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## Bioelectrohydrogenesis and inhibition of methanogenic activity in microbial electrolysis cells - A review

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Bioelectrohydrogenesis and inhibition of methanogenic activity in microbial

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#### 17 ABSTRACT

Microbial electrolysis cells (MECs) are a promising technology for biological hydrogen 18 production. Compared to abiotic water electrolysis, a much lower electrical voltage ( $\sim 0.2$  V) is 19 required for hydrogen production in MECs. It is also an attractive waste treatment technology as 20 21 a variety of biodegradable substances can be used as the process feedstock. Underpinning this 22 technology is a recently discovered bioelectrochemical pathway known as "bioelectrohydrogenesis". However, little is known about the mechanism of this pathway, and 23 numerous hurdles are yet to be addressed to maximize hydrogen yield and purity. Here, we 24 review various aspects including reactor configurations, microorganisms, substrates, electrode 25 materials, and inhibitors of methanogenesis in order to improve hydrogen generation in MECs. 26 *Keywords: Microbial electrolysis cell; Hydrogen; Methane; Methanogenesis; Inhibitor;* 27 Bioelectrohydrogenesis 28 29 \*Corresponding author. jwcwong@hkbu.edu.hk (Jonathan Woon-Chung Wong)

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#### 51 **1. Introduction**

52 Hydrogen is an important chemical feedstock for many industries, such as the fertilizer industry for ammonia synthesis, and the oil industry for the conversion of crude oils into transportation 53 fuels. It is a valuable energy carrier widely used to power hydrogen fuel cells (Logan 2004). 54 55 However, most of the hydrogen is conventionally derived from fossil fuel-based resources, primarily natural gas, via chemical refinery processes (Milbrant et al., 2009). Hence, its production 56 is generally considered as environmentally unsustainable. Biological production of hydrogen (bio-57 hydrogen) is a potentially more sustainable alternative, especially when organic wastes are used 58 as the process feedstock (Hallenbeck and Benemann 2002). 59

One promising option for bio-hydrogen production is via "bioelectrohydrogenesis" which 60 61 can be accomplished using an emerging technology platform known as bioelectrochemical systems (BESs) or microbial electrochemical technologies (METs) (Liu et al., 2005; Rozendal et al., 2006). 62 BESs have been developed for a wide range of applications, including wastewater treatment, fuel 63 64 gas production (H<sub>2</sub>, CH<sub>4</sub>), nutrient recovery, chemical synthesis, desalination and bioremediation (Sleutels et al., 2012). A key feature of this technology is that it employs microorganisms to 65 catalyze redox reactions at conductive electrode surfaces. The most widely studied BESs are either 66 microbial fuel cells (MFC), which aim to produce electricity; and microbial electrolysis cells 67 (MECs), which aim to produce biogas or value added chemicals (Logan et al., 2008; Clauwaert et 68 al., 2009; Chookaew et al., 2014). During the conversion of bio-waste into H<sub>2</sub>, exoelectrogenic 69 bacteria first oxidize (degrade) organic matter and transfer the electrons to a solid electrode 70 (bioanode) (Fig.2a). The electrons then travel through an external circuit and combine with protons 71 72 at an anaerobic cathode resulting in the generation of hydrogen (Logan et al., 2008). Typically, the reducing power attainable with a bioanode is insufficient to drive the hydrogen evolution reaction 73

74 (HER) at the cathode. However, by supplementing the process with a small voltage (normally ranging from 0.2 V to 1.0 V) the cathodic HER can be facilitated in a MEC (Reaction 1&2). Since 75 a much higher voltage ( $E^0 > 1.2$  V) is required in conventional water electrolysis (Fig. 2b) 76 processes (Reaction 3&4), using MEC for bio-hydrogen production is considered as an energy-77 efficient option. Indeed, it has been reported that the energy requirement for MECs is only about 78 0.6 kWh m<sup>-3</sup> (0.2 mol H<sub>2</sub> energy/mol-H<sub>2</sub> produced), whereas in water electrolysis 4.5-5 kWh m<sup>-3</sup> 79 is required (1.5-1.7 mol H<sub>2</sub> energy/mol-H<sub>2</sub> produced) (Logan et al., 2008, Cheng and Logan 2007). 80 Microbial Electrolysis: 81  $CH_3COO^- + 4H_2O \rightarrow 2HCO_3^- + 9H^+ + 8e^ E_{anode} = -0.279 V$ 82 (1) $2H^+ + 2e^- \rightarrow H_2$  $E_{cathode} = -0.414 \text{ V}$ (2) 83  $E^0 = E_{cathode} - E_{anode} = -0.135 V$ 84 Water Electrolysis: 85  $2H_2O \rightarrow O_2 + 4H^+ + 4e^ E_{anode} = 0.82 V$ 86 (3)  $2\mathrm{H}^+ + 2\mathrm{e}^- \rightarrow \mathrm{H}_2$  $E_{cathode} = -0.414 V$ 87 (4)  $E^0 = E_{cathode} - E_{anode} = -1.22 V$ 88 Further, waste materials other than fossil fuels are used as the feedstock to drive the HER, and the 89 H<sub>2</sub> production rate can be more than  $1 \text{ m}^3\text{H}_2\text{ m}^{-3}\text{ d}^{-1}(11 \text{ mol H}_2/\text{mol glucose})$ , which is three times 90 higher than dark fermentation (Logan et al., 2008; Wang and Ren 2013). 91

92 These features collectively make MECs a promising topic for research and development93 across the world, as reflected by the expanding volume of research outputs over the past decade

94 (Fig. 1). Nonetheless, only a few review articles have discussed the use of MEC for hydrogen production and methanogenesis (Logan et al., 2008; Geelhoed et al., 2010; Kundu et al., 2013; 95 Zhou et al., 2013; Zhang and Angelidaki 2014; Kadier et al., 2014; Jafary et al., 2015; Escapa et 96 al., 2016). A notable challenge to maximize hydrogen yields from MECs is the side production of 97 methane via methanogenesis. Herein we discuss the currently available methods for the inhibition 98 of methanogenesis in MECs, and highlight the use of chemical methanogenic inhibitors with the 99 focus on their mechanisms underpinning at the enzymatic level. We suggest options of using these 100 methanogenic inhibitors to improve the purity of the produced hydrogen from MECs. We also 101 102 discuss chemical inhibition strategies for other undesirable microbes such as sulfate reducers and acetogens. 103

104 **2. Reactor configurations** 

105 *2.1. Two-chamber MECs* 

The concept of bioelectrohydrogenesis was first demonstrated with a two-chamber MEC design 106 in 2005 (Liu et al., 2005). In this conventional design, the anode and cathode chambers are 107 108 separated by an ion (proton) exchange membrane (Fig. 2a). Liu et al. (2000) observed that over 90% of the organic substrate (acetate) in the anode chamber was degraded at the end of batch mode 109 with 78% coulombic efficiency (Fig. 3). However, the overall hydrogen production efficiency was 110 only 60-73%. This is largely due to losses of the produced hydrogen in unwanted processes within 111 the MEC, such as biomass production, conversion of substrate to polymers, and methanogenesis 112 from hydrogen and acetate. To increase the hydrogen production efficiency in MECs, preventing 113 hydrogen diffusion into the anode chamber is critical. Also, the internal resistance of the MEC 114 must be minimized by reducing the distance between the electrode pair. It was reported that a 115 higher rate of hydrogen (1.6 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>) could be obtained from two-chamber MECs using saline 116

catholyte, which provided high solution conductivity and hence lowered ohmic resistance (Nam
and Logan 2011). The use of a membrane is considered an effective way to minimize hydrogen
diffusion into the anode chamber, but it introduces complexity and cost to the process.
Nonetheless, in most cases the use of two-chamber MECs only enabled hydrogen production rates
ranging from 0.01 to 6.3 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup> (Cheng and Logan 2011).

