AQUEOUS LIQUID FEED ORGANIC FUEL CELL USING SOLID POLYMER ELECTROLYTE MEMBRANE

Inventors: Subbarao Surampudi, Glendora; Sekharipuram R. Narayanan, Altadena; Eugene Vamos, La Canada; Harvey A. Frank, Encino; Gerald Halpert, Pasadena; George A. Olah, Beverly Hills; G. K. Surya Prakash, Hacienda Heights, all of Calif.

Assignees: California Institute of Technology; University of Southern California, both of Pasadena, Calif.

ABSTRACT

A liquid organic fuel cell is provided which employs a solid electrolyte membrane. An organic fuel, such as a methanol/water mixture, is circulated past an anode of a cell while oxygen or air is circulated past a cathode of the cell. The cell solid electrolyte membrane is preferably fabricated from Nafion™. Additionally, a method for improving the performance of carbon electrode structures for use in organic fuel cells is provided wherein a high surface-area carbon particle/Teflon™-binder structure is immersed within a Nafion™-methanol bath to impregnate the electrode with Nafion™. A method for fabricating an anode for use in a organic fuel cell is described wherein metal alloys are deposited onto the electrode in an electro-deposition solution containing perfluorooctanesulfonic acid. A fuel additive containing perfluorooctanesulfonic acid for use with fuel cells employing a sulfuric acid electrolyte is also disclosed. New organic fuels, namely, trimethoxymethane, dimethoxymethane, and trioxane are also described for use with either conventional or improved fuel cells.

49 Claims, 13 Drawing Sheets

OTHER PUBLICATIONS


Primary Examiner—Stephen Kalafut

Attorney, Agent, or Firm—Fish & Richardson P.C.
IMMERSE THE CARBON ELECTRODE STRUCTURE IN 1% SOLUTION OF NAFION IN METHANOL FOR ABOUT 5 MINUTES TO ACHIEVE IMPREGNATION OF THE NAFION INTO THE ELECTRODE TO A LOADING OF 0.1 - 0.5 mg/cm².

REMOVE ELECTRODE FROM SOLUTION AND DRY IN VACUUM.

FIG. 7
**FIG. 11**

![Graph showing potential (volts vs. NHE) vs. current density (mA/cm²).]

**FIG. 8**

![Graph showing potential (volts vs. NHE) vs. current density (mA/cm²).]
PREPARE CARBON ELECTRODE STRUCTURES FROM A MIXTURE OF 200 m²/g HIGH SURFACE AREA CARBON PARTICLES AND TEFLON BINDER (15%) APPLIED TO A FIBER-BASE CARBON PAPER.

PREPARE A BATH OF HYDROGEN HEXACHLOROPALTINATE AND POTASSIUM PENTACHLOROQUORUTENIUM WITH A METAL ION CONCENTRATION IN THE RANGE OF 0.01-0.05 M DISSOLVED IN 1 M SULFURIC ACID.

ADD PERFLUOROOCTANESULFONIC ACID TO BATH WITH A CONCENTRATION IN THE RANGE OF 0.1-1.0 g l⁻¹

POSITION THE CARBON ELECTRODE IN THE BATH ALONG WITH A PLATINUM ANODE.

APPLY A VOLTAGE BETWEEN THE CARBON ELECTRODE AND THE ANODE FOR ABOUT 5-10 MINUTES TO ACHIEVE ELECTRODEPOSITION OF PLATINUM-RUTHENIUM TO A LOADING OF ABOUT 5 mg/cm².

REMOVE CARBON ELECTRODES FROM BATH AND WASH IN DEIONIZED WATER.

FIG. 9
FIG. 12

FIG. 13
FIG. 14

FIG. 15
**FIG. 16**

**FIG. 17**
**FIG. 18**

CELL VOLTAGE (VOLTS) vs. CURRENT DENSITY (mA cm²)

**FIG. 19**

POTENTIAL vs. NHE vs. CURRENT DENSITY (mA/cm²)
**FIG. 20**

Current Density (mA/cm²) vs. Potential vs. NHE

**FIG. 21**

Cell Voltage (Volts) vs. Current Density (mA/cm²)
BACKGROUND OF THE INVENTION

1. Technical Field

The invention generally relates to organic fuel cells and in particular liquid feed organic fuel cells.

2. Background Art

Fuel cells are electrochemical devices that convert the chemical energy of the fuel to electrical energy. Two types of fuel cells are generally known: direct and indirect.

A. Direct oxidation fuel cells

In a direct oxidation fuel cell, the fuel is oxidized directly at the anode of the fuel cell. The electrode structures for use in a liquid feed fuel cell are typically porous, allowing the liquid fuel to be fed into the cell. Direct oxidation fuel cells do not require a reforming stage, making them attractive for applications where fuel purity is not a concern.

B. Indirect or reformer fuel cells

In an indirect fuel cell, the fuel is first converted to hydrogen and/or carbon monoxide at a reformer, which is then oxidized at the anode of the fuel cell. This type of fuel cell is more efficient in converting chemical energy to electrical energy due to the higher combustion temperature available at the anode.

In direct methanol fuel cells, a specific problem is parasitic shunt currents, which can result in degradation of the fuel cell performance and affect the cost of the fuel cell. A parasitic shunt current is a current that flows around the intended path of current, bypassing the intended components.

3. Description of the Invention

The invention provides a liquid feed fuel cell that does not require sulfuric acid as an electrolyte. This is achieved by using a proton-conducting polymer electrolyte instead of sulfuric acid, which eliminates corrosion concerns and reduces the overall cost of the fuel cell.

4. Brief Description of the Drawings

Figure 1 illustrates a liquid feed direct methanol fuel cell using a solid polymer electrolyte. Figure 2 shows a liquid feed organic fuel cell using a solid polymer electrolyte.

5. Detailed Description of the Invention

The invention described herein was made in performance of work under a NASA contract, and is subject to the provisions of Public Law 96-517 (35 USC 202) in which the Contractor has elected to retain title.

In view of the aforementioned problems associated with using sulfuric acid as an electrolyte, it would be desirable to provide a liquid feed fuel cell that does not require sulfuric acid as an electrolyte.

In addition to the improvements in operational characteristics of the liquid feed fuel cell, the conventional method of fabricating high-surface-area electro-catalytic electrodes for use such fuel cells also needs to be improved. The existing method of fabrication of fuel cell electrodes is a fairly time-consuming and expensive procedure. Specifically, electrode fabrication requires that a high-surface-area carbon-supported alloy powder be initially prepared by a chemical method which usually requires about 24 hours.

Once prepared, the carbon-supported alloy powder is combined with a Teflon™ binder and applied to a carbon fiber-based support to yield a gas diffusion electrode. To volatilize impurities arising out of the Teflon™ binder and to obtain a fibrous matrix of Teflon™, the electrodes are heated to 200°-300° C. During this heating step, oxidation and sintering of the electrocatalyst can occur, resulting in a reduced activity of the surface of the electrode. Thus, the electrodes often require re-activation before use.

Also electrodes produced by conventional methods are usually of the gas-diffusion type and cannot be effectively used in liquid feed type fuel cells as the electrode is not adequately wetted by the liquid fuel. In general, the structure and properties of a fuel cell oxidation electrode (anode) for use in liquid feed type fuel cells are quite different from the gas/vapor feed fuel cells such as the hydrogen/oxygen fuel cell. The structure of anode in a liquid feed fuel cell should be very porous and the liquid fuel solution should have a pH of 5 or below. Carbon dioxide that is evolved at the fuel electrode should be effectively released from the zone of reaction. Adequate wetting of the electrodes is a major problem for liquid fuel cells—only those which use a sulfuric acid electrolyte.

As can be appreciated, it would be desirable to provide improved methods for fabricating electrodes, particularly for use in liquid fuel feed cells. It is also desirable to devise methods for modifying electrodes, originally adapted for gas-feed fuel cells, for use in liquid fuel feed cells.

In addition to improving the liquid feed fuel cell itself and for providing improved methods for fabricating the electrodes of fuel cell, it would be desirable to provide new effective fuels as well. In general, it is desirable to provide liquid fuels which undergo clean and efficient electrochemical oxidation within the fuel cell. The efficient utilization of organic fuels in direct oxidation fuel cells is, in general, governed by the ease by which the organic compounds are anodically oxidized within the fuel cell. Conventional organic fuels, such as methanol, present considerable difficulties with respect to electro-oxidation. In particular, the electro-oxidation of organic compounds such as methanol involves multiple electron transfer and is a very hindered process with several intermediate steps. These steps involve...
3 dissociative adsorption of the fuel molecule to form active surface species which undergo relatively facile oxidation. The ease of dissociative adsorption and surface reaction usually determines the facility of electro-oxidation. Other conventional fuels, such as formaldehyde, are more easily oxidized, but have other disadvantages as well. For example, formaldehyde is highly toxic. Also, formaldehyde is extremely soluble in water and therefore crosses over to the cathode of the fuel cell, thus reducing the performance of the fuel cell. Other conventional organic fuels, such as formic acid, are corrosive. Furthermore, many of the conventional organic fuels poison the electrodes of the fuel cell during electro-oxidation, thus preventing sustained operation. As can be appreciated, it would be desirable to provide improved fuels, particularly for use in liquid feed fuel cells, which overcome the disadvantages of conventional organic fuels, such as methanol, formaldehyde, and formic acid.

