.2	535.0	156.8	NA	2920.4	4405.4	2.15	NA
<b>Hamilton North</b>	8	10.5	30	2.93	7.9	11.0	2.3	23.1	18.8	13.6	13.6	NA	2640.9	2662.9	NA	NA
<b>Miller</b>	45	49.3	121	1.61	7.2	NA	0.6	0.0	41.6	30.0	NA	NA	1358.7	1471.9	0.02	NA
<b>Beryl</b>	49	36.0	101	1.88	6.1	5.6	0.4	0.0	20.8	90.0	NA	NA	1469.6	1717.9	0.05	1.9
<b>Judy</b>	NA	46.9	137	0.14-0.15	6.8	8.4	0.6	6.4	2.9	4.5	NA	0.002	117.4	131.7	0.11	NA
<b>(Andrew 1)</b>																
<b>Amethyst</b>	97	27.9	88	4.45	5.6	1.0	1.6	3.7	33.2	521.5	148.5	0.452	2673.9	4064.6	2.51	NA
<b>Rhyl</b>	NA	14.9	36	5.80	5.5	13.5	2.8	14.0	62.4	147.0	21.2	0.031	4777.0	5297.9	0.81	>1.2
<b>Dalton</b>	NA	28.8	91	0.26	5	0.9	1.0	1.8	15.6	5.5	5.5	NA	189.1	237.0	0.00	NA
<b>Davy</b>	6	28.2	88	3.87	6.8	6.5	NA	7.0	219.2	15.6	10.7	0.155	818.3	1142.7	0.66	NA
<b>Veslefrikk<sup>®</sup></b>	NA	29.8-35.0	67-114	0.29-0.72	6.5	8.4-17.2	NA	0.1-0.15	NA	NA	NA	NA	298.0-666.0	281.0-745.0	NA	2.2-8.1
<b>Average</b>						7.9	1.1	5.2	44.7	166.8	42.2	0.113	1473.7	1857.4	0.967	3.3

463 Growth of all investigated microbial groups occurs up to 72 °C (Fig. 1). Above 72°C, known  
464 homoacetogens will not grow, and at 80-94°C sulfate reducers cease to grow. Thirty-six  
465 cultivated SSRM and eleven methanogens have optimum growth temperature of  $\geq 80$  °C (Fig.  
466 1a and g) and will still grow, albeit at reduced rate, beyond their optimum temperatures. The  
467 maximum growth temperature for known methanogens and sulfur reducers is 122 °C and 113  
468 °C, respectively. The upper salinity limit that allows growth of all the major groups of  
469 investigated H<sub>2</sub>-oxidizing microorganisms is 3 M NaCl. The upper pH limit is 9.5 and the upper  
470 pressure limit for most mesophiles is 30-50 MPa.



**Figure. 2.** Critical temperature (without salinity stress) versus critical salinity (without temperature stress) for methanogens, homoacetogens and SSRM.

471

### 472 **2.3 Microbial growth regulation by competition and syntrophy**

473 Homoacetogenic bacteria are ubiquitous in anaerobic sediments [65, 169] and often co-exist  
474 with SSRM and methanogens [15, 170], as revealed by a combination of molecular (16S RNA

475 gene sequences) and culturing (e.g. metabolites, radiotracer) techniques. Few habitats have  
476 been identified in which homoacetogens compete with other H<sub>2</sub>-consumers (culturing studies)  
477 [169, 171]. Exceptions include a low-temperature and low-salinity petroleum reservoir where  
478 homoacetogens dominated over methanogens and SSRM (molecular study) [62], a granite  
479 groundwater at 400 m depth where cell numbers of methanogens and homoacetogens were  
480 balanced (molecular study) [83], and subsurface marine sediments where mixotrophic  
481 homoacetogenesis outperformed methanogenesis (culturing study) [82].

