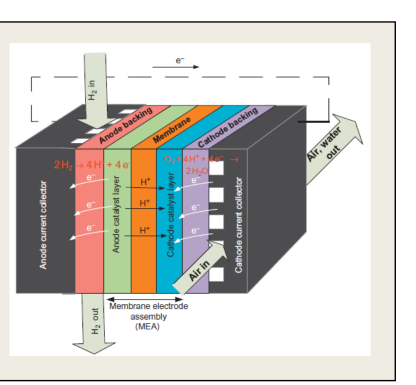


In a FC, fuel is fed continuously to the anode (negative electrode) and an oxidant (often oxygen in air) is fed continuously to the cathode (positive electrode). The electrochemical reactions take place at the electrodes to produce an electric current through an electrolyte, while driving a complementary electric current that performs work on the load.



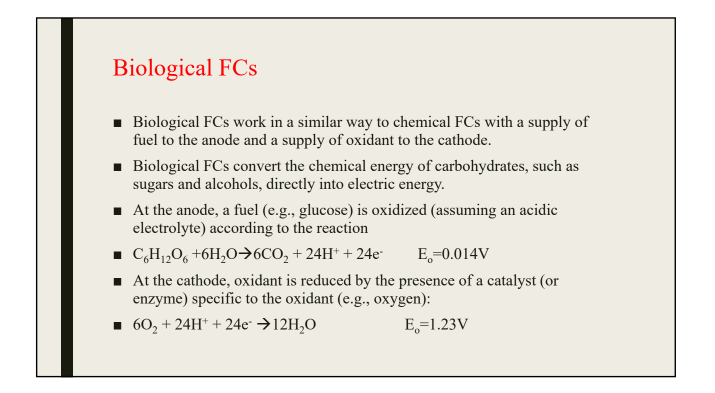
• At the anode of the FC, hydrogen gas ionizes, releasing electrons and creating H+ ion (protons), thereby releasing energy

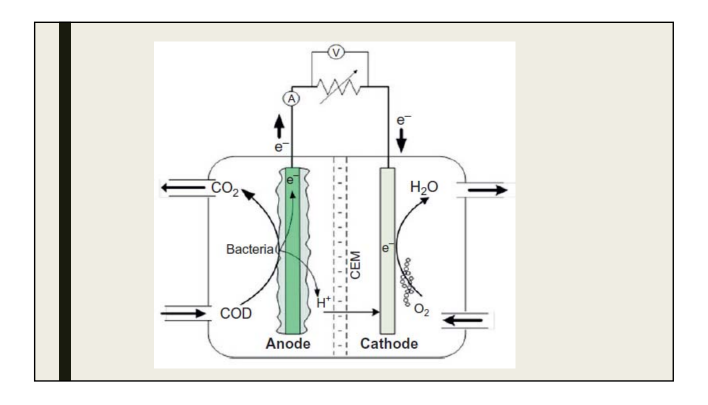
$2H_2 \rightarrow 4H^+ + 4e^-$

• At the cathode oxygen reacts with protons and electrons taken from the anode to form water

 $O_2 + 4H^+ + 4e^- \rightarrow 2H_2O$

- The electrons (negative charge) flow from anode to cathode in the external circuit and the H⁺ ions pass through the electrolyte.
- Importantly! the electrolyte should only allow proton transfer (or other ions in the case of other FC types) and not electron transfer (i.e., the electrolyte should be an electronic insulator). Otherwise the electrons would not pass around the external circuit and thus they would "short circuit" the cell and the function of the FC would be lost.





- The resultant electrochemical reaction creates a current as electrons and protons are produced from the oxidation of the fuel. The theoretical cell potentials for such reactions are similar to those of conventional FCs.
- The distinguishing feature, central to a biological FC, is the *use of the living organism* itself.
- In general a BioFCs functions in one of two ways, using biocatalysts, enzymes, or even whole cells.

1. The biocatalyst generates the fuel substrate for the electrochemical cell by a biocatalytic transformation or metabolic process. Thus the biocatalyst does not take part directly in electron transfer.

2. The biocatalyst participates in the electron transfer chain between the fuel and the anode.

- When enzymes are employed to achieve electrode activity, we have the so-called enzymatic biofuel cell;
- when microorganisms are responsible for the bioelectrocatalysis, we have the microbial fuel cell (MFC).
- Through the use of clean and renewable catalysts, MFCs provide a means to obtain renewable and sustainable energy and to treat wastewater, which is generally employed as the carbon source for the electrochemical system.

Types of biological FCs

■ In general, *microorganisms* can be used in *four ways* for producing electrical energy:

(i) To produce electrochemically active substances through fermentation or metabolism. The fuels are produced in separate reactors and pumped to the anode of a conventional FC, to generate electrical energy. In this configuration, the microbial bioreactor is kept separated from the FC; the system is not truly a BioFC.

(ii) The microbiological fermentation process proceeds directly in the anodic compartment of the FC.

(iii) The electron transfer mediators shuttle electrons between the microbial biocatalytic system and the electrode. The mediators accept electrons from the biological electron transport chain of the microorganisms and supply them to the anode of the biological FC.

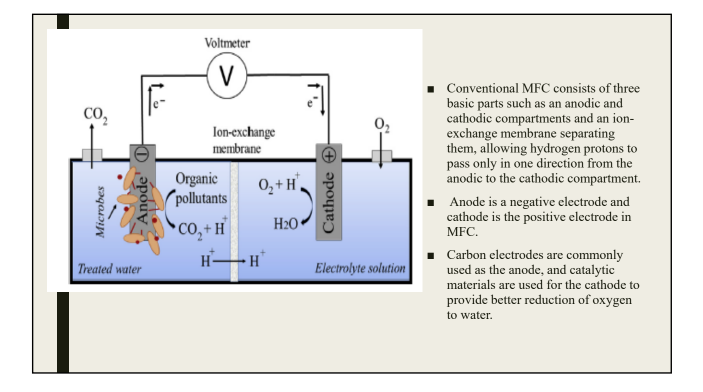
(iv) Metal-reducing bacterium, having cytochromes in the outer membrane that are able to directly communicate electrically with the electrode surface and create a mediatorless biological FC.

