Method and apparatus for methane production

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Abstract: In accordance with the invention there is presented herein a method of synthesizing methane by 1-step co-electrolysis of H2O and CO2 in the presence of an electrolyte comprising at least one superprotonic conducting acid compound of the general formula © : \( \text{M}_x \text{H}_y (\text{A}_2 \text{O}_z) \), wherein \( x = 1, 2, \) or \( 3; y = 1, 2, 3, 4 \) or \( 5; z = 1 \) or \( 2; M \) is at least one of Li, Na, Rb, Cs, Tl, NH4; and A is S, Se, P, As.

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METHOD AND APPARATUS FOR METHANE PRODUCTION

TECHNICAL FIELD

In the field of electrolytic conversion of carbon dioxide to methane there is suggested a method and an apparatus for 1-step electrolytic methane production by co-electrolysis of the reactants.

BACKGROUND

In recent years, many countries including Denmark, have invested heavily in alternative sources of energy in order to reduce national dependencies on fossilized fuels and to reduce emissions of carbon dioxide to the atmosphere. Particularly wind power has become an important energy source, in Denmark supplying up to more than 35% of the total energy consumption of the country.

Unfortunately, wind power has the drawback that power consumption in the society surrounding a wind power plant or wind power farm may not necessarily coincide with optimum wind conditions to the extent that many wind power plants are taken off-line from the power grid, even when optimal conditions are present for wind energy production, as end consumers are missing. This situation has led to a wide range of proposals aimed at storing the surplus energy generated by wind power plants for a later time.

One such proposal currently considered is to use surplus energy, preferably surplus electrical energy, for the reconversion of carbon dioxide into combustible carbon compounds such as e.g. methane or higher alkanes, by
reduction of carbon dioxide with water in electrolytic cells.

Traditionally, this electrolytic reconversion has been obtained in one of at least two possible two-step conversion processes, either by first obtaining \( \frac{3}{4} \) by electrolysis of water followed by subsequent conversion of \( \text{CO}_2 \) with \( \text{H}_2 \) in a Sabatier process, or co-electrolysis of \( \text{CO}_2 \) and \( \text{H}_2 \text{O} \) to form a syngas mixture of \( \text{CO} \) and \( \frac{1}{4} \), which is then subsequently reacted in a Fischer-Tropsch process to yield methane and water. However, as the overall energy efficiency of the two-step processes is low, research initiatives in recent years have sought to improve this deficiency.

One possible solution is the substitution of the 2-step conversion process with a 1-step co-electrolytic conversion process of \( \text{H}_2 \text{O} \) and \( \text{CO}_2 \) to \( \text{CH}_4 \) and \( \text{O}_2 \). The theoretical energy savings from a 1-step methanization process compared to a 2-step methanization from \( \text{CO}_2 \) is about 17% at atmospheric pressure and room temperature and due to increased reaction efficiency and changed conditions at increased temperatures, further gains of about 20% energy savings may be achievable. Hence, a 1-step co-electrolytic conversion process of \( \text{H}_2 \text{O} \) and \( \text{CO}_2 \) to \( \text{CH}_4 \) and \( \text{O}_2 \) is highly desirable over the 2-step methods of the prior art.

Ebbesen et al. report (Chem. Rev., 2014, 114, 10697-10734), that solid proton conducting electrolysis cells (SPCEC) have been applied for electrolysis of water combined with electrochemical conversion of \( \text{CO}_2 \). This process is often mentioned as co-electrolysis of \( \text{CO}_2 \) and \( \text{H}_2 \text{O} \), although in the mechanism reactions (1)-(3) only \( \text{H}_2 \text{O} \) is involved (in fact electrolyzed). Rather, \( \text{CO}_2 \) is electrochemically
converted to CH4 in a one-step process by reaction with protons released at the cathode, reaction (2). Also reported was the use of SPCECs for production of hydrogen with simultaneous upgrading of methane to higher hydrocarbons.

Using proton-conducting oxides as electrolyte and suitable BCZYZ electrodes (BaCe0.5Zr0.3Y0.16Zn0.04O3-δ) as electrolyte with an iron based composite fuel electrode (cathode), it was found that it was possible to convert 1-2% of the available CO2 into methane at 650°C following an overall 1-step co-electrolytic conversion reaction of the type:

(1) Anode reaction: \(4H_2O \rightarrow 2O_2 + 8H^+ + 8e^-\)
(2) Cathode reaction: \(CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O\)
(3) Overall reaction: \(CO_2 + 2H_2O \rightarrow CH_4 + 2O_2\)

However, as the equilibrium partial pressure of CH4 is low at 650°C, a part of the CO2 is converted into CO instead in accordance with the below reaction scheme:

(1a) Anode reaction: \(2H_2O \rightarrow 2O_2 + 4H^+ + 4e^-\)
(2a) Cathode reaction: \(2CO_2 + 4H^+ + 4e^- \rightarrow 2CO + 2H_2O\)
(3a) Overall reaction: \(2CO_2 \rightarrow 2CO + O_2\)

It is an aim of the present inventors to improve on these known processes, whereby methane is generated in a one-step process by co-electrolysis of CO2 and H2O in accordance with reactions (1)-(3), in particular to improve the conversion rate of methane over carbon monoxide, and to reduce the overall energy loss in the process.

To this aim, and with the present disclosure, the present inventors report a novel one-step electrolytic conversion
of CO₂ and H₂O to methane and oxygen in the presence of an electrolyte comprising a superprotonic conducting acid compound in solid or molten form of the general chemical formula \( \text{M}_x \text{H}_y (\text{A}_04)^z \), wherein \( x \) is 1, 2, or 3; \( y \) is 1, 2, 3, 4 or 5; \( z \) is 1 or 2; \( M \) is at least one of Li, Na, K, Rb, Cs, Tl, NH₄; and \( A \) is S, Se, P, As; which method is operative at temperatures down to only slightly above the boiling point of water, but preferably operated at temperatures in the range of from 250 to 450°C, and which method has the further benefit that methane is preferentially produced at the cathode via reactions (1) through (3) rather than carbon monoxide via reactions (1a) through (3a).

The use of a superprotonic conducting acid compound in solid or molten form of the general chemical formula \( \text{M}_x \text{H}_y (\text{A}_04)^z \) have previously been suggested in the art for use as superprotonic electrolytes in relation to H₂-production for fuel cells, cf. e.g. Gohi-Urtiaga et al., Int. J. Hydrogen Energy, 2012, 37, 3358-3372, wherein the chemistry and suitability of these compounds as superprotonic electrolytes in relation to electrolytic cells are thoroughly discussed.

In the context of the present disclosure, and in accordance with the usage in the scientific literature, a superprotonic electrolyte is an electrolyte which, at and above a certain temperature or, usually, at or above a narrowly defined temperature range, exhibits very low resistance to proton conduction. Figure 1 (Figure 6 of Gohi-Urtiaga et al.) displays an Arrhenius plot of the proton conductivity of various Cs-based superprotonic acid compounds suitable for use with the present invention. As can be observed from the figure, resistance to proton conduction in the electrolyte reduces for the three
detailed salts by at least 3 decades following a
temperature increase by less than 5°C (CSH2PO4, from about
230°C to about 235°C) at ambient pressure.

It is as yet not finally established in the scientific
literature, whether the transition from protonic
conductivity to superprotonic conductivity is a true phase
transition, and hence associated with a single well-defined
temperature, or not. For CSH2PO4, at least, the transition
correlates fully with an internal rearranging of the
constituent atoms from a monoclinic, P21/m crystal packing
to a cubic, Pm-3m packing, and hence at least for this
compound may constitute a true phase transition. The
superprotonic conductivity pertains also into the molten
phase of the acid compound, where at an upper, but ill-
defined temperature or temperature interval, the molten
phase loses superprotonic conductivity due to thermal
agitation. As expected from general principles,
pressurization of the system influences the temperature of
loss of superprotonic conductivity. Table 1 comprises a
list displaying apparent transition temperatures for
superprotonic conducting acid compound in solid or molten
form of the general chemical formula \((I) \ \text{M}_x\text{H}_y(\text{A04})_z\) as
defined above at atmospheric pressure and their resulting
proton conductivity after transition (c.f. Table 1 of Gohi-
Urtiaga et al. and Figure 2).

