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3	Correlation between circuital current, Cu(II) reduction and cellular
4	electron transfer in EAB isolated from Cu(II)-reduced biocathodes

# 5 of microbial fuel cells

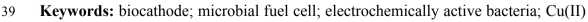
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# 18 Abstract

The performance of four indigenous electrochemically active bacteria (EAB) 19 (Stenotrophomonas maltophilia JY1, Citrobacter sp. JY3, Pseudomonas aeruginosa 20 21 JY5 and Stenotrophomonas sp. JY6) was evaluated for Cu(II) reduction on the cathodes of microbial fuel cells (MFCs). These EAB were isolated from well 22 adapted mixed cultures on the MFC cathodes operated for Cu(II) reduction. The 23 24 relationship between circuital current, Cu(II) reduction rate, and cellular electron transfer processes was investigated from a mechanistic point of view using X-ray 25 photoelectron spectroscopy, scanning electronic microscopy coupled with energy 26 dispersive X-ray spectrometry, linear sweep voltammetry and cyclic voltammetry. 27 JY1 and JY5 exhibited a weak correlation between circuital current and Cu(II) 28 reduction. A much stronger correlation was observed for JY3 followed by JY6, 29 demonstrating the relationship between circuital current and Cu(II) reduction for 30 31 these species. In the presence of electron transfer inhibitors (2,4-dinitrophenol or rotenone), significant inhibition on JY6 activity and a weak effect on JY1, JY3 and 32 JY5 was observed, confirming a strong correlation between cellular electron transfer 33 34 processes and either Cu(II) reduction or circuital current. This study provides evidence of the diverse functions played by these EAB, and adds to a deeper 35 understanding of the capabilities exerted by diverse EAB associated with Cu(II) 36 reduction. 37

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- 40 reduction; electron transfer inhibitor
- 41

#### 42 **1 Introduction**

Microbial fuel cells (MFCs) are emerging as new, sustainable and effective 43 technologies for the recovery of heavy metals from waste and wastewater [1]. 44 Among diverse heavy metals, Cu(II), which is common in the electroplating and 45 mining industries [2-3], has attracted significant attention due to the potential for its 46 47 recovery and simultaneous wastewater detoxification. The recovery of Cu(II) with abiotic cathode MFCs has been demonstrated over a wide range of operating 48 conditions and cell architectures [4-9]. However, abiotic cathodes often require the 49 use of costly noble or non-noble co-catalysts in addition to an acidic medium. 50 Therefore, the development of biocathodic MFCs, which are based on the catalysis 51 of self-regenerating electrochemically active bacteria (EAB) under a neutral 52 53 environment, may provide a sustainable and green alternative to abiotic systems. Biocathodic MFCs avoid the use of expensive materials and toxic organic reagents, 54 and reduce the consumption of energy and acids [10-14]. Biocathodes are also able 55 to reduce electrode overpotentials, the production of sludge, and the overall cost and 56 the maintenance of MFCs. MFCs, utilizing mixed cultures, have been shown to be 57 efficient in the recovery of Cu(II) from mixed metal influent and also a promising 58 59 system for the synthesis of copper [15]. In contrast, MFCs with biocathodes operated with pure cultures have been used to study specific EAB and their electrochemical 60 performance [10-14]. The exogenous EAB of Shewanella is one of the few examples 61 which is able to reduce a metal (Cr(VI)) in MFCs [16-17]. 62

Microorganisms possess endurance to aqueous Cu(II) through a variety of 63 mechanisms due to their intrinsic abilities and habitat sites [18-20]. Therefore, well 64 adapted microorganisms could provide new insights with regards to heavy metal 65 reduction in MFCs. A limited number of microorganisms exhibiting efficient rates of 66 Cu(II) reduction have been cultivated under Cu(II)-adaptive conditions [20-24]. 67 68 Similar considerations, in principle, hold true for their use in cathodic reductive environments in MFCs, under which the activities of different indigeneous EAB may 69 exhibit various Cu(II) reduction rates. In parallel, EAB immobilized on the surface 70 of cathodes may also be able to utilize cathodic electrons for their internal 71 metabolism [11,25]. The operation of MFCs at different concentrations of Cu(II) in 72 the catholyte is expected to clarify the correlation between the circuital current and 73 the rate of Cu(II) reduction associated with the use of isolated EAB. 74

