

United States Patent [19]

Ahmed et al.

[54] METHANOL PARTIAL OXIDATION REFORMER

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- [52] U.S. Cl. 422/111; 422/193; 429/20
- [58] **Field of Search** 429/19, 20; 423/651;
 - 422/109, 111, 193

[56] References Cited

U.S. PATENT DOCUMENTS

4,365,006	12/1982	Baker 429/17
4,522,894	6/1985	Hwang et al 429/19 X

[11] **Patent Number:** 5,939,025

[45] **Date of Patent:** Aug. 17, 1999

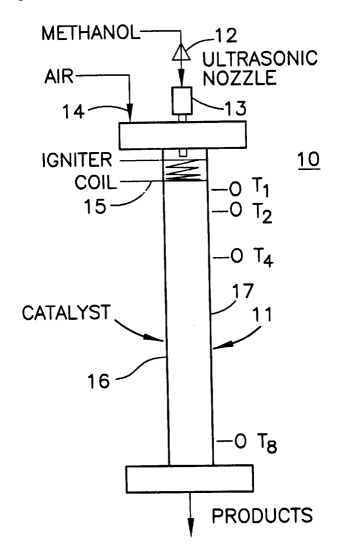
4,789,540	12/1988	Jenkins 422/111 X
4,902,586	2/1990	Wertheim 429/20
5,248,566	9/1993	Kumar et al 429/19

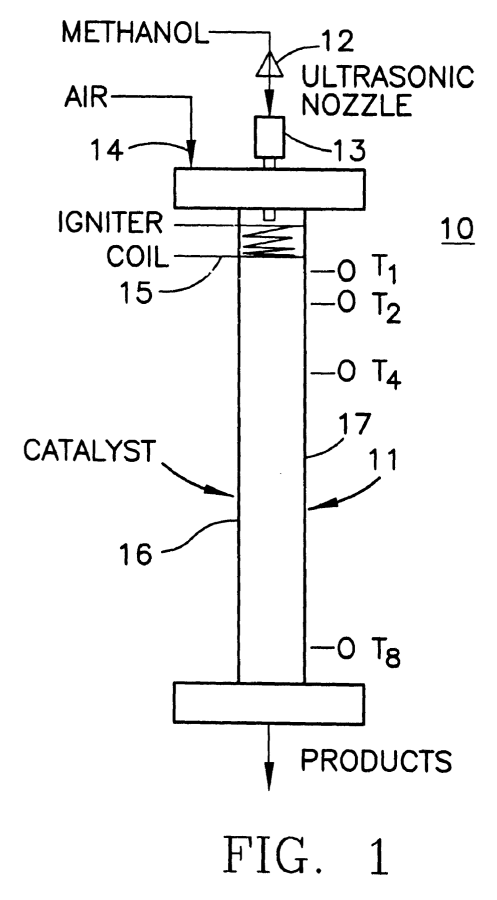
Primary Examiner—Stephen Kalafut Attorney, Agent, or Firm—Emrich & Dithmar