SUMMARY OF THE INVENTION

A general object of the invention is to provide an improved direct type liquid feed fuel cell. One particular object of the invention is to provide a direct type liquid feed fuel cell which does not require a sulfuric acid electrolyte. Another particular object of the invention is to achieve adequate wetting of electrodes for use in liquid feed fuel cells. Yet another particular object of the invention is to provide an improved method for wetting electrodes for use in fuel cells having sulfuric acid electrolytes. Still another particular object of the invention is to provide improved fuels for use in liquid feed fuel cells.

The object of providing an improved liquid feed direct fuel cell which does not require a sulfuric acid electrolyte is achieved in part by using a solid polymer electrolyte membrane in combination with a battery-type anode that is porous and is capable of wetting the fuel. In the improved liquid feed fuel cell, a battery-type anode structure and a cathode are bonded to either side of the solid polymer proton-conducting membrane forming a membrane-electrode assembly. A solution of methanol and water which is substantially free of sulfuric acid is circulated past the anode side of the assembly.

A solid polymer membrane is used, in part, because such membranes have excellent electrochemical and mechanical stability, high ionic conductivity, and can function both as an electrolyte and as a separator. Also, the kinetics of electro-oxidation of methanol and electro-reduction of air or oxygen are more facile at an electrode/membrane-electrolyte interface as compared to an electrode/sulfuric acid interface. The use of the membrane permits operation of the fuel cell at temperatures as high as 120°C. Since the fuel and water solution is substantially free of sulfuric acid, there is no need for expensive corrosion-resistant components in the fuel cell and its accessories. Also the absence of conducting ions in the fuel and water solutions, which can exist when a sulfuric acid electrolyte is employed, substantially eliminates the possibility of any parasitic shunt currents in a multi-cell stack.

The solid polymer electrolyte is preferably a proton-conducting cation-exchange membrane, such as the perfluorinated sulfonic acid polymer membrane, Nafion™. Nafion™ is a co-polymer of tetrafluoroethylene and perfluorovinyl ether sulfonic acid. Membranes of modified perfluorinated sulfonic acid polymer, polyhydrocarbon sulfonic acid and composites of two or more kinds of proton exchange membranes can also be used.

The anode is preferably formed from high surface area particles of platinum-based alloys of noble and non-noble metals. Binary and ternary compositions can be used for the electro-oxidation oxidation of organic fuels. Platinum-ruthenium alloy, with compositions varying from 10–90 atom percent of platinum, is the preferred anode electrocatalyst for the electro-oxidation of methanol. The alloy particles are either in the form of fine metal powders, i.e., "unsupported", or are supported on high surface area carbon material.

Conventional fuel cell anode structures (gas diffusion type) are not suitable for use in liquid feed type organic/air fuel cells. These conventional electrodes have poor fuel wetting properties. These conventional electrodes can be modified for use in liquid feed type fuel cells by coating them with substances that improve their wetting properties. Nafion™ with an equivalent weight of 1000 or higher is the preferred substance. The additive decreases interfacial tension of the liquid/catalyst interface and leads to the uniform wetting of the electrode pores and particles by the fuel and water solution, yielding enhanced utilization of the electro-catalyst. In addition to improving wetting properties, Nafion™ additive also provides iodine continuity with the solid electrolyte membrane and permits efficient transport of protons or hydronium ions generated by the fuel oxidation reaction. Further, the additive facilitates the release of carbon dioxide from the pores of the electrode. By using a perfluorinated sulfonic acid as the additive, anodic groups are not strongly absorbed on the electrode/electrolyte interface. Consequently, the kinetics of electro-oxidation of methanol are more facile than in sulfuric acid electrolyte. Other hydrophilic proton-conducting additives with the desired properties include montmorillonite clay, alkoxycelluloses, cycloextrins, mixtures of zeolites, and zirconium hydrogen phosphate.

The object of improving electrodes for operating in liquid feed fuel cells is achieved, in part, by using perfluorooctanesulfonic acid as an additive in an electro-desposition bath used in fabricating the electrode. An electro-desposition method using the perfluorooctanesulfonic acid additive comprises the steps of positioning a high-surface-area carbon electrode structure within a bath containing metallic salts, positioning an anode within the bath and applying a voltage between the anode and the cathode until a desired amount of metal becomes deposited onto the electrode. After deposition of the metal onto the electrode, the electrode is extracted from the bath and washed within deionized water.

Preferably, the metal salts include hydrogen hexachloro-platinate and potassium pentachloroaquoruthenium. The anode is composed of platinum. The carbon electrode structure includes high-surface-area carbon particles bound together by polytetrafluoroethylene, sold under the trade-mark Teflon™.

The object of providing for adequate wetting of an electrode within a liquid feed fuel cell having a sulfuric acid electrolyte is achieved by employing perfluorooctanesulfonic acid as an additive to the fuel mixture of the fuel cell. Preferably, the perfluorooctanesulfonic acid is added to the organic fuel and water mixture in concentrations from 0.001–0.1M.

The general objective of providing new fuels for use in organic fuel cells is achieved by using either trimethoxymethane, dimethoxymethane or trioxane. All three new fuels can be oxidized at a high rate into carbon dioxide and water within the fuel cell without poisoning the electrodes. Furthermore, neither trimethoxymethane, dimethoxymethane or trioxane are corrosive. Rates of oxi-
dation of the three new fuels are comparable to, or better than, oxidation rates of conventional organic fuels. For example, rates of oxidation for dimethoxymethane are higher than that of methanol at the same temperature. Trioxane achieves oxidation rates comparable to that of formaldehyde. However, trioxane has a much higher molecular weight than formaldehyde and, as such, molecules of trioxane do not cross-over to the cathode of the fuel cell as easily as molecules of formaldehyde.

Trimethoxymethane, dimethoxymethane and trioxane may be employed in a fuel cell having any of the improvements set forth above. However, the improved fuels may also be advantageously used within other organic fuel cells, including entirely conventional fuel cells.

Hence the various general objects of the invention set forth above are achieved. Other objects and advantages of the invention will be apparent from the detailed description set forth below.

BRIEF DESCRIPTION OF THE DRAWINGS

The objects and advantages of the present invention will become more readily apparent after reviewing the following detailed description and accompanying drawings, wherein:

FIG. 1 provides a schematic representation of an improved liquid feed organic fuel cell having a solid polymeric membrane configured in accordance with a preferred embodiment of the invention.

FIG. 2 provides a schematic representation of a multi-cell fuel system employing the improved liquid feed organic fuel cell of FIG. 1.

FIG. 3 is a graph illustrating the performance of a solid polymeric membrane electrolyte and a sulfuric acid electrolyte in liquid organic fuels.

FIG. 4 is a graph illustrating the performance of liquid feed fuel cell of FIG. 1 for methanol/air and methanol/oxygen combinations.

FIG. 5 is a graph illustrating the effect of fuel concentration on the performance of the liquid feed fuel cell of FIG. 1.

FIG. 6 is a graph illustrating the polarization behavior of the fuel electrode and cathode in the fuel cell of FIG. 1.

FIG. 7 is a block diagram illustrating a method for fabricating electrode containing hydrophilic proton-conducting ionomer additive for use in liquid feed cells.

FIG. 8 is a graph illustrating the polarization characteristics for methanol oxidation at electrodes containing the ionomer additive and prepared in accordance with the procedure shown in FIG. 7.

FIG. 9 is a block diagram illustrating a method for fabricating an electrode employing perfluorooctanesulfonic acid within an electro-deposition bath.

FIG. 10 is a schematic illustration of an electrochemical cell for use in performing the method of FIG. 9.

FIG. 11 is an illustrating polarization curves for an electrode fabricating using the method of FIG. 9.

FIG. 12 is a graph illustrating polarizations curves of a fuel cell using a sulfuric acid electrolyte and employing perfluorooctanesulfonic acid as a fuel additive.

FIG. 13 is a graph illustrating polarization curves of a fuel cell using dimethoxymethane as a fuel for various fuel concentration levels within a half cell having a sulfuric acid electrolyte.

FIG. 14 is a graph illustrating polarization curves of a fuel cell using dimethoxymethane as a fuel for differing temperature and concentrations within a half cell having a sulfuric acid electrolyte.

FIG. 15 is a graph illustrating cell voltage as a function of current density for the fuel cell of FIG. 1 using dimethoxymethane as a fuel.

FIG. 16 is a graph illustrating polarization curves of a fuel cell using trimethoxymethane as a fuel for various fuel concentration levels within a half cell having a sulfuric acid electrolyte.

