# 1 Title: Lifeless *Clostridia* stimulate methanogenesis on Fe<sup>0</sup> in an urban

# 2 lake corrosive community

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# 10 Abstract

11 Urban environments are webbed with iron-steel structures above and belowground. Underground, in nonsulfidic environments, it has been suggested that interspecies interactions cause Fe<sup>0</sup> corrosion. Particularly, 12 *Methanosarcinales* were assumed to interact syntrophically with acetogenic bacteria during Fe<sup>0</sup> corrosion. 13 14 Here we challenge this assumption and show that a community of methanogens (38% Methanosarcinales) prospers on Fe<sup>0</sup> especially after the demise of the acetogens. Acetogens were mainly represented by 15 16 Clostridium (81% of Bacteria). Methanogens were however more diverse including Methanosarcina (22% of 17 Archaea), Methanosaeta (17% of Archaea) and Methanothermobacter (22% of Archaea) as key groups. Surprisingly, acetogens started using electrons from  $Fe^0$  immediately, unchallenged by competing 18 19 methanogens. Acetogens were expected to be outcompeted by energy efficient methanogens with 20 comparatively lower H<sub>2</sub>-uptake thresholds. However, acetogens prevailed, perhaps because in contrast to 21 methanogens they contain [FeFe]-hydrogenases (encoded in the lake-Clostridia metagenome). [FeFe]-22 hydrogenases from *Clostridium* were previously shown effective at retrieving electrons from  $Fe^0$  for proton reduction. When acetogens thrived, methanogenic rates were low (25.2±8 µM/day) but increased appreciably 23 24 (62.5±5 µM/day) when acetogens collapsed. Acetate could not explain more than a third of the observed 25 methanogenic rates. The remaining methane could be explained by methanogens reducing their energy expenses while exploiting Clostridium remains such as readily fixed nitrogen and/or exuded [FeFe]-26 27 hydrogenases producing  $H_2$  on Fe<sup>0</sup>. This has implications on our understanding of viable interactions between autotrophic species retrieving electrons from  $Fe^0$  or other insoluble electron donors. 28

# 29 Implications

Corrosion damages to underground iron-structures distress both the local environment and the economy. Here we studied an urban lake community corroding  $Fe^{0}$ . The corrosive community was dominated by *Clostridium* and *Methanosarcinales*, which sequentially produced first more acetate and later methane. Here we bring evidence for an unusual interaction mechanism on  $Fe^{0}$ , showing that methanogenesis is enhanced by inactive acetogens. Alone, the *Clostridium*-acetogens discovered in this study, could retrieve electrons from  $Fe^{0}$ effectively, which makes them a good candidate for electrosynthesis of fossil-fuels' alternatives - a future biotechnological application.

# 37 INTRODUCTION

Steel infrastructure extends for billions of kilometers below ground enabling transport and storage of clean 38 39 water, chemicals, fuels, sewage, but also protection for telecommunication and electricity cables. Deep 40 underground, under anoxic, non-sulfidic conditions, steel was expected to persist unharmed for centuries (1-41 3). And yet, under such conditions, certain groups of anaerobes (methanogens and acetogens) strip electrons 42 off  $Fe^{0}$  leading to microbial induced corrosion (MIC) (4-8). Damages induced by MIC in the underground are 43 often discovered too late, leading to environmental and economic devastation. Thus, it is important to be able 44 to predict the lifespan of the material if exposed to microbial communities native to the site where steel 45 structures are located. This would lead to effective replacement strategies and recuperation of the metal prior to accidental spills that may be detrimental to the surrounding environment (1-3). 46

47 MIC in non-sulfidic environments is often linked to the presence of acetogens like Clostridium and 48 methanogens like Methanosarcinales on the surface of the corroded steel structure (4-9). It has been suggested 49 that Methanosarcinales were growing in a mutualistic relationship with the acetogens, and allegedly both 50 groups were gaining from the interaction (7, 8). This assumption was based on acetogens producing acetate, which would be then consumed by acetotrophic Methanosarcinales methanogens. Concurrently, acetogens 51 52 were expected to be favored by the removal of their metabolic product - acetate. However, apart from establishing a mutualistic interaction on Fe<sup>0</sup>, acetogens and methanogens may be interacting in two other ways: 53 54 1) by competing and 2) by establishing a succession to maximize access to electrons from  $Fe^{0}$  for reduction of 55  $CO_2$  to acetate (acetogens; reaction 1) or methane (methanogens; reaction 2).

