Application of Polypyrrole/Humic Acid Composite Electrode for Copper Ion Extraction from Drinking Water

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Copper is a metal commonly used to transport water systems due to its stability and advantageous properties. At the same time, corrosion may be a problem affecting these systems, for instance, causing the leaching of metal (Cu (II)) in water. In this study we have applied an electrochemical method for extracting this element. The use of humic acid and of a conductive polymer (polypyrrole) in a PIGE/HA/PPy modified electrode was evaluated for the electrochemical extraction of Cu(II) from drinking water. Synthetic water samples were contacted with Cu(0) pipes during 24 h (leaching yielded Cu(II)). Modified electrodes were prepared by mechanical transfer of humic acid (HA) to the base electrode (PIGE) and then depositing a polypyrrole (PPy) coating by potentiostatic electropolymerization. The modified electrode was subsequently utilized for Cu(II) electrochemical extraction from water samples using a chronocoulometric method, with a removal rate of up to 72%. Thus, the feasibility of decreasing Cu(II) concentration in drinking water systems was proven, corroborating the usefulness of the PIGE/HA/PPy surface modified electrode. Besides, the application in systems of this kind can be further improved by considering the different variables involved, e.g. PIGE electrode area, HA amount, PPy thickness, etc.

Keywords: Humic Acid, Ion Extraction, Drinking water, Modified Electrode

1. INTRODUCTION

Metallic Cu in contact with drinking water is oxidized by dissolved oxygen forming an adherent layer of cuprite (Cu$_2$O), where the inner film is thin and the outer film porous, facilitating ion
and molecule diffusion [1-4]. Cuprite remains in contact with the surface of metallic copper [5]. Oxidation of cuprous to cupric ion also occurs in Cu oxidation [6].

As for inorganic compounds present in drinking water and their effect on Cu corrosion [7,8], it is known that species such as bicarbonate, carbonate and dissolved carbonic acid, ammonium, sulfate and chloride are the principal inorganic species affecting the corrosion process. Variables such as time of stagnation, age of the pipes, among others, also influence the process of Cu release. Closely related to this, there is agreement that pH would be the most significant variable in Cu corrosion. Corrosion rates are observed to increase with pH decrease, [6, 9, 10]. At pH below 5, Cu corrodes quickly and evenly. At values above 9, Cu is usually protected. Between pH 5 and 9, if no protective film on Cu surface exists, localized corrosion is likely to occur, which is also significantly affected by this pH factor [11].

Controlling and decreasing Cu(II) concentration in drinking water, involves different types of treatments or strategies. The most used one is the adjustment of parameters such as pH, alkalinity and corrosion inhibitors that are regulated for various drinking water systems, usually working ex situ [12]. For low concentrations of Cu(II) present, the use of more sensitive electrochemical techniques, more sensitive than methods mentioned above, may be useful.

