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Electrochemical Production of Hydrogen and Sulfur by Low-Temperature Decomposition of Hydrogen Sulfide in an Aqueous Alkaline Solution

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ABSTRACT

Electrolysis of hydrogen sulfide to its constituents in a solution containing equimolar concentrations of NaOH and NaHS has been carried out at 80°C. In a double-compartment cell employing Naion® membrane as a separator, both crystalline elemental sulfur and high-purity hydrogen have been produced at high current efficiencies. Only minimal, if any, passivation of the anode by sulfur product was observed. According to solution composition, electrolysis could result in gas evolution at the anode, passivation of the anode by sulfur deposition, or oxidation of sulfide (S²⁻) or polysulfide (Sₓ²⁻) to sulfur oxyanions. However, in an optimized solution, electrolysis gave only anodic sulfur via bisulfide (HS⁻) and sulfur oxidation. Voltammetric and chronopotentiometric studies showed that sulfide, bisulfide, and polysulfide oxidation occurred at about the same potential.

Noring and Fletcher (1) pointed out some economic benefits that could arise from the production of elemental sulfur and hydrogen by electrolytic decomposition of hydrogen sulfide. To realize this goal, the problem of anode passivation by the sulfur produced must be solved. In indirect electrolysis, the use of either hot organic vapor to remove the sulfur from the electrode surface (2) or organic solvents to extract sulfur from the solution (3) have been suggested. In indirect electrolysis, hydrogen sulfide is first oxidized to sulfur dioxide and the oxidant is regenerated at the anode with simultaneous evolution of hydrogen at the cathode (4-7). Advantages of the direct approach involve a lower input power than that required to regenerate the reduced oxidant and that it will produce crystalline rather than colloidal sulfur, which is the normal product of indirect electrolysis.

Recently, direct electrolysis schemes without the use of solvents for sulfur have been proposed (8) or investigated (9). In one of these (9), it was reported that raising the temperature of electrolysis to 60°C can eliminate the blocking effect. This approach required a continuous flow of hydrogen sulfide through the electrolytic cell, thereby shifting the pH of the solution close to neutral values. In a separate publication (11), we have shown that these approaches involve problems such as low current efficiency, poor purity of the sulfur product and the formation of sulfur oxyanions. These will have to be overcome for the process to be commercially attractive. Further, analysis showed that the chemical, electrochemical, and mass transport processes involved in H₂S electrolysis under the conditions reported by Dandapani et al. (9) would be unlikely to result in sulfur precipitation from the anolyte, but would rather lead to anode passivation.

Our previous work (11) outlined the sequential steps (Fig. 1) which lead to complete electrolysis of hydrogen sulfide. Further, experimental results obtained from the electrolysis of hydrogen sulfide in alkaline solution are presented here. The conditions for efficient electrolysis are that the pH of the solution, thus the relative concentration of NaOH and NaHS, and the temperature are adequately controlled.

Experimental

Materials and materials preparation.—The solutions used in these experiments were prepared from deionized water and reagent-grade materials. High-purity hydrogen sulfide was obtained from Matheson Gas Company. The glass electrolytic cell, schematically shown in Fig. 2, consisted of two compartments separated by an ion-selective Naion® membrane. Anode materials employed included graphite, nickel, porous nickel-chromium alloy, as well as titanium. Cathode materials were either nickel or graphite. Two Hg/HgO (1M NaOH) reference electrodes were separately connected to the anode and cathode compartments via Luggin capillaries and were utilized for monitoring half-cell potentials.

Experimental procedures—Two groups of experiments are discussed in this work. In the first, hydrogen sulfide was bubbled continuously through the electrolytic cell before and during electrolysis. In the second, H₂S was absorbed in the NaOH scrubber, then introduced into a further compartment to adjust solution pH in the alkaline range. This solution was then introduced into the electrolytic cell. Conditions in the first group of experiments are the same as those reported by Dandapani et al. (9) where the electrolyte in the cell was essentially pure NaHS. In the second group of experiments, the electrolyte was NaHS plus NaOH, in different ratios. In addition to these solutions, Na,S was also examined as an electrolyte.