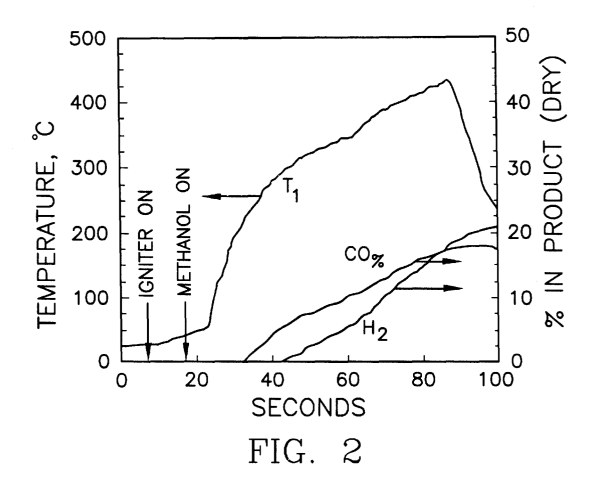
[57] ABSTRACT

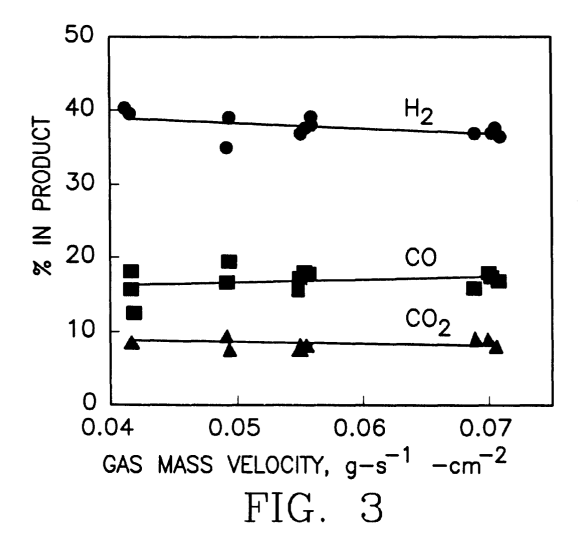
A partial oxidation reformer comprising a longitudinally extending chamber having a methanol, water and an air inlet and an outlet. An igniter mechanism is near the inlets for igniting a mixture of methanol and air, while a partial oxidation catalyst in the chamber is spaced from the inlets and converts methanol and oxygen to carbon dioxide and hydrogen. Controlling the oxygen to methanol mole ratio provides continuous slightly exothermic partial oxidation reactions of methanol and air producing hydrogen gas. The liquid is preferably injected in droplets having diameters less than 100 micrometers. The reformer is useful in a propulsion system for a vehicle which supplies a hydrogen-containing gas to the negative electrode of a fuel cell.

21 Claims, 7 Drawing Sheets









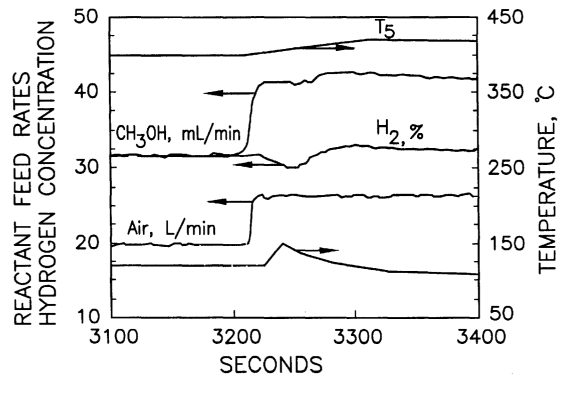
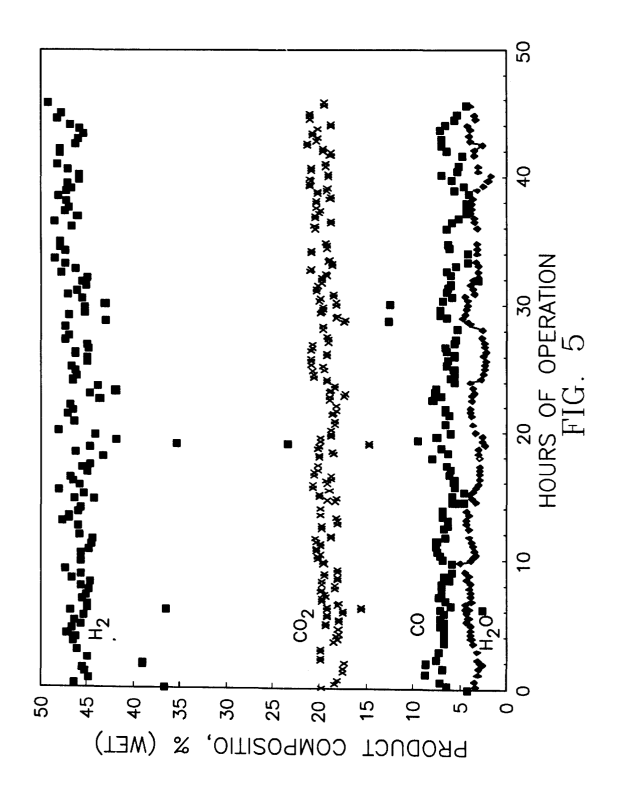
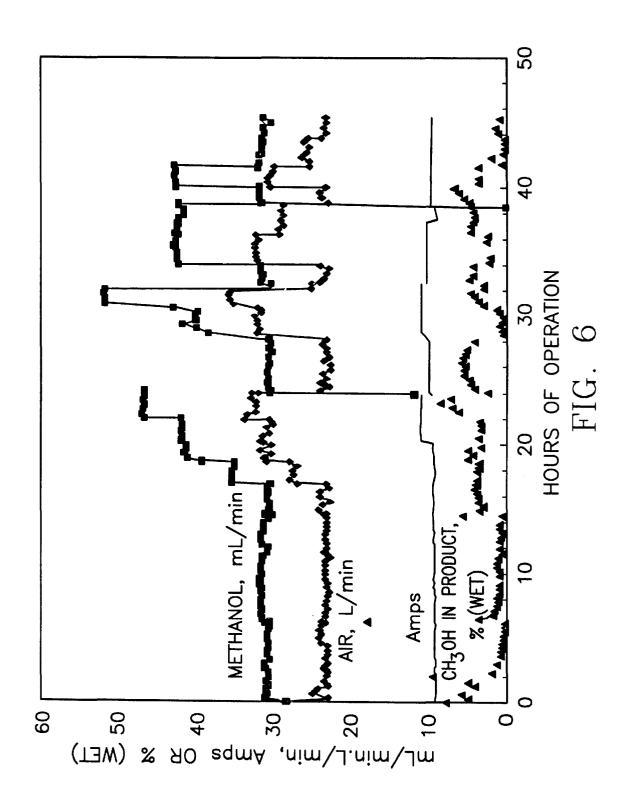
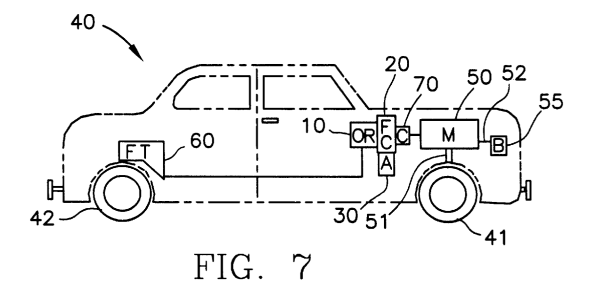