Conducting polymers are multifunctional materials, as Heeger mentions [13], these "offer a unique combination of properties not available from other known materials, which opens the possibility of use in a wide variety of applications". These properties make them useful for solid-phase microextraction, SPME, applications [14], where polypyrrole and polyaniline (PANI) [15, 16] were the most popular conductive polymers used for SPME fibers. These conducting polymers have p- or n-type doping-undoping possibility, a feature of most of these polymers that has been used by our research group for ion extraction [17, 18 to 20]. The results indicate that in fact the polymer could be used as extractant for arsenic, copper or mercury. Consequently, a simple electrode of rapid preparation has been recently developed using humic acid (HA) and polypyrrole films [18]. The results revealed that the presence of HA did not affect the electrical properties of the system, but indeed provoked morphological changes in the polymer making it more granular. In this context, other variables such as the nature and size of the anion enable the obtention of a surface displaying ion exchange properties [21, 22]. Chloride and acetate doped obtained polymers showed weak adhesion on the electrode surface, being unstable at high temperatures. Another important variable is the synthetic route used for electropolymerization. The potentiostatic route allows the attainment of a more highly cross-linked polymer that would favor the inclusion of anions such as arsenate [17]. The prepared modified electrode was evaluated utilizing As (V) solutions because this kind of polymer may undergo p-doping/undoping processes [18], which account for its good extractant abilities toward the analyte. Since humic acids are negatively charged, it would be expected that the presence of HA in the modified electrode was independent of its origin, its acid-base features and in addition of a very weak character. However, the differences between the two modified electrodes having HA in their composition enables postulating that interactions would be not so weak and an effect of the existing functional groups do exists. In this regard, as proposed for AsO$_4^{3-}$ functional groups such as phenol [23, 24], could interact with Cu(II) positive formal charge in the CuCl$_4^{2-}$ complex. This proposal is based on the fact that complexes with coordination number less than 6 may develop exchange
mechanisms associated to positively charged metal centers [24], as well as to the existence of phenolates in the HA. In this context, other variables, e.g. anion nature and size, exist that permit the obtention of a surface with ion-exchange properties [21,22]. On the other hand, polymers doped with chloride and acetate anions showed weak adhesion on the electrode surface, being unstable at high temperatures. Another important variable is the synthetic route used for the electropolymerization. The potentiostatic approach allows for a greater cross-linked polymer, which would favor the inclusion of anions such as arsenate [17]. The modified electrode prepared was evaluated utilizing As(V) solutions because this type of polymer can undergo p-doping – undoping processes [18], which accounts for its good extractant abilities toward the analyte. Since humic acids are negatively charged, it would be expected that the presence of HA in the modified electrode was independent of its origin, its acid-base features and in addition of a very weak character. However, the differences between the two modified electrodes bearing HA in their composition enable postulating that interactions would be not so weak and an effect of the existing functional groups do exists. In this regard, as proposed for AsO$_4^{3-}$ functional groups such as phenolics [23, 24], could interact with Cu(II) positive formal charge in the CuCl$_4^{2-}$ complex. This proposal is based on the fact that complexes with coordination number less than 6 may develop exchange mechanisms associated to positively charged metal centers [24] along with the existence of phenolates in the HA.

Therefore, considering problems of drinking water systems concerning to Cu(II) content as well as the described advantages of HA and PPy modified electrodes, a study of their applicability to drinking water samples is proposed in this work. At this earlier stage synthetic samples will be employed. The study also embraces the effect of potentiostatic polymer preparation on the efficiency of analyte extraction, considering that, although not unique, this is one of the relevant variables in the morphology of the modified electrode, consequently it may have a significant impact on its extraction capacity.

2. MATERIALS AND METHODS

2.1 Samples

2.1.1 Humic Substances

Humic acid (Sigma-Aldrich) was purified by precipitation as follows. A suitable concentrated solution was adjusted to pH 1 with HCl. The obtained precipitate was filtered off and then re-dissolved in NaOH. The precipitation was repeated and a second precipitate was collected, washed, air-dried, and used.

2.1.2 Drinking water

The synthetic water was prepared in a horizontal laminar flow hood, Factomet, model H24242, provided with di-octyl phthalate (DOP) aerosol penetration certificate and high efficiency particulate
air (HEPA) filters. Alkalinity adjustment was accomplished with NaHCO₃, while for pH gaseous carbon dioxide and/or oxygen (O₂, 32.0 g mol⁻¹) flushed directly into the solution was used. A suitable amount of sulfate, nitrate and sodium chloride was next added, to follow the composition quoted in Table 1. The different synthetic water samples prepared were poured into a Cu pipe system and kept during 24 h. This process is called stagnation. Subsequently to this process, triplicate samples in three different pipe cuts were collected [25].

