

# The performance of $\text{Cu}^{2+}$ as dissolved cathodic electron-shuttle mediator for $\text{Cr}^{6+}$ reduction in the microbial fuel cell

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## Research

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1 **The performance of  $\text{Cu}^{2+}$  as dissolved cathodic electron-shuttle mediator for  $\text{Cr}^{6+}$**   
2 **reduction in the microbial fuel cell**

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9

10 **Abstract**

11 The study investigates the performance of  $\text{Cu}^{2+}$  as dissolved cathodic electron-shuttle  
12 mediator (dcESM) for simultaneous  $\text{Cr}^{6+}$  reduction and electricity generation in a  
13 microbial fuel cell (MFC) at pH 2 and 4 conditions. The dcESM behavior of  $\text{Cu}^{2+}$  on  
14 carbon cloth (CC) catalyzes the reduction of  $\text{Cr}^{6+}$  into  $\text{Cr}^{3+}$  at pH 2 by undergoing redox  
15 reactions. However, at pH 4, a simultaneous reduction of  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$  was observed.  
16 Cyclic voltammetry studies were performed at pH 2 and 4 to probe the dcESM behavior  
17 of  $\text{Cu}^{2+}$  for  $\text{Cr}^{6+}$  reduction on CC electrode. Also, at pH 2, increasing the concentration  
18 of  $\text{Cu}^{2+}$  from 50 to 500  $\text{mg L}^{-1}$  favors the  $\text{Cr}^{6+}$  reduction by reducing the reaction time  
19 from 108 to 48 h and improving the current production from 3.9 to 6.2  $\text{mA m}^{-2}$ ,  
20 respectively. Nevertheless, at pH 4, the efficacy of  $\text{Cr}^{6+}$  reduction and electricity  
21 generation from MFC is decreased from 63 to 18% and 4.4 to 1.1  $\text{mA m}^{-2}$ , respectively,  
22 by increasing the  $\text{Cu}^{2+}$  concentration from 50 to 500  $\text{mg L}^{-1}$ . Furthermore, the  
23 performance of dcESM behavior of  $\text{Cu}^{2+}$  was explored on carbon felt (CF) and platinum  
24 (Pt) electrodes, and compare the results with CC. In MFC, at pH 2, with an initial  
25 concentration of 100  $\text{mg L}^{-1}$ , the reduction of  $\text{Cr}^{6+}$  in 60 h is 9.6  $\text{mg L}^{-1}$  for CC, 0.2  $\text{mg}$   
26  $\text{L}^{-1}$  for CF, and 51.3  $\text{mg L}^{-1}$  for Pt cathodes. The reduction of  $\text{Cr}^{6+}$  (initial concentration  
27 of 100  $\text{mg L}^{-1}$ ) at pH 4 in 120 h is 44.7  $\text{mg L}^{-1}$  for CC, 32.1  $\text{mg L}^{-1}$  for CF, and 70.9  $\text{mg}$   
28  $\text{L}^{-1}$  for Pt cathodes. Maximum power densities of 1659, 1509, and 1284  $\text{mW m}^{-2}$  were  
29 achieved when CF, CC, and Pt, respectively were employed as cathodes in the MFC.

30 **Keywords:** Microbial fuel cell (MFC); heavy metal removal; hexavalent chromium;  
31 copper; wastewater treatment; bioelectricity generation.

## 32 **1. Introduction**

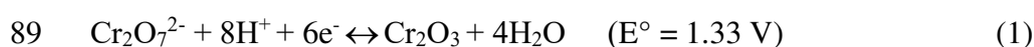
33           In recent years, hexavalent chromium ( $\text{Cr}^{6+}$ ) is exceedingly prevalent in  
34 various industrial effluents, and is often discharged from metallurgy, electroplating,  
35 leather tanning, and textile industries [1].  $\text{Cr}^{6+}$  is a well-known mutagen, teratogen, and  
36 carcinogen [2]. The discharge of  $\text{Cr}^{6+}$  to the environment is of critical concern because:  
37 (i) of its non-biodegradable nature; (ii) it undergoes various transformations and forms  
38 toxic, carcinogenic compounds; and (iii) it is bioaccumulative [3]. The existing  
39 traditional treatment techniques for the removal of  $\text{Cr}^{6+}$  are ion exchange,  
40 adsorption/biosorption, coagulation-flocculation, chemical precipitation,  
41 electrochemical method, biological reduction, and membrane filtration [4]. Although  
42 these techniques are highly promising, long-term applications are often hindered due to  
43 high operational/maintenance costs, additional energy requirement, and the formation  
44 of large secondary toxic sludge.

45           In the past decade, microbial fuel cells (MFCs) have received enormous  
46 attention as a promising technology for wastewater treatment coupled with electricity  
47 generation [5–7]. MFCs are devices that use exoelectrogenic bacteria to oxidize the  
48 organic matter in the anode chamber, thereby producing protons and electrons. The  
49 protons drift internally through a proton exchange membrane (PEM), while the  
50 electrons migrate externally to the cathode chamber, where they are reduced to form  
51 water by an appropriate catholyte [8, 9]. The anode chamber of the MFC is highly  
52 versatile to treat simple organic compounds like acetate [10] and glucose [11] to  
53 complex wastewater such as brewery [12], distillery [13], and starch [14]. Besides, the  
54 cathode chamber of the MFC is successfully employed for treating metal-laden  
55 wastewater containing single metal ions such as cobalt [15], copper [16], silver [17],

56 chromium [3, 18], and selenite [19]. Although MFCs offer promising solutions for the  
57 removal of  $\text{Cr}^{6+}$  [3, 18], the slow reaction kinetics and long operating time due to high  
58 cathodic overpotential hinder it from large-scale applications [20, 21]. Recently,  
59 Krishnani et al. [22] utilized various conductive polymers such as polyaniline,  
60 polypyrrole, polyaniline nanowires and palladium-decorated polyaniline for  $\text{Cr}^{6+}$   
61 reduction due to their electrical properties (like that of a semiconductor) and mechanical  
62 strength. Pang et al. [20] have reported an MFC employed with graphite felt coated  
63 conductive polypyrrole that reduces the cathodic overpotential and improves the  
64 electron shuttling at the cathode-catholyte interface for  $\text{Cr}^{6+}$  reduction. Nevertheless,  
65 their long-term applications are still limited due to complicated synthesis methods, poor  
66 dispersibility, weak stability, and low conductivity [23].

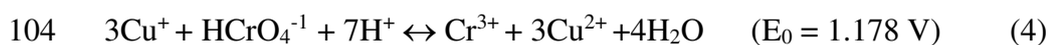
67           Recently, environmentally benign and cost-effective dissolved cathodic  
68 electron-shuttle mediators (dcESMs) have drawn wide attention as they mediate and  
69 expedite the reduction of  $\text{Cr}^{6+}$  in MFC. The dcESM can encourage electron transfer  
70 between the microbes or from cathode to the microbes, and/or from microbes to the  
71 electron-accepting compounds [24]. They exhibit reversible redox reactions and  
72 thereby improve the kinetics of  $\text{Cr}^{6+}$  reduction by diminishing the electrical repulsion  
73 between the negatively charged cathode and the electron acceptors ( $\text{Cr}_2\text{O}_7^{2-}$  or  $\text{CrO}_4^{2-}$ )  
74 [21, 25]. Also, due to their high solubility, they can quickly equilibrate the charges with  
75 cathode and electron-accepting substances. Liu et al. [26] reported improved reduction  
76 of  $\text{Cr}^{6+}$  using  $\text{H}_2\text{O}_2$  as a dcESM. However, its long-term operation was hindered due to  
77 poor oxygen reduction kinetics. Wang et al. [21] noticed  $\text{Fe}^{3+}$  as a dcESM, decreases  
78 diffusion resistance and cathodic overpotential, and hence enhances the  $\text{Cr}^{6+}$  reduction.  
79 Although  $\text{Fe}^{3+}$  improves the  $\text{Cr}^{6+}$  reduction, the power production was decreased by  
80 36% due to the loss of  $\text{Fe}^{3+}$  via reduction.

81 As an alternative, copper ( $\text{Cu}^{2+}$ ) can be used as a dcESM due to its excellent  
82 catalytic behavior.  $\text{Cu}^{2+}$  improves the biocathode performance in MFC by promoting  
83 the electron transfer between cathode and microbes [27–29]. Recently, Li and Zhou  
84 [25] have explored  $\text{Cu}^{2+}$  as a sole dcESM for  $\text{Cr}^{6+}$  reduction in the abiotic cathode of  
85 the MFC. When  $\text{Cu}^{2+}$  is employed as a dcESM for  $\text{Cr}^{6+}$  reduction, the high reduction  
86 potentials of  $\text{Cr}^{6+}$  and  $\text{Cu}^{2+}$  (at 25 °C; Eqs. (1) and (2), respectively) lead them  
87 electrochemically reduce to chromium oxide ( $\text{Cr}_2\text{O}_3$ ) and  $\text{Cu}^0$ , respectively under  
88 closed-circuit conditions of the MFC [2, 25, 30].



91 However, the dcESM behavior of  $\text{Cu}^{2+}$  is precisely influenced by cathode  
92 potential ( $E_h$ ) and pH of the catholyte [16]. The  $E_h$  and pH of  $\text{Cu}^{2+}$  mediated  $\text{Cr}^{6+}$   
93 reduction is calculated (calculations are shown in the Supplemental Materials) and is  
94 shown in Fig. 1a. When pH drops below a critical value ( $\text{pH} \leq 3.2$ ; Fig. 1a),  $\text{Cu}^{2+}$   
95 behaves as a dcESM by undergoing a redox process which expedites the reduction of  
96  $\text{Cr}^{6+}$  as shown in Eqs. (3) and (4) (at 25 °C). For instance,  $\text{Cu}^{2+}$  that are reduced to  $\text{Cu}^{1+}$   
97 acts as the electron donor for the reduction of  $\text{Cr}^{6+}$  to  $\text{Cr}^{3+}$ , while at the same time, the  
98  $\text{Cu}^{1+}$  is oxidized back to  $\text{Cu}^{2+}$  (Eqs. (3) and (4)). This is because, when pH drops below  
99 a critical value, the oxidation state of Cu is +2 (Fig. 1a); hence, it undergoes the redox  
100 process and exhibits dcESM phenomenon. While the valence state of Cr is +6 and it  
101 thermodynamically favors the reduction of  $\text{Cr}^{6+}$  to  $\text{Cr}^{3+}$  in the presence of an ideal  
102 electron donor,  $\text{Cu}^{2+}$ .