FIG. 4







METHANOL PARTIAL OXIDATION REFORMER

CONTRACTUAL ORIGIN OF THE INVENTION

The United States Government has rights in this invention 5 pursuant to Contract No. W-31-109-ENG-38 between the U.S. Department of Energy and The University of Chicago representing Argonne National Laboratory.

BACKGROUND OF THE INVENTION

This is an improvement in the invention described in U.S. Pat. No. 5,248,566 issued Sep. 28, 1993, the disclosure of which is herein incorporated by reference.

Fuel cells are being developed for use in automotive propulsion systems as alternatives for the internal combustion engine in buses, vans, passenger cars and other four wheel vehicles. The major motivations for developing fuel cell powered vehicles are low emissions of pollutants, high fuel energy conversion efficiencies, superior acceleration, low noise and vibration and the possible use of coal or 20 biomass derived alcohols rather than petroleum-base fuels. Although petroleum based fuels can also be used. The present invention is directed most specifically to systems for using methanol as a fuel.

The two most important operational requirements for a 25 stand-alone fuel cell power system for a vehicle are the ability to start-up quickly and the ability to supply the necessary power and demand for the dynamically fluctuating load. The rapid start-up requirement is obvious.

Alcohols such as methanol are likely fuels for use in fuel 30 cells for transportation applications. Methanol is a commodity chemical that is manufactured from coal, natural gas and other feed stocks, while ethanol is often produced from grain. For use in a fuel cell, however, alcohol must first be converted (reformed) to a hydrogen rich gas mixture. The 35 desired features for such a fuel reformer include rapid start-up, good dynamic response, fuel conversion, small size and weight, simple construction and operation and low cost.

Methanol has been used in steam reforming for providing engines, see Koenig et al. U.S. Pat. No. 4,716,859 and water, as a reaction product from a fuel cell, has been recycled for use in steam reforming of methanol, see Baker U.S. Pat. No. 4,365,006. Steam reforming of methanol is endothermic and complicates, by its energy requirements, its use in a vehicle. 45 reformer; Supplying the hydrogen rich gas on demand in an intermittent variable demanding environment is also a difficult requirement to meet and has been addressed by Ohsaki et al. U.S. Pat. No. 4,988,580 but this suggestion is not applicable to a small, mobile system. The catalytic exothermic partial 50 oxidation-reforming of fuels to produce hydrogen-rich gas streams is known, see Rao U.S. Pat. No. 4,999,993. The use of a partial oxidation reformer had not been used in a vehicle to accomplish the purposes of this invention prior to the disclosure of the Kumar et al. '566 patent which is satis- 55 factory for its intended purposes, but was based on theoretical considerations.

The subject invention is an improvement of that disclosed in the Kumar et al. '566 patent and relates to the use of specific reactor designs and catalyst along with mechanism 60 for controlling the oxygen to methanol mole ratio to control operating temperatures to produce a commercially viable system.

SUMMARY OF THE INVENTION

This invention relates to a partial oxidation reformer which has a longitudinal extent at least 1.5 times its diameter and which contains an oxide catalyst for partially oxidizing and reforming mixtures of water, air and methanol into a hydrogen containing gas.

