Palladium Recovery through Membrane Capacitive Deionization from Metal Plating Wastewater

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Supporting Information

ABSTRACT: The potential application of membrane capacitive deionization (MCDI) for recovery of palladium (Pd) ions from catalyst solution wastewater generated from the plating industry was investigated in this study. Several major issues were explored in this work to verify the suitability of MCDI for Pd recovery from a practical perspective: adsorption and desorption efficiencies, desorption mechanisms into high concentration of Pd concentrate, and its sustainability in long-term operation. The lab-scale MCDI operation achieved satisfactory and highly competitive Pd removal (99.07–99.94% removal with 1.42–1.52 of Pd selectivity over ammonium ions) showing that Pd can be effectively collected from plating industry wastewater. A high concentration of Pd concentrate (64.77 and 919.44 mg/L of Pd from the 10 and 100 mg/L Pd containing catalyst solution, respectively) was obtained through successive five operation cycles of adsorption/desorption phases. However, it is significant to note that the desorption efficiency was inversely proportional to the concentration of Pd concentrate which is likely due to the Pd ions discharged from carbon electrode toward Pd solution against the enhanced concentration gradient. The long-term operation results suggest that scaling could reduce the MCDI efficiency during Pd recovery (0.17% decrease in Pd removal for every cycle on average) and hence may require an adequate electrode cleaning regime.

KEYWORDS: Membrane capacitive deionization (MCDI), Electrosorption, Palladium concentration, Ion desorption, Wastewater from plating industry

INTRODUCTION

As a result of large-scale exploitation of minerals in contemporary industry and increasing concerns over environmental issues caused by wastes in electrical and electronic equipment (WEEE), reusing and recycling precious metals from secondary resources (i.e., industrial wastewater) are deemed to be important for sustainable utilization.1,2 With the significant increase in amount of WEEE and the ever-growing concerns of contamination by toxic and precious metal sources, a “WEEE directive” has been issued by The European Union, forcing EEE industries to take responsibility for the collection of, recovery, and recycling of WEEE.3 Printed circuit boards (PCBs), which is a key part of electronic devices, have been gaining interest as one of the largest sources of heavy metals.
In PCB manufacturing processes, palladium (Pd) complexes are widely used as catalysts to activate electro- and electroless-plating processes, which are regarded as very important steps for fabricating fine products. Electrowinning, the reclamation of used Pd solution from the plating process by electrodeposition of metals from wastewater, is known as the most effective and commonly used metal recovery system.\(^5\)\(^6\) However, recovering the wasted Pd by electrowinning is not feasible due to its limited recovery and energy-efficiency especially at low concentration, and a significant amount of low Pd concentration wastes is disposed by dilution with other wastewater sources to satisfy the regulations of industrial wastewater discharge. Several new technologies, such as precipitation, solvent extraction, conjugate absorbers, and ion exchange resins, have been studied as an alternative to the electrowinning for high Pd recovery with better efficiency.\(^6\)\(^−\)\(^9\) Concentrating the Pd-containing solution prior to electrowinning with an energy-efficient technology is expected to be one of the most possible options to improve the energy efficiency and total Pd recovery rate. Therefore, concentrating the Pd containing wastewater from the plating industry by means of capacitive deionization (CDI) with low energy use could be done to facilitate feasible and sustainable Pd recovery.

CDI is a technology that has been mainly applied for desalination to produce pure water through the extraction of charged species from water by applying an electrical potential difference over two porous electrodes. It has been one of the most alternative and feasible technology in recent years for sustainable fresh water production from unconventional water resources.\(^10\)\(^−\)\(^11\) This electrosorption process has especially shown to have better energy efficiency than reverse osmosis (RO) for deionization at low salt concentration levels in water (i.e., salt concentration $\leq 30$ mM), such as municipal wastewater and brackish water.\(^12\) Previous research on the development of advanced materials of electrodes, effective process designs, and energy recovery have drastically improved this process, making this technology feasible for potable water production.\(^12\)\(^−\)\(^14\)

Aside from the use of CDI for water production, this technology has also been further studied for other applications including recovery of resources from the water.\(^15\)\(^−\)\(^28\) Since there is a rising global demand for vastly depleted natural resources and nutrients, reclaiming valuable resources such as nitrogen, phosphorus, and mineral resources from municipal wastewaters and saline water sources has emerged as a great environmental solution.\(^29\)\(^−\)\(^31\) Since CDI is based on the principle of ions and charge separation, it has shown to be potentially effective in the recovery of charged resources by highly removing/concentrating or selectively collecting present in minute amounts in low salinity water.