122

#### 123 2.2. Single-chamber MECs

It is accepted that hydrogen evolution occurs due to the cathodic reduction reaction in MECs. The 124 cathodic conversion efficiency (CCE) can be calculated from the ratio of e<sup>-</sup> equivalent donated to 125 hydrogen formation and e<sup>-</sup> equivalent transferred from anode to cathode (Logan et al., 2008). A 126 CCE of less than 100% could be attributed to the diffusion of hydrogen to the anode surface, or to 127 128 biological oxidation. It was inferred that hydrogen diffusion would decrease the CCE by up to 33% in two-chamber MECs (Tartakovsky et al., 2008). To maximize the overall efficiency of a MEC 129 for bioelectrohydrogenesis, the e<sup>-</sup> equivalent liberated from the anodic substrate must first be 130 131 efficiently captured by the bio-anode, and subsequently dissipated at the cathode exclusively as hydrogen gas for external collection. Indeed if the produced hydrogen gas could be rapidly 132 harvested to avoid hydrogen diffusion to the anode, the use of membrane may be omitted... 133

In fact, the use of single-chamber MECs for bioelectrohydrogenesis has been the subject of many earlier studies (Rozendal et al., 2007; Call and Logan 2007; Hu et al., 2008; Tartakovsky et al., 2009). A key attractive feature of single chamber MECs is that both the anode and cathode are housed within one chamber. This single chamber MEC system could be more compact with a lower capital cost. Further, single chamber MECs often exhibit a lower internal resistance. Such systems generally have low ohmic loss and concentration overpotential due to the nonexistence of detrimental pH gradient between the anolyte and catholyte.(Rozendal et al., 2007; Call and Logan
2007; Hu et al., 2008; Tartakovsky et al., 2009). Call et al., (2008) also found that the bio-hydrogen
production rate recorded from their single-chamber MEC was more than double (3.12 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup> at
an applied voltage of 0.8V) as compared to that obtained from a two-chamber MEC under identical
operating conditions.

- 145
- 146 *2.3. Continuous flow MECs*

Like most other waste treatment bioprocesses, MECs are often characterized for their ability to treat their feedstock in a continuous fashion (Fig. 3). When operated in continuous mode, the organic stream is continuously loaded into the MEC at a defined flow rate. Often, the liquid electrolyte within a continuous flow system is recirculated to maximize mass transfer. The hydraulic turbulence created as such may help to minimize the accumulation of stagnant hydrogen gas in the porous electrode matrix (e.g. granular graphite bed), which may help to avoid any undesirable biological oxidation (loss) of hydrogen in the reactor.

154 Both organic loading rate (OLR) and applied potential are significant parameters to determine the yield of hydrogen from continuous flow MECs, and so these parameters are often 155 selected for process optimization (Cusick et al., 2011; Escapa et al., 2012; Rader et al., 2010). For 156 instance, Escapa et al. (2012) reported a Monod-type relationship between OLR and hydrogen 157 production rate (0.3 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>) in their continuous flow domestic waste water (DWW) fed MECs. 158 They found that the increase in hydrogen production rate reached a plateau, when the OLRs of 159 DWW were above 2000 mgCOD m<sup>-3</sup> d<sup>-1</sup>. In addition, the energy consumption for pumping the 160 solution should also be accounted. The produced H<sub>2</sub> and the energy consumption for pumping may 161 162 vary depending on the pumping flow rate. For instance, Kim and Logan (2011) noted that 4 x 10<sup>-</sup>

<sup>5</sup> W was required for pumping flow rate at 0.8 ml min<sup>-1</sup>. This was however, negligible (1%) compared to the energy produced as  $H_2$  (3.8 x 10<sup>-3</sup> W) (Kim and Logan 2011).

Most of the MECs were operated with a single pair of electrodes, and only rarely multi-165 electrode pair equipped MECs were used (Rader et al., 2010). Rader et al. (2010) evaluated a multi-166 electrode MEC equipped with eight separate pairs of graphite fiber anodes and stainless steel 167 cathodes (with a working capacity of 2.5 L) for bioelectrohydrogenesis. They found that similar 168 to single pair systems, the hydrogen production rate in their multi-electrode system was also 169 directly proportional to the cathode surface area, yielding a hydrogen production rate of up to 0.53 170 m<sup>3</sup>m<sup>-3</sup>d<sup>-1</sup> (Rader et al., 2010). The first pilot scale (1000 L) bio-hydrogen producing MEC was also 171 operated with the use of multiple electrode pairs in continuous mode for about 100 days using 172 winery wastewater as the feedstock (Cusick et al., 2011). Although the gas production of the pilot 173 system could reach up to 0.19 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>, the main component of the produced gas was methane 174 (86%) suggesting that most of the cathodically produced hydrogen was consumed by the 175 methanogens. Hence, to increase hydrogen yield, an effective method to prevent methanogenesis, 176 177 and to efficiently extract the hydrogen from the cathode is required. Other factors such as enrichment of exoelectrogenic biofilms, optimization of electrolyte pH and electrode arrangements 178 are also paramount at a pilot scale level. 179

Further, to improve the hydrogen production efficiency from MEC reactors, a suitable electrode configuration should be adopted. The planar electrodes (plate type) and flow through or porous electrodes (3D type) are more common electrode types used in MEC reactors. The planar electrode (e.g. graphite plate) has advantages such as high conductivity, chemical stability, low cost and surface accessibility, and ease of placement (Zhou et al., 2011). However, it is difficult to increase the surface area of the planar electrode. Gil-Carrera et al., (2011) increased the surface 186 area of the planar electrode by sandwiching the anode between a pair of cathodes. They found that 187 the sandwich electrode only increased the current density rather than hydrogen production due to the activity of hydrogenotrophic methanogens. 3D type electrodes (e.g. graphite granules, graphite 188 189 fiber brush, and reticulated vitreous carbon) have also been shown to have increased surface area as well as large relative porosity, and good electrical conductivity. Their major limitations are 190 relatively high cost, clogging and biofouling that leads to large resistivity. Also, the main 191 disadvantage of 3D electrode configuration in the MEC is the mass transport limitation at the anode 192 matrix (Zhou et al., 2011; Escapa et al., 2016) 193

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#### **3. MEC components**

Understanding the role of various components of a MEC system is critical to optimize the
bio-hydrogen production rate. Table 1 summarizes the bio-hydrogen production performances and
characteristics of some key components such as applied potential, substrates, microorganisms, and
electrode materials in various MEC studies.

200 *3.1. Effect of anode materials* 

The anode materials for MECs must be chosen based on several features such as - i. non-201 corrosive nature with electrolytes, ii. good electrical conductivity, iii. lack of toxicity to 202 microorganisms, iv. ability to support the adherence and proliferation of microorganisms, v. high 203 surface to volume ratio, vi. feasible electron transfer from a microorganism, vii. low overpotential, 204 205 viii. ease of fabrication, and ix. low cost and scalability (Logan et al., 2008; Logan 2008). The anode materials can be broadly classified as carbon or non-carbon based materials. Typically, 206 carbon-based materials such as carbon cloth and carbon paper are more widely used in MEC 207 208 systems (Liu et al., 2005; Cheng and Logan 2007; Rozendal et al., 2007; Call and Logan 2008; Hu

et al., 2008). High current densities  $(0.05 \text{ mA cm}^{-2})$  were obtained with graphite granules (Cheng 209 and Logan 2007; Ditzig et al., 2007; Freguia et al., 2007), graphite felt (Rozendal et al., 2006; 210 Rozendal et al., 2007), and graphite brushes (Call and Logan 2008) based MECs due to the large 211 212 porosity and surface specificity of these materials (Sleutels et al., 2011). Therefore, graphite is considered a good material of choice for anodes. Using granular graphite bed (528 cm<sup>2</sup>), hydrogen 213 production has been reported to reach 3.5 mol H<sub>2</sub> per mol acetate with a coulombic efficiency (CE) 214 of 88% (Cheng and Logan 2007). Further improvement of the CE (92%) could be achieved by 215 modifying the electrode with a positively charged ammoniacal compound as reported by Call and 216 Logan (2008), who observed that with their modified anode, there was more bacterial adhesion, a 217 faster start-up period and an overall more efficient electron transfer during the MEC process. The 218 application of conducting polymers and metal nanoparticles (Fe, Au, Pd) for electrode 219 220 modification has also been attempted to improve substrate oxidation, and electron transfer 221 efficiency in MEC (Xu et al., 2012; Fan et al., 2011). The structural strength of the electrode also appeared to be important. For instance, it was found that using a more structurally robust carbon 222 223 material (activated carbon) resulted in higher  $(3\times)$  current density than with a relatively fragile material (carbon cloth) (Wang et al., 2010; Li et al., 2009). 224