A major problem, as has been repeatedly reported in the
literature, in using the acid compounds of formula \((I)\) as
superprotonic electrolytes in electrolytic cells is that
the acid compounds of formula \((I)\) are inherently instable
at elevated temperatures and decompose according to the
general stoichiometric equation \((4)\) given below (c.f. Gohi-
Urtiaga et al. equation \((11)\) ):
wherein the reaction is driven to the right hand side by the evaporation of gaseous water. Under moist conditions, as is known, the reaction is reversible; however, the presence of moisture in the acid compounds, such as when these are subjected to gaseous water, leads to a loss of superprotonic conductivity. The electrolytic systems comprising acid compounds of the general formula \( \text{(I)} \) as studied hitherto have been shown to lack long-term stability due to degradation of the acid compound in accordance with reaction (4). However, the present inventors through their work has established that this observation by Gohi-Urtiaga et al. is unbalanced with respect to the metal-ions, \( M \).

Rather, the present inventors observe that a number of the mentioned acid compounds may decompose by stepwise polymerization by condensation and may irreversibly yield compounds of the general formula \( M_x \text{H}_y \delta \text{A}_z \delta \text{O}_4 (s) \) (formed by the reaction \( M_x \text{H}_y \delta \text{A}_z \delta \text{O}_4 (s) + \delta \text{H}_2 \text{O} (g) \)) or ultimately MAO3 as the end product of the dehydration process, which products further are known by the present inventors to restrict the use of the acid compounds contemplated by the present inventors.

The present inventors have now surprisingly discovered that by maintaining particular levels of water partial pressures and using "dry", that is non-condensing superheated steam, i.e. superheated steam, which is not saturated with \( \text{H}_2 \text{O} \), the decomposition reactions, can be suppressed and a functional electrolyte maintained. These discoveries have allowed the present inventors to construct an electrolytic
cell wherein 1-step methanization of CO2 is possible without loss of electrolyte activity over time.

The present inventors have found that following the instructions according to the present invention, the results obtainable by the invention are not influenced by the physical state, whether solid or molten, of the acid compounds for use with the invention, as long as these are in their superprotonic conducting state. Likewise, the present inventors have found that although the temperature at which superprotonic conductivity is attained for a given acid compound displays a dependency on the external pressure of the electrolytic system, the general teaching of the present invention is not affected thereby. Rather, it is considered that the skilled person will know how to establish if superprotonic conductivity has been attained at a given combination of temperature and pressure for a given acid compound, e.g. by establishing an Arrhenius-plot in accordance with Figure 1 for the particular combination of acid compound, temperature and pressure, to the extent that the data cannot be obtained from the pertinent prior art.

As will be discussed below, it is in general preferable to operate the electrolytic cell well above a minimum temperature just sufficient to establish superprotonic conductivity for a given combination of acid compound, temperature and pressure to obtain the best results for the invention.

Recently, particular focus has been given to obtaining energy in the form of biogas from reusable sources. A drawback to the production of biogas is that considerable amounts of CO2 (up to 40% w/w) and some sulfur-containing
compounds are byproducts of the biogas-production, but attempts have been made (e.g. WO 2012/003849) to utilize the formed carbon dioxide for electrolytic reforming into syngas. However, as the presented system is vulnerable to sulfur-poisoning of the employed catalysts, and electrolytes, additional steps need to be taken to separate the sulfur-containing compounds from the syngas. Surprisingly, the system presented by the present inventors does not suffer from the above drawback within the timeframe of the performed experimental studies; which makes the electrolytic cell and the associated method of the present invention ideally suited for utilizing biogas as a source of carbon dioxide for one-step methanization.

The solutions to the above technical problems suggested by the present inventors are detailed in the below description and in the claims and are presented with respect to the below discussed definitions of terms and parametric limitations and of the associated figures.

**SUMMARY OF THE INVENTION**

In accordance with the invention there is presented herein a method of synthesizing methane by 1-step co-electrolysis of H2O and CO2 in the presence of an electrolyte comprising at least one superprotonic conducting acid compound of the general formula (I)

\[
M_{x}H_{y}(A04)^{z}, \text{ wherein } x \text{ is } 1, 2, \text{ or } 3; y \text{ is } 1, 2, 3, 4 \text{ or } 5; z \text{ is } 1 \text{ or } 2; M \text{ is at least one of } Li, Na, K, Rb, Cs, Tl, NH_{4}; \text{ and } A \text{ is } S, Se, P, As;
\]

the method comprising: a) providing a pressurized electrolytic cell comprising an anode and a cathode,
wherein the electrolyte is arranged between the anode and the cathode; and wherein hydrogen ions are produced at the anode according to an anode reaction (1)

\[
4H_2O \rightarrow 2O_2 + 8H^+ + 8e^- ;
\]

and wherein methane is produced at the cathode according to a cathode reaction (2)

\[
CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O ;
\]

the method further comprising operating the electrolytic cell by: b) providing an electrical current applied across the electrolytic cell sufficient to induce anodic and cathodic reactions according to the anode (1) and cathode (2) reactions; c) maintaining the electrolytic cell at an operating water partial pressure from 1 to 30 bar; d) maintaining the electrolytic cell at an operating temperature sufficient to induce superprotonic conduction in the electrolyte of hydrogen ions produced at the anode; e) providing a flow of gaseous H2O as superheated steam as an anode reactants flow to the anode at a flow rate sufficient to allow the anodic reaction; f) providing a flow of CO2 as a cathode reactants flow to the cathode at a flow rate sufficient to allow the cathodic reaction; g) collecting a cathode products flow from the cathodic reaction; and h) optionally, isolating methane from the cathode products flow.

Also, and likewise in accordance with the present invention (cf. Figure 3), there is disclosed an electrolytic cell system (10) comprising an electrolytic cell (11) enclosed in a pressure chamber (12); the electrolytic cell comprising an anode compartment (20) comprising an anode
and an insulating anode enclosure (24) wherein the anode is embedded, and a cathode compartment (30) comprising a cathode (31) and an insulating cathode enclosure (34) wherein the cathode is embedded, and an electrolyte (40) arranged in contact with and between the anode (21) and the cathode (31), and wherein the anode compartment (20) and the cathode compartment (30) are separated by the electrolyte (40) and an insulating gasket (41) enclosing the electrolyte; the electrolyte (40) comprising at least one superprotonic conducting acid compound of the general formula (I):

$$M^xH_y(\text{A04})_z, \text{ wherein } x \text{ is } 1, 2, \text{ or } 3; y \text{ is } 1, 2, 3, 4 \text{ or } 5; z \text{ is } 1 \text{ or } 2; M \text{ is at least one of Li, Na, K, Rb, Cs, Tl, NH}_4; \text{ and A is } S, \text{ Se, P, As};$$

wherein the anode compartment (20) further comprises an anode flow path (22) for gaseous H2O as superheated steam to the anode (21) arranged for contacting the superheated steam as an anodic reactant with the anode; and an anode products flow path (23) from the anode (21) comprising means (15) for maintaining a constant water partial pressure across the anode; wherein the cathode compartment (30) further comprises a cathode flow path (32) for gaseous CO2 to the cathode (31) arranged for contacting the CO2 as a cathodic reactant with the cathode; and a cathode products flow path (33) from the cathode (31), optionally comprising means (19) for isolating methane from a gas flow; the electrolytic cell system (10) further comprising means for circulating an electric current across the electrolytic cell (11), and respective means (13,17,18) for passing a respective gas flow through a respective anode or cathode compartment (20,30). In the experiments reported herein, the means (19) for isolating methane from a gas
flow was implemented, proving the successful production of methane according to the methods of the present invention.

Further aspects and embodiments of the invention are detailed herein in the description, in the claims and in the figures.

**BRIEF DESCRIPTION OF THE DRAWINGS**

Figure 1: Arrhenius plot of the proton conductivity of various Cs-based acid compounds suitable for use with the invention.

Figure 2: Apparent transition temperature to superprotonic conducting state for various salts.

Figure 3: Schematic of the electrolytic cell system of the invention.

Figure 4: Measured XRD pattern of CSH2PO4 electrolyte with matrix of SnP2O7 after co-electrolysis and electrolyte rearrangement and calculated diagrams of three reference compounds.

Figure 5: GC-data showing 1-step methanization in KH2PO4-electrolyte (dried gas flow).

Figure 6: GC-data showing 1-step methanization in CSH2PO4-electrolyte (dried gas flow).

Figure 7: Polarization of electrolytic cell with CSH2PO4-electrolyte at two temperatures.

