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29	Estimating	Microbial	Hydrogen	Consumption	in
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Hydrogen Storage in Porous Media as a Basis for

Site Selection

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ABSTRACT

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

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

Highlights

- Review of the most important hydrogen-utilizing microorganisms in the underground.
- Elucidation of the growth criteria for 480 strains of the mayor hydrogenutilizers.

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

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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⁻¹)
- 80 MPa Megapascal

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1. Introduction

Zero-carbon energy generation from renewable sources can help mitigate carbon emissions and abate climate change [1-3]. One of the most significant challenges for renewable energy is the imbalance between supply and demand [3, 4]. The generation of hydrogen (H₂) via electrolysis of water during periods of renewable energy oversupply and subsequent H₂ storage is one way of overcoming this imbalance, as H₂ can be recovered and used for electricity generation during periods of renewable energy shortage [1, 5]. Subsurface storage of H₂ in salt caverns, depleted gas or oil fields or saline aquifers is being considered as an alternative to expensive purposebuilt storage containers [6]. However, the artificial elevation of the H₂ concentration in the subsurface may stimulate the growth of H₂-oxidizing (hydrogenotrophic) bacteria and archaea, here collectively referred to as microorganisms, with possible adverse implications for gas

injectivity and withdrawal via permeability reduction, H₂ volume loss and corrosion of metal infrastructure [4, 7]. Understanding the controls on microbial H₂ metabolism is therefore highly important.

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

Hydrogen plays a central role in the energy metabolism of subsurface life [9]. Yet, a quantitative assessment of the consumption of H₂ by deep microbial communities in the context of the global H₂ cycle is lacking [26]. In underground gas storage sites and oil reservoirs the most abundant H₂-oxidizers are hydrogenotrophic sulfate reducers, that couple H₂-oxidation to sulfate reduction to produce hydrogen sulfide (H₂S); hydrogenotrophic methanogens that reduce carbon dioxide (CO₂) to methane (CH₄) by oxidizing H₂; and homoacetogens that couple H₂ oxidation to carbon dioxide (CO₂) reduction producing acetate [7, 24, 27]. These

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

A recent review addressed the many possible abiotic and biotic H₂-producing and H₂-consuming processes in the subsurface [7]. However, it lacked a quantitative assessment of the processes of microbial growth and H₂ consumption relevant for H₂ storage. Strobel et al. [29] summarized the concept and potential of underground methanation using experimental data from the Sun Storage project. These authors highlighted controls on the growth of methanogens and changes in gas composition due to methanogenesis, but did not quantify microbial growth. Many studies report changes in gas composition, biofilm growth and clogging near injection wells but hardly any studies report quantitative figures on microbial growth or on permeability changes [30].

To date it remains unclear how subsurface microorganisms might react to elevated H₂ concentrations [7] and hence whether microbial growth is a concern for H₂ storage. Even in natural, non-engineered subsurface environments, there is little information on microbial H₂ turnover rates [31] and the behavior and population kinetics of microorganisms are not fully understood [29]. The majority of the available data on microbial H₂ turnover rates come from batch cultures at optimal growth conditions where the kinetics [31], the pace of life [32, 33], the physiological states and the prominent organisms may differ widely from the subsurface environment [7, 32]. A further complication arises from the fact that many microorganisms in the deep subsurface are not culturable with modern enrichment techniques [12, 34].

In this work, we review the state-of-the-art understanding of the controls of temperature, salinity, pH, pressure and nutrients and water on microbial growth on H₂ in the subsurface, with emphasis on the three major H₂-consuming processes methanogenesis, sulfate reduction and homoacetogenesis, to determine what reservoir conditions will be unfavorable to microbial

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

Physicochemical data from 42 depleted or close to depleted oil and gas fields (DOGF) of the British and Norwegian North Sea and the Irish Sea as well as five H₂ storage test sites provide the base for an evaluation of the number of sites where microbial growth of methanogens, sulfate reducers and homoacetogens can be expected. Using average nutrient contents of the microbial cells and site-specific dissolved ion concentrations, we calculate significant growth and a small H₂ consumption for growth-permitting DOGF.

2. State of the art understanding

2.1 Likely microbial hydrogen oxidation in hydrogen storage systems

Hydrogen oxidizing processes may be ranked according to the magnitude of their H_2 threshold and their standard free energy change ($\Delta G^{0'}$), two useful metrics to compare the likelihood of reactions to take place and the order at which they proceed (Table 1). The H_2 threshold defines the concentration of H_2 below which it is no longer consumed. Given all other factors are at optimum, the microbial population with the lowest H_2 threshold value is expected to be the most successful population in competing for H_2 [35].

The $\Delta G^{0'}$ marks the thermodynamic favorability of a reaction at ambient pressure and temperature, pH 7 and 1 M of all reactants. In oligotrophic (nutrient poor) high pressure and temperature environments, the order of the $\Delta G^{0'}$ may be used to determine which reaction is more energetically favorable. As can be seen from Table 1, more negative $\Delta G^{0'}$ values (more available free energy) are generally accompanied by lower H₂ thresholds. Not included in Table 1 are the kinetics which describe the rate of the electron transfer in the redox reaction.

Table 1. Biotic H₂-consuming processes ranked according to their free energy yield (ΔG^{0}) and measured H₂ threshold. Not included are Vanadium, Cobalt, Techneticum, Uranium and Selenium reduction, due their limited relevance for H₂ storage. NA= not available.

H ₂ - oxidizing process	Reaction (number)		H2 threshold (nM)	ΔG ⁰ ' (KJ mol H ₂ -1)	Typical ambient [H ₂] (nmol L ⁻¹)	Relevance for H ₂ storage
Chromate reduction	$\frac{1}{2}H_2 + \frac{1}{3}CrO_4^{2-} + \frac{5}{3}H^+ \to \frac{1}{3}Cr^{3+} + \frac{4}{3}H_2O$	(1)	<0.1 ^[36]	NA	NA	low
Aerobic hydrogen oxidation (Knallgas)	$H_2 + \frac{1}{2}O_2 \rightarrow H_2O$	(2)	$0.051^{[7]}$	-237 ^[7, 36]	NA	low
Denitrification	$H_2 + \frac{2}{5}H^+ + \frac{2}{5}NO_3^- \rightarrow \frac{1}{5}N_2 + \frac{6}{5}H_2O$	(3)	<0.05-0.5 ^[7]	-240.1 ^[7, 36] -224 ^[4, 37]	<0.05 ^[4, 35, 36]	low
Halorespiration	H_2 + halogenated compounds \rightarrow dehalogentated compounds + HCl	(4)	$0.05 - 0.27^{[36]}$ < $0.3^{[38]}$ $0.27 - 2^{[7]}$	-230 to -187 ^[7]	NA	low
Iron (III) reduction	$H_2 + ferric(oxy)hydroxides \rightarrow ferrous ir H_2O$	on + (5)	<0.11-0.8 ^[36, 38]	-228.3 ^[7, 38] -182.5 ^[36] -114 ^[4]	$0.2^{[4,35]} \\ 0.2 - 1^{[36]}$	intermediate
Manganese (IV) reduction	$2H_2 + MnO_2 \rightarrow Mn(OH)_2 + 2H_2O$	(6)	<0.05 ^[35]	-163 ^[4, 35]	<0.05 ^[4, 35]	low
Arsenate reduction	$H_2 + HAsO_4^{2-} + 2H^+ \rightarrow H_3AsO_3 + H_2O$	(7)	$0.03 \text{-} 0.09^{[36]}$	-162.4 ^[36]	$0.4 \text{-} 0.7^{[36]}$	low
Ammonification	$4H_2 + 2H^+ + NO_3^- \rightarrow NH_4^+ + 3H_2O$	(8)	$0.015 0.025^{[38,}$	-150 ^[4, 38]	<0.05 ^[4, 35]	low
Fumarate reduction	H_2 + fumarate \rightarrow succinate	(9)	$0.015^{[38, 39]}$	-86.2 ^[38]	NA	low
Hydrogenotrophic sulfate reduction	$4H_2 + SO_4^{2-} + H^+ \rightarrow HS^- + 4H_2O$	(10)	1-15 ^[38, 39]	-38 ^[7, 38] -48 ^[36] -57 ^[4]	1-2 ^[4, 35]	high
Hydrogenotrophic methanogenesis	$4H_2 + CO_2 \rightarrow CH_4 + 2H_2O$	(11)	0.4-95[38-40]	-34 ^[4, 38] -43.9 ^[36]	5-10 ^[4, 35] 7-13 ^[36]	high
Sulfur reduction	$H_2 + S \rightarrow HS^- + H^+$	(12)	$2500^{[7]}$	-33.1 ^[7]	NA	intermediate
Homoacetogenesis	$4H_2 + 2CO_2 \rightarrow CH_3COOH + 2H_2$	(13)	328-3640 ^[38, 39]	-26 ^[4, 38] -36.1 ^[36]	$100^{<[4]}, 117-150^{[36]}$	high

Abiotically, most of the H_2 -oxidizing reactions are very slow but mediated by microbial enzymes the processes are catalyzed [37, 41].

The three main microbial processes with implications for H_2 storage, hydrogenotrophic sulfate reduction, hydrogenotrophic methanogenesis (for simplicity from now on just referred to as sulfate reduction, and methanogenesis unless otherwise specified) and homoacetogenesis, require the highest threshold $[H_2]$ and are among the processes with lowest $\Delta G^{0'}$ (Table 1). Nevertheless, e.g. sulfate reduction is instantaneous in most geologic settings [42] possibly due to fast kinetics [37] and/or a relatively high availability of sulfate.

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

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

include non-hydrogenotrophs (e.g. [51, 52]). In non-engineered environments rich in Fe oxides and organic carbon, IRB may have a great advantage over SSRM, methanogens and homoacetogens, due to a very high affinity for H₂ [47]. We evaluate this process as of intermediate relevance for H₂ storage in DOGF.

Many IRB and a few SSRM can also couple H₂ oxidation to reduction of a variety of other trace metal oxides, e.g. MnO₄²⁻/MnO₂, CrO₄²⁻, Co, SeO₄²⁻, UO₂², TcO₄⁻, AsO₃⁻, and VO₄⁻ [43, 53, 54]. After Fe, the most abundant metal in sedimentary environments is Mn (~10 % of Fe abundance) [47, 53]. Due to the trace content of these compounds in the environment, their reduction has low relevance for H₂ storage.

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

Halogenated compounds are common in aquifers, and may arise from contamination or via natural processes in sediment [60, 61]. However, the concentrations of these compounds are extremely low: In aquifers of 167-1000 m depth, chloroflourocarbons reach maximum concentrations of \leq 1.1 µg L⁻¹ [61] and for pristine aquifers 0.003-0.007 µg L⁻¹ of chlorinated hydrocarbons were measured [60]. We evaluate the relevance of this process to H₂ storage as negligible.

Literature on the importance of anaerobic fumarate respiration using H₂ is scarce. Fumarate may be used as an alternative electron acceptor by SSRM [43, 62] and homoacetogens [63-66]. In the non-engineered subsurface, readily metabolizable organic matter, like fumarate, is rare [67]. Oil fields being rich in organic C compounds may contain more fumarate. Payler et al. [12] confirmed the presence of fumarate reductase, the key enzyme in fumarate reduction, in three out of five metagenomes from subsurface brines within sandstone. However, the

metagenomes belonged primarily to non-H₂ utilizing bacteria (*Halorubrum*) and fumarate concentrations were not reported. Acknowledging the lack of data in this field, we evaluate this process as being of low relevance for H₂ storage.

2.2 Factors governing microbial growth

Microbial growth and H₂ consumption rates vary with nutrient availability and environmental variables (e.g. [17, 68].) Each strain is adapted to an optimum set of nutrients and environmental conditions where potentially the greatest growth rates occur. Beyond the optimum conditions, organisms may grow but at reduced rate or they become dormant. In this section, we discuss the requirements for nutrients and water, and the overall impact of temperature, salinity, pH and pressure on the growth of the major microbial H₂—oxidizers in DOGFs, in the ranges relevant to H₂ storage. The specific activity of microbial strains grown at optimum conditions varies as well (reviewed in [69]) but the elucidation of differences between strains is beyond the scope of this review.