The invention, in one aspect, combines a particular oxidation reformer with a fuel cell for using the hydrogencontaining gas put out by the partial oxidation reformer and air to produce d.c. power which operates an electric motor in a transportation vehicle.

Another aspect of the invention is that the partial oxida-10 tion reformer is provided with mechanism for introducing methanol and water in small droplets and intimately mixing the droplets with air prior to partial oxidation.

The invention consists of certain novel features and a combination of parts hereinafter fully described, illustrated in the accompanying drawings, and particularly pointed out in the appended claims, it being understood that various changes in the details may be made without departing from the spirit, or sacrificing any of the advantages of the present invention.

BRIEF DESCRIPTION OF THE DRAWINGS

For the purpose of facilitating an understanding of the invention, there is illustrated in the accompanying drawings a preferred embodiment thereof, from an inspection of which, when considered in connection with the following description, the invention, its construction and operation, and many of its advantages should be readily understood and appreciated.

FIG. 1 is a schematic diagram of the partial oxidation reformer of the present invention;

FIG. 2 is a graphical representation of the relationship of temperature and gas concentrations with time during ignition;

FIG. 3 is a graphical relationship of product gas distributions at steady state as a function of reactant mass velocity:

FIG. 4 is a graphical illustration of the relationships of the a hydrogen rich gas stream from mobile combustion 40 transient behavior of the reformer: hydrogen concentration and temperatures during a step change in processing rates with the Oxygen/Methanol Mole Ratio of 0.25;

> FIG. 5 is a graphical representation of the product compositions with hours of operation via partial oxidation

> FIG. 6 is a graphical representation of the relationship between the feed rates of methanol and air, the current to the igniter and the methanol and the product with the hours of operations; and

> FIG. 7 is a schematic illustration of a vehicle incorporating the partial oxidation reformer and a fuel cell combination.

DESCRIPTION OF THE PREFERRED EMBODIMENT

In a fuel cell system operating on methanol, the overall reaction is:

$$CH_3OH + \frac{3}{20_2} \rightarrow CO_2 + 2H_2O \tag{1}$$

but the methanol must first be converted to hydrogen by either the steam reforming reaction,

$$CH_{3}OH(1)+H_{2}O(1) \rightarrow 3H_{2}+CO_{2}\Delta H_{298}=+131 \text{ kJ}$$
 (2)

or the partial oxidation reaction,

$$CH_{3}OH(1) + \frac{1}{2}O_{2} \rightarrow 2H_{2} + CO_{2}\Delta H_{298} = -155 \text{ kJ}$$
 (3)

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or some combination of the two. For either of these overall reforming reactions, the reaction mechanism involves several steps.

The steam reforming reaction is endothermic, i.e., requires the input of thermal energy via indirect heat transfer, leading to the need for a relatively large heat exchange surface area and consequently a large reformer. On the other hand, the partial-oxidation reaction is exothermic. No indirect heat transfer is required within the reactor. Further, since the reformer is heated up to its operating 10 can also be achieved by operating at low oxygen-totemperature by direct oxidation of the fuel, it starts much more rapidly than a steam reformer. Since no heat transfer is required, the reactor is compact and lightweight. Moreover, the system design is simple and is dynamically responsive to load changes.

The advantages of the partial oxidation reformers are compactness, a simple system design, rapid start-up capability, and good dynamic response, are especially attractive for fuel cell systems which have one or more of the following: space/weight limitations, frequent start-up and 20 shut down, and operation at loads that vary often and rapidly. Transportation fuel cell systems represent an application where all of these features are common.

Although reaction (3) shows the partial oxidation reaction taking place with an oxygen-to-methanol mole ratio of 0.5, 25 which may be preferred at start-up conditions, this reaction can be conducted at a lower oxygen-to-methanol ratio in which case the product also contains some carbon monoxide. Reaction (4) shows an example with an oxygen-tomethanol mole ratio of 0.25, 30

The carbon monoxide formed is subsequently converted to carbon dioxide and additional hydrogen via the water gas

$$CO+H_2 \rightarrow CO_2+H_2$$
 (5)

At an oxygen-to-methanol ratio of 0.25, the reaction is only marginally exothermic; indeed, the reaction becomes thermally neutral at an oxygen-to-methanol ratio of 0.23. Operating at a low oxygen-to-methanol ratio is also advantageous in reducing the amount of nitrogen introduced into the system.