There are several studies that investigated the suitability of CDI for the recovery of nutrients and minerals. Especially, most CDI studies have noted on the recovery of nitrogen\(^35\)\(^−\)\(^38\) and phosphorus\(^39\)\(^−\)\(^41\) from municipal wastewaters as one of the most possible applications. Works on copper removal revealed that CDI can be used to extract Cu\(^{II}\) from wastewater with high selectivity over NaCl and natural organic matter (NOM) in a competitive environment.\(^22\)\(^23\) Studies on lithium recovery using CDI show excellent removal performance with faster adsorption/desorption than conventional lithium recovery process, and durability over several adsorption and desorption cycles.\(^24\)\(^−\)\(^26\) A binder-free carbon nanotube (CNT) electrode was found to provide a much increased electrochemical surface area, leading to enhanced removal of Cr\(^{VI}\) and Cr\(^{III}\) ions\(^27\)

whereas Pb\(^{II}\) ions were more easily captured by an air-plasma treated CNT electrode in another study.\(^28\)

The main objective of this work is to investigate the potential application of CDI for the recovery of Pd resources from the plating industry wastewater. The performances of the CDI for Pd recovery under different feed conditions and the influence of various operating parameters were studied using lab-scale membrane CDI (MCDI) unit, which is currently one of the most advanced systems in this field.\(^32\) The issues of the MCDI performances including the adsorption and desorption efficiencies in long-term operations and its improvement to attain higher concentration of Pd concentrate were discussed. Following up the long-term runs, the sustainability of CDI as a recovery process has been discussed through investigating the scaling behavior caused by Pd. Lastly, we introduced a new process for the efficient Pd recovery from plating industry wastewater integrating CDI stage and also suggested improvements of CDI for its enhanced sustainability and feasibility through simplifying the Pd recovery process.

### EXPERIMENTAL SECTION

#### Model Palladium and Palladium Catalyst Solution for Electroless Plating.

Tetraamminepalladium(II) sulphate (Alfa Aesar, MA) was chosen as representative Pd species, since it can be found in Pd catalyst solutions. Three different concentrations of Pd in model solution were used for the MCDI tests: 1, 10, and 100 mg/L, which are within the range of actual Pd catalyst wastewater concentration observed in the PCB manufacturing process. A bulk palladium catalyst solution for electroless plating was also provided by MSC Co., Ltd. (Republic of Korea) to assess the performance of MCDI as a real palladium recovery process. The major compositions of the Pd catalyst solution under different concentrations of Pd species are listed in Table 1. As oxidation of carbon electrodes caused by dissolved oxygen in feedwater is regarded to be insignificant in the MCDI system, deaeration of the Pd feed solution was not performed in this study.\(^33\)

#### Carbon Electrodes and Ion Exchange Membranes.

Two porous carbon electrodes manufactured by Siontech Co. (Republic of Korea) were used. The electrodes consisted of a graphite sheet body coated with activated carbon P-60 (Kuraray Chemical Co., Japan) and PVDF (Inner Mongolia 3F-Wanhao Fluorine Chemical Co., Ltd., China) blended carbon slurry. The total carbon mass on each 10 $\times$ 10 cm$^2$ sized electrode was reported to be 0.8 g. Commercial anion and cation exchange membranes (Neosepta AFN and CMB, respectively) were obtained from Astom Corporation (Japan).