75 The bacteriological reduction of Cu(II), in the absence of a circuital current, is believed to be associated with cellular electron transfer processes through the 76 cytoplasmic membrane and with the flux of protons through ATP-synthase [26-27]. 77 78 Such mechanisms can be unravelled using cellular electron transfer inhibitors, such 79 as 2,4-dinitrophenol (DNP) and rotenone ( $C_{23}H_{22}O_6$ ). DNP dissipates the proton motive force associated with cellular electron transfer processes, and decreases 80 hydrogen production by inhibiting the ATP synthesis by photophosphorylation, in 81 the absence of a circuital current [28]. In contrast, rotenone inhibits the activity of 82 NADH-dehydrogenase and blocks the reduction of U(VI) by facultative anaerobic 83 bacteria in the absence of a circuital current [29]. The utilization of rotenone and 84 DNP, in the presence and absence of a circuital current passing through the EAB 85

cathodes of MFCs is thus expected to establish whether the circuital current and
 Cu(II) reduction are associated with cellular electron transfer processes in the EAB
 species.

89 In this study, we elucidate from a mechanistic point of view the performance and impact of four indigenous different EAB on Cu(II) reduction in MFCs. The EAB 90 91 tentatively identified as Stenotrophomonas maltophilia JY1, Citrobacter sp. JY3, Pseudomonas aeruginosa JY5 and Stenotrophomonas sp. JY6, were isolated from 92 well adapted mixed cultures grown on the surface of MFC cathodes operated for 93 Cu(II) reduction [15]. X-ray photoelectron spectroscopy (XPS), scanning electronic 94 95 microscopy (SEM) coupled with energy dispersive X-ray spectrometry (EDS), linear sweep voltammetry (LSV) and cyclic voltammetry (CV) were used to investigate the 96 97 effect of circuital current on the speciation of the deposited copper, the morphologies 98 of the cathodes and the cathodic redox reactions for each EAB species. The 99 relationship between circuital current, the rate of Cu(II) reduction and the cellular electron transfer processes associated with each of the four EAB were clarified 100 through the system response to a step change of Cu(II) concentration in the catholyte 101 and reaction mechanisms elucidated in the presence or absence of DNP or rotenone 102 103 electron transfer inhibitors.

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#### 105 2 Materials and Methods

#### 106 2.1 EAB isolation, incubation and identification

107 Bacterial isolates were obtained from mixed culture cultivated in the Cu(II)-reduced biocathodes of MFCs [15]. The isolation and incubation processes 108 109 are detailed in the Supplementary Material (SM). The DNA of these EAB isolates 110 was extracted using a Oubit2.0 DNA kit (Sangon Biotech (Shanghai) Co. Ltd., China) according to the manufacturer's procedure. The 16S rRNA gene was amplified by 111 112 PCR using universal primers 518 F (5' CAGAGTTTGATCCTGGCT3') and 1540R (5'AGGAGGTGATCCAGCCGCA3'), as described in SI. The sequence data were 113 compared with the GenBank database using the Blast server at NCBI 114 (http://www.ncbi.nlm.nih.gov/BLAST/) to accurately identify the bacterial strains. 115 All tests were performed in duplicate. 116

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#### 118 2.2 MFC reactors setup and operation

