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## Microbial fuel cell is emerging as a versatile technology: a review on its possible applications, challenges and strategies to improve the performances

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# Microbial fuel cell is emerging as a versatile technology: a review on its possible applications, challenges and strategies to improve the performances

## Abstract

Microbial fuel cells (MFCs) are emerging as a versatile renewable energy technology. This is particularly because of the multidimensional applications of this eco-friendly technology. The technology depends on the electroactive bacteria, popularly known as exoelectrogens, to simultaneously produce electric power and treat wastewater. Electrode modifications with nanomaterials such as gold nanoparticles and iron oxide nanoparticles or pretreatment methods such as sonication and autoclave sterilization have shown promising results in enhancing MFC performance for electricity generation and wastewater treatment. The MFC technology has been also investigated for the removal of various heavy metals and toxic elements, and to detect the presence of toxic elements in wastewater. In addition, the MFCs can be modified into microbial electrolysis cells to generate hydrogen energy from various organic matters. This article provides a comprehensive and state-of-the-art review of possible applications of the MFC technology. This also points out the various challenges that limit MFC performance. Finally, this article identifies the strategies to improve MFC performance for different applications.

## Disciplines

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1 **Microbial fuel cell is emerging as a versatile technology: A review on its**  
2 **possible applications, challenges, and strategies to improve the**  
3 **performances**

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11 **SUMMARY**

12 Microbial fuel cells are emerging as a versatile renewable energy technology. This is  
13 particularly because of the multidimensional applications of this eco-friendly technology. The  
14 technology depends on the electroactive bacteria, popularly known as exoelectrogens to  
15 simultaneously produce electric power and treat wastewater. The electrode modifications  
16 with nanomaterials such as gold nanoparticles, iron oxide nanoparticles or pre-treatment  
17 methods such as sonication and autoclave sterilization have shown promising results to  
18 enhance the MFC performance for electricity generation and wastewater treatment. The MFC  
19 technology has been also investigated for the removal of various heavy metals and toxic  
20 elements, and to detect the presence of toxic elements in wastewater. In addition, the MFCs  
21 can be modified into microbial electrolysis cells to generate hydrogen energy from various  
22 organic matter. This article provides a comprehensive and state-of-the-art review of possible

1 applications of the MFC technology. This also points out the various challenges that limit the  
2 MFC performance. Finally, this article identifies the strategies to improve MFC performance  
3 for different applications.

4 **KEYWORDS:** microbial fuel cell; electricity generation; wastewater treatment;  
5 bioremediation; biosensor; hydrogen production

## 6 **1. INTRODUCTION**

7 Depletion of non-renewable energy resources and environmental pollution are critical threats  
8 facing us. Extracting energy from organic or inorganic wastes can provide an efficient means  
9 to solve the energy and environmental problems simultaneously. Many anaerobic  
10 fermentation technologies have been combined with other purification techniques to generate  
11 alternative energy fuels such as hydrogen and methane [1-3]. However, a sustainable energy  
12 collection must include a diversity of carbon-neutral and renewable energy technologies.  
13 Microbial fuel cell (MFC) technology has attracted an increased number of researchers in the  
14 recent years due to its potential particularly for bioenergy production and wastewater  
15 treatment. This is reflected by the number of articles published in last five years that has  
16 increased successively from year to year, as shown in Fig. 1. MFC technology has become an  
17 attractive technology today because of its capability to convert the chemical energy present in  
18 organic/inorganic wastes into electrical energy. It links microbial metabolism with  
19 electrochemical reactions [3-5]. Consequently, the technology can be used for electricity  
20 generation, wastewater treatment, bioremediation of heavy metals/toxic compounds and other  
21 niche applications. The general principle of an MFC is given in Fig. 2. MFCs are the  
22 bioelectrochemical devices that typically consist of two chambers i.e. the anode chamber  
23 (anaerobic; contains an electrode, microorganisms and anolyte) and the cathode chamber  
24 (aerobic/anaerobic; an electrode, electron acceptor and a catalyst), separated by a proton

1 exchange membrane (PEM) e.g., nafion [6-8]. The microorganisms are used as the  
2 biocatalysts to oxidize the substrate in the anode chamber, and have been denoted as the  
3 power house of MFCs. The electrons are transferred to the anodic (an electrode) surface,  
4 which are then directed to the cathode through an electrical connection [9, 10]. In the  
5 cathode, the electrons combine with protons and oxygen to form water. A catalyst e.g.,  
6 platinum is generally used to catalyse the reduction reaction in the cathode; alternatively, a  
7 microorganism can also be used to replace such costly catalyst [11, 12].

8         The advantage of MFCs mainly lies in the use of microorganisms as the biocatalysts  
9 at the anode and the cathode chambers of MFCs. The exceptional characteristic of the  
10 microorganisms used in MFCs is their self-potential to mediate the electrons (generated from  
11 the oxidation of the substrates) from their outer cell membrane to the surface of an electrode  
12 (in anode) and to accept the electrons from the electrode surface (in cathode) to catalyse the  
13 reduction of electron acceptors e.g., oxygen reduction [5, 9, 12]. The microorganisms that  
14 contain a molecular machinery to transfer the electrons to an electron acceptor without any  
15 external assistance or to accept the electrons from the electrode surfaces are usually called as  
16 exoelectrogens. Due to this unique characteristic of exoelectrogens the MFC technology has  
17 been experimented for a number of applications. The most widely studied application of  
18 MFC technology is electricity generation. In the anode chamber of an MFC, the oxidation of  
19 organic matter by exoelectrogens results into a low redox potential while in the cathode  
20 chamber, reduction of an electron acceptor e.g., oxygen results into a higher redox potential.  
21 This difference in the redox potentials drives the electrons to flow from the anode to the  
22 cathode, which consequently results in bioelectricity generation. Many different designs have  
23 been utilized to produce electric current in various optimized parameters [10-12]. A pure  
24 culture (e.g., *G. sulfurreducens* and *Shewanella oneidensis*) or a mixed culture (from

1 anaerobic sludge or primary wastewater) can be used to generate electric current [13-18].  
2 Many attempts have been made to increase the electric output in MFCs. Out of these, anode  
3 surface modifications with nanomaterials and bacterial gene modification are the most  
4 prevalent approaches that have been employed to improve the MFC performances [19-21,  
5 22]. For example, nitrogen doped carbon nanoparticles were coated on carbon cloth  
6 electrodes, which increased the power density more than three times as compared to untreated  
7 electrodes [21]. Alternatively, a synthetic flavin biosynthesis pathway from *Bacillus subtilis*  
8 was expressed in *S. oneidensis* MR-1, which secreted a very high amount of flavins than the  
9 wild type, consequently, increasing the power output ~13 folds as compared to wild *S.*  
10 *oneidensis* [22]. Because bacteria can degrade the organic matter present in the wastewater,  
11 the technology can be used to remove the pollutants and generate electricity from wastewater.  
12 Several wastewaters ranging from low-strength to high-strength have been utilized in MFCs  
13 for their treatment and electricity generation simultaneously [23-28, 29-34]. In addition,  
14 MFC can be modified into microbial electrolysis cell (MEC) to produce hydrogen gas, but  
15 unlike MFC, electricity is provided in the MEC to produce hydrogen [35]. Generally, a  
16 voltage of 0.2 to 0.8 V is required to reduce the protons to form hydrogen [10]. Such low  
17 voltage is easily achievable in the MFC. Therefore, an MFC can be used to supply the voltage  
18 to the MEC for hydrogen production.

19 The aim of this review article is to critically analyse the routes of MFC applications  
20 and the strategies to improve their performances. Many review articles have been published  
21 describing specific aspects of the MFCs such as the substrates used in MFCs [3], assessment  
22 of MFC configurations [1], and specific application of MFCs like wastewater treatment [4],  
23 and bioremediation [6]. However, that the current review provides a comprehensive  
24 understanding of the MFC applications, their basic principles, challenges and the strategies to

1 improve their performances. The primary applications of MFCs i.e., electricity generation,  
2 wastewater treatment, bioremediation, biosensors, and hydrogen production have been  
3 covered. A special focus has been given to the strategies to improve the MFC performance,  
4 making the technology scalable in the real world to compete with commercialized green  
5 energy technologies.

## 6 **2. THE 'MOLECULAR MACHINERY' OF EXOELECTROGENS**

7 It is important to get an idea about the unique characteristic of MFC technology because of  
8 which this technology has become the centre of attraction among the renewable technologies.  
9 All the applications of MFC technology are particularly interesting because of the molecular  
10 machinery of the bacteria that helps in transferring the electrons to an electrode surface and  
11 vice-versa. The molecular machinery means the biomolecules, proteins or the genes that help  
12 to donate or accept the electrons between bacterial and electrode interface, which chiefly lies  
13 between the inner and the outer membrane of the bacteria. So far, only two bacteria namely,  
14 *Geobacter spp.* and *Shewanella spp.* have been extensively investigated to explore the  
15 extracellular electron transfer (EET) mechanisms. Two types of EET mechanisms have been  
16 confirmed in both the bacteria [5]. The first is direct electron transfer (DET) mechanism and  
17 the second is mediated electron transfer (MET) mechanism. The molecular machinery  
18 comprising the known pathways and hypothetical pathways is presented in Fig. 3.

19 *G. sulfurreducens* is the most studied and explored exoelectrogen in MFCs. It forms  
20 highly thick biofilms on the electrode surfaces and can utilize the various carbon sources as a  
21 substrate for bioenergy production. It has been found that *G. sulfurreducens* in its initial  
22 stages of biofilm formation relies on MET for electron transport. The exoelectrogen secretes  
23 flavin molecules such as riboflavin in the single layer biofilms. The riboflavin combines with  
24 outer membrane c-type cytochromes (OM c-Cyts) to make a complex that furthers the

1 electron transfer to the electrode surface [5,22]. As the biofilm grows  
2 *G. sulfurreducens* adapts to DET for extracellular electron transport. In a multi-layered  
3 biofilm, *G. sulfurreducens* active adjacent to electrode surface utilizes OM c-Cyts (essentially  
4 OmcZ) for extracellular electron transfer while the bacteria respiring distant from the  
5 electrode produce conductive nanowires (type IV pili) that assist in transporting the electrons  
6 inside the biofilm and finally onto the electrode surface [5].

7 The other exoelectrogen studied extensively for MFC applications is  
8 *Shewanella oneidensis*. The bacterium is the most versatile exoelectrogen in the MFCs  
9 because it exhibits the potential to reduce a variety of electron acceptors [36, 37]. Earlier  
10 *S. oneidensis* MR-1 was thought to produce conductive nanowires like type IV pili of  
11 *G. sulfurreducens*. But it is now confirmed that *S. oneidensis* does not contain nanowires and  
12 these nanowires like structures are the extensions of periplasmic and outer membrane  
13 multiheme cytochromes associated with outer membrane vesicles [38]. This exoelectrogen  
14 secretes mainly two types of flavin molecules. The first is riboflavin (RF) and the second is  
15 flavin mononucleotide (FMN). These flavin molecules act as cofactors for the cytochromes  
16 such as OmcA and MtrC. It has been found that RF acts as a cofactor for OmcA while FMN  
17 contains the binding sites for MtrC. [39]. These complexes, RF-OmcA and FMN-MtrC  
18 further promote the electron transfer to the electrode surfaces [39]. The various known  
19 proteins or genes from different exoelectrogens involved in EET mechanisms are depicted in  
20 Fig. 3. To date, some proteins or genes are well known to participate in EET mechanisms that  
21 function in a specific pathway. However, the functional role of other proteins/genes in EET  
22 mechanisms is still under debate and demands a deep investigation to validate their role and  
23 ability to mediate the electrons transfer.



### 1 3. MFCs FOR ELECTRICITY GENERATION

2 The MFCs are chiefly used for the application of electric current generation and many efforts  
3 have been made to ameliorate the current density such as electrode modifications, MFCs  
4 designs, use of metal catalysts at the anode as well as at the cathode etc. [1, 6, 8, 9, 12].  
5 Recent studies reporting high current densities even from reactors as small as 14 ml are  
6 encouraging [2]. Evidently, Bruce E. Logan and his colleagues at The Pennsylvania State  
7 University, United States America (USA) successfully ran a small fan using an MFC with a  
8 working volume of two litres, (<http://www.engr.psu.edu/mfccam/>). If a two litre - MFC can  
9 run a small fan, then we can conceptually expect higher current output from an MFC of  
10 higher volume capacity e.g., of 2000 litres or even more. But it is unlikely to be materialized  
11 in the near future because of obstacles including very high cost of the materials used in MFCs  
12 (electrodes, PEM), high internal resistance, costly catalysts (e.g., platinum) used in cathode  
13 for oxygen reduction, and limited availability of exoelectrogens in the environment.  
14 However, researchers from all around the world continue to contribute to the technology to  
15 make it a viable alternative for renewable energy generation.

16 The first step in MFCs towards current generation is the acclimatization of the  
17 exoelectrogens in the anode chamber and subsequent biofilm formation on the electrode  
18 surface (anode). Consequently, the exoelectrogens form a conductive biofilm on the anode  
19 surface. The biofilm thickness may be a few tens of micrometre, for example, ~30  $\mu\text{m}$  or ~50  
20  $\mu\text{m}$  [36, 37]. Biofilm formation by exoelectrogens is a unique characteristic and differs from  
21 other bacteria or microorganisms. The development of biofilm on the electrode surface from  
22 the single bacterial cell is stimulated by the assembly of adhesins and extracellular matrix  
23 components [38, 39]. Later, some pivotal proteins specifically pili and outer membrane c-  
24 type cytochromes (OMC c-Cyts) e.g., OmcZ, OmcS etc. also promote the biofilm formation

1 [40, 41]. *Geobacter sulfurreducens* is unable to form biofilm in the absence of pili and OMC  
2 c-Cyts [40]. The formation of thick biofilm is taken as an important parameter in MFCs for  
3 efficient performance. Usually, optimal biofilm thickness is preferred in MFCs for higher  
4 current densities, as highly thick biofilms also confine the electron passage [41]. In addition,  
5 the selection of suitable bacterial inoculum (pure culture or mixed culture) with preferred  
6 substrate can be highly beneficial to extract more energy for the current generation. For  
7 example, *Geobacter sulfurreducens* can reduce acetate with ~100% electron recovery to  
8 generate electricity [42].

9         After the establishment of a suitable biofilm, the exoelectrogens transfer the  
10 metabolically generated electrons from their outer cell membrane to the anode surface. There  
11 are two known electron transfer mechanisms i.e. direct electron transfer (DET) and mediated  
12 electron transfer (MET), which have been observed in case of *Geobacter* species and  
13 *Shewanella* species [43-45]. In *Geobacter sulfurreducens*, DET involves OMC c-Cyts (e.g.,  
14 OmcZ, OmcB) for the short-range electron transfer during the initial development of biofilms  
15 and pili (type IV) for long-range electron transfer in multilayer biofilms [17, 19]. In MET  
16 process flavin molecules such as riboflavin (RF) plays a key role in electron transfers [46]. In  
17 *Shewanella oneidensis*, the complex of cytochromes-flavins mediates the extracellular electron  
18 transfer mechanism. For example, flavin mononucleotide (FMN) acts as a cofactor for  
19 cytochrome MtrC and RF for cytochrome OmcA [47].

