Quantification of Full Range Ethanol Concentrations by Using pH Sensor

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This work was sported by Institute of Bioprocess Engineering at Erlangen University where the work was done

Abstract
A differential pH measurement device was used to achieve operation conditions of alcohol dehydrogenase reaction. Optimum operating conditions were temperature of 30°C, 10 µl of alcohol dehydrogenase enzyme volume (with a final activity of 563.75 units ml⁻¹) per 50 µl of sample, NAD⁺ concentration of 0.05 mM and 20 mM glycine-pyrophosphate buffer solution of pH 9.1. In this method a range of ethanol concentrations from 0 - 99.985 %, which means 0.000001714 - 17.14 M, were used. The maximum obtained change in pH, delta pH, was (-33) mR. A calibration curve of logarithmic values of ethanol concentrations against change in pH for standard ethanol samples was done. Since this calibration curve is a linear with a correlation coefficient (R) of 0.998, this calibration curve can be used in quantification of ethanol concentration. End point of equilibrium concentrations of reactants and products of ethanol oxidation reaction was measured within spectrophotometer. The results indicated 100 seconds of process time is required to reach the end point for all ethanol standard samples. This required time was satisfied with results of measuring change in pH within differential pH analyzer system.

Keywords: Ethanol analysis, Oxidation enzymatic reaction, Ethanol quantification, Differential pH measurement device

1. Introduction
Ethanol is one of the most important substances with different application. For instance, bioethanol is rapidly rising its market share because of application of new technologies i.e. flexible fuel technology. However, for any application of ethanol it is necessary to detect and quantify ethanol with high accuracy. Therefore many methods were used to determine the concentration of ethanol in case when it is either a main product or a byproduct. The available techniques of ethanol detection include gas chromatography, electrochemical and enzymatic assay. The most important methods can be found in previous literatures (Goodman and Jacksonville, 1975; Bauer and Magers, 1985; Watanabe et al 1985; Laccheri, 1987; Moldowan, 1987; Stefan and Luc, 1997; Kempa, 2004; Hernandez, 2005; Olt, 2007; Beutler, 1984; Bernt and Gutman, 1974; Majki and Berkec, 1980).

Many difficulties and disadvantages are accompanied these methods. Some of them, especially the non-enzymatic methods, are complex and time consuming. They require previous separation process (distillation, pervaporation), expensive instrumentation and trained operators. Such disadvantages can be overcome by the use of enzymatic methods. In such enzymatic methods, most enzyme-catalyzed reactions can be followed by simple, widely available spectroscopic or electrochemical methods (Azevedo et al 2005).

Optical methods require an apparatus, such as a spectrophotometer. Moreover, such methods cannot be applied to turbid samples, such as blood or food samples. Furthermore, for carrying out the enzymatic reaction, a troublesome procedure involving dissolution of various reagents and distribution of the resulting solutions into reaction vessels in amounts specified beforehand is required, and the reagent solutions, such as prepared NAD⁺ solution have poor storage stability. For these and other reasons, those methods are not suitable for general use. In order to overcome
these difficulties, the combined use of an enzymatic method and an electrochemical method for determining alcohols have been desired (Watanabe et al 1985). The optical density in such assay must be read at 340 nm, precluding the use of widely available and relatively inexpensive photoelectric colorimeters that do not allow precise determination in the UV range (Rodionov et al 2002). Additionally, in order to increase accuracy, there is a need to dilute the samples, due to the limitation of Lambert-Ber-Law’s linearity.

Other spectroscopic methods (colorimetric, chemiluminescent and fluorescent methods) can be used to detect the production of H₂O₂ during the oxidation of ethanol. Most spectrometric methods are based on a bienzymatic system, comprising AOX and a peroxidase enzyme. The need for fast, cheap, sensitive and continuous analyzing methods with a high sample throughput led to application of immobilized enzyme reactors into flow system based analysis (Rodionov et al 2002).