225

#### 226 *3.2. Effect of cathode materials*

Cathodic hydrogen production on plain carbon materials is often associated with a high overpotential, which could limit the hydrogen production efficiency of a MEC. To address this issue, metal-based catalysts could be used for catalyzing the HER. Platinum (Pt) has been a commonly used noble-metal based catalyst in MECs (Logan et al., 2008). However, it has been suggested that about 47% of the capital cost of a MEC was associated with the use of noble-metal based cathodic 232 catalysts (Rozendal et al., 2008). Alternatively, some of the metal catalysts such as Co/FeCo (Cheng and Logan 2008), NiMo/NiW (Hu et al., 2009), Fe/Fe<sub>3</sub>C (Li et al., 2012), Nickel powder 233 (Selembo et al., 2010), Pd nanoparticles (Huang et al., 2011), MoS<sub>2</sub> (Tokash and Logan 2011; 234 Tenca et al., 2013), carbon nanotubes (MWCNT) (Wang et al., 2012), and WC (Tungsten carbide) 235 (Harnisch et al., 2009) were investigated to replace Pt catalyst. Metal alloys such as 236 NiFeMo/CoMo (Jeremiasse et al., 2011), Ni-W-P/Ni-Ce-P (Wang et al., 2011), NiFe, NiFeP and 237 NiFeCoP (Mitov et al., 2012) were also investigated for HER in MECs under neutral/mild alkaline 238 conditions. The alloy cathodes NiMo, NiFeMo or CoMo showed superior catalytic activity 239 towards HER (at pH 7) compared with cathodes coated with only Ni (Mitov et al., 2012). These 240 findings suggest that Ni-based cathodes or cathodes modified with nanomaterials are promising 241 cathode materials for HER in MECs (Mitov et al., 2012). High surface area Ni foam cathodes (128 242  $m^2 m^{-2}$  projected area) were constructed to produce high volumetric hydrogen production (50 m<sup>3</sup>) 243 m<sup>-3</sup> d<sup>-1</sup> at 1.0 V) in continuous flow MEC using an anion exchange membrane. This effect was due 244 to a lower cathode overpotential (Ni foam cathode) than for Pt-based cathode. However, the 245 246 performance of the Ni foam cathode was unstable, and often associated with an increase of overpotentials over time (Jeremiasse et al., 2010). On the other hand, stainless steel is another 247 widely used cathode material for MECs due to low cost, high current density and low cathodic 248 overpotential (Zhang et al., 2010; Ambler and Logan 2011; Munoz et al., 2010; Selembo et al., 249 2009b). A high hydrogen production rate of up to 4.9 L h<sup>-1</sup> m<sup>-2</sup> (with 0.8 V applied voltage) was 250 obtained from a MEC equipped with a stainless steel (AISI 316 L) cathode (Munoz et al., 2010). 251 Alternatively, biocathodes are increasingly being considered for HER in MECs due to low 252 cost and high operational sustainability. Though the concept of a biocathode was discovered in the 253

1960s, it has not received much attention (He and Angenent 2008). It was found that

255 microorganisms that contain hydrogenase enzyme could catalyze hydrogen production in various 256 environments (Schwartz and Friedrich 2006). In recent years, further research using biocathodes has shown that they have many advantages over chemical cathodes for HER in MECs (He and 257 Angenent 2008). For instance, it was reported that a biocathode developed from a selected 258 electrochemically active mixed microbial culture could efficiently drive HER in a cathodic half-259 cell. The biocathode was poised at a potential of -0.7 V vs. Ag/AgCl, and the corresponding 260 hydrogen production rate was up to 0.6 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>, which is 3.6 times higher than the abiotic control 261 (0.08 m<sup>3</sup>m<sup>-3</sup>d<sup>-1</sup>) (Rozendal et al., 2007). A similar finding was reported by Jeremiasses et al. (2010), 262 who found that compared with an abiotic control, the biocathode increased HER by 21% (up to 263 0.11 L for 52 h). Microorganisms in the biocathode consisted of 46% Proteobacteria, 25% 264 Firmicutes, 17% Bacteroidetes, and 12% related to other phyla (Croese et al., 2011). Considering 265 266 that biocathodes could potentially be a low-cost substitute to metal-based catalysts, further understanding and development of biocathodes for HER is crucial. 267

268

#### 269 *3.3. Membrane options*

In general, most MECs are equipped with a cation exchange membrane or proton exchange 270 membrane (PEM) such as Nafion<sup>®</sup> 117 type PEM (Dhar and Lee 2013). The use of a membrane 271 separator in a MEC helps to prevent substrate crossover between the two half-cells, thereby 272 minimizing the loss of hydrogen (Logan et al., 2008). However, the membranes in wastewater-273 treating MECs often leads to the so-called pH splitting limitation due to the magnitudes higher 274 concentration of other ions such as Na<sup>+</sup>, K<sup>+</sup>, NH4<sup>+</sup>, and Ca<sup>2+</sup> compared with H<sup>+</sup> in wastewater 275 (nearly 10<sup>5</sup> times higher than that of proton H<sup>+</sup>) (Rozendal et al., 2006). As a result, the analyte 276 277 can easily become acidified, suppressing the microbial activity of substrate oxidation (Liu et al.,

278 2005), and the catholyte to become more alkaline, which is unfavorable for the hydrogen evolution 279 reaction. Recently, a sulfonated polyether ketone-based novel nanofiber reinforced PEM (NFR-PEM) was developed as a proton conductor for MECs, which showed lower gas and fuel 280 crossovers with higher proton conductivity compared with Nafion<sup>®</sup> membrane (Chae et al., 2014). 281 Membrane electrode assembly (MEA) cathode has also been developed to enhance hydrogen 282 production efficiency (maximum hydrogen efficiency of 41% with an applied voltage of 1.2 V) in 283 MECs (Jia et al., 2012). However, the use of membrane would incur significant capital cost. It has 284 been estimated that the cost of ion exchange membrane accounted for 38% (400  $\in$  m<sup>-2</sup>) of the 285 capital cost of a laboratory –scale H<sub>2</sub>-MEC, suggesting that nearly half of the total cost of MEC 286 was associated with the use of membrane (Rozendal et al., 2008). 287

On the other hand, avoiding the use of membranes could prevent the pH splitting limitation 288 289 and reduce capital costs. This may also allow the design of simpler reactor configurations (Call and Logan 2008). However, the membrane free MECs were also found to be problematic due to 290 diffusion of hydrogen from cathode to anode, where hydrogen may become available to 291 292 hydrogenotrophic methanogens leading to methane production. It was found that at an applied voltage of 0.2 V, methane concentrations in the product gas increased up to 28% due to the long 293 cycle time of the reactor. The high cathodic hydrogen recoveries  $(78 \pm 1\% \text{ to } 96 \pm 1\%)$  and lower 294 methane (1.9±1.3%) were achieved in a membrane free MEC with applied voltages ranging from 295 0.3 to 0.8 V, and with a solution conductivity of 7.5 mS cm<sup>-1</sup>(Call and Logan 2008). 296

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298 *3.4. Substrate versatility* 

MEC can produce hydrogen from a wide range of simple and complex organic substrates. Table 1 summarizes hydrogen production rate (in decreasing order) with different amounts of substrate 301 (mM or g/L) such as acetate, glucose, trehalose, glycerol, bovine serum lignocellulose and 302 different mixed waste stream from domestic and industrial sources. Indeed, the selection of substrates used in MEC can influence many process parameters such as current density (I,  $A/m^3$ ), 303 applied voltage (V); overall H<sub>2</sub> recovery ( $R_{H_2}$ , %); and energy efficiency relative to electrical input 304 (nE, %). Particularly, the selection of substrate can remarkably affect the hydrogen production rate 305  $(Q, m^{3}H_{2}/m^{3}d)$  (Kadier et al., 2014). Typically, fermentation end products such as acetate have 306 most commonly been used as MEC feedstocks. In fact, the most efficient MEC (hydrogen 307 production rate of 50 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>) reported thus far were fed with acetate (Jeremiasse et al., 2011). 308 Many other substrates have also been used for bioelectrohydrogenesis, including glucose (1.23 m<sup>3</sup> 309  $m^{-3} d^{-1}$ ), butyric acid (0.45  $m^3 m^{-3} d^{-1}$ ), lactic acid (1.04  $m^3 m^{-3} d^{-1}$ ), propionic acid (0.72  $m^3 m^{-3} d^{-1}$ ), 310 valeric acid (m<sup>3</sup>m<sup>-3</sup>d<sup>-1</sup>) (Cheng and Logan 2007), P-glycerol (0.8 m<sup>3</sup>m<sup>-3</sup>d<sup>-1</sup>) (Selembo et al., 2009b), 311 312 B-glycerol (0.41 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>) (Selembo et al., 2009b) and Trehalose (0.25 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>) (Xu et al., 2014a). However, it should be noted that because the anodic substrate oxidation and cathodic hydrogen 313 production take place at different locations within a MEC, bioelectrohydrogenesis rates of MECs 314 can vary remarkably, even when the systems are loaded with the same substrate. For example, 315 hydrogen production rates ranging from 0.01 to 50 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup> were recorded from various acetate-316 fed MECs. Therefore, other operational factors such as substrate concentration, applied voltage, 317 electrode materials, microbes and reactor configuration should also be considered (Kadier et al., 318 2014). 319

Using particulate, complex substrates such as sewage sludge directly as the feedstock for bioelectrohydrogenesis is uncommon due to the low concentration of soluble organic carbon (Ntaikou et al., 2010). To facilitate the treatment of these substrates, feedstock pretreatment could be an effective option. For instance, the bioelectrohydrogenesis rate of a MEC fed with an alkalinepretreated waste activated sludge (WAS) was 16-fold higher than the control without pretreatment (0.91 vs. 0.056 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup>) (Lu et al., 2012c). It was also found that bifrequency ultrasonic solubilization pretreatment could significantly increase the solubilization of carbon (mainly as short chain fatty acids) from WAS, leading to an improved bio-hydrogen yield (Liu et al., 2012). Their results showed that >90% of acetate and ~90% of propionate were effectively converted to hydrogen, followed by the utilization of n-butyrate and n-valerate. This finding suggested that cascade utilization of fermentative products occur during bioelectrohydrogenesis in a MEC.

