Removal of copper from water using a thermally regenerative electrodeposition battery

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HIGHLIGHTS

• We proposed a method that relies on waste heat to produce electrical power.
• The system was adapted to remove copper from water.
• Both electrical power production and treatment were achieved.
• The system obtained the highest copper removal of 77% and power production of 31 W/m².

GRAPHICAL ABSTRACT

ABSTRACT

A thermally regenerative ammonia battery (TRAB) recently developed for electricity generation using waste heat was adapted and used here as a treatment process for solutions containing high concentrations of copper ions. Copper removal reached a maximum of 77% at an initial copper concentration (C i) of 0.05 M, with a maximum power density (P) of 31 W m⁻²-electrode area. Lowering the initial copper concentration decreased the percentage of copper removal from 51% (C i = 0.01 M, P = 13 W m⁻²) to 2% (C i = 0.002 M, P = 2 W m⁻²). Although the final solution may require additional treatment, the adapted TRAB process removed much of the copper while producing electrical power that could be used in later treatment stages. These results show that the adapted TRAB can be a promising technology for removing copper ions and producing electricity by using waste heat as a highly available and free source of energy at many industrial sites.

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1. Introduction

Wastewaters from industries that are involved in mining, electroplating, or smelting operations contain various dissolved metals,
which can be harmful to the environment when released without treatment [1,2]. Various techniques have been used to treat these industrial wastewaters, including chemical precipitation [3], absorption [4,5], biosorption [6–9], photocatalytic removal [10–12], ion exchange [13], membrane separation [14,15], bioelectrochemical systems [16,17], electrodeposition/reduction [18,19], and electrocoagulation (EC) [20–22]. Electrocoagulation is a water treatment process that uses an electric current applied across one or two metal electrodes to remove metals such as copper, nickel, zinc and chromium [23,24]. Electrocoagulation offers many benefits over other technologies, including simple operation, high removal efficiency, and low sludge formation [25–27]. However, a major disadvantage of EC is that it requires substantial electrical power, in proportion to the initial concentration of heavy metals [27]. Therefore, reducing the concentration of heavy metals prior to EC treatment would decrease overall electrical power demands.

A method was recently proposed to produce electricity from low-grade waste heat, called a thermally regenerative ammonia battery (TRAB) [28,29]. In a TRAB, electrical power is obtained from the formation of metal ammine complexes, which are produced by adding ammonia to the anolyte, but not to the catholyte, of a battery consisting of two copper electrodes in a copper-nitrate electrolyte. The added ammonia generates a potential difference between the electrodes according to the reactions [30]:

\[
\begin{align*}
\text{Cu}^{2+}(aq) + 2e^- &\rightarrow \text{Cu}(s) \quad E^0 = +0.34 \text{V} \\
\text{Cu}(s) + 4\text{NH}_3(aq) &\rightarrow \text{Cu(NH}_3)_4^{2+}(aq) + 2e^- \quad E^0 = -0.04 \text{V}
\end{align*}
\]

where \(E^0\) is the standard reduction potential in V vs. the standard hydrogen electrode (SHE). After discharging the cell and generating electrical power, ammonia is separated from the anolyte using conventional distillation with low-grade waste heat, as described in our previous investigations (Fig. S1) [31,32]. The distilled ammonia is then added to the other electrolyte chamber for the next discharge cycle. While discharging the battery results in copper loss from the anode, the electrode can be regenerated when the ammonia is added to the other chamber, where copper will be re-deposited back onto the electrode.

A new approach for copper removal based on the TRAB was examined here to simultaneously remove copper and generate electricity. In this process, which we refer to as a thermally regenerative electrodeposition battery (TREB), solutions containing copper ions is introduced to both chambers, then ammonia is added only to the anolyte, and not the cathode chamber (treatment chamber). During battery discharge, copper ions are reduced to copper metal at the cathode and are removed from solution, accomplishing treatment of the water in the cathode chamber. Once the electrical power is discharged, the treated waste is collected into a low concentration tank either for reuse, or additional treatment in an EC cell to reduce copper concentrations to that allowable for discharge. The electricity generated by the TREB can be used as a power source for the EC cell, making the overall process more economical. Because the anode corrodes occurs during this process, the anolyte will have an increased concentration of copper. However, this high copper ion concentration anolyte can be collected and further used in a typical TRAB to harvest additional electrical energy from waste heat. It is also possible that copper could be extracted from this high concentration stream in the copper electroplating industrial units. The electrodes are alternatively operated as cathode and anodes during the treatment to maintain the copper on the electrodes (Fig. 1).