#### Bench-Scale MCDI Setup.

The MCDI tests were carried out in a bench-scale flow-through system as depicted in our previous study.\(^34\) The dimensions of the rectangular feed channel of the test cell were 10 cm in width and 10 cm in length, providing an effective ion adsorption/desorption area of 100 cm$^2$. There were two carbon electrodes placed within the test cell, whereas the anion and cation exchange membranes were inserted between those two electrodes. A nonelectrically conductive nylon spacer was placed between the two ion exchange membranes to ensure the water flow and prevent short circuit. A 1 cm diameter sized hole was punched in the center of carbon electrodes, ion

<table>
<thead>
<tr>
<th>parameter</th>
<th>1 mg/L of Pd</th>
<th>10 mg/L of Pd</th>
<th>100 mg/L of Pd</th>
</tr>
</thead>
<tbody>
<tr>
<td>total dissolved solids (μS/cm)</td>
<td>12.23</td>
<td>46.4</td>
<td>337</td>
</tr>
<tr>
<td>SO$_4^{2−}$ (mg/L)</td>
<td>0.90</td>
<td>9.03</td>
<td>90.27</td>
</tr>
<tr>
<td>NH$_4^+$ (mg/L)</td>
<td>4.36</td>
<td>13.93</td>
<td>37.04</td>
</tr>
<tr>
<td>pH</td>
<td>8.22</td>
<td>9.63</td>
<td>10.32</td>
</tr>
</tbody>
</table>

### Table 1. Major Characteristics of 1, 10, and 100 mg/L of Diluted Pd Catalyst Solutions
exchange membranes, and a nylon spacer to enable the solution to be completely in contact with the electrodes and provide flow within the entire system cell.\textsuperscript{35}

A 100 mL portion of feed solution was recirculated throughout the MCDI system with the flow rate of 30 mL/min by a peristaltic pump (Cole-Palmer, IL). The electrical potential supplied through the MCDI test unit was adjusted at a constant level by a potentiostat, WPG-100 (WonATech Co., Republic of Korea). The MCDI system was stabilized at the desired voltage after the feedwater was circulated by repeating five cycles of 2 min for ion adsorption followed by 2 min of ion desorption. All the tests were carried out in duplicate.

**Multiple Cycles Operation for Concentration of Pd.** The five or ten cycles operation procedure of MCDI is depicted in Figure 1. A 100 mL portion of feed solution containing 100 mg/L Pd circulated through the MCDI system for 8 min at 0.9 V adsorption condition. To desorb the Pd ions attached onto the carbon electrode, 50 mL of fresh DI water was made to circulate through the system at the same conditions as adsorption by applying reverse potential. The desorbed Pd ions from the MCDI system were collected onto the concentrate solution after the primary desorption process for all cycles. To completely remove residual Pd ions remaining on the carbon electrode, an additional desorption step (secondary desorption) in the MCDI system was carried out with newly replaced 50 mL fresh DI water at the same time and electric potential conditions by applying reverse potential. The water containing the residual Pd ions was then discarded. These adsorption, desorption, and secondary desorption processes were performed for five or ten cycles, with 100 mL of fresh feed solution and 50 mL of Pd concentrated water for each test. Complete desorption of ions from the electrodes was ensured by employing a further desorption step for 20 min at −1.2 V after the secondary desorption in each cycle and thereby no discharge of ions during this step was detected by a conductivity meter confirming that there are no remaining ions that could be released by electrical force. The mass balance on the adsorbed and desorbed Pd ions was also calculated to make sure that ions were completely released at the end of each cycle.

After the long-term operation, anode, cathode, and anion and cation exchange membranes in the MCDI system were disassembled and prerinsed with DI, followed by desiccation for over 24 h to ensure complete drying of the samples. Then the surface of samples was analyzed by scanning electron microscopy (SEM) and energy-dispersive spectroscopy (EDS) in order to investigate the effect of long-term operation on MCDI performances.

**Measurement of Water Quality.** The Pd concentrations of feed, permeate and concentrate waters were analyzed by inductively coupled plasma-mass spectrometer (ICP-MS), NexION 300D ICP-MS (PerkinElmer Co., MA). The water samples were diluted 100 times for accurate quantification of palladium ion. The calibration of ICP-MS was performed before measurement by external calibration with deionized water and four standard palladium solutions (0.625, 1.25, 2.5, and 5 mg/L).