20         The transferred electrons on the anode surface are transported to the cathode surface  
21 via an electrical connection. The electrons at the cathode surface react with protons and an  
22 electron acceptor. If the electron acceptor is oxygen the end product will be water, resulting  
23 maximum open circuit voltage (OCV) at the cathode of ca. 0.805 V. Generally, the cathode  
24 surface is bound with a catalyst to increase the oxygen reduction rate. The most commonly

1 used catalyst is platinum [1]. The carbon/platinum electrodes are commercially available with  
2 different concentrations of platinum e.g., carbon cloth with 0.2 mg/cm<sup>2</sup>, 0.5 mg/cm<sup>2</sup>.  
3 Alternatively, a microorganism can also be used for the oxygen reduction to make the fuel  
4 cell more cost-effective. Electron acceptors other than oxygen, such as ferricyanide,  
5 potassium permanganate are also useful alternatives [10].

6 The selection of exoelectrogens, substrate (electron donor), and the final electron  
7 acceptor are the pivotal factors in MFC technology. Different MFCs have used pure cultures  
8 as wells as mixed cultures for bioelectricity generation. Some examples of the MFC studies  
9 with pure cultures and mixed cultures are given in Table 1 and Table 2, respectively. The  
10 performance of similar MFCs with different inoculum can be compared to find which  
11 inoculum is more favourable to generate high power density. Some studies report that mixed  
12 cultures produce high power density than pure cultures [5]. However, a few other studies  
13 showed that pure cultures can also generate high current [40]. For example, in a continuous  
14 flow ministack MFC using carbon cloth for both the electrodes, fed with acetate, *G.*  
15 *sulfurreducens* produced higher power density than the mixed cultures using a similar reactor  
16 and operational conditions [40]. The study achieved a maximum power density of 1900  
17 mW/m<sup>2</sup>, which was approximately 21% more than the mixed cultures (sewage sludge  
18 inoculum) [40]. The selection of the inoculum in a particular growth phase (exponential  
19 phase) is also useful to attain high current in MFCs. It has been found that the bacteria in lag  
20 phase form thin biofilms and contain fewer amounts of c-type cytochromes while the bacteria  
21 in exponential phase form thicker biofilms and contain the higher number of c-type  
22 cytochromes, consequently generating higher electrical current [48]. Moreover, a selective  
23 inoculum of mixed culture referred as controlled inoculum (of known bacteria e.g.  
24 *Pseudomonas aeruginosa*, *Azospira oryzae*, *Acetobacter peroxydans* and *Solimonas*

1 *variicoloris*) has shown to produce a higher power density than unknown inoculum [49]. A  
2 study from our group revealed that such controlled inoculum can produce 100% more power  
3 than anaerobic sludge (inoculum) in a double chamber MFC [49]. Further, some pre-  
4 treatment methods of inoculum can also be employed to enhance the power output in the  
5 MFCs [5].

6         The microbial community structure in an MFC is affected by the type of substrates  
7 used in the anode chamber, which could be simple substrates that are easily fermentable or  
8 complex substrates that are non-fermentable [3]. For example, acetate is commonly used in  
9 MFCs, and the exoelectrogens such as *Geobacter* and *Shewanella* spp. readily use acetate for  
10 electricity production [5]. Therefore, the abundant availability of acetate in the anode can  
11 exclude the effect of other fermentable bacteria. But wastewaters may contain simple as well  
12 as complex organic contents. Hence, pre-acclimation strategies can be employed to hydrolyse  
13 and ferment the wastewaters. For example, three pre-acclimation strategies were employed to  
14 evaluate the response of microbial community for electricity generation in an air-cathode  
15 MFC inoculated with anaerobic sludge from domestic wastewater [50]. In the first strategy,  
16 the MFC was pre-acclimated with glucose and acetate; in the second, with glucose before  
17 adding domestic wastewater and in the third strategy, the wastewater was directly used  
18 without any pre-acclimation [50]. The results revealed a great variation in the microbial  
19 community due to the pre-acclimation strategies. The MFC with first strategy was abundant  
20 with bacteria belonging to phylum *Chloroflexi* and genus *Gemmobacter* while the MFC pre-  
21 acclimated with second strategy contained predominantly *Enterobacter* and *Escherichia*. On the  
22 other hand, the MFC with third strategy was dominant with *Dechloromonas* and *Anaerolinaceae*.  
23 Moreover, the MFC with first strategy generated maximum current density and achieved  
24 maximum COD removal as compared to the other MFCs [50].

1           The researchers engaged in the MFC studies around the globe have endeavoured  
2 many innovative efforts to increase the power output of the fuel cells. Many of them are  
3 developing new MFC designs using different effective materials for the electrodes and  
4 membrane, operating MFCs at specific conditions (e.g., setting electrode potentials,  
5 maintaining pH of the electrolytes, pre-treatment of membranes and electrodes), treatment of  
6 the inoculum, and nanomodification of the electrodes. Some methods used to increase the  
7 electricity generation in the MFCs are discussed in the following section.

8           Electrode modification with metal catalyst or nanoparticles or chemical treatment has  
9 become a new trend to improve the performance of MFCs. The main purpose to modify the  
10 electrodes in MFCs is to increase the power outputs, in the anode by providing high surface  
11 area for the biofilm formation and to increase the exocellular electron transfer (EET)  
12 mechanisms. The cathode modifications are the centre of attraction to replace the highly  
13 costly platinum catalyst by cheaper catalysts of nearly or same catalytic properties [12]. Most  
14 of the studies regarding electrode modifications also claimed to decrease the internal  
15 resistance of the system as well as start-up time of the reactor. In the anode, different  
16 approaches have been employed to modify the electrodes to increase the power outputs either  
17 by simple modification methods such as heat-treated electrodes and nitrogen-doped  
18 electrodes or by some sophisticated tools such as by coating some highly effective catalysts  
19 (e.g., gold nanoparticles, graphene, carbon nanotubes (CNT) etc.) on the electrodes [51-55].  
20 Interestingly, almost every kind of metal nanoparticles or other carbon nanoparticles have  
21 been used in the MFCs. Therefore, the researchers are now using electrode with different  
22 composite materials (e.g., CNT–gold–titania nanocomposites) to improve the performance  
23 [53]. Another effective method includes the use of nitrogen doped carbon nanoparticles to  
24 modify the electrode to enhance the EET mechanism. For example, nitrogen doped carbon

1 nanoparticles were coated on carbon cloth electrodes in a two-chamber MFC inoculated with  
2 *Shewanella oneidensis* MR-1. The study revealed that the treated electrodes absorbed more  
3 electron mediators (flavins) secreted by the organism that subsequently increased the electron  
4 transfer rate. Consequently, the power density also increased more than three times as  
5 compared to untreated electrodes [55]. The anode can also be modified with metal or non-  
6 metal nanoparticles (with different morphologies as well) to influence the EET and thus the  
7 performance of the MFCs. In a study, CNT powder was directly added to the anode chamber  
8 to increase the biofilm growth of *G. sulfurreducens* in a two-chamber MFC using plain  
9 carbon paper as the electrode material in both the chambers [52]. The addition of CNT  
10 powder in the anode chamber reduced the internal resistance of the system as well as the  
11 start-up time of the MFC. The shortened start-up time could be attributed to the promotion of  
12 the bacterial adhesion to the electrode material with the addition of CNT powder in the anode  
13 chamber [52]. The performance of the anode can be further improved by using different  
14 morphologies of the material that can provide more active sites and enhance biocompatibility  
15 with the electrode material. In a double chamber MFC, the anode (carbon cloth) was  
16 modified with bamboo-like carbon nanotubes that produced ca. four times higher power  
17 density than the MFC using plain carbon cloth as the anode [52].

18         It is evident that Fe (III) oxide exhibits high affinity for c-type cytochromes such as  
19 OmcA and MtrC present on the outer surface of *Shewanella* species [38, 39]. Therefore, it is  
20 more favourable for the bacteria to mediate the electrons from its outer surface to Fe (III)  
21 oxide. Moreover, it has been also revealed that *Shewanella* species are more attractive to iron  
22 oxide surfaces [5]. In other words, iron oxide surfaces enhance the microbial growth and  
23 increase the extracellular electron transfer, increasing the biofilm metabolic activity which  
24 can be advantageous for improving the performance of MFC-centred applications. For

1 example, Song et al., utilized graphene/  $\text{Fe}_3\text{O}_4$  nanocomposites coated carbon paper as the  
2 anode electrode to improve the bacterial activity in a two-chamber MFC inoculated with  
3 *Shewanella oneidensis* MR-1 [56]. The results showed that the start-up time of the MFC was  
4 significantly decreased with increase in  $\text{Fe}_3\text{O}_4$  concentration, indicating a faster attachment of  
5 bacteria onto the anode surface, which can be attributed to the high affinity of outer  
6 membrane c-type cytochromes to iron oxide [56]. In addition, the MFC with modified anode  
7 achieved a maximum current density of  $1800 \text{ mA/cm}^2$ , which was ~6 times higher than the  
8 bare anode (carbon paper) [56]. In another study,  $\text{Fe}_3\text{O}_4$ -carbon cloth was used as an anode to  
9 examine the beverage wastewater treatment and electricity generation [57]. The MFC  
10 produced a maximum current density that was 100% higher than the bare cathode and a COD  
11 reduction of ~52% was achieved [57]. The iron oxide layers can be prepared on the electrode  
12 surfaces to make them more biocompatible for enhanced microbial growth and functions. For  
13 example, stainless steel electrodes can be heat-treated to generate a layer of iron oxide on its  
14 surface. Evidently, Guo et al., prepared heat-treated stainless steel electrodes which generated  
15 a layer of iron oxide as confirmed by X-ray photoelectron spectroscopy [58]. This  
16 modification further improved the biofilm formation and enhanced the extracellular electron  
17 transfer as expected. Consequently, the current density was significantly increased. The MFC  
18 generated a maximum current density of  $1.5 \text{ mA/cm}^2$ , which was seven times higher than the  
19 bare electrode [58]. Previously, stainless steel mesh was modified with flame synthesis of  
20 carbon nanostructures on its surface, which increased its BET surface by 300 times as  
21 compared to the bare stainless steel mesh electrode. The microscopy results revealed that the  
22 addition of carbon nanostructures onto stainless steel mesh enhanced the biofilm formation.  
23 As a result, the MFC with modified anode produced a power density of  $187 \text{ mW/m}^2$ , which  
24 was 60 times higher than the bare anode [59].

1           The cathode modification is chiefly focused on to replace the platinum by some other  
2 cost-effective catalysts [12, 60-62]. Cobalt oxide and manganese oxide have shown the  
3 potential to substitute the platinum in MFCs. Specifically, cobalt oxide (with other materials  
4 e.g., iron phthalocyanine or nickel) has been repetitively experimented as a cathode catalyst  
5 for oxygen reduction reaction (ORR) [12]. Such MFCs with modified cathode electrode  
6 produced effective results but slightly lower than the MFCs with platinum (as cathode  
7 catalyst). An MFC using cobalt oxide-iron phthalocyanine as a cathode catalyst for oxygen  
8 reduction produced a maximum power density of ca.  $655 \text{ mW/m}^2$ , which was 37% higher  
9 than the MFC with iron phthalocyanine, indicating the effective potential of oxygen reduction  
10 activity of cobalt oxide for ORR [61]. In contrast, the MFC with a carnation-like manganese  
11 dioxide coated cathode produced 1.5 times higher power density than the plain electrode [62].  
12 Alternatively, some bacteria (pure cultures or even mixed cultures) have also been used as  
13 cathode catalyst for oxygen reduction but could not produce satisfactory electric outputs [24].  
14 Moreover, the overpotential obtained for ORR was also higher in the study due to the poor  
15 bacterial activity at the cathode, neglecting the choice of biocathode in real large scale MFC  
16 applications.

17           The electricity generated from MFCs can be further used to power electric  
18 instruments or machines. MFCs have been successfully applied to operate robots. Such robots  
19 are usually termed as "Gastrobots", which means robots with a stomach. These kinds of  
20 robots can metabolize the natural food or can be sustained by water or air. These robots  
21 digest the substrate fuel and convert it into electricity, which is usually stored in the batteries  
22 fitted in the robots, making them an autonomous power system. Evidently, MFCs were  
23 utilized to power a robot named as "Gastronome". Gastronome is thought to be the first robot  
24 that utilized biomass driven energy conversion technology [63]. Gastronome was built by



1 joining train-like three wheeled wagons, as shown in Fig. 4. A stack of six MFCs was used in  
2 the robot and Ni-Cd batteries were utilized, which were charged by the electric output of the  
3 MFCs [63]. Ecobot-II is another example of a robot that was completely driven by MFCs for  
4 environmental monitoring [64]. A picture of Ecobot-II is shown in Fig. 5. The robot was  
5 connected to a wireless transmitter that was further connected to a sensor (which can be for  
6 temperature, toxicity, humidity etc.) [64]. In addition, the robot was packed with eight MFCs  
7 and utilized raw foodstuffs such as rotten fruits as substrate fuel. The authors also claimed  
8 that Ecobot-II was the first robot in the world powered by MFCs that was utilized for  
9 environmental monitoring [64]. In an alternative study, the MFCs were successfully used to  
10 power wireless sensors to detect the changes in temperature. The diagram of the sensor and  
11 telemetry system powered by the MFC is given in Fig. 6. In this study, the MFC was  
12 connected with a highly efficient electronic circuitry to provide a stable power for wireless  
13 sensor [65]. The electricity produced by the MFC was further stored in a capacitor and was  
14 used to power the telemetry system. However, the voltage generated by the system was lower  
15 (2.1 V) than needed for a commercial electronic circuit (3.3 V). Therefore, a DC-DC  
16 converter was utilized to increase the potential and to power the transmitter that received the  
17 data from the sensor and transmitted to the receiver [65]. Further, Tender et al. demonstrated  
18 the application of MFC for the first time in the world to power a meteorological buoy [66].  
19 They used benthic type of MFCs and the meteorological buoy to measure air temperature,  
20 pressure, relative humidity, and water temperature. The results from this study are shown in  
21 Fig. 7.

#### 22 **4. MFCs FOR WASTEWATER TREATMENT**

23 The process of wastewater treatment involves safe disposal or recycling of water which is  
24 highly polluted or contains toxic substances. Wastewater discharged from different industries

1 can be particularly hazardous. According to an astounding report by Lux Research,  
2 governments and water utilities across the world spent approximate \$28 billion in year 2012  
3 to develop their existing wastewater treatment infrastructure that provided a surplus global  
4 wastewater treatment capacity of 16.3 million cubic metres (m<sup>3</sup>) per day. MFC technology  
5 has the potential to provide an effective platform for the treatment of highly polluted  
6 industrial wastewater or urban wastewater and can curb the financial expenditure, which can  
7 be further used for other development programs of a country.