56  $4Fe^{0} + 2CO_{2} + 4HCO_{3} + 4H^{+} \rightarrow 4FeCO_{3} + CH_{3}COOH + 2H_{2}O (\Delta G^{0}) = -388 \text{ kJ/mol}; \text{ Reaction 1})$ 

57  $4Fe^{0} + CO_{2} + 4HCO_{3} + 4H^{+} \rightarrow 4FeCO_{3} + CH_{4} + 2H_{2}O$  ( $\Delta G^{0} = -446 \text{ kJ/mol}$ ; Reaction 2)

Theoretically, under standard thermodynamic conditions, methanogens should have an advantage over 58 acetogens when provided with  $Fe^0$  as sole electron donor (Reactions 1 & 2). Especially, since methanogens, 59 unlike acetogens, are more effective at retrieving abiotic  $H_2$  (formed on Fe<sup>0</sup>) due to their low H<sub>2</sub>-uptake 60 thresholds (10, 11). Several groups of methanogens could corrode Fe<sup>0</sup> independent of acetogenic bacteria, 61 62 including species of Methanosarcina (12-14), Methanobacterium (14-16) and Methanococcus (6, 13, 14, 17). The mechanism by which methanogens corrode Fe<sup>0</sup>, has been debated and includes reports which suggest they 63 retrieve abiotic-H<sub>2</sub> off the Fe<sup>0</sup> surface (12), retrieve electrons directly using an unknown electron-uptake 64 65 mechanism (15, 18) or use extracellular enzymes, which stimulate enzymatic  $H_2$ -evolution on the Fe<sup>0</sup>-surface 66 (17). The later mechanism was especially relevant for *Methanococcus* species which harbored an unstable

genomic island encoding [NiFe]-hydrogenases and formate dehydrogenases enzymes additional to thosealready encoded by their standard genome (19).

And yet, oftentimes acetogens dominate corrosive communities, outcompeting methanogens when concentrations of H<sub>2</sub> are high and temperatures are low, presumably due to the higher kinetics ( $V_{max}$ ) of their hydrogenases (10). Moreover, unlike methanogens, acetogens contain [FeFe]-hydrogenases (20), which could retrieve electrons directly from Fe<sup>0</sup> for proton reduction to H<sub>2</sub> possibly (21-23).

73 In this study, we were interested to understand the dynamics of acetogens and methanogens in an urban area 74 where pipelines for infrastructure are buried underground. We studied corrosion of Fe<sup>0</sup> by an urban lake 75 community from nearby a construction site, on the island of Fyn, Denmark. Alike our previous findings in a 76 coastal-sea environment (24), the corroding microbial community was dominated by Clostridium and 77 Methanosarcinales. We used a combination of physiological experiments, process inhibition and whole 78 metagenomic analyses to study the interactions of acetogens and methanogens during Fe<sup>0</sup> corrosion. In contrast 79 to our previous report on a corrosive coastal community (24), the acetogens and methanogens in this lake did 80 not compete for retrieval of electrons from Fe<sup>0</sup>, instead methanogens appeared to take advantage of metabolites 81 and enzymes released by lifeless acetogens.

#### 82 **RESULTS AND DISCUSSION**

83 Methanogens, attached to corroded structures, were assumed to be indirectly involved in corrosion of  $Fe^{0}$  (7, 84 8). Here we show that methanogens from an urban lake may corrode  $Fe^{0}$  alone, as effectively as a complex 85 community of acetogens and methanogens together. However, within a complex community methanogenesis 86 was stimulated by deactivated, but abundant co-occurring *Clostridium*. We therefore propose an unusual 87 interaction, where methanogens may take advantage of acetogenic proton-reducing enzymes, which are set 88 free during the collapse of the acetogenic community.