**RESULTS AND DISCUSSION**

**Basic Performance of MCDI for Pd Collection.** To assess the capacity of MCDI in Pd recovery and optimize operating conditions, the Pd removal in this system was tested under varying operating conditions such as applied voltage, duration, and Pd concentrations with or without other ions in the adsorption stage.

**Pd Removal in MCDI under Different Operating Conditions.** The basic removal of Pd by MCDI was first evaluated using the synthetic Pd solution to observe the effect of voltage and time on Pd removal only, without the interference of other coexisting ions. Figure 2 shows the Pd removals from the 100 mg/L Pd solution at four different potentials during adsorption, ranging from 0.3 to 1.2 V applied voltage. As shown in the results, the MCDI process exhibited a very high efficiency for Pd removal throughout all the applied voltages. The removal efficiency and specific removal was 98.38% and 11.99 mg/g, respectively, even at a small applied voltage of 0.3 V, reaching 99.84% (12.18 mg/g specific removal) at 1.2 V when the adsorption was operated for 8 min. The highly efficient adsorption of Pd ions on carbon electrode might be driven by both electroosorptive and physical adsorption, such as the sorption behavior of other heavy metal ions in MCDI.\textsuperscript{36} A further adsorption test was carried out employing 0 V for 8 min using the synthetic Pd solution containing 100 mg/L of Pd to clarify the physisorption behavior during Pd removal in MCDI (Figure S1). A significant amount of Pd removal attained by physisorption (24.88%) implies that both electroosorption and physisorption play important roles adsorbing the Pd species during Pd removal in MCDI, leading to high Pd removal. However, this unanticipated behavior brings concern of deterioration of performance of MCDI. The hampered
or feed electrical conductivity. However, when the feed TDS linearly increases with the feed concentrations limits the movement and electrosorption of Pd ions but the electrical conductivity and higher resistance because of which due to the fact that, lower TDS or Pd concentration has poor solutions containing 1 and 2 mg/L of Pd, respectively. This is solution and 98.99 and 99.59% from catalyst solution, for feed Meanwhile, the Pd ions indicated slightly lower removal at 99.74% for Pd concentration between 5 and 100 mg/L. 

The Pd removal at different operation times for 0.9 V applied voltage in Figure 2 shows that the Pd ions were highly extracted by the MCDI system achieving 96.76% of Pd removal (11.80 mg/g specific Pd removal) within just 2 min and above 99.06% (12.08 mg/g) in about 4 min. To sum up, the operation of MCDI at 0.9 V for 8 min is likely to be the optimum condition for Pd recovery. Similar observations were made in the previous work optimizing the salt adsorption rate in MCDI, which showed similar trends with different operating times and voltages.

**Effect of Feedwater Concentration and Composition on Pd Removal.** Figure 3a presents the removal of Pd ions from the synthetic Pd and Pd catalyst solutions as a function of Pd concentration in feedwater. The Pd removals of both the synthetic and catalyst solutions were consistently more than 99.74% for Pd concentration between 5 and 100 mg/L. Meanwhile, the Pd ions indicated slightly lower removal at low feed concentration: 98.65 and 99.49% from synthetic solution and 98.99 and 99.59% from catalyst solution, for feed solutions containing 1 and 2 mg/L of Pd, respectively. This is due to the fact that, lower TDS or Pd concentration has poor electrical conductivity and higher resistance because of which limits the movement and electrosorption of Pd ions but the electrosorption linearly increases with the feed concentrations or feed electrical conductivity. However, when the feed TDS increases more than 150 mg/L of Pd, the sorption efficiency stabilizes or gradually decreases. 