8 In the late nineteenth century, Habermann and Pommer (1991) used MFCs for  
9 continuous treatment of wastewaters for nearly 5 years [67]. They used sodium sulphate  
10 solution (different concentrations (%), 0.5-5) as the electrolyte in the anode, sulphate reducing  
11 microorganisms such as *Proteus vulgaris*, *Escherichia coli*, *Pseudo-monas aeruoinosa* and *P.*  
12 *fluorescens*, and two types of wastewaters (sewage works effluent and landfill effluent). The  
13 results showed that the MFC achieved a COD reduction of 35% with sewage works effluent  
14 and 75% with landfill leachate [67]. In addition, a maximum anodic current density of 150  
15 mA/cm<sup>2</sup> at a potential of -50 mV was also obtained in the demonstration [67].

16 In the later years, different types of wastewaters were used in MFCs for its treatment  
17 and bioenergy production [54-60, 68, 69, 70-75]. On one side of the picture, MFC technology  
18 can be used to treat the wastewater while on the other side, the wastewater can be used to  
19 provide substrate as the carbon source for the bacterial growth and hence for the end products  
20 of the oxidation process i.e. electrons and protons for sustainable bioelectricity generation  
21 [3]. Primary wastewater from an industry such as chocolate industry wastewater [29] or palm  
22 oil mill effluent (POME) [34] can be used to provide the inoculum or the biocatalysts for the  
23 substrate oxidation. Moreover, defined bacterial culture (pure or mixed) can be isolated from  
24 the wastewater that can be further used as inoculum for the MFCs [5]. The wastewater can be

1 used as catholyte as well though it may contain some minerals that can act as electron  
2 acceptors [29]. Though our review is focused on the performance of MFCs for wastewater  
3 treatment, the next section of the article reviews some studies that demonstrated the  
4 efficiency of MFCs for wastewater treatment and some approaches employed to improve the  
5 wastewater treatment efficiency of the MFCs.

6 The effect of different parameters on MFC performance has been studied. These  
7 primarily include chemical oxygen demand (COD), biochemical oxygen demand (BOD),  
8 total solids, total dissolved solids, acidity etc. Usually, standard methods are adopted to  
9 evaluate the wastewater treatment efficiency of the MFCs. Typically, COD test is performed  
10 (or is sufficient) to examine the performance of MFC toward wastewater treatment. Some  
11 examples of MFC studies demonstrated for wastewater treatment are given in Table 3. The  
12 MFCs have achieved up-to 98% COD removal from wastewater [55, 56]. Almost all the  
13 studies demonstrated for wastewater treatment are coupled with the foremost application of  
14 MFCs i.e. electricity production.

15 Animal wastewaters contain high organic content and high concentrations of  
16 phosphate and nitrate in wastewater, the latter causing eutrophication of surface water. A few  
17 studies have demonstrated the use of animal wastewater in different MFCs for its treatment  
18 and bioenergy production. A study using swine wastewater in different MFCs (two  
19 chambered MFC and single chamber MFC) achieved maximum 92% COD removal and  
20 approximately 83% ammonia reduction after operation of the MFC for around 100 hours  
21 [26]. Another study treated animal carcass wastewater (ACW) with high organic content in  
22 an up-flow tubular MFC [68]. The disposed animal carcasses can be further hydrolysed with  
23 alkaline treatment (sodium hydroxide or potassium hydroxide) into smaller constituents like  
24 amino acids, sugars and minerals forming a sterile solution referred as ACW (of BOD-70 g/l,

1 COD-105 g/l and ammonia-1 g/l). The maximum COD reduction obtained in the  
2 demonstration was more than 50% and the nitrate removal efficiency of MFC was nearly  
3 80% [68].

4 Food wastewater or food industry wastewater is non-toxic but exhibits high BOD and  
5 is rich in sugars and starch as compared to other industrial wastewaters. A study using cereal  
6 wastewater in a double-chambered MFC achieved more than 95% COD removal. The initial  
7 COD of the feed wastewater was 595 mg/l [69]. The production of starch foodstuffs (for  
8 example, potato chips) in food industries requires great usage of water, consequently releases  
9 large quantities of wastewater to the environment. Such starch processing wastewater (SPW)  
10 comprises high contents of proteins, carbohydrates, cellulose, vitamins and other nutrients.  
11 An MFC demonstration used SPW to evaluate the treatment efficiency of a double  
12 chambered MFC. The MFC achieved 98% COD reduction after an operation of 140 days.  
13 This was accompanied by an ammonia-nitrogen removal efficiency of 91% [27]. In another  
14 study involving potato processing wastewater (PPW), 91% of COD reduction was achieved  
15 [33]. Similarly, another organic-rich, nontoxic wastewater i.e. chocolate wastewater was used  
16 in a double chambered MFC by Patil et al. [29]. The results showed that maximum 75% COD  
17 was removed after the MFC operation in batch-mode. The BOD removal and total solid  
18 removal was ca. 65% and 68%, respectively [29].

19 Conventional wastewater treatment techniques cannot effectively treat the  
20 wastewaters containing lignocellulosic biomass (e.g. cellulose, hemicellulose and lignin)  
21 However, Huang and Logan used paper recycling wastewater in a single chamber MFC  
22 (sMFC) for its treatment and electricity generation. The results suggested that the MFC, after  
23 nearly three weeks of operation, achieved more than 76% COD removal while ca. 96% of  
24 cellulose was removed by the bacteria [11]. This indicates that the microbial community in

1 the MFC not only degraded the lignocellulose biomass and converted it to simpler sugars but  
2 also extracted energy from such wastewaters to generate electricity.

3 The brewery wastewater has been widely investigated in different MFCs for its  
4 treatment and bioenergy production. The brewery wastewater exhibits high COD, up to 5000  
5 mg/l. Moreover, it contains high levels of carbohydrates or sugars that can be used as electron  
6 donors in the MFCs. Here we present two examples of the studies that used brewery  
7 wastewater in MFCs. In the first example, air cathode sMFC was used with different  
8 concentrations of the wastewater and was operated in fed-batch mode [32]. When the  
9 wastewater with less COD value was used in the MFC, low COD removal was obtained and  
10 vice-versa. When COD concentration was 84 mg/l and 1600 mg/l, the COD removal was  
11 ~58% and 98%, respectively [32]. In the second study, sMFC was operated in continuous  
12 mode with a hydraulic resistance time (HRT) of 2.13 hours. The wastewater was diluted with  
13 deionized water and the COD ranged between 600 mg/l and 660 mg/l. The sMFC achieved  
14 43% and 46% COD removal, respectively [31].

15 The effect of temperature on treatment efficiency of MFCs was investigated by Ahn  
16 and Logan using air-cathode sMFC [69]. They operated the fuel cell (batch mode and  
17 continuous mode) at two different temperatures i.e. ambient temperature ( $23 \pm 3^\circ \text{C}$ ) and  
18 mesophilic temperature ( $30 \pm 1^\circ \text{C}$ ). The results showed that the % COD removal, as well as  
19 the COD removal rate was higher in the MFCs operated at mesophilic temperature than the  
20 ambient temperature. Moreover, ca. 10% more nitrogen removal was achieved from the  
21 MFCs operated at higher temperature. Overall, the MFCs in the fed-batch mode removed  
22 more than 2.5 times COD as compared to MFCs operated in continuous mode [70].

23

1 Treatment of wastewaters from different other mills (agro-industries and oil  
2 industries) have been also investigated in MFCs. Such wastewaters show high COD and are  
3 toxic. For example, cassava mill effluent can have a COD over 16000 mg/l and a cyanide  
4 concentration of ca. 86 mg/l [71]. A 30 L double chambered MFC achieved nearly 90% COD  
5 removal after 120 hours of operation [71]. Palm oil industries release large amount of highly  
6 toxic wastewater, referred to as palm oil mill effluent (POME). POME exhibits COD and  
7 BOD as high as 50000 and 25000 mg/l, respectively [34]. Cheng et al. treated POME in an  
8 upflow membrane less MFC (UML-MFC) coupling MFC and up-flow anaerobic sludge  
9 blanket (UASB) reactors. This integrated system achieved 96% COD and 94% nitrogen  
10 removal [34].

11

12 Usually, the MFCs produce more power density with wastewater of high COD values.  
13 However, the highly concentrated substrate can cause fouling of the PEM, resulting in the  
14 restriction of protons, which consequently leads to the accumulation of protons in the anode  
15 chamber (low pH) and less availability of protons in the cathode (high pH). Therefore,  
16 concentrated wastewaters are sometimes diluted to maintain proper functioning of the MFCs.  
17 Furthermore, some pre-treatment methods can be employed to change the physiochemical or  
18 biological properties of the wastewater for enhanced performance of the MFCs. For example,  
19 the wastewater can be autoclaved to kill the methanogens (the anaerobic bacteria that  
20 yield methane as a metabolic by-product) that otherwise use the organic matter to produce  
21 methane instead of protons and electrons. A study showed that MFC with the autoclaved  
22 wastewater produced ca. 5% more power density than with raw wastewater [26]. Another  
23 pre-treatment method i.e. sonication was shown to be useful to increase the performance of  
24 the MFCs considerably. This approach was employed using raw wastewater that produced ca.  
25 16% more power density and increased the COD removal efficiency by nearly 5%. The

1 sonication process may improve the performance of the MFC by altering the biodegradability  
2 of the organic matter present in the wastewater or changing the molecular weight or particle  
3 size spectra of the organic matter. Moreover, wastewater stirring has also shown marginal  
4 improvement in the COD removal in the MFC [26]. However, some of these pre-treatment  
5 options are energy-intensive and may not be ideal for scale up.

6         Compared to industrial wastewaters, domestic wastewater is more biodegradable.  
7 Domestic wastewater can be a promising substrate for bioenergy production by MFCs. This  
8 approach can be utilized to make eco-friendly public toilets, which can generate electricity  
9 and can help to keep the surrounding environment neat and clean. For example, a single  
10 chamber air-cathode MFC (3-stage MFC/struvite extraction process system) was utilized to  
11 treat human urine with simultaneous extraction of struvite ( $\text{NH}_4\text{MgPO}_4 \cdot 6\text{H}_2\text{O}$ ), which is an  
12 eco-friendly fertilizer. Struvite crystals are generally present in human urine; thus, these can  
13 be extracted from urine using MFCs [77]. The anode was inoculated with anaerobic sludge.  
14 Human urine, supplemented with 0.5% yeast extract and 1% tryptone, was used as the  
15 substrate. The MFC achieved a power of  $14.32 \text{ W/m}^3$  after first stage, which reduced to  $11.76$   
16  $\text{W/m}^3$  after the third stage [77]. Also, the MFC enhanced urea hydrolysis during the  
17 operation, which was advantageous for struvite precipitation process. In their successive  
18 study, they added sea salts in the human urine (substrate) that increased the electricity  
19 generation as well as the struvite extraction [78]. After the addition of sea salts the power  
20 output increased by 10%, while the struvite extraction enhanced from 21 to 94%. Besides, the  
21 COD removal also improved from 16% to 18% [78]. In addition, the research group of  
22 Ioannis Ieropoulos at University of the West of England, Bristol (UK) had a successful field-  
23 trial on the MFC-based public toilets in Glastonbury Music Festival. A special urinal was  
24 fabricated and the collective urine was fed in the stack of MFCs connected in parallel, as

1 shown in Fig. 8 [79]. The MFCs were directly connected to LED lights to monitor the  
2 electricity generation. The trial was run for approximately 3 months and 2.5 – 5 L of urine  
3 was converted daily to power. For a period of 5 weeks, an average power of 75 mW was  
4 achieved each day and a maximum of 98% COD reduction was observed during the trial [79].  
5 In addition to human urine, human feces have been also used in MFCs to generate electricity.  
6 For example, a two chamber MFC was fed with human feces wastewater for electricity  
7 generation and its treatment. The wastewater was firstly fermented prior to use in MFCs to  
8 enhance the power generation. The results showed that the MFC achieved a maximum power  
9 density of 70.8 mW/m<sup>2</sup> and the total COD reduction was 78% after an operation of 190 h  
10 [79].

11 In MFCs, the wastewater treatment efficiency can be further improved by operating  
12 the fuel cells for longer periods. For example, an MFC (air-cathode) was operated for four  
13 cycles; each cycle lasted for approximately 35 days. The results suggested that the COD  
14 removal after the first cycle was ca. 95% which increased to more than 98% after the end of  
15 four cycles (after 140 days of MFC operation) [27]. This can be attributed to the longer  
16 duration available for the microorganisms to degrade the complex substrates completely into  
17 simpler substances. However, the coulombic efficiency achieved in the demonstration was ca.  
18 7%, indicating that most of the substrates did not convert to electricity, which could be due to  
19 the following reasons: (i) oxygen diffusion, (ii) production of fermented products, (iii)  
20 oxidization of other electron acceptors, and (iv) biomass production [27]. The integration of  
21 MFCs with other wastewater treatment technologies can extract more energy, thereby further  
22 improving the pollutant removal efficiency. Generally, the bacteria in MFCs effectively  
23 degrade the simpler or low-strength wastewaters whereas bioreactors such as anaerobic  
24 digester (AD) or UASB treat high-strength wastewaters [2]. Therefore, the wastewaters with



1 complex composition (e.g. POME) can be subjected to the fermentation in UASB that can  
2 provide more suitable or simpler substrates for electricity generation in MFCs. Moreover, the  
3 residual organics present in the effluent of UASB can be further removed in the MFCs.

4 Generally, the MFCs with smaller volumes (10-100 ml) are used in the laboratory  
5 with synthetic wastewaters. However, a significant number of efforts have been made to  
6 scale-up the MFC technology. For example, Zhu et al., constructed a 2-L MFC with  
7 staggered and inline electrode system using graphite rods [80], demonstrating faster start-up  
8 and higher power output as compared to the MFC with inline electrode array. Evidently, the  
9 former MFC produced a maximum power density of  $23.8 \text{ W/m}^3$  and the latter MFC generated  
10 a maximum power density of  $19.1 \text{ W/m}^3$  [80]. This higher power density can be accredited to  
11 the improved mass transfer in staggered electrode array. Besides, the MFC also achieved a  
12 84% COD reduction [80]. In another study, the MFC was further scaled-up to 20-L to treat  
13 brewery wastewater [81]. No catalyst and ion exchange membrane was used in this study.  
14 This MFC was operated for one year and a stable 75% COD removal performance was  
15 observed during the first five months [81]. Moreover, a maximum of ~94% of COD reduction  
16 was achieved at a flow rate of 1 ml/ min (hydraulic retention time=313) when the MFC was  
17 connected to an external resistor of  $10 \Omega$  [81]. In a subsequent demonstration, a MFC with  
18 90-L capacity (stacked with five modules) was fabricated by Dong et al. [82]. This was  
19 operated in an energy self-sufficient mode for approximately 180 days to treat brewery  
20 wastewater (diluted and real wastewater) [82]. A schematic diagram of the 90-L MFC is  
21 shown in Fig. 9. The results suggested that the MFC obtained a maximum COD reduction of  
22 ~87% and 85% with diluted and real wastewater, respectively. Besides, the MFC with real  
23 wastewater obtained higher energy production ( $0.097 \text{ kWh/m}^3$ ) as compared with diluted  
24 wastewater ( $0.056 \text{ kWh/m}^3$ ) [82]. Therefore, it can be concluded that the scale-up of MFC

1 technology has shown substantial improvements for wastewater treatment as well as for  
2 bioenergy production, which may pave the way for commercialization of MFCs in the near  
3 future.