There are two basic types of biological FCs; namely,

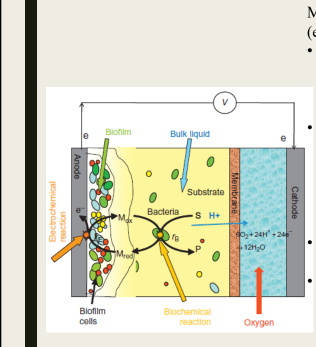
- microbial fuel cells (MFCs) and
- Enzymatic FCs.
- Electron mediators (relays) are used for the electrical connection of the biocatalyst and the electrode.
- Several methods have thus been used to functionalize the electrode surface with layers consisting of redox enzymes, electrocatalysts, and biocatalysts that promote electrochemical transformation at the electrode interface.

Microbial fuel cell (MFC)

- Microbial fuel cell (MFC) is a part of microbial electrochemical Technologies where cathode and anode combined with microorganisms serve for different purposes.
- MFC is an emerging biotechnological device that converts the chemical energy of organic matter into electricity by means of microorganisms.
- Similar to fuel cells MFC is theoretically highly efficient device able to produce electrical energy.
- However, in contrast to the fuel cell running on hydrogen or methanol, MFC can use wastewater simultaneously treating it and producing electric power.
- MFC is based on the metabolism of the bacteria and its ability to reduce redox active compounds.
- MFC is the only technology allowing generation of electricity directly from the solid and liquid organic wastes.
- Bacteria play the role of biocatalysts.



- Microorganisms that produce electricity are placed into the anodic compartment, wherein anaerobic conditions are maintained.
- The cathode is kept under aerobic conditions, which is provided by oxygen bubbling through the cathodic compartment or keeping the compartment and cathode exposed to the air.
- Microorganisms attached to the surface of anodic compartment (biofilm) receive a carbon source of energy (nutrients) in the form of wastewater, for example, that is necessary for them to grow and sustain life.
- Because bacteria are isolated in electrode compartment, the only way for them to survive is to process the organic substrate, which is fed for them and perform anaerobic respiration through the electrode.
- The principle of MFE operation is the detachment of electrons from the nutrient by microorganisms and electron transfer to the anode.
- Anode is connected with the cathode by a wire/electric circuit. Because of the difference of redox potentials, electrons start to move toward the cathode, where oxygen reduction occurs to form water.
- Electrons moving from the negative electrode to the positive generate the electric current produced by MFC.
- When electrons are detached from the nutrient, hydrogen protons are formed in the anodic compartment. The generated hydrogen ions pass from the anodic compartment through the ion-exchange membrane into the cathodic one where they are combined with oxygen to form water.

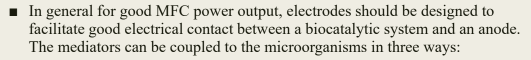


Microbes in the anode chamber oxidize fuel (electron donor) generating electrons and protons.

- The generation of current is due to the microorganisms, which transfer electrons from a reduced electron donor to an electron acceptor at a higher electrochemical potential.
- Anode-respiring bacteria, which have accumulated as an anode biofilm, carry out an oxidation (half-cell) reaction of organic matter, producing protons (one proton for every electron) and thus an electrical current from biomass.
- Carbon dioxide may eventually be obtained as an oxidation product.
- Electrons and protons are consumed in the cathode chamber, reducing oxygen to water and generating electricity.

Anode microbial behavior

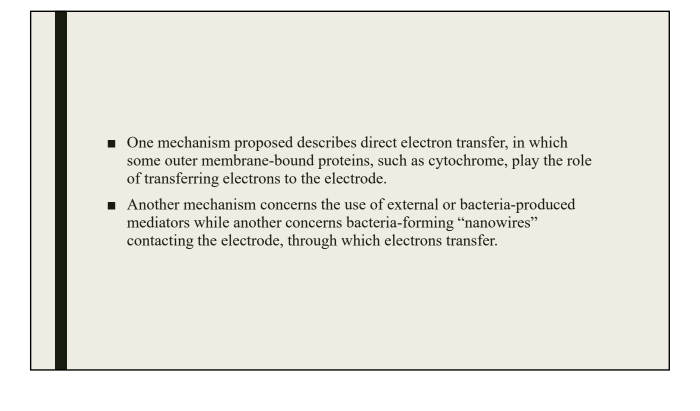
- The MFC is reliant upon the organism's own generation of electrons from the electron transport chain.
- The electron transport chain involves the cell wall of a microorganism, where there exist proteins responsible for energy generation.
- In the case of aerobic respiration, the core reaction is oxidation of an energy-rich compound, such as glucose, to allow the reduction of nicotinamide adenine dinucleotide (NAD+) to NADH by the electrons donated from the glucose. In turn NADH is oxidized and its electrons are transferred to adenosine triphosphate (ATP).
- However the cells can be connected (wired) to the electrode surface using low molecular weight redox species, called mediators. The mediators assist the shuttling of electrons between the intracellular bacterial space and electrode.
- A variety of organic compounds have been used as mediators for electron transfer between bacteria and electrodes, including thionine and organic dyes.

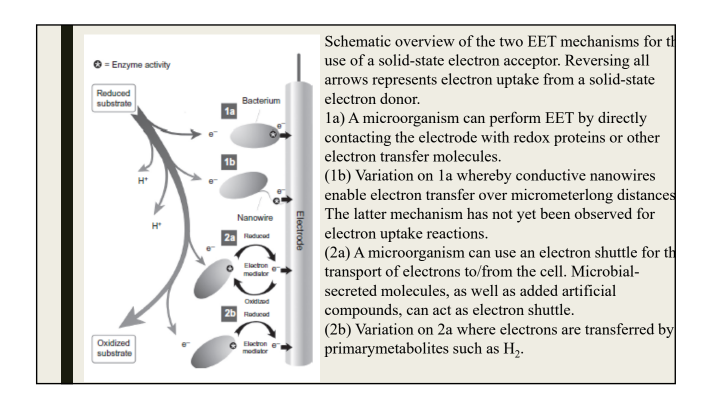


- (i) diffusion mediator shuttling between the microbial suspension and the anode surface,
- (ii) diffusion mediator shuttling between the anode and microbial cells covalently linked to the electrode. The microbial cells can be covalently linked to the electrode surface having – COOH groups, through amino groups of the microbial membrane resulting in the formation of amide bond. Organic reagents such as carbodiimide and acetyl chloride can be used to link the microbial cells to the surface, and
- (iii) mediator adsorbed on the microbial cells providing electron transport from the cells to the anode.