105 On the other hand, at pH above a critical value ( $\text{pH} \geq 3.2$ ),  $\text{Cu}^{2+}$  undergo reduction to  
106 stable cuprous oxide ( $\text{Cu}_2\text{O}$ ) (Eq. (5) and Fig. 1a), hindering the dcESM performance  
107 of  $\text{Cu}^{2+}$  on  $\text{Cr}^{6+}$  reduction.



109 Although the mechanism of  $\text{Cu}^{2+}$  in mediating  $\text{Cr}^{6+}$  reduction has been  
110 demonstrated, the dcESM behavior of  $\text{Cu}^{2+}$ , particularly in acidic conditions, has not  
111 yet been reported in any literature. Hence, the present study intends to: (i) demonstrate  
112 the dcESM behavior of  $\text{Cu}^{2+}$  on  $\text{Cr}^{6+}$  reduction at pH 2 and 4 (the two pH conditions  
113 were selected based on the above theoretical evidence) using carbon cloth (CC) as the  
114 electrode in MFC and elucidate the mechanism by cyclic voltammetry (CV) analysis;  
115 (ii) determine the effect of  $\text{Cu}^{2+}$  concentration for  $\text{Cr}^{6+}$  reduction and electricity  
116 production in the MFC; and (iii) compare the performance of  $\text{Cu}^{2+}$  as dcESM on CC  
117 with carbon felt (CF) and platinum (Pt) cathodes in the MFC for  $\text{Cr}^{6+}$  reduction and  
118 electricity generation.

## 119 **2. Materials and methods**

### 120 *2.1 MFC construction*

121 The experimental setup of an MFC is shown in Fig. 1b. A two-chambered  
122 reactor (each chamber having dimensions  $10.5 \times 10 \times 12 \text{ cm}$ ; 500 mL capacity; 300 mL  
123 working volume) was made with a plexiglass acrylic tube. The chambers were arranged  
124 directly adjacent to each other by a PEM (Nafion 117; Sigma-Aldrich; projected surface  
125 area of  $50.24 \text{ cm}^2$ ), and a square-ring with rubber gaskets were held in between the

126 PEM to maintain air-tight condition. PEM was subsequently pretreated with 30% H<sub>2</sub>O<sub>2</sub>,  
127 deionized water, 0.5 M H<sub>2</sub>SO<sub>4</sub>, and deionized water again, for 1 h each [31].

128           The anode was made up of a low molecular heterocyclic aminopyrazine  
129 (Apy)-reduced graphene oxide (r-GO) hybrid coated CC (r-GO-Apy-CC; 25 cm<sup>2</sup>). The  
130 anode material was selected based on one of our previous studies [23], where the r-GO-  
131 Apy-CC electrode was found to exhibit excellent bioelectrocatalytic activity for both  
132 bacterial adhesion and current generation. Electrodes such as CC (25 cm<sup>2</sup>; Synergic  
133 India Pvt. Ltd. India), CF (25 cm<sup>2</sup>; Synergic India Pvt. Ltd. India) and Pt (25 cm<sup>2</sup>; Kevin  
134 Scientific, Chennai) were used as cathodes, and are connected externally to the anode  
135 by a copper wire. The electrodes were previously soaked in deionized water for 24 h,  
136 subsequently dried in the oven at 100 °C for 15 min and were placed at 5 cm apart on  
137 either side of the PEM. The anode and cathode chambers were continuously purged  
138 with nitrogen to maintain anaerobic conditions. The anodic and cathodic pH was  
139 continuously monitored by employing an online pH sensor provided with a multi-  
140 channeled data acquisition system (Aqua controller Model 980 AP, Adsensor; India).  
141 Fluctuations in the pH of the two chambers were monitored by using online indicators  
142 (Fig. 1b).

## 143 2.2 *Inoculation*

144           The anode chamber of the MFC was inoculated with anaerobic sludge  
145 collected from anaerobic digester of the sewage treatment plant, Nesapakkam, Chennai,  
146 India. The sludge was washed with 0.85% NaCl (w/v) solution and subjected to heat  
147 shock pretreatment (100 °C, 2 h) to suppress the activity of methanogens [32]. Sodium  
148 acetate (pH value 7.0) was used as the carbon source in the anode chamber of the MFC.  
149 For inoculation, 50 mL of dewatered anaerobic sludge was added to 250 mL of

150 synthetic wastewater containing macronutrients as  $\text{NH}_4\text{Cl}$ ,  $125 \text{ mg L}^{-1}$ ;  $\text{NaHCO}_3$ ,  $125$   
151  $\text{mg L}^{-1}$ ;  $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ ,  $51 \text{ mg L}^{-1}$ ;  $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$ ,  $300 \text{ mg L}^{-1}$ ;  $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ ,  $6.25 \text{ mg L}^{-1}$   
152 and  $1.25 \text{ mL L}^{-1}$  of trace metal solution as reported in Lovley and Phillips [33]. Nitrogen  
153 gas was purged continuously to maintain anaerobic conditions in the anode and cathode  
154 chambers. Synthetic electroplating wastewater was used as a catholyte and was  
155 prepared by mixing an appropriate quantity of potassium dichromate ( $\text{K}_2\text{Cr}_2\text{O}_7$ ; 99%;  
156 Sigma-Aldrich) and copper sulfate ( $\text{CuSO}_4$ ; 99%; Sigma-Aldrich) with deionized  
157 water. The initial concentrations of  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$  were maintained at  $100 \text{ mg L}^{-1}$ . To  
158 study the effect of  $\text{Cu}^{2+}$  concentration on  $\text{Cr}^{6+}$  reduction, the initial concentration of  $\text{Cr}^{6+}$   
159 was maintained at  $100 \text{ mg L}^{-1}$ , and the different concentrations of  $\text{Cu}^{2+}$  of 50, 100, 300,  
160  $500 \text{ mg L}^{-1}$  were added. The conductivity of the catholyte was improved by adding  $\text{NaCl}$   
161 ( $11.7 \text{ g L}^{-1}$ ). The pH of the influent solution was adjusted with  $\text{H}_2\text{SO}_4$  (0.1 M) and  
162  $\text{NaOH}$  (0.1 M). All the experiments were carried out at ambient temperature.

### 163 2.3 *Measurements and analyses*

164 Hexavalent chromium was analyzed by UV-Vis spectrophotometer (UV-  
165 1800 PC, Shimadzu) at 540 nm. Total copper was measured using Atomic Absorption  
166 Spectrophotometry (AAnalyst 700, Perkin Elmer) after sampling at regular intervals.  
167 The removal efficiency was calculated using Eq. (6).

$$168 \text{ Removal efficiency (\%)} = \frac{(A-B)}{A} \times 100 \quad (6)$$

169 where A and B are the initial and observed concentrations in  $\text{mg L}^{-1}$ .

170 A precise 4-channel potentiostat (VSP 300 Biologic; India) was employed to  
171 investigate the electrochemical characteristics of the system. Current (I) was calculated  
172 according to Ohm's law,  $I = V/R$ , where R is the external resistance. Power (P) is the

173 product of voltage  $V$  and  $I$  ( $P = IV$ ). Power and current densities were calculated by  
174 dividing the respective terms by the cathode surface area ( $m^2$ ). The polarization study  
175 was performed by employing the I-V characterization technique using a three-electrode  
176 system, where the working electrode was a cathode, the counter electrode was an anode,  
177 and reference electrode was saturated Ag/AgCl (+ 0.197 V vs. SHE), that was placed  
178 close to the cathode.

179 The electrochemical activity of the electrode was examined by CV analysis  
180 in a separate single-cell system. The experiment was performed in a high purity quartz  
181 glass beaker (500 mL) with an airtight polytetrafluoroethylene cap, mounted over  
182 silicon encapsulated polytetrafluoroethylene ring. The cell system consists of a counter  
183 electrode of standard Pt, reference electrode of Ag/AgCl, and the working electrode of  
184 CC. CV analysis was conducted in the cell system deployed with  $Cu^{2+}$  mediated  $Cr^{6+}$   
185 solution (1:1), and the performance was compared with a controlled solution of  $Cr^{6+}$   
186 and  $Cu^{2+}$  solution. During CV analysis, the working electrode potential in controlled  
187  $Cu^{2+}$  and  $Cu^{2+}$  mediated  $Cr^{6+}$  solutions were linearly scanned from +1 V to -1 V at a  
188 scan rate of  $10\text{ mV s}^{-1}$ . In controlled  $Cr^{6+}$  solution, the potential was linearly scanned  
189 from +0.5 to +1.5 V for pH 2, and +0.5 to 1.0 V for pH 4 at a scan rate of  $10\text{ mV s}^{-1}$ .  
190 High-resolution scanning electron microscopy (FEI Quanta 200 FEG) equipped with  
191 energy-dispersive x-ray spectroscopy was employed to confirm  $Cr^{3+}$  and  $Cu_2O$   
192 monolayer formation on cathode surfaces for a specific period.