#### 4 **5. MFCs FOR BIOREMEDIATION OF SPECIFIC CONTAMINANTS**

5 The exoelectrogens produce electrons from their metabolism in the anode chamber of an  
6 MFC, which need to be reduced at the cathode chamber. Therefore, an electron acceptor is  
7 provided at the cathode to overcome the potential losses. In addition, a catalyst can also be  
8 used to increase the reduction reaction rate. Usually, the electron acceptors that exhibit a high  
9 redox potential, faster kinetics, a low cost and easy availability are significant and of great  
10 interest in MFC applications. For example, oxygen is one of the promising and widely used  
11 electron acceptors in the MFCs. In MFC system, various organic and inorganic toxic  
12 elements or compounds can be utilized as the electron acceptor in the cathode chamber for its  
13 removal or reduction to less toxic form and simultaneously for the electric current generation.  
14 For examples, metal ions, perchlorate, nitrobenzene, azo dyes, nitrate ( $\text{NO}_3^-$ ) etc. have been  
15 used as electron acceptors in different MFCs to explore the bioremediation potential of this  
16 technology. Some examples of MFC performance for bioremediation application are given in  
17 Table 4.

18 The high concentration of toxic heavy metals (e.g. cadmium, mercury, lead, arsenic,  
19 chromium etc.) in industrial effluents is harmful to the cellular metabolism of the flora and  
20 the fauna living on our planet. Therefore, the wastewaters that contain high concentration of  
21 toxic heavy metals need to be reduced into nontoxic form before they are discharged into the  
22 environment. MFCs have shown a great potential for the reduction of heavy metals both  
23 when used in the anode as well as the electron acceptor in the cathode chamber [72-75].  
24 Generally, the heavy metals with a high redox potential are of great interest to act as the

1 electron acceptor, to achieve higher power output from the cell. Before discussing the MFC  
2 potential for the removal of heavy metals let us get an idea about the processes that are  
3 responsible for heavy metal removal/reduction in MFCs.

4 Various heavy metals have been investigated in the anode chamber as well as in the  
5 cathode chamber of MFCs for their eco-friendly removal. For anodic removal, generally, a  
6 specific concentration of a heavy metal or toxic element is added in the anolyte  
7 (supplemented with carbon source and bacterial inoculum). On the other hand, a heavy metal  
8 with a high redox potential can be used as the electron acceptor in the cathode chamber. A  
9 few mechanisms have been demonstrated that are responsible for the removal of heavy  
10 metals or other toxic elements during MFC operation. The first mechanism is biosorption that  
11 has been widely recognized for the removal of toxic elements in the MFCs [73]. Biosorption  
12 is a combined term for the processes such as microprecipitation, complexation, chelation,  
13 coordination, and ion exchange. The biomolecules like polysaccharides, proteins and lipids  
14 contain the functional groups such as amine, sulfate, carboxylate, hydroxyl, and phosphate  
15 that help in the biosorption process to remove the heavy metals or toxic pollutants. These  
16 biomolecules may be present in the anolyte or on the bacterial cell walls, which play a major  
17 role in the removal of toxic pollutants. Moreover, some processes like biological oxidation,  
18 chemical oxidation, volatilization, anode electrode adsorption have been found responsible  
19 for the sulfide removal during the MFC operation.

20 A single chamber air-cathode MFC demonstrated for the removal of cadmium (Cd)  
21 and zinc (Zn) showed high removal efficiencies i.e. 90% and 97%, respectively [72].  
22 Moreover, in a dual chamber MFC, vanadium containing wastewater was employed as the  
23 cathodic electron acceptor for its simultaneous removal. The fuel cell after 10 days' operation  
24 achieved ca. 70 % removal of V (V) with a maximum power density of ca. 970 mW/m<sup>2</sup> [73].

1 In another study, a two-chamber MFC obtained a maximum power of ca. 431 mW/m<sup>2</sup> with  
2 more than 99.5% removal of Hg<sup>2+</sup>, which was used as an electron acceptor in the fuel cell  
3 [74]. Also ammonia–copper (II) complexes have been substantially recovered from  
4 wastewater using MFC technology. Cu (NH<sub>3</sub>)<sub>4</sub><sup>2+</sup> complexes can be reduced to Cu or Cu<sub>2</sub>O. In  
5 a study, 96% copper was successfully removed after 12 hours of operation of an MFC at a  
6 pH-of 9.0 [75].

7 Different types of dyes are used for the colouring purpose in the textile industry that  
8 results in the generation of a colossal volume of dye wastewater per year around the world.  
9 Dye wastewater contains many toxic and recalcitrant organic molecules and carcinogenic  
10 chemicals [6]. The discharge of such wastewater is threatening to the environment, animals  
11 as well as to the plants. Therefore, treatment of such hazardous wastewater is essential before  
12 its discharge to the environment. MFC technology provides an eco-friendly alternative for the  
13 treatment of dye wastewater and simultaneous bioelectricity generation. MFCs use  
14 microorganisms, therefore, the dyes can be reduced by different decolorization mechanisms  
15 involving enzymes, low molecular weight redox mediators, and chemical reduction by  
16 biogenic reductants. In the MFCs, the dye decolorization occurs in the anode chamber  
17 biologically under anaerobic conditions. For example, the azo bond of congo red dye was  
18 broken into the intermediates such as aromatic amines that can be completely degraded  
19 abiotically in cathode chamber [76].

20 An sMFC with bioanode and biocathode was demonstrated to decolorize an azo dye  
21 congo red, after the operation of the fuel cell for approximately one day. More than 98%  
22 congo red decolourization was achieved in that study [76]. Transfer of electrons from anode  
23 microorganisms and protons through PEM leads to the degradation of azo bond (–N=N–) in  
24 the cathode. Reduction of azo bond results in the formation of colourless and biodegradable

1 aromatic amines [76]. Dechlorinating microorganisms can be used in MFCs for the  
2 bioremediation of pentachloroethene (PCE) and trichloroethene (TCE) to reduce them into  
3 non-toxic end product ethene. Strycharz *et al.* successfully used *Geobacter lovleyi* and  
4 graphite electrodes (as the electron donor) for reductive dechlorination of PCE [83]. A  
5 consortium of anaerobic and aerobic bacteria in the cathodic chamber of dual chamber MFC  
6 demonstrated efficient degradation of pentachlorophenol (PCP). In the study, degradation rate  
7 for PCP was investigated at different pH values and variant temperatures. The most effective  
8 degradation rates achieved at a constant temperature of 50° C and pH 6 were 0.52 mg/L-h and  
9 0.36 mg/L-h, respectively [83]. In addition, *Geobacter species* have shown the tendency to  
10 reduce aqueous, soluble U (VI) into an insoluble form as U (IV). Multiple lines of evidence  
11 suggest that *G. sulfurreducens* entails the outer-surface *c*-type cytochromes for U (VI)  
12 reduction but do not require pili for the same purpose [84]. Further investigation revealed that  
13 *G. sulfurreducens* strain lacking the *pilA* gene reduced U (VI) to the parallel extent to wild  
14 type strain. Similarly, *c*-type cytochromes are also indispensable for *S. oneidensis* to reduce U  
15 (VI). Gene deletion studies demonstrated the importance of outer membrane, decaheme  
16 cytochrome MtrC in the electron transport to U (VI), as the strains deficient in *mtrC* and/or  
17 *omcA* were unable to reduce U (VI) [85]. Moreover, MFCs utilizing anaerobic biocathodes  
18 have shown the ability to reduce highly toxic Cr (VI) to much less toxic Cr (III) and  
19 subsequent precipitation to Cr (OH)<sub>3</sub> with simultaneous electricity generation [86]. The MFC  
20 with set biocathode potentials reduced Cr (VI) with increased reduction rate of 19.7 mg/L-d.  
21 Further, use of *Shewanella oneidensis* MR-1 (produced riboflavin, an electron shuttle  
22 mediator to transfer electrons) as a biocatalyst in the cathode under aerated conditions in the  
23 presence of lactate showed increased reduction rate for Cr (VI) [87]. An MFC fed with  
24 sulfide and glucose and predominated by *Firmicutes* obtained sulfide removal efficiencies of  
25 up-to 85% and a power output of 572.4 mW/m<sup>2</sup> at a current density of 1094.0 mA/m<sup>2</sup> [88].

1 Recently, analysis of 16S rRNA revealed that a strain showing similarity to *Klebsiella sp.* is  
2 capable of bioremediation of cyanide-containing wastewater in MFC. That study achieved  
3 more than 99.5% removal of cyanide and ca. 88% COD removal rate [89]. The investigations  
4 described in this section reflect that the MFC technology is a promising alternative for the  
5 bioremediation of hazardous contaminants.

## 6 **6. MFCs AS BIOSENSORS**

7 The online water-monitoring system is indispensable to maintain the proper usage of  
8 wastewaters from industries or municipal to conserve the aquatic environment as well as the  
9 public health. The MFC has been proven a successful biosensor to detect the organic  
10 compounds and contaminants in the wastewaters [90-92]. The conventional biosensors  
11 usually require a transducer whereas MFC in itself acts as a transducer, therefore MFC can  
12 prove to be a cost-effective biosensor. In the MFC-based biosensor, the exoelectrogens in the  
13 anode chamber serve as a signal generator or biological recognition element whereas  
14 electrodes and PEM (if used) acts as the transducer. The main advantage of the MFC-  
15 biosensor is its long-term stability. This is because the exoelectrogenic biofilms extend the  
16 lifespan of sensing element and curtail the replacement of sensing elements.

17 The basic principle of MFC-based biosensor is presented in Fig. 10. Generally, a toxin  
18 (or a sample to be detected) is provided at the anode chamber and its effect on the voltage  
19 output is measured. A sudden change in the voltage i.e. either fall or rise in the voltage is  
20 taken as the signal for toxin detection. For example, if a toxic element (i.e. chromium) is  
21 injected in the anode chamber, a sudden or slow fall in the voltage can be expected because it  
22 inhibits growth and activity of the exoelectrogens and, consequently, decreases the voltage  
23 [93]. On the other hand, if a carbon source (i.e. acetate) is injected in the anode chamber, a  
24 rise in the voltage is anticipated because it accelerates the growth and activity of the

1 exoelectrogens and, therefore, increases the voltage [93]. The results from this study are  
2 depicted in Fig. 11, which demonstrates different MFCs as the biosensors using low and high  
3 concentrations of different types of contaminants. Typically, the demonstration used three  
4 samples i.e. chromium (acute toxin), iron (non-toxic metal) and acetate (organic substrate) at  
5 different concentrations (chromium-1 mg/L and 8 mg/L, iron-1 mg/L and 48 mg/L, acetate-  
6 200 mg/L) in separate MFCs. The injection of acute toxic and non-toxic metal suddenly  
7 decreased the voltage marginally at low concentrations and severely at high concentrations.  
8 On the other hand, the addition of carbon substrate increased the voltage [93].

9         The MFC sensors can be operated in two modes. The first is flow-through and the  
10 second is flow-by electrodes. In the first mode, the water sample moves through the porous  
11 electrode, while in the second mode, the water sample flows parallel to the electrode surface  
12 [94]. The operation of MFC sensor in a flow-through mode can improve the diffusion of ions  
13 and the electrolytes, thereby increasing the sensitivity of the MFC-based toxicity sensors.  
14 Moreover, a study reported that flow-through anode in an MFC sensor also enhanced the  
15 diffusion of protons through anodic biofilm, improving the biocatalysis of the substrates by  
16 the exoelectrogens [95]. Evidently, the sensitivity of an MFC-based toxicity sensor was  
17 increased approximately 40 times by using a flow-through anode as compared to the flow-by  
18 anode [96].

19         According to the Michaelis-Menten equation, the biocatalytic activity of  
20 exoelectrogens in the anode chamber depends on the concentration of dissolved organic  
21 matter and it keeps increasing until the concentration of the organic matter reaches a  
22 saturation point [97]. MFC sensors are usually operated in turn-off mode for toxicity  
23 monitoring, and the metabolic activity of exoelectrogens can be suppressed by adding a certain  
24 concentration of a toxic pollutant in the anolyte, resulting a certain change in the electric  
25 output [94, 96]. The biological toxicity of the target toxic pollutants is generally measured by

1 correlating the concentration of the toxin to the electric signal output. Therefore, current  
2 change ( $\Delta I$ ) and inhibition ratio (IR) can be evaluated. Further,  $\Delta I$  can be utilized to obtain  
3 the sensitivity of the MFC-based toxic sensor by normalizing the  $\Delta I$  to the concentration of  
4 the toxic agent. On the other hand, IR represents the amplitude of the electric signal output  
5 and can be used to evaluate the toxicity of pollutants [98]. However, it is still unclear what  
6 maximum concentration of the toxic agent is required to obtain a signal output for toxicity-  
7 monitoring.

8         In the conventional MFC-based sensors, the sensitivity of toxic agents depends on the  
9 bioanode in the system or we can say bioanode acts as a sensing element in the MFC sensor  
10 to monitor the water toxicity. But recently, Yong et al. designed an MFC sensor with  
11 biocathode as the sensing element. The results revealed that the MFC sensor with biocathode  
12 showed better sensitivity than the MFC sensor with bioanode [99]. Such MFC sensors could  
13 be advantageous in comparison to bioanode because they do not need organic matter  
14 supplementation for baseline signal output and can reduce the negative effects of combined  
15 shock of toxicity and organic matter. Moreover, the signal output of an MFC sensor is greatly  
16 dependent and influenced by the performance of the anode and the cathode. Therefore, the  
17 modifications can be done in both the chambers to reduce the response time and increase the  
18 detection capacity. For example, the anode potential of the MFC sensor significantly affects  
19 the biosensor sensitivity and, therefore, can be optimized using a potentiostat. The anode  
20 potential usually determines the energy level of the electrons that get transferred from the  
21 surface of exoelectrogens to the anode surface and, hence, affects the electron transfer rate  
22 and the electric output signal [94]. A study revealed that the MFC sensor operated at a  
23 constant anode potential (-1.5 V) showed the highest sensitivity and an unbiased  
24 measurement of toxicity as compared to the MFC sensor without applying anode potential



1 [96]. Similarly, the cathode of MFC sensor can be altered to improve its water-monitoring.  
2 The performance of cathode (stability and catalysis) can affect the amplitude and the  
3 accuracy of output signal under non-toxic conditions as well as toxic conditions. In a study, a  
4 cathode-based MFC sensor array was designed like a bioanode MFC sensor array to detect  
5  $\text{Cu}^{2+}$  and acidic toxicity. An immediate voltage drop was observed when the MFC was  
6 injected with  $\text{Cu}^{2+}$  (2-6mg/L) and the pH was decreased from 6 to 4 [100]. Results are given  
7 in Fig. 12.