MFC bacteria

- The behavior of microorganisms in MFCs can involve several complex processes.
- This complexity will in particular be most apparent in the use of mixed microbial communities that would be present in most waste streams.
- Which species and how they populate the anode in MFCs will have a great bearing on both waste treatment (COD/BOD removal) as well as power generation.
- In addition to microorganisms that can transfer electrons to the anode, the presence of other organisms may influence the MFC performance, both positively and negatively.
- Mixed species may result in faster feed substrate removal without generation of power.





Terminology available in literature to describe microorganisms interacting with electrodes

■ Electrochemically active bacteria (EAB)/ electroactive bacteria (EAB)

Microorganisms able to donate electrons to or accept electrons from an electrode via a direct contact mechanism or self-secreted mediators.

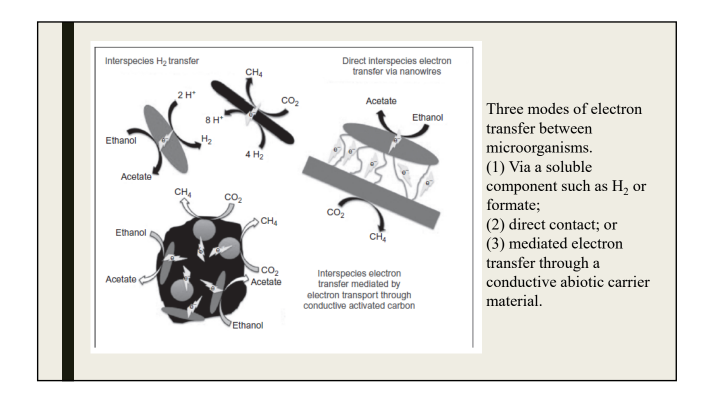
Electron donating microorganisms

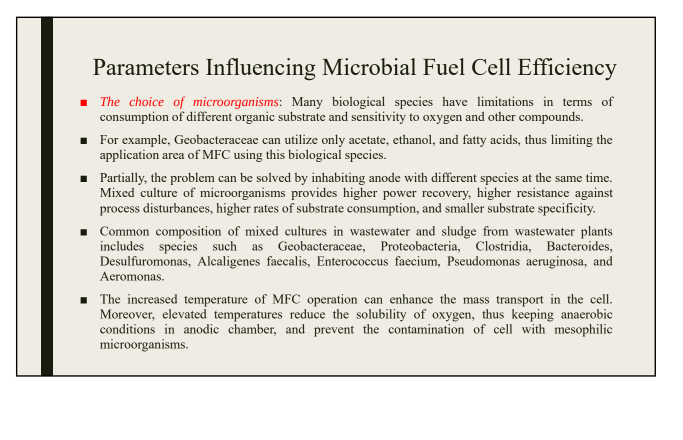
Anode-reducing bacteria (ARB): Microorganisms able to donate electrons to an anode

Anode-respiring bacteria (ARB): Microorganisms able to conserve energy by respiration with an anode as electron acceptor

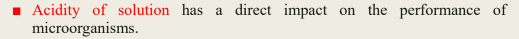
Electron accepting microorganisms

Cathode oxidizing bacteria: Microorganisms able to draw electrons from a cathode





- The distance between electrodes:
- Too close placement of electrodes in undivided MFCs (membrane free MFCs) leads to power reduction due to interruption of bacteria activity by oxygen generated at the cathode.
- In such cells, a cloth separator for example could reduce the effect of oxygen crossover.
- Electrodes should not be placed at a distance where oxygen interaction with anode is minimum, however not far beyond that point, because resistance of solution will rise too high and decrease power production efficiency.



- The best conditions for microbial growth are in the range of neutral pH.
- However, slightly alkaline conditions are also acceptable and even can improve pollutant degradation and power generation due to the presence of free OH ions, which improve the conductivity of solution.
- High salinity of wastewater also contributes to the greater removal efficiencies for the same reason.

Anode Materials

- Aside from other factors affecting the MFC performance, the anode material is a deciding factor for overall performance of MFCs because its
- composition,
- morphology, and
- surface properties

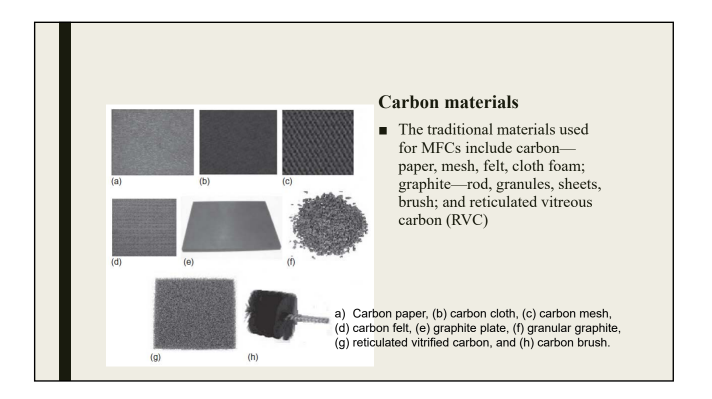
directly affect

- microbial attachment,
- electron transfer, and
- substrate oxidation.

Ideal anode materials require *high electrical conductivity, enhanced biocompatibility, high chemical and physical stabilities of electrodes,* and *low-cost materials.*

- Material of electrode plays an important role in efficiency of the process.
- For example anodes with higher specific surface area, i.e., surface area per volume of the electrode, such as graphite brush, carbon felt, and foam provides more stable power generation comparing to flat electrodes.
- Moreover, electrodes with high specific surface area are less prone to fouling.
- This can be explained by the fact that oxygen is consumed by bacteria on the surface of such electrodes and bacteria, which are located inside thick electrodes and stay in anaerobic conditions, thus processing more organic matter and transferring electrons.
- Cathode materials should have low overpotential toward oxygen reduction to water and have surface area significantly greater than surface area of anodes.