Figure 8: Comparison of polarization of electrolytic cell between KH2PO4 and CsH2P04-electrolytes.

Figure 9: Voltammetric curves obtained on a Ni electrode in CSH2PO4 at 350°C.
In accordance with the invention there is presented herein a method of synthesizing methane by 1-step co-electrolysis of H2O and CO2 in the presence of an electrolyte comprising at least one superprotonic conducting acid compound of the general formula (I)

$$M^{x+}A_0^{y-}$$

wherein x is 1, 2, or 3; y is 1, 2, 3, 4 or 5; z is 1 or 2; M is at least one of Li, Na, K, Rb, Cs, Tl, NH4; and A is S, Se, P, As;

the method comprising: a) providing a pressurized electrolytic cell comprising an anode and a cathode, wherein the electrolyte is arranged between the anode and the cathode; and wherein hydrogen ions are produced at the anode according to an anode reaction (1)

$$4H_2O \rightarrow 2O_2 + 8H^+ + 8e^-$$

and wherein methane is produced at the cathode according to a cathode reaction (2)

$$CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O$$

the method further comprising operating the electrolytic cell by: b) providing an electrical current applied across the electrolytic cell sufficient to induce anodic and cathodic reactions according to the anode (1) and cathode (2) reactions; c) maintaining the electrolytic cell at an operating water partial pressure from 1 to 30 bar; d) maintaining the electrolytic cell at an operating temperature sufficient to induce superprotonic conduction in the electrolyte of hydrogen ions produced at the anode;
e) providing a flow of gaseous H2O as superheated steam as an anode reactants flow to the anode at a flow rate sufficient to allow the anodic reaction; f) providing a flow of CO2 as a cathode reactants flow to the cathode at a flow rate sufficient to allow the cathodic reaction; g) collecting a cathode products flow from the cathodic reaction; and h) optionally isolating methane from the cathode products flow.

In one embodiment, also the products of the anodic reaction are collected, preferably as an anode products flow from the anodic reaction.

In one embodiment H2O, CO2 are also individually or, respectively, both isolated from the cathode products flow and individually, respectively, both recirculated into the respective flows of anode and/or cathode reactants.

Also, and likewise in accordance with the present invention (cf. Figure 3), there is disclosed an electrolytic cell system (10) comprising an electrolytic cell (11) enclosed in a pressure chamber (12); the electrolytic cell comprising an anode compartment (20) comprising an anode (21) and an insulating anode enclosure (24) wherein the anode is embedded, and a cathode compartment (30) comprising a cathode (31) and an insulating cathode enclosure (34) wherein the cathode is embedded, and an electrolyte (40) arranged in contact with and between the anode (21) and the cathode (31), and wherein the anode compartment (20) and the cathode compartment (30) are separated by the electrolyte (40) and an insulating gasket (41) enclosing the electrolyte; the electrolyte (40) comprising at least one superprotonic conducting acid compound of the general formula (I):

$$I$$
MxHy(A04)z, wherein x is 1, 2, or 3; y is 1, 2, 3, 4 or 5; z is 1 or 2; M is at least one of Li, Na, K, Rb, Cs, T1, NH4; and A is S, Se, P, As; wherein the anode compartment (20) further comprises an anode flow path (22) for gaseous H2O as superheated steam to the anode (21) arranged for contacting the superheated steam as an anodic reactant with the anode; and an anode products flow path (23) from the anode (21) comprising means (15) for maintaining a constant water partial pressure across the anode; wherein the cathode compartment (30) further comprises a cathode flow path (32) for gaseous CO2 to the cathode (31) arranged for contacting the CO2 as a cathodic reactant with the cathode; and a cathode products flow path (33) from the cathode (31), optionally comprising means (19) for isolating methane from a gas flow; the electrolytic cell system (10) further comprising means for circulating an electric current across the electrolytic cell (11), and respective means (13, 17, 18) for passing a respective gas flow through a respective anode or cathode compartment (20, 30). An illustrative embodiment of the electrolytic cell system (10) is disclosed in Figure 3.

In general, and preferably, the means (19) for isolating methane from a gas flow will be included into the electrolytic cell system (10), in order to isolate the methane of the cathodic reaction. Nevertheless, as the products flow following the methods detailed herein is reasonably concentrated in methane, in many situations the products flow from the cathode is directly usable in downstream processes, such as e.g. combustion processes, permitting the omission of the aforesaid means (19).
In the optional embodiment comprising means (19) for isolating methane, preferably the means (19) for isolating methane is connected to the means (13) for passing a gas through an anode compartment (20) via a flow path (19a) for recirculating condensed water produced via reaction (2). In a further embodiment, the means (19) for isolating methane is connected to the means (17) for passing CO2 through the cathode compartment (20) via a flow path (17a) for recirculating unreacted CO2 obtained from the means (19) for isolating methane. This is particularly advantageous, as the product stream of unreacted CO2 often will comprise a quantity of methane, which then rather than being discarded as waste is recycled and recovered in a subsequent pass. At steady state, when recycling via flow path (17a) is enabled, methane recovery is significantly increased.

The electrolytic cell system (10) and electrolytic cell (11) described herein has proven particularly suitable for executing the method of the invention. In some embodiments, the electrolytic cell system (10) further comprises a controller configured to execute a method of the invention and of its embodiments.

The present invention relies on the surprising discovery by the present inventors that the drawback to the prior art use of superprotonic conducting acid compounds of the general formula (I) in electrolytic cells can be overcome, thereby preventing reversible and irreversible conversion of the superprotonic conducting acid compounds to non-superprotonic conducting compounds, by providing and maintaining gaseous H2O as superheated steam at an operating water partial pressure from 1 to 30 bar across the
electrolytic cell during the 1-step co-electrolysis of H2O and CO2 to CH4 and O2.

The present inventors speculate, without considering themselves bound by this theory, that the surprising dual action of providing and maintaining gaseous H2O as superheated steam at an operating water partial pressure of from 1 to 30 bar across the electrolytic cell during the 1-step co-electrolysis of H2O and CO2 to CH4 and O2, is due to a combined effect of the high water partial pressure, which serves to drive reaction (4) to the left, while at the same time the superheated steam (i.e. steam heated to above the condensation point of water in the steam) serves to assure, that water does not condense in the electrolytic cell, thereby preventing irreversible conversion of the electrolyte or unwanted dissolution of the electrolyte.

Preferably, the operating water partial pressure is at least 2 bar, at least 4 bar, at least 6 bar, preferably at least 8 bar, and even more preferably at least 10 bar. For the acid compounds of formula (III) (cf. below), optimally the operating water partial pressure shall be at least 8 bar.

In the electrolytic cell system (10) of the invention, the method of the invention is implemented by providing a water-to-steam evaporator (14) arranged on the anode flow path (22) for providing gaseous H2O as superheated steam to the anode prior to the electrolytic cell, a means (15) for maintaining a constant water partial pressure across the anode, preferably in the form of a heated backpressure valve, and by operating the electrolytic cell (11) at a temperature which is higher than the temperature of the
evaporator (14), thereby creating superheated steam in the electrolytic cell.

In the test cell described below, the operating temperature of the electrolytic cell (11) is about 50% higher than the exit temperature from the water-to-steam evaporator (14). Also the back pressure valve is heated in order to prevent condensation along the entire length of the anode flow paths (22, 23). Naturally, superheated steam can in an embodiment be prepared prior to entering the electrolytic cell; and likewise heating to avoid condensation in the back pressure valve (15) or along the anode products flow path (23) can be decoupled from the back pressure valve (15) itself.

The use of an electrolyte comprising at least one superprotonic conducting acid compound of the general formula (I) constitutes an essential element of the present invention. In the absence of such superprotonic conducting acid compounds, the reverse anode reaction dominates and the advantages of the invention cannot be realized.