FIG. 17 is a graph illustrating polarization curves of a fuel cell using trimethoxymethane as a fuel for differing temperatures and concentrations within a half cell having a sulfuric acid electrolyte.

FIG. 18 is a graph illustrating cell voltage as a function of current density for the fuel cell of FIG. 1 using trimethoxymethane or methanol as a fuel.

FIG. 19 is a graph illustrating polarization curves of a fuel cell using trioxane as a fuel for various fuel concentration levels within a half cell having a two molar sulfuric acid electrolyte.

FIG. 20 is a graph illustrating polarization curves of a fuel cell using trioxane as a fuel for differing fuel cells employing fabricating electrode containing hydrophilic proton-conducting ionomer additive for use in liquid feed cells.

FIG. 21 is a graph illustrating cell voltage as a function of current density for the fuel cell of FIG. 1 using trioxane as a fuel.

DETAILED DESCRIPTION OF THE INVENTION

Referring to the figures, the preferred embodiments of the invention will now be described. Initially, an improved liquid feed organic fuel cell using a solid polymeric electrolyte membrane and a ionomeric anode additive is described, primarily with reference to FIGS. 1-6. Then, a method for fabricating the anode having the ionomeric additive is described with reference to FIGS. 7-8. A method for achieving improved wetting by fabricating an electrode containing perfluorooctanesulfonic acid is described with reference to FIGS. 9-11. A fuel cell employing perfluorooctanesulfonic acid as a fuel additive is described with reference to FIG. 12. Fuel cells employing dimethoxymethane, trimethoxymethane and trioxane as fuels are described with reference to FIGS. 13-21.

Fuel Cell Employing Solid Proton Conducting Electrolyte Membrane.

FIG. 1 illustrates a liquid feed organic fuel cell having a housing, an anode, a cathode and a solid polymer proton-conducting cation-exchange electrolyte membrane. As will be described in more detail below, anode, cathode and solid polymer electrolyte membrane are preferably a single multi-layer composite structure, referred to herein as a membrane-electrode assembly. A pump is provided for pumping an organic fuel and water solution into an anode chamber housing. The organic fuel and water mixture is withdrawn through an outlet port and is vented into a cathode chamber through a port within tank. Carbon dioxide formed in the anode compartment is vented through a port within tank. An oxygen or air compressor is provided to feed oxygen or air into a cathode chamber within housing. FIG. 2, described below, illustrates a fuel cell system incorporating a stack of individual fuel cells including the re-circulation system. The following detailed description of the fuel cell of FIG. 1
Anode is preferably formed from unsupported or supported platinum or platinum-ruthenium alloy particles either as fine metal powders, i.e. "unsupported", or dispersed on high surface area carbon, i.e. "supported". The high surface area carbon may be a material such as Vulcan XC-72A, provided by Cabot Inc., USA. A carbon fiber sheet backing (not shown) is used to make electrical contact with the particles of the electrocatalyst. Commercially available Toray™ paper is used as the electrode backing sheet. A supported alloy electrocatalyst on a Toray™ paper backing is available from E-Tek, Inc., of Framingham, Mass. Alternatively, both unsupported and supported electrocatalysts may be prepared by chemical methods, combined with Teflon™ binder and spread on Toray™ paper backing to produce the anode. A film and time-saving method of fabrication of electro-catalytic electrodes is described in detail herein below.

Platinum-based alloys in which a second metal is either tin, iridium, osmium, or rhenium can be used instead of platinum-ruthenium. In general, the choice of the alloy depends on the fuel to be used in the fuel cell. Platinum-ruthenium is preferable for electro-oxidation of methanol. For platinum-ruthenium, the loading of the alloy particles in the electrocatalyst layer is preferably in the range of 0.5-4.0 mg/cm². More efficient electro-oxidation is realized at higher loading levels, rather than lower loading levels.

Cathode 16 is a gas diffusion electrode in which platinum particles are bonded to one side of membrane 18. Cathode 16 is preferably formed from unsupported or supported platinum bonded to a side of membrane 18 opposite to anode 14. Unsupported platinum black (fuel cell grade) available from Johnson Matthey Inc., USA or supported platinum materials available from E-Tek Inc., USA are suitable for the cathode. As with the anode, the cathode metal particles are preferably mounted on a carbon backing material. The loading of the electrocatalyst particles onto the carbon backing is preferably in the range of 0.5-4.0 mg/cm². The electrocatalyst alloy and the carbon fiber backing contain 10-50 weight percent Teflon™ to provide hydrophobicity needed to create a three-phase boundary and to achieve efficient removal of water produced by electro-reduction of oxygen.

During operation, a fuel and water mixture (containing no acidic or alkaline electrolyte) in the concentration range of 0.5-3.0 mole/liter is circulated past anode 14 within anode chamber 22. Preferably, flow rates in the range of 10-500 milliliters/min. are used. As the fuel and water mixture circulates past anode 14, the following electrochemical reaction, for an exemplary methanol cell, occurs releasing electrons:

Anode: \( \text{CH}_3\text{OH}+\text{H}_2\text{O} \rightarrow \text{CO}_2+\text{H}_2 \pm \text{e}^- \) (1)

Carbon dioxide produced by the above reaction is withdrawn along with the fuel and water solution through outlet 23 and separated from the solution in a gas-liquid separator (described below with reference to FIG. 2). The fuel and water solution is then re-circulated into the cell by pump 20. Simultaneous with the electrochemical reaction described in equation 1 above, another electrochemical reaction involving the electro-reduction of oxygen, which captures electrons, occurs at cathode 16 and is given by:

Cathode: \( \text{O}_2+4\text{H}^++4\text{e}^- \rightarrow 2\text{H}_2\text{O} \) (2)

The individual electrode reactions described by equations 1 and 2 result in an overall reaction for the exemplary methanol fuel cell given by:

Cell: \( \text{CH}_3\text{OH}+1.5\text{O}_2 \rightarrow \text{CO}_2+2\text{H}_2\text{O} \) (3)

At sufficiently high concentrations of fuel, current densities greater than 500 mA/cm² can be sustained. However, at these concentrations, a crossover rate of fuel across membrane 18 to cathode 16 increases to the extent that the efficiency and electrical performance of the fuel cell are reduced significantly. Concentrations below 0.5 mole/liter restrict cell operation to current densities less than 100 mA/cm². Lower flow rates have been found to be applicable at lower current densities. High flow rates are required while operating at high current densities to increase the rate of mass transport of organic fuel to the anode as well as to remove the carbon dioxide produced by electrochemical reaction. Low flow rates also reduce the crossover of the fuel from the anode to the cathode through the membrane.

Preferably, oxygen or air is circulated past cathode 16 at pressures in the range of 10 to 30 psig. Pressures greater than ambient improve the mass transport of oxygen to the sites of electrochemical reactions, especially at high current densities. Water produced by electrochemical reaction at the cathode is transported out of cathode chamber 28 by flow of oxygen through port 30.

In addition to undergoing electro-oxidation at the anode, the liquid fuel which is dissolved in water permeates through solid polymer electrolyte membrane 18 and combines with oxygen on the surface of the cathode electrocatalyst. This process is described by equation 3 for the example of methanol. This phenomenon is termed "fuel crossover". Fuel crossover lowers the operating potential of the oxygen electrode and results in consumption of fuel without producing useful electrical energy. In general, fuel crossover is a parasitic reaction which lowers efficiency, reduces performance and generates heat in the fuel cell. It is therefore desirable to minimize the rate of fuel crossover. The rate of crossover is proportional to the permeability of the fuel through the solid electrolyte membrane and increases with increasing concentration and temperature. By choosing a
solid electrolyte membrane with low water content, the permeability of the membrane to the liquid fuel can be reduced. Reduced permeability for the fuel results in a lower crossover rate. Also, fuels having a large molecular size have a smaller diffusion coefficient than fuels which have small molecular size. Hence, permeability can be reduced by choosing a fuel having a large molecular size. While water soluble fuels are desirable, fuels with moderate solubility exhibit lowered permeability. Fuels with high boiling points do not vaporize and their transport through the membrane is in the liquid phase. Since the permeability for vapors is higher than liquids, fuels with high boiling points generally have a low crossover rate. The concentration of the liquid fuel can also be lowered to reduce the crossover rate. With an optimum distribution of hydrophobic and hydrophilic sites, the anode structure is adequately wetted by the liquid fuel to sustain electrochemical reaction and excessive amounts of fuel are prevented from having access to the membrane electrolyte. Thus, an appropriate choice of anode structures can result in the high performance and desired low crossover rates.

Because of the solid electrolyte membrane is permeable to water at temperatures greater than 60°C, considerable quantities of water are transported across the membrane by permeation and evaporation. The water transported through the membrane is condensed in a water recovery system and fed into a water tank (both described below with reference to FIG. 2) so that the water can be re-introduced into anode chamber 22.