In this study, we examined the feasibility of removing copper using the TREB. Industrial wastewater from processes such as electroplating can produce copper concentrations ranging from 0.3 M to 0.0001 M [33,34]. Based on this range of copper concentrations, we examined the impact of copper concentrations within this range on copper removal and power generation by adding soluble copper to final concentrations of 0.1 M to 0.002 M. Previous TREB tests examined power generation using copper and nitrate salts, but nitrate would not be feasible for wastewater treatment as high concentrations of nitrate cannot be safely discharged to the environment [35,36]. Therefore, a sulfate salt was used here for all tests as it is a safe ion naturally present in drinking water, it naturally exists at high concentrations in seawater, and because there is no primary drinking water standard for sulfate in the US [37]. To show the utility of the TREB process for treatment of copper solutions, we examined the performance of the TREB in terms of copper removal and power generation.

2. Experimental section

2.1. TREB construction and operation

The TREB was constructed as previously described [28,29]. The battery consisted of a cathode and an anode chamber, each 4 cm long and 3 cm in diameter, separated by an anion exchange membrane (AEM; Sellemion AMV, Asahi glass, Japan) with effective surface area of 7 cm\(^2\). Two 0.8 \(\pm\) 0.05 cm \(\times\) 2 \(\pm\) 0.05 cm pieces of copper mesh (50 \(\times\) 50 mesh; McMaster-Carr, OH) connected by copper wire were used as the electrodes, with each electrode placed 1 cm away from the membrane. In order to monitor the electrode potentials, two Ag/AgCl reference electrodes (+0.211 V vs. SHE; RE-SB; BASI) were inserted 1 cm away from each electrode (2 cm away from the membrane). The catholyte was mixed using a magnetic stirrer (6.4 \(\times\) 15.9 mm; VWR) at 600 rpm. Previous investigations showed that stirring the catholyte improved the power generation, due to better mass transfer of ions to the electrode [28].

The copper solution was prepared by dissolving different concentrations of CuSO\(_4\) (Sigma-Aldrich) in deionized water. The supporting electrolyte, (NH\(_4\))\(_2\)SO\(_4\) (Sigma-Aldrich), was added to increase the solution’s conductivity. Ammonium hydroxide (2 M final concentration; 5 N solution, Sigma-Aldrich) was added to the anolyte to form the copper ammonia complex and create the potential difference between the cathode and anode chambers. The initial anolyte pH was 9.7, consistent with that expected for the addition of a 2 M solution of ammonia. The cell was operated with a fixed external resistance that produced the highest power (based
on polarization data) for a whole batch cycle, which ended when the cell voltage decreased to <10 mV. All experiments were conducted at 30 °C in a constant temperature room.

2.2. Measurements and calculations

In the cathode chamber, the removal of copper occurs due to deposition on the copper electrode. Copper removal (%) was calculated based on the mass change of the electrode during discharge of the TREB as:

$$\text{Removal} = \frac{(m_{t,e} - m_{0,e})}{C_{i}MV_{t}} \times 100$$

where $m_{0,e}$ and $m_{t,e}$ are cathode electrode masses before and after treatment, $C_{i}$ is the initial concentration of Cu(II) in the solution, $M$ is the molecular weight of copper (63.55 g mol⁻¹), and $V_{t}$ is the volume of cathode chamber (28 ± 1 cm³). Coulombic efficiency of the cathode (CCE) was calculated as the ratio between actual produced charge and the theoretical amount of charge based on the mass change of the electrode, as:

$$\text{CCE(%) } = \frac{(m_{t,e} - m_{0,e})}{QM} \times 100$$

where $Q$ is the total charge transferred (C), and $F$ is the Faraday constant [96485 C mol⁻¹]. The mass was measured using an analytical balance with a precision of 0.0001 g. Considering the uncertainties of the volume and analytical balance, the uncertainty of Removal was 2%, while that for CCE were less than 0.01%.

