Demonstration of potentiometric pH

determination using the filament of an

incandescent lightbulb as indicator electrode

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Abstract

One of the earliest concepts of chemistry taught in school is pH. It is usually in-

troduced during the discussion of acids and bases. Students are taught to detect the

acidity of substances using pH indicators, usually in the form of indicator paper. While

this method is convenient and easy for students to relate to, it does not give any deeper

insight into the meaning of pH. To truly comprehend pH, students need to learn to

approach the concept from an electroanalytical point of view. The aim of this paper is

to avail teachers of an interactive and demonstrative experiment in order to facilitate

this transition.

We demonstrate the fabrication of a simple and economical metal/metal-oxide type

pH indicator electrode using simple household items. The sensing element is made from

the tungsten alloy filament of an incandescent lightbulb, that is sandwiched between

two pieces of Plexiglass using superglue.

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Keywords

pH, tungsten electrode, light-bulb, potentiometry

Introduction

pH is one of the most important concepts in chemistry, originally defined by Sørensen in 1909¹ as the negative logarithm of the concentration of hydrogen ion:

$$pH = -\log_{10} c_{H^+} \tag{1}$$

At first glance, it looks like a deceptively simple concept. Complications around this definition start to arise when we replace concentration with activity, taking into account that hydrogen ions have charge, therefore their behaviour correlates more closely with activity than concentration:

$$pH = -\log_{10} a_{H^{+}} = -\log_{10} c_{H^{+}} \gamma_{H^{+}}$$
(2)

where a_{H^+} is the activity and γ_{H^+} is the activity coefficient of hydrogen ions. This definition however raises another problem. It includes the activity coefficient of a single ion, which cannot be separated from the activity coefficient of counterions. Without knowing the contribution of individual ions to the activity coefficient, only the mean activity coefficient can be measured, for example in the case of HCl solution:

$$\gamma_{H^+,Cl^-} = \sqrt{\gamma_{H^+}\gamma_{Cl^-}} \tag{3}$$

Even though the Debye–Hückel equation can be used to calculate a theoretical activity coefficient for a single ion, the prevailing view is that if it cannot be measured, then a pH definition that is based on that activity coefficient is an unmeasurable quantity. This logical difficulty was summarized by Bates and Guggenheim in 1960.²

Their recommended solution was a convention, which became known as the *Bates–Guggenheim* convention. According to the convention, first the activity coefficient of chloride ions was calculated using the extended Debye–Hückel equation:

$$-\log_{10}\gamma = \frac{Az^2\sqrt{I}}{1 + Ba\sqrt{I}}\tag{4}$$

where A and B are constants, a is ionic radius and I is ionic strength calculated with the following equation:

$$I = \frac{1}{2} \sum_{i=1}^{n} c_i z_i^2 \tag{5}$$

where c_i is and z_i is the concentration and charge of the *i*th ion, and *n* is the total number of ionic species in the solution. Once the activity coefficient of chloride ion was calculated, the activity coefficient of another ion could be determined by measuring the mean activity coefficient of an electrolyte that it forms with chloride ions. Then, for instance, the activity coefficient of hydrogen ion can be calculated as:

$$\gamma_{H^+} = \frac{\gamma_{H^+,Cl^-}^2}{\gamma_{Cl^-}} \tag{6}$$

In this way, the contribution of chloride ions to the mean activity coefficient is accepted as correct by definition, which in turn makes the determination of individual activity coefficient of other ions possible. This convention has been the basis for the IUPAC (International Union of Pure and Applied Chemistry) recommendation to define pH in 1985³ and 2002,⁴ which is currently the most up to date definition from IUPAC:

$$pH = -\log_{10} a_{H^+} = -\log_{10} (m_{H^+} \gamma_{m,H^+} / m^{\theta})$$
(7)

where m_{H^+} is the molality and γ_{m,H^+} is the molality based activity coefficient of hydrogen ions, m^{θ} is the standard molality (1 mol kg⁻¹). The activity coefficient of hydrogen ions in

the above equation is calculated using the Bates-Guggenheim convention.

The electrochemical cell used in the measurement of the primary pH standards is known as the *Harned Cell*. ⁵ To measure the pH in such a cell, a conventional procedure was developed at NBS (National Bureau of Standards) ⁶ and recommended at present by the last IUPAC recommendations. ⁷ NIST (National Institute of Standards and Technology) in the U.S. and PTB (Physikalisch-Technische Bundesanstalt) in Germany have presented pH values using the Harned Cell.