482 Kinetic advantages of SSRM and methanogens (i.e. a higher affinity for H<sub>2</sub>, expressed as a  
483 low Michaelis-Menten constant,  $K_M$ , or Monod half saturation constant,  $K_S$  (H<sub>2</sub> concentration  
484 at which growth rate reaches half maximum growth rate), and a higher maximum  
485 growth/reaction rate,  $V_{max}$  or  $\mu_{max}$  for Michaelis-Menten kinetics and Monod kinetics,  
486 respectively) were proposed as the underlying cause for the few examples of the poor  
487 competitiveness of homoacetogens [172]. Very limited information on the H<sub>2</sub> consumption  
488 kinetics of homoacetogenic bacteria is available in literature [173]. The available data show  
489 that  $\mu_{max}$  differs by one order of magnitude between strains (0.02-0.5 h<sup>-1</sup>) [4, 173]. This may or  
490 may not be lower than the  $\mu_{max}$  for SSRM 0.057-5.5 h<sup>-1</sup> [4, 40, 174] and methanogens 0.032-  
491 1.4 h<sup>-1</sup> [40, 174]. Krumholz et al. [172] showed that homoacetogens were not able to compete  
492 effectively for H<sub>2</sub> in the presence of SSRM in a subsurface sandstone ecosystem at 30 °C  
493 regardless of  $pH_2$ , and despite significant homoacetogenesis at excess H<sub>2</sub>. Findings by Berta et  
494 al. [4] for a groundwater sediment held under excess  $pH_2$  and 20 °C contrasts this as  
495 homoacetogenesis rates were up to 21 times higher than SSR.

496 Environmental conditions may be a crucial determinant for the competitiveness of  
497 homoacetogens, as low temperatures (~15 °C) [175, 176] and low pH values [64, 93] favor  
498 their growth over methanogens. Under excess  $pH_2$ , homoacetogenic strains with high  $\mu_{max}$  such



499 as *Acetobacterium bakii* will outcompete methanogens [176]. The outstanding metabolic  
500 flexibility of homoacetogens for utilizing a vast variety of substrates may additionally explain  
501 why homoacetogens can compete with more specialized microorganisms like SSRM or  
502 methanogens [65, 176, 177].

503 As for the competitiveness of methanogens and SSRM, the H<sub>2</sub> thresholds of methanogens  
504 may be comparable (1-15 nM) or higher (>15-95) than for sulfate reducers and significantly  
505 lower than for sulfur reducers (<<2500 nM; Table 1), indicating an advantage of sulfate  
506 reducers over methanogens and sulfur reducers in most non-engineered, low p<sub>H<sub>2</sub></sub>  
507 environments. In line with this, Lackner [178] recently reviewed that sulfate reducers  
508 outcompete methanogens for H<sub>2</sub> in most studies. However, at excess H<sub>2</sub>, methanogens and  
509 sulfate reducers would be expected to process equal shares of the in situ H<sub>2</sub> pool [174]. Also,  
510 since concentrations of sulfate are much lower than bicarbonate in non-marine natural  
511 environments [40] (Table 2), the growth of sulfate reducers at excess H<sub>2</sub> will be limited by  
512 availability of their electron acceptor, making it possible for methanogens to compete [40]. As  
513 a general rule pH values below 7 favor the growth of methanogens over sulfate reducers [130].  
514 Above pH 7.5, sulfate reducers grow faster than methanogens and would be expected to  
515 outcompete them [130].

516 Syntrophic relationships between different functional groups have been documented frequently  
517 (whereby the metabolic products of one group serve as substrates for the other). For example,  
518 SSRM and homoacetogens were shown to participate cooperatively in microbial induced  
519 corrosion of steel where SSRM grew on acetate produced by homoacetogenesis [70]. Substrate  
520 provision by the co-culturing *Desulfovibrio vulgaris* enhanced growth of the dehalogen  
521 *Dehalococcoides ethenogenes* 195 by 24 % and caused three times higher dechlorination rates  
522 [179]. Syntrophy may also explain the detection of a combination of the SSRMs *Desulfovibrio*

523 and the homoacetogens *Acetobacterium* in petroleum and subsurface CO<sub>2</sub> reservoirs [62, 180],  
524 and the presence of H<sub>2</sub>-producing heterotrophs along with methanogens in petroleum reservoirs  
525 where the latter rely on H<sub>2</sub>-transfer by the former [181].