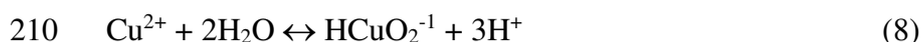
### 193 **3. Results and discussion**

#### 194 *3.1 The dcESM behavior of $Cu^{2+}$ for $Cr^{6+}$ reduction*

195 The dcESM behavior of  $Cu^{2+}$  on  $Cr^{6+}$  reduction (initial concentrations of 100  
196  $mg\text{ L}^{-1}$ ; 1:1 ratio) was characterized on CC cathode in MFC at pH 2 and 4. The temporal

197 behavior of  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$ , and corresponding variations in pH were monitored at pH 2  
198 and 4. The experimental result at pH 2 displays a large reduction in  $\text{Cr}^{6+}$  to  $\text{Cr}^{3+}$  with  
199 an efficiency of 99.9% at 84 h of operating time (Fig. 2a). In contrast, the reduction of  
200  $\text{Cu}^{2+}$  was not significant at pH 2. A slight decrease (25%) in the  $\text{Cu}^{2+}$  concentration was  
201 noticed within the initial few hours, probably, due to the reduction of  $\text{Cu}^{2+}$  to  $\text{Cu}^{1+}$  (Eq.  
202 (3)). However, at pH 4, a considerable reduction of  $\text{Cu}^{2+}$  along with  $\text{Cr}^{6+}$  was observed.

203           The results were interpreted that the reduction of  $\text{Cr}^{6+}$  in the presence of  $\text{Cu}^{2+}$   
204 involves heterogeneous reactions. At pH 2,  $\text{Cr}^{6+}$  and  $\text{Cu}^{2+}$  are highly protonated by  
205 surrounding the  $\text{H}^+$  ions and predominantly exist in the form of  $\text{HCrO}_4^{-1}$  and  $\text{HCuO}_2^{-1}$ ,  
206 respectively as shown in the reactions Eqs. (7) and (8). During the initial 10 h, a  
207 decrease in the pH was noticed from the initial pH (pH 2) due to the hydrolysis of  $\text{Cu}^{2+}$   
208 which involves an increase in the  $\text{H}^+$  ions in the solution as shown in Eq. (8).



211           Besides, a slight reduction in the  $\text{Cu}^{2+}$  (25%) was observed at this stage,  
212 probably due to the cathodic reduction of  $\text{HCuO}_2^{-1}$  to  $\text{Cu}^+$  in the presence of  $\text{H}^+$  ions as  
213 explained in Eq. (3) [34]. However, a steady-state reduction in  $\text{Cr}^{6+}$  concentration was  
214 observed throughout the experiment. This could be correlated with Eqs. (3) and (4) and  
215 explained as: the  $\text{Cu}^{2+}$  that is reduced to  $\text{Cu}^{1+}$  acts as an electron donor for the reduction  
216 of  $\text{Cr}^{6+}$  to  $\text{Cr}^{3+}$ , while at the same time, the  $\text{Cu}^{1+}$  is oxidized back to  $\text{Cu}^{2+}$ .  
217 Correspondingly, the pH of catholyte was increased owing to the consumption of  $\text{H}^+$   
218 ions as indicated in Eqs. (3) and (4). The color of the wastewater was changed from  
219 orange-yellow to greenish-yellow after 48 h, indicating, the complete reduction of  $\text{Cr}^{6+}$   
220 to  $\text{Cr}^{3+}$ .

221 A separate CV analysis was conducted to confirm the dcESM behavior of  
222  $\text{Cu}^{2+}$  on  $\text{Cr}^{6+}$  reduction at pH 2 using a single cell system with CC as a working electrode  
223 (Fig. 2b). At pH 2, CV analysis of controlled  $\text{Cu}^{2+}$  solution exhibits well-defined  
224 reduction and oxidation peaks at -0.36 and +0.36 V, respectively, which could be  
225 attributed to the dcESM behavior of  $\text{Cu}^{2+}$  on CC electrode (Fig. 2b) [35]. In a mixture  
226 of  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$  solution, CV analysis was repeated for 70 cycles at a scan rate of 10  
227  $\text{mV s}^{-1}$ . During the forward scan, a large cathodic current was drawn after 0.23V was  
228 due to the combined reduction of  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$ , and on the reverse scan, the peak  
229 observed at -0.34 V was correlated with the oxidation of  $\text{Cr}^{2+}$  to  $\text{Cr}^{3+}$  (Cycle 5; Fig. 2b)  
230 [36]. The additional peak observed at -0.01 V (Cycle 30; Fig. 2b) can be attributed to  
231 the oxidation of  $\text{Cu}^{1+}$  to  $\text{Cu}^{2+}$ . While continuing the scan up to 70 cycles, the peak at -  
232 0.34 V was observed to be diminished due to the stable and insoluble  $\text{Cr}^{3+}$  formation.  
233 Concurrently, the peak observed at -0.01 V was found to increase due to the increases  
234 in  $\text{Cu}^{1+}$  concentration. The CV study confirms the dcESM behavior of  $\text{Cu}^{2+}$  on  $\text{Cr}^{6+}$   
235 reduction at pH 2 and complements with the reported experimental results.

236 The proposed mechanism for the reduction of  $\text{Cr}^{6+}$  and  $\text{Cu}^{2+}$  at pH 2 and 4  
237 are shown in Fig. 3. At pH 4, simultaneous reduction of  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$  was observed  
238 with removal efficiencies of 71 and 56%, respectively (Fig. 4a). At this pH condition,  
239 dcESM behavior of  $\text{Cu}^{2+}$  was not observed; instead, simultaneous reduction of  $\text{Cr}^{6+}$  and  
240  $\text{Cu}^{2+}$  was noticed. This could be due to the stable form of  $\text{Cu}_2\text{O}$  which hinders the  
241 dcESM behavior of  $\text{Cu}^{2+}$  for  $\text{Cr}^{6+}$  reduction. However, simultaneous reductions of  $\text{Cu}^{2+}$   
242 and  $\text{Cr}^{6+}$  were occurred owing to the high cathodic potential as presented in Eqs. (1)  
243 and (5). The pH was observed to increase from 4 to 5.54 and the color of the effluent  
244 was changed from orange-yellow to pale yellow. In CV analysis, the peak observed at

245 -0.34 and -0.01 V at pH 2 conditions were absent at pH 4 and can be attributed to the  
246 irreversibility of the  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$  as stable  $\text{Cu}_2\text{O}$  and  $\text{Cr}_2\text{O}_3$ , respectively (Fig. 4b).

### 247 3.2 *Effect of $\text{Cu}^{2+}$ concentration on $\text{Cr}^{6+}$ reduction and electricity production*

248 Studies were performed by varying the  $\text{Cu}^{2+}$  concentration at pH 2 and 4 to  
249 understand the effect of  $\text{Cu}^{2+}$  and its dcESM behavior on  $\text{Cr}^{6+}$  reduction and  
250 bioelectricity generation. The results elucidate that, at pH 2, the presence of  $\text{Cu}^{2+}$  is  
251 highly favorable for the reduction of  $\text{Cr}^{6+}$ . Increasing the  $\text{Cu}^{2+}$  concentration from 50 to  
252  $500 \text{ mg L}^{-1}$  reduces reaction time from 108 to 48 h for the complete reduction of  $\text{Cr}^{6+}$   
253 (Fig. 5a). Similarly, the current production was improved from  $3.94 \text{ mA m}^{-2}$  by  
254 increasing the  $\text{Cu}^{2+}$  concentration from 50 to  $500 \text{ mg L}^{-1}$  (Fig. 5b). However, at pH 4,  
255 the presence of  $\text{Cu}^{2+}$  decreases the reduction of  $\text{Cr}^{6+}$ . By increasing the  $\text{Cu}^{2+}$   
256 concentration from 50 to  $500 \text{ mg L}^{-1}$ , the reduction efficiency of  $\text{Cr}^{6+}$  was observed to  
257 decrease from 63 to 18% (Fig. 5c). Correspondingly, the response in the current density  
258 was decreased from  $4.4 \text{ mA m}^{-2}$  (616 mV) to  $1.1 \text{ mA m}^{-2}$  (155 mV), respectively (Fig.  
259 5d).

260 At pH 2, increasing the  $\text{Cu}^{2+}$  concentration improves the reduction of  $\text{Cr}^{6+}$   
261 and electricity generation from MFC. This could be due to the dcESM behaviour of  
262  $\text{Cu}^{2+}$  improves the kinetics of  $\text{Cr}^{6+}$  reduction by diminishing the electrical repulsion  
263 between the negatively charged cathode and the  $\text{Cr}_2\text{O}_7^{2-}$  anions in the catholyte [21, 25].  
264 As the rate of electron transfer improves, the cathodic over potential decreases resulting  
265 in an improvement in the generation of electricity. On the other hand, at pH 4,  
266 increasing the concentration of  $\text{Cu}^{2+}$  decreases the reduction kinetics of  $\text{Cr}^{6+}$  as well as  
267 the responses in the current density. At pH 4,  $\text{Cu}^{2+}$  was electrochemically reduced to  
268  $\text{Cu}_2\text{O}$  (Eq. (5) and Fig. 1a) and deposited over the electrode surface (Fig. 7b). The

269 deposition of Cu<sub>2</sub>O increases with increase in the Cu<sup>2+</sup> concentration. This results in  
 270 higher cathodic overpotential at the cathode-catholyte interface that hinders the kinetics  
 271 of Cr<sup>6+</sup> reduction and current production at pH 4. In both the pH conditions, the  
 272 temporal response of the current density was found to be decreased as the experiment  
 273 progresses. The trend can be correlated with the decrease in the catholyte concentration  
 274 due to the reduction of Cr<sup>6+</sup> and can be theoretically explained by the Nernst equation as  
 275 in Eqs. (9) and (10).

$$276 \quad E_{\text{cathode}} - E_0 = -\frac{2.303 RT}{nF} \log \frac{[Cr^{3+}]}{[Cr^{6+}]} \quad (9)$$

$$277 \quad \eta_c = -\frac{2.303 RT}{nF} \log \frac{[Cr^{3+}]}{[Cr^{6+}]} \quad (10)$$

278 According to the Eq. (10), overpotential ( $\eta_c$ ) becomes more negative as the  
 279 concentration of Cr<sup>3+</sup> increases, causing a decrease in the cell potential [38, 39].  
 280 Eventually, the response in the current density decreases as the experiment progresses.  
 281 When the rate of reduction of Cr<sup>6+</sup> reaches an asymptotic state, the response of the  
 282 current density exhibits an analogous behavior to the concentration of Cr<sup>6+</sup>.