Use of enzyme electrode or biosensor to detect changes in either oxygen concentration or H₂O₂ concentration in reactions, catalyzed by immobilized alcohol oxidase, suffers from some difficulties. Measurements based on oxygen have practical inconveniences and limitations. The response is low, and the dependency on oxygen can reduce the accuracy and reproducibility of the device. Moreover, because of a high background signal, the minimum detectable concentration is not very low. The detection of H₂O₂ is the most commonly used alternative to overcome these drawbacks. These techniques, however, usually suffer from low sensitivity (Rodionov et al 2002; Majki and Berkec, 1980). It has been reported that when ethanol was oxidized just in buffer by AOX, immobilized in 31 µl bioreactors, the conversion started to decrease after 3 h of continuous operation and after 4 h more than 80 % of the initial conversion were lost. Also one of the main factors that affect the performance of a biosensor is the enzyme immobilization procedure itself (Azvedo et al 2005).

There seems to be no fully satisfactory method for wide range ethanol concentrations quantification available yet. The above mentioned methods are not only limited for a certain range of ethanol concentration, but also they have difficulties or problems as mentioned above. Furthermore, some of them are expensive.

Therefore, the aim of this study was to develop a simple enzymatic method for detection and quantification of a full range of ethanol concentrations without diluting of samples. During this method a reduction in the cost of the ethanol assay will be done by using a small quantity of one enzyme and very low NAD⁺ concentration. The measurements will be achieved by using a simple device, a differential pH analyzer system CL10.

The basic principle of this method is to assay ethanol sample by measuring pH variation produced in the medium during the enzymatic oxidation of ethanol to acetaldehyde. This reaction is achieved in presence of alcohol dehydrogenase enzyme. The following reaction represents reaction mechanism:

\[
\text{CH}_3\text{CH}_2\text{OH} + \text{NAD}^+ \leftrightarrow \text{CH}_3\text{CHO} + \text{NADH} + \text{H}^+ \quad (1)
\]

2. Experimental Procedure

2.1 Instruments

2.1.1 Differential pH analyzer system CL10 (Eurochem, Italy).

This device mainly consists of one mixing chamber, five peristaltic pumps and two capillary glass electrodes of pH sensor. This apparatus is controlled by using the CL10 Manger program. Figure 1 shows the schematic diagram of this apparatus. Details on the principle of its work, measurement and control can be found in different publications (Luzzana et al 1983; Rovida et al 1984; Luzzana et al 2001).

2.1.2 Spectrophotometer

In order observe the enzymatic reaction of ethanol oxidation to acetaldehyde; spectrophotometer type (Specord 205 from Analytic Jena AG, Germany) was used. A rise in absorbance, which is stoichiometric with amount of formed NADH, can be measured within spectrophotometer.

2.1.3 Curtipot program

A program was designed in order to execute pH and acid-base titration curves: Analysis and Simulation (Gutz, 2008).

2.2 Materials

β-nicotinamide-adeninedinucleotide (NAD⁺), Alcohol dehydrogenase enzyme ADH (EC 1.1.1.1), NADH were purchased from Sigma-Aldrich Company. High purity grade ethanol (99.985%) was used in preparing standard ethanol samples. These samples of different concentrations (0.0001714 - 17.14 M) were obtained by blending ethanol with bidest water of 0.005 microsiemens (µS). Buffer solution with pH of 9.1 at 25° C was composed from 0.02 M of glycine and 0.02 M of sodium pyrophosphate. Solution of alcohol dehydrogenase enzyme with a final
activity of (563.75 units/ml) was prepared by dissolving enzyme in a mixture of 25 % of glycerol/water. Magic N50 solution was prepared by blending 50 µl of 50 mM NAD⁺ with 100 ml of the used buffer and 850 ml of pure water. This NAD⁺ solution was called by authors Magic N50 solution because the effective of its low concentration in analyzing full ranges of ethanol concentrations. KIT assay (K-ETOH, Megazyme International Ireland Ltd, 2006) was used in order to make economic comparison.