2.2.1 Nutrients

Apart from water of sufficient thermodynamic activity (see Section 2.2.4), hydrogenotrophs require H_2 as a source of electrons (energy), an electron acceptor and a carbon source for cell division, together with a set of macro and trace elements as well as various organic nutrients [70]. Microorganisms can only access $H_{2(aq)}$ and hence the solubility of $H_{2(g)}$ is of direct relevance for all H_2 -consuming reactions. Given a gas phase of ~100 % H_2 in an H_2 storage system, the equilibrium solubility of H_2 exceeds the highest threshold value of an H_2 -consuming microorganism of 3.6 μ M (Table 1) by ~3 orders of magnitude at ambient pressure and temperature and under static conditions (Fig. A.1a), with further increase at higher pressures (Fig. A.1b). While under non-static conditions hydrogenotrophs will consume part

- of the H₂, these figures suggest no limitation by the H₂ solubility on microbial growth under
- 237 H₂ storage conditions.
- 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
- additionally require different vitamins (e.g. lipoic acid, biotin, riboflavin, folic acid, thiamine,
- etc.), yeast extract, coenzyme M, aromatic acids and phospholipids or a combination of these
- 243 (e.g. [8, 65, 73-75]).
- Nutrients may be assimilated from the solution or directly from minerals (e.g., [76-79]), the
- 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
- 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 CO₂ alone (autotrophy) or from organic carbon
- compounds (heterotrophy) [81]. Methanogens and homoacetogens can grow autotrophically or
- heterotrophically, and several can grow mixotrophically (e.g. [66, 82, 83]). SSRM typically
- grow heterotrophically but some grow autotrophically or mixotrophically [84, 85]. Nitrogen
- 254 may be assimilated from ammonia and nitrate or by nitrogen-fixation (diazotrophy).
- Diazotrophy is common amongst SSRM, methanogens and homoacetogens [86-89], though
- 256 homoacetogens often inhabit ammonia-rich environments [88].
- 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
- requirement for iron (1.8*10⁻⁶ M) than is usually observed for microorganisms [90] while

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

Literature on when nutrients become limiting is also scarce. Sulfate reducing SSRM require a minimum sulfate and phosphorus concentrations of ~3 mM and ~3.2-320*10⁻⁵ mM, respectively, for growth [92, 93]. Methanogens of the order *Methanosarcinae* require 29.6 mM Mg for optimum growth and growth ceases at 15.8 mM (reviewed in [91]). When grown under optimum conditions, the growth rate of autotrophs may be limited by the rate of transfer of H₂ and CO₂ from gas to liquid, as was shown for the methanogen *Methanobacterium* thermoautotrophicum [94] and for the sulfate reducers within *Desulfotomaculum sp* [95].

Carbon is unlikely to be limiting in the hydrocarbon-rich DOGF [56, 96, 97] but this is not a given in saline aquifers with no history of oil or gas. Sulfate is present in significant concentrations in most DOGF (Table 2) but H₂ injection can cause sulfate depletion due to accelerated growth of SSRM [98]. Nitrogen in the form of ammonia may be limiting in DOGF [51, 56, 90] but nitrate levels may be elevated [51], often due to contamination by drilling fluid [57-59].

2.2.2 Temperature

Temperatures of 22.5–80 °C or 20–100 °C have been suggested for H₂ storage based on a recommended depth range of 500- 2000 m for H₂ storage in DOGF and saline aquifers [99-101]. Microorganisms are classified according to their preferred growth temperature: psychrophiles grow optimally below 20 °C, psychotrophs grow optimally at or above 20 °C and may tolerate temperatures below 5 °C, mesophiles grow between 20 and 45 °C, thermophiles grow above 45-50 °C, and hyperthermophiles show optimal growth at temperatures of 80 °C or above [102, 103].

High temperatures alter the energetic properties (e.g., vibrational modes) of biomolecules in their aqueous solvent, change the substrate solubility or viscosity and the ionization of the aqueous medium [104]. Adverse effects of high temperature include DNA denaturing or damage, decreased protein stability, hydrolysis of ATP and ADP, amongst others [104, 105]. The metabolic strategies of thermophiles are highly diverse. For a discussion, the reader is referred to [106]. Thermophiles and hyperthermophiles are challenged by increased reaction rates at elevated temperature which can imply that abiotic reaction rates are so fast that there is no benefit to the microorganism if it catalyzes the reaction [41]. High-temperature-adapted microorganisms are therefore thought to produce enzymes with faster reaction rates [107]. Most cultivated hydrogenotrophic methanogens are mesophiles but known optimal growth temperatures for methanogens range from 15 to 98 °C (Fig. 1a). A considerable number of methanogens favor temperatures above 60 °C (Fig. 1a). The highest temperature that a methanogen was found to grow under is 122 °C (Methanopyrus kandleri) (Fig. 1b) [108]. Cultivated SSRM typically have optimum growth temperatures of 20-30 °C or 50-70 °C where sulfur reducing archaea have higher optimum growth temperatures than sulfur and sulfate reducing bacteria. The full range for optimum growth of SSRM spans 10-106 °C (Fig. 1i). The critical temperature for growth of cultivated SSRM is 113 °C (Pyrolobus fumarii) [109]. Homoacetogens typically have optimum growth temperatures between 20-30 °C (85 % of the here gathered cultivated strains; Fig. 1e). Thermophilic growth temperatures ≥60 °C have been reported for eight strains, only (e.g. Moorella mulderi, Thermoanaerobacter kivui, Acetogenium kivui) [110-112]. Corresponding upper limits for growth are 70-72 °C (Fig. 1f) [110-112].

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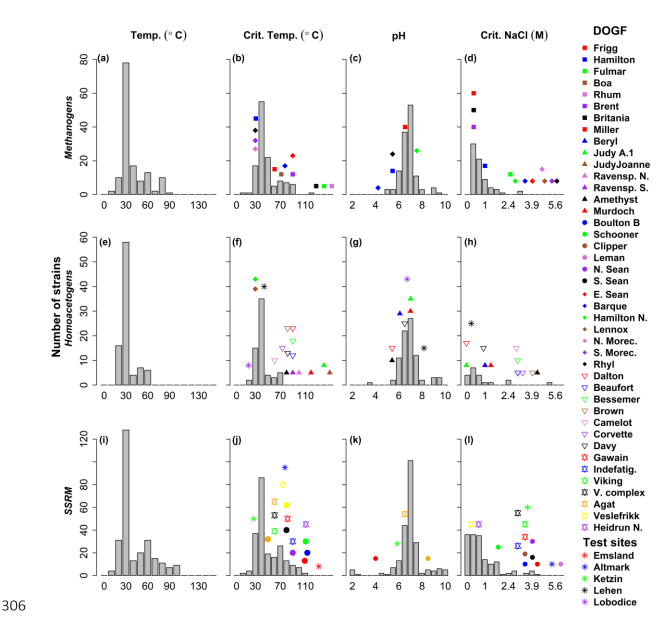


Figure. 1. Distribution of optimum growth temperature, critical growth temperature, optimum pH values and critical salinity for 123-140 methanogens (a-d), 21-91 homoacetogens (e-h) and 151-255 sulfur species reducing microorganisms (SSRM) (i-l). Distributed between the graphs for the different groups of H₂-oxidizers are the temperatures, pH values and salinities of 42 depleted oil and gas fields (DOGF) and five test sites for H₂ injection. Where ranges of a parameter were given (see Table A.4), the lower end value was plotted.

2.2.3 Salinity

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- The relevant salt concentration range for H₂ storage is 0-5 M NaCl [100], at which highly diverse prokaryote communities can be found [113]. Microorganisms are classified according to their salt tolerance: Non-halophilic microorganisms grow up to 0.2 M NaCl, slight halophile grow at 0.2–0.5 M NaCl, moderate halophile between 0.5–2.5 M NaCl, and extreme halophile
- High salt concentrations exert osmotic stress [114], requiring any microorganism living at high salt concentrations to maintain its intracellular environment at least isosmotic with the

that grow best in hypersaline media containing 2.5–5.2 M NaCl [113].

- environment [113]. Commonly, salt tolerance/requirement is enhanced at increased
- 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,
- fourteen SSRM (all mesophiles) have upper salinity limits for growth of ≥ 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).
- The salt tolerance of homoacetogens is poorly investigated. The majority of homoacetogens
- have low optimum salinities of >0-0.4 M NaCl. However, a few strains, i.e. Natroincola
- 337 histidinovorans, Sporohalobacter lortii and Halanaaerobium praevalens, grow optimally at
- >1.4 M NaCl and will tolerate salinities up to 2.6-5.1 M (Fig. 1h) [121-123].

2.2.4 Brine complexity

Natural brines contain dissolved ions whose interaction is extremely complex and may cause physicochemical stressors to brine habitability such as low water activity (a_w), high ionic strength, chaotropy (ability to disrupt the network of H₂-bonds between water molecules) or a combination of these [12]. Most bacteria grow well at an a_w around 0.98 (the approx. a_w for sea water) but relatively few species can grow at a_w of 0.96 or lower [124]. Halophilic microorganisms, including halophilic methanogens are one exception; several can grow at a_w as low as 0.75 [125] in [124]; [126]. Steinle et al. [127] challenged these limits by detecting SSR in a nearly MgCl₂ saturated brine with a_w of ~0.4.

There are indications of a more important role of chaotropy over a_w in limiting microbial life [126]. Chaotropic agents include MgCl₂, CaCl₂, FeCl₃, KI, LiBr, LiCl while examples of kosmotropic agents are NaCl, KCl, Na₂SO₄, MgSO₄, K₂SO₄, FeSO₄ [128]. As such one may speculate that most subsurface brines due to their dominance of NaCl and richness in sulfate are kosmotropic and albeit also stress-inducing, more permissive of microbial growth [12, 126]. Meanwhile, the interactions between chao- or kosmotropic agents, a_w and other physicochemical properties of brines may be very complex and hitherto not understood [12]. The further elucidation of this topic is subject to more research and beyond the scope of this paper.

2.2.5 pH

The brine pH may affect the growth of microorganisms via 1) a direct effect on the growth metabolism, and 2) an effect on the redox reaction. With respect to the former, most methanogens, homoacetogens and SSRM are adopted to a pH of 6.5-7.5 (Fig. 1c, g, k). Most methanogens and SSRM cannot grow outside the pH range 4–9.5 [29, 129, 130] (Fig. 1c, k). 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].

At the other end of the spectrum, ten known methanogens can endure acidic conditions of pH

4, e.g. the Methanoregula boonei, the Methanothermococcus okinawensis, the Methanosarcina

spelaei and the Methanocaldococcus bathoardescens [132-135].

367 Eighteen known SSRM are adapted to highly alkaline environment >pH 10, e.g. the

Desulfonatronovibrio hydrogenovorans, the Desulforispira natronophila and the

Desulfovibrio vietnamensis [136-138]. Thirteen known SSRM grow down to a pH of 4. Nine

known SSRM, all of them sulfur reducers, grow down to a pH of 1, e.g. the *Thiobacillus caldus*,

the Sulfolobus acidocaldarius, the Acidianus infernus and brierleyi, and the Stygiolobus

372 *azoricus* [139-142].

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373 Seven known homoacetogenic strains have high critical pH values up to 10.0-10.7, e.g.

Clostridium ultunense, Natrionella acetigena, Fuchsiella alkaliacetigena and ferrireducens,

Natronoincola histidinovorans, Peptostreptococcus productus B-52 and Moorella sp HUC22-

376 I [121, 122, 143-146]. The Clostridium drakai, ljundahlii, scatologenes, coccoides and

termoautrophicum are the most acidophilic known strains; they can tolerate pH as low as 3.6-

378 4.5 [147-151].

2.2.6 Pressure

Pressure ranges for H₂ storage of 5-20 MPa [99] or 1-50 MPa [100] have been reported. Life at high pressure requires homeostatic changes [103]. The high pressures encountered in pore spaces in the crust are generally less inhibitory to microbial cellular activity than the high temperatures, partly because of the relatively high osmotic pressure of cytoplasm [102], in particular in thermophiles and hyperthermophiles [41]. DNA synthesis and protein synthesis are among the most pressure-sensitive cellular processes [103, 152]. Protection against pressure includes biofilm [18] or spore formation [103].

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

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A temperature dependence of the pressure response was reported for the SSRM *Desulfovibrio*indonesiensis which has similar growth rates at high and ambient pressure 45 °C but reduces

its growth rate at 20 °C and 30 MPa relative to at 0.1 MPa [156]. Elevated pressure may

increase the maximum growth temperature by 2-12 °C relative to lower pressure (0.1-3 MPa)

[105, 108, 157].

2.2.7 Inhibitors

makinawite [46].

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- Exposure to hydrogen sulfide, H₂S, and its bisulfide ion, HS⁻, causes damage to microbial proteins and coenzymes [91, 158]. It remains unclear whether H₂S or HS⁻ is responsible for the toxicity effect but there is general consensus that H₂S can penetrate the microbial cell membrane more easily than HS⁻ [158]. Hydrogen sulfide dissociates with a pK₁ of 6.99 at 10 MPa and 25 °C to form >99 % HS⁻ at pH 8.5 [159].
- Growth of SSRM and methanogens is adversely affected at concentrations of H₂S >3.8-4.0 mM [160-162]. At 5.0-6.3 mM H₂S growth is completely inhibited for SSRM [160, 162], without however stopping all metabolic activity [160]. For methanogens and homoacetogens 3.8-7.5 mM H₂S and total sulfide concentrations of 3.3 mM, respectively, stop the growth [158, 162]. In systems with circumneutral pH and ferric ion concentrations above 1 mM, the concentrations of H₂S are predicted to be kept below toxic levels due to its precipitation in
- Carbon dioxide pressure above 1 bar can be toxic for microorganisms as shown for the SSRM

 Desulfotomaculum geothermicum and the methanogen Methanothermococcus

 thermolithotrophicus [163]. For many anaerobes like methanogens and homoacetogens,

 oxygen is toxic too [64, 102].
- Nitrate inhibits homoacetogenesis [164], and ammonium [165] and sulfate inhibit methanogenesis (reviewed in [166]), with minimum inhibitory concentrations varying

depending on the environment [162, 165]. For instance, sulfate concentrations as low as 2*10⁻⁴ M were shown to inhibit methanogenesis for 10 hours in lake sediments, possibly by competition with SSRM for available H₂ and C-substrate [161] (see section 2.3.9). Under H₂ storage conditions however, sulfate is likely not to affect methanogenesis, because sulfate inhibition was shown to be reversed by addition of H₂ [161]. For a discussion of an inhibitory effect of H₂, see section 2.5.