Protons generated at anode 14 and water produced at the cathode 16 are transported between the two electrodes by a proton-conducting solid electrolyte membrane 18. The maintenance of high proton conductivity of membrane 18 is important to the effective operation of an organic/air fuel cell. The water content of the membrane is maintained by providing contact directly with the liquid fuel and water mixture. The thickness of the proton-conducting solid polymer electrolyte membranes should be in the range from 0.05 to 0.5 mm to be dimensionally stable. Membranes thinner than 0.05 mm may result in membrane electrode assemblies which are poor in mechanical strength, while membranes thicker than 0.5 mm may suffer extreme and damaging dimensional changes induced by swelling of the polymer by the liquid fuel and water solutions and also exhibit excessive resistance. The ionic conductivity of the membranes should be greater than 1 ohm⁻¹ cm⁻¹ for the fuel cell to have a tolerable internal resistance. As noted above, the membrane should have a low permeability to the liquid fuel. Although a Nafion™ membrane has been found to be effective as a proton-conducting solid polymer electrolyte membrane, perfluorinated sulfonic acid polymer membranes such as Aci-plex™ (manufactured by Asahi Glass Co., Japan) and polymer membranes made by Dow Chemical Co., USA, such as XU/S13204.10 which are similar in properties to Nafion™ are also applicable. Membranes of polyethylene and polypropylene sulfonic acid, polystyrene sulfonic acid and other polyhydrocarbon-based sulfonic acids (such as membranes made by RAI Corporation, USA) can also be used depending on the temperature and duration of fuel cell operation. Composite membranes consisting of two or more types of proton-conducting cation-exchange polymers with differing acid equivalent weights, or varied chemical composition (such as modified acid group or polymer backbone), or varying water contents, or differing types and extents of cross-linking (such as cross linked by multivalent cations e.g., Al³⁺, Mg²⁺ etc.) can be used to achieve low fuel permeability. Such composite membranes can be fabricated to achieve high ionic conductivity, low permeability for the liquid fuel and good electrochemical stability.

As can be appreciated for the foregoing description, a liquid feed direct oxidation organic fuel cell is achieved using a proton-conducting solid polymer membrane as electrolyte without the need for a free soluble acid or base electrolyte. The only electrolyte is the proton-conducting solid polymer membrane. No acid is present in free form in the liquid fuel and water mixture. Since no free acid is present, acid-induced corrosion of cell components, which can occur in current-art acid based organic/air fuel cells, is avoided. This offers considerable flexibility in the choice of materials for the fuel cell and the associated subsystems. Furthermore, unlike fuel cells which contain potassium hydroxide as liquid electrolyte, cell performance does not degrade because soluble carbonates are not formed. Also by the use of a solid electrolyte membrane, parasitic shunt currents are avoided.

Referring now to FIG. 2, a fuel cell system employing a stack of fuel cells similar to the fuel cell shown in FIG. 1 will now be described. The fuel cell system includes a stack 25 of fuel cells, each having the membrane/electrode assembly described above with reference to FIG. 1. Oxygen or air is supplied by an oxidant supply 26 which may be, for example, a bottled oxygen supply, an air-blowing fan or an air compressor. An air and water or oxygen and water mixture is withdrawn from stack 25 through an outlet port 30 and conveyed to a water recovery unit 27. Water recovery unit 27 operates to separate the air or oxygen from the water. A portion of the air or oxygen separated by unit 27 is returned to oxidant supply 26 for re-entry into stack 25. Fresh air or oxygen is added to supply 27. Water separated by unit 27 is fed to a fuel and water injection unit 29 which also receives an organic fuel, such as methanol, from a storage tank 33. Injection unit 29 combines the water from the recovery unit 27 with the organic fuel from tank 33, yielding a fuel and water solution with the fuel dissolved in the water.

The fuel and water solution provided by injection unit 29 is fed into a circulation tank 35. A fuel and water mixture containing carbon dioxide is withdrawn through port 23 from stack 25 and is fed through a heat exchanger 37 and into circulation tank 35. Hence circulation tank 35 receives both a fuel and water solution from injection unit 29 and a fuel and water solution containing a carbon dioxide gas from heat exchanger 37. Circulation tank 35 extracts carbon dioxide from the fuel and water mixture and releases the carbon dioxide through a vent 39. The resulting fuel and water solution is fed through pump 20 and into stack 25. Circulation tank 35 can also be located between stack 25 and heat exchanger 34 so as to remove the carbon dioxide before the heat exchanger and thereby improve performance of the heat exchanger.

The operation of the various components illustrated in FIG. 2 will now be described in greater detail. Circulation tank 35 is a tower having a large head space. The liquid fuel and water mixture received from injection unit 29 is added into a top of the tower. The fuel and water mixture having carbon dioxide therein is fed into a bottom portion of the tower. Carbon dioxide gas released from the fuel and water mixture is allowed to accumulate in the head space and is ultimately vented. Alternately, the fuel and water mixture containing the carbon dioxide can be passed through a cluster of tubes of a microporous material such as Collard™ or GoreTex™ which allows gases to be released through walls of the tubes of the microporous material, while the liquid fuel flows along an axis of the tubes. Collard™ and GoreTex™ are registered trademarks of Celanese Corp. and Gore Association, USA.
A static re-circulation system (not shown) can be employed within an anode chamber of stack to separate carbon dioxide from the fuel and water mixture such that an external circulation tank need not be provided. With such a system, bubbles of carbon dioxide, due to innate buoyancy, tend to rise vertically within the anode chamber. Viscous interaction with the liquid fuel mixture surrounding the gas bubbles drags the liquid fuel upwards in the direction of outlet port. Once outside the anode chamber, the liquid releases the gas, exchanges heat with the surroundings and cools, thereby becoming denser than the liquid in the cell. The denser liquid is fed into the bottom of the anode chamber through an inlet port. Instead of expending electrical energy on the pump, the static re-circulation system takes advantage of the heat and gas produced in the cell. The aforementioned process forms the basis of the static re-circulation system which will not be described in further detail. It should be noted that the use of a static re-circulation system may restrict the orientation at which the fuel cell can be operated and may be viable only for stationary applications.

Test results for fuel cell having a Nafion™ electrolyte membrane.

The kinetics of electro-oxidation of methanol for a sulfuric acid electrolyte and Nafion™ electrolyte have been studied by galvanostatic polarization measurements in electrochemical cells (not illustrated but similar to an electro-deposition cell illustrated below in FIG. 10). The cells consist of a working electrode, a platinum counter electrode and a reference electrode. The working electrode is polarized within a solution containing the chosen electrolyte and liquid fuel. The potential of the working electrode versus the reference electrode is monitored.

FIG. 3 illustrates the polarization curve, i.e. polarization versus current density in milliamperes/cm² (mA/cm²), for the kinetics of methanol oxidation in the Nafion™ and sulfuric acid electrolytes, with curve 41 illustrating polarization for 0.5M sulfuric acid electrolyte and with curve 43 illustrating polarization for a Nafion™ electrolyte. Polarization is represented in potential versus NHE, wherein NHE stands for normalized hydrogen electrode. The curves represent measured data for a fuel consisting of a 1M mixture of methanol in water at 60°C. As can be seen from FIG. 3, the polarization losses are lower when the electrode is in contact with Nafion™ rather than sulfuric acid. Hence, it can be concluded that the kinetics of electro-oxidation of methanol are more facile when the electrolyte is Nafion™. These observations may be explained by the fact that strong adsorption of sulfate ions occurs at an electrode/sulfuric acid interface at positive potentials which hinders the kinetics of electro-oxidation. Such adsorption does not occur when Nafion™ is employed as an electrolyte since no such ions are produced. Also, it is believed that the kinetics of electro-reduction of oxygen or air are enhanced at an electrode/Nafion™ interface, in comparison to an electrode/sulfuric acid interface. This latter effect may be due to the higher solubility of oxygen in Nafion™ and the absence of strongly adsorbed anions. Therefore, the use of the proton-conducting solid polymer membrane as electrolyte is beneficial to the kinetics of both the electrode reactions and overcomes the disadvantages of a sulfuric acid electrolyte.

Also, sulfuric acid electrolytes suffer degradation at temperatures greater than 80°C. Products of degradation can reduce the performance of the individual electrodes. The electrochemical stability and thermal stability of a solid polymer electrolyte such as Nafion™ is considerably higher than that of sulfuric acid and the solid polymer electrolyte can be used at temperatures as high as 120°C. Therefore the use of the proton-conducting solid polymer membrane permits long term fuel cell operation at temperatures as high as 120°C, which provides an additional advantage since the kinetics of electro-oxidation of fuels and electro-reduction of oxygen occur with greater facility as the temperature is increased.

FIG. 4 illustrates the performance of the fuel cell shown in FIG. 2 when operated at 65°C for both a methanol oxygen combination and a methanol/air combination. In FIG. 4, voltage of the fuel cell is illustrated along axis and current density in mA/cm² is illustrated along axis. Curve indicates performance of the methanol/oxygen combination while curve illustrates performance of the methanol/air combination. As can be seen, the use of pure oxygen provides slightly better performance than air.