The removal of Pd in the catalyst solution was slightly lower than that in the synthetic Pd solution, within the range of 5 to 150 mg/L of Pd in feed solution. The lower Pd removal could be explained due to competitive adsorption from the several other ions present in the plating feed solution, such as ammonium. The ammonium and other minor ions present in the plating solution would competitively occupy the pore space within the carbon electrode, resulting in the decreased rate of Pd removal. However, it should be noted that the Pd ions were still highly removed by MCDI even in the presence of other ions in the catalyst solution. This is likely due to the selectivity preference for Pd ions in feedwater during MCDI, as shown in Figure 3b. The ion selectivity during Pd recovery via MCDI from Pd catalyst wastewater was ranged from 1.42 to 1.52 as the Pd concentration in the feed solution increased from 1 to 100 mg/L. In general the ion selectivity during the electrosorption process is strongly dependent on the initial concentration, ionic charge valency, and hydrated radius of ions. First, the ions with high concentration is preferentially adsorbed by the carbon electrode. More specifically, cations of higher valence are energetically favorable to screen the surface charge of electrodes. 

On account of the governing number of Pd ions contained in the catalyst solution and those higher charge (divalent) than ammonium and other ions, Pd was the most preferred species to be collected by the carbon electrode. And the selectivity was shown to be increased as the ratio of Pd over NH4 increased in higher concentration of catalyst solution.

Furthermore, Pd ions were shown to be more adsorbed in the catalyst solution under low concentrations such as 1 and 2 mg/L of Pd. This is because the coexisting ammonium ions increases the total ion mobility toward the carbon surface, leading to higher adsorption capacity under low TDS condition. Since the addition of other ions in Pd catalyst solution offers slightly higher TDS in the low concentration feedwater, the adsorption rate reaches higher than that of the synthetic Pd solution. This clearly shows that the coexistence of other ions such as ammonium in the catalyst solution could rather support the Pd removal performance of MCDI than competitively taking place of the pore and surface of electrode.

Therefore, it can be concluded that Pd could be sufficiently extracted through MCDI, and an outstanding Pd removal could be accomplished even though other co-ions existing in the Pd catalyst solution affect the efficiency of the process by competitive adsorption behavior. It is generally known the removal efficiencies of ions involved in the feed solution can be influenced by the initial pH of feedwater due to adsorption of hydrogen and hydroxide ions on the surface of electrodes. In our study, however, pH of the feed solution did not play a significant role in the Pd removal though the initial pHs of catalyst solutions involving 10 and 100 mg/L of Pd were 9.63 and 10.32, respectively.

**Five-Cycle Operation of MCDI to Obtain High Pd Concentration.** Enhancing Pd Concentration through Multiple MCDI Adsorption and Desorption Cycles. The
adsorption/desorption in MCDI was repeated for five cycles to capture Pd ions into the concentrate using a mixture of desorbed solution and original Pd feed solution. Figure 4 shows that the Pd concentration increases linearly in the concentrate solution after each cycle for initial feed Pd catalyst solutions containing 1, 10, and 100 mg/L of Pd in Figures 4a, b, and c, respectively. As shown in Figure 4a, the concentration of collected Pd ions in the concentrate after the fifth MCDI cycle reached 925.48 mg/L, suggesting that MCDI could successfully recover and concentrate Pd ions. Theoretically, the Pd concentration in the concentrate was supposed to be 997.76 mg/L assuming a fixed Pd removal of 99.78% and the adsorbed ions on the carbon electrodes were completely discharged into the concentrate during the first desorption phase. Perhaps this slightly lower amount of experimentally collected Pd ions in the concentrate is likely due to experimental errors.

Figure 4b and c also show the potential of MDCI in concentrating Pd ions from a feed with very low Pd concentrations to achieve higher Pd concentration such as in Figure 4a. In fact a somewhat logarithmic increase was observed with repeated cycles. For example, when the MCDI was operated with 10 mg/L of Pd catalyst solution, 19.36 mg/L of Pd was desorbed and collected in the first cycle, while this reduced to only 4.27 mg/L of Pd in the fifth cycle. Likewise for 1 mg/L of Pd catalyst feed solution, 1.91 mg/L of Pd was desorbed in the first cycle against 0.40 mg/L in the fifth cycle. Assuming a stable adsorption and full regeneration in the desorption step throughout the cycles, the Pd concentration in the concentrate was expected to be 99.92 and 9.90 mg/L after the fifth cycle for 10 and 1 mg/L Pd catalyst feed solutions, respectively. The apparent decrease in the desorption of Pd ions from the electrodes in the successive MCDI cycles shows that the desorption efficiency was deteriorated, and this MCDI process would hardly concentrate the Pd species in wastewater under a very low concentration in feed solution such as 1 mg/L, resulting in unfeasible Pd recovery.