Referring to FIG. 1, there is a partial oxidation reformer 11 with a total volume of 0.8 L. A liquid pump 12 delivers the liquid methanol to an ultrasonic nozzle 13 located at the top of the reactor 11. The nozzle 13 creates a mist of fine liquid methanol droplets and sprays it into the reactor 11. The air stream is fed in tangentially through inlet 14 from the side (similar to a cyclone separator), just above the nozzle 13 tip.

Injecting the liquid methanol as fine droplets permits uniform distribution of the methanol across the reactor cross-section. It also prevents local cold spots that might 55 otherwise be formed by deposition of a large drop of liquid on a surface. The tangentially fed air stream creates a turbulence which facilitates a mixing between the liquid methanol droplets and air.

The methanol droplets are carried down by the air stream 60 past an electrical igniter coil 15. The igniter 15 provides sufficient heat to vaporize part of the methanol to enable the partial-oxidation reaction in the subsequently catalyst section. The igniter mechanism may also be an electrical hot wire, a Pd catalyst, a spark plug or a glow plug.

The catalyst section consists of a ceramic honeycomb 16 (400 channels per square inch) disks coated with the copper zinc oxide catalyst. The honeycombed supports offer the advantages of uniform flow distribution and a low pressure drop within the reactor. The product stream emerges at the bottom of reactor 11. There may be a layer 17 of zirconia pellets (to serve as thermal mass) above the catalyst layer 16.

The air feed rate is used to control the reactor temperatures and hydrogen concentration in the product. The reactor temperatures increase with increasing oxygen-to-methanol mole ratio. Higher hydrogen concentration in the reformer methanol ratios. A ratio of 0.5 or higher is used for rapid start of the reformer; as the reformer heats up to the operating temperatures, the oxygen-to-methanol ratio is reduced to between 0.23-to about 0.4.

The reactor 11 was operated at various catalyst loadings and at various methanol and air feed rates. The reactor performance was monitored by recording flow rates, temperatures and gas compositions. The product gas was analyzed for hydrogen, carbon monoxide and carbon dioxide with the help of on-line detectors. A gas chromatograph was used to periodically analyze the gas stream for all major components (nitrogen, oxygen, carbon oxides, hydrogen, methane, water and methanol).

The reformer 10 illustrated in FIG. 1 consists of a cylindrical pipe 11 with an internal diameter of 5.1 cm (2) in.). Liquid methanol (or a methanol/water mixture) was sprayed into the top of the reactor 11 with an ultrasonic nozzle 13. Air was introduced tangentially at inlet 14 and mixed with the liquid mist as they travelled downward toward the catalyst 16. A nichcrome igniter 15 is used to start up the reformer 10.

A copper zinc oxide catalyst was supported on honeycomb disks. 5-cm in diameter by 2.5-cm high, weighing ~30 g each. A number of thermocouples T_i-T_8 were used to shift reaction. Reaction (5) is the water gas shift reaction: 35 record the temperature along the length of the reactor. Part of the reformer product stream was cooled to remove all condensables and then analyzed for hydrogen, carbon monoxide, and carbon dioxide by on-line detectors. Samples of the reformate gas were also analyzed by a gas chromatograph. The reactor has been operated at methanol liquid flow rates of 25-60 ml/min and air flow rates of 20-54 L/min. At 60 ml/min methanol and 54 L/min of air, the gas mass velocity based on the reactor cross-section is 0.1 gs⁻¹ cm⁻², and the space velocity was 1.8 s⁻¹. The steady-state data 10 for methanol. It is a compact cylindrical reaction vessel 45 reported here were obtained at oxygen-to-methanol ratios of ~0.25. This lower (than 0.5) ratio was maintained to achieve higher hydrogen concentrations in the product and, to prevent the high catalyst temperatures (detrimental to catalyst) that result from oxygen-rich feeds.



Each test was started with the reactor 10 at room temperature. The air flow was first established at the desired rate. The igniter power was then switched on, and the coil 15 was allowed to heat up for 1-10 s before the methanol flow was started. A typical plot of temperature and gas compositions versus time is shown in FIG. 2. The temperature just above the catalyst section (3 cm below the igniter coil) is given by curve T_1 . The concentrations of hydrogen and carbon oxides ($CO_x = CO_{--}CO_2$), on a dry basis, are also shown in FIG. 2. The T_1 curve shows that the ignition takes place within 5 seconds after the methanol flow is started. The first traces of carbon oxides appear 7 seconds later, and the first trace of hydrogen is observed 23 seconds after the methanol flow is started. The delay between the CO_x and H_2 curves is partly due to the hydrogen analyzer having a slower response time than the infrared analyzers for the carbon oxides.