#### 89 Corrosion under electron acceptor limited conditions

From the anoxic sediments of an urban lake near the university of Southern Denmark (**Fig. 1**), we enriched a methanogenic community for four successive transfers, strictly using Fe<sup>0</sup> as electron donor and CO<sub>2</sub> as sole electron acceptor. Under these conditions, the community was highly corrosive as confirmed by gravimetric and product accumulation analyses. Thus, the community utilized  $9.5 \pm 0.6$  mg Fe<sup>0</sup>, and induced 41% more weight loss than cell-free controls ( $6.7 \pm 0.05$  mg Fe<sup>0</sup>; n=3; p=0.01) (**Fig. 2**). Since CO<sub>2</sub> is the only electron acceptor available, two metabolisms are possible: methane production via CO<sub>2</sub>-reductive methanogenesis (reaction 3) and acetate production via acetogenesis (reaction 4). 97  $CO_2 + 8e^- + 8H^+ \rightarrow CH_4 + 2H_2O$  (reaction 3)

98  $2CO_2 + 8e^- + 8H^+ \rightarrow CH_3COOH + 2H_2O$  (reaction 4)

We determined if either of these processes, methanogenesis or acetogenesis, could be corrosive in isolation. Bacteria (acetogens) were inhibited with a cocktail of antibiotics (kanamycin and ampicillin), in order to favor only methanogens. Alone, methanogens were as corrosive  $(9.1\pm0.4 \text{ mg Fe}^0)$  as the whole community (n=3, p=0.35) (**Fig. 2**). Methanogens were inhibited with 2-bromoethanesulfonate (a coenzyme A analogue) in order to favor only acetogens. To our surprise acetogens were significantly more corrosive alone ( $10.6 \pm 0.4 \text{ mg Fe}^0$ ; n=3; p=0.04) than within a mixed community (**Fig. 2**).

105 To better understand how methanogens and acetogens corroded  $Fe^0$  we corroborated gravimetric 106 determinations with electron recoveries in metabolic products and functional metagenomics of the corrosive 107 community.

# 108 Electron recoveries exceeded estimated recoveries from abiotic H<sub>2</sub>

109 Electron recoveries revealed that this lake corrosive community does not rely on the abiotic  $H_2$  chemically generated at the Fe<sup>0</sup> surface (2e<sup>-</sup> + 2H<sup>+</sup>  $\rightarrow$  H<sub>2</sub>). As such, the community produced 3.3 times more methane (3.5 110  $\pm$  0.1 mM) than expected (1.1 $\pm$ 0.2 mM) from abiotic H<sub>2</sub> (Fig. 3). Interestingly, within the community, 111 112 methanogens started slowly (27 $\pm$ 5.7  $\mu$ M/day), not rivaling highly productive acetogens (68 $\pm$ 1.8  $\mu$ M/day) for 113 the first month (Fig. 3). At the end of the incubation, acetogenesis ceased, whereas methanogens sped up accomplishing rates of methanogenesis (62.5 $\pm$ 5.1  $\mu$ M/day), twofold above those predicted via acetoclastic 114 methanogenesis (28±7.3  $\mu$ M/day): CH<sub>3</sub>COOH  $\rightarrow$  CO<sub>2</sub> + CH<sub>4</sub> (Fig. 3). Perhaps, methanogens were non-115 competitively accessing electrons from Fe<sup>0</sup> in the absence of the acetogens, during the last month of incubation. 116 117 To further address this, we verified the effectiveness of methanogenesis after the eradication of bacteria with 118 the help of antibiotics. Bearing in mind that methanogens were as corrosive as the mixed community (Fig. 2), 119 we anticipated this would translate in a highly productive methanogenic community once bacteria were 120 rendered ineffective by antibiotics. To our surprise, methanogens unaccompanied by bacteria produced 121 significantly less methane  $(1.3\pm0.1 \text{ mM})$  than within the mixed community  $(3.5\pm0.1 \text{ mM})$  (Fig. 3). On the 122 other hand, acetogens accumulated significantly more acetate (2.5±0.05 mM), than they did within the mixed 123 community  $(2.0\pm0.03 \text{ mM})$  (Fig. 3). However, the mixed community showed overall better electron recoveries 124 than the summed electron recoveries of unaccompanied acetogens and methanogens put together (Fig. 3). We foresee three different scenarios to why the community was 15% more effective in utilizing Fe<sup>0</sup> as electron 125 126 donor than adding up the corrosive activities of solitary acetogens and solitary methanogens.

First, the collapse of the acetogens may release useful enzymes for the methanogens. During the last month of incubation, when the acetogenic population collapsed, we hypothesize that dying acetogens released enzymes (e.g. [FeFe]-hydrogenase), which boost  $H_2$ -production and consequently  $H_2$ -dependent methanogenesis. This supposition is backed by previous reports, which showed that acetogenic [FeFe]hydrogenases in isolation reduce protons to hydrogen when provided with Fe<sup>0</sup> as electron donor (21-23).