#### 526 **2.4 Microbial ecology in natural gas and petroleum reservoirs**

527 Recent years have seen a considerable effort in describing deep subsurface microbial  
528 communities, including those from gas and petroleum reservoirs. Isolated hydrogenotrophic  
529 microbes from these habitats are from the SSRM families *Archaeoglobaceae* [182],  
530 *Desulfomicrobiaceae*, *Desulfobulbaceae*, *Peptococcaceae*, *Desulfobacteraceae*,  
531 *Desulfovibrionaceae*, *Desulforobacteriaceae*, *Sulfurospirillaceae*, *Rhodobacteraceae*,  
532 *Ectothiorhodospiraceae*, *Hydrogenothermaceae* [27, 56, 97, 98, 183-187], the  
533 *Eubacteriaceae* and *Sporomusaceae* families which host homoacetogenic strains [97, 186,  
534 188], and the methanogen families *Methanosarcinaceae*, *Methanobacteriaceae*,  
535 *Methanomicrobiaceae*, *Methanopyraceae*, *Methanococcaceae*, *Methanocalculaceae* and  
536 *Methanosaetaceae* [98, 115, 186] in addition to uncultured microbial taxa [56, 184, 185, 189].  
537 Our collection of hydrogenotrophs (Fig. 1) lists many examples of the above microbial  
538 families, including the strain that holds the highest critical temperature for a methanogen,  
539 *Methanopyrus kandleri*. Sulphur reducing families that define the upper temperature limits for  
540 SSRM like *Thermoproteaceae* and *Pyrodictiaceae* were not reported. The cause for their  
541 absence may be a predominance of mesophilic and thermophilic sites but may also reflect a  
542 generally stronger growth of sulfate reducers over sulfur reducers in oil and gas reservoirs.  
543 Ranchou-Peyrouse et al. [98] showed that the microbial community in 35 out of 36 subsurface  
544 wells from seven natural gas storage sites was dominated by sulfate reducers.

#### 545 **2.5 Effect of high hydrogen concentrations on the microbial metabolism and community** 546 **structure**

547 A range of studies investigated the metabolism of methanogens at excess H<sub>2</sub> and ambient  
548 pressure, with unambiguous results. Conrad et al. [190] demonstrated that excess H<sub>2</sub> stimulated  
549 methanogenesis and growth rates in a paddy soil (species not specified). Opposed to this,  
550 results by Topcuoglu et al. [181] and Stewart et al. [191] suggest an inhibitory effect of high  
551 partial pressures of H<sub>2</sub>,  $p\text{H}_2$ , expressed as a ~10-fold drop in the growth yield (cells per mole  
552 CH<sub>4</sub>) of *Methanocaldococcus jannaschii* and a slight drop of ~0.1-0.7 h<sup>-1</sup> in the growth rate.  
553 Similar observations were made for *Methanothermobacter thermoautotrophicus* [192].  
554 However, within the excess H<sub>2</sub> experiment, higher H<sub>2</sub> concentrations stimulated growth [181],  
555 suggesting a complex influence of  $p\text{H}_2$ . Methanogens seem to express a  $p\text{H}_2$ -dependent change  
556 in their ecological strategy, i.e. maximum growth rate vs. maximum growth yield, as a means  
557 to cope with different environmental conditions [181]. Indeed, *M. jannaschii* is capable of  
558 sensing subtle changes in dissolved H<sub>2</sub> concentration and restraining the energy-intensive  
559 growth of flagella to H<sub>2</sub>-limiting conditions whereas at excess H<sub>2</sub> cells are mostly flagella  
560 devoid [193].

561 Only few studies investigated microbial H<sub>2</sub> turnover at high  $p\text{H}_2$  of up to 1.5-24.8 MPa [4, 194,  
562 195]. Methanogens (*M. jannaschii*) showed a strong inhibitory effect at high  $p\text{H}_2$  [194].  
563 However, the authors added CO<sub>2</sub> at a pressure of at least 0.2 MPa to the hydrogen gas mixture  
564 which at  $p\text{CO}_2 > 0.1$  MPa can be toxic methanogens [163]. Hence it is not clear whether H<sub>2</sub> or  
565 CO<sub>2</sub> performed the toxic action. For homoacetogens and SSRM, the H<sub>2</sub> consumption was  
566 shown not to change in response to different  $p\text{H}_2$  of 0.1-3.5 MPa [4, 195], indicating neither  
567 stimulation nor toxicity at different levels of excess H<sub>2</sub>. The comparison to limiting H<sub>2</sub>  
568 conditions was not made.

569 Apart from microbial metabolism, the microbial community may also change in response to  
570 high  $p\text{H}_2$ . Given a perturbation by H<sub>2</sub> injection it can be anticipated that other types of

571 microorganisms, e.g. the in hydrocarbon reservoirs, common fermenters [24, 96, 98, 183] will  
572 decrease in abundance while hydrogenotrophs will increase [7], in line with the Baas Beeking  
573 principle [196]. An increase in hydrogenotrophs in response to H<sub>2</sub> addition was recently  
574 confirmed for soils, however H<sub>2</sub> consumption increased in only one of the investigated soils,  
575 suggesting a pronounced influence of the indigenous microbial community [197]. Bioreactor  
576 experiments support a decrease in microbial diversity in response to high  $pH_2$  as well [198,  
577 199]. Puente-Sanchez et al. [200] were the first to report differences in the subsurface H<sub>2</sub>-  
578 consuming community in response to varying  $pH_2$  within the Iberian Pyrite Belt. Ranchou-  
579 Peyruse et al. [98] showed that town gas storage with more than 50 % H<sub>2</sub> changed the microbial  
580 community from a predominantly sulfate reducing community to a dominance of methanogens,  
581 and this balance was active even decades after injection stopped, possibly via H<sub>2</sub> trapping in  
582 the microporous system [98]. It was suspected that all sulfate was initially used up by SSRM  
583 following increased growth of methanogens [98].

