Tuning Electrochemical Bistability by Surface Area Blocking in the Cathodic Deposition of Copper

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ABSTRACT: We report herein a precise control of the electrochemical bistability induced by surface area changes during the cathodic deposition of copper. Small additions of 1,10-phenanthroline (Phen) in the reaction media present an intriguing effect on the global rate mainly due to the adsorption of protonated Phen. The increase of its concentration results in a shrinkage of the bifurcation (saddle-node) diagram and shifts it to less negative potentials. The dynamic bistability is verified by impedance measurements, and a negative hysteresis is clearly found. We calculated the apparent molar mass of the adsorbed molecules using surface gravimetric monitoring in the electrochemical experiments, and the results indicate that mass changes occur mainly due to the adsorption of copper from bivalent ions dissolved in the reaction media. Importantly, the adsorption of protonated Phen molecules does not show a considerable contribution in mass variations but prevents the formation of a copper course grained morphology over the surface. Imaging analysis indicates finer nodulations at the lower branch compared to the upper branch in the bistability domain. On the basis of these observations, a kinetic mechanism is proposed and a good agreement is obtained between the apparent molar mass extracted from experiments and the theoretical values. Altogether, our results contribute to a detailed physical chemical description of the nonlinear behavior, bringing new insights about this reaction and pointing out the possibility to design switchable surface electrodes by taking advantage of the bistable behavior.

INTRODUCTION

Nonlinear behavior in chemical systems far from equilibrium may appear in the form of dynamic instabilities. One very frequently observed is the bistability (BS) (i.e., two stable steady states at the same set of controlling parameters), induced by a saddle-node bifurcation, when positive feedback loops act as regulatory steps in the overall mechanism. As a result of these self-amplifying processes, a hysteretic profile arises, and consequently, chemical switches can be designed and finely tuned by the imposition of external perturbations. For spatially extended systems, the appearance of collective switching between states has been based on the bistable nature of the individuals, but it can also emerge from collective behavior even if the constituents are not bistable. This co-existence of different steady states has been verified in homogeneous and heterogeneous systems.

Electrochemical instabilities occur by the interaction of chemical properties of the solid/liquid interface (faradaic impedance) with electrical parameters of the equivalent circuit (applied voltage, current, and resistance), resulting in the formation of the so-called negative differential resistance (NDR). According to Koper, this condition is satisfied when an increase of the double layer potential promotes a decrease of the available electrode area, the electrochemical rate constant, or the interfacial concentration of the electro-active species. Examples attending to the three cases have been reported in the literature. Then, in the presence of the NDR, BS arises mainly due to the electrical autocalysis in the electrode potential as the essential positive feedback variable [N-shaped NDR (N-NDR)] or by chemical autocatalysis stemming from a reaction intermediate [S-shaped NDR (S-NDR)].

The emergence of BS has been observed in different electrochemical systems, such as the electroreduction of complexes of Ni(II) at a streaming mercury electrode, in the electroreduction Hg(I)/Hg(II) ions coupled with convection, in the anodic dissolution of V electrodes in acidic media, in the electro-oxidation of small organic molecules, and so forth. Overall, oscillations appear along with BS depending on the controlling parameters. It reveals a close correspondence between both nonlinear behaviors and corroborates with the cross-shaped phase diagram analysis. The confirmation of the BS in these systems, under
potentiostatic control, for instance, can be confirmed by impedance measurements in which zero imaginary impedance occurs in zero frequency of perturbation. 41 Although, detecting more than two branches in electrochemical systems are very rare, tristability has been observed as well.33,42

Considering the requirements to obtain a NDR, Nakanishi et al.45 have demonstrated that additions of small amounts of 1,10-phenanthroline (Phen) in the cathodic deposition of Cu can easily induce potential and current oscillations. This study extended the previous work of Schlitter et al.46 by a proposal of a mechanism for the electrochemical oscillations, and in short, the autocatalytic variable was recognized as the effective area of the electrode surface. To the best of our knowledge, these are the only two papers dedicated to investigate dynamic instabilities in this system. Furthermore, a more detailed study devoted to a deep understanding of the kinetics underlying the nonlinear behavior is of great interest, particularly due to the possibility to control electrochemical switches based on surface modifications.

Herein, we report the presence of BS in the electro-deposition of Cu generated by the addition of Phen in the reaction media. Two different surface deposits were observed in the same set of controlling parameters. We interpreted this type of instability mechanistically with the aid of the electrochemical quartz crystal nanobalance (EQCN), the electrochemical impedance spectroscopy (EIS), and scanning electron microscopy (SEM) images of the surface, contributing to a comprehensive physical chemical description of the nonlinear behavior. These experimental results open the possibility to design new chemical switches based on the formation of micro- or nanostructured surfaces in which the surface morphology would play an important role.

■ EXPERIMENTAL SECTION

The electrochemical experiments were carried out using a counter electrode as a high area platinized-platinum foil, a reference electrode as a reversible hydrogen electrode, and a working electrode (WE) as a polycrystalline platinum sheet with an electrochemical area of 1.4 cm² inferred by the charge in the hydrogen underpotential deposition.47,48 In case of the EQCN measurements, the WE was a disk-shaped quartz crystal 5 MHz AT cut, consisting of a Au film evaporated over a thin Ti layer—Ti/Au polished with a geometric area of 0.385 cm². Frequencies of the quartz were converted in mass variations by the Sauerbrey equation49 with $-5.816 \times 10^{-9}$ g Hz⁻¹ as a constant. All glassware apparatus were cleaned by the immersion in sulfonitic solution for 12 h followed by exhaustive rinses and, at least, one boil with ultrapure water (Milli-Q Millipore, 18.2 MΩ cm).