FIG. 5 illustrates the effect of fuel concentration on cell performance. Fuel cell potential is illustrated along axis while current density in mA/cm² is illustrated along axis. Curve 44 indicates performance for a 2.0 molar methanol solution at 150 degrees Fahrenheit (F). Curve 46 illustrates performance for a 0.5 molar methanol mixture at 140 degrees F. Curve 48 illustrates performance for a 0.4M methanol mixture at 160 degrees F. As can be seen, the 2.0M methanol mixture provides the best overall performance. Also, FIG. 5 illustrates that the fuel cell can sustain current densities as high as 300 mA/cm² while maintaining reasonably high voltage. In particular, the 2.0 molar methanol mixture provides a voltage of over 0.4 volts at nearly 300 mA/cm². The performance illustrated in FIG. 5 represents a significant improvement over the performance of previous organic fuel cells.

Polarization behavior of the anode and cathode of the fuel cell are illustrated in FIG. 6 as a function of current density in mA/cm², with voltage shown along axis and current density along axis. Curve 50 and current density at 150 degrees F. Curve 56 illustrates the polarization behavior for the fuel while curve 58 separately illustrates polarization behavior for the oxygen.

Anode Structures for Liquid Feed Type Fuel Cells.

The anode structure for liquid feed fuel cells must be quite different from that of conventional fuel cells. Conventional fuel cells employ gas diffusion type electrode structures that can provide gas, liquid and solid equilibrium. However, liquid feed type fuel cells require anode structures that are similar to batteries. The anode structures must be porous and must be capable of wetting the liquid fuel. In addition, the structures must have both electronic and ionic conductivity to effectively transport electrons to the anode collector (carbon paper) and hydrogen/hydronium ions to the Nafion™ electrolyte membrane. Furthermore, the anode structure must help achieve favorable gas evolving characteristics at the anode.

Electrodes required for liquid feed type fuel cells can be fabricated specifically or conventional gas diffusion electrodes available commercially can be modified with suitable additives. Electrode Impregnation with Ionomeric Additive.

The electrocatalyst layer and carbon fiber support of anode are preferably impregnated with a hydrophilic proton-conducting polymer additive such as Nafion™. The additive is provided within the anode, in part, to permit efficient transport of protons and hydronium produced by the electro-oxidation reaction. The ionomeric additive also promotes uniform wetting of the electrode pores by the liquid fuel/water solution and provides for
better utilization of the electrocatalyst. The kinetics of methanol electro-oxidation by reduced adsorption of anions is also improved. Furthermore, the use of the ionomeric additive helps achieve favorable gas evolving charactersitics for the anode.

For an anode additive to be effective, the additive should be hydrophilic, proton-conducting, electrochemically stable and should not hinder the kinetics of oxidation of liquid fuel. Nafion™ satisfies these criteria and is a preferred anode additive. Other hydrophilic proton-conducting additives which are expected to have the same effect as Nafion™ are montmorillonite clays, zeolites, alkoxycelluloses, cyclodextrins, and zirconium hydrogen phosphate.

FIG. 7 is a block diagram which illustrates the steps involved in impregnation of the anode with an ionomeric additive such as Nafion™. Initially, a carbon electrode structure is obtained or prepared. Commercially available high surface area carbon electrode structures which employ a mixture of high surface area electrocatalyst and Teflon™ binder applied on Toray™ paper may be used. High surface area catalyst particles and Toray™ paper, both available from E-Tek, Inc., using TFE-30™, an emulsion of polytetrafluoroethylene. Although these structures can be prepared from the foregoing component materials, prefabricated structures may also be obtained directly from E-Tek in any desired dimension.

At step 302, the electrodes are impregnated with an ionomeric additive, such as Nafion™, by immersing the electrocatalyst particles in a solution containing 0.5-5% of the ionomeric additive (by appropriate dilution, with methanol or isopropanol, of solutions supplied by Aldrich Chemical Co., or Solution Technologies Inc.) for 5-10 minutes. The electrode is then removed, at step 304, from the solution and dried in air or vacuum at temperatures ranging from 20°-60° C. to volatilize any higher alcohol residues associated with the Nafion™ solution. The impregnation steps 302-304 are repeated until the desired composition (which is in the range of 2-10% of the weight of the electrocatalyst) is achieved. A loading of 0.1 to 0.5 mg/cm² is exemplary. Electrode compositions with additive in excess of 10% may result in an increased internal resistance of the electrode and poor bonding with the solid polymer electrolyte membrane. Compositions with less than 2% of the additive do not typically result in improved electrode performance.

To form impregnated electrodes from electrocatalyst particles, the electrocatalyst particles are mixed in with a solution of Nafion™ diluted to 1% with isopropanol. Then the solvent is allowed to evaporate until a thick mix is reached. The thick mix is then applied onto a Toray™ paper to form a thin layer of the electrocatalyst. A mixture of about 200 weight/g gram high surface area particles applied to the Toray™ paper is exemplary. Note here that the electrocatalyst layer so formed has only Nafion™ and no Teflon™. Electrodes so prepared are then dried in a vacuum at 60° C. for 1 hour to remove higher alcohol residues, after which they are ready for use in liquid feed cells.

A commercially available high-surface area platinum-tin electrode was impregnated with Nafion™ according to the procedure described above. FIG. 8 compares the performance of a Nafion™-impregnated electrode with a non-impregnated electrode as measured within a half cell similar to the cell of FIG. 10 (below) but containing a sulfuric acid electrolyte. In particular, FIG. 8 illustrates the polarization measurements in liquid formaldehyde fuel (1 molar) with sulfuric acid electrolyte (0.5 molar). The current density in mA/cm² is illustrated along axis 306 and the potential in volts along axis 308. Curve 310 is the galvanostatic polarization curve for a platinum-tin electrode that does not include Nafion™. Curve 312 is the galvanostatic polarization curve for a platinum-tin electrode not impregnated with Nafion™.

It can be seen from FIG. 8 that far greater current densities are achieved with the Nafion™-impregnated electrode than with the non-impregnated electrode. Indeed with the non-impregnated electrode, very little oxidation of formaldehyde occurs. The addition of Nafion™ thus provides a dramatic improvement. In addition, the absence of any hysteresis in the galvanostatic polarization curves suggest that these coatings are stable.

What has been described thus far is an improved liquid feed fuel cell anode impregnated with an ionomeric additive. A method for fabricating the anode to include the ionomeric additive has also been described. The remaining sections of the Detailed Description provide a description of the use of perfluorooctanesulfonic acid as an additive within an electro-deposition bath used for fabricating electrodes and as a direct additive within a fuel. New fuels are also described. Electro-deposition of Electrodes using Perfluorooctanesulfonic Acid Additive

With reference to FIGS. 9-11, a method for fabricating an electrode for use in a organic fuel cell will now be described in detail. The method is advantageously employed for fabricating a cathode for use in the liquid organic fuel cell described above. However, electrodes prepared by the method of FIGS. 9-11 may alternatively be used in a variety of organic fuel cells.

Referring first to FIG. 9, the steps of a method for fabricating the anode will now be described. Initially, at 200, a carbon electrode structure is prepared by applying a mixture of high-surface-area carbon particles and a Teflon™ binder to a fiber-based carbon paper. Preferably, the carbon particles have a surface area of 200 meters²/g (m²/g). A suitable carbon particle substrate, referred to Vulcan XC-72, is available from E-Tek Inc. The Teflon™ binder is preferably added to achieve a percentage, by weight, of 15%. The fiber based carbon paper is preferably Toray™ paper, also available from E-Tek Incorporated. The carbon structure may be prepared from the foregoing component materials. Alternatively, commercial prefabricated structures are available in 2 inch by 2 inch squares directly from E-Tek Inc.

At step 202, an electro-deposition bath is prepared by dissolving hydrogen hexachloropaltinate (IV) and potassium pentachloroaquoruthonium (III) within sulfuric acid. Preferably, the resulting metal-ion concentration is within the range of 0.01-0.05M. Also, preferably, the sulfuric acid has the concentration of 1M. The forgoing compound is employed for obtaining platinum-ruthenium deposits on the carbon electrode structure. Alternative solutions may be employed. For example, to obtain platinum-tin deposits, a stannic chloride compound is dissolved in a sulfuric acid instead.

The metallic ion salts are dissolved in the sulfuric acid. Primarily to prevent hydrolysis of the solution. For ruthenium deposition, the resulting solution is preferably de-aerated to prevent the formation of higher oxidation states. High purity perfluorooctanesulfonic acid (C-8 acid) is added to the bath at step 204. C-8 acid is preferably added to a concentration in a range of 0.1-1.0 grams/liters. C-8 acid is provided to facilitate complete wetting of the carbon particles. C-8 acid is electro-inactive and does not specifically adsorb at metal sites within the structure. Therefore, C-8 acid is innocuous to subsequent electro-deposition processes. The addition of C-8 acid has been found to be highly
beneficial, and perhaps necessary for successful electro-
deposition onto the electrodes.