132 Secondly, the collapse of the acetogens results in methanogens co-feeding on  $Fe^{0}$  and acetate. 133 Alternatively, when acetogens collapse, acetoclastic methanogens may start utilizing the  $Fe^{0}$  directly, but can 134 only do so after they were provided with enough acetate for biomass synthesis. This supposition is backed by 135 reports that acetoclastic methanogens retrieve electrons from electrogenic bacteria directly or via conductive 136 particles, while also requiring acetate for incorporation into biomass (25, 26). Moreover, many of the strict H<sub>2</sub>-137 utilizing methanogens, require or are stimulated by acetate (27, 28), likely because it alleviates the need to 138 endogenously produce acetate for biomass synthesis.

139 Thirdly, the collapse of the acetogens may lead to fixed  $N_2$  accessible to methanogens corroding  $Fe^0$ . 140 Methanogens were more effective after the collapse of the acetogenic community, and yet they did require the 141 collapse of the acetogens to achieve the highest methanogenic rates. Some acetogens, including several 142 *Clostridium* species, are effective  $N_2$  fixers (29).  $N_2$  fixation catalyzed by nitrogenase enzymes is an energy demanding process requiring 16 ATPs for one fixed  $N_2$  (30, 31). We hypothesize that the collapse of the 143 144 acetogens leads to significant release of fixed-dinitrogen, now easily accessible to the methanogens. This 145 supposition was further supported by functional community analyses, which showed a high  $N_2$  fixation 146 capacity within the bacterial community but not the archaea (see below).

147 Thus, to better understand how acetogens and methanogens interact with each other inducing higher electron 148 recoveries as a community than alone (**Fig. 3**), we examined the functional metagenome of a community 149 residing on  $Fe^0$  for four successive transfers.

# 150 Acetogens

151 *Clostridium* species (especially 'Lachnoclostridium' *saccharolyticum*) dominated the corrosive community 152 (81.3% of all bacteria; 81.2% of all prokaryotes; **Fig. 4**) according to whole genome sequence analyses. Most 153 *Clostridium* have never been tested for autotrophic acetogenesis including '*L*.' *sachharolyticum* (32, 33). 154 Nevertheless, we could find the entire Wood Ljungdahl pathway for acetogenesis in its readily available 155 genome (NC\_014376.1). Using metagenome analyses we also reconstructed the Wood Ljungdahl pathway 156 almost in its entirety for our corrosive-*Clostridium*, which carried acetogenesis from Fe<sup>0</sup> and CO<sub>2</sub> (**Fig. 5**). 157 This acetogenic pathway has been intensively studied for biotechnological applications such as microbial

electrosynthesis, where electrodes are provided as electron donor instead of Fe<sup>0</sup> (34, 35). Our corrosive *Clostridium* was highly effective at generating acetate from Fe<sup>0</sup> at room temperature with rates (ca. 84  $\mu$ M/day; **Fig. 3**) comparable to those observed for various acetogens incubated at higher temperatures on poised electrodes (36) or Fe<sup>0</sup> (5, 37). Moreover, *Clostridium* acetogens, often associated with corrosion, have been recently reported to corrode Fe<sup>0</sup> as effectively as sulfate reducers (38).

163 In our enrichments on Fe<sup>0</sup>, *Clostridium*-acetogens were faster than the methanogens, initiating and completing 164 their activity two months before the methanogens (Fig. 2). Acetogens are faster not because the Wood-165 Ljungdahl pathway is more energy effective than methanogenesis, as acetogenesis generates less ATPs (0.3 166 ATPs/acetate; (39)), than CO<sub>2</sub>-reductive methanogenesis (0.5 ATPs/methane; (40)). Instead, these *Clostridium* 167 acetogens may be quicker at accessing electrons from Fe<sup>0</sup> because they contain [FeFe] proton-reducing hydrogenases, which are absent in methanogens (20). For example, the proton-reducing [FeFe] hydrogenases 168 169 from C. acetobutylicum corrodes Fe<sup>0</sup> effectively by drawing electrons for the reaction:  $2H^+ + 2e^- \rightarrow H_2$  (21-23). Clostridial [FeFe] hydrogenases are operative at H<sup>+</sup>-reduction compared to methanogenic [NiFe]-170 171 hydrogenases which are rather operative in the opposite direction doing  $H_2$ -oxidation, but less effective doing 172 proton reduction (20). In the environmental metagenome of these lake-*Clostridia* we did find [FeFe]-173 hydrogenases (EC. 1.12.7.2 ferredoxin hydrogenase), indicating they have the genetic potential to effectively carry proton-reduction. However, we still do not understand the exact mechanism how this lake *Clostridium* 174 establishes contact with Fe<sup>0</sup> for successful electron retrieval by the [FeFe]-hydrogenase enzymes. 175