### 283 3.3 The dcESM behavior of Cu<sup>2+</sup> on CF and Pt electrodes

284 The dcESM behavior of Cu<sup>2+</sup> on Cr<sup>6+</sup> reduction (initial concentrations of 100  
 285 mg L<sup>-1</sup>; 1:1 ratio) was characterized on Pt and CF in MFC at pH 2 (Fig. 6a) and pH 4  
 286 (Fig. 6b), and the results were compared with CC electrode. It was observed that, at pH  
 287 2, the reduction of Cr<sup>6+</sup> to Cr<sup>3+</sup> in 60 h is 99.8% for CF cathode, 48% for Pt cathode,  
 288 and 90% for CC cathode. As expected, not much reduction of Cu<sup>2+</sup> occurred in all the  
 289 electrodes at pH 2 (Fig. 6a). The color of the wastewater was changed from orange-  
 290 yellow to greenish-yellow, and subsequently, a stable blue color solution when CC and

291 CF were employed as the cathode material. However, in the case of Pt as a cathode, the  
292 color of the wastewater was changed moderately from orange-yellow to pale yellow.  
293 At pH 4, the simultaneous reduction of  $\text{Cr}^{6+}$  and  $\text{Cu}^{2+}$  was observed for Pt and CF  
294 electrodes. The reduction of  $\text{Cr}^{6+}$  at pH 4 in 120 h is 68% for CF cathode, 29% for Pt  
295 cathode, and 56% for CC cathode (Fig. 6b). Similarly,  $\text{Cu}^{2+}$  reduction in 120 h is 75%  
296 for CF cathode, 50% for Pt cathode, and 71% for CC cathode. The results indicate that  
297 the dcESM behavior of  $\text{Cu}^{2+}$  on  $\text{Cr}^{6+}$  reduction is exhibited not only on CC but also in  
298 CF and Pt cathodes in MFC.

299 The surface morphological characteristics of the  $\text{Cr}^{3+}$  and  $\text{Cu}_2\text{O}$  on the surface  
300 of Pt, CC, and CF were analyzed by performing the SEM analysis (Fig. 7a, 7b, and 7c,  
301 respectively), and the image clearly shows the nucleation of  $\text{Cr}^{3+}$  or  $\text{Cr}_2\text{O}_3$  and/or  $\text{Cu}_2\text{O}$   
302 on the cathode surface. Furthermore, polarization studies were performed with CC, CF  
303 and Pt to understand the efficiency of electrode material for power production (Fig. 7d).  
304 The maximum power densities of  $1659 \text{ mW m}^{-2}$  ( $4.9 \text{ mA m}^{-2}$ ),  $1509 \text{ mW m}^{-2}$  ( $5.5 \text{ mA}$   
305  $\text{m}^{-2}$ ), and  $1284 \text{ mW m}^{-2}$  ( $5.1 \text{ mA m}^{-2}$ ) were achieved when CF, CC, and Pt were  
306 employed as cathode materials. The study confirms that carbon-based electrode  
307 materials are ideal for bioelectricity generation from MFC.

### 308 3.4 *Practical implication to scale up MFC technology*

309 In summary, MFCs apply a simple redox principle in which the sole  
310 influencing factor for the entire electrochemical reductions is the degradation of organic  
311 matter (The details are provided in Supplementary Materials) by exoelectrogenic  
312 microorganisms at the anode chamber. At the cathode,  $\text{Cr}^{6+}$  electrochemically reduces  
313 to  $\text{Cr}^{3+}$  by accepting the electrons from the anode chamber. The use of dissolved  
314 electron–shuttle mediators reduces the activation energy at the cathode–electrolyte

315 interface and improves the cathode performance for  $\text{Cr}^{6+}$  reduction and bioelectricity  
316 generation. In fact,  $\text{Cu}^{2+}$  is a potential contaminant, and extreme consumption of copper  
317 leads to severe toxicological concerns, such as nausea, contractions, convulsions, or  
318 even death [16, 40]. Hence, incorporating  $\text{Cu}^{2+}$  as a dissolved mediator in MFC is a  
319 sustainable approach because  $\text{Cu}^{2+}$  not only enhances the reduction of  $\text{Cr}^{6+}$  but also  
320 reduces to its most stable,  $\text{Cu}_2\text{O}$  form, simultaneously. The findings demonstrated in  
321 the present study are highly significant to the electroplating industry where a  
322 combination of  $\text{Cr}^{6+}$  and  $\text{Cu}^{2+}$  are discharged at high acidic conditions from washing,  
323 rinsing, batch dumps, and processing and/or operational units. Presently, chemical  
324 coagulation/precipitation is the most widely adopted treatment technique for  
325 electroplating wastewater. However, this technique involves high operational costs due  
326 to the consumption of large amounts of chemicals and the generation of a huge quantity  
327 of sludge. Instead, the use of  $\text{Cu}^{2+}$  as a dcESM for  $\text{Cr}^{6+}$  reduction is a cost-effective  
328 method as it does not require any addition of chemicals, and both often co-exist in the  
329 effluents discharged from electroplating or mining industries. However, further studies  
330 are required to evaluate the long-term operation conditions and process economy of  
331 MFC over influent. Also, pilot/full-scale studies are needed for the practical  
332 implementation of this technology in industries.

#### 333 **4 Conclusions**

334 In the present work, dcESM phenomenon of  $\text{Cu}^{2+}$  on  $\text{Cr}^{6+}$  reduction in MFC  
335 is reported using CC, CF, and Pt electrodes. The dcESM behavior of  $\text{Cu}^{2+}$  for  $\text{Cr}^{6+}$   
336 reduction is highly influenced by  $E_h$  and pH of the catholyte in MFC. In acidic  
337 conditions, when the pH is below a critical point of 3.2,  $\text{Cu}^{2+}$  improves the reduction of  
338  $\text{Cr}^{6+}$  to 99.9% at 84 h of operating time. On the other hand, above the critical pH point,

339 simultaneous reduction of  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$  was observed with removal efficiencies of 71  
340 and 56%, respectively. Hence, the  $\text{Cu}^{2+}$  and its role as dcESM in MFC is highly  
341 advantageous as it not only enhances the  $\text{Cr}^{6+}$  reduction but also undergoes  
342 electrochemical reduction into non-toxic  $\text{Cu}_2\text{O}$ . Since  $\text{Cu}^{2+}$  and  $\text{Cr}^{6+}$  often co-exist in the  
343 wastewater from electroplating or mining industries, and due to their synergy, the study  
344 provides an entirely new concept of ‘using a pollutant to treat another one’ with  
345 simultaneous generation of energy.

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348 A/ET-1017/2014] in carrying out this research work.

#### 349 **Authors’ contributions**

350 Praveena Gangadharan planned and conducted the experiments. Indumathi M Nambi  
351 supervised the project.

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#### 354 **Availability of data and materials**