As stated, the at least one superprotonic conducting acid compound has the general formula (I):

\[
\text{(I) } \text{MxHy(A04)z, wherein } x \text{ is } 1, 2, \text{ or } 3; \text{ y is } 1, 2, 3, 4 \text{ or } 5; \text{ z is } 1 \text{ or } 2; \text{ M is at least one of Li, Na, K, Rb, Cs, Tl, NH}_4; \text{ and A is } S, \text{ Se, P, As;}
\]

In an embodiment of the invention, there is disclosed a method, wherein the at least one superprotonic conducting acid compound has the general formula (IB):

\[
\text{(IB) }
\]
In an embodiment of the invention, there is disclosed a method, wherein the at least one superprotonic conducting acid compound has the general formula (II):

$$\text{(II) } \text{M}_x\text{H}_y\text{A}_0 \text{A}_4; \text{ wherein M is Li, Na, K, Rb, Cs, NH}_4; \text{ A is S, Se, P, As; and } x \text{ is 1 or 2; y is 1 or 2; and } x+y \text{ is 2 when A is S or Se and 3 when A is P or As.}$$

In a further and preferred embodiment of the invention, there is disclosed a method, wherein the superprotonic conducting acid compound has the general formula (III):

$$\text{(III) } \text{MH}_2\text{PO}_4; \text{ wherein M is Li, Na, K, Rb, Cs, NH}_4.$$  

As documented in the experimental section, the present inventors have obtained very good results for superprotonic conducting acid compounds wherein A is phosphor (P) and M is selected from periodic system group 1 metals, Li, Na, K, Rb, and Cs, or NH4, in particular for K and Cs. Hence the invention in particular and most preferred relates to a method according to any embodiment, wherein the at least one superprotonic conducting acid compound is either KH2PO4 or CSH2PO4.

In one embodiment of the invention, the electrolyte comprises a mixture of at least two of the superprotonic conducting acid compounds selected according to any of the above embodiments.

In one embodiment of the invention, the at least one superprotonic conducting acid compound is either solid,
molten or a mixture of solid and molten. As was discussed in the background section above, the superprotonic conducting acids of formulas (I), (IB), (II) and/or (III), in particular KH₂PO₄ or CSH₂PO₄, transit from a non-superprotonic conducting state to a superprotonic conducting state at a relatively narrow region of temperature, potentially even at a single temperature in correspondence with a true phase transition. As mentioned, the superprotonic conductivity pertains also into the molten phase of the acid compound, where at an upper, but ill-defined temperature or temperature interval, the molten phase loses superprotonic conductivity due to thermal agitation and/or decomposition. The present inventors have investigated KH₂PO₄ and CSH₂PO₄ as electrolytes both in solid, in molten and in solid/molten state and found that the invention can be realized over the entire superprotonic conductivity range of these two compounds without discrimination.

In one embodiment, the at least one superprotonic conducting acid compound has the general formula (IV):

\[
(IV) \quad MH_5(AO_4)_2; \text{ where in} \\
M \text{ is Li, Na, K, Rb, Cs, } NH_4; \text{ and} \\
A \text{ is S, Se, P, As.}
\]

In one embodiment, the at least one superprotonic conducting acid compound has the general formula (V):

\[
(V) \quad MH_5(P_0_4)_2; \text{ where in } M \text{ is Li, Na, K, Rb, Cs; } NH_4
\]

It has further been found that it is advantageous to include a support material for the at least one superprotonic conducting acid compound into the electrolyte. Accordingly, the invention further relates to a method, wherein the
electrolyte further comprises a support material for the at least one superprotonic conducting acid compound.

In one embodiment of the invention, the support material comprises a pyrophosphate compound of the general formula (VI):

\[(VI) \quad M_t(P_2O_7), \text{ wherein } M_t \text{ can be Si, Ge or Sn.}\]

The present inventors have found (cf. Figure 4) that the pyrophosphate compounds according to formula (VI) interact particularly favorably with the compounds of formula (III) and (IV) and, when A is phosphor (P), forming as a subspecies of the compounds of formula (I), the superprotonic conducting diphosphate compounds of the general formula (V). The formed diphosphates are highly stable under the experimental conditions employed while retaining their superprotonic conducting properties as desired.

Particularly suited are compounds of formula (VI) in combination with a polybenzimidazole (PBI) fiber reinforcement layer in the electrolyte. In particular Sn(P2O7) or Sn(P2O7) reinforced with fibers of polybenzimidazole (PBI) were found to stabilize the compounds of formulas (III), (IV) and (V) and are accordingly preferred. Alternatively, fibers of Zr02, S1O2 and/or SiC were found to be suitable alternatives.

Surprisingly, periodic system group 4 dioxides, TiO2, Zr02, and Hf02, preferably Zr02, were found at least to retard, but (likely) to prevent, the conversion of the phosphates of formula (III) into the diphosphates of formula (V). The effect was observed for the oxides alone or in combination
with polybenzimidazole (PBI) fiber reinforcement. In a preferred embodiment of the invention, the addition of at least one of SiO₂, SnO₂, TiO₂, ZrO₂, or HfO₂ to the electrolyte as a support material is contemplated.

In general, and when a support material is present, the overall concentration in the electrolyte of the at least one superprotonic conducting acid compound is at least 10% w/w, at least 20% w/w, at least 30% w/w, at least 40% w/w, at least 50% w/w, at least 60% w/w, at least 70% w/w, at least 80% w/w, or at least 90% w/w based on total electrolyte weight.

In accordance with the invention, hydrogen ions are produced at the anode according to an anode reaction (1)

\[
4H_2O \rightarrow 2O_2 + 8H^+ + 8e^-;
\]

In the art, a range of elements, alloys and substances have been proven to permit electrochemical dissociation of water according to the anode reaction (1), in particular metal oxides. In relation to the present invention, the present inventors have found that oxides of iridium, in particular IrO₂ and/or Ir₂O₃, is/are advantageously comprised in the anode.

In accordance with the invention, methane is produced at the cathode according to a cathode reaction (2)

\[
CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O;
\]

In the art, a range of elements, alloys and substances have been proven to permit electrochemical association of CO₂ and H⁺ to methane according to the cathode reaction (2).
The inventors have found that both ceramics, such as ceramics comprising at least one of WC, MOC2 and/or COB2 are suitable as cathode material, but in particular it has been found that the cathode should comprise at least one of Ni, alloyed Ni, Co, Ru, Rh, or a combination thereof. Preferably, Ni or alloyed Ni is advantageously comprised in the cathode.

According to the invention, an electrical current applied across the electrolytic cell is provided, which is sufficient to induce anodic and cathodic reactions according to the above-mentioned anode (1) and cathode (2) reactions. As documented in Figures 5-8, the electrical current applied across the electrolytic cell (11) used in the experiments and sufficient to induce anodic and cathodic reactions according to the anode (1) and cathode reactions (2) should be at least 5 mA/cm², at least 10 mA/cm², at least 20 mA/cm², at least 25 mA/cm², at least 30 mA/cm², preferably at least 40 mA/cm², preferably at least 50 mA/cm², more preferably at least 80 mA/cm², and even more preferably at least 100 mA/cm².

Preferably, the operating water partial pressure is at least 2 bar, at least 4 bar, at least 6 bar, preferably at least 8 bar, at least 10 bar, at least 15 bar, at least 20 bar, and even more preferably at least 25 bar. For the acid compounds of formula (III), the operating water partial pressure optimally shall be at least 8 bar. In a particularly preferred embodiment, the operating water partial pressure is from 8 to 25 bar, preferably from 10 to 20 bar, and even more preferably from 20 to 25 bar.

The present inventors have observed that while it is possible to produce CH4 while maintaining the electrolytic
cell at a minimum operating temperature just sufficient to induce superprotonic conduction in the electrolyte of hydrogen ions produced at the anode, it is preferable to operate the electrolytic cell at a temperature at least 50°C above the same minimum operating temperature.

This effect is e.g. visible in Figure 7, wherein an electrolytic cell operating temperature of 250°C (squares) about 15°C above the narrow transition interval for CSH2PO4 from non-superprotonic to superprotonic conduction of between 230-235°C, exhibits considerable polarization of the electrodes with a strong dependency on the current, whereas raising the temperature to 300°C (circles) lowers the polarization considerably and nearly eliminates the dependency of the same polarization on the current.

Preferably, the operating temperature is at least 25°C, at least 50°C, at least 75°C, at least 100°C, at least 125°C, preferably at least 150°C and most preferably at least 175°C above the same minimum operating temperature, which is just sufficient to induce superprotonic conduction in the electrolyte of hydrogen ions produced at the anode. For the compounds of formula (III), in particular for KH2PO4 and CSH2PO4, the optimal temperature range is from at least 275°C, from at least 320°C, or from at least 340°C to 400°C corresponding to from about 50°C to 175°C above the transition temperature from non-superprotonic to superprotonic conduction for these compounds.

Contrary to this, the present inventors have not observed a negative effect of approaching the upper temperature or temperature range) at which a given acid compound exhibits superprotonic conductivity, as long as the upper temperature (or temperature range) is not exceeded. Rather
the rate of reaction for the formation of methane is increased.