A Solartron Programmable Electrochemical Interface, Model 1286, was used as the power source for the cyclic

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† On saturation of a NaOH solution with hydrogen sulfide, NaHS is formed as a product via the reaction: NaOH + H₂S \rightarrow NaHS + H₂O. Results obtained from solutions of reagent-grade NaHS in this work were identical to those from solutions of NaHS produced by the reaction of H₂S and NaOH.
The anode potential rose sharply to 1.8V with accompanying gas evolution. The sharp rise in overpotential may result from chemical reaction to give polysulfide species, as is the case at higher pH values, i.e., higher NaOH:NaHS ratios.

Electrolyte concentration ratio (NaOH:NaHS) = 1:1. The color of this solution was yellow, turning to light orange upon continued electrolysis at 20 mA/cm². The electrode potential at this current density was more positive than ~0.4V vs. Hg/HgO and was constant until ca. 60% of the total sulfur content of the solution was oxidized, when it increased sharply to ~0.5V vs. Hg/HgO. Stirring restored the potential to the previous value, suggesting that the electrolyte at the interface was depleted of the electroactive species. On continued electrolysis, no further rise in potential was observed, so long as the solution was continuously agitated, so no surface blocking or coating by sulfur occurred. When the solution was not stirred, electrolysis was carried out at 5 mA/cm² to prevent the anode potential from reaching values where sulfur oxyanions might form. After ca. 85% of the total sulfur content was oxidized, the current was again increased to 20 mA/cm², when yellow precipitates formed on the electrode surface and continuously fell from the electrode. This process was accompanied by a slight increase in the electrode potential. Continuous application of current resulted in a sharp rise of the potential to about +1.4V vs. Hg/HgO, giving a colorless solution and gas evolution. Electrolysis was then terminated. Very little increase in potential occurred during electrolysis, therefore blocking effects were eliminated. The pH of the anolyte changed from 13.4 to 9.6 during electrolysis.

Figure 3 shows polarization curves for the anode at different stages of electrolysis, corresponding to the oxidation of sulfide species to sulfur. The lowest potential for this conversion corresponded to ca. 25% conversion of the initial sulfur present. Further electrolysis resulted in an increase in anode potential, possibly attributable to concentration polarization and pH changes.

The anolyte was filtered, and the yellow precipitate was characterized using XRD and EDS techniques. The x-ray diffraction pattern is shown in Fig. 4, and indicates a crystalline orthorhombic structure, i.e., α-sulfur. Figure 5 is a representative EDS spectrum at 15 kV illumination, showing a strong sulfur peak with a trace amount of silicon. The purity of the product was confirmed using powder x-ray diffraction. A Picker 638 XRD generating CuKα radiation interfaced with a PDP 11/23 computer under an RSX llM operating system with in-house control and data gathering software was used for the analysis.

Since previous analysis (11) showed that sulfur can be precipitated near the anode from an equimolar NaOH:NaHS solution by continued electrolysis, different ratios of NaOH to NaHS were used as anolytes with 1M NaOH catholyte. The initial volumes of both before electrolysis was 25 ml. Electrolysis was carried out galvanostatically.

Results and Discussion

Galvanostatic studies.—Electrolyte concentration ratio (NaOH:NaHS) = 0:1.—The as-prepared 1M NaHS solution was slightly yellow. The steady-state open-circuit potential of the anode was ~0.37V vs. Hg/HgO (1M NaOH). Upon electrolysis at 20 mA/cm², the overpotential gradually increased from an initial value of ~0.4V and the solution became deeper yellow. With continued electrolysis, the potential rose sharply to 1.8V with accompanying gas evolution. The sharp rise in overpotential may result from precipitation of an insulating sulfur layer on the electrode surface. At this point only a very low current density (5 mA/cm²) could be applied at a low overpotential. Electrolysis was terminated when a black coating formed on the anode and the overpotential rose to a value where gas evolution again occurred. The total amount of charge passed during this time was equivalent to only about 50% of the total sulfur content in the anolyte solution.