In practice, pH is most commonly measured with a glass electrode. It has been known for more than a hundred years that the potential difference measured across a glass membrane depends on the ratio of the hydrogen ion activity in the two solutions that the membrane separates. ^{8,9} The glass electrode is usually combined with a reference electrode for convenience. Such an electrode pair is called the "combined glass electrode", and this the most common type the students might encounter. The glass electrode is the best electrochemical sensor and one of the best sensors ever made, with a linear response of over more than 13 orders of magnitude and excellent selectivity. However, it's not perfect. One of the imperfections is that although to a negligible extent, it responds to cations other than hydrogen ion. This behaviour is described by the Nikolsky–equation that takes the effect of interfering ions into account with the so-called "selectivity coefficient": ¹⁰

$$E = E^{\theta} + \frac{RT}{z_i F} \ln \left[a_i + \sum_j \left(k_{ij} a_j^{z_i/z_j} \right) \right]$$
 (8)

There are several other type of electrodes that can be used to measure pH. One of them is the ionophore based hydrogen ion selective electrode. Its function is based on an organic ionophore that selectively complexes hydrogen ions. When such an ionophore is embedded in a PVC based membrane, the crossmembrane potential difference depends on the activity ratio of hydrogen ion at the two opposing sides of the membrane. ¹¹

There are certain applications where the use of a glass electrode might be challenging or impossible. These include measuring pH at high temperature or in hydrogen–fluoride solution, or application in the food industry. Intensive research has been conducted for several decades to find alternatives to the glass electrode. One such alternative is to use metal/metal-oxide electrodes. One of the most often used type of these is the Ir/IrO₂ electrode. ¹² The oldest is certainly the Sb/Sb₂O₃ electrode, its initial characterization dating back to 1923. ¹³ It is pH sensitive because hydrogen ions participate in the equilibrium between antimony and its oxide on the surface:

$$2 \operatorname{Sb}_{(s)} + 3 \operatorname{H}_2 O \Longrightarrow \operatorname{Sb}_2 O_3 + 6 \operatorname{H}^+ + 6 \operatorname{e}^-$$
(9)

Another very popular metal/metal-oxide electrode used for pH measurements is the tungsten electrode. Its function is based on the intercalation and deintercalation of hydrogen ions into the oxide layer that is followed by a change in the oxidation state of tungsten oxide: ¹⁴

$$WO_3 + xH^+ + xe^- \Longrightarrow H_xWO_3$$
 (10)

It is well known that the filament of incandescent lightbulbs is made of an alloy called "tungsten bronze" that is mostly tungsten. ^{15–17} It means that it might be possible to fabricate a pH sensitive indicator electrode out of a traditional lightbulb.

In this paper the construction and usage of a simple and inexpensive tungsten pH sensitive electrode is presented, in which the sensing element is a tungsten filament salvaged from an incandescent lightbulb. This electrode could provide an interesting teaching aid like the pH sensitive coverslip glass electrode developed and characterized by Yong et al. ¹⁸ or other similar, but ionohore based indicator electrodes ^{11,19} The electrode can be assembled in a highschool or undergraduate chemistry laboratory by students and it can be used as an aid to demonstrate pH and ion-selective electrodes. The excercise can also be useful as an introduction to the IUPAC definition of pH, since it is closely intertwined with the potentiometric determination of pH.

Materials and Methods

Electrode fabrication. To obtain the sensing element of the electrode, a 100 W incandescent lightbulb (Tungsram brand, purchased at local hardware store) was carefully wrapped in cloth and broken with a mallet. The contact wire that still had the filament attached was cut near the stem (Fig. 1a). The electrode body was prepared from two, 4 mm thick, ≈ 0.5 cm $\times \approx 2$ cm, identically shaped Plexiglass pieces (Fig. 1b). In one of them, a ≈ 1 mm groove was cut longitudinally, in which the filmanet with stem was layed (Fig. 1c). Then, superglue was applied onto the Plexiglas surface that contained the groove with the filament and stem. The other Plexiglass piece was pressed onto the superglued surface, and was held firmly until the superglue cured (Fig. 1d). The electrode was sanded and polished from all sides except where the electrical lead, the stem was protruding (Fig. 1e).

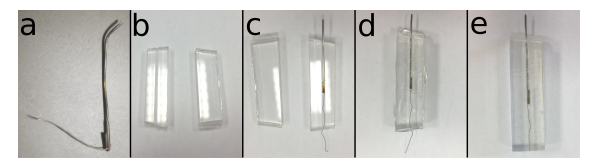


Figure 1: The fabrication process of the tungsten filament electrode.

Calibration. The fabricated tungsten filament electrode was calibrated in a series of Britton–Robinson buffers with the following pH values: 2.19, 3.04, 4.10, 5.12, 6.09, 7.19, 8.08, 8.89, 9.87, 10.88, 11.81. The pH was measured with a WTW SenTix combined pH electrode connected to a WTW inoLab pH 740p high input impedance pH meter (Xylem Analytics Germany Sales GmbH & Co. KG, Weilheim Germany), that was calibrated with three pH buffers (4.00, 7.00, 10.00 at 20 °C, Scharlab,S.L. Barcelona, Spain). The glass and the tungsten filament electrodes were calibrated by measuring their potential against the internal reference electrode of the glass electrode in the buffer series. The potential values

were recorded when the instrument indicated a steady reading. This ensured that they were taken at the same point of the response curve.

Selectivity study. Since the primary ion for the glass and the tungsten filament electrodes is already present in water at an activity that would influence the selectivity study, a buffer had to be used to lower the hydrogen ion activity. Most of the common pH buffers contain alkaline metals, and they could not be used to carry the selectivity study, because alkaline and alkaline earth metals are the most common interferences. For this reason, a pH = 8 TRIS buffer was used as a solvent to create the dilution series using KCl, NaCl, CaCl₂ and MgCl₂. In each cases, the activities ranged from 10^{-6} M to 10^{-1} M with tenfold increments in activity. The electrodes were immersed in each of the solutions and the resulting potential differences were measured and recorded with the already mentioned pH meter.