### 584 **3. Evaluating the potential hydrogen consumption in DOGFs**

#### 585 **3.1 Calculation of the microbial growth**

586 We screened 42 DOGF in the North Sea and the Irish Sea and five H<sub>2</sub> storage test sites for  
587 temperature, salinity, pH and pressure data (Fig. 1, Table A.4). We discovered significant  
588 differences in the salinity for the DOGF reported by sources [201] and [168]. Because we relied  
589 on the solution compositions for the calculation of the potential microbial growth in the fields,  
590 which are available from [168], we chose to use the salinity data from the same source.

591 The environmental data from the DOGF and H<sub>2</sub> storage test sites were aligned with the  
592 constraints for growth of methanogens, homoacetogens and SSRM (Fig. 1-2) to select in which  
593 fields growth can be expected. For the few fields that fulfil the growth constraints of all

594 investigated microorganisms, we calculated a first-order estimate of the microbial growth using  
595 the elemental cell composition as a proxy for the nutrient requirement [202, 203] (Text A.1).

596 Our calculations assumed that the supply of N and C are covered by diazotrophic and  
597 autotrophic growth, respectively. Requirements for trace elements were neglected in the  
598 calculation due to a lack of information on the relevant trace element contents in the reservoirs.  
599 Where a nutrient for a specific field was not available we used the average value from the fields  
600 given in Table 2. Any effect of the  $pH_2$  on microbial growth was neglected. We assumed that  
601 cells neither die nor are removed, and that nutrients are not replenished by inflow, re-  
602 mineralization from decaying biomass or mineral dissolution. Simultaneous growth by  
603 different microorganisms was not considered.

604 Percentages of nutrients in the cells (Text A.1) were converted to mass using a wet cell mass  
605 of  $1.77 \cdot 10^{-12}$  g for methanogens [204],  $3.2\text{-}6.2 \cdot 10^{-13}$  g for homoacetogens and  $7.81 \cdot 10^{-13}$  g for  
606 SSRM. The cell wet weight of homoacetogens was calculated by dividing the cell volume of  
607  $1.62\text{-}3.14 \mu\text{m}^3$  for the subsurface mixotrophic homoacetogen *Acetobacterium psammolithicum*  
608 [172] with an assumed bacterial density of  $1 \cdot 10^{-12}$  g  $\mu\text{m}^{-3}$  [205]. The cell wet weight of SSRMs  
609 was calculated using a cell dry weight of  $3.125 \cdot 10^{-13}$  g for *Desulfovibrio desulfuricans* [206]  
610 and dividing this with a general bacterial dry weight to wet weight ratio of 0.4 [207].  
611 Subsequently, the concentrations of C, H, O, Ca, K, Na, S, Mg, P and Fe in the DOGF (Table  
612 2) were divided by the mass of the respective cell nutrients per microbial cell calculated above.  
613 This resulted in the maximum cell count within each microbial group,  $G$ , that could potentially  
614 be created based on a single nutrient, where the lowest  $G$  indicated the limiting nutrient for cell  
615 growth. For an example of those calculations, see Text A.1.

### 616 **3.2 Estimation of the cell-specific hydrogen consumption**

617 Hydrogen may be consumed at rates of  $0.7\text{-}5.0 \cdot 10^5$  nM h<sup>-1</sup> for homoacetogens,  $0.008\text{-}5.8 \cdot 10^5$   
618 nM h<sup>-1</sup> for methanogens and  $0.005\text{-}130 \cdot 10^5$  nM h<sup>-1</sup> for SSRM (Tables A.1-A.3), the latter  
619 considering sulfate concentrations in the range of  $0\text{-}2.3 \cdot 10^{-2}$  M in the DOGF (Table 2). In a  
620 few studies, the microbial H<sub>2</sub> consumption was related to growth (Tables A.1-A.3), enabling  
621 the calculation of the H<sub>2</sub> consumption per synthesized cell and the time for when the microbial  
622 cell count  $G$  would be reached (Text A.2).