8         The application of an MFC-based BOD sensor with municipal or industrial  
9 wastewater could be more challenging in real-world applications because the wastewaters  
10 contain easily degradable organic matters as well as toxic pollutants. During the operation of  
11 an MFC-based BOD sensor, sudden changes in BOD and toxicity could simultaneously occur  
12 [101]. In a MFC-based sensor, the current density decreases with respect to the toxicity of the  
13 toxic agents, while the current density increases with rise in BOD [101]. Therefore, the  
14 sudden variation in BOD might wane the responses of MFC sensor for toxicity. Evidently, a  
15 study demonstrated that a combined shock of BOD and toxicity affected the signal output  
16 when using the MFC sensor for the detection of Cr (IV) [102]. In other words, it can be stated  
17 that signal interference is caused by the combined shock of BOD and toxicity when MFC  
18 sensor is used for water-monitoring. Recently, Yong et al. studied the effect of organic matter  
19 concentration (in anode) on toxicity monitoring to avoid the signal interference by the  
20 combined shock of BOD and toxicity [103]. The study revealed that the background organic  
21 matter concentration should be fixed at a high level of oversaturation for maximizing the  
22 signal output when the ' $\Delta I$ ' is selected relative to the concentration of a toxic agent. On the  
23 other hand, IR should be fixed to a lower value near to the detection limit to maximize the  
24 signal output [103]. The results of this study are shown in Fig. 13.

1           The passage of oxygen into the anode chamber affects the metabolic activity of  
2 anaerobic microorganisms in MFC-based biosensors, thereby, affecting the biosensor  
3 sensitivity. Therefore, it is important to solve this limitation to improve the performance of  
4 these kinds of biosensors. The oxygen diffusion can be diminished by placing an ion  
5 exchange membrane between the cathode and the anode that is less permeable to oxygen.  
6 Generally, nafion is used as the proton exchange membrane in MFCs, but it shows high  
7 oxygen permeability [94]. Recently, a sulfonated ketone ether membrane was applied in a  
8 MFC-based biosensor replacing nafion. The MFC with the new membrane showed better  
9 sensitivity results as compared to nafion [104]. The better performance was attributed to the  
10 lower oxygen permeability of the membrane [104]. The other challenges include its long  
11 response time and detection reliability to replace the commercialized real water-monitoring  
12 systems. However, the longer response time for detection of contaminants can be minimized  
13 by modifying the MFC sensor structure. For example, in a study, the response time was  
14 significantly reduced from 36 min to 5 min by decreasing the volume of anode from 25 ml to  
15 5 ml in the MFC [104]. On the other hand, the detection reliability can be further ameliorated  
16 by connecting various MFCs in parallel. Such MFC array has been reported for effective  
17 water quality monitoring [94].

18           A few MFC-based biosensors have been commercialized. One such product is named  
19 Biomonitoring system (HATOX-2000), which has been invented by a Korean company and  
20 can be utilized for online monitoring of water toxicity. More detailed information of this  
21 product can be accessed from elsewhere ([www.ecotrade.org](http://www.ecotrade.org)).

22

23

## 1 **7. MICROBIAL ELECTROLYSIS CELLS FOR HYDROGEN PRODUCTION**

2 An MFC produces electricity from organic waste while a microbial electrolysis cell (MEC)  
3 produces hydrogen gas. The working principle of an MEC is similar to an MFC as the  
4 electrons generated by the exoelectrogens in the anode combines with protons at the cathode  
5 to produce hydrogen gas as the final product. But unlike MFC, electricity is provided in the  
6 MEC to produce hydrogen. Theoretically a voltage of 0.2 to 0.8 V is required to reduce the  
7 protons to form hydrogen. Such low voltage is easily achievable in the MFC. Therefore, an  
8 MFC can be used to supply the voltage to the MEC for hydrogen production. The electrode  
9 material used in the MFCs can be employed in the MECs as well. Moreover, the  
10 exoelectrogens are also required to produce hydrogen gas in MECs. In MECs, similar to  
11 MFCs, a cathode catalyst such as platinum is used to overcome the overpotentials to drive  
12 hydrogen production. Unlike MFCs, the MECs require strictly anaerobic conditions for  
13 hydrogen production. However, the higher concentration of hydrogen gas promotes the  
14 growth of methane-producing microorganisms. Subsequently, the hydrogen gas is  
15 contaminated by methane and the resultant hydrogen output is decreased. Different types of  
16 organic sources and wastewater can be applied in MEC for hydrogen production. Notably,  
17 MEC has shown higher hydrogen yields than that obtained with fermentation. For example,  
18 the maximum theoretical yield of 7 mol-H<sub>2</sub>/mol-glycerol by oxidation is achievable. The  
19 hydrogen yields reported in some studies using fermentation vary from 0.05-1.05 mol-  
20 H<sub>2</sub>/mol-glycerol [106, 107], but a hydrogen yield of 3.9 mol-H<sub>2</sub> /mol-glycerol has been  
21 achieved using MEC [108]. In addition, a hydrogen yield of 7.2 mol-H<sub>2</sub> /mol-glucose was  
22 also obtained in the study against the maximum theoretical yield of 12 mol-H<sub>2</sub>/mol- glucose  
23 [108].

1           There are some obstacles that limit the application of MECs at the large scale. For  
2 example, a single MFC generally produces an OCV of approximately 0.8 V and a resultant  
3 working voltage of ~0.5 V can be achieved in an MFC [109]. This decrease in voltage could  
4 be due to higher internal resistance in the MFC system, energy utilization by bacteria, and  
5 electrode overpotentials [109]. Therefore, three or five MFCs can be connected in series to  
6 increase the resultant voltage output. But the voltage reversal can reduce the voltage output  
7 over the long-term [109]. This problem was resolved by Hatzell et al. by using a capacitor in  
8 the circuit to prevent the voltage reversal. In this study, the MFCs were connected in a  
9 parallel configuration to charge the capacitors. Then the capacitors were connected in series  
10 to discharge the voltage to the MECs. Such a system increased the hydrogen production rate  
11 approximately 2.3 times as compared to coupled systems without capacitors [110]. Another  
12 major limitation in MECs is the consumption of hydrogen by methanogens to produce  
13 methane, which consequently reduces the hydrogen generation. Many approaches have been  
14 used to inhibit the methanogens in MECs. For example, the cathode can be exposed to  
15 oxygen or ultraviolet radiation to inhibit the methanogens. In a demonstration, the exposure  
16 of cathode to air decreased the methane concentration from 3.4% to less than 1% [111]. On  
17 the other hand, the exposure of ultraviolet (UV) radiation in the MEC maintained high  
18 concentrations of hydrogen (91%), while without UV irradiation, methane concentrations  
19 increased significantly [112]. Recently, the use of antibiotics has shown the potential to  
20 inhibit the methanogens [113]. In a study, Catal et al. used different concentrations of four  
21 antibiotics (neomycin sulfate, 2-bromoethane sulfonate, 2-chloroethane sulfonate, and 8-aza-  
22 hypoxanthine) to measure the inhibition of methanogenesis on a mixed culture community to  
23 improve the hydrogen production. The results showed that the increasing concentrations of  
24 the antibiotics decreased the concentration of methane effectively that resulted in a  
25 comparatively higher hydrogen production [114]. The third major problem that hinders the

1 use of MEC at pilot scale is the necessity of a catalyst at the cathode. Usually, platinum is  
2 used as the cathode catalyst in MECs that is very expensive. Moreover, it can be easily  
3 poisoned by sulfide present in the water. Therefore, its replacement with a catalyst which is  
4 cost-effective and has similar catalytic properties is required to launch the technology at a  
5 large scale. Some catalysts have been already experimented in MECs to replace platinum. For  
6 example, Yang et al. recently used polyaniline/multi-walled carbon nanotube as the cathode  
7 catalyst in a single chamber MEC. The results suggested that a maximum hydrogen  
8 production rate of  $1.04 \text{ m}^3/\text{m}^3/\text{day}$  was achieved with the catalyst, which was comparable to  
9 the performance with platinum [114]. The same catalyst was further used in a different study  
10 with biocathodes that achieved a maximum hydrogen production rate of  $0.67 \text{ m}^3/\text{m}^3/\text{day}$   
11 [115]. Moreover, nano-Mg (OH)<sub>2</sub>/graphene composites at different concentrations were  
12 demonstrated as the cathodic catalyst in MEC to improve hydrogen production. The cathodic  
13 hydrogen recovery and hydrogen production rate obtained with the catalyst were ca. 84% and  
14  $0.63 \text{ m}^3/\text{m}^3/\text{day}$ , which were higher as compared to the Pt/C cathode [116].

## 15 **8. CONCLUSIONS AND CHALLENGES**

16 The MFCs provide a suitable, eco-friendly alternative to produce energy and to treat  
17 wastewater simultaneously. Several wastewaters ranging from low-strength to high-strength  
18 have been utilized in MFCs for their treatment and electricity generation simultaneously.  
19 However, the power outputs achieved in the MFCs are low and can be enhanced by the  
20 following approaches; 1) a suitable design that results in low internal resistance; 2) using  
21 nanoparticles that increase the electron transfer mechanisms; 3) use of genetically engineered  
22 microorganisms; 4) addition of pre-treated inoculum or control inoculum; 5) decreasing the  
23 start-up time of the MFC. For example, graphene/ Fe<sub>3</sub>O<sub>4</sub> nanocomposites coated carbon paper  
24 as the anode electrode decreased the start-up time and achieved a maximum current density

1 of 1800 mA/cm<sup>2</sup>, which was ~6 times higher than the bare anode [56]. The electricity  
2 generated from MFCs can be further used to power electric instruments or machines. As  
3 noted earlier, MFCs have been successfully applied to operate the "Gastrobots" for bioenergy  
4 production and environmental monitoring.

5 Further efficient treatment of wastewater can be achieved by operating the fuel cells  
6 at mesophilic temperatures. Moreover, the MFCs integrated with other anaerobic  
7 fermentation technologies such as with UASB, have shown enhanced COD removal  
8 efficiency. Significant efforts have been made to scale-up the MFC technology. For example,  
9 a MFC with 90-L capacity obtained a maximum COD reduction of ~87% with brewery  
10 wastewater [82].

11 MFCs have shown a great potential for the reduction of heavy metals or toxic  
12 pollutants when used in the anode as well as in the cathode chamber as the electron acceptor.  
13 The heavy metals with a high redox potential are of great interest to act as the electron  
14 acceptor, to achieve higher power output from the cell. The biomolecules that may be present  
15 in the anolyte or on the bacterial cell walls contain the functional groups, which play a major  
16 role in the removal of toxic pollutants. MFCs have achieved heavy metal removal of even  
17 upto 99.5% (Hg<sup>2+</sup>) and 97% (Zn). The MFCs can also be applied as a BOD or COD sensor to  
18 detect the availability of a toxic pollutant in the wastewater. The voltage drop/rise is taken as  
19 the signal for the detection of the toxin or the sample. The change in voltage is usually  
20 proportional to the concentration of the toxin. The low sensitivity and detection reliability are  
21 the main challenges in MFC-based biosensors. The sensitivity of an MFC-based toxicity  
22 sensor can be improved by operating them in a flow-through mode. A study showed that the  
23 sensitivity of the biosensor increased approximately 40 times by using a flow-through anode  
24 as compared to the flow-by anode [96]. In addition, an MFC can be amended to an MEC to

1 produce another biofuel i.e., hydrogen energy, while the MFC may be a substantial  
2 alternative to supply the required voltage. One of the major limitations in MECs is the  
3 consumption of hydrogen by methanogens to produce methane, which consequently reduces  
4 the hydrogen generation. However, the use of antibiotics and exposure of ultraviolet  
5 radiations have shown the potential to inhibit the methanogens [113]. The results showed that  
6 the increasing concentrations of the antibiotics decreased the concentration of methane  
7 effectively, resulting a higher hydrogen production [114].

8         The MFC technology has been used for various applications, however, there are some  
9 challenges that need to be addressed to make the technology economically viable. The first  
10 prime hurdle is a feasible design for upscaling the MFC. The previous designs exhibit some  
11 drawbacks such as high internal resistance, electrode spacing, exchange of anolyte and  
12 catholyte across the PEM etc. when we think to scale up them for long-term operations.  
13 However, some designs have already been introduced but have not been explored at the  
14 industrial scale. The second challenge is to provide cost-effective electrode materials and  
15 PEM (if used) for MFCs. For scale up, the available electrode materials such as carbon paper  
16 and carbon cloth would be very expensive. Another obstacle is the choice of an electron  
17 acceptor at the cathode. Oxygen is abundantly available and is the preeminent choice for the  
18 electron acceptor. But continuous sparging of oxygen at the cathode can also affect the  
19 activity of anaerobic microbial community at the anode during long-term operations since  
20 oxygen can diffuse through the PEM to the anode. Platinum is most commonly used for  
21 oxygen reduction reaction, but it is very expensive, and a cheaper alternative is required. For  
22 example, at the small scale (MFC of 250 ml capacity), commercially available 0.5 mg/cm<sup>2</sup>  
23 20% platinum on carbon paper of 20 cm<sup>2</sup> costs ~250 US\$ (Fuel Cell Earth, USA). If we want  
24 to scale up the MFC reactor, we need larger electrode and obviously, a large amount of

1 platinum. This makes the use of platinum uneconomical at the large scale. Moreover,  
2 platinum turns poisonous when it reacts with certain elements/chemicals in the water such as  
3 sulphide, making the use of platinum impractical for wastewater treatment application.  
4 Therefore, the replacement of platinum is the must in scaling-up the MFCs.

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## **Highlights**

- The state-of-the-art information on major applications of MFCs and strategies to improve them is provided in this article.
- The basic principles of all the applications are thoroughly discussed.
- The obstacles that limit the technology to use in real world applications are reported.
- Many approaches such as electrode modification, genetic engineering etc. can be utilized to improve the MFC performances.

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Table 1. Performance of microbial fuel cells for bioelectricity generation using pure cultures

Inoculum	Type of MFC	Substrate	Electrode materials	Current density/Power density	References
<i>Klebsiella pneumonia</i>	Single-chamber MFC	Glucose	Carbon cloth	199 mA/m <sup>2</sup>	25
<i>Desulfovibrio desulfuricans</i>	Double-chamber MFC	Wastewater	Graphite felt	233 mA/m <sup>2</sup>	26
<i>Escherichia coli</i>	Double-chamber MFC	Glucose	<sup>1</sup> PAN/ TiO <sub>2</sub> composite-anode Carbon cloth–cathode	3390 mA/m <sup>2</sup>	27
<i>Saccharomyces cerevisiae</i>	Single-chamber MFC	Synthetic wastewater	Graphite plates	282 mA/m <sup>2</sup>	28
<i>Thermincola ferriatica</i>	Double-chamber MFC	Acetate	Graphite carbon fibres	12000 mA/m <sup>2</sup>	14
<i>Lysinbacillus sphaericus</i>	Double-chamber MFC	Glucose	Graphite felt	85 mW/m <sup>2</sup>	30
<i>Citrobacter sp.</i>	Single-chamber MFC	Acetate	Carbon cloth	205 mA/m <sup>2</sup>	31
<i>Ochrobactrum sp.</i>	Double-chamber MFC	Xylose	Carbon fibres brush	2625 mW/m <sup>3</sup>	32
<i>Shewanella putrefaciens</i>	Single-chamber MFC	Lactate	Carbon cloth	4920 mW/m <sup>3</sup>	33
<i>Scenedesnum</i>	Double-chamber MFC	Acetate	Carbon fiber brush-anode Carbon cloth-cathode	1926 mW/m <sup>2</sup>	34
<i>Shewanella oneidensis</i>	Mini-MFC	Lactate	Graphite-felt	3000 mW/m <sup>2</sup>	35

<i>Cyanobacteria</i>	Single-chamber MFC	Domestic wastewater	Graphite felt-anode, Carbon cloth-cathode	114 mW/m <sup>2</sup>	36
<i>Chlorella vulgaris</i>	Double-chamber MFC	Wastewater	Carbon felt-anode Carbon cloth-cathode	2485 mW/m <sup>3</sup>	37
<i>Rhodopseudomonas palustris</i>	Single-chamber MFC	Wastewater	Carbon paper-anode Carbon cloth-cathode	2720 mW/m <sup>2</sup>	38
<i>Coriolus versicolor</i>	Double-chamber MFC	<sup>2</sup> ABTS	Carbon fibres	320 mW/m <sup>3</sup>	39
<i>Geobacter metallireducens</i>	Double-chamber MFC	Domestic wastewater	Carbon paper	40 mW/m <sup>2</sup>	40
<i>Geobacter sulfurreducens</i>	Double-chamber MFC	Acetate	Carbon fibres	1.9 mW/m <sup>2</sup>	17

Note:- <sup>1</sup>PAN= Polyaniline

<sup>2</sup> ABTS = 2, 2'-Azino-bis (3-ethylbenzthiazoline-6-sulfonic acid)

Units of surface power density are given in milliwatts per square meter; volume power density in watts per cubic meter; and current density in milliampere per square meter.