Lu et al., (2010) examined the possibilities of using proteins as the substrate for 331 332 bioelectrohydrogenesis in MECs. Using bovine serum albumin (BSA), they found that hydrogen was produced at a rate of 0.42 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup> with a yield of 21 mmol H<sub>2</sub> g-COD<sup>-1</sup> (applied voltage 0.6 333 V) in single chamber MECs. However, with the same operational condition a substantially lower 334 performance (0.05 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup> and 2.6 mmol H<sub>2</sub> g-COD<sup>-1</sup>) was obtained when a more complex protein 335 (peptone) was used as the substrate. Lignocellulose waste biomass such as corn stover, sugarcane 336 bagasse, straw, sawmill and paper mill discards could be a promising feedstock for the bio-337 338 hydrogen production in MECs (Lalaurette et al., 2009). Lalaurette et al., (2009) investigated a twostage process by combining dark-fermentation and electrohydrogenesis process that produces the 339 overall hydrogen yield of 9.95 mol-H<sub>2</sub>/mol-glucose using cellobiose. Similarly, the integrated 340 hydrogen production process from cellulose by combining dark fermentation, MFC, and MEC 341 yielded a higher maximum of 14.3 mmol H<sub>2</sub>/g cellulose with a rate of 0.24 m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup> (Wang et al., 342 2011). 343

344

#### 345 4. Interference of methanogens in H<sub>2</sub>-MEC

A vast diversity of microbes can be co-enriched within a MEC. These microbes include 346 extracellular electron transferring bacteria such as Geobacter sulfurreducens, Shewanella 347 putrefaciens, Rhodoferax ferrireducens, Rhodopseudomonas palustris DX-1, and Ochrobactrum 348 anthropi YZ-1 (Fedorovich et al., 2009). Additionally, methanogenic archaea, e.g. 349 hydrogenotrophic methanogen orders Methanobacteriales (MBT) and Methanomicrobiales 350 acetoclastic families 351 (MMB), and methanogen Methanosarcinaceae (MSC) and Methanosaetaceae (MST) within the order Methanosarcinales may also be present in these MECs 352 (Lu et al., 2012b). These microorganisms were generally found in most of the mixed inoculums of 353 354 bioelectrochemical systems (MEC/MFC). The activity of methanogens in H<sub>2</sub> producing MECs severely suppresses hydrogen yield and the purity of the produced hydrogen (Tice et al., 2014). 355

The co-production of methane with hydrogen has been observed in MECs fed with acetate, 356 glucose and complex organic matter (Call and Logan 2008; Chae et al., 2010; Hou et al., 2014; 357 358 Chae et al., 2008; Wagner et al., 2009). Because most MEC processes are operated under fully anaerobic conditions, methanogenesis can also take place when acetate or H<sub>2</sub> are available as 359 substrates. Acetoclastic methanogens convert acetate to methane (reaction 5) whereas 360 hydrogenotrophic methanogens can utilize carbon dioxide and hydrogen to form methane (reaction 361 6) (Wang et al., 2009; Chae et al., 2010). In H<sub>2</sub> producing MECs, the processes that lead to 362 hydrogen and methane production are shown below, 363

364 Hydrogen production by ARB,

365 Anode: 
$$CH_3COOH + 2H_2O \rightarrow 2CO_2 + 8H^+ + 8e^-$$
 (3)

366 Cathode:  $8H^+ + 8e^- \rightarrow 4H_2$ 

367 Co-production of CH<sub>4</sub> by methanogens,

(4)

$$368 \qquad CH_3COOH \rightarrow CH_4 + CO_2 \qquad (5)$$

$$369 \qquad 4H_2 + CO_2 \rightarrow CH_4 + 2H_2O \tag{6}$$

370 Hence, the production of hydrogen at the cathode would be tremendously hampered by methanogenic activity due to the consumption of acetate or hydrogen for methane production (Lu 371 et al., 2012a). Ultimately, acetoclastic methanogens would decrease the efficiency of electron 372 373 transfer from the substrate (electron donor) to the anode (reaction 5). In other words, acetoclastic methanogens would compete with exoelectrogens (ARB) for substrates such as acetate thus 374 reducing the columbic efficiency of bioelectrohydrogenesis. Hydrogenotrophic methanogens 375 directly consume H<sub>2</sub> produced on the cathode (reaction 6), decreasing the cathodic hydrogen 376 recovery (Lu et al., 2011). Thus, to maximize the electron efficiency and cathodic hydrogen 377 378 recovery, it is critical to suppress methanogenic activity in  $H_2$  producing MECs.

# 379 4.1. Methanogenesis control methods and inhibition of methanogenesis by targeting Methyl 380 Coenzyme M reductase (MCR)

To improve hydrogen yields in the MEC reactor we need to inhibit acetate and hydrogen utilizing methanogens, sulfate reducers and homoacetogens. The use of chemical inhibitors targeting specific groups of microbes may potentially address the challenge of low  $H_2$  yields, as well as methane and sulfide contamination in  $H_2$  producing MECs. To control the activity of methanogens for undesirable biological metabolisms in  $H_2$  producing MECs, specific inhibitors should be used for acetate utilizing sulfate reducers, acetoclastic methanogens, hydrogen utilizing sulfate reducers, hydrogenotrophic methanogens, and homoacetogens (Fig. 4).

In general, anti-microbial compounds compete with the target enzymes involved in the biochemical pathways for methane formation (Chae et al., 2010; Catal et al., 2015). It is understood 390 that halogenated hydrocarbons (e.g. CHCl<sub>3</sub> or CHX<sub>3</sub>) can inhibit the production of methane from H<sub>2</sub>/CO<sub>2</sub> and acetate. This is due to the complete blocking of corrinoid enzymes. To inhibit methyl-391 coenzyme M reductase in hydrogenotrophic and acetoclastic methanogens, 2-bromomethane 392 sulfonate (2-BES) and Lumazine are often used as methanogenic inhibitors (Liu et al., 2011). 2-393 BES is a structural analog of CoM. Hence, it can block methane formation catalyzed by methyl-394 CoM reductase. Similarly, Lumazine is a structural analogue of methanopetrin and it can inhibit 395 methanogenesis. Due to the specificity of these chemicals, they are considered specific inhibitors 396 for methanogens. 397

For example, it has been reported that for complete inhibition of methanogenesis in a thermophilic anaerobic digestion process, a very high concentration (50 mM) of 2-BES is required (Zinder et al., 1984). In a separate study, a much lower concentration of 2-BES (10 mM) was found to be effective at suppressing methanogenesis in a similar anaerobic digestion system (Siriwongrungson et al., 2007). In soil systems, the effective inhibitory concentrations of 2-BES were reported to range from 5 to 20 mM, whereas <1 mM 2-BES was required to inhibit rumen methanogens (Wüst et al., 2009; Ungerfeld et al., 2004).

The specific inhibitor sodium molybdate (5 mM) can be effectively used as to inhibit sulfate reducing bacteria (Scholten et al., 2000) to control hydrogen sulfide formation. Also, halogenated aliphatic hydrocarbon compounds (e.g. CHCl<sub>3</sub>) can inhibit the activity of methanogenic archaea as well as of homoacetogenic bacteria and acetate/hydrogen-utilizing sulfate-reducing bacteria (Scholten et al., 2000; Liu et al., 2011).