Polarization tests were performed using a potentiosat (model 1470E, Solatron Analytical, Hampshire, England) to measure the cell voltage ($U$), and each electrode potential in a constant temperature room. External resistances were switched every 5 min from 40.9 to 1 Ω [for 0.1 and 0.05 M Cu(II)], 61.2 to 8.2 Ω [for 0.01 M Cu(II)], and 200 to 60 Ω [for 0.002 M Cu(II)] in decreasing order. Both current density ($i = U/R_A$, A m⁻²), and power density ($P = U^2/R_A$, W m⁻²) were normalized to a single electrode projected surface area of 1.6 cm² [38]. Before connecting the external resistance to the cell, the open circuit potential of the cathode was recorded for 5 mins. The potential of the cathode chamber was calculated using the Nernst equation at various initial concentrations of Cu(II) in the catholyte (Equation S1; Fig. S2). To calculate the theoretical potential by the Nernst equation, the Cu(II) ion activities were estimated using the OLI Studio software (Cedar Knolls, NJ).

The total charge transferred over the entire copper removal process was calculated by integrating the current–time profile $Q = \int i t$, where $Q$ is the total charge (C), $i$ the current (A), and $t$ time (s). The energy density, normalized to the total electrolyte volume ($E$, Wh m⁻³), was calculated as $E = \int U t h/V$, where $U$ is the voltage (V), $i$ the current (A), $t$ the cycle time (h), and $V$ the total volume (2 x 28 mL). Based on possible variations in the volume of the solutions, the experimental errors for the calculated energy densities were less than 4%.

After discharging the TREB, the battery was recharged by separating ammonia from the anolyte effluent and re-dissolving it in the catholyte. In practice, low-grade waste heat (<100 °C) could be used for this process, for example by using a distillation column with a reboiler and condenser temperature of 70.4 and 43.3 °C, respectively. Previous simulation results indicated that 97% of ammonia could be recovered during this separation [28,29].

**Fig. 2.** Copper removal from cathode chamber (striped columns), and cathodic coulombic efficacy (filled columns) of TREBs with various copper concentrations, and 1 M (NH₄)₂SO₄ supporting electrolyte. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.).

3. Results and discussion

3.1. Copper removal at different concentration of copper in the solution

The copper removal efficiency (%) strongly depended on the initial Cu(II) concentration. Copper removal was >50% when initial Cu(II) concentrations were ≥0.01 M using a 1 M (NH₄)₂SO₄ electrolyte for both the anolyte and catholyte. The highest removal was achieved at a concentration of 0.05 M (77%). Treatment of solutions containing 0.002 M or less Cu(II) had <2% copper removal, resulting in a very low rate of copper deposition and copper recovery under these conditions (Fig. 2).

Low recoveries of copper from the catholyte at relatively low Cu(II) concentrations were due to unwanted side reactions. The cathodic coulombic efficiency (CCE), which is the ratio of the produced current to the theoretical amount of current based on the change in the mass of the electrode, can be used to evaluate the relative importance of the electrodeposition reaction for copper ions compared to other possible side reactions in the cathode chamber. A CCE <100% (range of 2 to 89%) was obtained in all experiments, confirming side reactions occurred in the catholyte (Fig. 2). We expected that the main side reactions involved the formation of a Cu(OH)₂ precipitate, and the formation of copper ammine complexes [Cu(NH₃)₂]²⁺, Cu(NH₃)₃²⁺, and Cu(NH₃)₄²⁺ in the cathode chamber. These reactions and their relevant equilibrium constants are [39]:

$$\text{Cu(OH)₂(s)} = \text{Cu}^{2+} + 2\text{OH}^- , \quad \text{pK}_{sp} = 19.36$$  \hspace{1cm} (5)

$$\text{NH}_4^+ = \text{NH}_3 + \text{H}^+ , \quad \text{pK}_a = 9.25$$  \hspace{1cm} (6)