### 623 **3.3 Calculation of the hydrogen consumption in a hydrogen storage system**

624 We calculated the minimum H<sub>2</sub> consumption for the DOGF Frigg and Hamilton by dividing  
625 the H<sub>2</sub> consumption per synthesized cell with the microbial cell count. The calculation of the  
626 moles of H<sub>2</sub> the in aquifer anticipated equal volumes of H<sub>2</sub> and water and used the ideal gas  
627 law and the field size, temperature and pressure data in Table 2 and Table A.3. The percentage  
628 of H<sub>2</sub> that was consumed as a function of growing and resting microbial cells was calculated  
629 by dividing the potential H<sub>2</sub> consumption with the H<sub>2</sub> concentration in the reservoir. Text A.3  
630 shows our calculations for the Frigg reservoir and methanogens.

## 631 **4. Results and discussion**

### 632 **4.1 Characterization of the likelihood for growth in 42 DOGF**

633 Using the environmental limits constraining microbial growth on H<sub>2</sub>, we analyzed the  
634 physicochemical parameters for 42 DOGF in the British and Norwegian North Sea and the  
635 Irish Sea and five H<sub>2</sub> storage test sites (Fig. 1, Table A.4). Of the 47 fields, five fields have a  
636 temperature of 122 °C or higher and may be considered sterile with respect to H<sub>2</sub>-consuming  
637 microorganisms. Thirty-two fields have a temperature >72 °C, implying that homoacetogenesis  
638 cannot take place. Twenty fields have a temperature ≥90 °C implying that homoacetogenesis  
639 and sulfate reduction cannot take place. Fourteen DOGF have a temperature >90 °C and <122  
640 °C and pressures of 18.2-44 MPa where (piezophile) methanogens and SSRM will grow.

641 Of the fifteen sites with temperatures  $<72\text{ }^{\circ}\text{C}$  where all investigated groups of microorganisms  
642 will grow, only six fields (Frigg, Hamilton, Veslefrikk, Ketzin, Lehen and Lobodice) fulfill the  
643 remaining pressure and salinity requirements for growth. Two fields, Leman and Rhyl, have  
644 salinities  $\geq 5.8\text{ M}$  where no significant microbial growth can be expected. This finding is  
645 supported by stable gas compositions at the similarly saline  $\text{H}_2$ -storage test sites of the  
646 H2STORE project, Emsland and Altmark (Fig. 1, Table A.4), though a low microbial  
647 population of  $\sim 10^2\text{ cells ml}^{-1}$  was present [208]. Lennox, North Morecambe and South  
648 Morecambe with temperatures of  $30\text{-}33\text{ }^{\circ}\text{C}$  and salinities of  $4.6\text{-}5.1\text{ M}$ , could permit the growth  
649 of the halophile homoacetogenic strain *Halanaerobium praevalens*, only. Hamilton North,  
650 Camelot and The V gas field complex with salinities of  $2.9\text{-}5.0\text{ M}$  may permit the growth of  
651 sulfate reducers and *Halanaerobium praevalens*. The Viking field has temperature of  $65\text{-}80\text{ }^{\circ}\text{C}$   
652 and a salinity of  $3.8\text{ M}$  and so is likely to host only mesophilic SSRM, although pressures  $>30$   
653 MPa that could become growth inhibiting. The  $\text{H}_2$ -storage test site Ketzin has similar salinity  
654 to the Viking field but a lower pressure ( $4.0\text{ M NaCl}$ ,  $35\text{ }^{\circ}\text{C}$ ,  $6\text{ MPa}$ ). Here SSRM were  
655 suspected to cause a  $2\text{-}4\%$  decrease in  $\text{H}_2$  and a reduction in the concentration of sulfate from  
656  $22\text{ to }8 \times 10^{-3}\text{ M}$  [208].