Identical two-chamber MFCs with cylindrical chambers 4.0 cm long by 3.0 cm 119 in diameter were used in all experiments. The anodic and cathodic chambers were 120 121 separated by a cation exchange membrane (CEM) (CMI-7000 Membrane International, Glen Rock, NJ) with a projected surface area of 7.1 cm<sup>2</sup>. Both the 122 anode and cathode were filled with graphite felt  $(1.0 \text{ cm} \times 1.0 \text{ cm} \times 0.5 \text{ cm}, 8 \text{ pieces}, 1.0 \text{ cm} \times 0.5 \text{ cm})$ 123 Sanye Co., Beijing, China) and carbon rods were used as current collectors in both 124 anode and cathode. For each of the duplicate reactors three replicate experiments 125 were performed. The MFCs were operated at a fixed external resistance of 510  $\Omega$ . 126

The anodes were inoculated with suspended bacteria collected from a previous acetate-fed MFC reactor and an equivalent volume of nutrient solution containing acetate (1.0 g/L) was added [30-31]. The cathodes were fed using the same medium,

except acetate was replaced by NaHCO<sub>3</sub> (10 mg/L), with further addition of Cu(II) 130 (5 mg/L) to the cathodic chambers. Cathodes were quantitatively inoculated with the 131 isolates with a total of  $3 \times 10^8$  colony forming unit. The analyte and catholyte were 132 sparged with N<sub>2</sub> gas for 15 min prior to adding the solutions into the electrode 133 chambers. No measurable Cu(II) in the anolyte and acetate in the catholyte were 134 135 observed during each cycle operation, excluding the possibility of Cu(II) and acetate diffusion between the chambers, although the retention of Cu(II) on the ion 136 exchange membrane could not be precluded [15]. Other catholyte and operational 137 conditions are described in SM. 138

The performance of the biocathodes was evaluated against four control experiments, including (i) the operation of the reactors under the open circuit condition (OCC), (ii) the operation with a closed circuit but without the inoculation of the EAB (abiotic control), (iii) the cathodes covered with the EAB but in the absence of Cu(II), and (iv) the cathodes tested in the absence of both EAB and Cu(II). The last two control experiments were used to evaluate the CVs performance of the cell.

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### 147 2.3 Measurement, analysis and calculation

The circuital current, dissolved oxygen, biomass, organics and Cu(II)
concentration were determined according to the methodology reported in SM. The
rate of Cu(II) removal and charge distribution were calculated as detailed in SM.

Maximum power was obtained by running LSV at a scan rate of 0.1 mV/s 151 [30-31]. Power and current densities were normalized to the projected surface area of 152 153 the membrane. EAB cathode redox behavior was studied using CV (CHI 650, 154 Chenhua, Shanghai). The potential was scanned between -0.36 V and +0.46 V (vs SHE) at a scan rate of 1.0 mV/s using a standard three-electrode arrangement with the 155 biocathode as the working electrode, platinum plate as the counter electrode, and 156 Ag/AgCl as the reference electrode. One-way ANOVA in SPSS 19.0 was used to 157 analyze the statistical variation of the data, and all of the data indicated significance 158 levels of p < 0.05. 159

160 The surfaces of the Cu-laden biomass were analyzed by X-ray photoelectron 161 spectroscopy (XPS, Thermo Fisher Scientific, ESCALAB 250, US) with a Mono Al 162 K $\alpha$  X-ray source (1486.6 eV of photons). The X-ray source was run at a reduced 163 power of 150 W. The morphology of the electrodes after Cu(II) reduction were 164 examined with a SEM (QUANTA450, FEI company, USA) equipped with an EDS 165 (X-MAX 20 mm<sup>2</sup>/50 mm<sup>2</sup>, Oxford Instruments, UK) according to the method 166 described previously [15,30].