2.2.8 Summary of environmental growth constraints

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

Figure 2 shows the critical temperature versus critical salinity for 269 cultivated strains and reveals that salt tolerances up to 1-1.7 M are widely distributed over the entire temperature range while salt tolerances >1.7 M are mainly found at a critical temperature tolerances of 40-50 °C. Hence, from the point of view of minimizing microbial impacts on H₂ storage, sites with temperatures >50 °C and salinities >1.7 M are preferred.

Table 2. Reservoir conditions for selected depleted, or soon to be depleted oil and gas fields. Except where otherwise indicated, the data are from [168]. Fereference [167]. The salinity was calculated from the chloride concentration and the concentrations of dissolved N₂ was estimated from the mol percentage in the gas phase, neglecting any effect of salinity. NA= not analyzed. See Table A.4 for extended data.

Field name	Area (Km²)	P (MPa)	Temp (°C)	Salinity (M)	pН	HCO ₃ - (mM)	N ₂ (mM)	SO ₄ ² - (mM)	K ⁺ (mM)	Ca ⁺² (mM)	Mg ⁺² (mM)	P (mM)	Na ⁺ (mM)	Cl ⁻ (mM)	Fe ⁺² (mM)	Organic acids (mM)
Frigg	100	19.5	61	0.07-0.53	6.5-	16.3	0.4	NA	26.3-	0.4-2.0	1.9-7.1	NA	75.2-	58.7-	0.04-	NA
88					7.4				31.2				534.8	490.3	0.27	
Hamilton	15	9.6	30	1.59-4.18	5.8	4.8	2.1	0.6-	8.4-	72.8-	19.5-	0.012-	1354.8-	1453.3-	4.03	NA
								7.4	29.7	720.0	37.6	0.028	2210.9	3700.7		
Barque	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
Hamilton	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
North																
Miller	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
Beryl	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
Judy	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
(Andrew 1)																
Amethyst	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
Rhyl	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
Dalton	NA	28.8	91	0.26	5	0.9	1.0	1.8		15.6	5.5	NA	189.1	237.0	0.00	NA
Davy	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
Veslefrikk [®]	NA	29.8-	67-114	0.29-0.72	6.5	8.4-	NA	0.1-	NA	NA	NA	NA	298.0-	281.0-	NA	2.2-8.1
		35.0				17.2		0.15					666.0	745.0		
Average						7.9	1.1	5.2	44.7	166.8	42.2	0.113	1473.7	1857.4	0.967	3.3

Growth of all investigated microbial groups occurs up to 72 °C (Fig. 1). Above 72 °C, known homoacetogens will not grow, and at 80-94 °C sulfate reducers cease to grow. Thirty-six cultivated SSRM and eleven methanogens have optimum growth temperature of ≥80 °C (Fig. 1a and g) and will still grow, albeit at reduced rate, beyond their optimum temperatures. The maximum growth temperature for known methanogens and sulfur reducers is 122 °C and 113 °C, respectively. The upper salinity limit that allows growth of all the major groups of investigated H₂-oxidizing microorganisms is 3 M NaCl. The upper pH limit is 9.5 and the upper pressure limit for most mesophiles is 30-50 MPa.

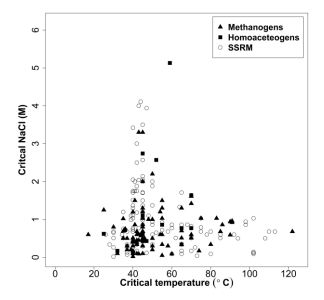


Figure. 2. Critical temperature (without salinity stress) versus critical salinity (without temperature stress) for methanogens, homoacetogens and SSRM.

2.3 Microbial growth regulation by competition and syntrophy

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

gene sequences) and culturing (e.g. metabolites, radiotracer) techniques. Few habitats have been identified in which homoacetogens compete with other H₂-consumers (culturing studies) [169, 171]. Exceptions include a low-temperature and low-salinity petroleum reservoir where homoacetogens dominated over methanogens and SSRM (molecular study) [62], a granite groundwater at 400 m depth where cell numbers of methanogens and homoacetogens were balanced (molecular study) [83], and subsurface marine sediments where mixotrophic homoacetogenesis outperformed methanogenesis (culturing study) [82].

Kinetic advantages of SSRM and methanogens (i.e. a higher affinity for H₂, expressed as a low Michaelis-Menten constant, K_M , or Monod half saturation constant, K_S (H₂ concentration at which growth rate reaches half maximum growth rate), and a higher maximum

low Michaelis-Menten constant, K_M , or Monod half saturation constant, K_S (H₂ concentration at which growth rate reaches half maximum growth rate), and a higher maximum growth/reaction rate, V_{max} or μ_{max} for Michaelis-Menten kinetics and Monod kinetics, respectively) were proposed as the underlying cause for the few examples of the poor competitiveness of homoacetogens [172]. Very limited information on the H₂ consumption kinetics of homoacetogenic bacteria is available in literature [173]. The available data show that μ_{max} differs by one order of magnitude between strains (0.02-0.5 h⁻¹) [4, 173]. This may or may not be lower than the μ_{max} for SSRM 0.057-5.5 h⁻¹ [4, 40, 174] and methanogens 0.032-1.4 h⁻¹ [40, 174]. Krumholz et al. [172] showed that homoacetogens were not able to compete effectively for H₂ in the presence of SSRM in a subsurface sandstone ecosystem at 30 °C regardless of pH₂, and despite significant homoacetogenesis at excess H₂. Findings by Berta et al. [4] for a groundwater sediment held under excess pH₂ and 20 °C contrasts this as homoacetogenesis rates were up to 21 times higher than SSR.

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

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

As for the competitiveness of methanogens and SSRM, the H₂ thresholds of methanogens may be comparable (1-15 nM) or higher (>15-95) than for sulfate reducers and significantly lower than for sulfur reducers («2500 nM; Table 1), indicating an advantage of sulfate reducers over methanogens and sulfur reducers in most non-engineered, low *p*H₂ environments. In line with this, Lackner [178] recently reviewed that sulfate reducers outcompete methanogens for H₂ in most studies. However, at excess H₂, methanogens and sulfate reducers would be expected to process equal shares of the in situ H₂ pool [174]. Also, since concentrations of sulfate are much lower than bicarbonate in non-marine natural environments [40] (Table 2), the growth of sulfate reducers at excess H₂ will be limited by availability of their electron acceptor, making it possible for methanogens to compete [40]. As a general rule pH values below 7 favor the growth of methanogens over sulfate reducers [130]. Above pH 7.5, sulfate reducers grow faster than methanogens and would be expected to outcompete them [130].

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

and the homoacetogens *Acetobacterium* in petroleum and subsurface CO₂ reservoirs [62, 180], and the presence of H₂-producing heterotrophs along with methanogens in petroleum reservoirs where the latter rely on H₂-transfer by the former [181].

2.4 Microbial ecology in natural gas and petroleum reservoirs

Recent years have seen a considerable effort in describing deep subsurface microbial communities, including those from gas and petroleum reservoirs. Isolated hydrogenotrophic microbes from these habitats are from the SSRM families Archaeoglobaceae [182], Desulfomicrobiaceae, Desulfobulbaceae, Peptococcaceae, Desulfobacteracceae, Desulfovibrionaceae, Desulforobacteriaceae, Sulfurospirillaceae, Rhodobacteraceae, Ectothiorhodospiraceae, Hydrogenothermaceae [27, 56. 97. 98. 183-1871, Eubacteriaceae and Sporomusaceae families which host homoacetogenic strains [97, 186, 188], families Methanosarcinaceae, Methanobacteriaceae, and the methanogen Methanomicrobiaceae, Methanopyraceae, Methanococcaceae, Methanocalculaceae and Methanosaetaceae [98, 115, 186] in addition to uncultured microbial taxa [56, 184, 185, 189]. Our collection of hydrogenotrophs (Fig. 1) lists many examples of the above microbial families, including the strain that holds the highest critical temperature for a methanogen, Methanopyrus kandleri. Sulphur reducing families that define the upper temperature limits for SSRM like Thermoproteaceae and Pyrodictiaceae were not reported. The cause for their absence may be a predominance of mesophilic and thermophilic sites but may also reflect a generally stronger growth of sulfate reducers over sulfur reducers in oil and gas reservoirs. Ranchou-Peyrouse et al. [98] showed that the microbial community in 35 out of 36 subsurface wells from seven natural gas storage sites was dominated by sulfate reducers.

2.5 Effect of high hydrogen concentrations on the microbial metabolism and community

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A range of studies investigated the metabolism of methanogens at excess H2 and ambient pressure, with unambiguous results. Conrad et al. [190] demonstrated that excess H₂ stimulated methanogenesis and growth rates in a paddy soil (species not specified). Opposed to this, results by Topcuoglu et al. [181] and Stewart et al. [191] suggest an inhibitory effect of high partial pressures of H₂, pH₂, expressed as a ~10-fold drop in the growth yield (cells per mole CH₄) of Methanocaldococcus jannaschii and a slight drop of ~0.1-0.7 h⁻¹ in the growth rate. Similar observations were made for Methanothermobacter thermoautotrophicus [192]. However, within the excess H₂ experiment, higher H₂ concentrations stimulated growth [181], suggesting a complex influence of pH_2 . Methanogens seem to express a pH_2 -dependent change in their ecological strategy, i.e. maximum growth rate vs. maximum growth yield, as a means to cope with different environmental conditions [181]. Indeed, M. jannaschii is capable of sensing subtle changes in dissolved H₂ concentration and restraining the energy-intensive growth of flagella to H₂-limiting conditions whereas at excess H₂ cells are mostly flagella devoid [193]. Only few studies investigated microbial H₂ turnover at high pH₂ of up to 1.5-24.8 MPa [4, 194, 195]. Methanogens (M. jannaschii) showed a strong inhibitory effect at high pH₂ [194]. However, the authors added CO₂ at a pressure of at least 0.2 MPa to the hydrogen gas mixture which at pCO₂ >0.1 MPa can be toxic methanogens [163]. Hence it is not clear whether H₂ or CO₂ performed the toxic action. For homoacetogens and SSRM, the H₂ consumption was shown not to change in response to different pH₂ of 0.1-3.5 MPa [4, 195], indicating neither stimulation nor toxicity at different levels of excess H₂. The comparison to limiting H₂ conditions was not made. Apart from microbial metabolism, the microbial community may also change in response to

high pH₂. Given a pertubation by H₂ injection it can be anticipated that other types of

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microorganisms, e.g. the in hydrocarbon reservoirs, common fermenters [24, 96, 98, 183] will decrease in abundance while hydrogenotrophs will increase [7], in line with the Baas Becking principle [196]. An increase in hydrogenotrophs in response to H_2 addition was recently confirmed for soils, however H_2 consumption increased in only one of the investigated soils, suggesting a pronounced influence of the indigenous microbial community [197]. Bioreactor experiments support a decrease in microbial diversity in response to high pH_2 as well [198, 199]. Puente-Sanchez et al. [200] were the first to report differences in the subsurface H_2 -consuming community in response to varying pH_2 within the Iberian Pyrite Belt. Ranchou-Peyruse et al. [98] showed that town gas storage with more than 50 % H_2 changed the microbial community from a predominantly sulfate reducing community to a dominance of methanogens, and this balance was active even decades after injection stopped, possibly via H_2 trapping in the microporous system [98]. It was suspected that all sulfate was initially used up by SSRM following increased growth of methanogens [98].

3. Evaluating the potential hydrogen consumption in DOGFs

3.1 Calculation of the microbial growth

We screened 42 DOGF in the North Sea and the Irish Sea and five H₂ storage test sites for temperature, salinity, pH and pressure data (Fig. 1, Table A.4). We discovered significant differences in the salinity for the DOGF reported by sources [201] and [168]. Because we relied on the solution compositions for the calculation of the potential microbial growth in the fields, which are available from [168], we chose to use the salinity data from the same source.