#### 657 **4.2 Microbial growth estimates for three low-temperature and low-salinity DOGF**

658 Our first order approach for calculating microbial growth in the Veslefrikk reservoir yielded a  
659 maximum  $1 \times 10^8$  methanogenic cells  $\text{mL}^{-1}$ ,  $2 \times 10^8$  SSRM cells  $\text{mL}^{-1}$  or  $5 \times 10^8$  homoacetogenic  
660 cells  $\text{mL}^{-1}$ . The Frigg reservoir a maximum of  $1 \times 10^8$  methanogenic cells  $\text{mL}^{-1}$ ,  $1 \times 10^8$  SSRM  
661 cells  $\text{mL}^{-1}$  or  $2 \times 10^8$  homoacetogenic cells  $\text{mL}^{-1}$ . The Hamilton reservoir could host a maximum  
662 of  $1 \times 10^7$  methanogenic cells  $\text{mL}^{-1}$ ,  $2 \times 10^7$  SRCM cells  $\text{mL}^{-1}$  or  $6 \times 10^7$  homoacetogenic cells  $\text{mL}^{-1}$ .  
663 <sup>1</sup>. These cell counts describe a maximum cell growth for each hydrogenotrophic group because  
664 simultaneous growth of hydrogenotrophs was not considered. The higher growth of  
665 homoacetogens over SSRM and methanogens results from a lower wet cell mass that causes a

666 lower nutrient demand per cell (see Text A.1). Our calculations are in line with total cell  
667 concentrations of  $10^5$ - $10^{15}$  cells/ mL<sup>-1</sup> in oil reservoirs [209], and equal to or up to four order  
668 of magnitudes higher than cell counts from gas reservoirs ( $0.001$ - $1.2 \cdot 10^7$  cells mL<sup>-1</sup>)[51, 97,  
669 185]. The range of the literature data reflects that our simple methodology to calculate  
670 microbial numbers may serve as a first approximation to estimating microbial growth in the  
671 subsurface, only.

672 Acknowledging that trace elements were not accounted for in our calculation, N and P are the  
673 first limiting nutrients in the reservoirs Frigg, Hamilton and Veslefrikk. However, this does not  
674 imply that microbial growth is N and P limited, as many microorganisms may use of  
675 ammonium (not measured) as N-source, and in the Hamilton reservoir the C:P ratio was  
676 between 59:1 and 158:1, whereas the limiting C:P ratio for microbial growth is in the range of  
677 400:1 to 800:1 (reported for the SSRM *D. desulfuricans*) [92]. At moderately acidic pH values  
678 such as the pH of 5.8 in the Hamilton reservoir, P may further be continuously replenished by  
679 mineral buffering with apatite.

#### 680 **4.3 Hydrogen consumption in three low-temperature and low-salinity DOGF**

681 The H<sub>2</sub> consumption in the Frigg reservoir by homoacetogens constitutes <0.01- 3.2 % of the  
682 H<sub>2</sub> in the aquifer, <0.01- 1.3 % for methanogens and <0.01- 1.3 % for SSRM. In the Hamilton  
683 reservoir, the rates are <0.01- 2.0 %, <0.01- 2.3 % and <0.01- 0.5 % for homoacetogens,  
684 methanogens and SSRM, respectively. For actively growing cells these consumption rates may  
685 be reached after only 0.1-19.1 days, which is the time it takes for the microorganisms to grow  
686 up to their maximum cell counts, based on the dissolved nutrient concentrations. Resting cells,  
687 i.e. cells that undergo no or only very little cell division, need 2.5-3.5 months (SSRM) or up to  
688 3.6-6.6 years (methanogens) to reach the maximum cell count and consume the given  
689 percentage H<sub>2</sub>.



690 In a real aquifer system, nutrients are likely to at least partly be replenished by decaying cells,  
691 mineral weathering and inflowing brine, and cells will continue to consume H<sub>2</sub> beyond the time  
692 it takes to reach the maximum cell count (maintenance). As such our H<sub>2</sub> consumption estimates  
693 may be regarded as minima. On the other hand, considering that, with the exception of one  
694 study (Berta et al. [4]), our calculations employ laboratory H<sub>2</sub> consumption rates at optimal  
695 nutrient supply and optimal physicochemical conditions (Tables A.1-A.3), the H<sub>2</sub> consumption  
696 in the oligotrophic subsurface is likely overpredicted. Comparing the employed laboratory H<sub>2</sub>  
697 consumption rates to H<sub>2</sub> consumption rates by SSR and methanogenesis in oil and natural gas  
698 reservoirs of ~0.05-351 nM h<sup>-1</sup> and 0-1185 nM h<sup>-1</sup>, respectively (SO<sub>4</sub><sup>2-</sup>: 8.3-805\*10<sup>-5</sup> M; HCO<sub>3</sub><sup>-</sup>  
699 : 3.5-246\*10<sup>-4</sup> M) [51, 183], shows that the field H<sub>2</sub> consumption by SSR is 1.5 times to eight  
700 orders of magnitude lower, and 0.7 times to 7 orders of magnitude lower for methanogenesis.  
701 Within the operation and injection wells of a natural gas reservoir, H<sub>2</sub> consumption rates by  
702 SSR and methanogenesis were up to 2544 and 4533 nM h<sup>-1</sup>, respectively, [51], which falls  
703 within the lower range of the values reported from laboratory studies. Acknowledging the  
704 unknown but presumably low *p*H<sub>2</sub> in above experiments, and that maintenance requirements  
705 were not included in our H<sub>2</sub>-consumption calculations, we expect the actual H<sub>2</sub> consumption  
706 in a H<sub>2</sub> storage system to lie within the higher range of our calculated values.