- In the anode compartment of MFC,
- microorganisms adhere to the anode,
- generate electrons by substrate oxidation, and
- then transfer the electrons to anode.
- The surface characteristics of the anode directly affect the microbial adhesion and therefore the electrical performance of MFC.
- Increased specific surface area and more positive charge on electrode surface, which interact with negatively charge bacteria, aided the microbial adhesion and electron transfer on the anode surface.
- Thus, anode modification is an effective strategy for enhanced characteristics of the electrode.



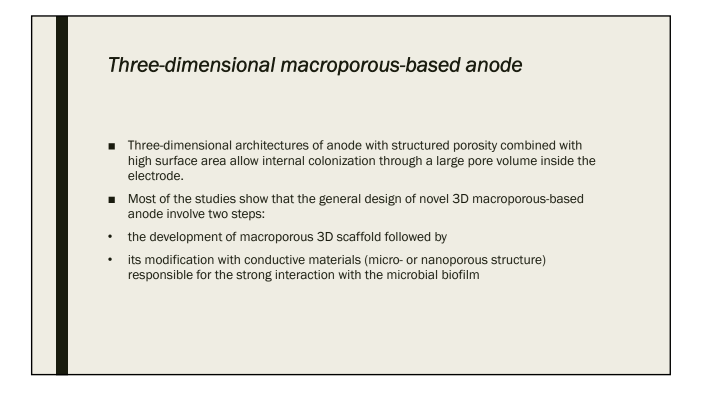
- One of the main limitations of using traditional carbon materials as MFC anodes is related to the electrocatalytic activity for the electrode microbial reactions, because the pores or spaces became clogged with the biofilm and lose efficiency.
- Furthermore, in some cases, the low specific area, lack of durability, brittleness, and high cost of carbonaceous materials inhibit the application of these electrodes in large-scale MFCs.

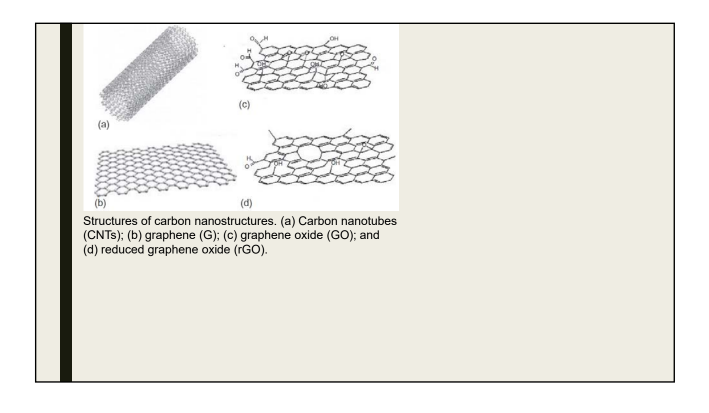
Metal materials

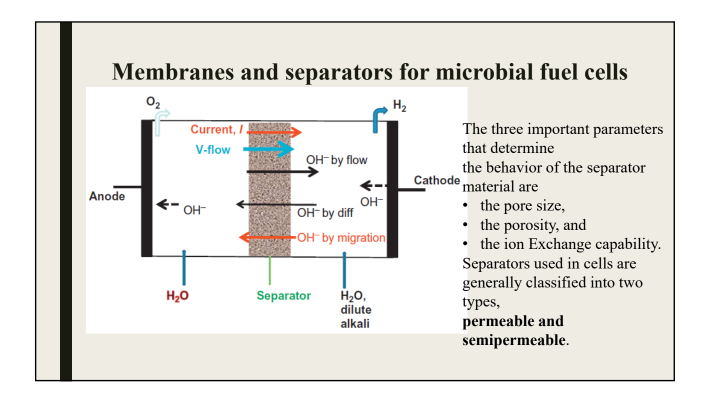
- Several metal materials have been used as MFC anode due to the higher conductivity of metal materials over carbon materials.
- The noncorrosive requirement of MFC anode limited the choice of metal materials used for MFC applications.
- Stainless steel and titanium are common metals proposed as potential anode materials in several studies. Several studies investigated stainless steel as MFC anode, because of the better mechanical properties expected for long-time operation and scale-up applications compared with carbon electrode
- However, copper was investigated as anode for MFC. MFCs with Cu anodes showed that copper is an unsuitable anode material, due to corrosion and its toxicity to microorganisms, even if it initially showed high current generation comparable to that of carbon electrode.
- Noble metals as electrode materials in MFC reduce the internal resistance of the cell, but the high cost and week adhesion of bacteria prevent their wide-scale integration in MFC