Effect of Concentration of Concentrate Solution on Pd Desorption Efficiency in MCDI. After every adsorption/desorption cycle at −0.9 V for 8 min, a secondary desorption step was further conducted and ions were released in a newly replaced 50 mL of DI water in each cycle at the same operating condition to unload any residual ionic species that probably still remained on the electrode. Figure 5 presents the concentration of further desorbed Pd into the DI water in each cycle termed in this study as "residual water". The number of additionally discharged Pd ions in the secondary desorption increased with observed in Figure 4a. In fact a somewhat logarithmic increase was observed with repeated cycles. For example, when the MCDI was operated with 10 mg/L of Pd catalyst solution, 19.36 mg/L of Pd was desorbed and collected in the first cycle, while this reduced to only 4.27 mg/L of Pd in the fifth cycle. Likewise for 1 mg/L of Pd catalyst feed solution, 1.91 mg/L of Pd was desorbed in the first cycle against 0.40 mg/L in the fifth cycle. Assuming a stable adsorption and full regeneration in the desorption step throughout the cycles, the Pd concentration in the concentrate was expected to be 99.92 and 9.90 mg/L after the fifth cycle for 10 and 1 mg/L Pd catalyst feed solutions, respectively. The apparent decrease in the desorption of Pd ions from the electrodes in the successive MCDI cycles shows that the desorption efficiency was deteriorated, and this MCDI process would hardly concentrate the Pd species in wastewater under a very low concentration in feed solution such as 1 mg/L, resulting in unfeasible Pd recovery.

Figure 4. Concentration of Pd in concentrate solution after desorption of ions from the carbon electrodes. After adsorption of ions from new catalyst solutions containing (a) 100, (b) 10, and (c) 1 mg/L of Pd, the adsorbed ions on the electrode surface were discharged in the concentrate water for 8 min at 0.9 V. A new feed solution was then replaced to initiate another Pd recovery cycle, whereas the concentrate solution kept being used over every cycle.
the number of MCDI cycles, clearly indicating that the Pd ions were not completely freed from the carbon electrode during the primary desorption stage, whereas this system showed a fairly stable adsorption performance over every cycle. There are several possibilities that caused degraded desorption performance with the repeated cycles of MCDI, such as electro-oxidation, physical adsorption of ions, and scale formation.36,42,43 However, those reasons seem not to be the main cause of the significantly reduced desorption efficiency with the repeated cycles in this study, whereas the adsorption capacity of the carbon electrode was quite stable. The desorption efficiency in the primary desorption step in each cycle was notably decreased as the concentration of the Pd concentrate increased with repeated cycles. Most of the unreleased ions were quickly released in DI water in each secondary desorption step implying that this incomplete discharge of the ions in the first desorption phase is presumably owing to the effect of enhanced ionic concentration of the concentrate solution as explained by the

Figure 5. Concentration of Pd in residual solution after secondary desorption in each cycle. After adsorption/desorption using new catalyst solutions containing (a) 100, (b) 10, and (c) 1 mg/L of Pd, the remaining ions on the electrode surface were further released in new DI water (residual water) for 8 min at 0.9 V. The new feed solution was then replaced to initiate another Pd recovery cycle, whereas the concentrate water was used over every cycle. The new DI water was then replaced to measure the concentration of residual water after each cycle.