Suitable flowrates for the gasses used in the reactions are from 1 ml/min/cm² to 25 ml/min/cm² based on the size of the electrode. Preferably, the flowrates for the gasses used shall be from 2 ml/min/cm² to 20 ml/min/cm², from 3 ml/min/cm² to 15 ml/min/cm², from 5 ml/min/cm² to 12.5 ml/min/cm², or, from 7.5 ml/min/cm² to 10 ml/min/cm².

In preferred embodiments of method according to the invention, the anode and/or the cathode reactant flow is/are supplied to the electrolytic cell at a pressure of at least 50% of the operating water partial pressure and at a temperature lower by 50°C than the aforementioned operating temperature with the proviso that pressure and temperature must be sufficient to allow water to be present as steam. The advantage of supplying, in particular the water, at a lower temperature and pressure to the electrolytic cell is that, with the increase in temperature inside the cell, superheating of the steam is assured and water condensation is prevented.

Carbon dioxide for use in the method is readily available from many sources; however, it is preferably that the source of carbon dioxide is a source of high concentration carbon dioxide, such as e.g. the exhaust from a fossil fuel fired power plant, a production site for cement, or a production site for biogas. In one embodiment, it is preferred that the source providing the flow of CO₂ as a cathode reactants flow shall be a flow of carbon dioxide obtained from biogas. In another embodiment, it is preferred that the source of carbon dioxide is a cement plant. In one embodiment it is preferred that the source
of electrical energy is a wind plant, in another embodiment it is preferred that the source of electrical energy is a hydro-electrical plant, electrical energy from solar cells, or combinations thereof.

In an embodiment, wherein the source of carbon dioxide comprises a sulfuric content, the method further comprises to cease providing a flow of anode and cathode reactants, while the electrolytic cell is subjected to heating, flushing with an inert gas stream, and a reverse current for removing sulfuric compounds deposited during methane production in the electrolytic cell.

According to the invention, an anode products flow is can be collected from the anode, and a cathode products flow can be collected from the cathode. Methane can then be isolated from the cathode products flow in a final process step, particularly isolated from a cathode products flow comprising methane, unreacted CO2 and small amounts of formed H2O and/or CO. The invention does not rely on a specific means of methane isolation, but generally considers the isolation of methane from a gas flow within the common general knowledge of the skilled person. The present inventors have found that when the process of the invention is conducted with sufficient skill following the directions of the present disclosure, only little CO is formed and the majority of CO2 is consumed, such that the isolation of methane mostly consists in condensation of formed water. In the inventors' experiments, a gas-chromatography-column was fully sufficient for completely separating methane from the further gaseous components of the product flow.
Referring now to Figure 3, wherein an illustrative embodiment of the electrolytic cell system (10) is disclosed:

An electrolytic cell system (10) comprising an electrolytic cell (11) enclosed in a pressure chamber (12); the electrolytic cell comprising an anode compartment (20) comprising an anode (21) and an insulating anode enclosure (24) wherein the anode is embedded, and a cathode compartment (30) comprising a cathode (31) and an insulating cathode enclosure (34) wherein the cathode is embedded, and an electrolyte (40) arranged in contact with and between the anode (21) and the cathode (31), and wherein the anode compartment (20) and the cathode compartment (30) are separated by the electrolyte (40) and an insulating gasket (41) enclosing the electrolyte; the electrolyte (40) comprising at least one superprotonic conducting acid compound of the general formula (I):

$$\text{MxHy}(\text{A04})z$$

wherein $x$ is 1, 2, or 3; $y$ is 1, 2, 3, 4 or 5; $z$ is 1 or 2; M is at least one of Li, Na, K, Rb, Cs, Tl, NH$_4$; and A is S, Se, P, As;

wherein the anode compartment (20) further comprises an anode flow path (22) for gaseous H$_2$O as superheated steam to the anode (21) arranged for contacting the superheated steam as an anodic reactant with the anode; and an anode products flow path (23) from the anode (21) comprising means (15) for maintaining a constant water partial pressure across the anode; wherein the cathode compartment (30) further comprises a cathode flow path (32) for gaseous CO$_2$ to the cathode (31) arranged for contacting the CO$_2$ as a cathodic reactant with the cathode; and a cathode products flow path (33) from the cathode (31), optionally
comprising means (19) for isolating methane from a gas flow; the electrolytic cell system (10) further comprising means for circulating an electric current across the electrolytic cell (11), and respective means (13,17,18) for passing a respective gas flow through a respective anode or cathode compartment (20, 30). As discussed, when present the means (19) for isolating methane is preferably connected to the means (13) for passing a gas through an anode compartment (20) via a flow path (19a) for recirculating condensed water produced via reaction (2). Also preferably, the means (19) for isolating methane is connected to the means (17) for passing CO2 through the cathode compartment (20) via a flow path (17a) for recirculating unreacted CO2 obtained from the means (19) for isolating methane.

In accordance with the invention, the electrolytic cell system (10) comprises an electrolytic cell (11) enclosed in a pressure chamber (12). In one embodiment, only the electrolytic cell (11) is enclosed in the pressure chamber, in other embodiments, the pressure chamber (12) may enclose further elements of the electrolytic cell system. In one embodiment, the pressure chamber (12) encloses the further components of the electrolytic cell system fully. This e.g. can be realized by placing the elements comprised in the electrolytic cell system (10) inside an autoclave, which allows external pressure control over the entire electrolytic cell system. In this manner, a transportable electrolytic cell system can be provided.

In accordance with the invention, the electrolytic cell (11) comprises an anode compartment (20) comprising an anode (21) and an insulating anode enclosure (24) wherein the anode is embedded; and a cathode compartment (30)
comprising a cathode (31) and an insulating cathode enclosure (34) wherein the cathode is embedded; and an electrolyte (40) arranged in contact with and between the anode (21) and the cathode (31); and wherein the anode compartment (20) and the cathode compartment (30) are separated by the electrolyte (40) and an insulating gasket (41) enclosing the electrolyte. Thereby an electrically insulated cell (11) is provided.

Preferably, the electrically insulated cell (11) can be assembled and disassembled to allow access to the components of the cell between productions according to the methods of the invention.

In an embodiment, the anode compartment (20) and the cathode compartment (30) can be manufactured from stainless steel, alloys of copper and aluminum, or brass. The compartments (20,30) preferably are manufactured from the same material for improved seal and temperature response, and can be provided, in embodiments, with through passages for insulated bolts or screw threads for securing a tight connection between the compartments during production according to the methods of the invention. E.g., improvements to the connection were achieved between the experiments detailed in Figures 5 and 6, by using the above embodiments, thus allowing for reducing the contribution of atmospheric air to the measured product flow.

In accordance with the invention, the electrolyte (40) comprises at least one superprotonic conducting acid compound of the general formula (I):
MxHy (A04)z, wherein x is 1, 2, or 3; y is 1, 2, 3, 4, or 5; z is 1 or 2; M is at least one of Li, Na, K, Rb, Cs, Tl, NH₄; and A is S, Se, P, As; or of formulas (IB), (II), (III) and/or (V), as detailed above with regards to the methods of the invention in general.

In order to provide a flow of gaseous H₂O as superheated steam to the anode (21), the anode compartment (20) further comprises an anode flow path (22) for gaseous H₂O as superheated steam to the anode (21) arranged for contacting the superheated steam as an anodic reactant with the anode. It is preferable to maximize the contact of the superheated steam with the electrolyte (40), which can e.g. be effectuated as detailed below, by providing the anode in the form of a powder, preferably embedded in a meandering flow path (26) for the steam, whereby steam contact to the anode (21) is also maximized and steam contact to the electrolyte (40) is minimized such that the protons from the water can be absorbed in the electrolyte and the electrons transferred to the anode and the oxygen liberated. Thereby the risk of irreversible conversion of the electrolyte to a non-superprotonic conducting salt is reduced.

Also in accordance with the electrolytic cell system (10) of the invention, the cathode compartment (30) further comprises a cathode flow path (32) for providing gaseous CO₂ or a CO₂ containing gas stream to the cathode (31) arranged for contacting the CO₂ as a cathodic reactant with the cathode; preferably by contacting the CO₂ with the electrolyte (40) to pick up the protons of reaction (2) and the cathode (31) to let the cathode products comprising
mostly methane and water free to enter into the cathode flow path (33). The cathode products flow path (33) from the cathode (31) optionally comprises means (19) for isolating methane from a gas flow. As mentioned, the invention does not rely on a specific means (19) of methane isolation, but generally considers the isolation of methane from a gas flow within the common general knowledge of the skilled person. In the inventors' experiments, a gas-chromatography-column was fully sufficient for completely separating methane from the further gaseous components of the product flow.