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## 168 **3 Results and Discussion**

### 169 3.1 EAB Isolation

Four EAB were successfully isolated from well adapted mixed cultures grown on
the surface of MFC cathodes operated for Cu(II) reduction [15] (Table S1). All
bacteria were gram-negative and major opportunistic to facultative or survived from

an anaerobic environment. JY1 matching Stenotrophomonas maltophilia, is known to

and has never been reported in MFCs. JY3 closely related to Citrobacter sp., is able to remove Cu(II) in a medium of  $SO_4^{2-}$  through the formation of CuS precipitate in the absence of cathodic electrons [20]. It has been isolated previously from anodic MFC biofilms in the absence of Cu(II) [32]. JY5 matching Pseudomonas aeruginosa, is able to remove Cu(II) in the absence of cathodic electrons [21]. It has been reported in

180 either denitrification autotrophic biocathodes of MFCs [33], or in MFCs operated for 181

convert Cu(II) into Cu(0) on the cell surface, in the absence of cathodic electrons

[22-24]. S. maltophilia has been isolated previously from a copper polluted area [22],

the degradation of phenol and glycerol through mediated electron transfer [34-37]. 182

Finally, JY6 corresponding to Stenotrophomonas sp., can tolerate high concentrations 183 of metals including Ag, Cu, Cd, Hg and Mn [38-39] and has been isolated previously 184

185 from a wide range of environments.

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187 3.2 EAB activity assessment

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### Here Fig. 1

All four EAB played a significant role in the reduction of Cu(II) in the MFCs and 189 displayed rates of Cu(II) reduction higher than those found in both OCC and abiotic 190 CCC controls (Fig. 1A). Previous studies with mixed culture biocathodes using 191 192 nitrobenzene, pentachlorophenol, Cr(VI), Co(II) or Cu(II) as an electron acceptor [31,40-44] support these findings. The Cu(II) reduction rates in the range  $(0.82 \pm 0.05)$ 193  $-1.11 \pm 0.01$  mg/L/h) were close to those found in mixed cultures (1.07 ± 0.01 194 mg/L/h) at the same Cu(II) concentration (5 mg/L) [15], reflecting the robust capacity 195 196 of these isolates for reducing Cu(II).

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### Here Fig. 2

198 The results in Figs. 1 and 2 show that the EAB catalytic process produced higher circuital currents (Fig. 1B), open circuit potentials (Fig. 2A and B) and maximum 199 power production (Fig. 2C and D) in comparison with the values under abiotic control, 200 irrespective of the Cu(II) concentrations of 5 mg/L (Figs. 1, 2A and 2C) and 20 mg/L 201 (Figs. 1, 2B and 2D). The background circuital current, voltage output, power 202 production and electrode potential in the absence of Cu(II) (Figs. 1B, 2A and 2C, and 203 Fig. S1) were attributed to the reduction of residual dissolved oxygen [15,43]. 204

Higher circuital currents and rates of Cu(II) reduction were observed by 205 increasing the concentration of Cu(II) in the cathodic chamber from 5 mg/L to 20 206 207 mg/L (Fig. 1). This effect was more significant for JY3 and JY6, demonstrating the 208 significant impact of circuital current on the rate of Cu(II) reduction for these two species. Conversely, the circuital currents observed with JY1 and JY5 were 209 insignificant (Fig. 1B), in relation to the significant increase observed in the rate of 210 Cu(II) reduction (Fig. 1A). The reduction of Cu(II) can therefore be ascribed to the 211 effect of the abiotic cathodes only and to the effect of the individual JY1 or JY5 212 bacteria in response to the increase in Cu(II) concentration (Fig. 1A). These results 213 214 illustrate the weak correlation between circuital current and the rate of Cu(II)

reduction for both JY1 and JY5, a result that will be further supported by the XPS 215

analyses.