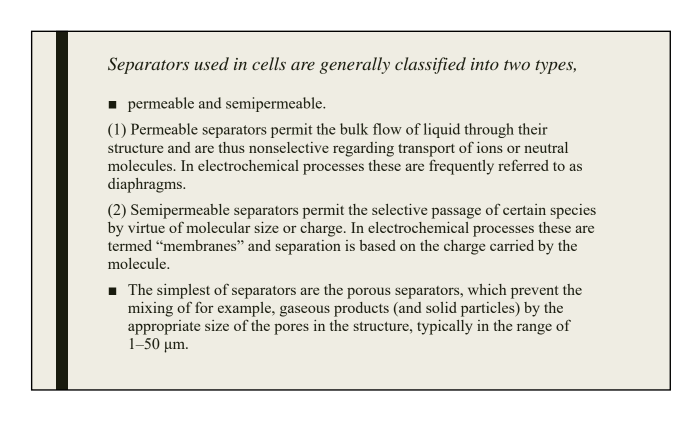
Anode materials	MFC efficiency	References	
	MFC efficiency 40 W m ⁻² based on anode area; the higher power density must either be a calculation error, or due solely to copper corrosion with the solution used in the tests (Zhu and Logan, 2014) 2 ± 0.5 mW m ⁻² based on anode area 23 mW m ⁻² based on anode area Maximum current density of stainless steel electrode was around 2.4 A m ⁻² vs. 1.1 A m ⁻² of graphite electrode (constant potential chronoamperometry) Maximal current densities reached under constant polarization at -100 mV vs. SCE: 3.1 A m ⁻² for plain stainless steel, 5.9 A m ⁻² for plain graphite, and 8.2 A m ⁻² for stainless steel grid, each of 25 cm ² projected area No date of power density of Ti because daily current densities of Ti was far low ~0.1 A m ⁻² than that of flat graphite, roughened graphite, and Pt-coated Ti with values between 2 and 2.5 A m ⁻² 0.8 mW m ⁻² of Al electrode vs. 1.8 mW m ⁻² carbon fiber cloth electrode 0.2 mW m ⁻² of Al electrode vs. 1.8 mW m ⁻² carbon fiber cloth electrode	References Kargi and Eker (2007) Zhu and Logan (2014) Dumas et al. (2007) Dumas et al. (2008) Erable and Bergel (2009) ter Heijne et al. (2007) Ouitrakul et al. (2007) Ouitrakul et al. (2007)	Metal materials applied as anode MFC and their performance in MFCs
Stainless	0.1 mW m ⁻² vs. 1.8 mW m ⁻² carbon fiber cloth	Ouitrakul	
steel	electrode	et al. (2007)	
Pt	78 W m ⁻² for Pt nanoelectrode was connected to the wire covered with nanoparticle Ti vs. 35.08 W m ⁻² of simple electrode Pt to the simple Ti wire	Torabiyan et al. (2014)	
Au	Maximum power density of 0.15 µW cm ⁻²	Qian et al. (2009)	
	Maximum power density of 95 $\mu W~cm^{-2}$ vs. results from Qian et al. (2009)	Choi and Chae (2013)	

Anode materials	te materials	References	
Mn ⁴⁺ -graphite anode	10.2 mW m ⁻² electrode was obtained with a Mn ⁴⁺ graphite anode 787.5 mW m ⁻² maximum power density of Mn ⁴⁺ -graphite anode vs. 5.32 mW m ⁻² of NR-covalently linked woven graphite anode and 0.65 mW m ⁻² of woven graphite anode with anaerobic sewage sludge as biocatalyst; and 91 mW m ⁻² maximum power density of Mn ⁴⁺ -graphite anode vs. 1.20 mW m ⁻² of NR-covalently linked woven graphite anode and 0.44 mW m ⁻² of woven graphite anode with <i>E. coli</i> as biocatalyst	Park and Zeikus (2002) Park and Zeikus (2003)	Composite materials applied as anode MFC and their efficiency as MFC anode
Graphite-ceramic composite anodes containing Mn ²⁴ and Ni ²⁺ Graphite paste with an incorporated Sb(V) complex Aluminum-alloy mesh	Maximum power density of \sim 105 mW m ⁻² of composite anode vs. \sim 20 mW m ⁻² of graphite anode Maximum power density of \sim 115 mW m ⁻² of composite anode vs. \sim 25 mW m ⁻² of graphite anode 2966 mW m ⁻² maximum power density	Lowy et al. (2006) Lowy and Tender (2008) Chen et al.	
composite carbon cloth electrode	obtained with composite anode 22.4 mW m ⁻² of aluminum-alloy mesh anode	(2013)	
Composite graphite/PTFE electrodes	760 mW m ⁻² of graphite/PTFE composite electrode with optimized PTFE content (e.g., 30%, w/w) for <i>E. coli</i> -catalyzed MFC vs. Park and Zeikus (2003)	Zhang et al. (2007)	









Semipermeable membranes: ion-exchange membranes

 The main properties required of ion-exchange membranes for them to be successful in technical processes are

1. low electrical resistance, to minimize the membrane internal resistance (IR) losses,

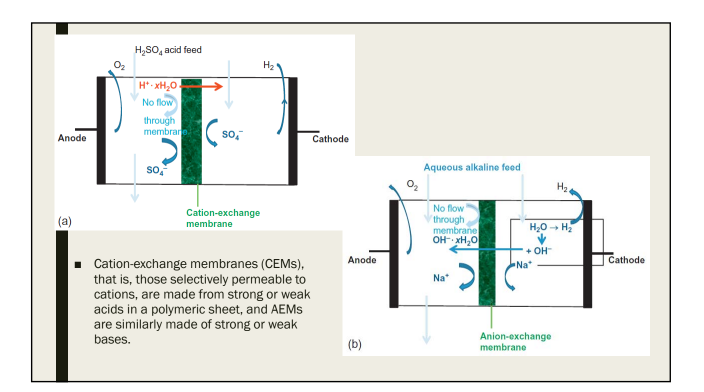
2. high permselectivity; that is, it should be highly permeable for an ion of one charge, but should be highly impermeable to coions, and to nonionised molecules and solvents,

3. good mechanical and dimensional stability,

4. good chemical stability, over a wide pH range and in the presence of oxidizing agents,

5. good operating characteristics; that is, it should be capable of operation over a wide range of current densities and under varying conditions of temperature, current density, pH, etc.

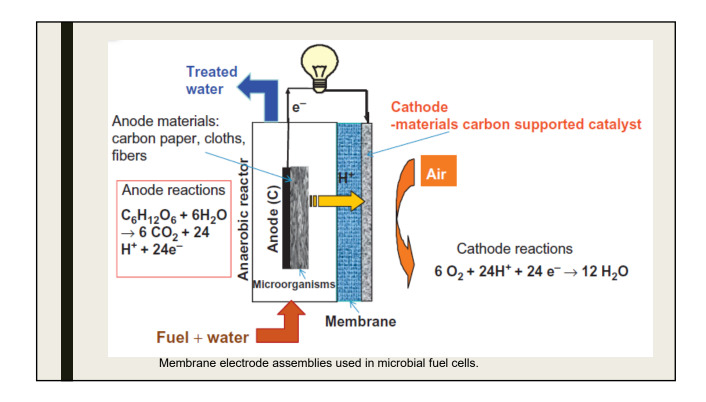
- The properties of ion-exchange membranes are determined by two parameters; that is, the basic polymer matrix and the type and concentration of the fixed ionic group.
- The electric properties and the selectivity of ion-exchange membranes are determined mainly by the type and the concentration of the fixed ionic charges in the polymer matrix.
- There are a series of different cationic and anionic groups that can be introduced into a basic polymer matrix