2.3 Procedure

The differential pH device was set to a temperature of 30°C, a waiting time of 4 seconds and maximum reaction time (cycle time) of 300 seconds. First runs were done with buffer only as system check-up. For ethanol quantification, 50 µl of sample were well mixed with 10 µl of alcohol dehydrogenase enzyme to get 60 µl of enzyme-ethanol complex in glass tube. 250 µl of Magic N50 solution was injected into the mixing chamber which contained 1090 µl of buffer. After 40 seconds the measurement cycle was started and about 315 µl of this solution were then automatically distributed into electrodes 1 and 2. 50 µl of enzyme-ethanol complex solution were added to 775 µl of remaining solution in mixing chamber. The difference in pH of electrode 1 and electrode 2 was observed. The pH deviation is a result of ethanol conversion to acetaldehyde.

By the same procedure, differences in pH were measured for buffer solution devoid of sample and for buffer with only 250 µl of Magic N50. These measurements were done in order to see the substances effect on system measurement.

Additionally, the reaction was observed by measuring the absorption of solution caused by the consumption of NAD⁺, or the formation of NADH during the conversion of ethanol to acetaldehyde within spectrophotometer at 340 nm of ultraviolet light.

3. Results and discussion

The differential pH analyzer system CL 10 was achieved detection of differences in pH between solutions. Thus, this device was used to measure change in pH, which is caused by enzymatic alcohol dehydrogenize reaction, for samples containing 0 – 99.985 % of alcohol.

After construction of delta pH value of buffer, the obtained delta pH values were in range from -0.891 mpH to -33.269 mpH for a full ethanol concentrations range of 0.00001714 M - 17.14 M respectively. These obtained delta pH values are sufficient to be used in quantifying any ethanol concentration in sample. The required time to reach constant delta pH value, end point, is inversely proportional to the ethanol concentration and it is not exceed 100 seconds. These results can be shown in figure 2.

The results indicated a nonlinear relationship between the alcohol concentration and the obtained change in pH as shown in figure 3. The non linearity was because of change in ethanol concentration which caused a change in the enzymatic reaction order from first to zero order. As approved, the order of reaction will be changed from first order reaction for low substrate concentration to zero order for high substrate concentration (Dunn et al 2003; Murry, 2003; Lee 2001). Such nonlinear plot can not be used as a calibration curve for ethanol determination.

Since ethanol concentration is stoichiometric to liberated H⁺ according to equation1 and pH of solution is a logarithmic function of [H⁺], a calibration curve was done by plotting the logarithmic value of ethanol concentrations against change in pH as shown in figure 4. This figure shows a linear relationship which can be used in quantification of ethanol concentration. Figure 4 consist of two parts a and b. Part a can be used for obtained change in pH up to -2.7 m pH while part b can be used for obtained change in pH starting from -3 up to -33 m pH as shown in figures (4.a & 4.b). The correlation coefficient of the linear regression of calibration curve, R, was 0.988 which means that the difference between predicted values and observed values of ethanol concentration is too low. This suggestive of a good model fit. In addition to that Chi square test was applied in order to evaluate the calibration curve precision. Degree of freedom, n-1, for seven standard samples of ethanol is 6. Therefore, for 6 degree of freedom and 0.05 probability, Chi critical table value is 12.592 (Snedecor and Corchran, 1989).The Chi square value was calculated 0.002 as shown in table 1. Since Chi square value is less than Chi critical table value, null hypothesis is accepted. That means there is no significant difference between the expected and observed values. In order to estimate the precision and the accuracy of the measurements, statistical evaluation for the results was made. In table 2, maximum standard deviation and maximum coefficient of variation for three replicates of measurements were 0.7 and 5.8% respectively. This low coefficient of variation range demonstrates that the measurements had little dispersion from the mean value which makes the results more precise and confidential.

To value the results, the oxidation reaction of ethanol to acetaldehyde was observed by measuring the change in absorption within spectrophotometer at 340 nm. The change in absorption of solution was caused by the formation of NADH from NAD⁺ during the conversion of ethanol to acetaldehyde. The absorption values can be shown in
figures 5 and 6. The results showed that the time required reaching end point of the reactions for standard ethanol samples was not exceeding 100 seconds. These results are satisfied the results obtained by measuring change in pH within differential pH analyzer system.