The majority of the EAB cathodes displayed significant overshoots in voltage 217 output and power density in comparison with the abiotic controls, at a Cu(II) 218 concentration of 20 mg/L (Fig. 2B and D). This suggests that the demand for electrons 219 in these experiments exceeded the rate of electrons supplied by microbial activity, and 220 221 this resulted in the depletion of electrons and ions in the catholyte. Such observations 222 have also been reported in other MFC studies using  $O_2$  as electron acceptor [45-47]. The cathode with bacterial JY3 displayed the highest performance with the highest 223 circuital current (Fig. 1B) and most importantly without displaying an overshoot in 224 power density, indicating that the microbial activity with this species was significant. 225

The cathodic potentials of all EAB were found to vary much more than the anodic potentials, over the current density range investigated (Fig. S2A, initial Cu(II) of 5 mg/L; Fig. S2B, initial Cu(II) of 20 mg/L). In consequence, the performance of the MFCs was controlled by the reduction of copper at the biocathode, rather than the microbial phenomena that occurred at the anode. The presence of the EAB at the cathode, therefore, had a significant impact on the propensity of the cathodes for electricity generation and voltage output.

Increasing the concentration of Cu(II) in the catholyte diverted a larger fraction 233 of electrons from the anodic oxidation of organics towards the reduction of Cu(II), 234 235 with JY6 showing the highest value among the different EAB ( $11.2 \pm 0.1\%$  from a total of 2.9 C and at a Cu(II) of 20 mg/L). The remaining  $0.4 \pm 0.0\%$  was used for 236 oxygen reduction,  $52.3 \pm 0.0\%$  for organics production, and  $33.5 \pm 0.0$  for biomass 237 growth (Table S2). A large fraction of the cathodic electrons from  $2.6 \pm 0.1\%$  for JY6 238 239 up to  $43.2 \pm 0.4$  for JY3 were involved in other parallel reactions or unknown 240 processes.

The abiotic cathodes displayed reductive peak potentials of 0.033 V at a Cu(II) 241 of 5 mg/L (Fig. S3A and Table S3) and 0.037 V at 20 mg/L (Fig. S3B and Table S3), 242 reflecting the positive shift of reductive peak potential at a higher Cu(II) 243 concentration. Reductive onset potentials for all EAB cathodes were more positive 244 than the abiotic controls (0.272 - 0.297 V vs. 0.264 V at a Cu(II) of 5 mg/L and245 0.278 - 0.328 V vs. 0.267 V at a Cu(II) of 20 mg/L) (Table S3) and the shift was 246 more significant for the reductive peak potentials of all EAB cathodes with respect 247 to the abiotic controls  $(0.040 - 0.063 \text{ V vs. } 0.033 \text{ V at a Cu(II) of 5 mg/L}, 0.043 - 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 - 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 - 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 - 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 - 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 - 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 - 0.043 \text{ V at a Cu(II) of 5 mg/L}, 0.043 \text{ V at$ 248 0.079 V vs. 0.037 V at a Cu(II) of 20 mg/L). These results, in combination, 249 demonstrate the varying degree of influence exerted by each EAB on the catalytic 250 251 activity toward Cu(II) reduction, which is consistent with the effect of EAB on the 252 reduction of Co(II) and chloramphenicol on mixed culture cathodes [43,48]. The JY1 and JY5 bacteria exhibited much lower reductive onset potentials than JY3 and JY6, 253 implying a weaker interaction between cathodic electrons and the reduction of Cu(II) 254 through the mediation of these two species. 255

The reductive peak currents, at a Cu(II) concentration of 5 mg/L, observed on the cathodes covered by the EAB were lower than values registered with the abiotic cathode (0.973 mA) with the exception of JY6 (Table S3), suggesting some degree of mass transfer inhibition for Cu(II) due to the contact between the EAB and the electrode surface [47]. At a higher concentration of Cu(II) (20 mg/L), the mass transfer inhibition became less significant, as expected. Similar observations have been reported for the reduction of other electron acceptors such as Cr(VI) and Co(II) in mixed culture cathodes [41,43].

The lack of a significant difference observed between CVs of the abiotic controls and the biotic cathodes in the absence of Cu(II) (Fig. S4), in concert, reflected the importance of Cu(II) ions on the occurrence of reduction reactions on the EAB, despite the expected variability due to the redox species present on the surfaces of these EAB cells (Insets in Fig. S4).