A carbon electrode structure resulting from step
200 is placed within the electro-deposition bath resulting
from step 204. A platinum anode is also positioned within
the bath. For the deposition of other metal ions, an alternate
anode material may be employed.

A voltage is then applied between the carbon electrode
structure and the platinum anode at step 208. The voltage is
applied for about 5 to 10 minutes to achieve electro-
deposition of platinum-ruthenium onto the carbon electrode
to a loading of about 5 mg/cm². Preferably, a voltage of
approximately −0.8 V vs mercury sulfate reference electrode
is applied.

After a desired amount of metal is deposited onto the
carbon electrode, the electrode is removed, at step 210, and
washed in deionized water. Preferably, the electrode is
washed at least three times in circulating de-ionized water
for 15 minutes each time. The washing step is provided
primarily to rid the surface of the carbon electrode of
absorbed chloride and sulfate ions. The washing step has
been found to be highly desirable, and perhaps necessary, for
yielding an effective electrode for use in an organic fuel cell.

Electrodes, resulting from the fabrication method of step
206, have been found to have very uniform “cotton-ball-
shaped particles, with a significant amount of fine structure.
Average particle size has been found to be on the order of 0.1
microns.

A deposition setup for use in implementing the method of
FIG. 9 is illustrated in FIG. 10. Specifically, FIG. 10
illustrates a three-electrode cell 212 which includes a single
carbon-structure electrode 214, a pair of platinum counter-
electrodes (or anodes) 216 and a reference electrode 218. All
electrodes are positioned within a bath 220 formed of the
aforementioned metallic/C-8 acid solution. Electrical contacts
222 and 224 are positioned on interior side surfaces of
cell 212 above bath 220. A magnetic stirrer 226 is positioned
within bath 220 to facilitate stirring and circulation of the
bath. A circulating water jacket 228 is provided around cell
212 for use in regulating the temperature within the cell. The
platinum anodes are positioned within fine glass frits 230,
the glass frits being provided to isolate the anodes from the
cathode to prevent the oxidation products of the anode from
diffusing into the cathode.

Reference electrode 214 is a mercury/mercuric-sulfate
reference electrode. The reference electrode is provided to
monitor and control the electrical potential of carbon elec-
structure 214. Preferably both potentiostatic and gal-
vanostatic control methods are employed. The composition of
the alloy deposit is controlled by choosing a bath com-
position, summarized above, and by performing electrode-
deposition at current densities well above any limiting
current densities for the metal deposition. When selecting an
appropriate bath composition it is important to normalize for
the electrochemical equivalence of the metals within the
composition.

During operation, the quantity of charge passed from
anode to cathode is detected and employed to monitor the
quantity of material deposited. In this regard, an amount of
charge used in any hydrogen evolution reaction must be
subtracted from each measurement of total charge.

Accurate electro-deposition typically occurs within a
period of five to ten minutes, depending upon the operating
conditions and the catalyst loading desired.

The monitoring equipment for use in monitoring and
controlling the electrode potential are not illustrated in FIG.
10 as the function and operation of such devices are well
known to those skilled in the art.

FIG. 11 illustrates the performance of an exemplary
electrode deposited using the method of FIG. 9 within the
A gasoline-carbon electrode cell of FIG. 7. In FIG. 11, potential in
volts versus NHE is provided along axis 240 whereas
current density in mA/cm² is provided along axis 242. Curve
246 illustrates the galvanostatic polarization curve for a
carbon-supported platinum-ruthenium alloy electrode pre-
pared in accordance with the forgoing for a loading of 5
gm/cm². Curve 246 illustrates galvanostatic polarization for an
electrode having a 1 mg/cm² loading. In each case, the
electrode was employed within a sulfuric acid electrolyte in
a half-cell. The fuel cell included an organic fuel composed
of 1 molar methanol and 0.5 molar sulfuric acid, operated at
60°C. At the loading of 5 mg/cm², the electrode sustains a
continuous current density of 100 mA/cm² at 0.45 volts
versus NHE.

The results illustrated in FIG. 11 are exemplary of per-
formance which may be achieved using an electrode fabric-
ated in accordance with the method of FIG. 9. Further
performance enhancement may be achieved with appro-
priate optimization of the electro-deposition conditions and the
alloy composition. Hence, the particular conditions and
concentrations described above are not necessarily optimal
but merely represent a currently known best mode for fabricating electrodes.

Perfluorooctanesulfonic Acid (C-8 Acid) as a Fuel Additive

The use of C-8 acid as an additive within an electro-
deposition bath was described above. It has also been
determined that C-8 acid may be advantageously applied as
an additive within the fuel of a liquid feed fuel cell employ-
ing a sulfuric acid electrolyte. In particular, it has been found
that straight chain C-8 acid, having the molecular formula
C₈F₁₇SO₃H, in concentrations from 0.001 to 0.1M is an
excellent wetting agent within a liquid feed fuel cell.

FIGS. 12 illustrates results of experiments which contrast
the use of C-8 acid as an additive with fuel cells lacking the
additive. In particular, FIG. 12 illustrates the results of
half-cell experiments using a Teflon™ coated high-surface
area carbon-supported platinum and platinum alloy elec-
rode mounted within a sulfuric acid electrolyte. The results
were obtained using a half-cell similar to the cell illustrated
in FIG. 10. FIG. 12 illustrates potential versus NHE along
the vertical axis 400 and current density in mA/cm² along a
horizontal axis 402. Four curves are provided illustrating
polarization for a fuel containing no additive (curve 404),
0.0001M additive (curve 406), 0.001M additive (curve 408) and
0.01M additive (curve 412).

As can be seen from FIG. 12, the addition of the C-8
additive decreases the polarization rather significantly.
Although not shown, the oxidation of methanol has also
been investigated using 0.1M pure C-8 acid solutions with-
out any sulfuric acid. Polarization curves (not shown) indi-
cate that the kinetics are not effected by the presence of the
perfluorooctanesulfonic ion.

Thus, FIG. 12 demonstrates that the use of C8 acid as an
additive in the concentration range of 0.0001M or greater
is beneficial to liquid fuel solutions when employing commer-
cial available Teflon™ coated fuel cell electrodes, at least
for fuel cells employing sulfuric acid as an electrolyte.

With reference to the remaining figures, three new fuels
for use in liquid feed fuel cells are described. The fuels are
dimethoxymethane, trimethoxymethane, and trioxane.
Dimethoxymethane as a fuel for a liquid feed fuel cell

FIGS. 13-15 illustrate the results of experiments con-
ducted using dimethoxyethane (DMM) as a fuel for an
organic direct liquid feed fuel cell. In use, DMM is mixed
with water to a concentration in the range of about 0.1 to 2M
and fed into a fuel cell. Other concentrations may also be effective. The fuel cell may be of conventional design or may include one or more of the improvements described above. Within the fuel cell, the DMM is electro-oxidized at the anode of the cell. The electro-oxidation of DMM involves a series of dissociative steps followed by surface reaction to form carbon dioxide and water. The electrochemical reaction is given by:

\[
\text{(CH}_3\text{OCH}_2\text{CH}_2\text{OH})+4\text{H}^+ + 3\text{e}^- \rightarrow \text{CO}_2 + 2\text{H}_2\text{O}
\]

Experiments testing the electro-oxidation of DMM have been performed in half-cells of the similar to the cell shown in FIG. 10 with temperature control using a 0.5M sulfuric acid electrolyte with Pt-Sn or Pt-Ru electrocatalyst electrodes. The galvanostatic polarization curves shown in FIG. 13 illustrate the electro-oxidation characteristics of DMM for platinum-tin electrodes for several different fuel concentrations. The platinum-tin electrodes are of the gas-diffusion type consisting of 0.5 mg/cm² total metal supported on Vulcan XC-72 obtained from Etek, Inc., Framingham, Mass. In FIG. 13, current density is along axis 500 and polarization (in terms of potential vs NHE) is provided along axis 502. Curves 504, 506, 508 and 510, respectively, illustrate polarization for DMM concentrations of 0.1M, 0.5M, 1M and 2M. FIG. 13 shows that increased concentration improves the kinetics of oxidation of DMM. The curves of FIG. 13 were measured in a half cell employing 0.5M sulfuric acid as an electrolyte, along with 0.1M C-8 acid. The measurements were conducted at room temperature.

It has been found that DMM can be oxidized at potentials considerably more negative than methanol. Also, temperature has been found to significantly influence the rates of oxidation. However, DMM has a low boiling point of 41° C. Hence, difficulties may arise in attempting to use DMM in a liquid feed fuel cell for temperatures higher than the boiling point.