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Drinking water</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>pH</strong></td>
<td>Pre-stagnation</td>
<td>6.5±0.0</td>
<td>6.5±0.1</td>
<td>7.8±0.0</td>
<td>7.8±0.0</td>
</tr>
<tr>
<td>Alkalinity-CaCO₃ (mg L⁻¹)</td>
<td>54.5±0.1</td>
<td>308.7±1.1</td>
<td>53.3±0.2</td>
<td>299.0±2.3</td>
<td></td>
</tr>
<tr>
<td>O₂ (%)</td>
<td>6.4±0.2</td>
<td>5.1±0.1</td>
<td>6.9±0.1</td>
<td>6.7±0.0</td>
<td></td>
</tr>
<tr>
<td>Anions (mg L⁻¹)</td>
<td>SO₄²⁻</td>
<td>257.9±3.2</td>
<td>251.9±2.1</td>
<td>259.5±5.0</td>
<td>261.0±3.0</td>
</tr>
<tr>
<td>Cl⁻</td>
<td>103.5±0.2</td>
<td>102.3±0.2</td>
<td>104.0±0.2</td>
<td>105.8±0.2</td>
<td></td>
</tr>
<tr>
<td>NO₃⁻</td>
<td>12.1±0.2</td>
<td>14.5±0.2</td>
<td>13.0±0.2</td>
<td>12.1±0.2</td>
<td></td>
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<tr>
<td>Cu²⁺</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
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<tr>
<th>Characteristics</th>
<th>Drinking water</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>pH</strong></td>
<td>Post-stagnation</td>
<td>6.8±0.1</td>
<td>7.0±0.1</td>
<td>7.8±0.0</td>
<td>8.2±0.0</td>
</tr>
<tr>
<td>Alkalinity-CaCO₃ (mg L⁻¹)</td>
<td>56.6±0.9</td>
<td>313.4±6.4</td>
<td>55.9±0.9</td>
<td>311.8±3.2</td>
<td></td>
</tr>
<tr>
<td>O₂ (%)</td>
<td>4.6±0.3</td>
<td>2.5±1.5</td>
<td>4.9±0.3</td>
<td>4.6±0.3</td>
<td></td>
</tr>
<tr>
<td>Anions (mg L⁻¹)</td>
<td>SO₄²⁻</td>
<td>255.4±2.6</td>
<td>259.8±3.8</td>
<td>259.5±0.4</td>
<td>264.8±2.1</td>
</tr>
<tr>
<td>Cl⁻</td>
<td>102.2±0.6</td>
<td>104.4±1.9</td>
<td>103.9±1.6</td>
<td>106.3±0.6</td>
<td></td>
</tr>
<tr>
<td>NO₃⁻</td>
<td>15.4±4.2</td>
<td>16.3±3.1</td>
<td>18.1±3.4</td>
<td>12.5±0.1</td>
<td></td>
</tr>
<tr>
<td>Cu²⁺ (mg L⁻¹)</td>
<td>1.0±0.4</td>
<td>2.62±0.51</td>
<td>0.99±0.2</td>
<td>1.47±0.1</td>
<td></td>
</tr>
</tbody>
</table>

2.2 Chemical characterization

2.2.1 Humic Substances

Functional groups determination was accomplished for total and carboxylic acidity, while phenolic acidity was obtained by difference [26, 27].

2.2.2 Drinking water

All pre- and post- stagnation (24 h) drinking water samples were characterized. Separation and determination of SO₄²⁻, Cl⁻ and NO₃⁻ was conducted by chromatography and a conductivity detector (Dionex ICS-1500 Ion Chromatograph) using NaOH as mobile phase at 1.5 mL min⁻¹ flow rate [28].
Cu (II) content was quantified after 24 h stagnation by atomic absorption spectrometry (GBC Scientific Equipment, Model SensAA), using EPA 7000B Method for traces. The samples were digested with 2-10% \( \text{HNO}_3 \) (m/m).

Alkalinity was determined by acid-base titrimetry following EPA 310.1 method [30]. pH was measured according to EPA 150.1 method and percent dissolved \( \text{O}_2 \) following EPA 360.1 method [30].

2.3 Electrochemical experiments

2.3.1 Apparatus

All electrochemical experiments were carried out on a potentiostat system VoltaLab PGZ100 in a three-compartment/three-electrode glass cell under high purity argon atmosphere. Paraffin-impregnated graphite (PIGE) was prepared and used as working electrode. The graphite exposed surface of the electrode was mechanically regenerated prior to each experiment with emery paper (grain size 2000 and 2500), and thorough washing with double-distilled water. A Pt wire (20 cm\(^2\)) was the counter electrode. All potentials quoted in this work are referred to an Ag|AgCl reference electrode.

All electrodic surfaces were studied in drinking water and solutions containing Cu(II) in 20 mM \( \text{HCl} + 100 \text{ mM KCl} \) (solution A), that were prepared by dissolving the appropriate amount of \( \text{CuCl}_2 \) in double-distilled water according to standard procedures. Pyrrole (Sigma-Aldrich) was purified by distillation, stored at 4º C and protected from light. All other chemicals used were analytical grade (Merck). All solutions were prepared and subsequently deaerated with high purity Ar for at least 20 min.