176 One possibility is that they retrieve electrons (i) directly via membrane-bound proteins as it has been proposed 177 for *Clostridium ljungdahlii* grown on electrodes (34, 35), or (ii) indirectly by exuding exocellular enzymes 178 like hydrogenases or formate dehydrogenases for effective electron uptake for enzymatic  $H_2$  production (17, 179 21-23). However, if enzymes were exocellular they would deliver  $H_2$  non-specifically for both the *Clostridium*-180 acetogens and hydrogenotrophic methanogens, resulting in competition for enzymatically-released  $H_2$ . Then the microorganisms with highest affinity for H<sub>2</sub> would be favored. Generally, *Clostridium* species have a lower 181 182 H<sub>2</sub>-threshold (minimum H<sub>2</sub>-concentration for growth) compared to most methanogens (41). Thus, if 183 Clostridium releases enzymes extracellularly it would not benefit. We therefore propose that during the first 184 phase *Clostridium* is most likely to directly retrieve electrons possibly using membrane bound [FeFe]-185 hydrogenases rather than expelling these enzymes extracellularly to be used by their competitors - the 186 methanogens (Fig. 6a).

#### 187 Methanogens

Methanogens were highly diverse comprising both acetoclastic *Methanosarcinales* (ca. 38% of all archaea;
including 22% *Methanosarcina* and 16% *Methanosaeta*), as well as hydrogenotrophic groups – mostly

represented by *Methanothermobacter* (ca. 22% of all Archaea) (**Fig. 4**). *Methanothermobacter* species are rarely associated with corroded structures (42) and were shown to be especially corrosive when provided with a partner bacterium (43). On the other hand, *Methanosarcinales* have been often associated with corroded structures (4-9) were capable of utilizing  $Fe^0$  as electron donor in pure cultures (12-14), but assumed to be indirectly involved in corrosion feeding on acetate delivered by acetogens (7, 8). Here we showed that methanogens alone could indeed corrode  $Fe^0$  as effectively as a mixed community of acetogens and methanogens (**Fig. 2**).

During the acetogenic period (**Fig. 6**), when *Clostridium* proliferates on Fe<sup>0</sup> and produces acetate, we expected *Methanosarcinales* to feed on the acetate produced by *Clostridium*. Our data confirmed that during this period acetate explained all the methane produced. Accordingly, when acetogens collapsed, the rate of acetate consumption ( $40.1 \pm 6 \mu$ M/day) surpassed that of methanogenesis ( $25.2 \pm 8 \mu$ M/day) (**Fig. 3**) likely due to incorporation in the biomass of the methanogens. During this acetogenic period, the survival of hydrogenotrophic groups like *Methanothermobacter* may depend on retrieval of abiotic-H<sub>2</sub> as well as H<sub>2</sub>produced by acetoclastic methanogens (44, 45).

204 During the methanogenic period (Fig. 6), when acetogens collapsed and methanogenic rates rose steeply from 205 ca. 25  $\mu$ M/day to ca. 63  $\mu$ M/day, methanogens appear to take advantage of inactivated acetogens. Methanogens 206 alone did not produce methane with the same rates (max. 17 µM/day) as they did together with acetogens (Fig. 207 3). Thus, methanogens appear to require the preceding occurrence of the acetogens. We propose this is due to 208 a combination of factors provided by 'dying' Clostridia such as: (i) leaked [FeFe]-proton reducing 209 hydrogenases; (ii) available acetate for methanogenic biomass production; and (iii) readily fixed nitrogen. Our 210 proposition was supported by metagenome analyses, which showed the presence of [FeFe]-ferredoxin 211 hydrogenases in the acetogens, and an abundance of Firmicutes-nitrogenase genes (86% of the nifH, within 212 the entire corrosive community), which were far less abundant in the metagenome of methanogens (0.2% nifH 213 from Methanosarcina; 0.9% from Methanosaeta; and 1.6% in Methanomicrobia).

In conclusion, we describe here a novel form of interspecies interaction between acetogenic *Clostridia* and *Methanosarcinales*-methanogens during Fe<sup>0</sup> corrosion. We observed the interaction between acetogens and methanogens was not limited to acetate-transfer and required the collapse of the acetogens for an advantageous succession of the methanogens.