Numerous reports have explored strategies to inhibit methanogens or suppress methane
 formation in H<sub>2</sub> producing MECs (Table 2). Typically, those strategies entail the manipulation of

412 the physiochemical conditions of the process, targeting the sensitive nature of methanogens to the imposed environmental stress. For example, Hu et al. (2008) examined three different suppression 413 strategies, namely (i) lowering the electrolyte pH to 5.8 with phosphate buffer: NaH<sub>2</sub>PO<sub>4</sub>, 25.4 414 g/L; Na<sub>2</sub>HPO<sub>4</sub>, 4.25 g/L; (ii) exposing the cathode to air for 15 min when the methane was found 415 to have accumulated in the MEC headspace; and (iii) boiling the anodes from MFCs at 100°C for 416 15 min before placing them in the MEC. Their results implied that lowering the pH in the MEC to 417 5.8 was immediately effective for suppressing methane production. However, methane production 418 (up to 5.5%) resumed after two batch cycles, suggesting that the acidic shock could only be a short-419 term solution to the problem (Hu et al., 2008). Similar findings were reported by Kim et al. (2004) 420 and Chae et al. (2010), who showed that acidification also led to inhibition of the exoelectrogen 421 and hence a reduced efficiency of H<sub>2</sub> production. Hence, using acidification to suppress 422 methanogenesis in MEC may not be suitable. 423

It has been demonstrated that a remarkable inhibition of methanogenesis was achieved by lowering the operating temperature to 15°C and 4-9°C (Liu et al., 2005; Lu et al., 2011). However, as most exoelectrogens and methanogens can tolerant a broad range of temperatures, lowering the temperature does not significantly contribute towards improving the hydrogen yield. Further, this method is not effective for suppressing methanogenic activity during long-term operation of H<sub>2</sub> producing MECs (Rader and Logan 2010).

430

431 Another effective strategy to suppress methane production is via optimization of applied 432 voltage. In general, increasing the applied voltage of a MEC increases  $H_2$  production and 433 concentration. It was shown that methane production was higher than  $H_2$  production with a 434 relatively low applied voltage of 0.4 V (22% H<sub>2</sub> and 68% CH<sub>4</sub>), whereas with a higher applied voltage of 0.7 V, methane production decreased to <4% (Wang et al., 2009). However, increasing 435 the applied voltage (at a given current density) would increase energy consumption, resulting in a 436 437 "trade-off" between H<sub>2</sub> production and energy consumption. In single chamber MECs inoculated with mixed cultures from wastewater, the combination of short operation cycles and higher applied 438 voltages could further reduce the methane production to 3%, albeit the methane production was 439 not completely eliminated (Wang et al., 2009). Nam et al. (Nam et al., 2011) reported that there 440 was lower methane production at the anode set potential of -0.2V (vs. Ag/AgCl) compared with 441 other set potentials (-0.4 V, 0 V and 0.2 V vs. Ag/AgCl). However, the improved hydrogen yield 442 (68% H<sub>2</sub> and 21% CH<sub>4</sub>) was only transient (i.e. during the initial 38 days), and the composition of 443 the produced biogas after 39 days became significantly enriched with methane (55% H<sub>2</sub> and 34% 444 CH<sub>4</sub>) (Nam et al., 2011). 445

446 The use of methanogenic inhibitors in MECs may offer several advantages over other physicochemical methods. The use of 2-bromoethane sulfonate (2-BES) to inhibit methane 447 generation in MECs has been well studied. For example, it was reported that the addition of 2-BES 448 449 (286  $\mu$ M) reduced methane generation from 145.8 ± 17.4  $\mu$ mol-CH<sub>4</sub> to 10.2 ± 1.2  $\mu$ mol-CH<sub>4</sub>, reducing the electron loss (as CH<sub>4</sub>) from  $36 \pm 4.4$  % to  $2.5 \pm 0.3$  % in a mixed culture H<sub>2</sub> producing 450 MECs (Chae et al., 2010). The acetate-fed MEC achieved an overall hydrogen efficiency from 56 451  $\pm$  5.7 % to 80.1  $\pm$  6.5 % (equal to 3.2 mol-H<sub>2</sub>/mol-acetate). Also, it was found that in an MFC, a 452 significant fraction (35-56 %) of removed soluble chemical oxygen demand (sCOD) was used by 453 methanogenesis or other undesired biological processes leading to low coulombic efficiency (0.7-454 455 8 %). However, after adding 6 mM 2-BES to the MFC bioreactor, no methane was detected and the power density of the MFC increased by 25% (He et al., 2005). 456

Recently, improved hydrogen production was demonstrated in single chamber MECs with the addition of 5% chloroform to inhibit methanogens for up to 11 cycles (Zhang et al., 2016). The maximum hydrogen production obtained was  $8.4\pm 0.2$  mol H<sub>2</sub> mol-glucose<sup>-1</sup> at a rate of  $2.39 \pm 0.3$ m<sup>3</sup> m<sup>-3</sup> d<sup>-1</sup> with high energy efficiency (165 ±5%) (Zhang et al., 2016). Chloroform (CHCl<sub>3</sub>) blocks the activity of corrinoid enzymes and inhibits the activity of methyl-coenzyme M reductase in methanogenic archaea (Table 2).

Hari et al., (2016) examined that the chemical inhibitor 2-BES (10 mM) can effectively 463 suppress methanogenesis in MEC for bioenergy production using fermentable substrates like 464 propionate (Hari et al., 2016). The inhibition of methanogenesis increased coulombic efficiency to 465 about 84 % by encouraging new microbial interactions, which eventually diverted more electrons 466 to current conversion (Parameswaran et al., 2009 and 2010). Addition of Alamethicin (13 µM) can 467 also be used to suppress methanogenesis and promote acetogenesis in bioelectrochemical systems. 468 469 Alamethicin selectively suppressed the growth of methanogens in mixed-culture bioelectrochemical systems. Also, no methane was detected in the mixed-culture reactors treated 470 with alamethicin, and methane was detected without alamethicin at nearly 100% coulombic 471 472 efficiency. This indicates that alamethicin can effectively suppress methanogens and inhibit methanogenesis in MECs (Zhu et al., 2015). 473

Catal et al., (2015) demonstrated that methanogenesis can be controlled effectively in longterm by the addition of inhibitors in hydrogen producing MECs. The methanogenic inhibitors namely neomycin sulfate, 8-aza-hypoxanthine, 2-bromoethanesulfonate and 2-chloroethane sulfonate were used to examine the inhibition of methanogenesis. The application of antibiotics as methanogenic inhibitors in this study provides a novel approach to inhibit methanogenesis in MECs. Moreover, the methanogenic inhibition methods such as applied potential, rapid extraction

of H2, heat treated electrode, use of biocathode, addition of fatty acids, intermittent oxygen exposure, and use of microbial cultures enriched in the presence of the chemical inhibitor were only able to limit methane formation to a certain extent. In contrast, no methane was detected when methanogenic inhibitors were added directly into MECs (Table 2). Also, the methanogenic inhibitors specifically compete with MCR and inhibit methane generation in hydrogen producing MEC. The growth of methanogen in MECs is a known challenge and requires specific control strategies like methanogenic inhibitors (Table 2).

In general, methanogenic pathways use several cofactors, namely coenzyme M (CoM; 487 HSCH<sub>2</sub>CH<sub>2</sub>SO<sub>3</sub><sup>-</sup>), methanofuran (2-aminomethylfuran linked to phenoxy group), and 488 methanopterin (H<sub>4</sub>MPT;5,6,7,8-tetrahydromethanopterin) (Fig. 5). These cofactors act as C1 489 carriers in methanogenesis (Liu et al., 2011) and they are used by all methanogens. The terminal 490 step of the methanogenic pathway is methane formation, whereby the methyl group carried by 491 492 CoM is reduced to methane by an enzyme known as methyl-coenzyme M reductase (MCR). This 493 enzyme catalyzes the reaction of CH<sub>3</sub>-S-CoM (Methyl CoM) with CoB (CoenzymeB) to produce methane (CH<sub>4</sub>) and heterodisulfide CoM-S-S-CoB as presented in Fig. 6. 494

In the methanogenesis pathway, the terminal step is the reaction of CoM with N-7-495 496 mercaptoheptanoylthreonine phosphate (CoB). The main product of this terminal step is methane, although mixed disulfide (CoM-S-S-HTP) could also be formed (Ellermann et al., 1988). The 497 MCR enzyme was isolated from methanogens and tested for the inhibition. Enzyme inhibitors that 498 were selected had a terminal sulfonate  $(SO_3)$  and are structural analogues of CoM. Several 499 inhibitors have been investigated such as 1-butanesulfonate, 1-propanesulfonate, 2-500 501 azidoethanesulfonate, 2-bromoethanesulfonate, 3-azidopropanesulfonate, 3-bromopropane sulfonate, 3-bromopropionate, 3-chloropropanesulfonyl chloride, 3-fluoropropanesulfonate, 3-502