The environmental data from the DOGF and H₂ storage test sites were aligned with the constraints for growth of methanogens, homoacetogens and SSRM (Fig. 1-2) to select in which fields growth can be expected. For the few fields that fulfil the growth constraints of all

investigated microorganisms, we calculated a first-order estimate of the microbial growth using the elemental cell composition as a proxy for the nutrient requirement [202, 203] (Text A.1). Our calculations assumed that the supply of N and C are covered by diazotrophic and autotrophic growth, respectively. Requirements for trace elements were neglected in the calculation due to a lack of information on the relevant trace element contents in the reservoirs. Where a nutrient for a specific field was not available we used the average value from the fields given in Table 2. Any effect of the pH_2 on microbial growth was neglected. We assumed that cells neither die nor are removed, and that nutrients are not replenished by inflow, remineralization from decaying biomass or mineral dissolution. Simultaneous growth by different microorganisms was not considered. Percentages of nutrients in the cells (Text A.1) were converted to mass using a wet cell mass of $1.77*10^{-12}$ g for methanogens [204], $3.2-6.2*10^{-13}$ g for homoacetogens and $7.81*10^{-13}$ g for SSRM. The cell wet weight of homoacetogens was calculated by dividing the cell volume of 1.62-3.14 µm³ for the subsurface mixotrophic homoacetogen *Acetobacterium psammolithicum* [172] with an assumed bacterial density of 1*10⁻¹² g µm⁻³ [205]. The cell wet weight of SSRMs was calculated using a cell dry weight of 3.125*10⁻¹³ g for *Desulfovibrio desulfuricans* [206] and dividing this with a general bacterial dry weight to wet weight ratio of 0.4 [207]. Subsequently, the concentrations of C, H, O, Ca, K, Na, S, Mg, P and Fe in the DOGF (Table 2) were divided by the mass of the respective cell nutrients per microbial cell calculated above. This resulted in the maximum cell count within each microbial group, G, that could potentially be created based on a single nutrient, where the lowest G indicated the limiting nutrient for cell growth. For an example of those calculations, see Text A.1.

3.2 Estimation of the cell-specific hydrogen consumption

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Hydrogen may be consumed at rates of $0.7-5.0*10^5$ nM h⁻¹ for homoacetogens, $0.008-5.8*10^5$ nM h⁻¹ for methanogens and $0.005-130*10^5$ nM h⁻¹ for SSRM (Tables A.1-A.3), the latter considering sulfate concentrations in the range of $0-2.3*10^{-2}$ M in the DOGF (Table 2). In a few studies, the microbial H₂ consumption was related to growth (Tables A.1-A.3), enabling the calculation of the H₂ consumption per synthesized cell and the time for when the microbial cell count *G* would be reached (Text A.2).

3.3 Calculation of the hydrogen consumption in a hydrogen storage system

We calculated the minimum H₂ consumption for the DOGF Frigg and Hamilton by dividing the H₂ consumption per synthesized cell with the microbial cell count. The calculation of the moles of H₂ the in aquifer anticipated equal volumes of H₂ and water and used the ideal gas law and the field size, temperature and pressure data in Table 2 and Table A.3. The percentage of H₂ that was consumed as a function of growing and resting microbial cells was calculated by dividing the potential H₂ consumption with the H₂ concentration in the reservoir. Text A.3 shows our calculations for the Frigg reservoir and methanogens.

4. Results and discussion

4.1 Characterization of the likelihood for growth in 42 DOGF

Using the environmental limits constraining microbial growth on H₂, we analyzed the physicochemical parameters for 42 DOGF in the British and Norwegian North Sea and the Irish Sea and five H₂ storage test sites (Fig. 1, Table A.4). Of the 47 fields, five fields have a temperature of 122 °C or higher and may be considered sterile with respect to H₂-consuming microorganisms. Thirty-two fields have a temperature >72 °C, implying that homoacetogenesis cannot take place. Twenty fields have a temperature ≥90 °C implying that homoacetogenesis and sulfate reduction cannot take place. Fourteen DOGF have a temperature >90 °C and <122 °C and pressures of 18.2-44 MPa where (piezophile) methanogens and SSRM will grow.

Of the fifteen sites with temperatures <72 °C where all investigated groups of microorganisms will grow, only six fields (Frigg, Hamilton, Veslefrikk, Ketzin, Lehen and Lobodice) fulfill the remaining pressure and salinity requirements for growth. Two fields, Leman and Rhyl, have salinities ≥5.8 M where no significant microbial growth can be expected. This finding is supported by stable gas compositions at the similarly saline H₂-storage test sites of the H2STORE project, Emsland and Altmark (Fig. 1, Table A.4), though a low microbial population of ~10² cells ml⁻¹ was present [208]. Lennox, North Morecambe and South Morecambe with temperatures of 30-33 °C and salinities of 4.6-5.1 M, could permit the growth of the halophile homoacetogenic strain Halanaerobium praevalens, only. Hamilton North, Camelot and The V gas field complex with salinities of 2.9-5.0 M may permit the growth of sulfate reducers and *Halanaerobium praevalens*. The Viking field has temperature of 65-80 °C and a salinity of 3.8 M and so is likely to host only mesophilic SSRM, although pressures >30 MPa that could become growth inhibiting. The H₂-storage test site Ketzin has similar salinity to the Viking field but a lower pressure (4.0 M NaCl, 35 °C, 6 MPa). Here SSRM were suspected to cause a 2-4 % decrease in H₂ and a reduction in the concentration of sulfate from 22 to 8*10⁻³ M [208].

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

Our first order approach for calculating microbial growth in the Veslefrikk reservoir yielded a maximum 1*10⁸ methanogenic cells mL⁻¹, 2*10⁸ SSRM cells mL⁻¹ or 5*10⁸ homoacetogenic cells mL⁻¹. The Frigg reservoir a maximum of 1*10⁸ methanogenic cells mL⁻¹, 1*10⁸ SSRM cells mL⁻¹ or 2*10⁸ homoacetogenic cells mL⁻¹. The Hamilton reservoir could host a maximum of 1*10⁷ methanogenic cells mL⁻¹, 2*10⁷ SRCM cells mL⁻¹ or 6*10⁷ homoacetogenic cells mL⁻¹. These cell counts describe a maximum cell growth for each hydrogenotrophic group because simultaneous growth of hydrogenotrophs was not considered. The higher growth of homoacetogens over SSRM and methanogens results from a lower wet cell mass that causes a

lower nutrient demand per cell (see Text A.1). Our calculations are in line with total cell concentrations of 10⁵-10¹⁵ cells/ mL⁻¹ in oil reservoirs [209], and equal to or up to four order of magnitudes higher than cell counts from gas reservoirs (0.001-1.2*10⁷ cells mL⁻¹)[51, 97, 185]. The range of the literature data reflects that our simple methodology to calculate microbial numbers may serve as a first approximation to estimating microbial growth in the subsurface, only.

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

4.3 Hydrogen consumption in three low-temperature and low-salinity DOGF

The H_2 consumption in the Frigg reservoir by homoacetogens constitutes <0.01- 3.2 % of the H_2 in the aquifer, <0.01- 1.3 % for methanogens and <0.01- 1.3 % for SSRM. In the Hamilton reservoir, the rates are <0.01- 2.0 %, <0.01- 2.3 % and <0.01- 0.5 % for homoacetogens, methanogens and SSRM, respectively. For actively growing cells these consumption rates may be reached after only 0.1-19.1 days, which is the time it takes for the microorganisms to grow up to their maximum cell counts, based on the dissolved nutrient concentrations. Resting cells, i.e. cells that undergo no or only very little cell division, need 2.5-3.5 months (SSRM) or up to 3.6-6.6 years (methanogens) to reach the maximum cell count and consume the given percentage H_2 .

In a real aquifer system, nutrients are likely to at least partly be replenished by decaying cells, mineral weathering and inflowing brine, and cells will continue to consume H₂ beyond the time it takes to reach the maximum cell count (maintenance). As such our H₂ consumption estimates may be regarded as minima. On the other hand, considering that, with the exception of one study (Berta et al. [4]), our calculations employ laboratory H₂ consumption rates at optimal nutrient supply and optimal physicochemical conditions (Tables A.1-A.3), the H₂ consumption in the oligotrophic subsurface is likely overpredicted. Comparing the employed laboratory H₂ consumption rates to H₂ consumption rates by SSR and methanogenesis in oil and natural gas reservoirs of $\sim 0.05-351$ nM h⁻¹ and 0-1185 nM h⁻¹, respectively (SO₄²⁻: 8.3-805*10⁻⁵ M; HCO₃⁻ : 3.5-246*10⁻⁴ M) [51, 183], shows that the field H₂ consumption by SSR is 1.5 times to eight orders of magnitude lower, and 0.7 times to 7 orders of magnitude lower for methanogenesis. Within the operation and injection wells of a natural gas reservoir, H₂ consumption rates by SSR and methanogenesis were up to 2544 and 4533 nM h⁻¹, respectively, [51], which falls within the lower range of the values reported from laboratory studies. Acknowledging the unknown but presumably low pH₂ in above experiments, and that maintenance requirements were not included in our H₂-consumption calculations, we expect the actual H₂ consumption in a H₂ storage system to lie within the higher range of our calculated values. Our lower-end results are in agreement with no H₂ consumption during storage operations of H₂ -rich town gas in Beynes, France [210]. Our upper end results are in agreement with a loss of ~3 %, presumably by methanogenesis, at the H₂ storage (SunStorage) test site in Lehen, Austria [211]. A H₂ consumption of 17 % by methanogens at the Lobodice town gas storage site over a time span of seven months [210, 212] seems exceptional in the light of our calculations and the reported SSR and methanogenesis rates from the field. With a very low salinity of 0.03 M, temperatures of 20-45 °C, a pH of 6.7 and 4 MPa pressure, Lobodice is

among the few sites which has highly favorable conditions for microbial growth considering

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all of these parameters (Table A.4). The high H₂ consumption at Lobodice highlights the importance of our site selection tool, as H₂ storage may face serious economical and technical problems if a site with growth-favoring conditions is selected.

As mentioned, Berta et al. [4] measured high H_2 consumption rates under excess H_2 and oligotrophic conditions (P< 9.7*10⁻⁷ M; SO₄²⁻² \leq 9.5*10⁻⁴ M; DOC= 2.6*10⁻⁴ M), indicating that nutrient scarcity does not imply low H_2 consumption. A comparison to the nutrient concentrations in the DOGF reveals that many of them have a higher nutrient status (P= 0.002-0.452*10⁻³ M; SO₄²⁻² up to 23.1*10⁻³ M; organic acids= 1.2-8.1*10⁻³ M, Table 2), implying that H_2 consumption in DOGF under excess H_2 conditions may be even higher than reported in [4]. The experiment by Berta et al. [4] is further highly relevant because cells were at steady state, i.e. at the predominant growth stage in nature, but still consumed vast amounts of H_2 . Indeed the H_2 consumption of cells at steady state or resting may be just as high or higher than for growing cells but growth is low or absent (Tables A.1-A.3).

4.4 Knowledge gaps and future research

More work is needed to predict the magnitude of microbial growth, H₂ consumption rates, and (not least) the mutual interaction of the microbial processes in DOGFs. The list of unknowns and uncertainties is long. To begin with are the poorly elucidated nutrient requirements of the microorganisms, especially in mixed cultures (e.g., [71]). Adding to this are the missing or incomplete datasets on the physical environment of certain reservoirs along with their gas phase and brine compositions, including chaotropy and kosmotropy characteristics. A better elucidation of the latter would allow the calculation of the dominating microbial processes via their free energies of the reaction. Combined with an analysis of the microbial community and metabolism this could give new insights into whether or not we can theoretically predict which microbial processes occur in DOGF and to which extend.

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

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

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

the microbial community structure due to the pertubation with elevated H₂. Protocols for the careful cultivation of nutrient-deprived deep subsurface cells need to be developed. More baseline research includes determinations of the critical salinities and pressure tolerances that to date are missing for many cultivated strains, as well as the study of the brine compositional effects on the microbial community and metabolism. Research employing already cultivated species can make use of the fact that the large majority of the cultivated species isolated from subsurface environments can be found in other near-surface marine and terrestrial geothermal environments [9, 183], and should employ chemostat studies that mimic the natural environment.

5. Conclusion

Here we presented a novel site selection tool for H₂ storage and demonstrated its application for 42 DOGF in the British and Norwegian North Sea and the Irish Sea and five H₂ storage test sites. Our results highlight the hard limits to the cultivable microbial life on H₂ and can –with some certainty- exclude life in several high-salinity or high-temperature, i.e. deeper reservoirs. For low-salinity and low-temperature reservoirs our calculations indicate significant microbial growth and a small but not insignificant H₂ consumption, both of which may further increase during repeated storage cycles, giving replenishment of nutrients by mineral weathering, decaying microbial cells and inflowing water. Hence, from the point of view of minimizing H₂ loss, clogging and corrosion, sites with more extreme conditions may be chosen over low-temperature and low-salinity reservoirs where the majority of microorganisms can proliferate. Yet, any storage operation will have to consider increased operational difficulties and costs with increased depth. Additional investigation on subsurface life on H₂ is encouraged to help manifest whether H₂ consumption in low-temperature aquifers is a threat to H₂ storage.

ASSOCIATED CONTENT

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Appendix. Figure A.1 shows the solubility of hydrogen as a function of temperature and pressure. Laboratories studies investigating homoacetogenesis, methanogenesis and SSR are listed in Tables A.1, A.2 and A.3, respectively. Table A.4 provides the reservoir conditions for 42 DOGF and five H₂ storage test sites. Text A.1 holds a discussion of the importance of other hydrogen oxidizing processes for hydrogen storage. A detailed calculation of the number SSRM cells that could grow based on the N content in the Frigg reservoir can be found in Text A.1. Text A.2 and A.3 hold an example calculation of the hydrogen consumption per synthesized cell and the calculation of the potential hydrogen consumption in a hydrogen storage system, respectively.

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**

The authors declare no competing financial interest.