2.3.2 Preparation of modified PIGE

As for PIGE, four electrodic surfaces were characterized by cyclic voltammetry: (1) PIGE; (2) PIGE modified with HA (Aldrich) (PIGE/HA); (3) PIGE electrochemically modified with pyrrole (PIGE/PPy); and (4) PIGE modified with HA and then electropolymerized with pyrrole (PIGE/HA-D/PPy; PIGE/HA/PPy). Humic acid was transferred and immobilized onto the electrode surface using the abrasive transfer technique described elsewhere [31].

PPy electrochemical coating was conducted on PIGE or PIGE/HA from 1.4 mM pyrrole + 50 mM \( \text{Na}_2\text{SO}_4 + 30 \text{ mM H}_2\text{SO}_4 \) solutions utilizing the potentiostatic method. To evaluate its effect on extraction efficiency, an initial potential of 0 V for 30 seconds, followed by 1 to 4 min at 1.0 V, was respectively applied.

2.3.3 Extractions

PIGE/HA/PPy electrode surface was studied in solutions containing 22.32 mM \( \text{CuCl}_2 + 20 \text{ mM HCl} + 100 \text{ mM KCl} \) (solution A, cell A) and 20 mM \( \text{HCl} + 100 \text{ mM KCl} \) (solution B, cell B) to select
the doping–undoping potentials to perform the extraction. The overall volume of electrolyte in each cell was always 10.0 mL. Each extraction was accomplished as follows. The modified PIGE/HA/PPy electrode was dipped into drinking water and potentiostatically perturbed at 0.4 V for 5 min. Subsequently the device was kept into solution B at -0.6 V for 5 min. Initially and after several extractions, the copper concentration in drinking water and solution B was determined by inductively coupled plasma/optical emission spectrometry (ICP–OES) on a Varian Liberty series II instrument.

3. RESULTS AND DISCUSSION

3.1 Chemical characterization

Acid-base humic acid characterization indicated that an important content of carboxylic (5.5 ± 0.2 mEq g⁻¹) and phenolic (1.9 ± 0.2 mEq g⁻¹) groups exist, validating thus the likely interactions with anions such as AsO₄³⁻ and CuCl₄²⁻ [18, 19], known as bridging metals.

Table 1 exhibits the results of the chemical characterization of drinking water samples pre- and post-stagnation. In general, pH and alkalinity remained without significant variation, while for O₂ (%) content a decrease occurred in all samples, probably due to its participation in corrosion processes during stagnation.

Regarding to anions, they remained nearly constant, with no significant changes. Hence, every drinking water sample maintained its original characteristics.

Table 1 demonstrates that Cu(II) exists only in post-stagnation samples, which is consistent with sample preparation, purity and quality of utilized reagents. Samples 2 and 4, which are considered highly corrosive towards Cu(0), presented the highest Cu(II) concentration. These results clearly showed that sample 2 conditions in terms of dissolved O₂, pH and alkalinity are sufficient to cause metal pipe largest corrosion.

According to results, pH variation between samples of higher alkalinity has a negative effect above pH 7 (to overcome the barrier of neutrality), while alkalinity increase would generate an increased presence of Cu(II). However, it is also evident that the combination of higher alkalinity and neutrality caused the greatest corrosion.

3.2 Electrochemical experiments

In Figure 1, the voltammetric profile of PIGE and PIGE/HA modified electrodes are shown. A slight current drop for the modified electrode is observed owing to the presence of HA at the electrode surface. This behavior means that HA incorporation into their active sites causes blockage of the surface properties of the electrode, reducing its conductivity [32]. Besides, an inflection peak at a less cathodic potential – 400 mV, due to H⁺ reduction and H₂ formation [33].
This is a consequence of the slow scan rate used in the characterization of both electrodes. This fact brings about low voltammetric response (Randles-Sevcik equation) and further growth of the diffusion layer at the surface, allowing the occurrence of redox phenomena [34].

In Fig. 2(a) and (b) voltammetric response of PIGE/HA/PPy and PIGE/PPy modified electrodes at different electropolymerization times (1 to 4 min) are depicted. As for this variable, a current increase with polymerization time is observed, but at higher times the electropolymerization response is not reproducible because of polymer fall off.