355 The data used to support the findings of this study are available from the corresponding  
356 author upon request.

#### 357 **Competing interests**

358 The authors declare that they have no competing interests.

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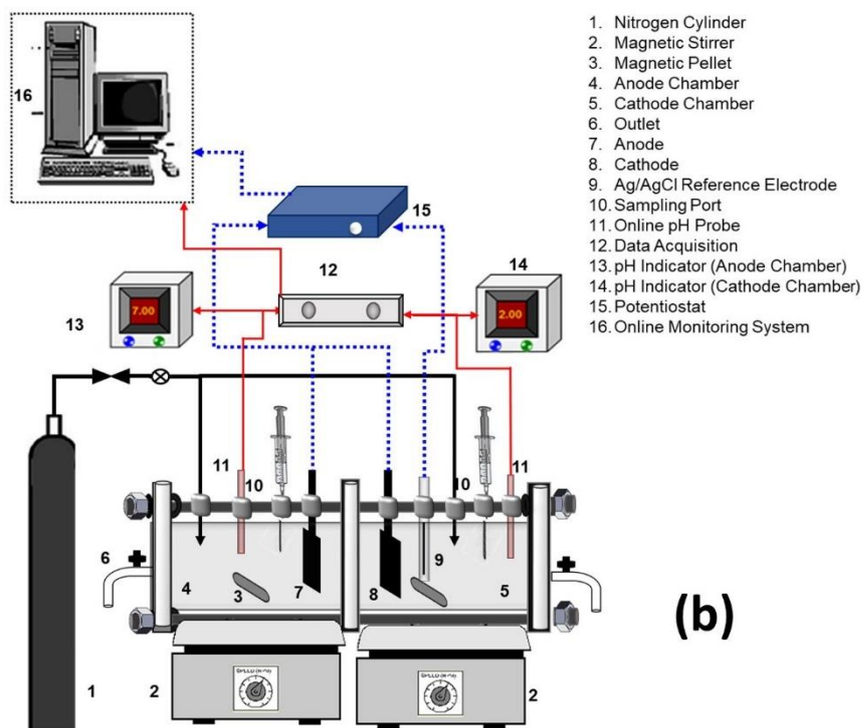
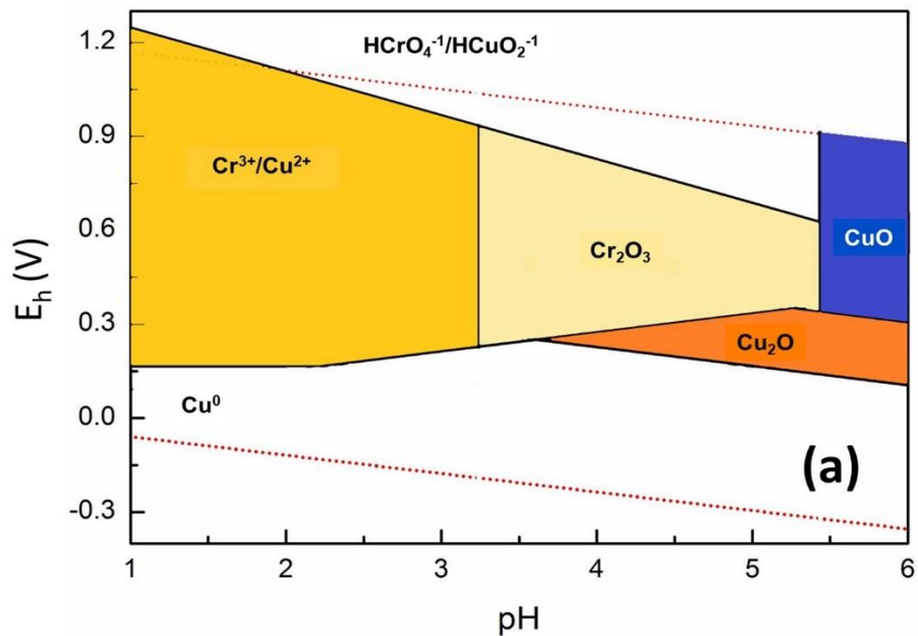
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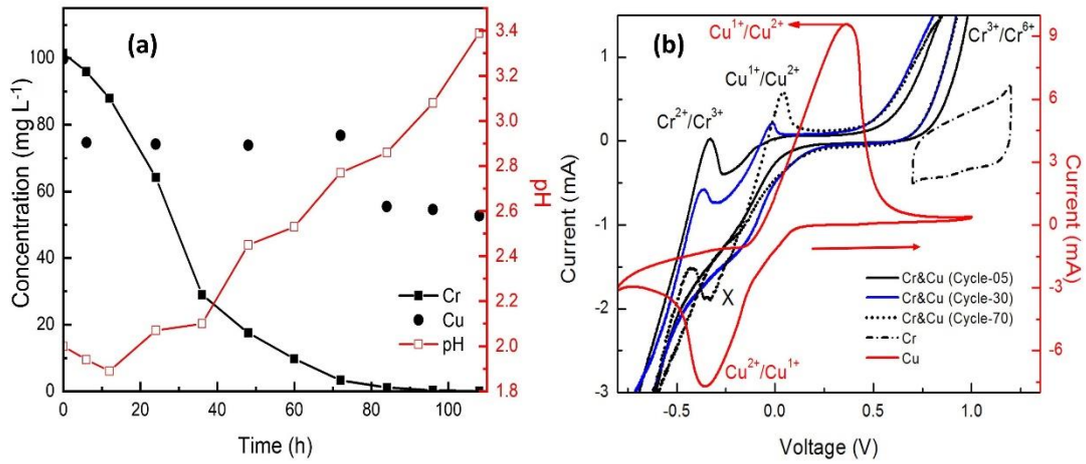
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472 **Fig. 1** (a) The equilibrium  $E_h$ -pH diagram of Cr-Cu-H<sub>2</sub>O system; (b) The

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experimental set-up of MFC.

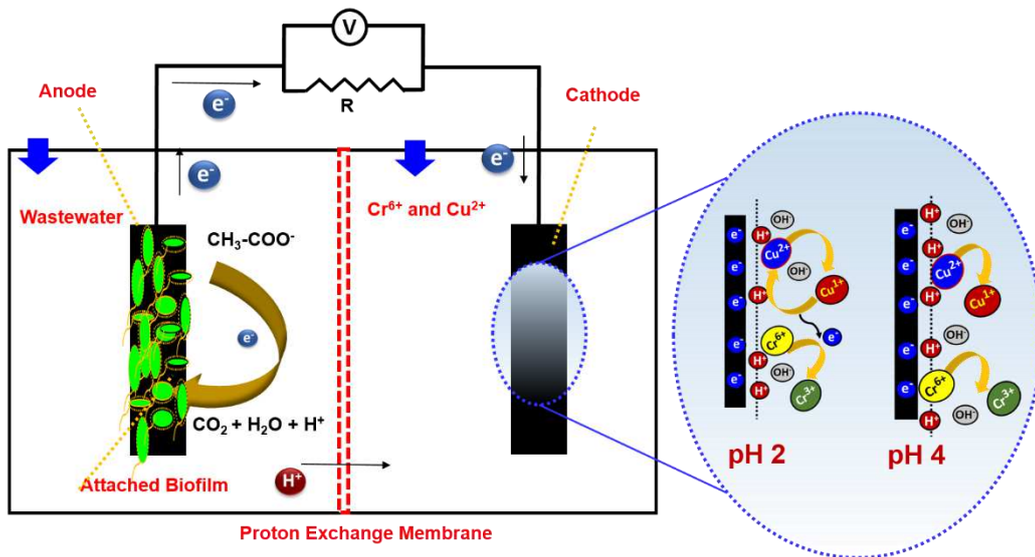
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476 **Fig. 2** (a) Temporal behavior of Cr<sup>6+</sup> and Cu<sup>2+</sup> and corresponding change in the pH  
 477 value at pH 2; (b) Cyclic voltammograms of CC electrode in Cu<sup>2+</sup>, Cr<sup>6+</sup>, and Cr<sup>6+</sup> &  
 478 Cu<sup>2+</sup> mediated solution at pH 2.

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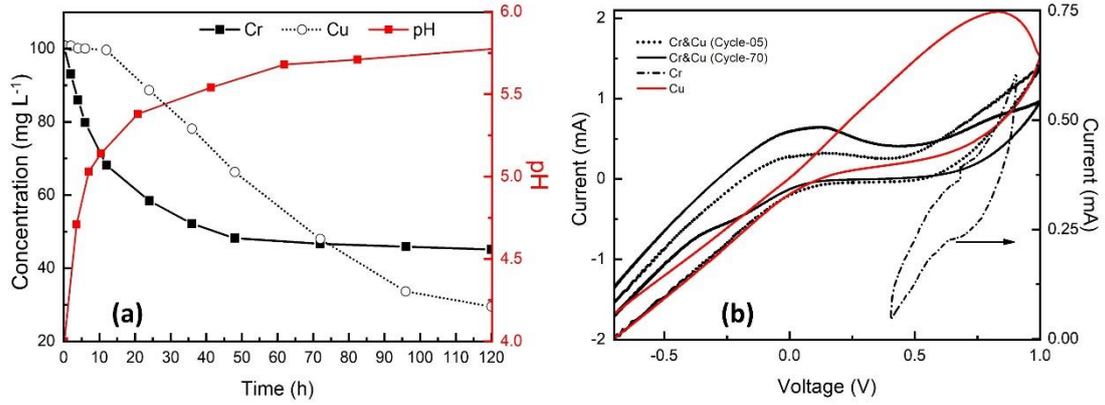
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481 **Fig. 3** The proposed mechanism for the reduction of Cr<sup>6+</sup> and Cu<sup>2+</sup> at pH 2 and 4.

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486 **Fig. 4** (a) Temporal behavior of Cr<sup>6+</sup> and Cu<sup>2+</sup> and corresponding change in the pH

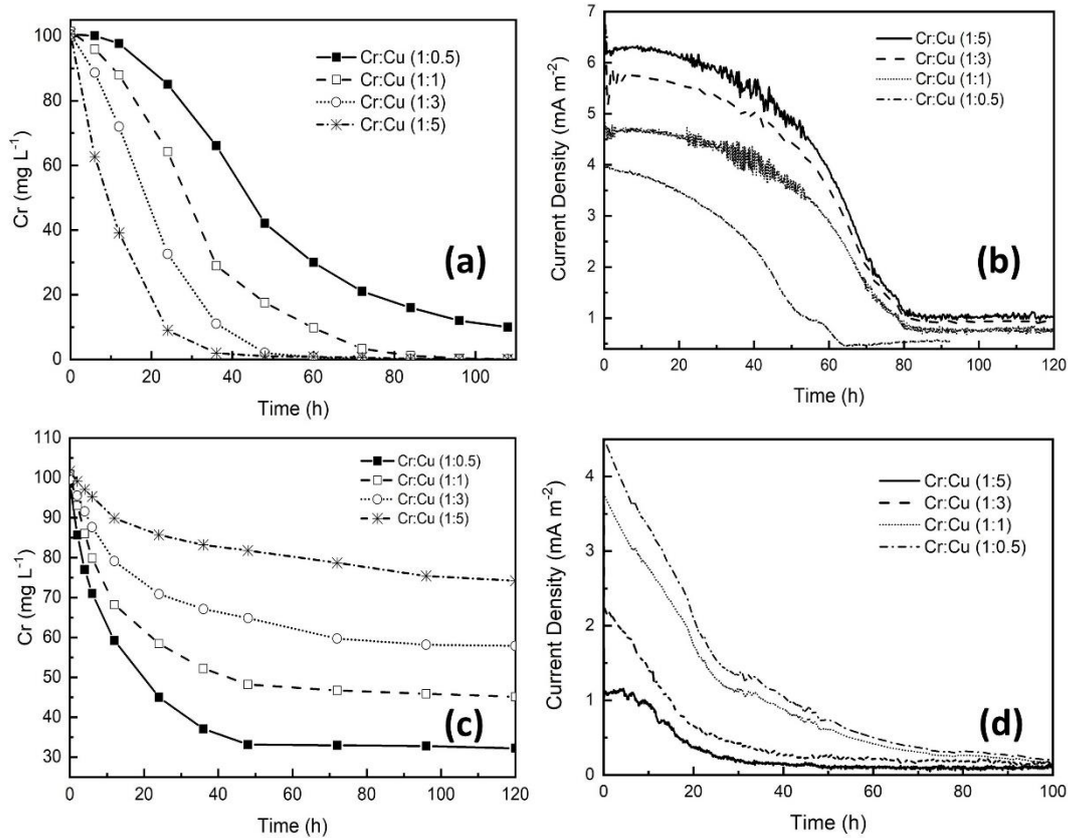
487 value at pH 4; (b) Cyclic voltammograms of CC electrode in Cu<sup>2+</sup>, Cr<sup>6+</sup>, and Cr<sup>6+</sup> &