The reactor was operated at steady-state conditions at various methanol and air flow rates. The concentrations of hydrogen, carbon monoxide, and carbon dioxide as a function of the reactant gas mass velocity are shown in FIG. 3. For each of these tests the oxygen-to-methanol molar ratio was maintained at ~ 0.25 . The data indicate that the gas compositions are slightly affected by the mass velocity: H₂ and CO₂ decrease, and CO increases at higher mass velocities. Over the range of mass velocities tested, hydrogen 10 various from 37% to 39%; carbon monoxide 16% to 18% and carbon dioxide 8% to 9%. The product stream was found to contain 1-4% methanol and an average of 1.5% methane. The product stream, when passed through a water-gas shift reactor, would convert all the unreacted methanol and the carbon monoxide to carbon dioxide and hydrogen. Thus, the shifted reformate (neglecting further conversion of the 1-5% unreacted methanol in the gas) would yield a product containing ~47% hydrogen (dry basis). This value is 9% lower than the theoretical maximum for these reaction 20 combination of the partial oxidation reformer 10 and the fuel conditions.

Transient Tests

Steady State Tests

An important requirement of reformers for transportation fuel cell systems is their ability to provide varying amounts of hydrogen on demand, while maintaining the product gas 25 compositions. The dynamic response of the reactor 10 was tested by imposing a step change in the reactant flow rates while maintaining a constant oxygen to methanol ratio. An example is shown in FIG. 4, where the methanol flow rate was increased by 33%, from 30 to 40 ml/min, accompanied 30 by a proportional increase in the air flow rate. The resulting effect on hydrogen concentration and the temperatures just above and just below the catalyst bed, T_4 and T_8 , respectively, are shown in the figure. If the hydrogen production rate had not increased after the step change, the 35 supported on silica. The copper oxide-zinc oxide is preferred increase in the air flow alone would have dropped the hydrogen concentration to 28%. The hydrogen curve in the figure, however, shows only a minor (1.5%) and temporary drop to 30%. This indicates that the amount of hydrogen produced also increased with the increase in feed rate. The hydrogen concentration returns to its original level in ~80 seconds. Among the temperature curves, T_8 shows a damped response, compared to T₄ because of the thermal mass (catalyst and walls) between the two thermocouples.

Referring to FIGS. 5 and 6, there is shown the relation- 45 ships for the methanol and air feed rate as well as the amperage to the ignition coil for almost fifty hours of operation and the product mix (FIG. 5) as well as the unreacted methanol in the product (FIG. 6). Of importance is the fact that the amount of hydrogen in the product stays 50 size distribution of the methanol droplets, it being preferred relatively constant between about 45 and 50% even during variations in the methanol feed rate and the amount of methanol in the product also remains relatively constant if under 10% showing that over 90% of the methanol is reacted in the reformer of the present invention.

Referring now to FIG. 7, there is illustrated an automobile 40 representative of the type of vehicles in which the subject invention is useful. The vehicle 40 is provided with a pair of front wheels 41, a pair of rear wheels 42, and an electric motor 50 connected to one of the pairs 41, 42 as by a drive shaft 51 and is electrically connected by suitable means 52 to a battery 55. The battery 55 can be used to start the vehicle 40 in the same manner batteries function with internal combustion engines and to run accessories. A partial oxidation reformer **10** of the type previously described, is in liquid 65 or gas communication with a fuel tank 60 and is connected to a fuel cell 20 of the type previously described. An

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afterburner **30** is connected to the off gases from the negative electrode or anode 21 of the fuel cell and is used to react the remaining hydrogen in the gas leaving the anode 21 to extract heat therefrom for either heating or cooling the passenger compartment of the automobile 40, as required. A motor controller 70 is interposed between the fuel cell 20 and the electric motor 50 and coordinates the dc power output from the fuel cell 20 and the variable speed requirements for the motor 50. Such motor controllers are well known in the art. Also in FIG. 7, there is illustrated a single feed from the fuel tank 60 to the partial oxidation reformer 10, it may be that multiple feeds will be provided to the partial oxidation reformer 10. Moreover, it is possible that the partial oxidation reformer 10 will be a multiple zone 15 reformer, with the final zone thereof containing either an oxidation catalyst for converting carbon monoxide to carbon dioxide or containing a methanation catalyst for converting carbon monoxide to methane.

The main object of this invention is obtained by the cell 20 which provides rapid response to variable acceleration demands by the motor 50 due to the exothermic reaction, thereby obviating the need for thermal energy input to the partial oxidation reformer 10 during periods of acceleration or increased power demand.