Time analysis enables stating; in general, that longer electropolymerization leads to a greater voltammetric response. This affords larger amount of PPy deposite on the electrode surface. Nonetheless, at times 3 and 4 min the above relationship no longer holds since no current increase was obtained. This may be ascribed to ions diffusion hindrance to get into the polymeric matrix to counteract the charge formed during polymer oxidation. Thus, a thicker polymer film will generate an impediment to the entry of anionic species [34].

As far as charge density of p-doping process (anionic charge) and p-undoping (cationic charge) are concerned, they are similar, suggesting a reversible p-doping process. Based on charge density saturation, owing to greater amount of PPy, and the observed charge difference between PIGE/PPy and PIGE/HA/PPy electrodes polymerized for 1 min, the latter was chosen as the optimum electropolymerization time.
Figure 2. Potentiodynamic response of PIGE/HA/PPy (a) and PIGE/PPy (b), in 20 mM HCl and 100 mM KCl at different electropolymerization time between -0.6 and 0.4 V at 0.1 V s⁻¹.

Figure 3 shows that the PIGE/ HA/Py modified electrode exhibits a higher current response than PIGE/Py, as well as greater anionic and cationic charge. PIGE previous studies, wherein Ppy was prepared using the potentiodynamic electropolymerization approach, demonstrated that the higher conductivity and voltammetric response is chiefly due to morphology changes of the polymer caused by the presence of HA in the polymeric matrix [31, 32].
The PIGE electrode surface morphology presents an irregular surface structure [32, 35] and PIGE/Ppy a nodular one [36, 37], whose porosity and roughness depend on coating thickness [37]. Therefore, the presence of HA affected both surfaces [31, 32].

HA in the PIGE/HA electrode is found as irregular aggregates of heterogeneous forms [32], affecting thus Ppy deposition on the PIGE/HA/Ppy electrode surface. This arises by the inclusion and surface blockage of the PIGE electrode high-energy sites [24, 32], generating further polymer growth, with a PPy film of less regularity and thickness at the time nucleation and electropolymerization take place [31, 32].

Hence, it can be stated that HA presence brings about a higher voltammetric response at the PIGE/HA/Ppy modified electrode versus the PIGE/Ppy one at identical polymerization time. Consequently, the surface becomes more conductive [18] and suitable for Cu(II) extraction from water synthetic solutions and real samples.

![Figure 3](image_url)

Figure 3. Comparative response of potentiodynamic electropolymerization of PIGE, PIGE/Ppy and PIGE/HA/Ppy in 20 mM HCl and 100 mM KCl during 1 min between 0.6 and 0.4 V at 0.1 Vs⁻¹.

3.3 Extractions

All Cu (II) extractions were accomplished using a PIGE/HA/Ppy electrode prepared by 1 min potentiostatic electropolymerization. Electrode characterization, Fig 3, shows that doping occurs at ca. 400 mV, while undoping proceeds at −500 mV. At an early stage of the extraction process, the
modified electrode was used in synthetic solutions, in order to assess its efficiency and reversibility after multiple successive extractions. The obtained results (not shown) exhibit high extraction accuracy, coefficient of variation 2.5%, and similar values of the transferred charge in the p-doping/undoping process, indicating that the extraction process is reversible.

Extraction results in different drinking water samples are summarized in Table 2, wherein extraction efficiency and transferred charges may be evaluated.

**Table 2.** Drinking water Cu(II) extraction and doping-undoping charges after 10 successive extractions utilizing a PIGE/HA/Ppy modified electrode.