Thus, during  $Fe^0$  corrosion, methanogenesis was stimulated by lysed co-occurring *Clostridium* and we provide support for three different scenarios: (i) the collapse of the acetogens releasing enzymes that boost methanogenesis, (ii) acetate contributing to increased biomass production and (iii) the collapse of the acetogens leading to more fixed N<sub>2</sub> accessible to the methanogens corroding Fe<sup>0</sup>. Moreover, when acetogens collapse,

methanogens don't have to compete anymore for electrons from  $Fe^0$ . Thus, the interaction is a form of opportunistic scavenging with methanogens prospering due to the demise of the *Clostridium*.

# 224 MATERIALS AND METHODS

#### 225 Sample collection and enrichment culture conditions

Sediment cores were sampled during the month of July 2016 from a small lake located near a construction site 226 on the campus of the University of Southern Denmark (SDU), Odense (Fig. 1). The salinity of the lake was 227 228 0.6 psu, and gas bubbles (including methane) were continuously released to the water surface while sampling. 229 Sediment cores were sliced in the laboratory, and the depth horizon 15-20 cm was used for downstream 230 enrichments in a DSM modified 120 media (modifications: 0.6g/L NaCl, without casitone, without sodium 231 acetate, without methanol, and without  $Na_2S \times 9H_2O$ ). The enrichment cultures were prepared in 50 mL blue 232 butyl-rubber-stoppered glass vials with an anoxic headspace of a CO<sub>2</sub>:  $N_2$  gas mix (20:80, v/v). Iron granules 233 (99.98% Thermo Fisher, Germany) or iron coupons ( $3 \text{cm} \times 1 \text{cm} \times 1 \text{mm}$ ) were the only source of electrons 234 over the course of five successive transfers. All incubations were performed in triplicate.

All enrichments were transferred when methane production reached stationary phase. DNA extractions, SEM analyses, and further experiments were performed at the fourth transfer, after 2 years of enrichment on Fe<sup>0</sup>. In addition, methanogen-specific coenzyme  $F_{420}$  auto-fluorescence was monitored via routine microscopy to confirm the presence or absence of methanogens. To evaluate the solitary corrosive potential of methanogens, we blocked all bacteria with an antibiotic cocktail 200 µg/mL of kanamycin and 100 µg/mL of ampicillin as done before (46). To evaluate the solitary corrosive potential of the acetogens, we inhibited all methanogens with 2 mM 2-bromoethanesulfonate (BES) as done before (47).

# 242 Chemical analyses

Methane concentrations were analyzed on a Thermo Scientific Trace 1300 gas chromatograph system coupled to a thermal conductivity detector (TCD). The injector was operated at  $150^{\circ}$ C and the detector at  $200^{\circ}$ C with 1.0 mL/min argon as reference gas. The oven temperature was constant at 70°C. Separation was done on a TG-BOND Msieve 5A column (Thermo Scientific; 30-m length, 0.53-mm i.d., and 20-µm film thickness) with argon as carrier gas at a flow of 25 mL/min. The GC was controlled and automated by a Chromeleon software (Dionex, Version 7). On our set-up the limit of detection for H<sub>2</sub> and CH<sub>4</sub> was 5 µM.

Acetate production was measured using the Dionex ICS-1500 Ion Chromatography System (ICS-1500) equipped with the AS50 autosampler, and an IonPac AS22 column coupled to a conductivity detector (31 mA).

For separation of volatile fatty acids, we used  $4.5 \text{ mM Na}_2\text{CO}_3$  with  $1.4 \text{ mM Na}\text{HCO}_3$  as eluent. The run was isothermic at 30°C with a flow rate of 1.2mL/min. The limit of detection for acetate was 0.1 mM.