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REFERENCES

- Kharel S, Shabani B. Hydrogen as a long-term large-scale energy storage solution to
- support renewables. Energies 2018;11(10):1-17.
- Duan HX. The public perspective of carbon capture and storage for CO₂ emission
- reductions in China. Energ Policy 2010;38(9):5281-9.
- 813 3. Beckingham LE, Winningham L. Critical knowledge gaps for understanding water-
- rock-working phase interactions for compressed energy storage in porous formations. Acs
- 815 Sustain Chem Eng 2020;8(1):2-11.
- 816 4. Berta M, Dethlefsen F, Ebert M, Schafer D, Dahmke A. Geochemical effects of
- millimolar hydrogen concentrations in groundwater: An experimental study in the context of
- subsurface hydrogen storage. Environ Sci Technol 2018;52(8):4937-49.
- Heinemann N, Booth MG, Haszeldine RS, Wilkinson M, Scafidi J, Edlmann K.
- 820 Hydrogen storage in porous geological formations onshore play opportunities in the
- Midland Valley (Scotland, UK). Int J Hydrogen Energy 2018;43(45):20861-74.
- 822 6. Zivar D, Kumar S, Foroozesh J. Underground hydrogen storage: A comprehensive
- review. Int J Hydrogen Energy 2020; in press.
- 7. Gregory SP, Barnett MJ, Field LP, Milodowski AE. Subsurface microbial hydrogen
- 825 cycling: Natural occurrence and implications for industry. Microorganisms 2019;7(53):1-27.
- 826 8. Fredrickson JK, McKinley JP, Bjornstad BN, Long PE, Ringelberg DB, White DC, et
- al. Pore-size constraints on the activity and survival of subsurface bacteria in a late cretaceous
- shale-sandstone sequence, northwestern New Mexico. Geomicrobio J 1997;14(3):183-202.
- 829 9. Colman DR, Poudel S, Stamps BW, Boyd ES, Spear JR. The deep, hot biosphere:
- Twenty-five years of retrospection. PNAS Perspective 2017;114(3):6895-903.
- 10. Lovley D, Chapelle FH. Deep subsurface microbial processes Rev Geophys
- 832 1995;33(3):365-81.

- Hallbeck L, Pedersen K. Characterization of microbial processes in deep aquifers of
- the Fennoscandian Shield. J Appl Geochem 2008;23:1796-819.
- Payler SJ, Biddle JF, Lollar BS, Fox-Powell MG, Edwards T, Ngwenya BT, et al. An
- ionic limit to life in the deep subsurface. Front Microbiol 2019;10:1-16.
- 837 13. Krumholz LR, McKinley JP, Ulrich GA, Suflita JM. Confined subsurface microbial
- communities in cretaceous rock. Nature 1997;386(6):64-6.
- 839 14. Methe BA, nelson KE, Eisen JA, Paulsen IT, Nelson W, Heidelberg JF, et al. Genome
- of Geobacter sulfurreducens: Metal reduction in subsurface environments. Science
- 841 2003;302(5652):1967-9.
- 842 15. Aüllo T, Ranchou-Peyruse A, Ollivier B, Magot M. Desulfotomaculum spp. and
- related gram-positive sulfate-reducing bacteria in deep subsurface environments. Front
- 844 Microbiol 2013;4:1-12.
- 845 16. Roh Y. Isolation and characterization of metal-reducing *Thermoanaerobacter* strains
- from deep subsurface environments of the Piceance Basin, Colorado. Appl Environ
- 847 Microbiol 2002;68(2):6013-20.
- Pedersen K. Microbial processes in radioactive waste disposal. Stockholm, Sweden;
- 849 2000.
- 850 18. Yin W, Wang Y, Liu L, He J. Biofilms: The microbial "protective clothing" in
- extreme environment. Int J Mol Sci 2019;20(3423):1-18.
- 852 19. Coombs P, Wagner D, Bateman K, Harrison H, Milodowski AE, Noy D, et al. The
- role of biofilms in subsurface transport processes Q J Eng Geol 2010;43:131-9.
- 854 20. Escudero C, Vera M, Oggerin M, Amils R. Active microbial biofilms in deep poor
- porous continental subsurface rocks. Nature Sci Rep 2018;8(1538):1-9.

- 856 21. Wu X, Pedersen K, Edlund J, Eriksson L, Aastroem M, Andersson AF, et al. Potential
- for hydrogen oxidizing chemolithoautotrophic and diazotrophic populations to iniate biofilm
- formation in oligotrophic, deep terrestrial subsurface waters. Microbiome 2017;5(37):1-13.
- 22. Daly RA, Roux S, Borton MA, Morgan DM, Johnston MD, Booker AE, et al. Viruses
- control dominant bacteria colonizing the terrestrial deep biosphere after hydraulic fracturing.
- 861 Nat Microbiol 2019;4:352-61.
- Frank YA, Kadnikov VV, Gavrilov SN, Banks D, Gerasimchuk AL, Mardanov AV,
- et al. Stable and variable parts of microbial community in Siberian deep subsurface thermal
- aquifer system revealed in a long-term monitoring study. Front Microbiol 2016;7:1-15.
- Pannekens M, Kroll L, Mueller H, Mbow FT, Meckenstock RU. Oil reservoirs, an
- exceptional habitat for microorganisms. N Biotechnol 2019;49:1-9.
- 25. Lerm S, Westphal A, Miethling-Graff R, Alawi M, Seibt A, Wolfgramm M, et al.
- Thermal effects on microbial composition and microbiologically induced corrosion and
- mineral precipitation affecting operation of a geothermal plant in a deep saline aquifer.
- 870 Extremophiles 2013;17(2):311-27.
- 26. Zgonnik V. The occurrence and geoscience of natural hydrogen: A comprehensive
- 872 review. Earth-Sci Rev 2020;203(103140):1-50.
- 873 27. Kleinitz W, Boehling E. Underground gas storage in porous media- operating
- experience with bacteria on gas quality 67th EAGE Conference and Exhibition SPE
- EUROPEC 13-16 June Madrid, Spain Society of Petroleum Engineers 2005 p. 1-6.
- 28. Loto CA. Microbiological corrosion: mechanism, control and impact—a review. Int J
- 877 Adv Manuf Tech 2017;92:4241-52.
- 878 29. Strobel G, Hagemann B, Huppertz TM, Ganzer L. Underground bio-methanation:
- 879 Concept and potential. Renew Sust Energ Rev 2020;123(109747):1-11.

- 880 30. Taylor SW, Jaffe PR. Biofilm growth and the related changes in the physical
- properties of a porous medium 1. Experimental investigation. Water Resour Res
- 882 1990;26(9):2153-9.
- Harris SH, Smith RL, Suflita JM. In situ hydrogen consumption kinetics as an
- indicator of subsurface microbial activity. Fems Microbiol Ecol 2007;60(2):220-8.
- Hoehler TM, Barker Joergensen B. Microbial life under extreme energy limitation.
- 886 Nat Rev 2013;11:83-94.
- 887 33. Maier RM, Pepper IL, Gerba CP. Environmental Microbiology Second Edition ed:
- 888 Academic Press 2009.
- 889 34. Parkes RJ, Sass H. Deep sub-surface In: Schaechter M, editor. Encyclopaedia of
- 890 Microbiology Elsevier Academic Press 2009.
- 891 35. Lovley D, Goodwin S. Hydrogen concentrations as an indicator of the predominant
- 892 terminal electron-accepting reactions in aquatic sediments. Geochim Cosmochim Acta
- 893 1988;52:2993-3003.
- Heimann A, Jakobsen R, Blodau C. Energetic constraints on H₂-dependent terminal
- 895 electron accepting processes in anoxic environments: A review of observations and model
- approaches. Environ Sci Technol 2010;44:24-33.
- 897 37. Appelo CAJ, Postma D. Geochemistry, groundwater and pollution second ed. Leiden
- 898 A.A.Balkema Publishers 2007.
- 899 38. Loeffler FE, Tiedje JM, Sanford RA. Fraction of electrons consumed in electron
- acceptor reduction and hydrogen thresholds as indicators of halorespiratory physiology. Appl
- 901 Environ Microbiol 1999;65(9):4049-56.
- 902 39. Cord-Ruwisch R, Seitz HJ, Conrad R. The capacity of hydrogenotrophic anaerobic
- bacteria to compete for traces of hydrogen depends on the redox potential of the terminal
- electron acceptor. Arch Microbiol 1988;149:350-7.

- 905 40. Karadagli F, Rittmann BE. Kinetic characterization of *Methanobacterium bryantii*
- 906 *M.o.H.* Environ Sci Technol 2005;39:4900-5.
- 907 41. Amend JP, Shock EL. Energetics of overall metabolic reactions of thermophilic and
- 908 hyperthermophilic archaea and bacteria. FEMS Microbiol Rev 2001;25:175-243.
- 909 42. Machel HG. Bacterial and thermochemical sulfate reduction in diagenetic settings:
- old and new insights. Sediment Geol 2001;140:143-75.
- 911 43. Muyzer G, Stams AJM. The ecology and biotechnology of sulphate-reducing bacteria.
- 912 Nat Rev 2008;6:441-54.
- 913 44. Findlay AJ. Microbial impact on polysulfide dynamics in the environment. Fems
- 914 Microbiol Lett 2016;363:1-12.
- 915 45. Hedderich R, Klimmek O, Kroeger A, Dirmeier R, Keller M, Stetter KO. Anaerobic
- 916 respiration with elemental sulfur and with disulfides. FEMS Microbiol Rev 1999;22:353-81.
- 917 46. Rickard D, Luther GW. Chemistry of iron sulfides. Chem Rev 2007;107:514-62.
- 918 47. Lovley D. Dissimilatory Fe(III) and Mn(IV) reduction. Microbiol Rev
- 919 1991;55(2):259-87.
- 920 48. Hernsdorf AW, Amano Y, Miyakawa K, Ise K, Suzuki Y, Ananharaman K, et al.
- 921 Potential for microbial H₂ and metal transformations associated with novel bacteria and
- archaea in deep terrestrial subsurface sediments. Nature 2017;11:1915-29.
- 923 49. Javaherdashti R. Microbially Influenced Corrosion . An Engineering Insight second
- 924 ed. Switzerland Springer; 2008.
- 925 50. Wiegel J, Hanel J, Aygen K. Chemolithoautotrophic thermophilic iron(III)-reducer.
- In: Ljungdahl LG, Adams MW, Barton LL, Ferry JG, Johnson MK, editors. Biochemistry
- and physiology of anaerobic bacteria. New York: Springer; 2003. p. 235-51.

- 928 51. Ivanova AE, Borzenkov IA, Tarasov AL, Milekhina EI, Belyaev SS. A
- 929 microbiological study of an underground gas storage in the process of gas extraction.
- 930 Microbiology 2007;76:461-8.
- 931 52. Slobodkin AI, Jeanthnon C, L'Haridon S, Nazina T, Miroshnichenko M, Bonch-
- 932 Osmoloskaya EA. Dissimilatory reduction of Fe(III) by thermophilic bacteria and archaea in
- 933 deep subsurface petroleum reservoirs of Western Siberia. Curr Microbiol 1999.
- 53. Kashefi K, Lovley D. Reduction of Fe(III), Mn(IV), and toxic metals at 100°C by
- 935 Pyrobaculum islandicum. Appl Environ Microbiol 2000;66(3):1050-6.
- 936 54. Jiang Y, Zhang B, He C, Shi J, Borthwick AGL, Huang X. Synchronous microbial
- 937 vanadium (V) reduction and denitrification in groundwater using hydrogen as the sole
- 938 electron donor. Water Resour Res 2018;141:289-96.
- 939 55. Wisotzksy F, Eckert P. Sulfat-dominierter BTEX Abbau im Grundwasser eines
- 940 ehemaligen Gaswerks-standortes. Grundwasser 1997;2:11-20.
- 941 56. Ranchou-Peyruse A, Gasc C, Guignard M, Auello T, Sequidt D, Peyret P, et al. The
- sequence capture by hybridization: a new approach for revealing the potential of mono-
- aromatic hydrocarbons bioattenuation in a deep oligotrophic aquifer. Microb Biotechnol
- 944 2017;10(2):469-79.
- 57. Zettlitzer M, Moeller F, Morozova D, Lokay P, Würdemann H. Re-establishment of
- 946 the proper injectivity of the CO₂-injection well Ktzi 201 in Ketzin, Germany. Int J Greenh
- 947 Gas Control 2010;4(6):952-9.
- 948 58. Bath A. Drilling fluid tracers Review and update of industry experience and issues
- 949 for RWMD site characterisation programme. Loughborough, UK: Nuclear Decommissioning
- Authority, Radioactive Waste Management Directorate; 2011.