The black coating from the electrode surface was heated in a glass ampul. Yellow vapor was observed to condense on its walls when the temperature was slightly above the melting point of sulfur.

The pH of the bulk anolyte decreased from an initial value of 12.6 to 9.4 when electrolysis was terminated, suggesting that the pH at the electrode/electrolyte interface was even lower during electrolysis. As a result, any sulfur-oxidized deposits on the electrode surface without further chemical reaction to give polysulfide species, as is the case at higher pH values, i.e., higher NaOH:NaHS ratios.

Electrolyte concentration ratio (NaOH:NaHS) = 0:1.—The color of this solution was yellow, turning to light orange upon continued electrolysis at 20 mA/cm². The electrode potential at this current density was more positive than ~0.4V vs. Hg/HgO and was constant until ca. 60% of the total sulfur content of the solution was oxidized, when it increased sharply to ~0.5V vs. Hg/HgO. Stirring restored the potential to the previous value, suggesting that the electrolyte at the interface was depleted of the electroactive species. On continued electrolysis, no further rise in potential was observed, so long as the solution was continuously agitated, so no surface blocking or coating by sulfur occurred. When the solution was not stirred, electrolysis was carried out at 5 mA/cm² to prevent the anode potential from reaching values where sulfur oxyanions might form. After ca. 85% of the total sulfur content was oxidized, the current was again increased to 20 mA/cm², when yellow precipitates formed on the electrode surface and continuously fell from the electrode. This process was accompanied by a slight increase in the electrode potential. Continuous application of current resulted in a sharp rise of the potential to about +1.4V vs. Hg/HgO, giving a colorless solution and gas evolution. Electrolysis was then terminated. Very little increase in potential occurred during electrolysis, therefore blocking effects were eliminated. The pH of the anolyte changed from 13.4 to 9.6 during electrolysis.

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relative intensities of the S:Si peaks should not be used as a measure of sample composition since sulfur deteriorates rapidly under the beam compared with Si. However, after 43s of beam exposure time, the nominal composition was 98.63:1.32:0.05 atom percent (a/o) (S:Si:Na). Silicon in the product may result from slight dissolution of the glass cell.

As expected, the conversion efficiency for sulfur production estimated gravimetrically is less than that estimated coulometrically. Table I lists the weights of the solid product and the estimated efficiencies for three different electrolysis runs from solutions of 1:1 NaOH:NaHS molar ratios. Estimated efficiency values are between 60 and 70% of the zero valence sulfur content of the initial anolyte solution. In comparison, however, the current efficiency estimated from the total coulombs passed during electrolysis was about 100%.

**Electrolyte concentration ratio (NaOH:NaHS) = 1.5:1.** Unlike the previous experiment, neither sulfur deposition nor gas evolution was observed at the anode on electrolysis, even when the amount of charge passed exceeded 150% of the total sulfur species in solution. A possible explanation is that further oxidation of polysulfide species to sulfur oxyanions had taken place. This is possible because of the high pH of the solution, which only changed from an as-prepared value of 13.7 to 13.3, when electrolysis was terminated.

In all of the above three anolyte solutions, the catholyte remained colorless during electrolysis, showing that transfer of sulfide species to the cathode compartment was small. Thus, the cathode process was reduction of water to hydrogen close to 100% efficiency. The cathode gas product was collected over water and analyzed using a gas chromatograph. Its normalized composition was 99.97% hydrogen.