707 Our lower-end results are in agreement with no H<sub>2</sub> consumption during storage operations of  
708 H<sub>2</sub> -rich town gas in Beynes, France [210]. Our upper end results are in agreement with a loss  
709 of ~3 %, presumably by methanogenesis, at the H<sub>2</sub> storage (SunStorage) test site in Lehen,  
710 Austria [211]. A H<sub>2</sub> consumption of 17 % by methanogens at the Lobodice town gas storage  
711 site over a time span of seven months [210, 212] seems exceptional in the light of our  
712 calculations and the reported SSR and methanogenesis rates from the field. With a very low  
713 salinity of 0.03 M, temperatures of 20-45 °C, a pH of 6.7 and 4 MPa pressure, Lobodice is  
714 among the few sites which has highly favorable conditions for microbial growth considering

715 *all* of these parameters (Table A.4). The high H<sub>2</sub> consumption at Lobodice highlights the  
716 importance of our site selection tool, as H<sub>2</sub> storage may face serious economical and technical  
717 problems if a site with growth-favoring conditions is selected.

718 As mentioned, Berta et al. [4] measured high H<sub>2</sub> consumption rates under excess H<sub>2</sub> and  
719 oligotrophic conditions (P < 9.7\*10<sup>-7</sup> M; SO<sub>4</sub><sup>2-</sup> ≤ 9.5\*10<sup>-4</sup> M; DOC = 2.6\*10<sup>-4</sup> M), indicating that  
720 nutrient scarcity does not imply low H<sub>2</sub> consumption. A comparison to the nutrient  
721 concentrations in the DOGF reveals that many of them have a higher nutrient status (P = 0.002-  
722 0.452\*10<sup>-3</sup> M; SO<sub>4</sub><sup>2-</sup> = up to 23.1\*10<sup>-3</sup> M; organic acids = 1.2-8.1\*10<sup>-3</sup> M, Table 2), implying  
723 that H<sub>2</sub> consumption in DOGF under excess H<sub>2</sub> conditions may be even higher than reported in  
724 [4]. The experiment by Berta et al. [4] is further highly relevant because cells were at steady  
725 state, i.e. at the predominant growth stage in nature, but still consumed vast amounts of H<sub>2</sub>.  
726 Indeed the H<sub>2</sub> consumption of cells at steady state or resting may be just as high or higher than  
727 for growing cells but growth is low or absent (Tables A.1-A.3).

#### 728 **4.4 Knowledge gaps and future research**

729 More work is needed to predict the magnitude of microbial growth, H<sub>2</sub> consumption rates, and  
730 (not least) the mutual interaction of the microbial processes in DOGFs. The list of unknowns  
731 and uncertainties is long. To begin with are the poorly elucidated nutrient requirements of the  
732 microorganisms, especially in mixed cultures (e.g., [71]). Adding to this are the missing or  
733 incomplete datasets on the physical environment of certain reservoirs along with their gas phase  
734 and brine compositions, including chaotropy and kosmotropy characteristics. A better  
735 elucidation of the latter would allow the calculation of the dominating microbial processes via  
736 their free energies of the reaction. Combined with an analysis of the microbial community and  
737 metabolism this could give new insights into whether or not we can theoretically predict which  
738 microbial processes occur in DOGF and to which extend.

739 A further complication is the non-cultivability of many microorganisms in the deep subsurface,  
740 including DOGF [12, 34, 56, 98, 183]. Considering tiny culturabilities of  $\leq 0.1\%$  of the total  
741 viable cell count in many subsurface environments [34], any attempts to assign sterile habitats  
742 or quantify microbial  $H_2$  consumption via cultivated microorganisms may seem in vain. In gas  
743 reservoirs, the percentage of cultured bacteria may be higher, ranging between 86-95% within  
744 each phylum [98]. Field-based metabolic activity measurements could circumvent any non-  
745 cultivability issues observed in laboratory experiments. Field studies should also be prioritized  
746 considering that microbial cell sizes and masses in nature are only 4-21 % of the laboratorial  
747 cell masses [213] which reduces the nutrient requirement per cell, thereby allowing more cells  
748 to proliferate on any given amount of nutrients.