503 hydroxypropanesulfonate, 3-iodopropane sulfonate, 3-mercapto-1-propanesulfonate, 4bromobutyrate, 4-bromobutyrate sulfonate, 7-bromoheptanoylthreonine phosphate (CoB 504 analogue), 4-chlorobutyrate and chloromethanesulfonate (Table 3). These inhibitors compete with 505 506 MCR and inhibit methane generation. It is known that MCR has cofactor 430 (F<sub>430</sub>), which has Ni(I) in its active site. This Ni(I) reacts with inhibitors and changes to the inactive Ni(III) state 507 (Kunz et al., 2006). The central nickel atom of  $F_{430}$  is coordinated by four planar tetrapyrrole 508 nitrogen atoms. For example, the methanogenic inhibitor, 1-bromoethane sulfonate (1-BES) can 509 interact with Ni(I)-MCR<sub>red</sub> and forms the inactive state of Ni(III)-MCR<sub>sulfonate</sub>, while in the absence 510 511 of inhibitor, Ni(I)-MCR<sub>red</sub> interacts with CH<sub>3</sub>-SCoM to form methane as depicted in the reaction scheme in Fig. 7. The use of inhibitors in H<sub>2</sub>-MECs offers an advantage of long-term inhibition. 512 However, the concentration of inhibitors can vary based on the field application and this can 513 514 influence cost of operation of the MECs. To address this challenge for practical applications, the inhibitors can be added only when needed. Another option could be by adopting feedback 515 inhibitor-dosing strategy based on the composition of biogas. Here, if H<sub>2</sub> partial pressure is lower 516 517 than a certain threshold, dosing of an inhibitor is triggered.

518

519 5. Conclusion and future prospects

To achieve large-scale implementation of MECs for hydrogen production, methanogenesis has to be controlled. Other issues that can also influence H<sub>2</sub>-MEC performance are those relating to the bioanode. These include the pH sensitivity of biofilms. Bioelectrohydrogenesis is a microbial process. Therefore, a better understanding of microbial electron transfer mechanisms will certainly be important from a process stability perspective. Reactor design also plays an important role for scaling up of MEC. For example, single chamber MECs that lack a membrane 526 always showed the production of methane with lower hydrogen yields. As discussed, most MEC 527 studies were conducted with small-scale laboratory systems (Table 1). Only few pilot scale plants with capacities between 20 L and 1000 L were trialed, and the performance of these plants was 528 529 affected by technical challenges such as influent flocculation, water leakage, electrochemical losses and production of unfavorable products (Wang et al., 2013). Cusik et al. (2011) developed 530 the first pilot scale (1000 L) single chamber continuous flow membrane-less MECs for 531 bioelectrohydrogenesis. However, their process failed to produce hydrogen due to formation of 532 methane via hydrogenotrophic methanogenesis. It is now accepted that using membrane-less 533 MECs for hydrogen production is practically challenging. To maximize the yield and purity of 534 hydrogen, effective and implementable strategies should be identified to reduce the formation of 535 methanogenic growth and to promote hydrogen formation. As reviewed here, it is feasible to select 536 537 suitable inhibitor(s) to prevent methane formation (Fig. 8). Future research should be devoted towards developing robust, combinatorial and specific anti-microbial approaches to bring the 538 technology towards practical application. 539

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## 875 Figure captions

876	Fig. 1. (A) Year-wise publication of journal papers on MECs and (B) country wise distribution
877	of publications on MECs. Source: "Web of Science" search with "Microbial electrolysis
878	cell" as the research paper topic as in June 2017. (others- Saudi Arabia, Germany,
879	Sweden, Mexico, Denmark, Taiwan, Iran, Wales, Switzerland, Malaysia, Hungary,
880	Greece, Finland, Turkey, Singapore, Qatar, Israel, Ireland, Bulgaria, U Arab Emirates,
881	Thailand, South Africa, Scotland, Russia, Poland, Nigeria, New Zealand, Ecuador,
882	Austria, Vietnam, Romania, Portugal, Morocco, Lebanon, Kuwait, Indonesia, Czech
883	Republic, Chile, Brazil, and Argentina)
884	Fig.2 Operational principle of microbial electrolysis cell (a) and water electrolysis cell (b);
885	Acetate - organic substrate for exoelectrogenic bacteria (Biofilm), Anode- positive
886	terminal electrode that accept e <sup>-</sup> from Exoelectrogenic bacteria, Cathode - negative
887	terminal electrode that donate $e^-$ for $H_2$ evolution; Potentiostat or power supply -
888	Electrical device to control applied cell potential for hydrogen evolution reaction, and
889	PEM- proton exchange membrane (optional)
890	Fig. 3. Hydrogen producing microbial electrolysis set up; (A) H - shaped two chamber MEC —
891	320 mL (Liu et al., 2005) (B) two chamber MEC - 32 mL (Cheng and Logan 2007), (C)
892	single chamber MEC - 28 mL (Calland Logan 2008), (D) Single chamber MEC in round
893	bottom flasks - 250 mL (Brown et al., 2014), (E) single chamber MEC in borosilicate
894	glass serum vials -100 mL (Hu et al., 2008), F) single chamber MEC in borosilicate glass
895	serum tubes - 28 mL (Hu et al., 2009), (G) continuous flow MEC with multi-electrodes -
896	2.4 L, 1.67 mL min <sup>-1</sup> (Rader and Logan 2010), (H) pilot-scale continuous flow MEC fed
897	with winery wastewater — 1000 L, 1 L d <sup>-1</sup> (Cusik et al., 2011).

898	<b>Fig. 4.</b> Inhibition of undesirable biological metabolisms in H <sub>2</sub> producing MECs by selective
899	methanogenic inhibitors (CHCl <sub>3</sub> , 2-BES, CH <sub>3</sub> F, Na <sub>2</sub> MoO <sub>4</sub> , etc.,) additions to augment
900	electrohydrogenesis in MECs.
901	Fig. 5. Hydrogenotrophic methanogenesis and acetoclastic methanogenesis pathways.
902	Hydrogenotrophic methanogenesis starts with stepwise (1-7) reduction of CO <sub>2</sub> to
903	methane via coenzyme-bound intermediates. Acetoclastic methanogenesis starts with the
904	activation of acetate to acetyl-CoA. (H4MPT, tetrahydromethanopterin; CoA, Co enzyme
905	A; CH <sub>3</sub> COSCoA, acetyl-CoA)
906	Fig. 6. Terminal step of methanogenesis for methane generation.
907	Fig. 7. The mechanism of inhibition of the methanogenic enzyme, Methyl –Coenzyme M
908	Reductase (Mcr) by bromoethanesulfonic acid (BES).

