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# POTENTIOMETRIC TITRATION METHOD FOR QUANTITATIVE DETERMINATION OF HYDROGEN PEROXIDE

# TO ALL WHOM IT MAY CONCERN

BE IT KNOWN THAT (1) CHARLES J. PATRISSI, employee of the United States Government, and (2) RUSSELL R. BESSETTE, citizens of the United States of America, and residents respectively of (1) Newport, County of Newport, State of Rhode Island, and (2) Mattapoisett, County of Plymouth, Commonwealth of Massachusetts, have invented certain new and useful improvements entitled as set forth above of which the following is a specification:

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1	Attorney Docket No. 95764
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3	POTENTIOMETRIC TITRATION METHOD FOR QUANTITATIVE
4	DETERMINATION OF HYDROGEN PEROXIDE
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6	STATEMENT OF GOVERNMENT INTEREST
7	The invention described herein may be manufactured and used
8	by or for the Government of the United States of America for
9	governmental purposes without the payment of any royalties
10	thereon or therefore.
11	
12	CROSS REFERENCE TO OTHER RELATED APPLICATIONS
13	Not applicable.
14	
15	BACKGROUND OF THE INVENTION
16	(1) Field of the Invention
17	The present invention relates to titration methods, and
18	more specifically to a potentiometric titration method for a
19	quantitative determination of hydrogen peroxide.
20	(2) Description of the Prior Art
2.1	There continues to be a need for energy sources with a high
22	energy density. In particular, there is a need for high energy
23	density energy sources that can power unmanned undersea
24	vehicles. Such energy sources when used to power such vehicles

are required to have high energy density for long duration and
 quiet operation. Additionally, they must be relatively
 inexpensive, environmentally friendly, safe to operate,
 reusable, capable of a long shelf life and not prone to
 spontaneous chemical or electrochemical discharge.

6 The zinc silver oxide (Zn/AgO) electrochemical couple has 7 served as a benchmark energy source for undersea applications. 8 Because of its low energy density, however, it is not suitable 9 for unmanned undersea vehicles whose energy density requirements 10 are seven times those of the Zn/AgO electrochemical couple.

11 In an effort to fabricate power sources for unmanned undersea vehicle with increased energy density (over zinc-based 12 power sources), research has been directed towards semi fuel 13 cells (as one of several high energy density power sources being 14 considered). Semi fuel cells normally consist of a metal anode, 15 such as magnesium (Mg) and a catholyte such as hydrogen peroxide 16  $(H_2O_2)$ . In general the performance and health of these types of 17 semi fuel cells are a function of the quantity of hydrogen 18 19 peroxide in the catholyte. The key to achieving a high energy density for these types of semi fuel cells lies in the efficient 20 usage of the hydrogen peroxide. The electrochemical processes 21 22 during cell discharge are:

Anode: Mg  $Mq^{2+} + 2e^{-}$ 

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Cathode:  $H_2O_2 + 2H^+ + 2e^- ->$ 1 2H2O The voltage at the cathode and the total semi fuel cell voltage 2 are directly related to the concentration of hydrogen peroxide 3 in the catholyte according to the Nernst equation: 4  $E = E^{0} + (0.0591 \times \log ([H_2O_2] \times [H^{+}]^{2}))/2$ 5 where E is the half cell voltage at the cathode,  $E^0$  is the 6 standard voltage at unit activity of  $H_2O_2$  and  $H^+$ , and  $[H_2O_2]$  and 7 [H<sup>+</sup>] are the molar concentrations of peroxide and protons 8 respectively. The Nernst equation shows that as the peroxide 9 concentration decreases so does the cell voltage. 10 It is important to directly monitor and control the 11 hydrogen peroxide concentration  $[H_2O_2]$ , because the concentration 12 is used to assess the functional condition and performance of 13 the semi-fuel cell. For example, if the hydrogen peroxide 14 concentration differs significantly from expected levels for a 15 given semi fuel cell load, then the pump controlling the 16 hydrogen peroxide input can be directed to increase or decrease 17 the amount of hydrogen peroxide being pumped into the semi fuel 18 cell. 19 In a laboratory environment, measurement of hydrogen 20

21 peroxide concentration in a semi fuel cell is performed using a 22 colorimetric titration method. In this method, a solution of 23 unknown peroxide concentration is colored with a small amount of 24 indicator material such as iron(II) 1,10 phenanthroline. Then,