Figure 6. Enhanced concentration of Pd concentrate and the corresponding concentration of Pd in residual solutions after secondary desorption with increased (a) desorption time and (b) applied potential. The secondary desorption to release residual Pd ions from the carbon electrode was performed at 0.9 V and 8 min.
Gouy–Chapman Theory. Theoretically the electrosorptive capacity of the carbon electrodes in CDI is significantly affected by the electrical double-layer (EDL) capacity. Based on the Gouy–Chapman Theory, the electrolyte concentration in the aqueous solution is one of the major factors determining the EDL capacity on the carbon electrode surface in CDI.

Inversely, the increasing electrolyte concentration of the concentrate solution after each successive cycle leads to a lower or reverse ion concentration gradient between the carbon electrode and water media, disturbing the ion desorption from the porous carbon electrode surface. Since the concentration of concentrate solution increased as the operation cycle was repeated, the ions captured from the carbon electrode would be hardly released into the concentrate during the desorption stage, leading the Pd ions adsorbed on the carbon surface to be remained. The concentrate solution approaches saturation, and the lower or reverse concentration gradient is detrimental to desorption. For both desorption and adsorption, a higher concentration gradient is preferred to act as the driving force to overcome resistance to mass transfer of metal ions between the electrode and the solution. This incomplete desorption behavior could also be found in a previous study which investigated the effect of the concentration of concentrate on desorption performance of CDI.

Enhancing Desorption Efficiency through Increasing Desorption Time or Applied Potential for Higher Pd Concentration. As shown in the previous section, the adsorbed Pd ions were not completely released during each primary desorption stage as the Pd concentrations increased in the successive MCDI cycles. This section investigates how two main operating parameters—desorption time and applied voltage—affect the concentration rate of Pd ions in the concentrate water (Figure 6). The results show the final Pd concentration in the concentrate after the fifth cycle using an original catalyst solution of 100 mg/L of Pd where adsorption was performed at 0.9 V for 8 min and desorption was conducted with different potentials (−0.8 to −1.2 V) and times (6–12 min). Figure 6a shows that the concentration of Pd in the concentrate increased from 918.2 mg/L at 8 min desorption time to just 920.07 mg/L at 12 min. This indicates increasing desorption time longer than 8 min insignificantly enhances the concentration of Pd. However, a slight increase of Pd concentration in concentrate water was observed when a higher potential is used as shown in Figure 6b where the Pd concentration increased from 919.44 at 0.9 V to 929.75 mg/L at −1.2 V. The desorption process generally proceeds rapidly, and the majority of the adsorbed ions usually should be desorbed within about 2 min of applying a reverse potential, and thus, the increased desorption time significantly improved the efficiency of Pd collection in the concentrate. However, elevating the applied potential raised the desorption current, promoting better desorption and resulting in enhanced Pd concentration in the concentrate solution.

Deterioration of Pd Adsorption Performance under Longer Operation Cycles. During the MCDI operation with five cycles, it was shown earlier that the adsorbed Pd ions were not completely released from the carbon electrode in the primary desorption phase and additional desorption steps were employed in each cycle during the five cycle MCDI operation. Therefore the change of adsorptive capacity of the carbon electrode was investigated in terms of much longer cycles of ten cycles in total. The longer MCDI operation was carried out with adsorption at 0.9 V for 8 min using the initial feed of 100 mg/L Pd solution, whereas the primary desorption process was performed at −0.9 V for 8 min followed by a further desorption phase at −1.2 V for 20 min. No discharge of ions after those two desorption phases were confirmed in each cycle.

Figure 7 exhibits the Pd removal in each MCDI cycle. The Pd removal gradually decreased from 99.78% (12.17 mg/g) to 98.04% (11.96 mg/g) in the 10th cycle. Although this is a marginal decrease of only 1.74% in 10 cycles and does not appear to be significant, it should be a concern for the long-term operation of MCDI for Pd recovery as this indicates an average 0.17% decrease in the removal for every cycle. If this decreasing trend shown in Figure 7 is to continue, it seems likely that the removal in MCDI will reach zero after 555 cycles of operations, which is not at all desirable. This decrease in Pd removal is likely attributed to physisorption of Pd ions within the porous structure of carbon electrode instead of electrosorption. In common with the adsorption behavior of other metal ions such as Cd, Pb, and Cr on activated carbon, the adsorption of Pd ions on carbon electrode surface would be governed by a complicated adsorption process involving complex formation and surface adsorption mechanisms. The Pd ions in the aqueous solution migrate into the pores and interact with the interior carbon surface of pores. Those ions that are physically adsorbed and especially within the inner pores of porous carbon electrode are not likely to be easily detached by simply reversing the charges, resulting in reduced available surface area of the carbon electrode for further adsorption.