FIG. 14 illustrates polarization for two different concentrations at two different temperatures. Current density is provided along axis 512 and polarization (in terms of potential vs NHE) is provided along axis 514. Curve 516 illustrates polarization for a 1M concentration of DMM at room temperature. Curve 18 illustrates polarization for a 2M concentration of DMM at 55° C. As can be seen, improved polarization is achieved using a higher concentration at a higher temperature. Also, a comparison of curve 510 of FIG. 13 with curve 518 of FIG. 14 illustrates that an increase in temperature yields an improved polarization for the same concentration level. Hence, it is believed that an increase in temperature results in improved kinetics of electro-oxidation.

In addition to the half cell experiments illustrated in FIGS. 13 and 14, fuel cell experiments were also conducted to verify the effectiveness of DMM in a fuel cell. The direct oxidation of DMM in fuel cells was carried out in a liquid feed type fuel cell as illustrated above in FIGS. 1 and 2. Hence, the fuel cell employed a porous conducting solid polymer membrane (Nafion™ 117) as the electrolyte. The membrane electrode assembly consisted of a fuel oxidation electrode made of unsupported platinum-ruthenium catalyst layer (4 mg/cm²) and gas-diffusion type unsupported platinum electrode (4 mg/cm²) for the reduction of oxygen. The fuel cell used a 1M solution of DMM on the fuel oxidation side and oxygen at 20 psi on the cathode.

Analysis of the oxidation products of DMM show only methanol. Methanol is considered a possible intermediate in the oxidation of DMM to carbon dioxide and water. However, since the fuel cell system is compatible with methanol, the presence of methanol as an intermediate is not a concern since the methanol is also ultimately oxidized to carbon dioxide and water.

The current-voltage characteristics of a liquid feed direct oxidation fuel cell using DMM as a fuel is shown in FIG. 15. The fuel cell was operated at 37° C. In FIG. 15, current density in mA/cm² is provided along axis 520. Cell voltage in volts is provided along axis 522. Curve 524 illustrates cell voltage as a function of current density for a 1M DMM solution described above. As can be seen from FIG. 15, the cell voltages reached 0.25 V at 50 mA/cm² with DMM which is as high as that attained with methanol (not shown). By working at a higher temperature and using a Pt-Sn catalyst, even better performance may be achieved. The low boiling point of DMM also makes it a candidate for a gas-feed type operation.

Thus from both half-cell and full-cell measurements it has been found that DMM is capable of being oxidized at very high rates. Therefore, it is believed that DMM is an excellent fuel for use in direct oxidation fuel cells. Also, DMM is a non-toxic, low-vapor pressure liquid, permitting easy handling. In addition DMM can be synthesized from natural gas (methane) by conventional techniques.

Trimethoxymethane as a fuel for a liquid feed fuel cell FIGS. 16–18 illustrate the results of experiments conducted using trimethoxymethane (TMM) as a fuel for an organic direct liquid feed fuel cell. As with DMM described above, in use, TMM is mixed with water to a concentration in the range of about 0.1 to 2M and fed into a fuel cell. Other concentrations may also be effective. The fuel cell may be of conventional design or may include one or more of the improvements described above. Within the fuel cell, the TMM is electro-oxidized at the anode of the cell. The electrochemical oxidation of TMM is represented by the following action:

\[
\text{CH}_3\text{OCH}_2\text{CH}_2\text{OH} + \text{H}_2\text{O} \rightarrow 4\text{CO}_2 + 2\text{H}^+ + 2\text{e}^-
\]

Experiments verifying the electro-oxidation of TMM have been performed in half-cells similar to the cell shown in FIG. 10 with temperature control using a Pt-Sn electrode with a 0.5M sulfuric acid electrolyte including 0.01M C-8 acid. Results of these half-cell experiments are illustrated in FIGS. 16 and 17. FIG. 16 provides galvanostatic polarization curves for several different concentrations of TMM for the above-mentioned Pt-Sn electrodes. The Pt-Sn electrodes were of the gas-diffusion type and consisted of 0.5 mg/cm² of total metal supported on Vulcan XC-72 obtained from Etek, Inc., Framingham, Mass. In FIG. 16, current density in mA/cm² is provided along axis 600 and polarization (in terms of potential vs NHE) is provided along axis 602. Curves 604, 606, 608 and 610, respectively, illustrate polarization for TMM concentrations of 0.1M, 0.5M, 1M and 2M TMM. FIG. 16 shows that improved polarization is achieved at higher concentration levels. All measurements shown in FIG. 16 were obtained at room temperature.

It is found that TMM can be oxidized at potentials considerably more negative than methanol. Also, it has been found that temperature affects the oxidation rate of TMM. FIG. 17 illustrates polarization at two different concentrations and at two different temperatures. In FIG. 17, current density in mA/cm² is provided along axis 612 and polarization (in potential vs NHE) is provided along axis 614. Curve 616 illustrates polarization for a 1M TMM concentration at room temperature whereas curve 618 illustrates polarization for a 2M concentration of TMM at 55° C. The
curves of FIG. 17 were obtained using a Pt-Sn electrode in a 0.5M sulfuric electrolyte including 0.01M C-8 acid. As can be seen, improved polarization is achieved using a higher concentration at a higher temperature. A comparison of curve 618 of FIG. 17 with curve 610 of FIG. 16 illustrates that an increase in temperature yields an improved performance for the same concentration level. Although not shown, it has been found that at temperatures as high as 60°C, the rate of oxidation of TMM is twice that at 25°C.

In addition to the half cell experiments illustrated in FIGS. 16 and 17, full fuel cell experiments were also conducted to verify the effectiveness of TMM in a fuel cell. The direct oxidation of TMM in fuel cells was carried out in a liquid feed type fuel cell of the type illustrated above in FIGS. 1 and 2. Hence, the fuel cell used the proton conducting solid polymer membrane (Nafion® 117) as the electrolyte. The membrane electrode assembly of the fuel cell included unsupported platinum-ruthenium catalyst layer (4 mg/cm²) and gas-diffusion type unsupported platinum electrode (4 mg/cm²) for the reduction of oxygen. The fuel cell used a 2M solution of TMM on the fuel oxidation side and oxygen at 20 psi on the cathode.

As with DMM, an analysis of the oxidation products of TMM show only methanol and methanol is considered a possible intermediate in the oxidation of TMM to carbon dioxide and water. For fuel cells which are compatible with methanol, the presence of methanol as an intermediate product is not a concern because the methanol is ultimately oxidized to carbon dioxide and water.

The current-voltage characteristics of the above-described liquid feed direct oxidation fuel cell is shown in FIG. 18 for both TMM and methanol. Current density in mA/cm² is provided along axis 620 and cell voltage is provided along axis 622. Curve 624 shows cell voltage as a function of current density for a 1M concentration of TMM. Curve 626 illustrates the same for a 1M concentration of methanol. The measurements shown in FIG. 18 were obtained at 65°C. Although not shown, at 90°C, cell voltages can reach 0.52 V at 300 mA/cm² with TMM which is higher than that attained with methanol.

Thus from both half-cell and full-cell measurements it has been found that TMM, like DMM, is capable of being oxidized at very high rates. Also like DMM, TMM is a non-toxic, low-vapor pressure liquid, permitting easy handling, and can be synthesized from natural gas (methane) by conventional methods.

TMM as a fuel for a liquid feed fuel cell FIGS. 19-21 illustrate the results of experiments conducted using trioxane as a fuel for an organic direct liquid fuel cell. As with DMM and TMM described above, this cell was carried out in a liquid feed type fuel cell of the type described in FIGS. 1 and 2. Hence, the fuel cell used the proton conducting solid polymer membrane (Nafion® 117) as the electrolyte. The fuel cell used a 1M solution of trioxane on the fuel oxidation side and oxygen at 20 psi on the cathode.

As with DMM and TMM, an analysis of the oxidation products of trioxane show only methanol and methanol is considered a possible intermediate in the oxidation of TMM to carbon dioxide and water. For fuel cells which are compatible with methanol, the presence methanol as an intermediate product is not a concern because the methanol is ultimately oxidized to carbon dioxide and water.

The current-voltage characteristics of the above-described liquid feed direct oxidation fuel cell is shown in FIG. 21 for trioxane. Current density in mA/cm² is provided along axis 724 and cell voltage is provided along axis 726. Curve 728 shows cell voltage as a function of current density for a 1M concentration of trioxane. The measurements shown in FIG. 21 were obtained at 60°C. The performance illustrated in FIG. 21 may be improved considerably using platinum-tin electrodes, rather than Pt-Ru electrodes.
A measurement of crossover, not shown, in the trioxane/oxygen fuel cell suggests that a rate of crossover is at least 5 times lower than that in methanol fuel cells. The decreased rates of crossover are extremely desirable since, as described above, crossover affects the efficiency and performance of fuel cells.

Thus from both half-cell and full-cell measurements it has been found that trioxane, like DMM and TMM, is capable of being oxidized at very high rates.