- 951 59. Gittel A, Soerensen KB, Skovhus KI, Schramm A. Prokaryotic community structure
- and sulfate reducer activity in water from high-temperature oil reservoirs with and without
- 953 nitrate treatment. Appl Environ Microbiol 2009;75(22):7086-96.
- 954 60. Laturnus F, Lauritsen FR, Groen C. Chloroform in a pristine aquifer system: Toward
- an evidence of biogenic origin. Water Resour Res 2000;36(10):2999-3009.
- 956 61. Hoehener P, Werner D, Balsiger C, Pasteris G. Worldwide occurrence and fate of
- 957 chlorofluorocarbons in groundwater. Crit Rev Environ Sci Technol 2003;33(1):1-29.
- 958 62. Grabowski A, Nercessian O, Fayolle F, Blanchet D, Jeanthon C. Microbial diversity
- 959 in production waters of a low-temperature biodegraded oil reservoir. Fems Microbiol Ecol
- 960 2005;54:427-43.
- 961 63. Liu F, Conrad R. Chemolithotrophic acetogenic H₂/CO₂ utilization in Italian rice field
- 962 soil. Isme J 2011;5:1526-39.
- 963 64. Kuesel K, Drake HL. Acetogens In: Thiel JRV, editor. Encyclopedia of Geobiology:
- 964 Springer Science+Business Media B.V; 2011.
- 965 65. Schuchmann K, Mueller V. Energetics and Application of Heterotrophy in
- Acetogenic Bacteria. Appl Environ Microbiol 2016;82(14):4056-69.
- 967 66. Bengelsdorf FR, Beck MH, Erz C, Hoffmeister S, Karl MM, Riegler P, et al. Chapter
- 968 four- Bacterial anaerobic synthesis gas (syngas) and CO₂ + H₂ fermentation. Adv Appl
- 969 Microbiol 2018;103:143-221.
- 970 67. Esteve-Nunez A, Nunez C, Lovley DR. Preferential reduction of Fe(III) over fumarate
- by Geobacter sulfurreducens. J Bacteriol 2004;186(9):2897-9.
- 972 68. Eecke HCV, Akerman NH, Huber JA, Butterfield DA, Holden JF. Growth kinetics
- and energetics of a deep-sea hyperthermophilic methanogen under varying environmental
- 974 conditions. Environ Microbiol Rep 2013;5(5):665-71.

- 975 69. Freitag TE, Prosser JI. Correlation of methane production and functional gene
- 976 transcriptional activity in a peat soil. Appl Environ Microbiol 2009;75(21).
- 977 70. Usher K, Kaksonen A, Bouquet D, Cheng KY, Geste Y, Chapman PG, et al. The role
- 978 of bacterial communities and carbon dioxide on the corrosion of steel. Corros Sci
- 979 2015;98:354-65.
- 980 71. Choong YY, Norli I, Abdullah AZ, Yhaya MF. Impacts of trace element
- 981 supplementation on the performance of anaerobic digestion process: A critical review.
- 982 Bioresour Technol 2016;209:369-79.
- 983 72. Pedersen K, Karlsson F. Investigations of subterranean microorganisms. Their
- 984 importance for performance assessment of radioactive waste disposal. Swedish Nuclear Fuel
- and Waste Management Co; 1995.
- 986 73. Moench TT, Zeikus JG. Nutritional growth requirements for *Butyribacterium*
- 987 *methylotrophicum* on single carbon substrates and glucose. Curr Microbiol 1983;9:151-4.
- 988 74. Magot M, Basso O, Tardy-Jacquenod C, Caumette P. Desulfovibrio bastinii sp. nov.
- and Desulfovibrio gracilis sp. nov., moderately halophilic, sulfate reducing bacteria isolated
- 990 from deep subsurface oilfield water. Int J Syst Evol Microbiol 2004;54:1693-7.
- 991 75. Steinsbu BO, Thorseth IH, Nagakawa S, Inagaki F, Lever MA, Engelen B, et al.
- 992 Archaeoglobus sulfaticallidus sp. nov., a thermophilic and facultatively lithoautotrophic
- 993 sulfate-reducer isolated from black rust exposed to hot ridge flank crustal fluids. Int J Syst
- 994 Evol Microbiol 2010;60:2745-52.
- 995 76. Casar CP, Kruger BR, Flynn TM, Masterson AL, Momper LM, Osburn MR. Mineral-
- hosted biofilm communities in the continental deep subsurface, Deep Mine Microbial
- 997 Observatory, SD, USA. Geobiology 2020;18:508-22.
- 998 77. Samuels T, Bryce C, Landenmark H, Marie-Loudon C, Nicholson N, Stevens AH, et
- al. Microbial weathering of minerals and rocks in natural environments. In: Dontsova K,

- Balogh-Brunstad Z, Le Roux G, editors. Biogeochemical cycles: Ecological drivers and
- environmental impact: American Geophysical Union. John Wiley & Sons, Inc.; 2020.
- 1002 78. Wlodarczyk A, Lirski M, Fogtman A, Koblowska M, Bidzinski G, Matlakowska R.
- The oxidative metabolism of fossil hydrocarbons and sulfide minerals by the lithobiontic
- microbial community inhabiting deep subterrestrial kupferschiefer black shale. Front
- 1005 Microbiol 2018;9(972):1-14.
- 1006 79. Napieralski S, Buss HL, Brantley SL, Lee S, Xu H, Roden EE. Microbial
- 1007 chemolithotrophy mediates oxidative weathering of granitic bedrock. PNAS
- 1008 2019;116(52):26394-401.
- 1009 80. Huang J, Sheng X-F, Xi J, Lin-Yan H, Huang Z, Wang Q, et al. Depth-related
- changes in community structure of culturable mineral weathering bacteria and in weathering
- patterns caused by them along two contrasting soil profiles. Appl Environ Microbiol
- 1012 2014;80(1):29-42.
- 1013 81. Alber BE. Autotrophic CO₂ metabolism. In: Schaechter M, editor. Encyclopedia of
- 1014 Microbiology Elsevier Academic Press; 2009.
- 1015 82. Liu S, Suflita JM. H₂-CO₂-Dependent anaerobic O-demethylation activity in
- subsurface sediments and by an isolated bacterium. Appl Environ Microbiol
- 1017 1993;59(5):1325-31.
- 1018 83. Kotelnikova S, Pedersen K. Evidence for methanogenic archaea and homoacetogenic
- bacteria in deep granitic rock aquifers. FEMS Microbiol Rev 1997;20:339-49.
- 1020 84. Londry KL, Jahnke LL, Des Marais DJ. Stable carbon isotope rations of lipid
- biomarkers and biomass for sulfate-reducing bacteria grown with different substrates.
- 1022 Goldschmidt Conference 2001.
- 1023 85. Camacho A. Sulfur bacteria In: Likens GE, editor. Encyclopedia of inland waters 1:
- 1024 Elsevier Science 2009.

- Welsh DT, Bourges S, de Wit R, Herbert RA. Seasonal variations in nitrogen-fixation
- 1026 (acetylene reduction) and sulphate-reduction rates in the rhizosphere of Zostera noltii:
- Nitrogen fixation by sulphate-reducing bacteria Mar Biol 1996;125:619-28.
- Whitman WB, Bowen TL, Boone DR. The methanogenic bacteria. In: Balows A,
- 1029 Truper HG, Dworkin M, Harder W, Schleifer K-H, editors. The Prokaryotes New York:
- 1030 Springer-Verlag; 1992. p. 719-67.
- 1031 88. Drake HL. Acetogenesis London, United Kingdom Chapman & Hall; 2012.
- 1032 89. Kapili BJ, Barnett SE, Buckley DH, Dekas AE. Evidence for phylogenetically and
- catabolically diverse active diazotrophs in deep-sea sediment Isme J 2020;14:971-83.
- 1034 90. Herbert BN, Gilber PD, Stockdsle H, Watkinson RJ. Factors controlling the activity
- of sulphate-reducing bacteria In reservoirs during water injection. Society of Petroleum
- Engineers; 1985. Report No.: SPE-13978-MS Contract No.: SPE-13978-MS.
- 1037 91. Chen Y, Cheng JJ, Creamer KS. Inhibition of anaerobic digestion process: A review.
- 1038 Bioresour Technol 2008;99:4044-64.
- 1039 92. Okabe S. Rate and stoichiometry of sulfate reducing bacteria in suspended and
- biolfilm cultures. Montana, USA: Montana State University; 1992.
- 1041 93. Van Verseveld HW, Duine JA, editors. Proceedings of the 5th international
- symposium on microbial growth on C₁ compounds. International Symposium on microbial
- growth on C₁ compounds; 1986; University of Groningen, The Netherlands.
- 1044 94. Taylor GT, Pirt SJ. Nutrition and factors limiting the growth of a methanogenic
- bacterium (Methanobacterium thermoautotrophicum) Arch Microbiol 1977;113:17-22.
- 1046 95. van Houten RT, Yun SY, Lettinga G. Thermophilic sulphate and sulphite reduction in
- lab-scale gas-lift reactors using H₂ and CO₂ as energy and carbon source. Biotechnol Bioeng
- 1048 1997;55:807-14.

- 1049 96. Slobodkin AI, Slobodkina GB. Thermophilic prokaryotes from deep subterranean
- 1050 habitat. Mikrobiologiya 2014;83(3):255-70.
- 1051 97. Basso O, Lascourreges JF, Le Borgne F, Le Goff C, Magot M. Characterization by
- culture and molecular analysis of the microbial diversity of a deep subsurface gas storage
- 1053 aquifer. Res Microbiol 2009;160:107-9.
- 1054 98. Ranchou-Peyruse M, Auguet J-C, Maziere C, Restrepo-Ortiz CX, Guignard M,
- Dequidt D, et al. Geological gas-storage shapes deep life. J Environ Biol 2019;21(10):3953-
- 1056 64.
- 1057 99. Matos CR, Carneiro JF, Silva PP. Overview of large-scale underground energy
- storage technologies for integration of renewable energies and criteria for reservoir
- identification. J Energy Storage 2019;21:241-58.
- 1060 100. Shi Z, Jessen K, Tsotsis TT. Impacts of the subsurface storage of natural gas and
- hydrogen mixtures Int J Hydrogen Energy 2020;45(15):f8757-73.
- 1062 101. Hassanpouryouzband A, Joonaki E, Edlmann K, Heinemann N, Yang J.
- Thermodynamic and transport properties of hydrogen containing streams. Sci Data
- 1064 2020;1(1):1-14.
- 1065 102. Yen TF. Microbial enhanced oil recovery: Principle and practice Boca Raton, Florida:
- 1066 CRC Press; 1990.
- 1067 103. Salwan R, Sharma V. Physiological and biotechnological aspects of extremophiles.
- 1068 London, United Kingdom Elsevier 2020.
- 1069 104. Jaenicke R, Sterner R. Life at high temperatures. Prokaryotes 2006;2:167-209.
- 1070 105. Holden JF. Extremophiles: Hot Environments In: Schaechter M, editor.
- Encyclopaedia of Microbiology Elsevier Academic Press; 2009.

- 1072 106. Hosh S, Lepcha K, Basak A, Mahanty AK. Thermophiles and thermophilic
- hydrolases In: Salwan R, Sharma V, editors. Physiological and biotechnological aspects of
- extremophiles. London, United Kingdom Elsevier Acadamic Press; 2020.
- 1075 107. Miller JF, Nelson CM, Ludlow JM, Shah NN, Clark DS. High pressure-temperature
- bioreactor: assays of thermostable hydrogenase with fiber optics. Biotechnol Bioeng
- 1077 1989;34:1015-21.
- 1078 108. Takai K, Nakamura K, Toki T, Tsunogai U, Miyazaki M, Miyazaki J, et al. Cell
- proliferation at 122°C and isotopically heavy CH₄ production by a hyperthermophilic
- methanogen under high-pressure cultivation. PNAS 2008;105(31):10949-54.
- 1081 109. Pley U, Schipka J, Gambacorta A, Jannasch HW, Fricke H, Rachel R, et al.
- 1082 Pyrodictium abyssi sp. nov. represents a novel heterotrophic marine archaeal
- hyperthermophile growing at 110°C Syst Appl Microbiol 1991;14:245-53.
- 1084 110. Basen M, Geiger I, henke L, Mueller V. A genetic system for the thermophilic
- acetogenic bacterium *Thermoanaerobacter kivui*. Appl Environ Microbiol 2018;84(3):1-11.
- 1086 111. Leigh JA, Mayer F, Wolfe RS. Acetogenium kivui, a new thermophilic hydrogen-
- oxidizing, acetogenic bacterium. Arch Microbiol 1981;129:275-80.
- 1088 112. Balk M, Weijma J, Friedrich MW, Stams AJM. Methanol utilization by a novel
- thermophilic homoacetogenic bacterium, Moorella mulderi sp. nov., isolated from a
- 1090 bioreactor. Arch Microbiol 2003;179:315-20.
- 1091 113. Oren A. Life at high salt concentrations In: Dworkin M, Falkow S, Rosenberg E,
- 1092 Schleifer K-H, Stackebrandt E, editors. The Prokaryotes A handbook on the biology of
- bacteria Ecophysiology and Biochemistry. Volume 2. Singapore: Springer; 2006.
- 1094 114. Oren A. The bioenergetic basis for the decrease in metabolic diversity at increasing
- salt concentrations: implications for the functioning of salt lake ecosystems. Hydrobiologia
- 1096 2001;466(1-3):61-72.