**Optimal electrolyte concentration.** The results have demonstrated the feasibility of hydrogen sulfide electrolysis at high conversion efficiencies in an equimolar NaOH:NaHS solution to produce hydrogen and crystalline sulfur, the latter with minimal passivation. During electrolysis, water was transported across the membrane from the anodic to the cathodic compartment, as noted by a change in level. The formation of sulfur results in the migration of cation (Na⁺) from the anodic compartment, through the membrane, to the cathodic compartment to preserve charge neutrality. At the anode, OH⁻ species are consumed in the typical process

$$\text{NaHS} + \text{OH}^- \rightarrow \text{S} + \text{H}_2\text{O} + \text{(Na}^+\text{)} + 2\text{e}^-$$

The pH is therefore reduced at the anode, with production of water in the electrolyte. Na⁺ and H₂O migrate through the membrane to the cathode, where H₂O is consumed

$$\text{(Na}^+\text{)} + \text{H}_2\text{O} + \text{e}^- \rightarrow \text{H}_2 + \text{NaOH}$$

Thus, the catholyte volume increases. Figure 6 shows the catholyte volume increase as a function of charge passed in electrolysis for initial catholyte and anolyte volumes of 25 ml. The potential difference across the membrane, measured by two reference electrodes on either side on current interruption, shows the same trend. This is shown in Fig. 7 and is due to the concentration difference across the anode-cathode compartments.

**Cyclic voltammetric studies.** In this study, cyclic voltammetry on graphite electrodes was carried out after different elapsed electrolysis times. The electrodes were small enough to prevent changes in overall composition. The anolyte solutions used were 1M NaHS, 1M NaHS + 1M NaOH, 0.34M Na₂S₄, and 0.25M Na₂S₄ + 0.25M NaOH. Expressed in terms of NaOH molar ratio, these are 0:1(NaHS), 1:1(NaHS), 0:1(Na₂S₄), and 1:1(Na₂S₄), respectively. The catholyte was always 1M NaOH initially.

**Table I. Measured weight of product sulfur and corresponding current efficiency for sulfur production during H₂S electrolysis in an aqueous alkaline solution.**

<table>
<thead>
<tr>
<th>Run number</th>
<th>Weight of sulfur collected (g)</th>
<th>a Percent of initial anolyte content</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run 1</td>
<td>0.529</td>
<td>66</td>
</tr>
<tr>
<td>Run 2</td>
<td>0.481</td>
<td>60</td>
</tr>
<tr>
<td>Run 3</td>
<td>0.497</td>
<td>62</td>
</tr>
</tbody>
</table>

*This percentage is based on 25 ml of the anolyte used in the experiment, and translates for a 1:1 (NaHS:NaOH) molar solution to 0.8015g of sulfur.*
Cyclic voltammograms prior to electrolysis are shown in Fig. 8. They all exhibit one broad anodic peak and a narrow cathodic peak near their open-circuit potentials, with an indication of a current increase at more positive potentials. However, two major characteristics may be noted: (i) the cathodic peak for polysulfide reduction is smaller in NariS-containing solutions than in Na₂S₄-containing solutions, which may be attributed to the known absence of polysulfide species in the fresh NaHSH solutions; (ii) there is a sharp reduction in current after the anodic peak in solutions without NaOH due to the formation of a passivating layer, whereas in equimolar NaOH solutions, the decrease occurs over a wider potential range. No evidence of bubble formation was observed during the second anodic current rise, implying that this may be due to further oxidation of polysulfide species. From the peak width, the anodic process may result from at least two successive electrode reactions, as is shown by the two peaks resolved at low scan rates (Fig. 9). The cathodic process is due to a simple reduction of polysulfide species or absorbed sulfur to sulfide ions.

If the reactions occurring near the anodic peak convert sulfide or low polysulfide to higher polysulfides, then electrolysis at a low overpotential should consume only these reactant species. Thus, the height of the anodic peak should decrease with electrolysis time. This is shown in voltammograms obtained after a fraction of the total sulfur has been oxidized (Fig. 10). The cathodic peak, however, becomes larger, and Fig. 11 shows the anodic peak height-square root of sweep rate relationship, which is linear, showing that reaction is under diffusion control. To confirm whether the first peak height corresponds only to the concentration of sulfide species, the voltammograms were obtained after bubbling H₂S through the solution, which produced a higher first anodic peak (Fig. 11).