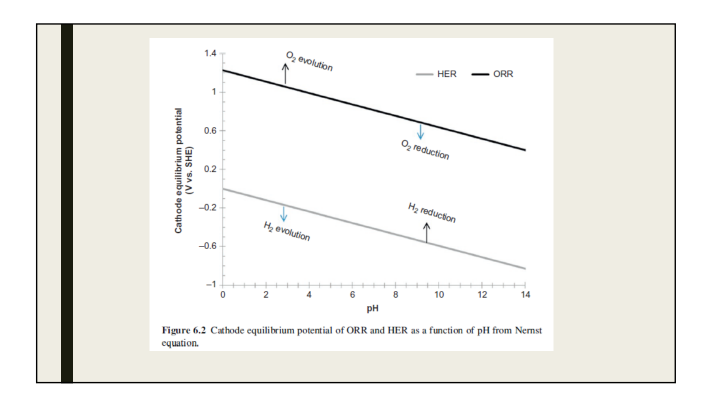
Cation		Anion function group	
Sulfonate,	Strongly	Quaternary ammonium, -N(CH ₃) ₃ ⁺ OH ⁻	Strongly
$-SO_3^- H^+$	acidic	Tri-ethylammonium	basic
		Trialkylbenzyl ammonium	
Carboxylate,	Weakly	Trimethyl amine, -N(CH ₃) ₂	Weakly
-COOH	acidic	Triethyl amine	basic
Phosphonate		DABCO (4-diazabicyclo[2.2.2]octane)	
		(triethylenediamine)	
		Dimethyl sulfide, CH ₃ –S–CH ₃	
		Diethyl sulfide	
		N, N, N', N'-Tetramethyl-1,6-hexanediamine	

The sulfonic acid groups are strong acids, completely dissociated over nearly the entire pH range, and carboxylic acid groups are weak acids and undissociated at values of pH of 6.

AEMs mostly use the quaternary ammonium (QA) group in the polymer matrix leading to a strongly basic membrane. Tertiary, secondary, and primary amines and to a lesser extent phosphonium and sulfonium groups, are used as less strongly basic functional groups.



Anode react		$E^{\circ\prime}$ at pH SHE)	17 (V	vs.	Catho de reaction	E [∞] at pH 7 SHE)	(V vs.		
				Microbial fuel cells					
AQDSH2/AQ HS ⁻ /SQ4 ²⁻		-0.18 -0.217	-0.15		Oxygen/H ₂ O ClO ₃ ⁻ /Cl ⁻	+0.82* +0.81	0.9		
HS ⁻ /S		-0.27			ClO ₄ ⁻ /Cl ⁻	+0.81	L		
Methane/HCt Oil and greas C ₈ H ₁₆ O/HCC	ase	-0.25 -0.29	-0.20	-	Fe ³⁺ /Fe ²⁺ NO ₃ ⁻ /N ₂	+0.77 +0.75	[‡] ₀₃ +		
Acetate/HCO Propionate/H Ethanol/HCC	03 ⁻ HCO3 ⁻	-0.29 -0.29 -0.33	-	-	C ₂ H ₄ Cl ₂ /C ₂ H ₄ C ₂ Cl ₄ /C ₂ HCl ₃ C ₂ HCl ₃ /cis-	+0.739 +0.574 +0.550	÷ 0.7		
Waste water C 10 H 140 3 N/F	/HCO ₃ -	-0.33	-0.25		C2H2Cl2 Oxygen/H2O	+0.51 ^b	- 0.E		
Protein C ₁₆ H HCO ₃ ⁻ Lactate/HCO		-0.333			C ₂ H ₃ Cl/C ₂ H ₄ NO ₃ ⁻ /N ₂ ⁻	+0.45	4 F		
Pynivate/HO		-0.37	-0.30 -		Cr2O72-/Cr3+	+0.365°	+ 05 m		
Methanol/HC	CO3-	-0.39			NO3-/NH4+	+0.36	ode		
H_2/H^+		-0.41	-0.35	node	[Fe(CN) ₆] ³⁻ / [Fe(CN) ₆] ⁴⁻	+0.36	F° Cathode		
Glucose/HC0	-	-0.43	-0.35	E " A	NO ₂ ⁻ /NO	+0.35	- U.4 - M		
NADH/NAD		-0.32			NO ₂ ⁻ /NH ₄ ⁺	+0.34	室		
Neutral red _{ex}	ad/neutral	-0.33			O2 (gas)/H2O2	+0.260	† 03		
Cysteine/cyst Methyl violo methyl violo	ogen _{md} /	-0.34 -0.44	-0.40	-	SeO42 ⁻ /Se HSeO3 ⁻ /Se	+0.322 +0.26	ł		
			-0.45	-	O ₂ (gas)/H ₂ O ₂ Fumarate ² -/ succinate ² -	+0.26 +0.03	• 0.2		
							• 0.1		
			-0.50				t.		



Other performance indicators	Metric parameter (to enable system comparisons)
Electrochemical	
Volumetric power density (W m ⁻³)	Typically based on the anodic chamber empty bed volume. This measure is often used to compare dissimilar systems, but extrapolation from laboratory-scale reactors to very large systems may be misleading
Power density (W m ⁻²)	Typically refers to power per specific electrode (anode or cathode) or membrane-projected areas
Electrochemical losses	A number of electrochemical techniques may be utilized to investigate and quantify system losses Polarization plots can be used to determine activation, ohmic and concentration polarization Charge transfer processes may be further investigated using Tafel plots or EIS to investigate charge and mass transport-related impedances

Other performance indicators	Metric parameter (to enable system comparisons)
ternal impedance (Ω) ternal capacitance	Internal resistance is indicative of the ohmic losses that may be incurred and allows for impedance matching with external loads for maximum power transference. However, in nonsteady operation, capacitive and inductive reactances may assume significance Capacitance may be identified by step or other
	perturbations and by using EIS in conjunction with assumed equivalent circuit models. Electrode type and biofilm structure determines anodic capacitive behavior and potentially influences power production