#### 253 DNA purification and metagenomic analyses

254 DNA was isolated as previously described before (24), using a combination of two commercially available 255 kits: MasterPure<sup>TM</sup> Complete DNA and RNA Purification Kit (Epicenter, Madison, Wi, USA), and the Fast 256 Prep spin MP<sub>tm</sub> kit for soil (Mobio/Quiagen, Hildesheim, Germany). DNA quality was verified on an agarose 257 gel, and DNA was quantified on a mySPEC spectrophotometer (VWR®/ Germany). Whole metagenome 258 sequencing was performed on a NovaSeq 6000 system, using an Illumina TrueSeq PCR-free approach via a 259 commercially available service (Macrogen/ Europe). Unassembled DNA sequences were merged, quality 260 checked, and annotated using the Metagenomics Rapid Annotation (MG-RAST) server (v4.03) with default 261 parameters (48). Illumina True Seq sequencing resulted in 3,723,388 high-quality reads of a total of 4,032,354 262 with an average length of 250 bp. For taxonomic analyses, the metagenomic data was compared with the 263 RefSeq (49) database available on the MG-RAST platform. Of a total of 2,664,384 OTUs, only 391 were not 264 classified in a domain. The rarefaction curve indicated that most of the prokaryotic diversity was covered in 265 our sample. To investigate genes involved in carbon and nitrogen fixation, sequencing reads were annotated 266 against the KEGG Orthology (KO) reference database. Both taxonomic and functional analyses were 267 performed with the following cutoff parameters: *e*-value of 1e–5, a minimum identity of 80%, and a maximum 268 alignment length of 15 bp.

#### 269 **Removal of corrosion crust and corrosion rates**

The corrosion crust from the iron coupons was removed with inactivated acid (10% hexamine in 2M HCl) (50). Then, the iron coupons were dried with  $N_2$  gas stream, weighted and anaerobically stored.

#### 272 Scanning electron microscopy

Fixation of cells on tron coupons was performed anaerobically by adding 2.5% glutaraldehyde in 0.1M phosphate buffer (pH 7.3) and incubating at 4°C for 12 h. The corroded coupons were then washed three times with 0.1 M phosphate buffer at 4°C for 10 min each. Dehydration was accomplished by a series of anoxic pure ethanol steps (each step 10 min; 35%, 50%, 70%, 80%, 90%, 95% and 100% v/v) (51). The coupons were chemical dried with hexamethyldisilazane under a gentle N<sub>2</sub> gas stream. Specimens were stored under N<sub>2</sub> atmosphere and analyzed within 18-24 h at the UMASS electron microcopy facility using the FEI Magellan 400 XHR-SEM with a resolution of 5kV.

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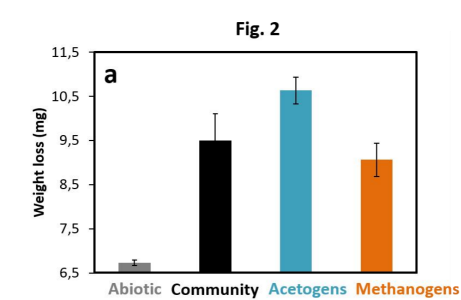
# 430 Figures

- 431 **Fig. 1.** The sampling site located in an urban construction area on the island of Fyn, Denmark (upper panel)
- 432 is a small lake near the university of Southern Denmark (lower panel). Anoxic lake sediment was
- 433 sampled with push cores during the summer of 2017, when ebullition due to increased
- 434 methanogenesis is commonly observed.





436	<b>Fig. 2.</b> Corrosion of Fe <sup>0</sup> by a microbial community (4 <sup>th</sup> successive transfer on Fe <sup>0</sup> ) as determined by weight
437	loss determination (a). Inhibition experiments revealed that acetogens were more corrosive than the
438	mixed community, whereas methanogens were as corrosive alone as within the mixed community (a).
439	(b) Visual observations of the $Fe^{0}$ -surface after exposure to a corrosive community for 5 months
440	Removal of the black crust revealed changes in surface roughness. (c) Scanning electron microscopy
441	of a mixed community after 5 months of incubation reveals cells are attached to the Fe <sup>0</sup> -surface.