Other objects of this invention have been attained by the novel partial oxidation reformer 10 disclosed having means for mixing droplets of methanol and air in a turbulent zone in the reactor, alone or in combination with a fuel cell suitable for a vehicle and in combination with the necessary components for a vehicle.

A variety of catalysts may be used, preferably the catalyst is an oxide catalyst and more preferably, the catalyst is one or more of copper oxide-zinc oxide, FeZnO and cobalt and, as previously indicated, it is preferred to be supported on a honeycomb support for the purposes and advantages hereinbefore set forth.

At initial start-up, it is preferred that the oxygen to 40 methanol mole ratio be higher than 0.5; however, a steady state is preferred that the oxygen to methanol mole ratio be in the range of from about 0.23 to about 0.4. By controlling the oxygen to methanol mole ratio, the temperature of the reactants in contact with the catalyst is controlled. Preferably, the temperature of the reactants in contact with the catalyst is less than about 500° C., and more particularly less than about 450° C. The preferred temperature is between about 350-400° C.

Another important aspect of the present invention is the that the size distribution be in the range of from about 20 to about 50 micrometers. It is also advantageous to have the droplet size be less than about 50 micrometers and less than about 100 micrometers will provide advantages.

Another important aspect of the invention is the turbulence generated by introducing the air through the inlet 14 tangentially to the fine droplets of methanol introduced through the ultrasonic nozzle 13. By introducing the air tangentially, turbulence is encouraged and therefore, intimate mixing of the air and methanol occurs and also helps prevent liquid droplet deposition on the reactor wall.

It is contemplated that a minor amount of water will be added with the methanol in order to facilitate conversion of the carbon monoxide via the water shift reaction previously set forth to carbon dioxide. The amount of water in the methanol is preferably less than about 50 volume percent and more preferably in 20-30% volume range.

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Because it is disadvantageous for large droplets of liquid to contact the catalyst bed 16, there may be inserted between the catalyst bed 16 and igniter coil 15, a bed of ceramic particles, for instance, zirconia. These particles may be 1/4 inch pellets of zirconia whose purpose is simply to insure that liquid drops of water and methanol do not intermittently contact the active catalyst. If the catalyst is contacted with too much liquid, the temperature of the catalyst drops and the catalyst may become temporarily inert, until it heats up again.

The present invention is particularly useful with fuel cells of the type described in FIG. 2 of the incorporated '566 patent, with the product from reactor 10 being directed to the anode of the fuel cell illustrated in the '566 patent.

The methanol partial oxidation reformer's advantages can 15 best be described relative to the other reformers being developed or used with fuel cell systems. These prior art reformers are based on the steam reforming reaction, where the reactants, methanol and water, are first vaporized and preheated before they can be fed into the reactor. The 20 inventive partial oxidation reformer requires no prevaporization or preheating.

The steam reforming reaction itself is strongly endothermic and so considerable heat must be provided in order to maintain the reactor temperatures. Since the heat transfer is 25 indirect, i.e., transferred from across a solid wall, the required rate of heat transfer is accomplished with a large heat transfer area. Consequently, the reactor designs become complex, large and heavy. Moreover, a burner is required to generate the heat that is to be provided to the reactor. The 30 partial oxidation reformer, in contrast, is a simple cylindrical piece of hardware 10. No heat exchange surfaces are required, and therefore, this type of reactor is compact and lightweight.

During cold starts, the steam reformer catalysts must be 35 heated by passing hot gases on the other side of a wall. This is a slow process and requires a long time. For example, the start-up times for the steam reformer in the DOE fuel cell takes -45 minutes to reach operating temperatures. The partial oxidation reformer operates on an exothermic reac-40 tion which heats up the gas stream and the catalyst simultaneously. This enables this reactor 10 to have very short start-up times. The methanol partial oxidation reformer developed herein has reached operating temperatures within 1 minute (FIG. 2).

Fuel cell systems using steam reformers are limited in their dynamic response. This is because during load changes both feed and heat input rates into the reformer need to change simultaneously. While the feed rate can be increased rapidly, the heat input is much slower because it occurs by 50 indirect heat transfer. The partial oxidation reformer requires no indirect heat exchange and therefore can respond to load changes as fast as the feed rates can be changed.

While there has been disclosed what is considered to be the preferred embodiment of the present invention, it is 55 said partial oxidation catalyst. understood that various changes in the details may be made without departing from the spirit, or sacrificing any of the advantages of the present invention.