488 Cu<sup>2+</sup> mediated solution at pH 4.

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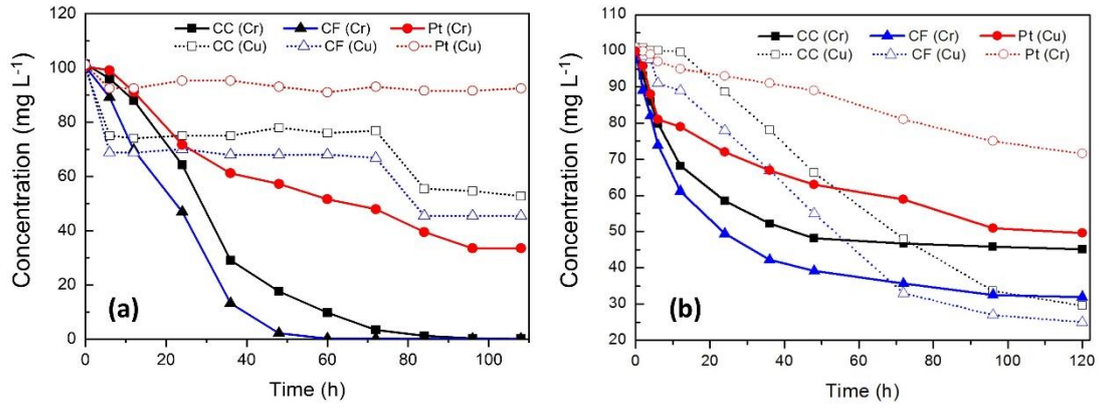
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493 **Fig. 5** Effect of  $\text{Cu}^{2+}$  concentration at pH 2: (a) temporal reduction of  $\text{Cr}^{6+}$ ; (b) response  
 494 in the current density. Effect of  $\text{Cu}^{2+}$  concentration at pH 4: (a) temporal reduction of  
 495  $\text{Cr}^{6+}$ ; (b) response in the current density.

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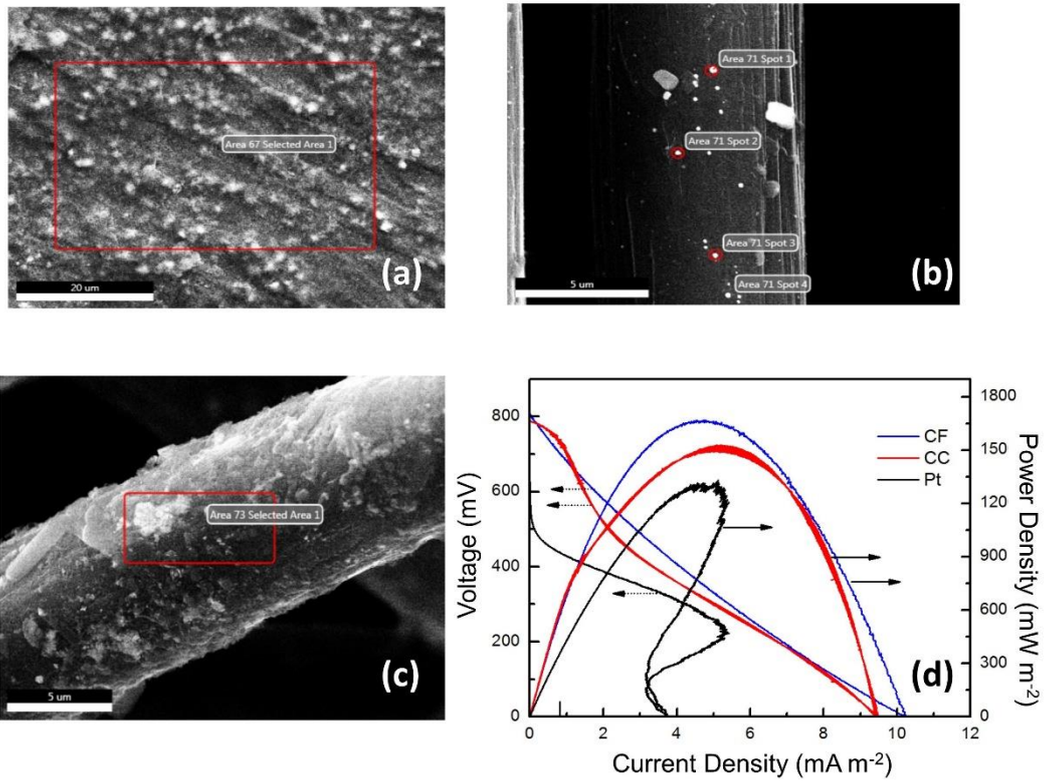
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500 **Fig. 6** dcESM behavior  $\text{Cu}^{2+}$  on CC, CF, and Pt: (a) at pH 2; (b) at pH 4.



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502 **Fig. 7** SEM image of Cr and Cu adsorbed on the surface of cathode: (a) Pt; (b) CC;

503 (c) CF; (d) Polarization curve.

# Figures

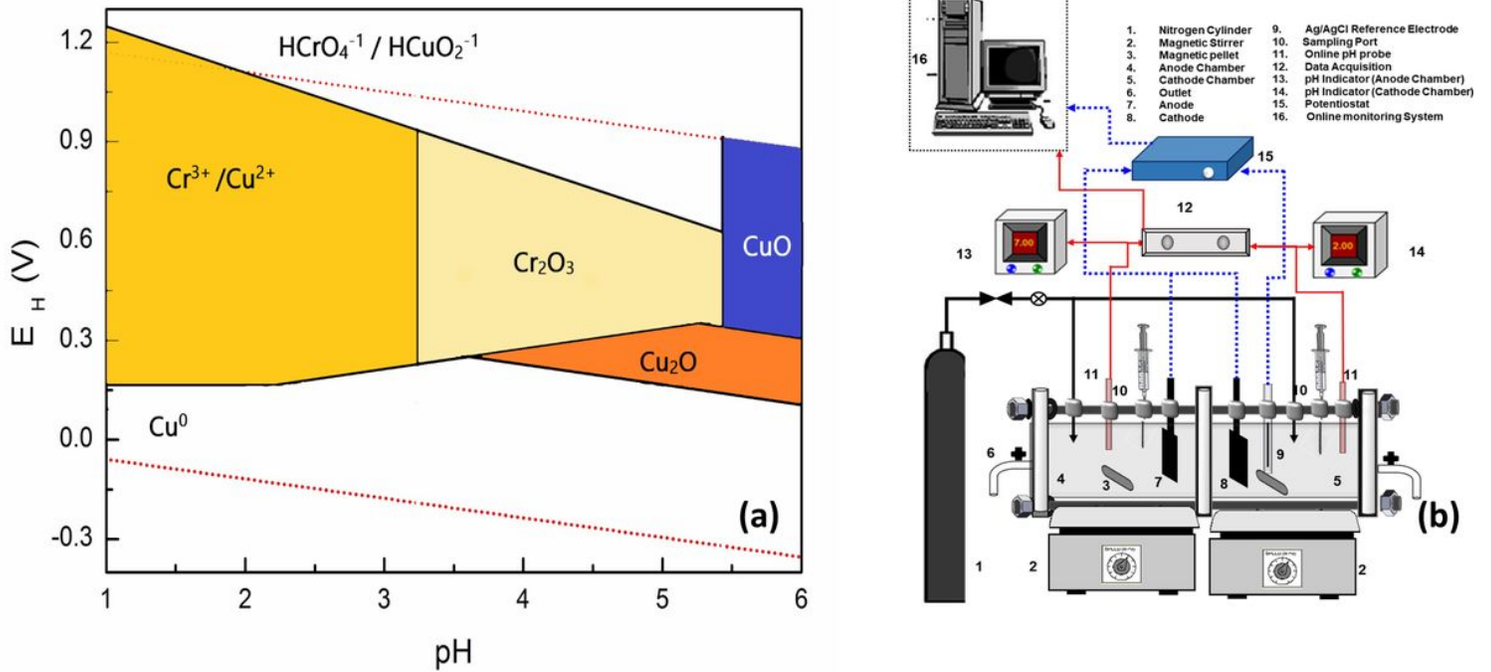


Figure 1

(a) The equilibrium Eh – pH diagram of Cr-Cu-H<sub>2</sub>O system. (b) The experimental set-up of MFC.