- 1097 115. Ollivier B, Fardeau M-L, Cayol J-L, Magot M, Patel BKC, Prensier G, et al.
- 1098 Methanocalculus halotolerans gen. nov., sp. nov., isolated from an oil-producing well. Int J
- 1099 Syst Bacteriol 1998;48:821-8.
- 1100 116. Zhilina TN, Zavarzina DG, Kevbrin VV, Kolganova TV. Methanocalculus
- 1101 natronophilus sp. nov., a new alkaliphilic hydrogenotrophic methanogenic archaeon from a
- soda lake, and proposal of the new family Methanocalculaceae. Microbiology
- 1103 2013;82(6):698-706.
- 1104 117. Krekeler D, Sigalevich P, Teske A, Cypionka H, Cohen Y. A sulfate-reducing
- bacterium from the oxic layer of a microbial mat from Solar Lake (Sinai), Desulfovibrio
- 1106 oxyclinae sp. nov. Arch Microbiol 1997;167:369-75.
- 1107 118. Ollivier B, Hatchikian G, Guezennec J, Garcia J-L. Desulfohalobium retbaense gen.
- 1108 nov. sp. nov. a halophilic sulfate-reducing bacterium from sediments of a hypersaline lake in
- 1109 Senegal. Int J Syst Bacteriol 1991;41(1):74-81.
- 1110 119. Wood AP, Kelly DP. Isolation and characterisation of Thiobacillus halophilus sp.
- 1111 nov., a sulphur-oxidising autotrophic eubacterium from a Western Australian hypersaline
- 1112 lake. Arch Microbiol 1991;156:277-80.
- 1113 120. Jakobsen RF, Kjeldsen KU, Ingvordsen K. Desulfohalobium utahense sp. nov., a
- moderately halophilic, sulfate-reducing bacterium isolated from Great Salt Lake. Int J Syst
- 1115 Evol Microbiol 2006;56:2063-9.
- 1116 121. Rosenberg E, DeLong EF, Lory S, Stackebrankt E, Thompson F. The Prokaryotes.
- Firmicutes and Tenericutes fourth edition ed. Heidelberg: Springer Science and Business
- 1118 Media 2014.
- 1119 122. Zhilina TN, Detkova EN, Rainey FA, Osipov GA, Lysenko AM, Kostrikina NA, et al.
- Natronoincola histidinovorans gen. nov., sp. nov., a new alkaliphilic acetogenic anaerobe.
- 1121 Curr Microbiol 1998;37:177-85.

- 1122 123. Zeikus JG, Hegge PW, Thompson TE, Phelps TJ, Langworthy TA. Isolation and
- description of *Haloanaerobium praevalens gen. nov.* and *sp. nov.*, an obligately anaerobic
- halophile common to Great Salt Lake sediments. Curr Microbiol 1983;9:225-34.
- 1125 124. Motamedi M, Karland O, Pedersen K. Survival of sulfate reducing bacteria at
- different water activities in compacted bentonite. Fems Microbiol Lett 1996;141(1):83-7.
- 1127 125. Kushner DJ. Microbial life in extreme environments. London: Academic Press 1978.
- 1128 126. Hallsworth JE, Yakimov MM, Golyshin PN, Gillion JLM, D'Auria G, de Lima ALves
- 1129 F, et al. Limits of life in MgCl₂-containing environments: chaotropicity defines the window.
- 1130 Environ Microbiol 2007;9(3):801-13.
- 1131 127. Steinle L, Knittel K, Felber N, Casalino C, de Lange G, Tessarolo C, et al. Life on the
- edge: active microbial communities in the Kryos MgCl₂- brine basin at very low water
- 1133 activity. Isme J 2018;12:1414-26.
- 1134 128. Cray JA, Russell JT, Timson DJ, Singhai RS, Hallsworth JE. A universal measure of
- chaotropicity and kosmotropicity. Environ Microbiol 2013;15(1):287-96.
- 1136 129. Yuan H, Chen Y, Zhang H, Jiang S, Zhou Q, Gu G. Improved Bioproduction of
- Short-Chain Fatty Acids (SCFAs) from Excess Sludge under Alkaline Conditions. Environ
- 1138 Sci Technol 2006;40:2025-9.
- 139 130. O'Flatherty V, Mahony T, O'Kennedy R, Colleran E. Effect of pH on growth kinetics
- and sulphide toxicity of a range of methanogenic, synthrophic and sulphate-reducing bacteria.
- 1141 Process Biochem 1998;33(5):555-69.
- 1142 131. Sorokin DY, Abbas B, Merkel AY, Riipstra EIC, Sinninghe Samste JS, Sukhacheva
- 1143 MV, et al. Methanosalsum natronophilum sp. nov., and Methanocalculus alkaliphilus sp.
- 1144 nov., haloalkaliphilic methanogens from hypersaline soda lakes. Int J Syst Evol Microbiol
- 1145 2015;65:3739-45.

- 1146 132. Stewart LC, Jung J-H, Kim Y-T, Kwon S-W, Park C-S, Holden JF.
- Methanocaldococcus bathoardescens sp. nov., a hyperthermophilic methanogen isolated
- from a volcanically active deep-sea hydrothermal vent. Int J Syst Evol Microbiol
- 1149 2015;65:1280-3.
- 133. Takai K, Inoue A, Horikoshi K. Methanothermococcus okinawensis sp. nov., a
- thermophilic, methane-producing archaeon isolated from a Western Pacific deep-sea
- hydrothermal vent system. Int J Syst Evol Microbiol 2002;51:1089-95.
- 1153 134. Ganzert L, Schirmack J, Alawi M, Mangelsdorf K, Sand W, Hillebrand-Voiculescu
- A, et al. Methanosarcina spelaei sp. nov., a methanogenic archaeon isolated from a floating
- biofilm of a subsurface sulphurous lake. Int J Syst Evol Microbiol 2014;64:3478–84.
- 1156 135. Braeuer SL, Cadillo-Quiroz H, Ward RJ, Yavitt JB, Zinder SH. Methanoregula
- boonei gen. nov., sp. nov., an acidiphilic methanogen isolated from an acidic peat bog. Int J
- 1158 Syst Evol Microbiol 2011;61:45-52.
- 1159 136. Zhilina TN, Zavarzin GA, Rainey FA, Pikuta EN, Osipov GA, Kostrikina NA.
- 1160 Desulfonatronovibrio hydrogenovorans gen. nov., sp. nov., an alkaliphilic, sulfate-reducing
- 1161 bacterium. Int J Syst Bacteriol 1997;47(1):144-9.
- 137. Sorokin DY, Muyzer G. Desulfurispira natronophila gen. nov. sp. nov.: an obligately
- anaerobic dissimilatory sulfur-reducing bacterium from soda lakes. Extremophiles
- 1164 2010;14:349-55.
- 1165 138. Nga DP, Ha DTC, Hien LT, Stan-Lotter H. Desulfovibrio vietnamensis sp.nov., a
- halophilic sulfate-reducing bacterium from Vietnamese oil fields. Anaerobe 1996;2:385-92.
- 1167 139. Hallberg KB, Lindstroem EB. Characterization of *Thiobacillus caldus* sp. nov., a
- moderately thermophilic acidophile. Microbiology 1994;140:3451-1456.

- 1169 140. Segerer A, Neuner A, Kristjansson JK, Stetter KO. Acidianus infernus gen. nov. sp.
- nov. and Acidianus brierleyi comb. nov.: facultatively aerobic, extremely acidophilic
- thermophilic sulfur-metabolizing archaebacteria. Int J Syst Bacteriol 1986;36(4):559-64.
- 1172 141. Segerer A, Trincone A, Gahrtz M, Stetter KO. Stygiolobus azoricus gen. nov., sp.
- 1173 nov. represents a novel genus of anaerobic, extremely thermoacidophilic archaebacteria of
- the order Sulfolobales. Int J Syst Bacteriol 1991;41(4):495-501.
- 1175 142. Fliermans CB, Brock TD. Ecology of sulfur-oxidizing bacteria in hot acid soils. J
- 1176 Bacteriol 1972;111(2):343-50.
- 1177 143. Schnuerer A, Schink B, Svensson BH. Clostridium ultunense sp. nov., a mesophilic
- bacterium oxidizing acetate in syntrophic association with a hydrogenotrophic methanogenic
- 1179 bacterium. Int J Syst Bacteriol 1996;46(4):1145-52.
- 1180 144. Inokuma K, Nakashimada Y, Akahoshi T, Nishio N. Characterization of enzymes
- involved in the ethanol production of Moorella sp. HUC22-1. Arch Microbiol 2007;188:37-
- 1182 45.
- 1183 145. Hirano S, Masuda N. Characterization of NADP-Dependent 7,B-Hydroxysteroid
- Dehydrogenases from Peptostreptococcus productus and *Eubacterium aerofaciens*. Appl
- 1185 Environ Microbiol 1982;43(5):1057-63.
- 1186 146. Zhilina TN, Zavarzina DG, Detkova EN, Patutina EO, Kuznetsov BB. Fuchsiella
- 1187 ferrireducens sp. nov., a novel haloalkaliphilic, lithoautotrophic homoacetogen capable of
- iron reduction, and emendation of the description of the genus *Fuchsiella*. Int J Syst Evol
- 1189 Microbiol 2015;85:2432-40.
- 1190 147. Wiegel J, Braun M, Gottschalk G. Clostridium thermoautotrophicum species novum,
- a thermophile producing acetate from molecular hydrogen and carbon dioxide. Curr
- 1192 Microbiol 1981;5:255-60.

- 1193 148. Tanner RS, Miller LM, Yang D. Clostridium ljungdahlii sp. nov., an acetogenic
- species in clostridial rRNA homology group I Int J Syst Bacteriol 1993;43(2):232-6.
- 1195 149. Kaneuchi C, Benno Y, Mitsuoka T. Clostridium coccoides, a new species from the
- 1196 feaces of mice. Int J Syst Bacteriol 1976;26(4):482-6.
- 1197 150. Kuesel K, Dorsch T, Acker G, Stackebrandt E, Drake HL. Clostridium scatologenes
- strain SL1 isolated as an acetogenic bacterium from acidic sediments. Int J Syst Evol
- 1199 Microbiol 2000;50:537-46.
- 1200 151. Gößner AS, Picardal F, Tanner RS, Drake HL. Carbon metabolism of the moderatey
- acid-tolerant acetogen Clostridium drakai isolated from peat. Fems Microbiol Lett
- 1202 2008;287:236-42.
- 1203 152. Abe F, Kato C, Horikoshi K. Pressure-regulated metabolism in microorganisms.
- 1204 Trends Microbiol 1999;7(11):447-53.
- 1205 153. Vikromvarasiri N, S. B, Pisutpaisal N. Comparative performance of *Halothiobacillus*
- 1206 Neapolitanus and Paracoccus Pantotrophus in sulphur oxidation. Energy Procedia
- 1207 2015;79:885-9.
- 1208 154. Cao J, Gayet N, Zeng X, Shao Z, Jebbar M, Alain K. Pseudodesulfovibrio indicus
- gen. nov., sp. nov., a piezophilic sulfate-reducing bacterium from the Indian Ocean and
- reclassification of four species of the genus Desulfovibrio. Int J Syst Evol Microbiol
- 1211 2016;66:3904-11.
- 1212 155. Takai K, Miyazaki M, Hirayama H, Nakagawa S, Querellou J, Godfroy A. Isolation
- and physiological characterization of two novel, piezophilic, thermophilic
- chemolithoautotrophs from a deep-sea hydrothermal vent chimney. J Environ Biol
- 1215 2009;11(8):1983-97.

- 1216 156. Fichtel K, Logemann J, Fichtel J, Rullkoetter J, Cypionka H, Engelen B. Temperature
- and pressure adaptation of a sulfate reducer from the deep subsurface. Front Microbiol 2015
- 1218 6:1-13.
- 1219 157. Kurr M, Huber R, Koenig H, Jannasch HW, Fricke H, Trincone A, et al.
- 1220 Methanopyrus kandleri, gen. and sp. nov. represents a novel group of hyperthermophilic
- methanogens, growing at 110 ° C*. Arch Microbiol 1991;156:239-47.
- 1222 158. Ntagia E, Chatzigiannidou I, Williamson AJ, Arends JBA, Rabaey K.
- Homoacetogenesis and microbial community composition are shaped by pH and total sulfide
- concentration. Microb Biotechnol 2020;13(4):1026-38.
- 1225 159. Suleimenov OM, Seward TM. A spectrophotometric study of hydrogen sulphide
- ionisation in aqueous solutions to 350°C. Geochim Cosmochim Acta 1997;61(24):5187-98.
- 1227 160. Kushkeyvych I, Dordevic D, Vitezova M. Toxicity of hydrogen sulfide toward
- sulfate-reducing bacteria *Desulfovibrio piger Vib-7*. Arch Microbiol 2019;201(3):389-97.
- 1229 161. Winfrey MR, Zeikus JG. Effect of sulfate on carbon and electron flow during
- microbial methanogenesis in freshwater sediments. Appl Environ Microbiol 1977;33(2):275-
- 1231 81.
- 1232 162. Choi E, Rim JM. Competition and inhibition of sulfate reducers and methane
- producers in anaerobic treatment. Water Sci Technol 1991;23:1259-64.
- 1234 163. Dupraz S, Fabbri A, Joulian C, Dictor M-C, Battaglia-Brunet F, Menez B, et al.
- 1235 Impact of CO₂ concentration on autotrophic metabolisms and carbon fate in saline aquifers –
- 1236 A case study. Geochim Cosmochim Acta 2013;119:61-76.
- 1237 164. Froestl JM, Seifritz C, Drake HL. Effect of nitrate on the autotrophic metabolism of
- the acetogens Clostridium thermoautotrophicum and Clostrium thermoaceticum J Bacteriol
- 1239 1996;178(15):4597–603.