Operational	
Loading rate/hydraulic retention time	Indicative of the capacity of the system to facilitate treatment to specified limits
Energy conversion efficiency	A consideration of conversion efficiencies in terms of total influent energy content and the energy content of the consumed COD can be useful, e.g., where BES are to be considered as part of integrated treatment process
Reactor recovery time	Robustness of the biocatalytic system to environmental and chemical shocks, and subsequent process recovery, may often be conveniently expressed as a time constant
Biofilm biomass concentration	Amount of active biofilm has a direct impact on biocatalytic activity and its development may be monitored by measuring biomass concentrations, i.e., through protein and DNA
Mean time to failure (MTTF), mean time between maintenance (MTBM)	The robustness of scaled-up systems should be assessed by measures such as these, considering statistical measure of time to/between failures/ maintenance during operation
Costs, operating and capital	Financial metrics associated with CAPEX (capital expenditure) and OPEX (operational expenditure)
Product yields	BES associated with product formation, e.g., MEC production of hydrogen, needs to be assessed in terms of product yields

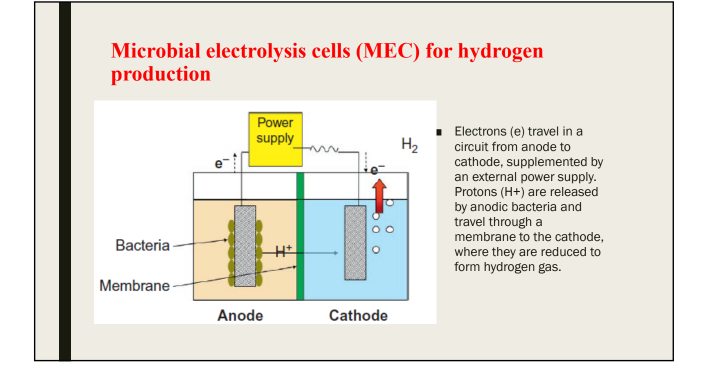
Applications of microbial electrochemical and fuel cells

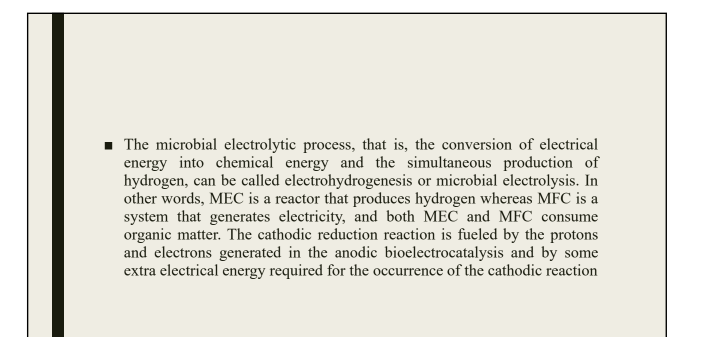
Wastewater treatment

Oxidation reactions (anode)	
Glucose : $C_6H_{12}O_6 + 12H_2O \rightarrow 6HCO_3^- + 30H^+ + 24e^- E^\circ = -0.429 V vs. SHE$	
Glycerol: $C_3H_8O_3 + 6H_2O \rightarrow 3HCO_3^- + 17H^+ + 14e^- E^\circ = -0.289V$ vs. SHE	
Malate : $C_4H_5O_5^- + 7H_2O \rightarrow 4H_2CO_3 + 11H^+ + 12e^- E^\circ = -0.289V$ vs. SHE	
Sulfur: $HS^- \rightarrow S^0 + H^+ + 2e^- E^\circ = -0.230 \text{ V vs. SHE}$	
Reduction reactions (cathode)	
$O_2 + 4H^+ + 4e^- \rightarrow 2H_2O_E^\circ = +1.230V$ vs. SHE	
$O_2 + 2H^+ + 2e^- \rightarrow H_2O_2 E^\circ = +0.269V \text{ vs. SHE}$	
$NO_3^- + 2e^- + 2H^+ \rightarrow NO_2^- + H_2O E^\circ = +0.433 V \text{ vs. SHE}$	
$NO_2^- + e^- + 2H^+ \rightarrow NO + H_2O_E^\circ = +0.350V$ vs. SHE	
$NO + e^- + H^+ \rightarrow 1/2N_2O + 1/2H_2O$ $E^\circ = +1.175V$ vs. SHE	
$1/2N_2O + e^- + H^+ \rightarrow 1/2N_2 + 1/2H_2O~{\it E^\circ} = +1.355V$ vs. SHE	
$2NO_3^- + 12H^+ + 10e^- \rightarrow N_2 + 6H_2O_E^\circ = +0.734V$ vs. SHE	
$Fe^{3+} + e^- + H^+ \rightarrow Fe^{2+} + 1/2H_2O \ E^\circ = +0.773V \text{ vs. SHE}$	
$MnO_2 + 4H^+ + 3e^- \rightarrow Mn^{2+} + 2H_2O E^\circ = +0.602 V vs. SHE$	