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## 270 *3.3 Electrode morphology and product analysis*

SEM examination showed that the EAB covered the surface of the cathodes only
sparsely (Fig. S5A, D, G, and J). EDS analysis confirmed the presence of Cu
precipitates on the surfaces of EAB cells (Fig. S5B, E, H and K) while no Cu, or very
little Cu precipitate, was observed on the bare surface of the electrodes (Fig. S5C, F, I
and L). EDS detection of carbon, oxygen, sodium and phosphorus was attributed to
the cellular components of EAB on the electrodes, while gold was associated with the
sample pretreatment.

XPS analyses (Fig. S6) indicated that the whole Cu2p region comprises 2p1/2
and 2p3/2 peaks, while only the Cu2p3/2 peaks could be used for the assignment of
copper chemical states [49]. The abiotic control reported the exclusive presence of
Cu(0) (Fig. S6A), shown with its characteristic peak at the Cu2p3/2 region of 932.4
eV [49], while under OCC only adsorbed Cu(II) was observed (Fig. S6B), as expected.
This result clearly demonstrates the importance of the effect of circuital current on the
reduction of Cu(II) to Cu(0) with the abiotic cathodes.

285

### Here Fig. 3

The XPS analysis on the EAB cathodes in the presence of a circuital current (Fig. 286 287 3), reported the exclusive presence of Cu(0), while under OCCs, both Cu(0) and Cu(II) were observed for JY1 (Fig. 4A) and JY5 (Fig. 4C), and only Cu(II) for both JY3 (Fig. 288 4B) and JY6 (Fig. 4D). JY5 had a net Cu(0) production of  $1.81 \pm 0.11$  mg and JY1 289 exhibited a  $1.45 \pm 0.05$  mg, both of which were nearly equivalent to the sum of  $1.17 \pm$ 290 0.07 mg in the abiotic controls and  $0.70 \pm 0.09$  mg (JY5) or  $0.44 \pm 0.03$  mg (JY1) in 291 292 the absence of a circuital current (Table S4). These results in concert confirm the weak correlation between circuital current and the rate of Cu(II) reduction for both 293 294 JY1 and JY5, which is consistent with the results shown in Fig. 1. The observation of 295 Cu(0) in JY1 in the absence of a circuital current is consistent with previous studies, where S. maltophilia converted Cu(II) into Cu(0) under facultative conditions [22-24]. 296

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299 3.4 Effect of electron transfer inhibitors (rotenone and DNP)

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Here Fig. 4

In the presence of either rotenone or DNP, the rate of Cu(II) reduction (Fig. 5A) 301 or the circuital current (Fig. 5B) registered in the MFCs with JY1 or JY5 EAB, 302 changed little, suggesting a weak correlation between the cellular electron transfer 303 processes and either Cu(II) reduction or the circuital current. Conversely, the cathode 304 with the JY6 species, returned an appreciable decrease in both the rate of Cu(II) 305 306 reduction and circuital current, suggesting a strong interaction between the cellular electron transfer processes in JY6 and either Cu(II) reduction or circuital current. This 307 strong interaction observed with JY6 rather than with JY1, JY3 and JY5, was also 308 supported by the apparent decrease in cathodic electrons used for Cu(II) reduction in 309 the former, in the presence of rotenone or DNP, while little change in was observed 310 with the other three EAB (Table S2). The negative effects of rotenone or DNP 311 312 electron transfer inhibitors on JY6 was also reflected by a decrease in the voltage 313 output (Fig. S7A and B), maximum power density (Fig. S7C and D) and cathode potential (Fig. S7E and F), in addition to the negative shifts observed in reductive 314 onset potential, reductive peak potential and reductive peak current (Table S3; Fig. 315 **S8**). In contrast, little change on the above characterization parameters was observed 316 for JY1, JY3 and JY5. These results further support the weak dependence between the 317 318 cellular electron transfer processes and either Cu(II) reduction or circuital current for JY1, JY3 and JY5, and confirm the significant correlation observed between the 319 320 cellular electron transfer processes and either Cu(II) reduction or circuital current for JY6. The findings of this study are summarized graphically in Fig. 6. 321