Accordingly, in preferred embodiments of the invention, the anode and/or the cathode is/are a powder. By providing the anode and/or the cathode as a powder, the contact surface between conversion gas and the electrodes is increased and conversion improved.

Further, and for facilitating the use of powder electrodes, the electrolytic cell (11) in some, preferred embodiments comprises at least one respective anodic and/or cathodic meandering flow path (26,36) arranged respectively in the anode (20) or the cathode (30) compartment in contact with a respective anode (21) or cathode (22) for increasing a residence time of a respective anodic or cathodic reactant at a respective anode or cathode.

It was found that the meandering flow paths (26,36) suitably could be manufactured in stainless steel, in particular in stainless steel coated with metal oxides wherein the metal of the coating metal oxides suitably are selected from periodic table groups 4, 5 or 6, preferably wherein the metal is selected from periodic table group 5,
and most preferably wherein the stainless steel is coated with a layer of tantalum and tantalum oxide.

A particularly important aspect of the electrolytic cell system (10) of the invention is, as previously described, that the anode products flow path (23) from the anode (21) shall comprise suitable means (15) for maintaining a constant water partial pressure across the anode. The present inventors have found the maintenance of a constant water partial pressure across the electrolytic cell (11) essential for the long-term stability of the electrolyte without which, the 1-step methanization reaction of the invention rapidly ceases to function due to irreversible decomposition of the electrolyte. In one embodiment of the electrolytic cell system (10) according to embodiments of the invention, the means (15) for maintaining a constant water partial pressure across the anode (21) is a backpressure valve (15), preferably a heated backpressure valve (15) to avoid water condensation in the valve.

Further, and for providing a functioning electrolytic cell system (10), the electrolytic cell system (10) comprises means for circulating an electric current across the electrolytic cell (11), and means (13,17,18) for passing a respective gas flow through a respective anode or cathode compartment (20,30), such means (13,17,18) typically being pumps or fans.

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention further comprises a water-to-steam evaporator (14) arranged on the anode flow path (22) for providing gaseous H2O as superheated steam to the anode.
In one embodiment, the electrolytic cell system (10) according to embodiments of the invention further comprises a water condenser (16) arranged on the anode products flow path (23) after the means (15) for maintaining a constant water partial pressure across the anode (21).

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the electrolyte further comprises a support material for the at least one superprotonic conducting acid compound.

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the support material comprises a pyrophosphate compound of the general formula (VI):

\[(VI) \, Mt(P_2O_7)\]

wherein Mt can be Si, Ge, or Sn; alone or in combination with a polybenzimidazole (PBI) fiber reinforcement.

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the support material is a either Sn(P2O7) or Sn(P2O7) with a polybenzimidazole (PBI) fiber reinforcement.

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the anode comprises at least one oxide of iridium, preferably IrO2 and/or Ir2C.

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the cathode comprises at least one of Ni, alloyed Ni, Co, Ru, Rh, or a combination thereof, preferably Ni.
In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the anode and/or the cathode is/are a powder.

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the at least one superprotonic conducting acid compound is selected from at least one of a superprotonic conducting acid compound according to at least one of the general formulas (I), (II), (III), (IV) or (V).

In one embodiment, the electrolytic cell system (10) according to embodiments of the invention, the electrolytic cell system (10) further comprises a controller for executing a method according to any of the herein detailed aspects and embodiments.

In a further aspect and embodiment of the invention there is disclosed the use of a superprotonic conducting acid compound of the general formula (I):

\[ M_x H_y (A_{04})_z \]

wherein
\[ x = 1, 2, \text{ or } 3; \]
\[ y = 1, 2, 3, 4, \text{ or } 5; \]
\[ z = 1 \text{ or } 2; \]
\[ M \text{ is at least one of } \text{Li, Na, K, Rb, Cs, Tl, NH}_4; \]
\[ \text{and } A \text{ is } S, \text{Se, P, As}; \]
as an electrolyte in a 1-step co-electrolytic conversion of H₂O and CO₂ to O₂ and CH₄.

Also disclosed herein are the uses according to the invention of compounds of formulas (I), (II), (III), (IV), and/or (V) as an electrolyte in a 1-step co-electrolytic conversion of H₂O and CO₂ to O₂ and CH₄.
EXAMPLES

Example 1:
X-ray diffractogram showing evidence of almost complete conversion of \( \text{CSH}_2\text{PO}_4 \) into \( \text{CsH}_5(\text{PO}_4)_2 \) in the electrolyte in the presence of \( \text{SnP}_2\text{O}_7 \), cf. Figure 4. Topmost spectrum (A): measured data. Lower three spectra (B) – (D) calculated diffractograms corresponding to respectively (B): \( \text{CsH}_5(\text{PO}_4)_2 \); (C): \( \text{CsH}_2\text{P}_0_4 \); and (D): \( \text{SnP}_2\text{O}_7 \). In particular the almost complete absence of the dominant 26-peaks for \( \text{CSH}_2\text{PO}_4 \); and \( \text{SnP}_2\text{O}_7 \), at respectively 22° and 19°, compared to the presence of all major and minor diffraction peaks for \( \text{CsH}_5(\text{PO}_4)_2 \) is noteworthy and strongly indicative of near complete conversion.

Example 2:
\( \text{KH}_2\text{P}_0_4-\text{SnP}_2\text{O}_7 \) (pellet) with \( \text{Ni} \)-pellet cathode, \( \text{IrO}_2 \)-pellet anode. No meandering flow pattern, 8 bar operating water partial pressure, \( 320^\circ \text{C} \), 40 mA/cm\(^2\), electrolytic cell electrode diameter 1.26 mm. Gas-chromatographic (GC) detection (c.f. Figure 5) confirmed by Raman spectroscopy. Figure showing enlarged CO-region compared to \( \text{CH}_4 \)-region.

Direct, 1-step electrolytic conversion of \( \text{CO}_2 \) to \( \text{CH}_4 \) is confirmed with hardly any conversion of \( \text{CO}_2 \) into \( \text{CO} \). Full consumption of \( \text{H}^+ \) as witnessed by absence of \( \frac{3}{4} \) within the detection sensitivity of the GC. Proportion of \( \text{O}_2 \) and \( \text{N}_2 \) matches atmospheric proportion consistent with these two molecules being inactive in the conversion process.

Example 3:
\( \text{CsH}_2\text{P}_0_4-\text{SnP}_2\text{O}_7 \) (pellet) with \( \text{Ni} \)-pellet cathode, \( \text{IrO}_2 \)-pellet anode. No meandering flow pattern, 8 bar operating water...
partial pressure, 320°C, 50, 80 and 100 mA/cm², electrolytic cell electrode diameter 1.26 mm. Gas-chromatography detection (cf. Figure 6) confirmed by Raman spectroscopy. Figure showing enlarged CO-region compared to CH₄-region.

Direct, 1-step electrolytic conversion of CO₂ to CH₄ is confirmed with hardly any conversion of CO₂ into CO. Observed increase of ClHU-production with increasing current. Also, increase in CO, N₂ and O₂ with increasing current. Full consumption of H⁺ as witnessed by absence of H₂ within the detection sensitivity of the GC. Proportion of O₂ and N₂ matches atmospheric proportion consistent with these two molecules being inactive in the conversion process. Reduction in levels of O₂ and N₂ compared to Example 1 due to improved seal.

Example 4:
Polarization curves for a Ni|CSH₂PO₄/PBI |IrO₂-electrolytic cell, (No meandering flow pattern, 8 bar operating water partial pressure, electrolytic cell electrode diameter 1.26 mm) in dependence of current at respectively 250°C (squares, 3 data points) and 300°C (circles, 10 data points), cf. Figure 7. Generally, the lowest possible polarization potentials are desirable, and hence operating at 300°C rather than 250°C is preferable for the exemplified system.

Example 5:
Comparison of KH₂PO₄-SnP₂0₇ and CsH₂PO₄-SnP₂0₇ electrolytes (pellets), electrolytic cell electrode diameter 1.26 mm, Ni-pellet cathode, IrO₂-pellet anode. No meandering flow pattern, 8 bar operating water partial pressure, 320°C, cf. Figure 8.
The experiments indicate that electrolytes comprising the higher atomic number metals in formula (I) are more efficient as electrolytes than the lower atomic number metals, wherefore in one embodiment, K, Cs, and Rb are preferred over Li, Na, and NH4.