1 a chemical of known concentration, typically cerium (IV) in sulfuric acid solution, (the titrant solution) is added that 2 reacts with peroxide. When the solution turns clear, all of the 3 hydrogen peroxide has been consumed. There is a 2:1 correlation 4 between the number of titrant reactant molecules consumed during 5 the titration and the number of hydrogen peroxide molecules 6 initially present in the solution when cerium (IV) is used. 7 The 8 concentration of hydrogen peroxide can be determined using this correlation. This method is not suitable for use in an unmanned 9 undersea vehicle, however, because it requires visible detection 10 11 of a color change by a human operator. Currently there is no 12 automated means for quantifying the concentration of hydrogen peroxide in a semi fuel cell onboard an unmanned undersea 13 vehicle. 14

What is needed is a method of quantifying the concentration of hydrogen peroxide in a semi fuel cell catholyte that is automated and can provide concentration data that can be interpreted by a digital processor.

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### SUMMARY OF THE INVENTION

It is a general purpose and object of the present invention to establish a method of quantifying the concentration of hydrogen peroxide in a semi fuel cell catholyte that is

automated and can provide concentration data that can be
 interpreted by a computer.

This object is accomplished by employing an electrochemical 3 potentiometric titration method. The method entails titration 4 of a known volume of a catholyte containing an unknown amount of 5 hydrogen peroxide in a titration cell having two electrodes, a 6 7 platinum working electrode and a silver/silver chloride reference electrode. A known concentration of a titrant is 8 added to the known volume of catholyte in the titration cell. 9 Simultaneously, as the titrant is added the potential between 10 the working electrode and the reference electrode is monitored. 11 The point at which all of the hydrogen peroxide has been 12 consumed is signaled when the cell potential changes abruptly. 13 14 Since the concentration of the titrant is already known, the amount of titrant added (concentration multiplied by volume) is 15 directly related to the amount of hydrogen peroxide consumed. 16 The concentration of hydrogen peroxide is calculated from the 17 volume of catholyte and the moles of hydrogen peroxide. 18

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#### BRIEF DESCRIPTION OF THE DRAWINGS

A more complete understanding of the invention and many of the attendant advantages thereto will be readily appreciated as the same becomes better understood by reference to the following

1 detailed description when considered in conjunction with the 2 accompanying drawings wherein:

FIG. 1 is a diagram illustrating the apparatus of thepresent invention;

5 FIG. 2 is a diagram illustrating flow injection analysis 6 system of the present invention;

FIG. 3 is a graph of dE/dV versus titrant as recorded by
the injection analysis system of the present invention.

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## DESCRIPTION OF THE PREFERRED EMBODIMENT

11 Referring now to FIG. 1 there is illustrated a diagram of the invention where an electrochemical titration cell 10 12 contains a certain volume of catholyte 12. Inside the titration 13 cell 10 are two electrodes, working electrode 14 and reference 14 15 electrode 16. Also contained inside the titration cell is a mechanical stir bar 18. In the preferred embodiment, the 16 17 titration cell will be relatively small to conserve volume when used onboard an unmanned undersea vehicle. In the preferred 18 embodiment, the working electrode 14 is made of platinum, and 19 the reference electrode 16 is made of silver/silver chloride. A 20 21 titrant solution 20 is introduced into the titration cell 10. 22 In the preferred embodiment, the titrant solution 20 is a solution of Ce4+. The potential between the working electrode 23 and the reference electrode is measured once the titrant 24

solution 20 is introduced. In the preferred embodiment, the
 potential is measured by means of a potentiostat/galvanostat 22.
 The chemical reactions occuring in the titration cell are
 shown in Equations 1-3:

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 $H_2O_2 \quad --> \qquad 2e^{-} + 2H^{+} + O_2 \qquad (1)$   $Ce^{4+} + e^{-} --> \qquad Ce^{3+}. \qquad (2)$ 

 $H_2O_2 + Ce^{4+} --> 2H^+ + O_2 + Ce^{3+}$  (3)

9 The addition of  $Ce^{4+}$  into the catholyte oxidizes the hydrogen 10 peroxide. During the addition of  $Ce^{4+}$ , the cell potential will be 11 controlled by the  $H_2O_2/O_2$  redox couple. Immediately following 12 consumption of all the peroxide, the cell potential will shift to 13 that of the  $Ce^{4+}/Ce^{3+}$  redox couple. This abrupt change in the cell 14 potential signals the end point of the titration and can be used by a 15 computer to calculate the molarity of the hydrogen peroxide.