#### 322

#### Here Fig. 6

#### 323 4 Conclusions

324 Metal reduction in MFCs is known to be dependent upon the availability of cathodic electrons and upon the composition of bacterial communities on the 325 biocathodes [11,15,40]. On the basis of the results obtained with indigenous EAB JY1, 326 JY3, JY5 and JY6 isolated from well adapted mixed cultures grown on the surface of 327 MFC cathodes used for Cu(II) reduction, this study demonstrates the close correlation 328 among circuital current, the rate of Cu(II) reduction and the cellular electron transfer 329 processes for JY6, and the weak dependence of these for JY1 and JY5 (Fig. 6). We 330 therefore suggest the possibility of a mechanism involving direct electron transfer 331 from the surfaces of the cathodes to JY6, followed by the subsequent cellular electron 332 transfer to Cu(II) for its reduction. This mechanism is not reflected in JY3, which only 333 334 exhibited a close correlation between circuital current and the rate of Cu(II) reduction 335 (Fig. 6). In summary, this study provides an evidence of the diverse functions played 336 by these EAB species in the mixed cultures and the corresponding effect on MFC characterization parameters. The results in this study add to a deeper understanding of 337 the capabilities exerted by diverse EAB associated with Cu(II) reduction, and provide 338 new insights into the potential application of metallurgical biocathode MFCs for Cu(II) 339 recovery at industrial scale. 340

341

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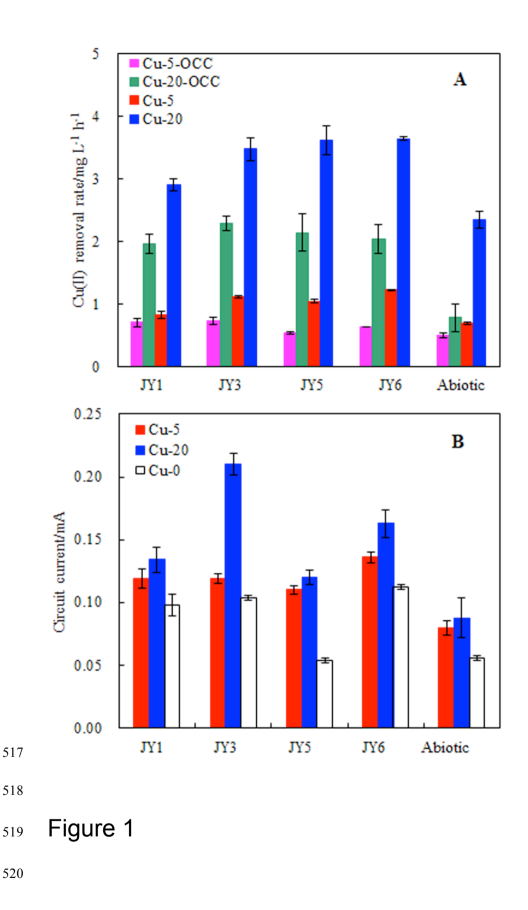
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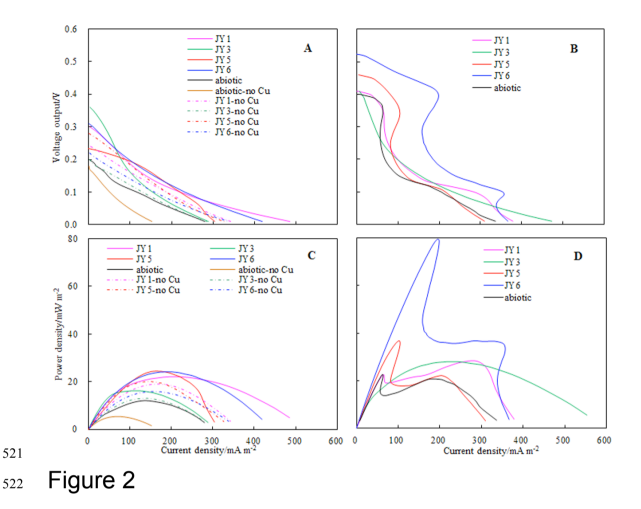
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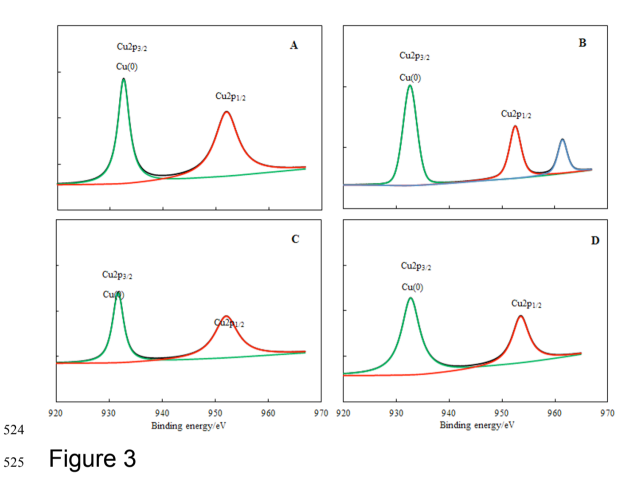
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## 502 Figure captions