**Example 6:**
In an experiment following a production run using a carbon dioxide source from biogas which comprised sulfides, it was tested if the electrodes and electrolyte (C\(\text{SH}_2\text{PO}_4\) and SnP\(\text{O}_2\)) could be restored to initial capacity without contamination with sulfuric compounds. 10 mA/cm\(^2\), anode/cathode-switch (counter current) for 15 min at 400°C, was found to restore the system for the tested electrolytic cell, electrode diameter 1.26 mm.

**Example 7:**
In an experiment (Figure 9) the hypothesis of a 1-step methanization mechanism was tested using voltammetry. Generally, the main features of an electrochemical reaction are: 1) It takes place at the interface between electronic (electrode) and ionic conductor (electrolyte), wherein this interface is called an electric double layer (EDL). 2) The oxidation and reduction parts of the electrochemical reaction are physically separated. 3) The electrons are the participants of electrochemical reaction and cross the EDL during the reaction (electron transfer). 4) The main characteristics of the electrochemical reaction are current (the reaction rate) and potential (defines the activation energy barrier of the reaction).

Commonly, electrochemical reactions have more complicated mechanisms than simple electron transfer. The mechanisms
of an electrochemical reaction can include chemical steps, adsorption steps etc. taking place at the same electrode.

In the present invention, it is speculated that the electrochemical reaction has a mechanism with an immediately following chemical reaction at the same electrode (EC mechanism). This hypothesis was tested by the present inventors using voltammetry.

In voltammetry, electrochemical reactions are studied at one electrode. During the measurement, the potential of the electrode is changed linearly with time, and the current response is measured. The results are presented in the form of current-potential plot. Each peak on this plot is conclusive of an electrochemical reaction.

The results obtained for negative polarization at Ni electrode in molten CsH₂PO₄ in the N₂, N₂-CO₂ (1:1); N₂-CO₂ (1:3) and CO₂ atmospheres at 350°C are given in Figure 9.

Weak electrochemical activity can be seen in pure nitrogen atmosphere. There is a reduction wave with maximum at -0.4V. The only electrochemically active compound in this system is water, which can be electrochemically reduced according to the reaction (2). It can be seen from Figure 1, that the result of the addition of CO₂ is a dramatic increase of the reduction current, i.e. acceleration of the electrochemical reaction. This fact can be explained by an EC mechanism of the electrochemical reaction. Moreover, in chemical reaction (3) water is produced; which is the reactant in electrochemical reaction (2). As this is a catalytic EC mechanism, this can be the reason of the electrochemical reaction acceleration in presence of CO₂. It can be also seen, that the initial reaction potential
moves in positive direction when CO2 is introduced into the system, which means that less energy is spent for polarization of the electrode. After the electrochemical experiment, the gaseous part of the electrochemical cell has been analyzed using Raman spectroscopy and methane was found to be present as one of the gaseous products.

In conclusion, the present investigation showed that methane is produced in a single electrochemical process step. Moreover, this process is sufficiently accelerated by CO2 because of the catalytic nature of the EC electrochemical reaction. Furthermore, the reaction potential becomes considerably more positive when CO2 is introduced into the system. These facts show the presence of a one-step methane formation process (the present invention); which is more energy efficient than the two-step processes hitherto known.

CLOSING COMMENTS

The term "comprising" as used in the claims does not exclude other elements or steps. The term "a" or "an" as used in the claims does not exclude a plurality. A single processor or other unit may fulfill the functions of several means recited in the claims. Although the present invention has been described in detail for purpose of illustration, it is understood that such detail is solely for that purpose, and variations can be made therein by those skilled in the art without departing from the scope of the invention.
1. A method of synthesizing methane by 1-step co-electrolysis of H2O and CO2 in the presence of an electrolyte comprising at least one superprotonic conducting acid compound of the general formula (1)

\[ M_x H_y (A_0)^z \], wherein

- x is 1, 2, or 3;
- y is 1, 2, 3, 4 or 5;
- z is 1 or 2;
- M is at least one of Li, Na, K, Rb, Cs, Tl, NH4;
- and A is S, Se, P, As;

said method comprising:

a) providing a pressurized electrolytic cell comprising an anode and a cathode, wherein said electrolyte is arranged between said anode and said cathode; and wherein hydrogen ions are produced at said anode according to an anode reaction (1)

\[ 4H_2O \rightarrow 2O_2 + 8H^+ + 8e^- \]; and

wherein methane is produced at said cathode according to a cathode reaction (2)

\[ CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O \];

said method further comprising operating said electrolytic cell by:

b) providing an electrical current applied across said electrolytic cell sufficient to induce anodic and cathodic reactions according to said anode (1) and cathode reactions (2);

c) maintaining said electrolytic cell at an operating water partial pressure from 1 to 30 bar;

d) maintaining said electrolytic cell at an operating temperature sufficient to induce superprotonic conduction in said electrolyte of hydrogen ions produced at said anode;
e) providing a flow of gaseous H2O as superheated steam as an anode reactants flow to said anode at a flow rate sufficient to allow said anodic reaction;

f) providing a flow of gaseous CO2 as a cathode reactants flow to said cathode at a flow rate sufficient to allow said cathodic reaction;

g) collecting a cathode products flow from said cathodic reaction; and

h) optionally, isolating methane from said cathode products flow.

2. A method according to claim 1, further comprising:

i) collecting an anode products flow from said anodic reaction.

3. A method according to either claim 1 or 2, wherein said at least one superprotonic conducting acid compound has the general formula (IB):

(IB) \[ M_{x}H_{y}A_{0}^{4} \]; wherein;

- \( M \) is Li, Na, K, Rb, Cs, NH4;
- \( A \) is S, Se, P, As;
- \( x \) is 1 or 2;
- \( y \) is 1 or 2; and
- \( x+y \) is 2 when \( A \) is S or Se, and 3 when \( A \) is P or As.

4. A method according to any of the claims 1 to 3, wherein said at least one superprotonic conducting acid compound has the general formula (II):

(II) \[ MH_{2}A_{0}^{4} \]; wherein

- \( M \) is Li, Na, K, Rb, Cs, NH4; and
- \( A \) is S, Se, P, As.
5. A method according to any of the preceding claims, wherein said superprotonic conducting acid compound has the general formula (III):

(III) \( \text{MH}_2\text{PO}_4 \); wherein

\( \text{M} \) is Li, Na, K, Rb, Cs, NH\(_4\).

6. A method according to any of the preceding claims, wherein said at least one superprotonic conducting acid compound is either KH\(_2\)PO\(_4\) or CS\(_4\)PO\(_4\).

7. A method according to either claim 1 or 2, wherein said at least one superprotonic conducting acid compound has the general formula (IV):

(IV) \( \text{MH}_3(\text{A}_4\text{O}_4)_2 \); wherein

\( \text{M} \) is Li, Na, K, Rb, Cs, NH\(_4\); and

\( \text{A} \) is S, Se, P, As.

8. A method according to claim 7, wherein said superprotonic conducting acid compound has the general formula (V):

(V) \( \text{MH}_5(\text{P}_0\text{O}_4)_2 \); wherein

\( \text{M} \) is Li, Na, K, Rb, Cs, NH\(_4\).

9. A method according to either claim 7 or 8, wherein said at least one superprotonic conducting acid compound is either KH\(_5\)(P\(_0\)O\(_4\))\(_2\) or CsH\(_5\)(P\(_0\)O\(_4\))\(_2\).

10. A method according to any of the preceding claims, wherein said electrolyte comprises a mixture of at least two of said superprotonic conducting acid compounds.
11. A method according to any of the preceding claims, wherein said at least one superprotonic conducting acid compound is either solid, molten or a mixture of solid and molten.

12. A method according to any of the preceding claims, wherein said electrolyte further comprises a support material for said at least one superprotonic conducting acid compound.

13. A method according to claim 12, wherein said support material comprises a pyrophosphate compound of the general formula (VI):

\[(VI) \quad \text{Mt}(P\text{2}0\text{7})\],

wherein Mt can be Si, Ge, or Sn; alone or in combination with a polybenzimidazole (PBI) fiber reinforcement.