749 The lack of knowledge about the changes in microbial ecology as a response to increased  $H_2$   
750 concentrations beyond the level of functional groups is one of the major hurdles in our attempt  
751 to understand of the effect of high  $H_2$  concentrations on the subsurface microbiology. Emerging  
752 evidence on the subject highlights species-specific responses to high  $pH_2$  [98, 198, 200], and  
753 that  $H_2$  injection may leave its fingerprint on the subsurface microbial community for decades  
754 [98]. Knowledge about the initial effect of a drastic increase in  $pH_2$  in the subsurface is lacking.  
755 One possibility is that more EPS will be produced as a response to the perturbation with  
756 increased  $H_2$ , as has been shown for other types of perturbation [18, 92, 214], and considering  
757 the toxicity of high  $pH_2$  on methanogens [181, 192, 194], with possible adverse effects on gas  
758 injectivity and withdrawal.

759 Future research should address the effect of high  $pH_2$  on the metabolisms of different functional  
760 groups and the EPS production in different geological settings and under changing nutritional  
761 supply and physicochemical conditions. Mixed culture studies at low and high  $pH_2$  can give  
762 insight into competitive and syntrophic relations under these conditions and reveal changes in

763 the microbial community structure due to the perturbation with elevated H<sub>2</sub>. Protocols for the  
764 careful cultivation of nutrient-deprived deep subsurface cells need to be developed. More base-  
765 line research includes determinations of the critical salinities and pressure tolerances that to  
766 date are missing for many cultivated strains, as well as the study of the brine compositional  
767 effects on the microbial community and metabolism. Research employing already cultivated  
768 species can make use of the fact that the large majority of the cultivated species isolated from  
769 subsurface environments can be found in other near-surface marine and terrestrial geothermal  
770 environments [9, 183], and should employ chemostat studies that mimic the natural  
771 environment.

## 772 **5. Conclusion**

773 Here we presented a novel site selection tool for H<sub>2</sub> storage and demonstrated its application  
774 for 42 DOGF in the British and Norwegian North Sea and the Irish Sea and five H<sub>2</sub> storage test  
775 sites. Our results highlight the hard limits to the cultivable microbial life on H<sub>2</sub> and can –with  
776 some certainty- exclude life in several high-salinity or high-temperature, i.e. deeper reservoirs.  
777 For low-salinity and low-temperature reservoirs our calculations indicate significant microbial  
778 growth and a small but not insignificant H<sub>2</sub> consumption, both of which may further increase  
779 during repeated storage cycles, giving replenishment of nutrients by mineral weathering,  
780 decaying microbial cells and inflowing water. Hence, from the point of view of minimizing H<sub>2</sub>  
781 loss, clogging and corrosion, sites with more extreme conditions may be chosen over low-  
782 temperature and low-salinity reservoirs where the majority of microorganisms can proliferate.  
783 Yet, any storage operation will have to consider increased operational difficulties and costs  
784 with increased depth. Additional investigation on subsurface life on H<sub>2</sub> is encouraged to help  
785 manifest whether H<sub>2</sub> consumption in low-temperature aquifers is a threat to H<sub>2</sub> storage.

786

## 787 ASSOCIATED CONTENT

788 **Appendix.** Figure A.1 shows the solubility of hydrogen as a function of temperature and  
789 pressure. Laboratories studies investigating homoacetogenesis, methanogenesis and SSR are  
790 listed in Tables A.1, A.2 and A.3, respectively. Table A.4 provides the reservoir conditions  
791 for 42 DOGF and five H<sub>2</sub> storage test sites. Text A.1 holds a discussion of the importance of  
792 other hydrogen oxidizing processes for hydrogen storage. A detailed calculation of the  
793 number SSRM cells that could grow based on the N content in the Frigg reservoir can be  
794 found in Text A.1. Text A.2 and A.3 hold an example calculation of the hydrogen  
795 consumption per synthesized cell and the calculation of the potential hydrogen consumption  
796 in a hydrogen storage system, respectively.

## 797 **Author Contributions**

798 The manuscript was written through contributions of all authors. All authors have given  
799 approval to the final version of the manuscript.

## 800 **Declaration of interest**

801 The authors declare no competing financial interest.

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