Table 2. Performance of microbial fuel cells for bioelectricity generation using mixed cultures

Source of inoculum	Type of MFC	Substrate	Electrode material	Current density/Power density / Voltage	Reference
Dairy manure wastewater	Single-chamber MFC	Dairy manure wastewater	Graphite fiber brush	190 mW/m <sup>2</sup>	42
Potato wastewater	Single-chamber MFC	Potato wastewater	Graphite fiber brush	217 mW/m <sup>2</sup>	42
Activated sludge	Double-chamber MFC	Acetate, glucose	Carbon paper	410 mV	43
Primary wastewater	Double-chamber MFC	Acetate	Graphite rods	152 mA/m <sup>2</sup>	44
Activated sludge	Single-chamber MFC	Acetate, glucose	Carbon cloth	1084 mW/m <sup>2</sup>	45
Activated sludge	Double-chamber MFC	<sup>1</sup> POME	Polyacrylonitrile carbon felt	107 mW/m <sup>2</sup>	46
Activated sludge	Single-chamber MFC	Glucose	Carbon cloth	68 mW/m <sup>2</sup>	47
Activated sludge	Single-chamber MFC	Acetate	Graphite coated with graphene -anode, carbon cloth-cathode	670 mW/m <sup>2</sup>	48
Primary wastewater	Single-chamber MFC	Acetic acid	Graphite fiber brushes-anode Carbon cloth-cathode	835 mW/m <sup>2</sup>	49
Primary wastewater	Single-chamber MFC	Ethanol	Graphite fiber brushes-anode Carbon cloth-cathode	820 mW/m <sup>2</sup>	49
Primary wastewater	Single-chamber MFC	Lactic acid	Graphite fiber brushes-anode Carbon cloth-cathode	739 mW/m <sup>2</sup>	49
Primary wastewater	Single-chamber MFC	Succinic acid	Graphite fiber brushes-anode Carbon cloth-cathode	444 mW/m <sup>2</sup>	49

Anaerobic sludge	Double-chamber MFC	Slaughterhouse wastewater	Carbon cloth-anode Titanium mesh-cathode	578 mW/m <sup>2</sup>	51
Anaerobic reactor effluent	Double-chamber MFC	Acetate	Carbon cloth-anode Granular active carbon-cathode	1200 mW/m <sup>3</sup>	52
Soil	Double-chamber MFC	Cellulose	Carbon paper	188 mW/m <sup>2</sup>	53

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Note: <sup>1</sup>POME =Palm Oil Mill Effluent

Units of surface power density are given in milliwatts per square meter, volume power density in watts per cubic meter, and units of voltage in millivolts.

Table 3. Performance of microbial fuel cells for wastewater treatment

Wastewater	Type of MFC	Electrode material	% COD reduction	Reference
Swine wastewater	Single-chamber MFC	Toray carbon paper as anode carbon cloth as cathode	92	54
Starch processing wastewater	Single-chamber MFC	Carbon paper	98	55
Real urban wastewater	Double-chamber MFC	Graphite electrodes	70	60
Olive mill wastewaters	Single-chamber MFC	Carbon cloth as electrodes	65	61
Protein-rich wastewater	Double-chamber MFC	Graphite rods as electrodes	80	4
Paper recycling wastewater	Single-chamber MFC	Graphite fibers-brush	76	11
Cassava mill wastewater	Double-chamber MFC	Graphite plates electrode	86	62
Food processing wastewater	Double-chamber MFC	Carbon paper electrodes	95	68
Domestic wastewater	Double-chamber MFC	Plain graphite electrodes	88	69
Chocolate industry wastewater	Double-chamber MFC	Graphite rods as electrodes	75	70
Biodiesel wastes	Single-chamber MFC	Carbon brush electrodes	90	71
Beer brewery wastewater	Single-chamber MFC	Carbon fibers	43	72
Brewery wastewater	Single-chamber MFC	Carbon cloth as electrodes	98	73
Potato Processing wastewater	Tubular MFC	Graphite particles as anode Graphite felt as cathode	91	74



Palm oil mill effluent	<sup>1</sup> UML-MFCs	Graphite granules, Carbon fiber felt	90	75
Animal carcass wastewater	Up-flow tubular MFC	Graphite felt as anode Carbon cloth as cathode	51	76
Food waste leachate	Double-chamber MFC	Carbon felt	85	83
Chemical wastewater	Double-chamber MFC	Graphite plates	63	84

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Note: <sup>1</sup>UML-MFCs = Up-flow membrane-less microbial fuel cell

Table 4. Performance of microbial fuel cells for bioremediation

Heavy metals/ Wastewater	Type of MFC	Electrode material	% Removal	Power density	Reference
Chromium (VI)	Double-chamber MFC	Graphite granules-cathode Graphite brush-anode	94	6.4 W/m <sup>3</sup>	85
Chromium (VI)	Double-chamber MFC	Carbon fiber felt	76	970 mW/m <sup>2</sup>	86
Sulfide	Double-chamber MFC	Carbon fiber felt	85	572.4 mW/m <sup>2</sup>	87
Cadmium	Single-chamber MFC	Carbon cloth	90	3600 mW/m <sup>2</sup>	88
Zinc	Single-chamber MFC	Carbon cloth	97	3600 mW/m <sup>2</sup>	88
Vanadium	Double-chamber MFC	Carbon fiber felt	68	970 mW/m <sup>2</sup>	89
Ammonia–copper (II)	Double-chamber MFC	Graphite felt-anode Graphite plate-cathode	96	140 mW/m <sup>2</sup>	90
Mercury (Hg <sup>2+</sup> )	Double-chamber MFC	Graphite felt-anode Carbon felt- cathode	99.5	433 mW/m <sup>2</sup>	91
Azo dye Congo red	Single-chamber MFC	Carbon brush	98.3	-	92
Cyanide	Double-chamber MFC	Carbon cloth	88.3	-	93
Copper (Cu <sup>2+</sup> )	Double-chamber MFC	Graphite felt electrodes	99.5	319 mW/m <sup>2</sup>	106
Chromium (VI)	Single-chamber MFC	Carbon brush-anode Carbon cloth-cathode	99	419 mW/m <sup>2</sup>	107
Nitrate	Single-chamber MFC	Graphite rods	30	3900 mW/m <sup>3</sup>	108
Nitrite	Single-chamber MFC	Graphite rods	37	3600 mW/m <sup>3</sup>	108

Note: Units of surface power density are given in milliwatts per square meter; volume power density in watts per cubic meter.

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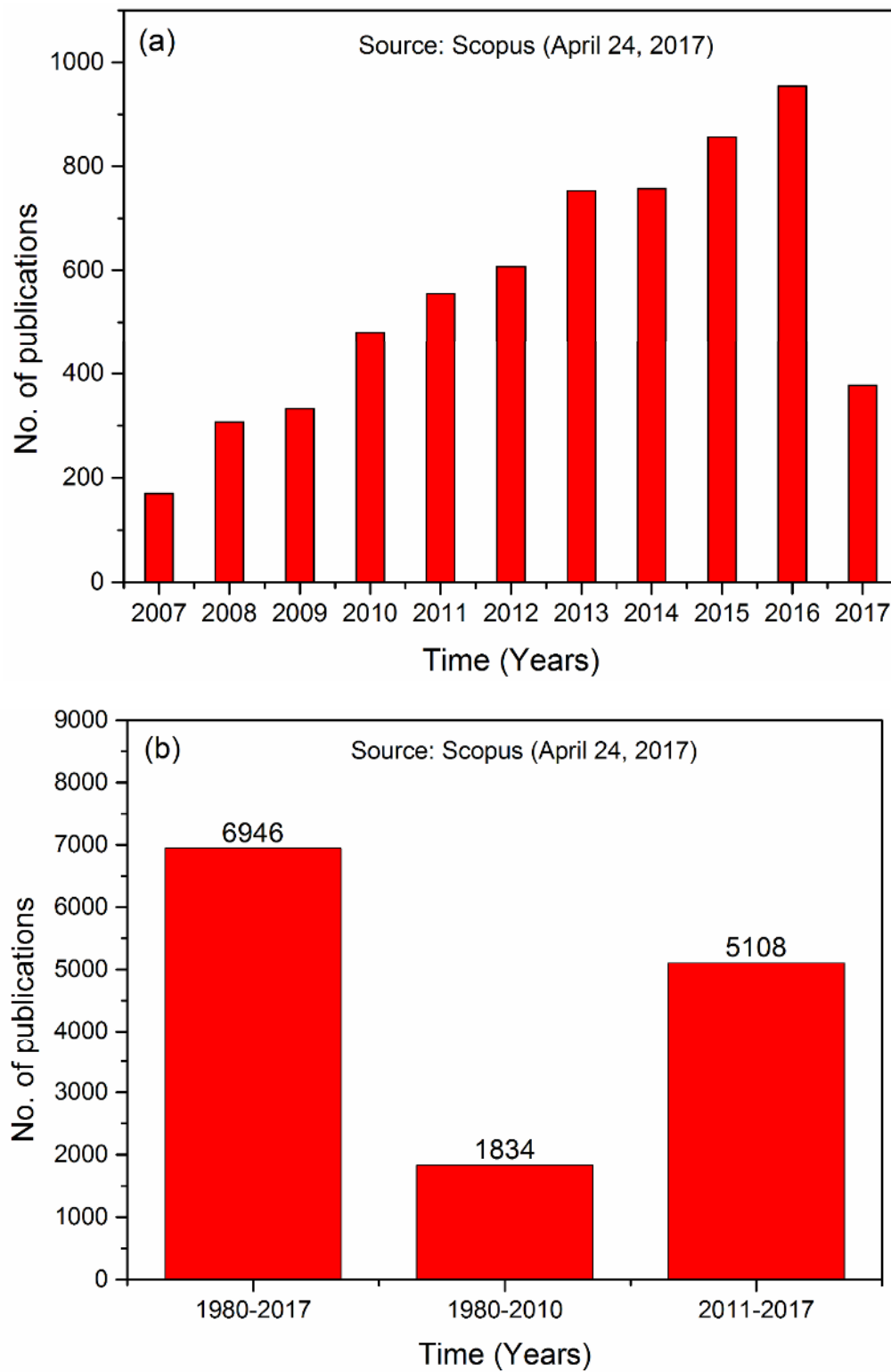
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3. Molecular Machinery for Extracellular Electron transfer mechanisms. Schematic image of the proposed EET of two metal respiring bacteria and their interactions with an electrode in a bioelectrochemical system. Dashed arrows indicate hypothetical electron flow and solid arrows indicate experimental proved electron flow. (A) Branched outer membrane cytochromes (OMCs) system of *Geobacter sulfurreducens*. Electrons can be transported between inner membrane, periplasm, outer membrane, and an electrode via a chain of cytochromes and menaquinones (MQ). Terminal OMCs can vary depending on the environmental conditions. (B) Unique Mtr-pathway and terminal reductases of *Shewanella oneidensis*. Quinones (Q) pass electrons to CymA or TorC, which transfer the electrons to terminal reductases or an MtrCAB complex. MtrCAB complex can interact with the electrode direct or via flavin molecules (FL). The figure is designed by Ms. Helena Reiswich and is a reproduction from [37], with permission from the publisher and the corresponding author.
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5. EcoBot II fully assembled with the wireless transmitter and temperature sensor on top [64].
6. (a) Block diagram of the telemetry system powered by the microbial fuel cell. (b) A sensor and a telemetry system powered by a microbial fuel cell [65].
7. Example 7-day time record of meteorological data transmitted from first generation BMFC-powered buoy [66].
8. (a) Pee power field trial in Glastonbury Music Festival, June 2015; (b) urinal assembly and MFC stack arranged in 12 modules [79].
9. Schematic diagram of the 90 L stackable baffled microbial fuel cell [82].
10. (a) Schematic diagram of an MFC. (b) Mechanism for MFC-based BOD monitoring. Increased BOD input provides more organic matter/fuel for the MFC, which in turn results in an increase in current output. (c) Mechanism for MFC-based toxicity monitoring. Increased toxin input will repress/inhibit the cell viability/metabolic activity, which directly reduces the current output [94].

11. Voltage responses of MFC-based biosensors to different samples. The figure shows the performance of four MFCs used to sense the addition of three samples (with different concentrations) in the anode chamber, resulting five shocks (a) to (e). **(a)** The MFC was injected with iron (non-toxic metal) of concentration 48 mg/L after 150 minutes of operation. The injection suddenly decreased the voltage from 121 mV to 67 mV. **(b)** The MFC was injected with chromium (acute toxin) of concentration 1 mg/L after 74 minutes of operation. After 134 minutes of the first fall (shock), the voltage decreased from the steady point (89 mV) to 81 mV. **(c)** After 74 minutes of operation, there was a steep fall in the voltage from 109 mV to 91 mV. **(d)** In another MFC, iron of concentration 1 mg/L was injected in the anode chamber after 30 minutes of operation. This low concentration decreased the voltage slightly from 121 mV to 118 mV, though higher concentration sharply decreased the voltage as mentioned earlier in (a). **(e)** The effect of carbon substrate was also sensed in the MFC, addition of 200 mg/L sodium acetate showed instant rise in the voltage from 102 mV to 114 mV after 2 minutes, which further increased to 122 mV after 4 minutes.

12. **(a)** The MFC array used for  $\text{Cu}^{2+}$  toxicity monitoring, **(b)** acidic toxicity monitoring [100].

13. The signal interference of an MFC sensor by the combined shock of biochemical oxygen demand (BOD) and toxicity in a continuous flow-through mode: **(a)** the MFC sensor operated with background acetate of 0.3 mM; **(b)** the MFC sensor operated with background acetate of 5 mM [103].