Curtipot program was used to estimate the capacity of the buffer and to measure the equivalent end point of traditional titration curve. The result indicated that the buffer of 0.02 M of glycine and 0.02 M of pyrophosphate will have one equivalent point, although it contained polyprotic acids, when it was titrated with 0.1 M of NaOH. This means it actually behaves like monoprotic acid and it is stable for the measurements as shown in figure 7.

In order to see the stability of differential pH analyzer system CL10, the following two experiments have been done in absent of ethanol oxidation reaction. The changes in pH were measured for systems consist of either buffer only or by adding 250 µl of Magic N50 solution to the buffer. The results indicated that glycine-pyrophosphate buffer was suitable to be used for this system as there was no noise in measurement. Also there was no noticeable influence of the Magic N50 solution on the measurement. The ability of these preparations to influence the measurement can be seen in figure 8.

Additional experiments had been done for samples of ethanol concentrations from 0.1714 to 17.14 M. In these experiments Magic N50 solution was prepared from NAD⁺ solution of 100mM. The obtained delta pH values were increased by a factor of 1.6 in comparison with those using 50 mM of NAD⁺ as shown in figure 9. The general behavior was still nonlinear. This indicates that there is no reason to increase the amount of NAD⁺ for this ethanol essay as that will lead to increase the essay costs and will be on expensive of time too. The set up of differential pH analyzer device CL10 has a limit of 300 seconds of maximum reaction time. However, Magic N50 solution of 100 mM NAD⁺ or more could be used if somebody wants a wide range of delta pH for his own standard calibration curve.

4. Economic aspect

In order to evaluate the economic aspect of this new method, a comparison with KIT method (K-ETOH) of Megazyme International Ireland Ltd was done. The results of this economic study (calculation is not showed) indicated that assaying 373 samples by using KIT method require using 1865mg NAD⁺, 5.6mg alcohol dehydrogenase (ADH) enzyme and 154.4mg aldehyde dehydrogenase (AL-DH) enzyme. However, applying the present method for assaying same number of samples utilizes 155mg NAD⁺ and 4.7mg alcohol dehydrogenase enzyme. Aldehyde dehydrogenase enzyme was not used in the present method.

The above calculation demonstrates the following:
1. There is a factor of 12 in the number of samples that can be assayed for the same amount of NAD⁺, which means a noticeable cost reduction.
2. In addition to that, KIT assay uses two enzymes, alcohol dehydrogenase ADH and aldehyde dehydrogenase AL-DH, while we just have used one enzyme, alcohol dehydrogenase. It could be seen that the amount of alcohol dehydrogenase enzyme for our method is slightly less than of that used in KIT method.

Additional cost by applying Kit assay is to be considered due to the use of NADH. The use of KIT assay also needs an additional calibration curve of NADH absorbent because of the necessity of accurate quantification within spectrophotometer.

5. Conclusions

It is possible to detect full range of ethanol concentration with lowering the cost. The recent method is a development for enzymatic methods by reduction the number of required enzymes to one. Also there is no need to dilute samples during the analysis. This will decrease the errors which are happened by users often. Using logarithmic values of ethanol concentrations in making calibration curve bring new approach in concentration quantification by using change in pH. This work offers the basis to start studies of full range concentrations quantification within differential pH measurement sensor device. Therefore, work in optimization is recommended.

Acknowledgements

This work was supported by Institute of Bioprocess Engineering at Erlangen University.

References


Table 1. Statistical evaluation of the calibration curve: Chi square worksheet direction