<table>
<thead>
<tr>
<th>Drinking Water</th>
<th>Initial Cu(II) (mg/L)</th>
<th>Extracted Cu(II) (mg/L)</th>
<th>Extracted Cu(II) (%)</th>
<th>Doping charge (\times 10^{-2} \text{C})</th>
<th>Undoping charge (\times 10^{-2} \text{C})</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.64</td>
<td>0.43</td>
<td>66.93</td>
<td>1.34 ± 0.26</td>
<td>1.36 ± 0.48</td>
</tr>
<tr>
<td></td>
<td>1.35</td>
<td>0.82</td>
<td>60.64</td>
<td>1.37 ± 0.28</td>
<td>1.24 ± 0.27</td>
</tr>
<tr>
<td></td>
<td>1.12</td>
<td>0.73</td>
<td>65.28</td>
<td>1.86 ± 0.23</td>
<td>1.94 ± 0.23</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>64.28 ± 1.16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>2.06</td>
<td>1.11</td>
<td>53.88</td>
<td>1.75 ± 0.28</td>
<td>2.09 ± 0.43</td>
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<tr>
<td></td>
<td>2.77</td>
<td>1.60</td>
<td>57.71</td>
<td>3.43 ± 0.46</td>
<td>2.46 ± 0.91</td>
</tr>
<tr>
<td></td>
<td>3.04</td>
<td>1.75</td>
<td>57.57</td>
<td>1.56 ± 0.14</td>
<td>1.08 ± 0.38</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>56.39± 2.17</td>
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<tr>
<td>3</td>
<td>0.78</td>
<td>0.54</td>
<td>69.23</td>
<td>1.46 ± 0.37</td>
<td>1.23 ± 0.26</td>
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<tr>
<td></td>
<td>1.15</td>
<td>0.73</td>
<td>63.55</td>
<td>1.54 ± 0.33</td>
<td>1.66 ± 0.21</td>
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<tr>
<td></td>
<td>1.04</td>
<td>0.59</td>
<td>56.76</td>
<td>1.63 ± 0.12</td>
<td>1.39 ± 0.20</td>
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<td>63.18 ± 6.25</td>
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<tr>
<td>4</td>
<td>1.49</td>
<td>1.00</td>
<td>67.71</td>
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<td>1.40</td>
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<td>0.97 ± 0.24</td>
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<td></td>
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<td>72.59 ± 5.10</td>
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</table>

From these data and their one way Anova analysis, all extraction percentages are significantly different among them \((P <0.0001)\). The largest percentage of extraction was for sample 4 and the lowest for sample 2, precisely the one with the highest Cu(II) content.

In this regard, pH is a critical parameter. A review of Cu(II) speciation shows, from a global analysis, that a pH variation from 7.0 to 8.0 causes the Cu(II), chiefly complexed as Cu(OH)+, Cu(OH)2 and Cu(OH)3−, to increase up to 99%. This has an impact on the extraction efficiency because during doping this modified electrode incorporates anionic species in order to counteract its charge while oxidation proceeds. On this matter, background exists \([20]\) that permits assuming that as the extracted species concentration decreases, a shift of the equilibrium of the involved species would take place, originating thus the high extraction rate of the analyte under survey. The extraction has an acceptable accuracy, with coefficient of variance in the range 2 -10 % \((n = 30)\).

On the other hand, from Cu(II) mass balance accomplished in the initial drinking water and in cells A and B used for the extraction, it may be ascertained that part of the extracted Cu(II) remained
in the modified electrode, existing non-proportionality in the release during undoping, with respect to the efficiency obtained for values of the transferred charge. This behavior might be ascribed to a synergistic effect of HA + PPy, as described elsewhere [18]. In fact, as HA bears, among others, carboxylic and phenolic functional groups, this makes these species very chelating amidst natural organic substances (partly responsible for cation exchange in soils and aquifer systems), conferring them additional affinity toward ions or metallic species to be extracted [39]. These functional groups allow interaction with Cu(II) positive formal charge, analogous to what happened in arsenate extraction conducted with the same modified electrode [32]. Hence, mass transfer would be compensated by anions present in solution, enabling reversibility of the transferred charges.

In addition, it has been shown that, despite Cu(II) retention, the electrode continued functioning properly, remaining stable during the performed extractions, which permitted postulating that it would be possible to improve extraction performance by increasing the number of successive extractions using the same electrode, or investigating the influence of other variables such as supporting electrode area.

4. CONCLUSIONS

The potentiostatic HA-PPy electrode modification made possible complexed Cu(II) extraction from drinking water synthetic samples with an efficiency of up to 72%. Besides, pH and alkalinity dependence of the assayed samples was verified. Thus, the high efficiency exhibited by this electrode permits regarding it as a likely solution to corrosion problems with copper pipes employed for potable water transportation. Consequently, it is justified to attempt optimization of its design and application to real samples.

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