14. A method according to either claim 12 or 13, wherein said support material is a either Sn(P2O7) or Sn(P2O7) with a polybenzimidazole (PBI) fiber reinforcement.

15. A method according to any of the preceding claims, wherein said anode comprises at least one oxide of iridium, preferably IrO2 and/or Ir2O3.

16. A method according to any of the preceding claims, wherein said cathode comprises at least one of Ni, alloyed Ni, Co, Ru, Rh, or a combination thereof, preferably Ni.

17. A method according to either of claims 15 or 16, wherein said anode and/or said cathode is/are a powder.
18. A method according to any of the preceding claims, wherein said operating water partial pressure is at least 8 bar.

19. A method according to any of the preceding claims, wherein said operating temperature is from 275°C to 400°C.

20. A method according to any of the preceding claims, wherein said electrical current applied across said electrolytic cell sufficient to induce anodic and cathodic reactions according to said anode (1) and cathode reactions (2) is at least 5 mA/cm², preferably 40 mA/cm².

21. A method according to any of the preceding claims, wherein said anode and/or said cathode reactants flow is/are supplied to said electrolytic cell at a pressure of at least 50% of said operating water partial pressure and at a temperature lower by 50°C than said operating temperature with the proviso that pressure and temperature must be sufficient to allow water to be present as steam.

22. A method according to any of the preceding claims, wherein said method is executed on an electrolytic cell system (10) according to any of the claims 23 to 36.

23. An electrolytic cell system (10) comprising an electrolytic cell (11) enclosed in a pressure chamber (12); said electrolytic cell comprising an anode compartment (20) comprising an anode (21) and an insulating anode enclosure (24) wherein said anode is embedded, and a cathode compartment (30) comprising a
cathode (31) and an insulating cathode enclosure (34) wherein said cathode is embedded, and an electrolyte (40) arranged in contact with and between said anode (21) and said cathode (31), and wherein said anode compartment (20) and said cathode compartment (30) are separated by said electrolyte (40) and an insulating gasket (41) enclosing said electrolyte; said electrolyte (40) comprising at least one superprotonic conducting acid compound of the general formula (I):

$$M_xH_y(AO_4)z$$

wherein

- $x$ is 1, 2, or 3;
- $y$ is 1, 2, 3, 4 or 5;
- $z$ is 1 or 2;
- $M$ is at least one of Li, Na, K, Rb, Cs, Tl, NH$_4$; and
- $A$ is S, Se, P, As;

wherein said anode compartment (20) further comprises an anode flow path (22) for gaseous H$_2$O as superheated steam to said anode (21) arranged for contacting said superheated steam as an anodic reactant with said anode; and an anode products flow path (23) from said anode (21) comprising means (15) for maintaining a constant water partial pressure across said anode;

wherein said cathode compartment (30) further comprises a cathode flow path (32) for gaseous CO$_2$ to said cathode (31) arranged for contacting said CO$_2$ as a cathodic reactant with said cathode; and a cathode products flow path (33) from said cathode (31), optionally comprising means (19) for isolating methane from a gas flow;

said electrolytic cell system (10) further comprising means for circulating an electric current across said electrolytic cell (11), and respective means (13,17,18) for passing a respective gas flow through a respective anode or cathode compartment (20,30).
24. An electrolytic cell system (10) according to claim 23, the system further comprising at least one respective anodic and/or cathodic meandering flow path (26,36) arranged in contact with a respective anode (21) or cathode (22) for increasing a residence time of a respective anodic or cathodic reactant at a said respective anode or cathode.

25. An electrolytic cell system (10) according to claim 23 or 24, the system further comprising a water-to-steam evaporator (14) arranged on said anode flow path (22) for providing gaseous H2O as superheated steam to said anode.

26. An electrolytic cell system (10) according to any of the claims 23 to 25, the system further comprising a water condenser (16) arranged on said anode products flow path (23) after said means (15) for maintaining a constant water partial pressure across said anode (21).

27. An electrolytic cell system (10) according to any of the claims 23 to 26, wherein said means (15) for maintaining a constant water partial pressure across said anode (21) is a heated backpressure valve.

28. An electrolytic cell system (10) according to any of the claims 23 to 27, wherein said pressure chamber (12) encloses said electrolytic cell system.

29. An electrolytic cell system (10) according to any of the claims 23 to 28, wherein said electrolyte further comprises a support material for said at least one superprotonic conducting acid compound.
30. An electrolytic cell system (10) according to claim 29, wherein said support material comprises a pyrophosphate compound of the general formula (VI):

\[(VI) \text{Mt}(P_{2}O_{7})\]

wherein Mt can be Si, Ge, or Sn; alone or in combination with a polybenzimidazole (PBI) fiber reinforcement.

31. An electrolytic cell system (10) according to either claim 29 or 30, wherein said support material is a either Sn(P2O7) or Sn(P207) with a polybenzimidazole (PBI) fiber reinforcement.

32. An electrolytic cell system (10) according to any of the claims 23 to 31, wherein said anode comprises at least one oxide of iridium, preferably IrO2 and/or Ir2O3.

33. An electrolytic cell system (10) according to any of the claims 23 to 32, wherein said cathode comprises at least one of Ni, alloyed Ni, Co, Ru, Rh, or a combination thereof, preferably Ni.

34. An electrolytic cell system (10) according to any of the claims 23 to 33, wherein said anode and/or said cathode is/are a powder.

35. An electrolytic cell system (10) according to any of the claims 23 to 34, wherein said at least one superprotonic conducting acid compound is selected from at least one of a superprotonic conducting acid compound according to at least one of the general formulas (I), (II), (III), (IV) or (V).
36. An electrolytic cell system (10) according to any of the claims 23 to 35, further comprising a controller for executing a method according to any of the claims 1 to 22.

37. Use of a superprotonic conducting acid compound of the general formula (I):

\[ M_xH_y(AO_4)^z, \]

wherein

- \( x \) is 1, 2, or 3;
- \( y \) is 1, 2, 3, 4, or 5;
- \( z \) is 1 or 2;
- \( M \) is at least one of Li, Na, K, Rb, Cs, Tl, NH_4;
- and \( A \) is S, Se, P, As;

as an electrolyte in a 1-step co-electrolytic conversion of H_2O and CO_2 to O_2 and CH_4.

38. Use of a superprotonic conducting acid compound according to claim 37, wherein said superprotonic conducting acid compound is selected from a superprotonic conducting acid compound of the general formula (II), (III), (IV), or (V).
Arrhenius plot of transition temperatures ranges for 3 salts from proton conducting to superproton conducting state

Figure 1
<table>
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<tr>
<th>Compound</th>
<th>Superionic T (°C)</th>
<th>Conductivity (S cm⁻¹)</th>
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<tr>
<td>CsHSO₄</td>
<td>141</td>
<td>8 × 10⁻³</td>
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<td>RbHSO₄</td>
<td>160–170</td>
<td>~ 10⁻⁴</td>
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<td>KHSO₄</td>
<td>150–180</td>
<td>~ 10⁻⁴</td>
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<td>CsHSeO₄</td>
<td>128</td>
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<td>RbHSeO₄</td>
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<tr>
<td>NH₄HSeO₄</td>
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<td>Cs₃H(SeO₄)₂</td>
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<td>Rb₃H(SeO₄)₂</td>
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<td>K₃H(SeO₄)₂</td>
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<td>10⁻³</td>
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<td>CsH₂PO₄</td>
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<td>RbH₂PO₄</td>
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<td>KH₂PO₄</td>
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<td>LiH₂PO₄</td>
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<td>KH₂AsO₄</td>
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<td>NH₄H₂AsO₄</td>
<td>125</td>
<td>2.5 × 10⁻³</td>
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Table 1: Apparent transition temperatures at atmospheric pressure for transition to superprotonic conducting state of various salts

Figure 2
Figure 3
Figure 5
Figure 7
Figure 9
**INTERNATIONAL SEARCH REPORT**

**PCT/DK2018/050221**

**A. CLASSIFICATION OF SUBJECT MATTER**


**ADDITIONAL INFORMATION**

According to International Patent Classification (IPC) or the national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

C25B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic database consulted during the international search (name of database and, where practicable, search terms used)

EPO-Internal, WPI Data

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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**Date of the actual completion of the international search**

23 November 2018

**Date of mailing of the international search report**

04/12/2018

**Name and mailing address of the ISA/Authorized officer**

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Ritter, Thomas
<table>
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<td>US 4897167 A</td>
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