**Figure 1.**



(c)

Source: Scopus (April 24, 2017)

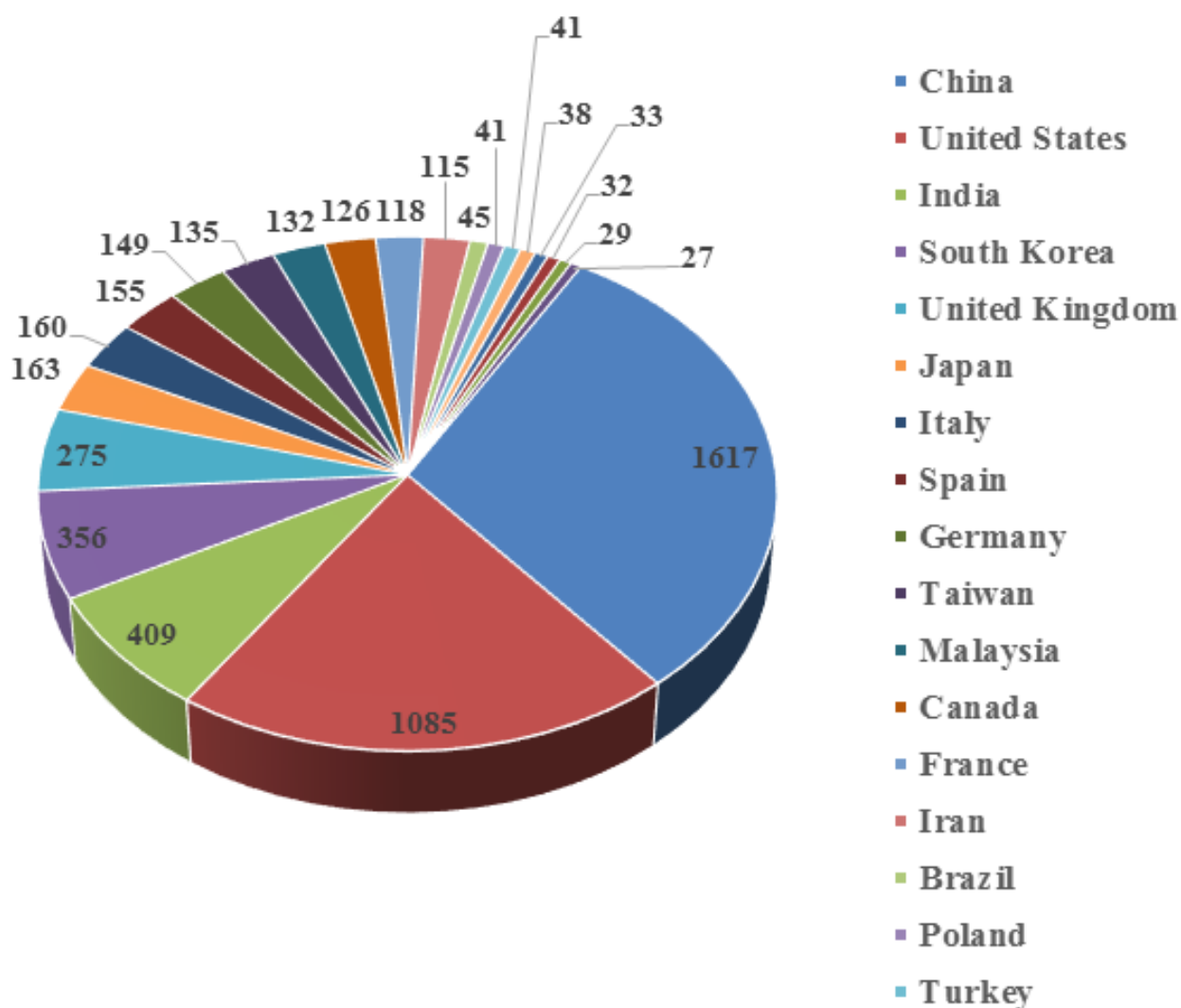


Figure 2.

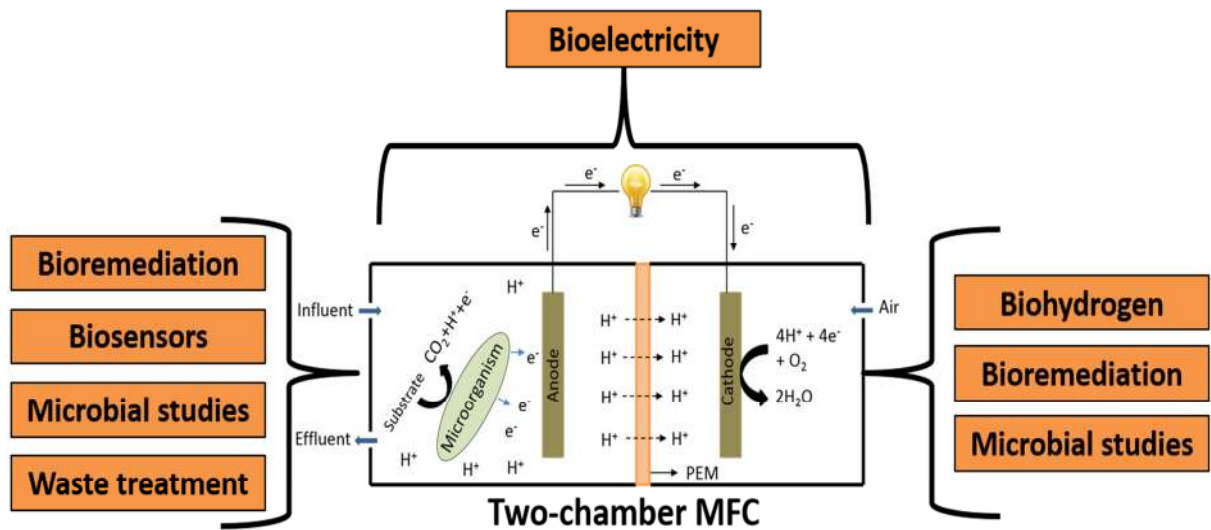
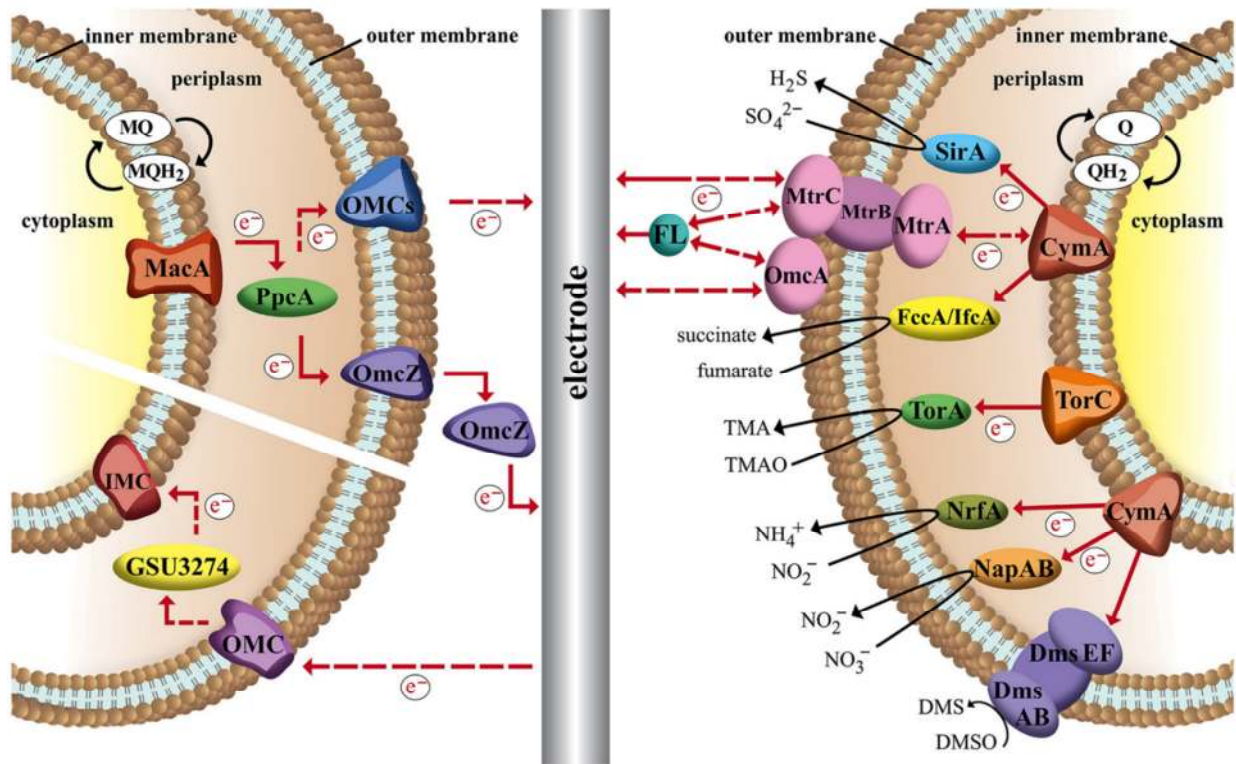
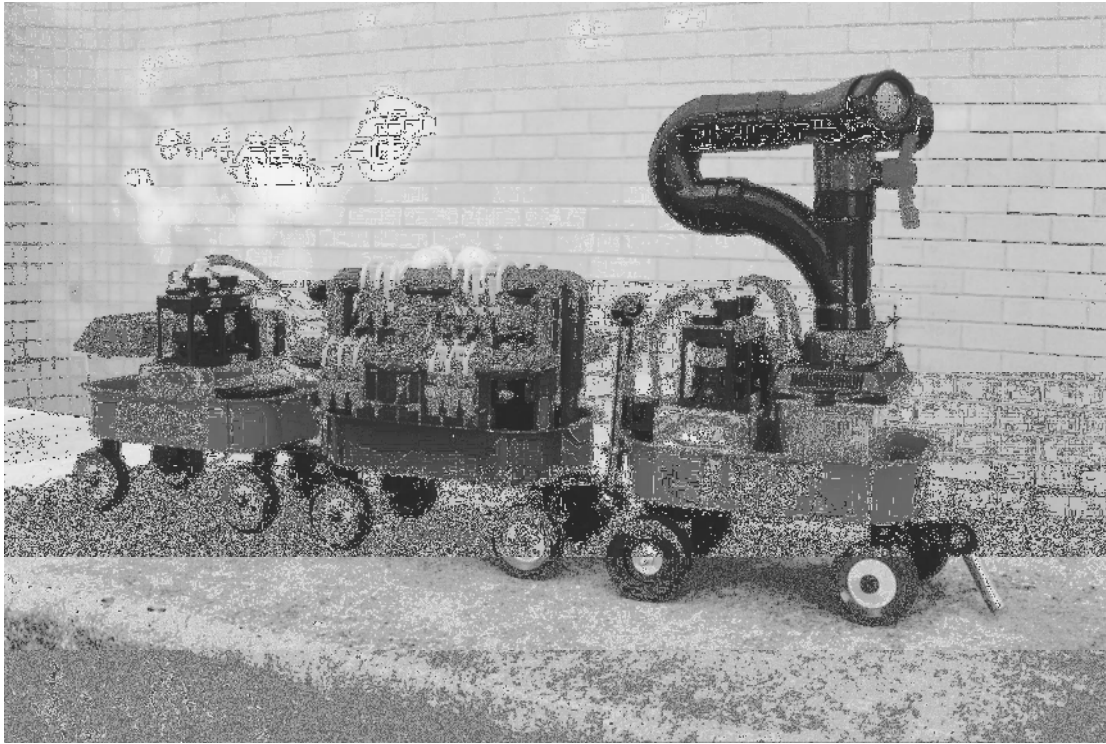


Figure 3.





**Figure 4.**



**Figure 5.**

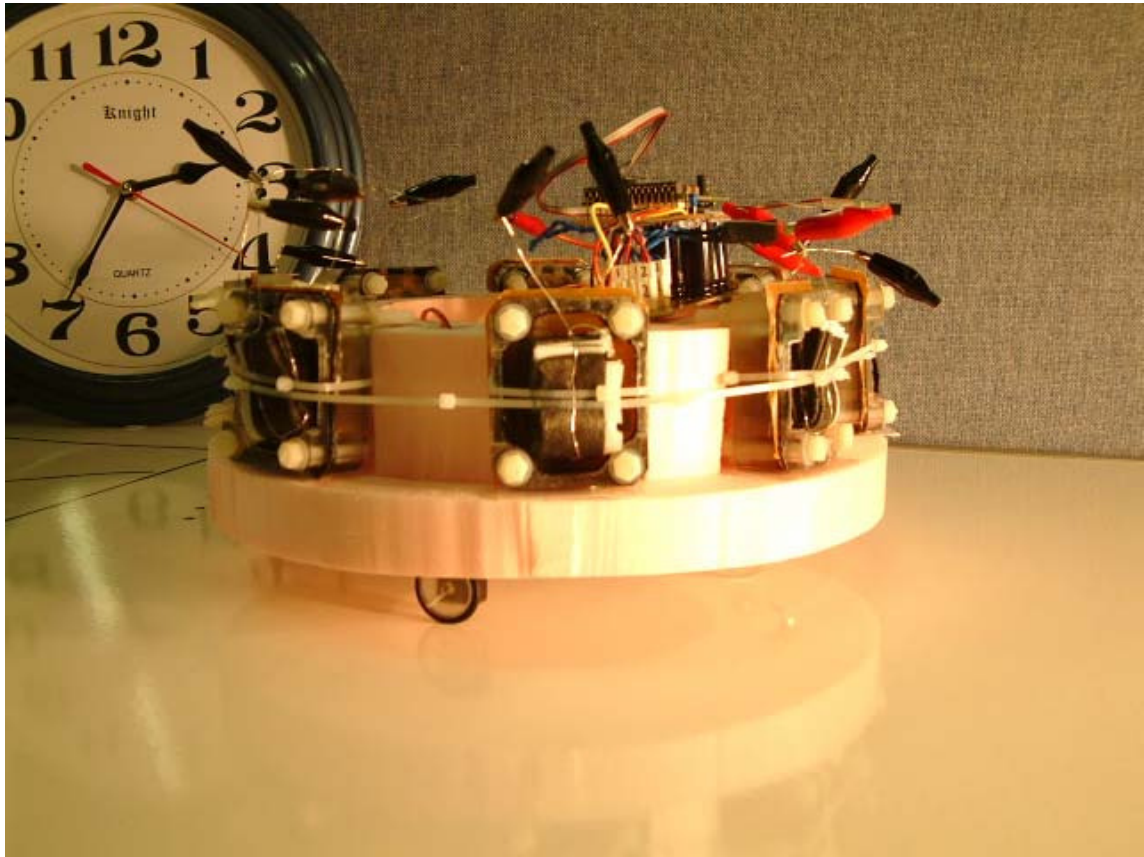
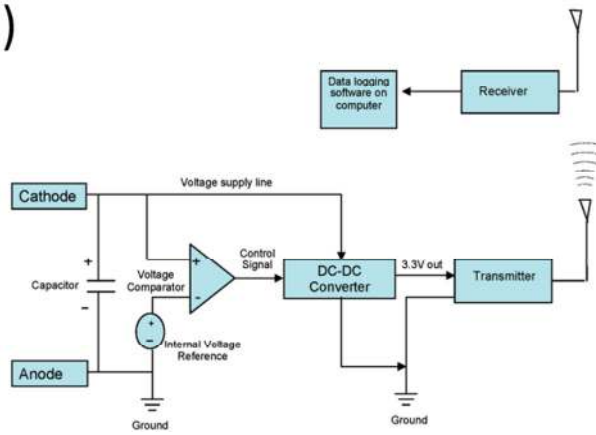


Figure 6.