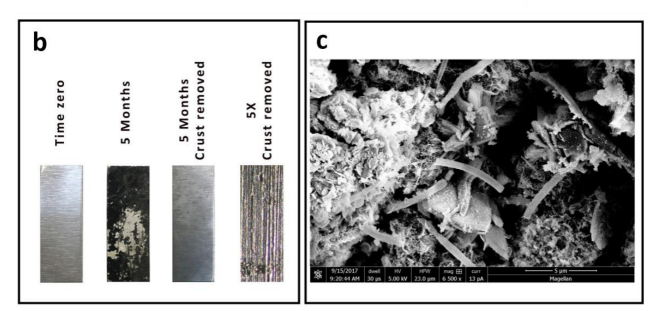


Fig. 3. Product formation using Fe<sup>0</sup> as sole electron donor. (a) An abiotic control which includes Fe0 445 exposed to the culture medium containing bicarbonate, will build up H<sub>2</sub> in the absence of cells. From 446 447 abiotic H<sub>2</sub> microorganisms could use four mols H<sub>2</sub> to produce a mol of products (acetate or methane) according to reactions 3 and 4. (b) A mixed community after four successive transfers solely with  $Fe^{0}$ 448 449 as electron donor, generated successively acetate with very high rates and later methane with 450 similarly high rates. The products generated could not be explained solely by abiotic- $H_2$  (c) 451 Acetogens were even more effective alone during the first month of incubation. The amount of acetate they produced could not be explained by abiotic-H<sub>2</sub>. (d) Methanogens alone although as 452 453 corrosive as the mixed community, they were not as effective at producing methane as they were 454 within the mixed community. (e) Total products as mM electron equivalents (eeq) produced on  $Fe^{0}$ 455 under four different conditions. The following values are considered for each product: 2 mM eeq per 456 mol H2, 8 mM eeq for each mol of methane or acetate (see reactions 3 and 4). All experiments are 457 run in triplicates (n=3). When error bars are not shown they were smaller than the symbols.

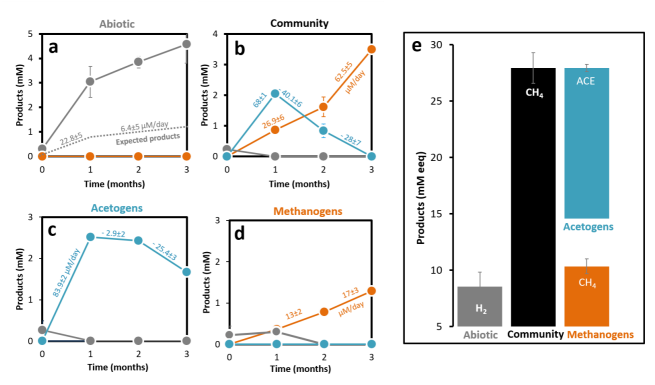


Fig. 3

458

460 Fig. 4. Whole genome sequence distribution for (a) bacterial and (b) archaeal genera representing a corrosive
 461 community enriched on Fe<sup>0</sup> after four subsequent transfers.



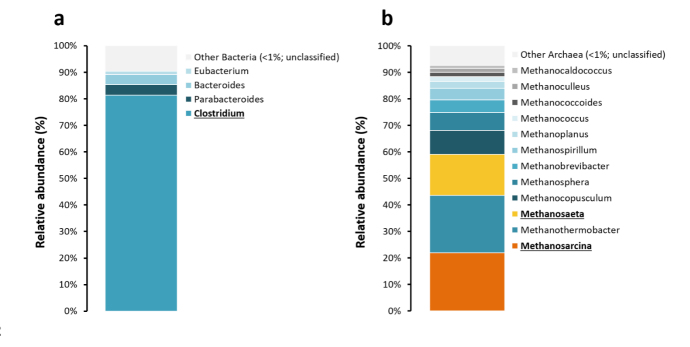
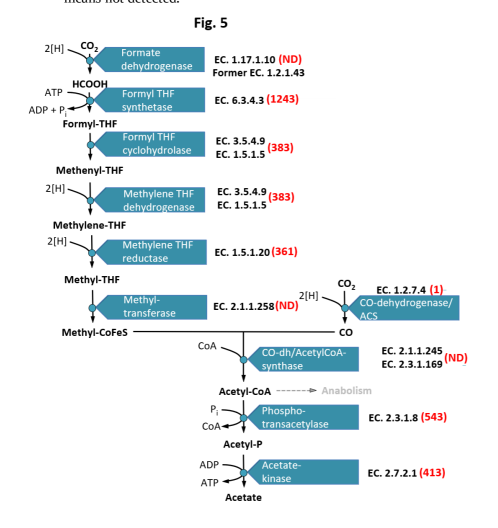


Fig. 5. Representative enzymes of the Wood-Ljungdahl pathway discovered in the metagenome of *Clostridia* from an Fe<sup>0</sup>-corrosive community enriched from an urban lake. The numbers are shown in red. ND
 means not detected.



- 467 Fig. 6. Modeled interactions between *Clostridium*-acetogens with acetoclastic *Methanosarcinales* and
- 468 hydrogenotrophic methanogens during the acetogenic phase when acetate production was highest (a)
- and (b) during the methanogenesis phase when methane production was highest.