The embodiments of the invention in which an exclusive property or privilege is claimed are defined as follows:

1. A partial oxidation reformer comprising a longitudinally extending chamber having a methanol, a water and an air inlet and an outlet, igniter mechanism in said chamber for igniting a mixture of methanol and air, a partial oxidation catalyst in said chamber spaced from said inlet for convert- 65 ing methanol and oxygen to carbon dioxide and hydrogen, and means for controlling the oxygen to methanol mole ratio

to provide continuous exothermic partial oxidation reactions of methanol and air producing hydrogen gas, controlling and maintaining the oxygen to methanol mole ratio in the range from between about 0.25 to about 0.4 for continuous operation and at higher values for start-up operation.

2. The partial oxidation reformer of claim 1, wherein said chamber is cylindrical having a ratio of the length to the diameter of not less than about 1.5.

3. The partial oxidation reformer of claim 2, and further comprising means for forming the methanol at the inlet into droplets having a diameter of less than 100 micrometers.

4. The partial oxidation reformer of claim 3, wherein the methanol droplets have a size less than about 50 micrometers

5. The partial oxidation reformer of claim 3, wherein the methanol droplets have a size distribution in the range of from about 20 to about 50 micrometers.

6. The partial oxidation reformer of claim 3, wherein said methanol droplets are introduced into said reactor parallel to the longitudinal axis of said chamber and said air inlet introduces air to said chamber tangentially to the longitudinal axis of said chamber.

7. The partial oxidation reformer of claim 1, wherein said means to control the oxygen to methanol mole ratio is operated to maintain a temperature of the reactants in contact with said catalyst in the reformer at less than about 500° C.

8. The partial oxidation reformer of claim 1, wherein said means for controlling the oxygen to methanol mole ratio is operated to maintain a temperature of the reactants in contact with said catalyst in the reformer of less than about 450° C.

9. The partial oxidation reformer of claim 1, wherein said means for controlling the oxygen to methanol mole ratio is operated to maintain a temperature of the reactants in contact with said catalyst in the reformer of less than about 400° C.

10. The partial oxidation reformer of claim 1, and further comprising means for forming the methanol into droplets and wherein air is mixed with the methanol droplets to form a mist at an end of the reactor away from said outlet.

11. The partial oxidation reformer of claim 10, wherein the catalyst is CuO—ZnO.

12. The partial oxidation reformer of claim 1, wherein the catalyst is an oxide.

13. The partial oxidation reformer of claim 12, wherein the weight of catalyst is not less than 11 kg per liter per minute of liquid methanol.

14. The partial oxidation reformer of claim 1, wherein said igniter mechanism is one or more of an electrical hot wire, a Pd catalyst, a spark plug or glow plug.

15. The partial oxidation reformer of claim 1, wherein said igniter is electrically operated.

16. The partial oxidation reformer of claim 1, wherein a bed of ceramic pellets is positioned between said inlets and

17. A partial oxidation reformer comprising a longitudinally extending chamber having a methanol, a water and an air inlet and an outlet, igniter mechanism in said chamber for igniting a mixture of methanol and air, a partial oxidation catalyst in said chamber spaced from said inlet for converting methanol and oxygen to carbon dioxide and hydrogen, and means for controlling the oxygen to methanol mole ratio to provide continuous exothermic partial oxidation reactions of methanol and air producing hydrogen gas, forming the methanol into droplets and mixing air with the methanol droplets to form a mist at an end of the reactor away from said outlet.

18. A partial oxidation reformer comprising a longitudinally extending chamber having inlets for liquid methanol, water and air and an outlet, mechanism for forming the methanol and water in droplets having diameters less than 100 micrometers, igniter mechanism for igniting a mixture of methanol and air, a partial oxidation catalyst selected from the group consisting of CuO—ZnO, FeZnO and Co supported on SiO₂ in said chamber spaced from said inlets for converting methanol and oxygen to carbon dioxide and hydrogen, and means for controlling the oxygen to methanol mole ratio to provide continuous exothermic partial oxidation reactions of methanol and air producing hydrogen gas.

19. The partial oxidation reformer of claim 18, wherein said igniter mechanism is one or more of an electrical hot

wire, a Pd catalyst, a spark plug or glow plug and said methanol droplets have a diameter less than about 50 micrometers.

20. The partial oxidation reformer of claim **19**, wherein said means for controlling the oxygen to methanol mole ratio is operated to maintain the ratio between about 0.25 to about 0.4 for continuous operation and at higher values for start-up operation.

21. The partial oxidation reformer of claim **20**, wherein 10 said methanol droplets are introduced into said reactor parallel to the longitudinal axis of said chamber and said air inlet introduces air to said chamber tangentially to the longitudinal axis of said chamber.

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