16 In a preferred implementation of the invention, a flow injection analysis system 24 is used for on-line analysis as illustrated in 17 FIG. 2. A micro-pump 26 will be connected to the catholyte chamber 18 28 of a semi fuel cell. The micro-pump 26 will remove a small fixed 19 volume of catholyte and fill a fixed volume sample loop 30. The loop 20 30 empties into a dilution chamber 32 containing a known volume of 21 electrolyte that does not contain  $H_2O_2$  to dilute the small fixed 22 volume of catholyte. A micro-pump 27 will then fill a second fixed 23 volume sample loop 34 with the diluted sample of the catholyte. This 24

1 diluted sample will be emptied into a titration cell 10 containing the reference electrode 16 and the working electrode 14. 2 The electrodes are connected to a combined programmable digital 3 processing unit 36a and high impedance voltmeter 36b. The digital 4 processing unit 36a also controls a micro-burette 38 that introduces 5 the titrant into the titration cell 10 at a fixed rate as a stirring 6 device 18 mixes the titrant and diluted sample. The digital 7 processing unit 36a receives the readings from the voltmeter 36b and 8 performs the calculation dE/dV, where dE is the change in cell 9 potential and dV is the change in volume of titrant from the previous 10 data point. The endpoint of the titration is signaled when the slope 11 of this graph changes from positive to negative as illustrated in FIG 12 3. At this point the digital processing unit 36a is programmed to 13 stop the micro-burette 38 from introducing any more titrant into the 14 15 titration cell 10. Based on the volume of titrant that was delivered up to the endpoint, and because all of the volumes are fixed, the 16 digital processing unit 36a is programmed to calculate the 17 concentration of hydrogen peroxide in the original catholyte sample. 18

In a laboratory experiment, 20 micro liters of catholyte was diluted to 20 milliliters in a 50-milliliter dilution chamber containing 40g/L of sodium chloride. The H<sub>2</sub>O<sub>2</sub> concentration in the catholyte was determined to be 0.105 moles per liter using the colorimetric cerium (IV) titration method. The diluted catholyte was then placed in a titration cell and the cell potential was measured.

A Ce<sup>4+</sup> titrant solution that was 0.001366 M was then titrated into a titration cell and the cell potential was measured 45 seconds after each addition of titrant. A graph of dE/dV versus titrant added is illustrated in FIG. 3. The graph shows the sharp change in dE/dV at 3.00 milliliters, which is the end point of the titration. The calculation of the hydrogen peroxide molarity is as follows:

7 (0.00300 L of  $Ce^{4+}$ ) \* (0.001366 moles of  $Ce^{4+}/one$ 8 liter of  $Ce^{4+}$  solution)\*(1 mole  $H_2O_2/2$  moles 9  $Ce^{4+}/20x10^{-6}$  L of catholyte = 0.102 moles of  $H_2O_2$ 10 per liter of solution.

11 The error of the measurement is acceptable at 2.9%.

The advantages of the present invention over the prior art are autonomous / automated control of hydrogen peroxide,  $H_2O_2$ , concentration to assess the functional condition (health) and performance of a hydrogen peroxide,  $H_2O_2$ , based fuel cell.

Obviously many modifications and variations of the present 16 invention may become apparent in light of the above teachings. 17 For example they include various reference electrodes such as 18 the saturate calomel electrode, various non corroding electrode 19 materials such as gold or palladium, various dilution ratios 20 depending on titration cell volume, expected peroxide 21 concentration, etc, various types of electronic instrumentation 22 to perform the measurement and acquire and process the data, and 23

different analysis methods to determine the endpoint such as second derivative plot.

In light of the above, it is therefore understood that within the scope of the appended claims, the invention may be practiced otherwise than as specifically described. 1 Attorney Docket No. 95764

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# POTENTIOMETRIC TITRATION METHOD FOR QUANTITATIVE DETERMINATION OF HYDROGEN PEROXIDE

### ABSTRACT OF THE DISCLOSURE

An electrochemical potentiometric titration method that 7 entails titration of a known volume of a catholyte containing an 8 unknown amount of hydrogen peroxide in a titration cell having 9 two electrodes, a platinum working electrode and a silver/silver 10 chloride reference electrode. A known concentration of a 11 titrant is added to the catholyte in the titration cell. 12 Simultaneously, as the titrant is added the potential between 13 the working electrode and the reference electrode is monitored. 14 15 The point at which all of the hydrogen peroxide has been consumed is signaled when the cell potential changes abruptly. 16 Since the concentration of the titrant is already known, the 17 amount of titrant added (concentration multiplied by volume) is 18 directly related to the amount of hydrogen peroxide consumed. 19 The concentration of hydrogen peroxide is calculated from the 20 volume of catholyte and the moles of hydrogen peroxide. 21