(a)



(b)

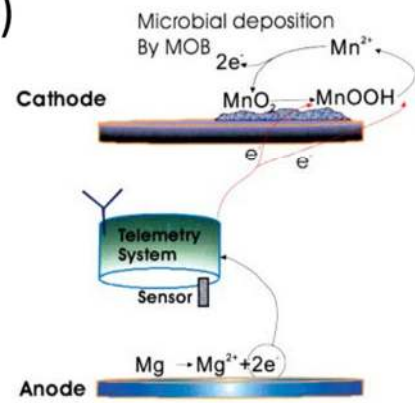


Figure 7.

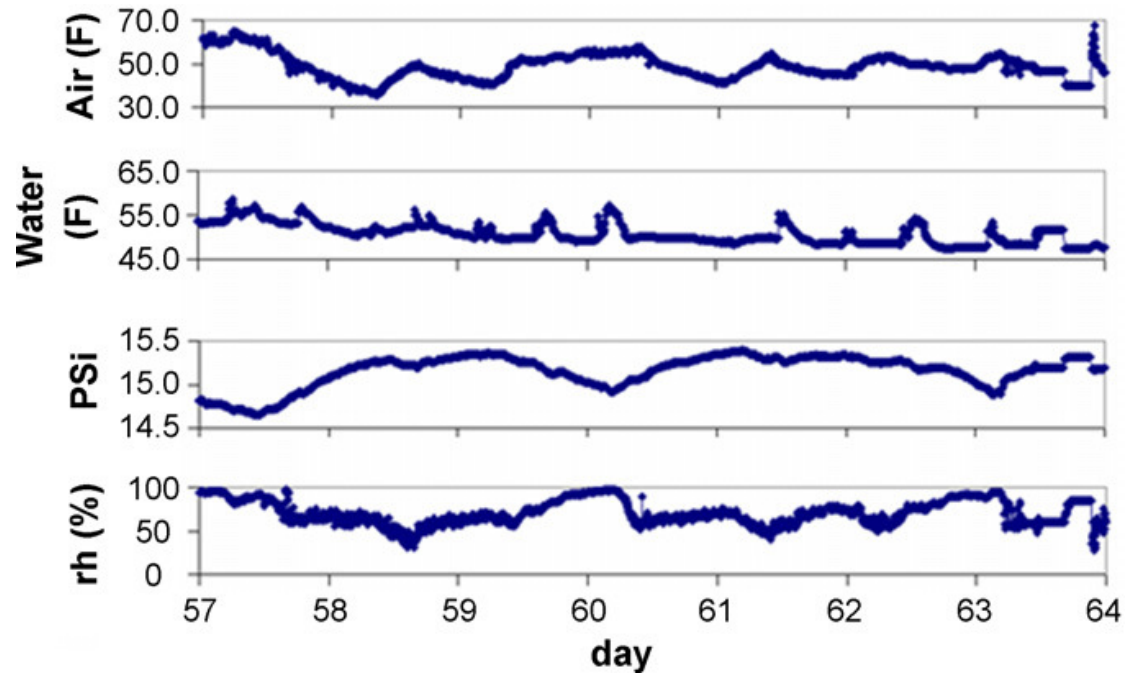


Figure 8.



Figure 9.

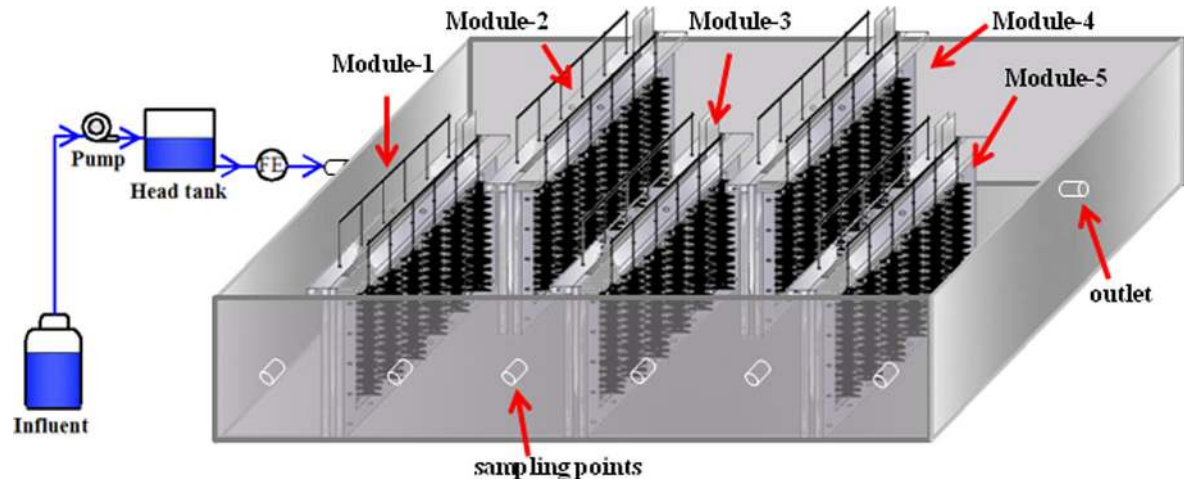


Figure 10.

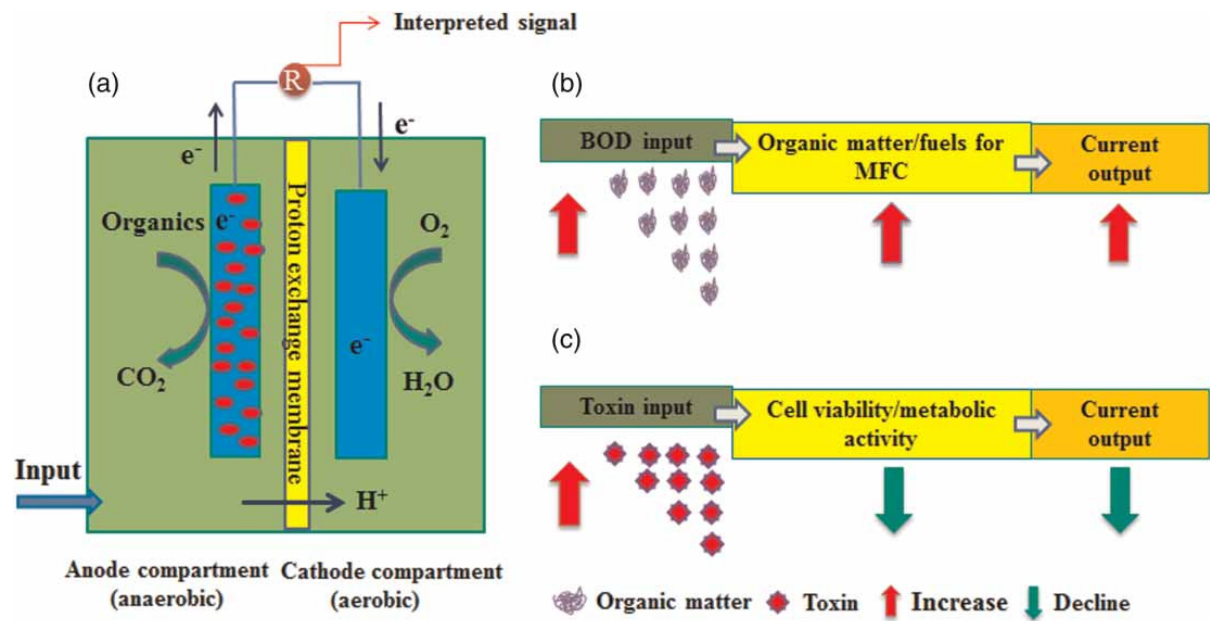


Figure 11.

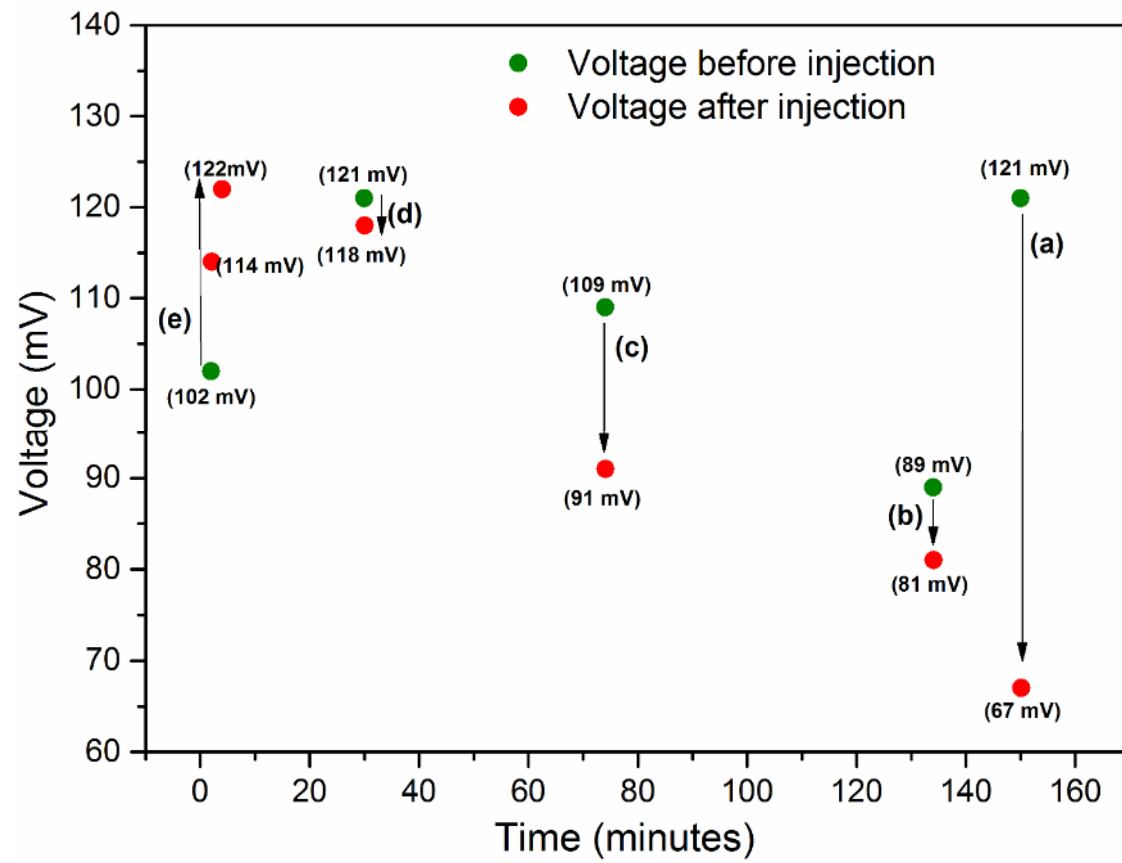




Figure 12.

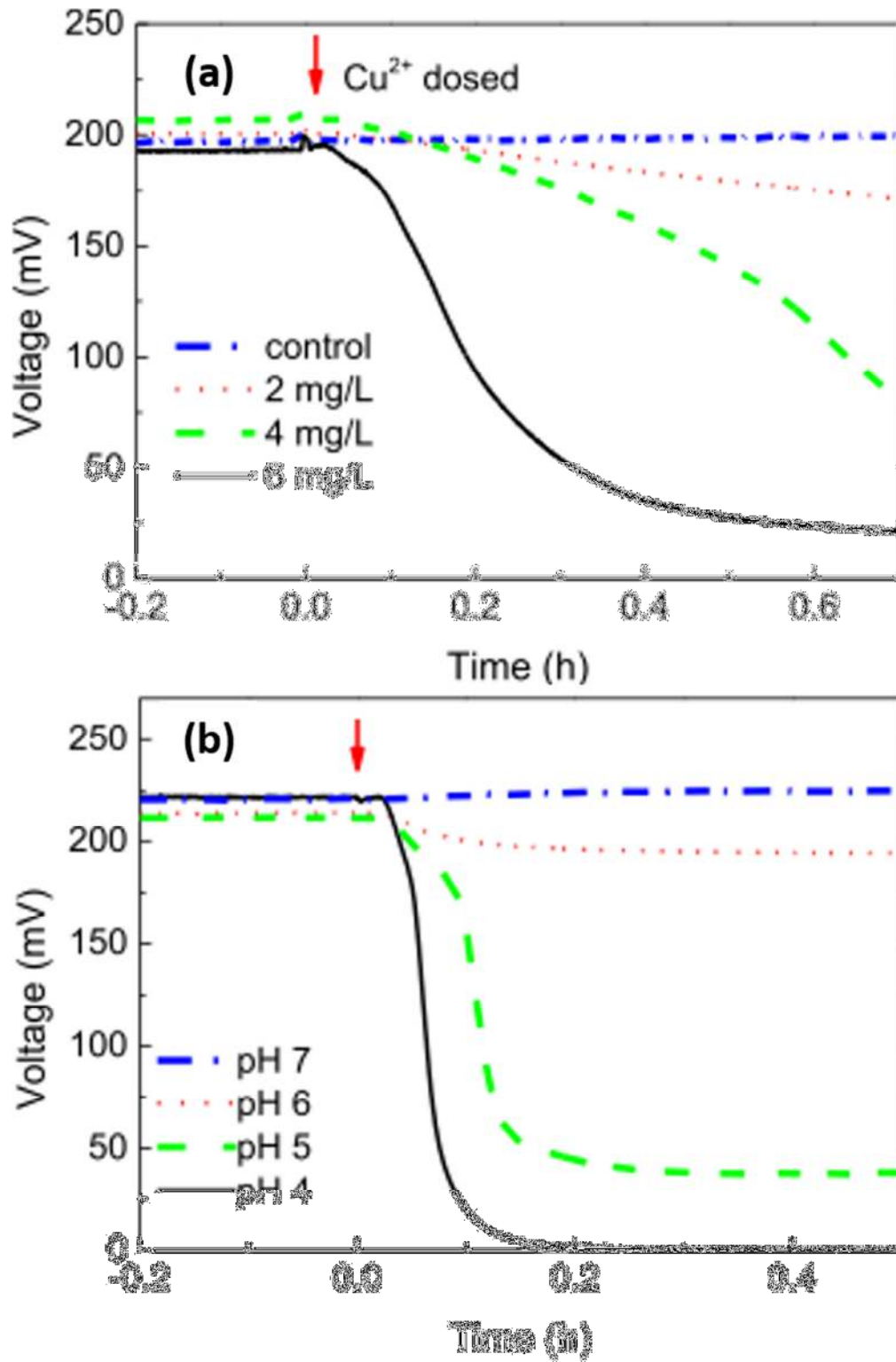


Figure 13.