Voltammograms in the cathodic compartment were identical at different stages of electrolysis in all cases. These results suggest that diffusion of polysulfide species from the anode to the cathode compartments through the membrane is negligible.
Chronopotentiometric studies.—Under constant current electrolysis conditions over an extended period of time, it was observed that the anode potential first decreases and then gradually increases (see for example, Fig. 3). This result is consistent with galvanostatic polarization measurements and may be attributed to a self-catalytic reaction. Short-term chronopotentiometric behavior of a graphite electrode, first in a freshly prepared 1:1 (NaOH:NaHS) solution (Fig. 12), and second, after several coulombs have been passed (Fig. 13), reveal characteristic features which indicate that the reaction involves parallel oxidation of two or more electroactive species. This indication is supported by the broadening of the anodic peaks in the voltammograms (Fig. 8) and also by the pH of the initial solution which suggests equilibrium concentrations of both HS⁻ and S²⁻.

Table II shows a generalized list of possible chemical and electrochemical reactions that might occur involving species that are pH and/or temperature dependent. Although formation of higher polysulfides is possible, currents at the initial stages of electrolysis are predominantly controlled by the oxidation of HS⁻, S²⁻, and S⁵⁻. At the onset of electrolysis, S⁵⁻ is absent and current results mainly from the oxidation of HS⁻ and/or S²⁻. A corresponding initial increase in anode potential results, which may be due to depletion of reactant species at the anode/
Electrolysis during continuous passage of H₂S through anolyte.—This section represents a summary of initial attempts (13) to reproduce the results of Dandapani et al. (9) for H₂S electrolysis in an alkaline environment. The anolyte and catholyte were either 1M or 6M NaOH solutions. High-purity H₂S gas was bubbled through the anolyte before and during electrolysis. The experimental arrangement was the same as before, except that a flat plate plexiglass cell containing 100 ml of solution in each compartment was used.

As electrolysis temperature was increased stepwise from 25°C to about 80°C, there was a marked increase in current at a constant potential or a similar decrease in anode overpotential at a constant current. These observations are consistent with the results of Dandapani et al. (9) and Allen and Hickling (14). However, prolonged electrolysis, even at 80°C, did not lead to sulfur precipitation. Rather, two distinct gas evolution processes accompanied by a slow increase in anode potential were observed. The first occurred a short time after the start of electrolysis at potentials more positive than -200 mV vs. Hg/HgO. This gas evolution process stopped when H₂S bubbling was discontinued and/or nitrogen was supplied. Since oxygen cannot be evolved at this potential, the process must be associated with sulfur-containing species. Possible gases are H₂S and/or SO₂, resulting from either, or a combination, of the following reactions

\[ 2\text{HS}^- \rightarrow \text{S} + \text{H}_2\text{S} + 2e^- \quad [1] \]

\[ \text{S}^2_2^- + 2\text{H}_2\text{O} \rightarrow \text{SO}_2^2^- + \text{H}_2\text{S} + 2\text{H}^+ + 4e^- \quad [2] \]

\[ 2\text{HS}^- + 2\text{H}_2\text{O} \rightarrow \text{SO}_2^2^- + \text{H}_2\text{S} + 2\text{H}^+ + 4e^- \quad [3] \]

In a neutral pH solution which is reminiscent of an H₂S-saturated solution, the standard potentials of reactions [2] and [3] are about -0.05 and -0.07V vs. NHE, respectively. Therefore, although all three reactions may possibly take place, the occurrence of [2] and [3] are more likely, especially since no further increase in the potential was observed on extended electrolysis.