Conclusion

What has been described are a variety of improvements to liquid feed fuel cells including improved electrolyte and electrode structures, improved methods for fabricating electrodes, additives for improving fuel performance and a set of three new fuels. The various improvements may be implemented separately or, for the most part, may be combined to achieve even more enhanced performance. It should be noted, however, that the above-described use of C-8 acid as an additive in a fuel is expected to be effective only for fuel cells employing an acid electrolyte such as sulfuric acid and may not be effective if employed using a fuel cell configured with a proton exchange membrane.

The methods, embodiments and experimental results shown herein are merely illustrative and exemplary of the invention and should not be construed as limiting the scope of the invention.

What is claimed is:

1. In a liquid feed direct fuel cell having an anode, a cathode, an electrolyte, means for circulating an organic fuel past the anode and means for flowing oxygen past the cathode, wherein the improvement comprises:
   - said electrolyte comprising a solid polymer hydrogen ion conductor membrane; and
   - wherein said organic fuel is free of an acid electrolyte.
2. The fuel cell of claim 1, wherein, said membrane is composed of a co-polymer of tetrafluoroethylene and perfluorocarbonyl sulfonic acid.
3. The fuel cell of claim 1, wherein said membrane is composed of a perfluorinated sulfonic acid polymer.
4. The fuel cell of claim 1, wherein said membrane is composed of a polyhydrocarbon sulfonic acid polymer.
5. The fuel cell of claim 1, wherein said membrane is composed of a composite of two or more hydrogen ion conductor membranes.
6. The fuel cell of claim 1, wherein said organic fuel is selected from a group consisting of methanol, formaldehyde, and formic acid.
7. The fuel cell of claim 1, wherein said organic fuel is selected from a group consisting of dimethoxymethane, trimethoxymethane and trioxane.
8. The fuel cell of claim 1, wherein said anode is an electrode impregnated with a hydrophilic proton conducting water-insoluble additive.
9. The fuel cell of claim 8, wherein said additive is a co-polymer of tetrafluoroethylene and perfluorovinylsulfonic acid.
10. The fuel cell of claim 8, wherein said additive is montmorillonite clay.
11. The fuel cell of claim 8, wherein said additive is an alkoxycellulose.
12. The fuel cell of claim 8, wherein said additive is a cycloextrin.
13. The fuel cell of claim 8, wherein said additive is a mixture of zeolites.
14. The fuel cell of claim 8, wherein said additive is zirconium hydrogen phosphate.
15. A liquid feed fuel cell, comprising:
   - an anode; a cathode; and a solid cation exchange polymer electrolyte membrane disposed between said anode and said cathode;
   - means for circulating a liquid organic fuel and water solution past said anode, a liquid organic and water fuel solution, said solution being free of sulfuric acid; and
   - means for flowing oxygen past said cathode.
16. A liquid fuel cell, comprising:
   - an anode coated with an amount of hydrophilic proton conducting water-insoluble additive to provide ionic and electronic conductivity, wetting and gas evolving properties to said anode; a cathode; and a cation exchange polymer electrolyte membrane disposed between said anode and said cathode;
   - means for flowing a liquid comprising organic fuel and water solution past said anode; and
   - means for flowing oxygen past said cathode.
17. The fuel cell of claim 16, wherein said membrane is a solid proton exchange membrane.
18. The fuel cell of claim 17, wherein said membrane is composed of a co-polymer of tetrafluoroethylene and perfluorovinylsulfonic acid.
19. The fuel cell of claim 17, wherein said membrane is composed of a perfluorinated sulfonic acid polymer.
20. The fuel cell of claim 17, wherein said membrane is composed of polyhydrocarbon sulfonic acid polymer.
21. The fuel cell of claim 17, wherein said membrane is composed of a composite of two or more proton exchange membranes.
22. The fuel cell of claim 16, wherein said organic fuel is selected from a group consisting of methanol, formaldehyde, and formic acid.
23. The fuel cell of claim 16, wherein said organic fuel is selected from a group consisting of dimethoxymethane, trimethoxymethane and trioxane.
24. The fuel cell of claim 16, wherein said additive is a co-polymer of tetrafluoroethylene and perfluorovinylether sulfonic acid.
25. The fuel cell of claim 16, wherein said additive is montmorillonite clay.
26. The fuel cell of claim 16, wherein said additive is an alkoxycellulose.
27. The fuel cell of claim 16, wherein said additive is a cycloextrin.
28. The fuel cell of claim 16, wherein said additive is a mixture of zeolites.
29. The fuel cell of claim 16, wherein said additive is zirconium hydrogen phosphate.
30. A liquid feed fuel cell, comprising:
   - a housing having an anode chamber and a cathode chamber;
   - a co-polymer of tetrafluoroethylene and perfluorovinylether sulfonic acid polymer electrolyte membrane mounted within said housing and separating said anode and cathode chambers; a cathode formed on a side surface of the membrane facing the cathode chamber; and
   - an anode formed on an opposing side of the membrane facing the anode chamber, said anode being impregnated with co-polymer of tetrafluoroethylene and perfluorovinylether sulfonic acid; and
   - means for circulating a liquid organic fuel and water solution past said anode;
means for flowing oxygen past said cathode;
means for withdrawing carbon dioxide from the anode chamber;
means for withdrawing unused fuel and water from the anode compartment; and
means for withdrawing unused oxygen and water from the cathode chamber.

31. An electrode comprising a metal alloy coated with an amount of hydrophilic proton conducting, water insoluble additive to provide ionic and electronic conductivity, wetting and gas evolving properties to the electrode.

32. A method for processing a carbon structure composed of carbon particles supported by a binder, said method comprising the steps of:
immersing the carbon structure within a bath containing a liquid perflourinated sulfonic acid polymer; and
removing and drying said carbon structure.

33. The method of claim 32, wherein polymer is a 1% co-polymer of tetrafluoroethylene and perfluorovinylether sulfonic acid solution and said polymer is dispersed in methanol.

34. The method of claim 32, wherein said step of immersing said carbon structure in a bath containing a liquid polymer is performed for 5 to 10 minutes.

35. A structure processed according to the method of claim 32.

36. An electro-deposition bath for use in fabricating an electrode for use in a liquid feed fuel cell, comprising perfluorooctanesulfonic acid.

37. A method for fabricating an electrode for use in a fuel cell, said method comprising the steps of:
providing a bath containing a solution of metallic salts dissolved in sulfuric acid;
adding perfluorooctanesulfonic acid to said bath;
positioning a carbon electrode structure within said bath;
positioning an anode within said bath; and
applying a voltage between said anode and said electrode until a desired amount of metal in the range of 0.5-4.0 mg/cm² becomes deposited onto said electrode.

38. The method of claim 37, wherein said metal salts include hydrogen hexachloroplatinate and potassium pentachloroauratothanium.

39. The method of claim 37, wherein said anode is composed of platinum.

40. The method of claim 37, wherein said carbon electrode structure includes carbon combined with a polytetrafluoroethylene binder.

41. The method of claim 37, wherein said carbon electrode includes carbon bound by a 15%, by weight, polytetrafluoroethylene binder and applied on a carbon based fiber layer.

42. The method of claim 37, wherein said acid is provided with a concentration in the range of 0.01-0.05M.

43. The method of claim 37, including the further steps of extracting said electrode from said bath and washing said electrode in deionized water.

44. A method for fabricating an electrode having metal ions deposited thereon for use in a liquid organic fuel cell, said method comprising the steps of:
providing a bath comprising a solution of hydrogen hexachloroplatinate and potassium pentachloroauratothanium dissolved in sulfuric acid, wherein said hydrogen hexachloroplatinate and potassium pentachloroauratothanium has a concentration in the range 0.01-0.05M;
adding perfluorooctanesulfonic acid to said bath, with a concentration within the range of 0.1-1.0 grams/liter;
positioning a carbon electrode structure into said bath, wherein said carbon electrode structure has a mixture of carbon particles with a surface area of about 200 meters/gram a polytetrafluoroethylene binder, with the mixture applied to a fiber-based carbon paper;
positioning a platinum anode into said bath; and
applying a voltage between said anode and said electrode until a desired amount of platinum and ruthenium in the range of 0.5-4.0 mg/cm² become deposited onto said electrode.

45. An electrode fabricated according to the method of claim 44.

46. A liquid feed fuel cell, comprising:
an anode;
a cathode;
a liquid organic fuel, water, acid electrolyte and perfluorooctanesulfonic acid additive solution;
means for flowing said solution past said anode; and
means for flowing oxygen past said cathode.

47. A liquid feed fuel cell, comprising:
an anode;
a cathode;
an electrolyte;
a liquid organic fuel selected from a group consisting of trioxane, dimethoxymethane, and trimethoxymethane;
means for flowing said liquid organic fuel past said anode; and
means for flowing oxygen past said cathode.

48. The fuel cell of claim 47, wherein said fuel is dissolved in water to a concentration of between 0.1 and 2.0M.

49. A method for generating energy comprising the steps of:
providing a liquid-feed fuel cell; and
operating the liquid-feed fuel cell using an organic fuel selected from a group consisting of trioxane, dimethoxymethane and trimethoxymethane.