- 1240 165. Wang H, Zhang Y, Angelidakis I. Ammonia inhibition on hydrogen enriched
- anaerobic digestion of manure under mesophilic and thermophilic conditions. Water Resour
- 1242 Res 2016;105:314-9.
- 1243 166. Conrad R. Contribution of hydrogen to methane production and control of hydrogen
- 1244 concentrations in methanogenic soils and sediments. Fems Microbiol Ecol 1999;28:193-202.
- 1245 167. Barth T, Riis M. Interactions between organic acid anions in formation waters and
- reservoir mineral phases. Org Geochem 1992;19(4-6):455-82.
- 1247 168. Oil and gas field data from the North Sea [Internet]. Oil and Gas Authority 2020
- 1248 [cited 19.5.2020]. Available from: https://www.ogauthority.co.uk/data-centre/.
- 1249 169. Hoehler TM, Albert DB, Alperin MJ, Martens CS. Acetogenesis from CO₂ in an
- anoxic marine sediment. Limnol Oceanogr 1999;44(3):662-7.
- 1251 170. Pedersen K. Microbial life in deep granitic rock. Fems Microbiol Ecol 1997;20:399-
- 1252 414.
- 1253 171. Breznak JA. Acetogenesis from carbon dioxide in termite guts. In: H.L. D, editor.
- 1254 Acetogenesis Chapman & Hall Microbiology Series (Physiology / Ecology / Molecular
- Biology / Biotechnology). Boston, MA: Springer; 1994.
- 1256 172. Krumholz LR, Harris SH, Tay ST, Suflita JM. Characterization of two subsurface H₂-
- 1257 utilizing bacteria, Desulfomicrobium hypogeium sp. nov. and Acetobacterium
- 1258 psammolithicum sp. nov., and their ecological role. Appl Environ Microbiol
- 1259 1999;65(6):2300-6.
- 1260 173. Phillips J. Extracellular electron uptake by acetogenic bacteria: Does H₂ consumption
- favor the H₂ evolution reaction on a cathode or metallic iron? Front Microbiol 2020;10:1-13.
- 1262 174. Robinson JA, Tiedje JM. Competition between sulfate-reducing and methanogenic
- bacteria for H₂ under resting and growing conditions. Arch Microbiol 1984;137:26-32.

- 1264 175. Fu B, Jin X, Conrad R, Liu H, Liu H. Competition between chemolithotrophic
- acetogenesis and hydrogenotrophic methanogenesis for exogenous H₂/CO₂ in anaerobically
- digested sludge. Front Microbiol 2019;10:1-9.
- 1267 176. Kotsyurbenko OR, Glagolev MV, Nozhevnikova AN, Conrad R. Competition
- between homoacetogenic bacteria and methanogenic archaea for hydrogen at low
- temperature. Fems Microbiol Ecol 2001;38:153-9.
- 1270 177. Lever MA. Acetogenesis in the energy-starved deep biosphere—a paradox? Front
- 1271 Microbiol 2012;2:1-18.
- 1272 178. Lackner N, Wagner AO, Ilmer P. Effect of sulfate addition on carbon flow and
- microbial community composition during thermophilic digestion of cellulose. Appl Microbiol
- 1274 Biotechnol 2020;104:4605-15.
- 1275 179. Men Y, Feil H, VerBerkmoes NC, Shah MB, Johnson DR, Lee PKH, et al.
- Sustainable syntrophic growth of Dehalococcoides ethenogenes strain 195 with Desulfovibrio
- vulgaris Hildenborough and Methanobacterium congolense: global transcriptomic and
- proteomic analyses. Isme J 2012;6:410-2.
- 1279 180. Freedman AJE, BoonFei T, Thompson JR. Microbial potential for carbon and nutrient
- cycling in a geogenic supercritical carbon dioxide reservoir. Environ Microbiol
- 1281 2017;19:2228-45.
- 1282 181. Topcuoglu BD, Meydan C, Nguyen TB, Lang SQ, Holden JF. Growth kinetics,
- carbon isotope fractionation, and gene expression in the hyperthermophile
- 1284 Methanocaldococcus jannaschii during hydrogen-limited growth and interspecies hydrogen
- transfer. Appl Environ Microbiol 2019;85(9):1-14.
- 1286 182. Stetter KO, Huber R, Bloechl E, Kurr M, Eden RD, Fielder M, et al.
- Hyperthermophilic archaea are thriving in deep North Sea and Alaskan oil reservoirs Nature
- 1288 1993;365:743-5.

- 1289 183. Bonch-Osmoloskaya EA, Miroshnichenko ML, Lebedinsky AV, Chernyh TN, Nazina
- TN, Ivoilov VS, et al. Radioisotopic, culture-based, and oligonucleotide microchip analyses
- of thermophilic microbial communities in a continental high-temperature petroleum reservoir.
- 1292 Appl Environ Microbiol 2003;69(10):6143–51.
- 1293 184. Tian H, Gao P, Chen Z, Li Y, Li Y, Wang Y, et al. Compositions and abundances of
- sulfate-reducing and sulfur-oxidizing microorganisms in water-flooded petroleum reservoirs
- with different temperatures in China. Front Microbiol 2017;8(143):1-14.
- 1296 185. Auello T, Berlendis S, Lascourreges JF, Dessort D, Duclerc D, Saint-Laurent S, et al.
- New bio-indicators for long term natural attenuation of monoaromatic compounds in deep
- terrestrial aquifers. Front Microbiol 2016;7(122):1-16.
- 1299 186. Mori K, Tsurumaru H, Harayama S. Iron corrosion activity of anaerobic hydrogen-
- consuming microorganisms isolated from oil facilities. J Biosci Bioeng 2010;110(4):426-30.
- 1301 187. Okpala GN, Chen C, Fida T, Voordouw G. Effect of thermophilic nitrate reduction on
- sulfide production in high temperature oil reservoir samples. Front Microbiol 2017;8:1-13.
- 1303 188. Balk M, Mehboob F, Gelder A, Riipstra WIC, Sinninghe Damste JS, Stams AJM.
- 1304 (Per)chlorate reduction by an acetogenic bacterium, *Sporomusa sp.*, isolated from an
- underground gas storage. Appl Microbiol Biotechnol 2010;88:595-603.
- 1306 189. Berlendis S, Lascourreges JF, Schraauwers B, Sivadon P, Magot M. Anaerobic
- biodegradation of BTEX by original bacterial communities from an underground gas storage
- 1308 aquifer. Environ Sci Technol 2010;44:3621-8.
- 1309 190. Conrad R, Schuetz H, Babbel M. Temperature limitation of hydrogen turnover and
- methanogenesis in anoxic paddy soil. Fems Microbiol Ecol 1987;45:281-9.
- 1311 191. Stewart LC, Algar CK, Fortunato CS, Larson BI, Vallino JJ, Huber JA, et al. Fluid
- geochemistry, local hydrology, and metabolic activity define methanogen community size
- and composition in deep-sea hydrothermal vents. Isme J 2019;13(7):1711-21.

- 1314 192. Enoki M, Shinzato N, Sato H, Nakamura K, Y. K. Comparative proteomic analysis of
- 1315 Methanothermobacter themautotrophicus DH in pure culture and in co-culture with a
- butyrate-oxidizing bacterium. Plos One 2011;6(8):1-10.
- 1317 193. Mukhopadhyay B, Johnson EF, Wolfe RS. A novel pH₂ control on the expression of
- flagella in the hyperthermophilic strictly hydrogenotrophic methanarchaeaon Methanococcus
- 1319 jannaschii. PNAS 2000;97(21):11522-7.
- 1320 194. Miller JF, Shah NN, Nelson CM, Ludlow JM, Clark DS. Pressure and temperature
- effects on growth and methane production of the extreme thermophile *Methanococcus*
- 1322 *jannaschi*. Appl Environ Microbiol 1988;54(12):3039-42.
- 1323 195. Schieche D, Murty MVS, Kermode RI, Bhattacharyya D. Biohydrogenation of
- fumarate using Desulfovibrio desulfuricans: Experimental results and kinetic rate modelling.
- 1325 J Chem Technol Biotechnol 1997;70(3):316-22.
- 1326 196. Baas Becking LGM. Geobiologie of inleiding tot de milieukunde. Den Hague, the
- Netherlands: W.P. Van Stockum & Zoon; 1934.
- 1328 197. Xu Y, Teng Y, Wang XB, Li R, Christie P. Exploring bacterial community structure
- and function associated with ploychlorinated biphenyl biodegradation in two hydrogen-
- amended soils Sci Total Environ 2020;745(140839):1-12.
- 1331 198. Braga Nan L, Trably E, Santa-Catalina G, Bernet N, Delgenes J-P, Escudie R.
- Biomethanation processes: new insights on the effect of a high H₂ partial pressure on
- microbial communities. Biotechnol Biofuels 2020;13(141):1-17.
- 1334 199. Treu L, Kogias PG, de Diego-Diaz B, Campanaro S, Bassani I, Fernandez-Rodriguez
- J, et al. Two-year microbial adaptation during hydrogen-mediated biogas upgradingprocess in
- a serial reactor configuration. Bioresour Technol 2018;264(140-147).

- 200. Puente-Sánchez F, Arce-Rodríguez A, Oggerind M, García-Villadangosa M, Moreno-
- Paza M, Blanco Y, et al. Viable cyanobacteria in the deep continental subsurface. PNAS
- 1339 2018;115(42):10702-7.
- 1340 201. Gluyas JG, Hichens HM. The United Kingdom oil and gas fields commemorative
- millennium volume Gluyas JG, Hichens HM, editors: Memoirs of the Geological Society of
- 1342 London; 2003.
- 202. Zhang Y, Zhang Z, Suzuki K, Maekawa T. Uptake and mass balance of trace metals
- for methane producing bacteria. Biomass Bioenerg 2003;25:427–33.
- 1345 203. Scherer P, Lippert H, Wolff G. Composition of the major elements and trace-elements
- of 10 methanogenic bacteria determined by inductively coupled plasma emission-
- 1347 spectrometry. Biol Trace Elem Res 1983;5(3):149-63.
- 1348 204. Amid A, Mignard D, Wilkinson M. Seasonal storage of hydrogen in a depleted
- natural gas reservoir. Int J Hydrogen Energy 2016;41(12):5549-58.
- 1350 205. Kettle H, Louis P, Holtrop G, Duncan SH, Flint HJ. Modelling the emergent
- dynamics and major metabolites of the human colonic microbiota. Environ Microbiol
- 1352 2015;17(5):1615-30.
- 1353 206. Littlewood D, Postgate JR. On the osmotic behaviour of *Desulphovibrio*
- desulphuricans J Gen Microbiol 1957;17:378-89.
- 1355 207. Bratbak G, Dundas I. Bacterial dry matter content and biomass estimations. Appl
- 1356 Environ Microbiol 1984;48(4):755-7.
- 1357 208. Würdemann H, Halm H, Lerm S, Kleyböcker A. Verbund-Forschungsvorhaben
- H2STORE: Untersuchung der geohydraulischen, mineralogischen, geochemischen und
- biogenen Wechselwirkungen bei der Untertage-Speicherung von H₂ in konvertierten
- Gaslagerstätten: Teilprojekt 4- Mikrobiologie: Abschlussbericht: Berichtszeitraum:

- 1361 01.08.2012 bis 31.12.2015. Potsdam: Helmholtz-Zentrum Potsdam Deutsches
- 1362 GeoForschungsZentrum GFZ; 2016.
- 1363 209. Nazina TN, Pavlova NK, Tatarkin YV, Shestakova NM, Babich TL, Sokolova DS, et
- al. Microorganisms of the carbonate petroleum reservoir 302 of the Romashkinskoe oilfield
- and their biotechnological potential. Microbiology 2013;82(2):190-200.
- 1366 210. Stolten D, Emonts B. Hydrogen Science and Engineering: Materials, Processes,
- 1367 Systems, and Technology, 2 Volume Set: Wiley-VCH; 2016.
- 1368 211. Bauer S. Underground Sun Storage. Final Report Vienna, Austria; 2017.
- 1369 212. Smigan P, Greksak M, Kozankova J, Buzek F, Onderka V, Wolf I. Methanogenic
- bacteria as a key factor involved in changes of town gas stored in an underground reservoir.
- 1371 Fems Microbiol Ecol 1990;73(3):221-4.
- 1372 213. Fagerbakke KM, Heldal M, Norland S. Content of carbon, nitrogen, oxygen, sulfur
- and phosphorus in native aquatic and cultured bacteria. Aquat Microb Ecol 1996;10 15-27.
- 1374 214. Mitchell AC, Phillips AJ, Hiebert R, Gerlach R, Spangler LH, Cunningham AB.
- Biofilm enhanced geologic sequestration of supercritical CO₂. Int J Greenh Gas Control
- 1376 2009;3:90-9.

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