Acetate	Dual	0.08 mA/cm ²	Min et al. (2008)	
Ethanol	Dual	0.025 mA/cm ²	Kim et al. (2007a,b)	
Glucose	Dual	283 mW/m ²	Rahimnejad et al. (2011)	
Synthetic acid-mine	Dual	0.064 mA/cm ²	Cheng et al. (2007)	
drainage water				Energy receivery from yorigue
Synthetic wastewater	Dual	0.086 mA/cm ²	Mohan et al. (2008)	Energy recovery from various
Synthetic wastewater	Single	0.017 mA/cm ²	Aldrovandi et al. (2009)	
Synthetic wastewater	Dual	0.008 mA/cm ²	Jadhav and Ghangrek ar	wastewater substrates in MFCs
			(2009)	
Synthetic	Dual	0.005 mA/cm ²	Kargi and Eker (2007)	
WW-molasses and urea				
Domestic wastewater	Single	1.7 and 3.7 W/	Liu et al. (2004) and Liu and	
		m ³	Logan (2004)	
Domestic wastewater	Dual	0.06 mA/cm ²	Wang et al. (2009)	
Domestic wastewater	Single	42 mA/m ²	Nimje et al. (2012)	
Primary effluent	Single	13 mA/m ²	Ishii et al. (2012)	
Influent from AD	Dual	$58 \pm 2 \text{ W/m}^3$	Aelterman et al. (2006)	
Effluent for AD	Dual	$42 \pm 8 \text{ W/m}^3$	Aelterman et al. (2006)	
Alcohol distillery	Dual	1000 mA/m ²	Ha et al. (2012)	
Agriculture wastewater	Single	13 mA/m ²	Nimje et al. (2012)	
Bad wine wastewater	Dual	3.8 W/m ³	Rengasamy and Berchmans	
			(2012)	
Brewery wastewater	Single	0.2 mA/cm ²	Feng et al. (2008)	
Brewery wastewater	Single	0.18 mA/cm ²	Wen et al. (2009)	
Brewery and bakery	Single	10 mA/m ²	Velasquez-Orta et al. (2011)	
Cheese whey	Dual	42 mA/m ²	Stamatelatou et al. (2010)	
Chocolate industry	Dual	0.302 mA/cm ²	Patil et al. (2009)	
wastewater		2		
Dairy wastewater	Single	25 mA/m ²	Velasquez-Orta et al. (2011)	
Dairy wastewater	Single	5.7 W/m ³	Ayyaru and Dharmalingam	
			(2011)	
Dairy/food wastewater	Single	15 mA/m ²	Nimje et al. (2012)	
Distillery wastewater	Single	245.3 mA/m ² 0.004 mA/cm ²	Mohanakrishna et al. (2012)	
Farm manure	Single		Scott and Murano (2007)	
Food processing	Dual	0.05 mA/cm ²	Oh and Log an (2005)	
wastewater				
Food waste	Single	207 W/m ³	Goud and Mohan (2011)	
Food (composite) waste	Single	107.9 W/m ³	Goud et al. (2011)	
Hospital wastewater	Dual	8±5 W/m ³	Rabaey and Verstraete	
The second s	Dual	1411 111-3	(2005)	
Hospital wastewater Human feces		14±1 W/m ³ 70.8 W/m ³	Aelterman et al. (2006)	
Human feces wastewater	Dual	/0.8 W/m	Du et al. (2011)	

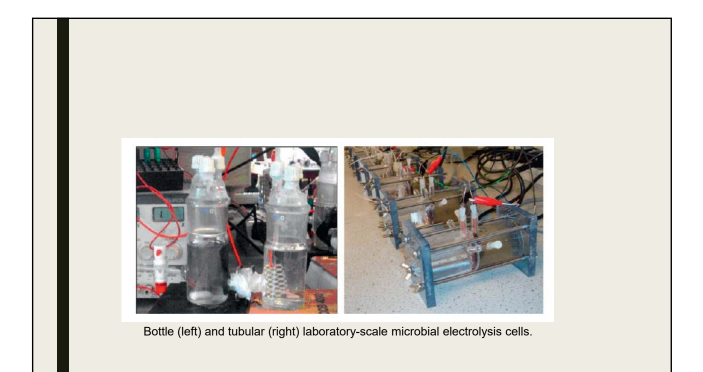
Meat processing	Single	0.115 mA/cm ²	Heilmann and Logan (2006)
wastewater			
Palm oil effluent with acetate	Dual	622 mW/m ²	Jong et al. (2011)
Paper recycling wastewater	Single	0.25 mA/cm ²	Huang and Logan (2008)
Paper wastewater	Single	125 mA/m^2	Velasquez-Orta et al. (2011)
Paper wastewater	Single	8 mA/m^2	Nimje et al. (2012)
Pharmaceutical	Single	177.36 W/m ³	Velvizhi and Mohan (2011)
Protein-rich waste water	Dual	0.008 mA/cm^2	Liu et al. (2009)
Real urban wastewater	Dual	0.018 mA/cm ²	Rodrigo et al. (2010)
Sewage sludge	Tubular	73 mA/m ²	Yuan et al. (2012)
Starch processing wastewater	Single	0.09 mA/cm ²	Lu et al. (2009)
Swine wastewater	Single	0.015 mA/cm ²	Min et al. (2005)

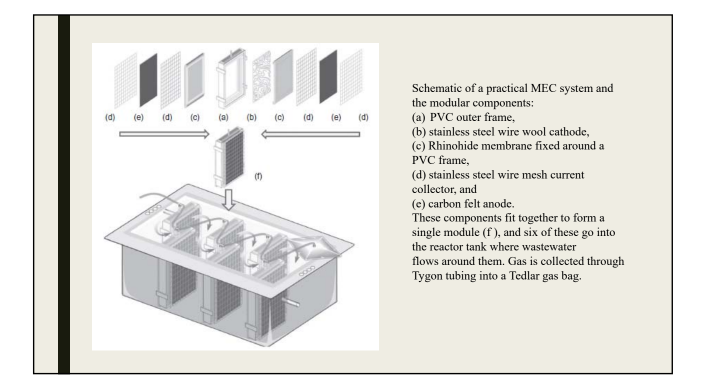




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- MECs make use of exoelectrogens (electrochemically active microorganisms) to harvest electrons (and protons) from organic matter in an anaerobic anode chamber.
- These electrons are transferred in a circuit from anode to cathode, producing an electrical current.
- In MECs, protons (H⁺ ions) are transferred (under acidic conditions) via a membrane from the anode to an anaerobic cathode (unlike in an MFC) and therefore oxygen is not present and not reduced to form water.
- The H⁺ ions can be used to form hydrogen at the cathode (or other useful products such as hydrogen peroxide), yet this will not occur spontaneously due to the energy required to drive the chemical reaction.
- Cathodic reaction under acidic conditions, $2H^+ + 2e \rightarrow H_2$
- Or under alkaline conditions, $2H_2O+2e \rightarrow H_2 + 2OH$





Advantages

- MECs are fully anaerobic. With an MEC both electrodes can be fully submerged in a tank, whereas with an MFC the cathode must be exposed to air either through aeration of the associated electrolyte or by direct exposure as an air cathode.
- MECs produce higher-value products. MECs produce products that are higher in value than the electricity produced by MFCs. It has been estimated that the financial value of hydrogen produced in an MEC could range from 3 times to almost 10 times that of electricity produced in an MFC.

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