Both the cation exchange membrane and carbon cathode were analyzed by SEM and EDS after the ten cycles of operation to further observe their possible physical deterioration due to repetitive operations. Figure 8c shows that several deposits were found attached including metallic crystals on the cation exchange membrane. The EDS results in Figure 8c and d confirm the presence of Pd crystals. These results show that the decline in the MCDI performance was also likely due to Pd complexation or crystal formation on both the cathode and cation exchange membrane.

Potential Integration of MCDI and Ion Selective Electrodes for Pd Recovery. Electrowinning, one of the most widely used Pd recovery process from wastewaters, is still regarded as an energy-intensive technology. Especially its energy efficiency is critically low for water sources with very low Pd concentrations, and for this reason, the wastewater containing low Pd concentrations is rather disposed. However, as shown in Figure

![Figure 7. Pd removal from catalyst solution for ten cycles.](image-url)
integrating MCDI with the existing electrowinning process is expected to supplement the technology for concentrating Pd ions in the wastewater. This integrated process could be feasible due to the energy efficiency of MCDI and enhanced recovery rate of Pd species. Besides, we can anticipate a more compact Pd recovery process consisting of MCDI only using a Pd selective electrode as presented in Figure 9b which could significantly enhance the energy efficiency of the Pd concentration.

CONCLUSIONS

The potential of MCDI for the recovery of Pd from Pd catalyst wastewater from plating industry has been explored under various operating conditions and solution compositions, and the issues to be overcome are addressed in this work. The performances were measured in terms of the adsorptive and desorptive behavior during multiple cycles of MCDI operations for enrichment of the Pd concentrate. The following are the major findings from this lab-scale study:

- Through adequate process optimization such as operation time and applied potential for a fixed electrode mass, MCDI can be quite effective in Pd removal from plating wastewater containing relatively high Pd concentrations for its recovery and reuse.
- It was demonstrated that Pd can be significantly concentrated through successive cycles of MCDI operation by accumulatively desorbing the ions onto the Pd concentrate. However, as the concentrate solution becomes saturated with ions as adsorption cycles are performed repeatedly, incomplete discharge of Pd ions from the electrodes during the primary desorption phase was observed. Applying higher reverse potential slightly enhanced the desorption efficiency and hence its ability to concentrate the recovered Pd from the feedwater. This unwelcoming effect is concerned to lead this process into unfeasible Pd recovery especially using very low Pd concentration wastewaters such as 1 mg/L of Pd.
- The electrosorptive capacity of the MCDI electrode decreased with the increase in the number of operation cycles because of the likely physical adsorption of Pd ions that cannot be easily released by reversing polarity of the carbon electrodes. SEM and EDS analysis further indicted the formation of a complex or crystal on both surfaces of cathode and cation exchange membrane which may be mitigated through proper physical and chemical cleanings.
- The study pointed out the potential for MCDI to be either integrated with the conventional electrowinning process to supplement Pd recovery from the plating industry. The other option is to apply MCDI using Pd ion selective electrode or ion-exchange membrane which could significantly enhance the MCDI energy efficiency.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acssuschemeng.7b02923.

- Pd removal by physical adsorption during the recovery process (PDF)

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Notes
The authors declare no competing financial interest.

Figure 8. SEM images of virgin (a) carbon cathode and (b) cation exchange membrane and used (c) carbon cathode and (d) cation exchange membrane after the ten-cycle MCDI test. Plus, the Pd/C ratio described in this figure was measured by EDS analysis.

Figure 9. Possible application of MCDI: (a) an integrated Pd recovery process consisting of MCDI and electrowinning and (b) single MCDI process using a Pd selective electrode material.
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