$$\text{Cu}^{2+} + \text{NH}_3 = [\text{Cu(NH}_3)_2]^{2+} , \quad \text{pK}_1 = -4.25$$  \hspace{1cm} (7)

$$[\text{Cu(NH}_3]_2^{2+} + \text{NH}_3 = [\text{Cu(NH}_3)_3]^{2+} , \quad \text{pK}_2 = -3.61$$  \hspace{1cm} (8)

$$[\text{Cu(NH}_3]_3^{2+} + \text{NH}_3 = [\text{Cu(NH}_3)_4]^{2+} , \quad \text{pK}_3 = -2.98$$  \hspace{1cm} (9)

$$[\text{Cu(NH}_3]_4^{2+} + \text{NH}_3 = [\text{Cu(NH}_3)_5]^{2+} , \quad \text{pK}_4 = -2.24$$  \hspace{1cm} (10)

When the solution had a low copper ion concentration of 0.002 M, the main side reaction would likely be formation of a Cu(OH)₂ precipitate, primarily due to the higher initial pH of the catholyte (Fig. S3). During experiments at this copper concentration, we observed formation of precipitates. Based on our calculations, the observed pale blue precipitates were likely Cu(OH)₂ (Table S1). A higher pH should also result in a shift in the NH₄⁺/NH₃ acid/base reactions towards NH₃ formation side, result-
ing in greater formation of different copper ammine complexes, as shown by equations 6–10. At higher copper concentrations (>0.01 M), however, copper ions are less involved in the side reactions, mainly due to the lower pH of catholyte. Therefore, the loss of free Cu(II) ions to ammonia complexes, as well as formation of a Cu(OH)₂ precipitate rather than current generation, resulted in limited copper ion removal from the lower concentration solutions (<0.01 M).

The highest CCE was obtained at a copper concentration of 0.05 M (~90%), confirming most of the current was consumed by Cu(II) ion deposition onto the cathode electrode. The highest CCE for a copper concentration of 0.05 M was the main reason for the highest percentage of copper removal at this copper concentration.

3.2. Power production with different concentration of copper ions

Increasing the initial Cu(II) concentration from 0.002 M to 0.1 M improved the power production from 2 W m⁻²-electrode area to 37 W m⁻²-electrode area (Fig. 3). This increase in power was due to more positive cathodic potentials during battery discharge. Power generation was examined over a complete discharge cycle (defined as a final cell voltage <10 mV), at the external resistance that produced the maximum power density (5.6 Ω for 0.1 M and 0.05 M, 20.6 Ω for 0.01 M, and 99.6 Ω for 0.002 M; Fig. 3). The reason the optimum external resistance varied was likely due to the different copper ion redox activity (i.e., electron transfer ability) at different concentrations of copper [40]. The discharge time was reduced by decreasing the initial Cu(II) concentration in the solution, mainly due to the Cu(II) side reactions as well as lower availability of copper ions in the catholyte (Fig. 4). When the initial concentration of copper ions was decreased, the solution pH increased from 3.8 for 0.1 M Cu(II) to 4.8 for 0.002 M Cu(II), which would result in an increased formation of Cu(OH)₂ precipitates during treatment due to the higher pH, consistent with visual observations of formation of a light blue solid in the reactor with these changes [28]. The differences in the power production curves were mostly caused by the changes in cathode potentials. The cathode potential initially increased and reached a maximum, where power density reached a maximum as well, and then it started to become more negative. In contrast, there was an insignificant change in the anode potential over time (Fig. 4b).

3.3. Total charge transfers and energy densities

Decreasing the Cu(II) initial concentration from 0.1 M to 0.002 M reduced the total charge transfer, based on coulombic recoveries decreasing from 400 coulombs (C) to 5 C (Fig. 5). This decrease was due to lower copper deposition rates at lower initial Cu(II) concentrations. The energy densities produced by the TREBs during the treatment increased with the Cu(II) initial concentration, similar to the increases in maximum power densities obtained in polarization tests. The highest energy density produced by the treatment system was 280 Wh m⁻³ at an initial Cu(II) concentration of 0.1 M, which is ~60% of the energy produced in a previous TRAB with nitrate salts in the electrolyte, at the same copper concentration.