**Fig. 8.** Perspective of single-chamber H<sub>2</sub> producing MECs with the addition of suitable inhibitors.

## **Table 1.** Summary of hydrogen production rate in various MEC systems.

MEC configuration / Working volume	Anode	Cathode	Microbial inoculum/ Source	Substrate	Applied voltage (V)	H <sub>2</sub> rate or Yield (m <sup>3</sup> H <sub>2</sub> m <sup>-3</sup> d <sup>-1</sup> )	H2 (%)	CH4 (%)	Ref.
Two chamber continuous flow at 2.6 mL min <sup>-1</sup> / 200 mL	Graphite felt	Co-Mo alloy	Mixed cultures / Waste water effluent	Acetate / 2.72 g L <sup>-1</sup>	1.0	50	NA	NA	Jeremiasse et al., 2011
Single chamber fed batch / 28 mL	Heat treated Graphite brush	Carbon cloth/Pt	Mixed cultures / Pennsylvania State University WWP	Acetate / 1.5 g L <sup>-1</sup>	0.6	3.6	68	35	(Nam et al., 2011)
Single chamber fed batch / 28 mL	graphite brush	Carbon cloth/Pt	Mixed cultures / enriched biofilm in MFC	Acetate / 1 g $L^{-1}$	0.8	3.12	96	1.9	(Call and Logan 2008)
Single chamber fed batch / 28 mL	Carbon cloth	Carbon cloth/Pt	Mixed cultures / Pennsylvania State University WWP	Acetate / 5 g L <sup>-1</sup>	0.6	2.3	85	>1%	(Hu et al., 2009)
Single chamber fed batch / 26 mL	Graphite brush	Carbon cloth	Mixed cultures / enriched biofilm in MFC	Fermentation effluent / 6.5 g L <sup>-1</sup>	0.6	2.11	96	NA	(Lu et al., 2009)
Single chamber fed batch / 28 mL	Ammonia treated graphite brush	Carbon cloth/Pt	Mixed cultures / WWP	P-Glycerol / 1 g L <sup>-1</sup>	0.9	2.01	88	1.2	(Selembo et al., 2009b)
Single chamber fed batch / 28 mL	Carbon cloth	Carbon cloth/NiMo	Mixed cultures / Pennsylvania State University WWP	Acetate / 5 g L <sup>-1</sup>	0.6	2.0	86	<1	(Hu et al., 2009)
Single chamber fed batch / 28 mL	graphite brush	Carbon cloth/Pt	Mixed cultures / enriched biofilm in MFC	Acetate / 1 g $L^{-1}$	0.6	1.99	78	28	(Call and Logan 2008)
Single chamber fed batch / 28 mL	Ammonia treated graphite brush	Carbon cloth/Pt	Mixed cultures / WWP	Glucose / 1 g $L^{-1}$	0.9	1.87	87	1.2	(Selembo et al., 2009b)
Single chamber fed batch / 28 mL	Heat treated graphite brush	Carbon cloth/Pt	Mixed cultures / enriched biofilm in MFC	Food processing waste water / 8.1 Kg m <sup>-3</sup>	0.9	1.8	32	55	(Tenca et al., 2013)
Single chamber / 28 mL	graphite fiber brush	SS brush	Mixed cultures/ ARB biofilm from MFC	Acetate / 1 g L <sup>-1</sup>	0.5	1.7	84	2.3	(Call et al.,2009)
Single chamber batch / 400 mL	Graphite granules	Ti tube/Pt	Mixed cultures / enriched biofilm in MFC	Acetate / 0.5 g	1.0	1.58	88	0.04	(Guo et al., 2010)

Single chamber fed batch / 28 mL	Carbon cloth	Carbon cloth/NiW	Mixed cultures / Pennsylvania State University WWP	Acetate / 5 g $L^{-1}$	0.6	1.5	75	<1%	(Hu et al., 2009)
Single chamber fed batch / 28 mL	graphite fiber brush	SS A286	Mixed cultures / enriched biofilm in MFC	Acetate / 1 g L <sup>-1</sup>	0.9	1.5	80	NA	(Selembo et al., 2009a)
Single chamber fed batch / 26 mL	Graphite brush	Carbon cloth	Mixed cultures / enriched biofilm in MFC	Buffered effluent / 6.5 g <sup>-1</sup>		1.41	83	NA	(Lu et al., 2009)
Single chamber fed batch / 28 mL	Ammonia treated Graphite brush	SS	Mixed cultures / Pennsylvania State University WWP	Acetate / 1 g L <sup>-1</sup>	0.9	1.4	91%	<1	Ambler and Logan 2011
Two chamber fed batch / 14 mL	Ammonia treated graphite granule	Carbon cloth/Pt	Mixed cultures / WWP	Glucose / 1 g $L^{-1}$		1.23	71	NA	(Cheng and Logan 2007)
Single chamber fed batch / 28 mL	Graphite fiber brush	Carbon cloth/Pt	Clostridium thermocellum enriched biofilm in MFC	Synthetic effluent / 5 g $L^{-1}$	0.5	1.11	63	120 mL g-COD <sup>-1</sup>	Lalaurette et al., 2009
Single chamber fed batch / 28 mL	Graphite fiber brush	Pt	Mixed cultures/ Swine farm WWP	Swine waste water/ 2g L <sup>-1</sup>	0.55	1	77	13	(wagner et al., 2009)
Two chamber fed batch / 14 mL	Ammonia treated graphite granule	Carbon cloth/Pt	Mixed cultures / WWP	Acetic acid / 1 g L <sup>-1</sup>	0.6	1.1	91	NA	(Cheng and Logan 2007)
Two chamber fed batch / 14 mL	Ammonia treated graphite granule	Carbon cloth/Pt	Mixed cultures / WWP	Lactic acid / 1 g L <sup>-1</sup>		1.04	91	NA	(Cheng and Logan 2007)
Single chamber fed batch / 28 mL	Graphite fiber brush	Carbon cloth/Pt	Clostridium thermocellum enriched biofilm in MFC	Cellobiose / 5 g $L^{-1}$	0.5	0.96	69	210 mL g-COD <sup>-1</sup>	Lalaurette et al., 2009
Two chamber fed batch / 26 mL	Graphite brush	Carbon cloth/Pt	Mixed cultures / WAS	Alkaline WAS / 2.4 g L <sup>-1</sup>	0.6	0.91	72	NA	(Lu et al., 2012c)
Single chamber fed batch / 28 mL	Ammonia treated graphite brush	Carbon cloth/Pt	Mixed cultures / WWP	Glucose / 1 g L <sup>-1</sup>	0.5	0.83	81	9.5	(Selembo et al., 2009b)
Single chamber fed batch / 28 mL	Ammonia treated graphite brush	Carbon cloth/Pt	Mixed cultures / WWP	P-Glycerol / 1 g L <sup>-1</sup>	0.5	0.80	80	9.5	(Selembo et al., 2009b)
Single chamber fed batch / 28 mL	Ammonia treated graphite brush	Carbon cloth/Pt	<i>Geobacter sp.,/</i> enriched biofilm in MFC	Potato waste water / 1.9-2.5 g L <sup>-1</sup>	0.9	0.74	73	13	(Kiely et al., 2011)

Two chamber fed batch / 14 mL	Ammonia treated graphite granule	Carbon cloth/Pt	Mixed cultures / WWP	Propionic acid / 1 g $L^{-1}$		0.72	89	NA	(Cheng and Logan 2007)
Single chamber fed batch / 28 mL	graphite fiber brush	SS 304	Mixed cultures / enriched biofilm in MFC	Acetate / 1 g L <sup>-1</sup>	0.9	0.59	77	NA	(Selembo et al., 2009a)
Single chamber fed batch / 28 mL	graphite fiber brush	SS420	Mixed cultures / enriched biofilm in MFC	Acetate / 1 g L <sup>-1</sup>	0.9	0.58	67	NA	(Selembo et al., 2009a)
Single chamber continuous flow at 0.88 mL min <sup>-1</sup> / 140 mL	Graphite granules	Carbon felt	Mixed cultures / ARB biofilm from an acetate-fed MFC having a <i>Geobacter</i> -rich community	Acetate / 10 mM	1.06	0.57	59	2	(Lee et al., 2009)
Two chamber fed batch / 14 mL	Ammonia treated graphite granule	Carbon cloth/Pt	Mixed cultures / WWP	Butyric acid / 1 g L <sup>-1</sup>		0.45	80	NA	(Cheng and Logan 2007)
Single chamber fed batch / 26 mL	Graphite brush	Carbon cloth/Pt	Mixed cultures / enriched biofilm of the Harbin Wenchang WWP in MFC	Bovine serum albumin / $0.7 \text{ g L}^{-1}$	0.6	0.42	34	<0.9 mM g- COD <sup>-1</sup>	(Lu et al., 2010)
Single chamber fed batch / 28 mL	Ammonia treated graphite brush	Carbon cloth/Pt	Mixed cultures / WWP	B-Glycerol / 1 g L <sup>-1</sup>	0.9	0.41	87	1.2	(Selembo et al., 2009b)
Single chamber fed batch / 28 mL	graphite fiber brush	SS316	Mixed cultures / enriched biofilm in MFC	Acetate / 1 g L <sup>-1</sup>	0.9	0.35	55	NA	(Selembo et al., 2009a)
Single chamber fed batch / 38 mL	Graphite brush	Carbon cloth/Pt	Mixed cultures / WAS	Trehalose / 50 mM	0.8	0.25	80	NA	(Xu et al., 2014a)
Single chamber fed batch / 28 mL	Heat treated graphite brush	$MoS_2$	Mixed cultures / enriched biofilm in MFC	Industrial waste water 4.1 Kg m <sup>-3</sup>	0.7	0.17	NA	70	(Tenca et al., 2013)
Two chamber fed batch / 14 mL	Ammonia treated graphite granule	Carbon cloth/Pt	Mixed cultures / WWP	Valeric acid / 1 g L <sup>-1</sup>	0.6	0.14	67	NA	(Cheng and Logan 2007)
Single chamber fed batch / 28 mL	Ammonia treated graphite brush	Carbon cloth/Pt	Mixed cultures / WWP	B-Glycerol / 1 g L <sup>-1</sup>	0.5	0.14	82	9.5	(Selembo et al., 2009b)
Single chamber fed batch / 28 mL	Heat treated graphite brush	SS304 sheet	Mixed cultures / enriched biofilm in MFC	Industrial waste water 4.1 Kg m <sup>-3</sup>	0.7	0.12	NA	62	(Tenca et al., 2013)
Two chamber fed batch / 14 mL	Ammonia treated graphite granule	Carbon cloth/Pt	Mixed cultures / WWP	Cellulose / 1 g $L^{-1}$	0.6	0.11	68	NA	(Cheng and Logan 2007)