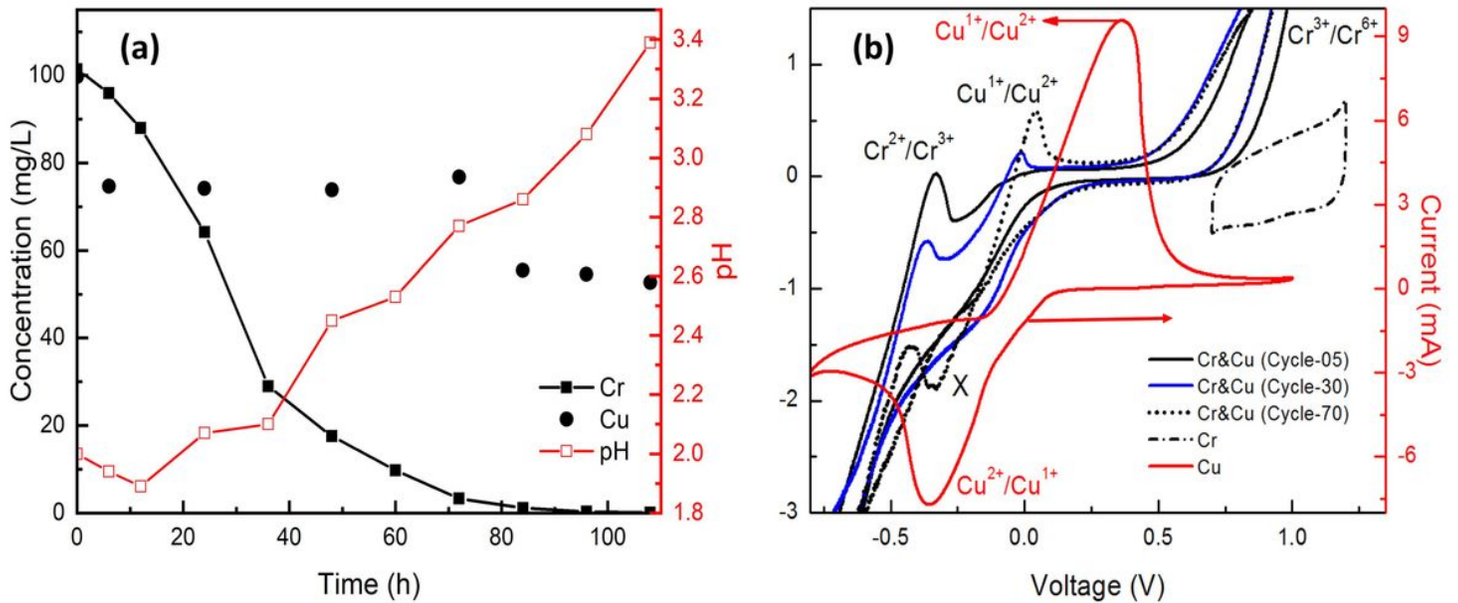


Figure 2

(a) Temporal behavior of Cr<sup>6+</sup> and Cu<sup>2+</sup> and corresponding change in the pH value at pH 2. (b) Cyclic voltammograms of CC electrode in Cu<sup>2+</sup>, Cr<sup>6+</sup>, and Cr<sup>6+</sup> & Cu<sup>2+</sup> mediated solution at pH 2.

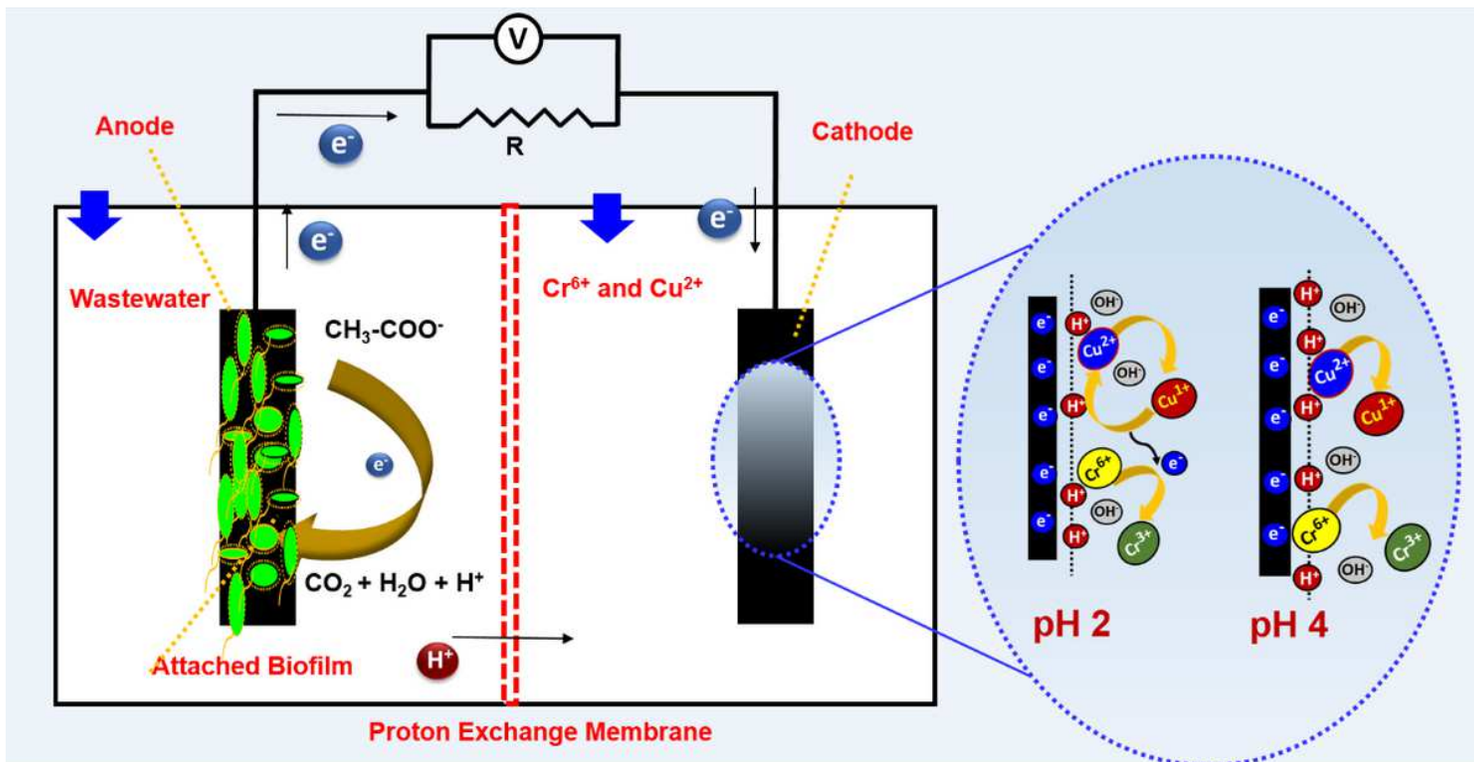


Figure 3

The proposed mechanism for the reduction of  $\text{Cr}^{6+}$  and  $\text{Cu}^{2+}$  at pH 2 and pH 4.

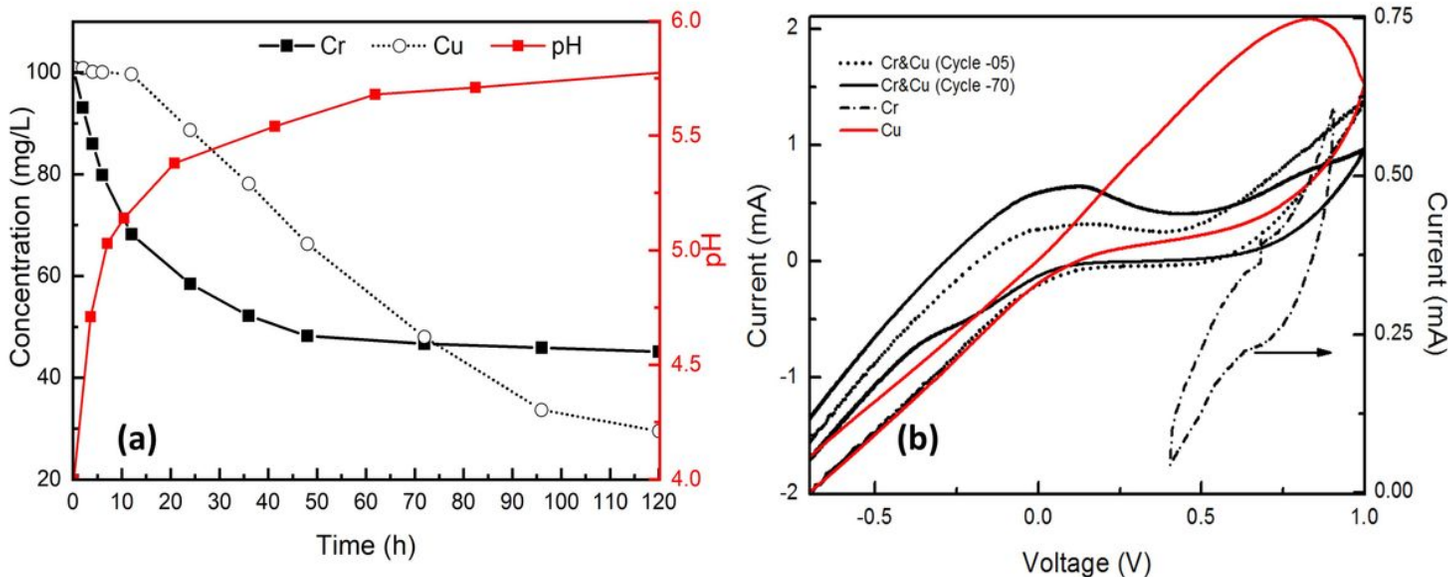
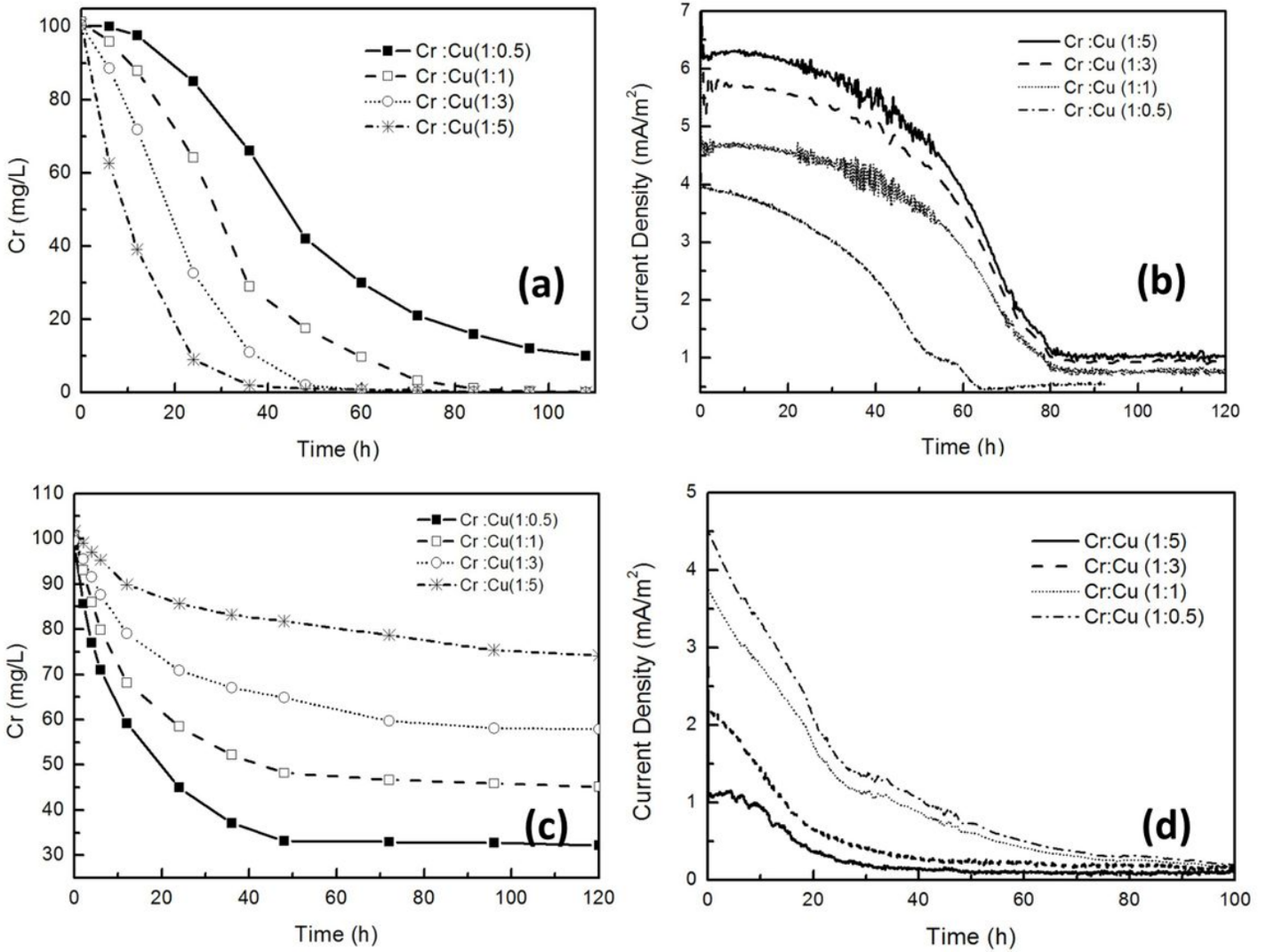