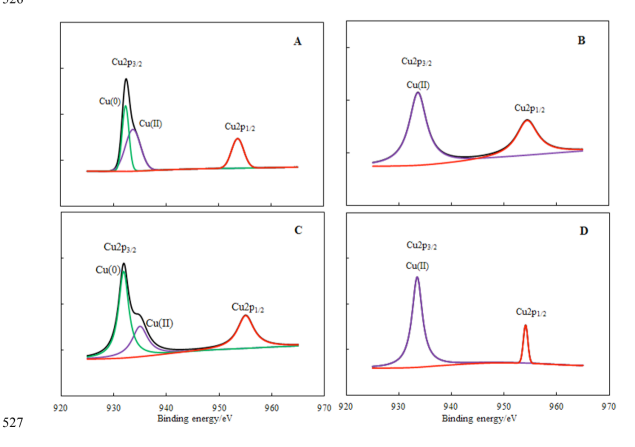
- Fig. 1 Comparison of Cu(II) removal rate (A) and circuital current (B) with various
  EAB at an initial Cu(II) of either 5 mg/L or 20 mg/L.
- Fig. 2 Voltage output (A and B), power density (C and D) as a function of current
  density with various EAB at an initial Cu(II) of 5 mg/L (A and C) or 20 mg/L (B and
  D).
- Fig. 3 XPS spectra of Cu precipitates on the cathodes catalyzed by EAB of JY1 (A),
  JY3 (B), JY5 (C) or JY6 (D) (initial Cu(II): 20 mg/L; 20 batch cycles).
- 510 Fig. 4 XPS spectra of Cu precipitates on the cathodes catalyzed by EAB of JY1 (A),
- 511 JY3 (B), JY5 (C) or JY6 (D) in the absence of circuital current (initial Cu(II): 20
- 512 mg/L; 20 batch cycles).
- **Fig. 5** Comparison of Cu(II) removal rate (A) and circuital current (B) in response to the inhibitor of rotenone or DNP (initial Cu(II): 20 mg/L).
- 515 **Fig. 6** Summary of correlation between circuital current, Cu(II) reduction and cellular
- electron transfer in EAB of JY1, JY3, JY5 and JY6.











528 Figure 4