Hazards

When salvaging the tungsten filament from the lighbulb, care must be taken. The lightbulb must be wrapped several times in a cloth before it is hit with a mallet. Care must be taken when the filament is picked up from the broken pieces of glass. When the Plexiglass body is prepared, a saw is used to cut the groove, then superglue is used to enclose the filament in the groove. During these steps, close supervision of the students is necessary to avoid any harm. Lastly, HCl and NaOH is used during the preparation of the pH buffer series. If these get in contact with skin or the eye, flush with plenty of water for several minutes.

Results and Discussion

To make sure that the measured pH values are accurate, the glass electrode was first calibrated with high accuracy pH buffers. The resulting slope was 57.19 mV / pH at 21 °C. Then, the pH values of the created Britton–Robinson buffer series were measured. Once

these were known with high accuracy, the same buffer series was used again to calibrate the tungsten filament electrode. The resulting calibration plot can be seen in Fig. 2B. The slope of the calibration equation is 40.05~mV / pH, which is significantly lower than that of the theoretically expected value from the Nernst–equation, or the one that was measured for the glass electrode in this study. The lower value might be explained by the alloying elements in the tungsten filament. It is well known, that the filaments of incandescent lightbulbs are not made of pure tungsten, but a so-called "tungsten bronze" alloy. The alloy consists of tungsten and potassium or sodium, which improves ductility that helps during the manufacturing process. $^{15-17}$ Even high purity metal/metal—oxide electrodes don't reach the nernstian slope, which is well documented in the literature Nonetheless, a slope of 40.05~mV / pH is enough to measure pH reliably. The electrode response was slighty non-linear, but it can be said that it works well within the tested pH range of 2–12.

Because of these alloying elements, the tungsten filament electrode was expected to be less selective towards hydrogen ions than the glass electrode. The selectivity study was performed in dilution series of KCl, NaCl, CaCl₂ and MgCl₂ from 10^{-1} M to 10^{-6} M with tenfold increases in concentration. A buffer was used to lower the hydrogen ion activity, and because most of the alkaline buffers contain an ion that interefes with hydrogen ion–selective electrodes, the TRIS buffer was selected at pH = 8. The TRIS buffer had a concentration of 1.25 M. This concentration was more than enough to increase the ionic strength to a value at which the activity coefficients for the members of the buffers series can be regarded as constant, and concentration can be used in the selectivity calculations instead of activity. The measured potential values for the glass and tungsten filament electrodes for the interfering ions can be seen in Fig. 2A and B, respectively. As expected, no interference can be observed for any of the tested ions when the glass electrode was used. However, when measured with the tungsten filament electrode, some of the tested ions interfered with the pH measurement at higher than 10^{-4} M concentration. Below this concentration the potential corresponding to pH = 8 buffer was observed (\approx -315 mV). Above 10^{-4} M, most of the ions were interfering

to a small extent. Interestingly, K^+ caused the potential to drop at higher concentration. This result was replicated several times. The selectivity coefficients were calculated with the separate solution method from activities corresponding to the same potential. The selectivity coefficient values are summarized in Table 1.

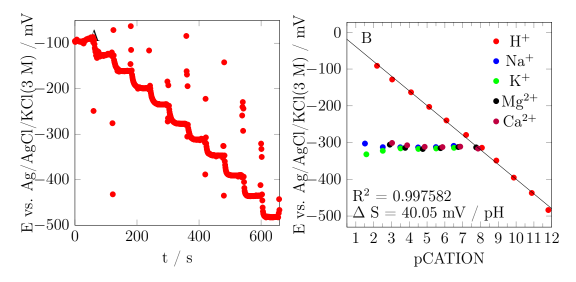


Figure 2: Calibration and selectivity study of the glass electrode (A), and the tungsten filament electrode (B).

Table 1: The selectivity coefficients of the tungsten filament electrode for the different cations.

Interfering ion	Selectivity coefficient
Na ⁺	$8.57 \cdot 10^{-7}$
K^{+}	$1.91 \cdot 10^{-7}$
Ca^{2+}	$9.48 \cdot 10^{-7}$
Mg^{2+}	$6.17 \cdot 10^{-7}$

Conclusion

The construction and usage of an economical pH sensitive electrode was presented that is highly selective towards hydrogen ions. It has been shown that the electrode performs very well, with a 40.05 mV / pH slope in the pH range 2–12. There was no significant interference from K^+ , Na^+ , Mg^{2+} and Ca^{2+} ions. The potentiometric cell utilizing the

electrode can be used as an engaging demonstration of the potentiometric determination of pH. The participating students showed high motivation during the electrode fabrication process. They were excited to use electrodes that they've prepared themselves to measure pH. As a result of the interactive demonstration and evaluation, they showed a deeper understanding of pH and its potentiometric determination.

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