The second gas evolution process occurred at a higher anode potential at a later stage of electrolysis and was attributed to sulfur blocking the electrode surface. It was reported earlier (9) that this blocking layer could be eliminated if electrolysis was carried out at a temperature higher than 60°C. Results at 80°C in this work did not confirm this. The potential on a graphite anode always sharply increased with time of electrolysis, and the blocking effect became more pronounced at lower pH when dissolution of sulfur from the electrode surface could not occur. This phenomenon was only observed when the low overpotential gas evolution process was avoided by starting electrolysis before complete anolyte saturation with H₂S, so that both gas-evolution processes were not successive. The second gas evolution process occurred at potentials higher than 1.9V vs. Hg/HgO and may be attributed to oxygen evolution.

These effects were not dependent on solution concentration, since the same behavior was observed in both 1M and 6M NaOH. However, when metal or metal-alloy anodes (Ni, Ni-Cr, and Ti) were used in place of graphite anode, the
high anode potential at the end of electrolysis was more pronounced.

Conclusions

The feasibility of an electrochemical process for the production of hydrogen and sulfur by the decomposition of hydrogen sulfide in an alkaline solution has been demonstrated. For this process to be successful, gas evolution at high anode potential at the end of electrolysis was more pronounced. For this process to be successful, gas evolution at high anode potential at the end of electrolysis was more pronounced. Results have demonstrated that the process previously proposed (11) is feasible by control of pH, temperature, and conditions of electrolysis. In particular, electrolysis in a saturated solution in the vicinity of equimolar NaOH:NaHS concentrations leads to hydrogen and crystalline sulfur production in quantitative yields (at least 95%). In particular, H2S must not be continuously bubbled into the anolyte during electrolysis so that pH can be properly controlled and anode passivation can be avoided.

In this work, sulfur was produced in a batch process. To be commercially viable, the process must be made continuous. One approach, described in more detail elsewhere (11), would be to continuously transfer the concentrated caustic, pumped through the membrane to the cathode compartment, to the scrubbing system, maintaining flow so that the solution leaves half-neutralized. Finally, in a practical system, the Naflon membrane may be replaced by an inexpensive microporous separator.

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Potential-pH Diagrams for Adsorbed Species

Application to Sulfur Adsorbed on Iron in Water at 25° and 300°C

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ABSTRACT

The principle of equilibrium potential-pH diagrams (Pourbaix diagrams) is extended to the case of two-dimensional species adsorbed on metals. Up to now, the only solid compounds taken into account in these diagrams have been three-dimensional compounds. However, species may be adsorbed in E-pH domains where three-dimensional compounds are unstable. The principle of the calculation of the E-pH relations existing at equilibrium in aqueous solution between the dissolved species and the species adsorbed on a metal is presented. The calculation is made for sulfur adsorbed on iron in water containing H2S, HS-, HSO3-, or SO42-. A similar calculation has also been done for sulfur adsorbed on platinum (5).

The principle and the use of equilibrium potential-pH diagrams, or Pourbaix diagrams, are well known (1). In these diagrams, potential-pH lines limit the domains of stability of the various chemical forms of an element in solution, at equilibrium, are calculated from the Nernst law. To our knowledge, the only solid compounds to be taken into account in E-pH diagrams have been three-dimensional compounds. However, the formation of a three-dimensional (bulk) solid compound MA (A may be H, O, N, or S) on a metal (M) is usually preceded by the formation of a two-dimensional phase of A adsorbed on the metal surface, which is more stable than the bulk compound (2). Thus, adsorbed layers may be formed in E-pH domains where a classical diagram does not predict such phases. This phenomenon must not be neglected because the presence of an adsorbed monolayer can induce marked changes in the reactivity of the metal. As an example, in corrosion, a monolayer of sulfur adsorbed on Ni or Ni-Fe alloys enhances the anodic dissolution and hinders the formation of the passive film, drastically affecting the corrosion resistance of the metal or alloy (3, 4).

The aim of this work is to establish the E-pH relations existing at equilibrium in aqueous solution between dissolved species of an element and the element adsorbed on a metal. The principle of the calculation is presented and thereafter applied to the adsorption of sulfur and oxygen on iron in water containing HS-, H2S, HSO3-, or SO42-. A similar calculation has also been done for sulfur adsorbed on platinum (5).