<table>
<thead>
<tr>
<th>Ethanol sample (mM)</th>
<th>Mean value of obtained change in pH(mpH)</th>
<th>Observed log[ethanol] (O)</th>
<th>Expected log[ethanol] (E)</th>
<th>(O-E)^2/E</th>
</tr>
</thead>
<tbody>
<tr>
<td>17147</td>
<td>33.269</td>
<td>4.234</td>
<td>4.226</td>
<td>0.008</td>
</tr>
<tr>
<td>1714.7</td>
<td>24.519</td>
<td>3.234</td>
<td>3.289</td>
<td>-0.055</td>
</tr>
<tr>
<td>171.47</td>
<td>13.640</td>
<td>2.233</td>
<td>2.124</td>
<td>0.110</td>
</tr>
<tr>
<td>17.147</td>
<td>5.876</td>
<td>1.233</td>
<td>1.292</td>
<td>-0.059</td>
</tr>
<tr>
<td>1.715</td>
<td>2.479</td>
<td>0.069</td>
<td>0.094</td>
<td>-0.026</td>
</tr>
<tr>
<td>0.172</td>
<td>1.623</td>
<td>-0.931</td>
<td>-0.983</td>
<td>0.052</td>
</tr>
<tr>
<td>0.017</td>
<td>0.892</td>
<td>-1.931</td>
<td>-1.904</td>
<td>-0.028</td>
</tr>
</tbody>
</table>

\[ \sum \frac{(O-E)^2}{E} = 0.002 \]

\[ = \text{Chi square value} \]

Mean values were multiplied by (-1) as the measurements were in (–mpH).

Table 2. Statistical evaluation of the measurements

<table>
<thead>
<tr>
<th>Ethanol sample (mM)</th>
<th>17147</th>
<th>1714.7</th>
<th>171.47</th>
<th>17.147</th>
<th>1.715</th>
<th>0.172</th>
<th>0.017</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean value of obtained change in pH(mpH)</td>
<td>33.269</td>
<td>24.519</td>
<td>13.64</td>
<td>5.876</td>
<td>2.479</td>
<td>1.623</td>
<td>0.892</td>
</tr>
<tr>
<td>Number of replicates</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Standard deviation</td>
<td>0.377</td>
<td>0.116</td>
<td>0.703</td>
<td>0.247</td>
<td>0.098</td>
<td>0.079</td>
<td>0.052</td>
</tr>
<tr>
<td>Coefficient of variation %</td>
<td>1.1</td>
<td>0.05</td>
<td>5.2</td>
<td>4.2</td>
<td>3.9</td>
<td>4.9</td>
<td>5.8</td>
</tr>
</tbody>
</table>

Mean values were multiplied by (-1) as the measurements were in (–mpH).

Figure 1. Schematic diagram of the differential pH analyzer system CL10.P1 to P5, peristaltic pumps; C, mixing chamber; M, stirring motor; N, magnetic stirrer; E1 and E2, class capillary electrodes pH sensors; A, differential amplifier (Luzzana et al 1983).
Figure 2. Delta pH (mpH) measured with time within differential pH analyzer system CL10 for different ethanol concentrations samples by using Magic N50 solution of 50mM NAD\(^+\). Each point is an average value of three measuring values.

\[ y = 18.197x^{0.2791} \]
\[ R^2 = 0.9873 \]

Figure 3. Delta pH (mpH) measured against different ethanol concentrations by Differential pH analyzer system CL10. Each point is an average value of three measuring values. The absolute values were obtained by multiplying the values of change in pH by (-1).

\[ \text{Absolute delta pH (mpH)} = 18.197^* [\text{Ethanol}]^{0.279} \]
\[ R^2 = 0.987 \]
Figure 4. Calibration curves between logarithmic values of ethanol concentrations against change in pH. Linear relationship was obtained. Change in pH is absolute value which was obtained by multiply the obtained change in pH, delta pH, by (-1).

(a) can be used for obtained change in pH up to -2.7 mpH

(b) can be used for obtained change in pH starting from -3 mpH.
Figure 5. Absorption of formed NADH against time within spectrophotometer during Enzymatic Oxidation of different ethanol concentrations

Figure 6. Absorption of formed NADH against time within spectrophotometer during Enzymatic Oxidation of different ethanol concentrations

Figure 7. Titration of pyrophosphate-glycine buffer of (20mL, 0.02mol/L) with (0.1 mol/L NaOH)
Figure 8. Observation of stability of differential pH analyzer system CL10. This measurement was devoid from ethanol oxidation reaction by using:

1. only buffer
2. adding 250µl Magic N50 solution to the buffer

Figure 9. Delta pH (mpH) measured with time within differential pH analyzer system CL10 for three different ethanol concentrations samples by using Magic N50 solution of 100mM NAD+.