Fig. 3. Power densities produced by copper removal using TREBs operated with various concentrations of copper, and 1 M (NH₄)₂SO₄ as the supporting electrolyte in both electrolytes, and 2 M ammonia in the anolyte.

Fig. 4. Whole batch cycle performance of copper removal from solutions with different copper concentrations: (a) power densities; (b) cathode (open symbols) and anode (filled symbols) potentials. The initial electrolyte contained a different concentration of Cu(II), 1 M (NH₄)₂SO₄, and additional 2 M NH₃ in the anolyte, and the cell was operated at its maximum power. The arrow shows when the anode electrode was replaced.

Fig. 5. Total charge and corresponding discharge energy densities produced by TREBs operated to remove copper from the catholyte, using 1 M (NH₄)₂SO₄ (supporting electrolyte) in both chambers, and additional 2 M NH₃ in the anolyte. The cell was operated for maximum power production.
solution [28], rather than sulfate salts used here (Fig. 5). The higher energy densities obtained here for higher copper concentrations were due to the higher discharge voltages and the total charge transferred ($W = QU$, where $W$ is energy, $Q$ the charge, and $U$ the voltage).

4. Reuse of the high concentration effluent

Another possible concern is the increased copper concentration in the anolyte. While the TREP process can produce an effluent catholyte with a reduced copper concentration, the anolyte copper concentration will increase over time. To minimize the volume of the anolyte used, the solution in the anode compartment could remain in the chamber for a several batches of catholyte (Fig. S4). To examine the reuse of the anolyte, a solution with a Cu(II) concentration of 0.05 M was examined in the TREP process, with the anolyte used for four batches of catholyte. For the third batch of catholyte, there was 68% copper removal and a maximum power density of 26 W m$^{-2}$, similar to that obtained in the first batch cycle ($Removal = 77\%$; $P = 33$ W m$^{-2}$). However, for the fourth batch cycle, both copper removal and power substantially decreased (23% removal, 12 W m$^{-2}$) (Fig. 6). After each batch, the concentration of copper ion in the anolyte increased, resulting in the enhancement of copper ammonia complex concentration (Cu(II)\(\text{NH}_3\)\(^2\)) and a decrease in the concentration of ammonia (Eq. (2)). The benefit of a higher copper complex concentration in the anolyte with each cycle, which could have increased the cell potential, was offset in the fourth batch cycle by a reduction in the ammonia concentration in the anolyte. As a result, there was a decrease in the power (Equation S2), and less copper removal.

5. Outlook

The TREP process was shown to simultaneously remove copper and generates power, but the copper was not reduced to levels sufficient for water discharge (50–80 μM) [41]. However, the copper remaining in the water following TREP treatment could be reduced to levels suitable for discharge using a standard electrocoagulation treatment system. The advantages of using the TREP process to first treat a wastewater is increased recovery of copper, the use of waste heat which recovers copper without the use of electrical power, electrical power generation, and less copper in a waste sludge. In demonstrating the concept of copper removal using TREP, only simple solutions were used here as they contained only a copper sulfate electrolyte. Actual industrial wastewaters will contain other constituents, notably chloride ions, which could impact treatment efficiency. The effect of these other anions and cations on the process will require further study.

6. Conclusions

The TREP process was shown to be a potential method of treatment, with >30% copper ion removal from solutions containing copper ions (>0.01 M), and production of up to 25 W m$^{-2}$ based on charging the cell with ammonia solutions prepared using low-grade waste heat. In addition, the produced high concentration effluent can be used for a several number of batches, minimizing the volume of anolyte used. Overall, using a low-grade waste-to-electricity TREP system for removal of copper ions from solution represents a promising approach for decreasing the energy requirements of current treatment plants and improved copper recovery.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jhazmat.2016.10.022.

References


Fig. 6. Copper removal and power density of four batch cycles of TREP fed with solution with a copper ion concentration of 0.05 M. The solution in the anode compartment remained in the chamber, while the catholyte were replaced by a new solution after each batch.