Figure 4

(a) Temporal behavior of  $\text{Cr}^{6+}$  and  $\text{Cu}^{2+}$  and corresponding change in the pH value at pH 4. (b) Cyclic voltammograms of CC electrode in  $\text{Cu}^{2+}$ ,  $\text{Cr}^{6+}$ , and  $\text{Cr}^{6+}$  &  $\text{Cu}^{2+}$  mediated solution at pH 4.



**Figure 5**

Effect of  $\text{Cu}^{2+}$  concentration at pH 2: (a) temporal reduction of  $\text{Cr}^{6+}$ ; (b) response in the current density.  
 Effect of  $\text{Cu}^{2+}$  concentration at pH 4: (a) temporal reduction of  $\text{Cr}^{6+}$ ; (b) response in the current density.

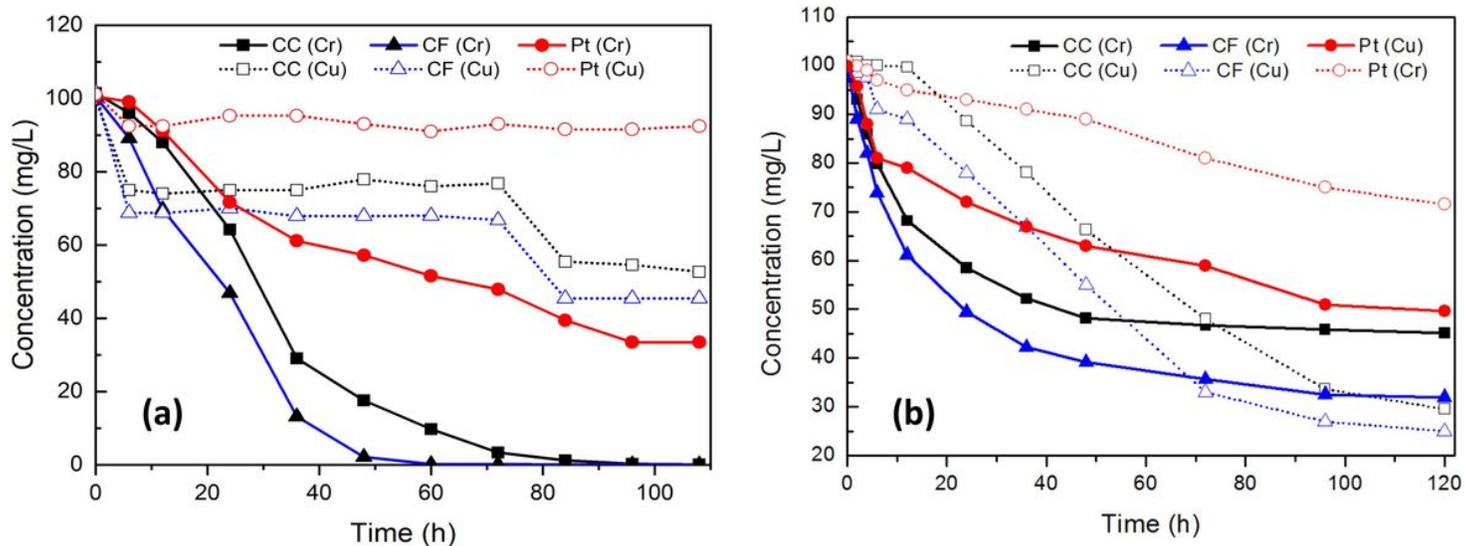
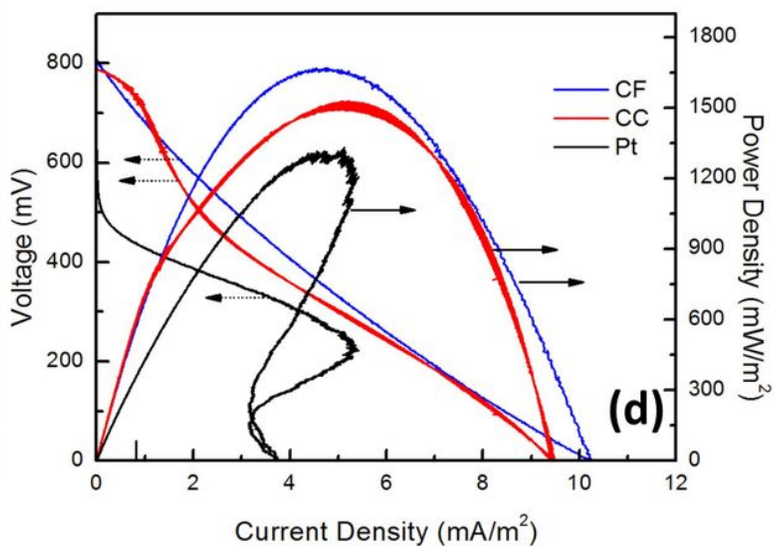
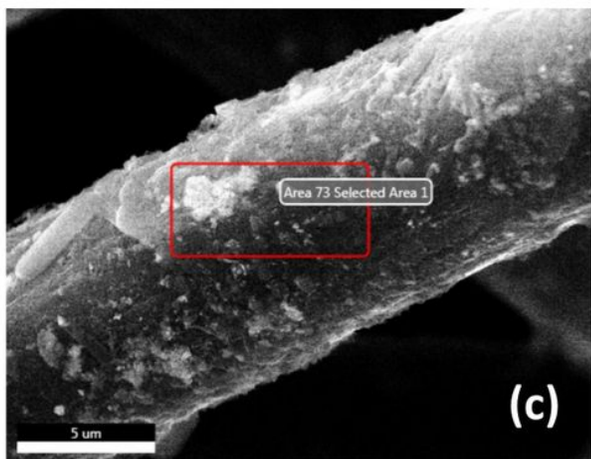
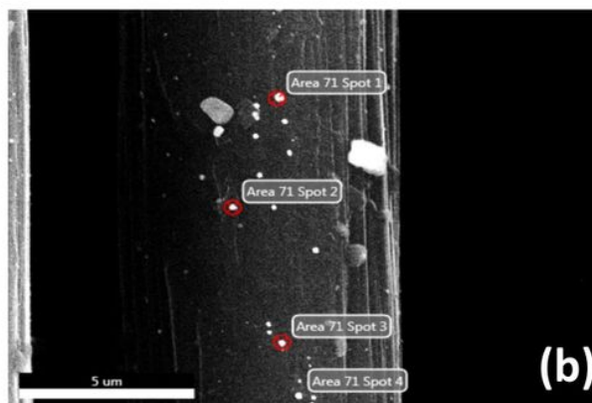
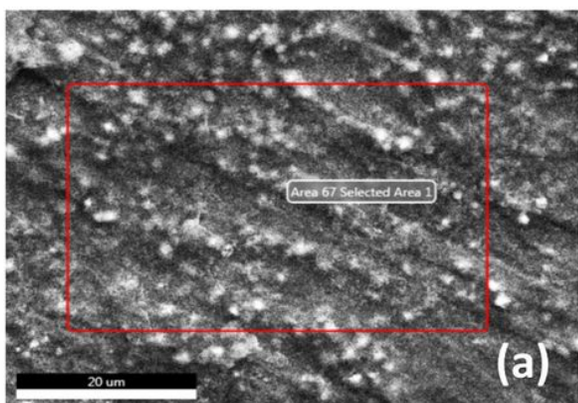


Figure 6

dcESM behavior  $\text{Cu}^{2+}$  on CC, CF, and Pt: (a) at pH 2; (b) at pH 4.



## Figure 7

SEM image of Cr and Cu adsorbed on the surface of cathode: (a) Pt; (b) CC; (c) CF. (d) Polarization curve.

## Supplementary Files

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- [SupplementaryMaterialFinal.docx](#)