Single chamber fed batch / 28 mL	Graphite fiber brush	Carbon cloth/Pt	Clostridium thermocellum enriched biofilm in MFC	Lignocellulose / 5 g L <sup>-1</sup>	0.5	0.11	68	120 mL g-COD <sup>-1</sup>	Lalaurette et al., 2009
Two chamber fed batch / 28 mL	Graphite felt	Ti plate/Pt	Pelobacter propionicus/ Anaerobic digested sludge	Acetate / 2 mM	0.8	0.052	97	2.5	(Chae et al., 2008)
Two chamber fed batch / 120 mL	Carbon brush	Pt/C	Mixed cultures / anaerobic sludge from WWP	Acetate / 1 g L <sup>-1</sup>	0.8	0.0231	32	NA	(Xiao et al., 2012)
Two Chamber fed batch / 6.6 L	Graphite felt	Ti/Pt	Mixed cultures / sludge from UASB reactor	Acetate / 10 Mm	0.5	0.02	NA	NA	(Rozendal et al., 2006)
Two chamber fed batch / 120 mL	Carbon brush	Fe/Fe3C @C	Mixed cultures / anaerobic sludge from WWP	Acetate / 1 g $L^{-1}$	0.8	0.0182	35	NA	(Xiao et al., 2012)
Two chamber fed batch / 200 mL	Carbon felt	Ti/Pt	Mixed cultures / Gwangju sewage treatment plant	Acetate / 1.5 g L <sup>-1</sup>	-	0.013	44	NA	(Lee et al., 2015)
Two chamber fed batch / 130 mL	Carbon brush	Carbon cloth/MoS <sub>2</sub> /CNT- 90	NA	Acetate / 1 g L <sup>-1</sup>	0.8	0.01	12.7	NA	(Yuan et al., 2014)
Two chamber fed batch / 120 mL	Carbon brush	CNT	Mixed cultures / anaerobic sludge from WWP	Acetate / 1 g L <sup>-1</sup>	0.8	0.0076	16	NA	(Xiao et al., 2012)
Two chamber fed batch / 120 mL	Carbon brush	CNT	Mixed cultures / anaerobic sludge from WWP	Acetate / 30 mM	1.06	NA	31	32	(Lee et al., 2009)
Two chamber fed batch / 120 mL	Carbon brush	CNT	Mixed cultures / anaerobic sludge from WWP	Acetate / 80 mM	1.06	NA	28	37	(Lee et al., 2009)
Single chamber fed batch / 130 mL	Graphite fiber brush	Carbon cloth/Pt	Mixed cultures/ Liede WWP	Acetate	0.8	3.7 mol H <sub>2</sub> /mol acetate	95	<0.6	(Hou et al., 2014)
Two chamber fed batch / 28 mL	Heat treated Graphite brush	SS/Pt	Mixed cultures/ Pennsylvania State University WWP	Acetate/ 1.5 g L <sup>-1</sup>	0.9	$3.2 \text{ mol } H_2/$ mol acetate	90	NA	(Nam and Logan 2011) (Nam and Logan 2011)
Two chamber continuous flow at 0.368 g L <sup>-1</sup> / 292 mL	Carbon paper	Carbon paper/Pt	Mixed cultures / enriched biofilm in MFC	Domestic waste water/ 1 g L <sup>-1</sup>	0.5	0.154 H <sub>2</sub> g- COD <sup>-1</sup>	42	NA	(Ditzig et al., 2007)

2 Note: WAS- waste activated sludge; WWP- waste water treatment plant; MFC – Microbial fuel cell; NA- data not available

- 3 Table 2. Methods used for the suppression of methanogens in microbial electrolysis cell for high
- 4 yield hydrogen production

Methanogenesis suppression method	Details	Hydrogen production rate	Remarks	Reference
		$(m^{3}H_{2}m^{-3}d^{-1})$		
Applied potential	0.8 V	-	Methane increased at below 0.8 V	Ding et al., 2016
Rapid H <sub>2</sub> extraction method	gas-permeable hydrophobic membrane and vacuum	1.58± 0.5	No methane	Lu et al., 2016
Heat treated electrode	Bioanode boiled at 100°C for 15 min	0.69	1% methane detected in head space	Hu et al., 2008
Biocathode	Hydrogen producing bioelectrode developed at -0.65 V	10	Methane detected at start up time	Rozendal etal., 2008
Effect of fatty acids	Acetic acid and propionic acid mixture	0.265	No Methane detected.	Ruiz et al., 2014
Oxygen exposure	Bio-anode exposed to air for 24 h	-	No Methane production for 12 h	Ajayi et al., 2010
Specific culture	Heat treated Clostridium <i>ljungdahlii</i> isolated from anerobic sludge treated with 2- bromoethanesulfonate	-	No methane detected over 300 days. Acetate along with hydrogen were produced from CO <sub>2</sub>	Bajracharya et al., 2017
Chemical inhibitor or methanogen	5% chloroform	$2.39\pm0.3$	No methane was detected in fed batch cycle	Zhang et al., 2016
	2-bromoethanesulfonate, 10 mM	$1.08 \pm 0.1$	No methane detected	Hari et al., 2016
	2-bromoethanesulfonate (286 μM)	-	No methane detected	Chae et al., 2010

2-bromoethanesulfonate (50 mM)	-	Methanogens were completed inhibited	Parameswaran et al., 2009
Alamethicin (13 µM)	-	No methane detected	Zhu et al., 2015
2-chloroethane sulfonate (20 mM), 2-bromoethane sulfonate (20 mM), 8-aza- hypoxanthine (3.6 mM)	-	Methane inhibited with increasing hydrogen production	Catal et al., 2015

### 8 **Table 3.** Inhibition of Methyl-Coenzyme M reductase (MCR) for different methanogens

Inhibitors	Apparent concentration (mM)	Organisms	References
1-butanesulfonate 1-propanesulfonate	70 mM	Methanothermobacter marburgensis -	(Kunz et al., 2006)
2-azidoethanesulfonate 2-bromoethanesulfonate	0.001 mM 4 µM	- Methanothermobacter thermautotrophicus, Methanothermobacter marburgensis	(Gunsalus et al., 1980)
3-azidopropanesulfonate	1 μΜ	Methanothermobacter thermautotrophicus	(Ellermann et al., 1989)
2-bromoethanesulfonate	0.004 mM	-	(Ellermann et al., 1988)
3-azidopropanesulfonate	0.04 mM competitive, reversible	Methanothermobacter thermautotrophicus	(Ellermann et al., 1989)
3-bromopropane sulfonate	0.00005 mM, irreversible, strong inhibitor and competitive substrate	Methanothermobacter marburgensis	(Goenrich et al., 2004)
3-Bromopropionate	irreversible	Methanothermobacter marburgensis	
3-chloropropanesulfonyl chloride	1mM	Methanothermobacter marburgensis	(Kunz et al., 2006)
3-fluoropropanesulfonate	-	Methanothermobacter thermautotrophicus	(Rospert et al., 1992)
3-hydroxypropanesulfonate	-	Methanothermobacter thermautotrophicus	(Ellermann et al., 1989)
3-iodopropane sulfonate	-	Methanothermobacter marburgensis	(Goenrich et al., 2004)
3-mercapto-1- propanesulfonate		Methanothermobacter marburgensis	(Kunz et al., 2006)

4-bromobutyrate	-	Methanothermobacter marburgensis	(Kunz et al., 2006; Goenrich et al., 2004)
4- bromobutanesulfonate	0.006 mM	Methanothermobacter marburgensis	(Kunz et al., 2006)
7-bromoheptanoylthreonine phosphate	-	Methanothermobacter thermautotrophicus	(Gunsalus et al., 1980)
4-Chlorobutyrate	-	Methanothermobacter marburgensis	(Kunz et al., 2006)
4-bromobutyrate sulfonate	-	Methanothermobacter marburgensis	(Dey et al., 2007)
Chloromethanesulfonate	-	Methanothermobacter thermautotrophicus	(Ellermann et al., 1989)

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26 Fig. 2.















Fig. 6





Fig. 8.