Before each cyclic voltammetry experiment, the electrodes were cycled at 1.0 V s⁻¹ between 0.05 and 1.50 V for about 500 times in order to assure a reproducible surface structure. The cleanliness of the apparatus was checked from successive cyclic voltammograms at 0.05 V s⁻¹ until the characteristic voltammetric profile for the polycrystalline Pt electrode was observed. The supporting electrolyte was prepared by H₂SO₄ (Sigma-Aldrich, 95–98%) and CuSO₄ (Sigma-Aldrich, ≥98%) solutions using ultrapure water with final concentrations of 0.5 mol L⁻¹. The surface inhibition was caused by the addition of Phen (Sigma-Aldrich, ≥99%) in the reaction media.

Before the experiments, the electrolyte was kept in an ultrasonic bath for, at least, 10 min in order to obtain a better dissolution of Phen and was purged for 15 min with molecular nitrogen (White Martins, 99.99%) to remove oxygen. The electrochemical system was controlled with a potentiostat (Autolab/Eco-Chemie, PGSTAT302N) equipped with SCAN250, FRA32M, and EQCN modules. EIS was conducted at steady state with a sinusoidal potential perturbation in the frequency range of 50 kHz to 0.5 mHz and an amplitude of ±10 mV. Surface images were obtained by SEM (JEOL, JSM-6360LV), operating at 20 kV. All experiments were carried out at room temperature.

■ RESULTS AND DISCUSSION

Electrochemical Experiments. Figure 1 depicts cyclic voltammograms at slow sweep rates of $dE/dt = 0.01$ V s⁻¹ of copper sulfate in sulfuric acid solution with a final concentration of 0.5 mol L⁻¹ for each reagent. The potential window is 0.05–1.30 V (Figure 1a), −0.10 to 0.30 V (Figure 1b, black line), and, after the addition of 2.0 mmol L⁻¹ of Phen, −0.10 to 0.30 V (Figure 1b, red line). The electrodeposition of Cu is a well-known reaction studied along several decades.50–55 As already reported in an acidic media,52 the reduction of Cu starts around 0.30 V, following an instantaneous nucleation mechanism at very first stage of the faradic reaction for higher concentrations of copper (e.g., 0.025 and 0.05 mol L⁻¹). The current density reaches ~20 mA cm⁻² at 0.05 V and still shows negative values of currents up to the crossover potential of 0.30 V. From this potential, in the positive direction, it is possible to observe an intense current density peak of 32 mA cm⁻² at 0.55 V, corresponding to the oxidation of the Cu deposits and, in sequence, falling down to
zero current. For $E > 0.62$ V, Cu dissolution and a surface passivation occur.

Focusing on the potential interval of $-0.10$ to $0.30$ V (Figure 1b), the electrodeposition of Cu takes place continuously (black line) with a small hysteresis along the positive and negative sweeps. There is a mean separation of 1.9 mA cm$^{-2}$ between the curves. However, when Phen molecules are added in the electrolyte, with a final concentration of 2.0 mmol L$^{-1}$ (red line), the deposition process is substantially suppressed for $E < 0.15$ V. This inhibition reached a minimum value of $-5.21$ mA cm$^{-2}$ at 0.1 V and produced a noteworthy separation of 8.6 mA cm$^{-2}$ between the current curves along the both directions of the sweeps in the interval of $0.03 - 0.15$ V. The suppression is more pronounced below $0.03$ V, but it manifests slightly above $0.15$ V as well. Nakanishi et al.\textsuperscript{45} have postulated that adsorbed metallic complexes $[\text{Cu(I)}(\text{Phen})_2]^{2+}(\text{ad})$ could be mainly responsible for the inhibition and the formation of the NDR. Overall, for $E < 0.03$ V and $E > 0.15$ V, the hysteresis is so small that can be considered negligible. Accordingly, we considered a BS domain around the interval of $0.03 - 0.15$ V.

At this point, the experimental conditions where BS can be found were, definitely, well determined. This knowledge allowed a safe variation in the controlling parameters without getting too far from the BS domain. Figure 2 shows the dependence of the upper and lower potential limits of the BS by the change of the external resistance ($R_{\text{ext}}$) connected in series to the WE (Figure 2a--d) at Phen = 2.0 mmol L$^{-1}$ and the variation of concentration of Phen in the reaction media (Figure 2e--h) at $R_{\text{ext}} = 28$ $\Omega$ cm$^2$. All mapping experiments were conducted in low sweep rates of $dE/dt = 0.01$ V s$^{-1}$. Similar observations were done using slow rates in the cyclic voltammetries by Orlik et al.\textsuperscript{37} Both concentration and external resistance are considered as experimentally accessible controlling parameters in electrochemical systems and can induce dynamic instabilities by subtle changes.\textsuperscript{24}

The increase of $R_{\text{ext}}$ is followed by a progressive increase on the BS domain and it is also accompanied by a pronounced shift to negative potentials, as verified in Figure 2a--d. For external resistances of 7, 14, 21, and 28 $\Omega$ cm$^2$, the difference between the upper and lower potential limits of the BS was found to be 0.10, 0.16, 0.21, and 0.26 V, respectively (we adopted the turning points in the $j$ vs $E$ curves for the calculation). These regions do not depend on the vertex potentials in the cyclic voltammetries (Figure S1 in the Supporting Information). Indeed, the process that results in the abrupt current changes and generates the BS seems to occur in particular potentials values. This is a strong indicative that the adsorption of the inhibiting species happens in those specific potentials and remains on the surface, desorbing completely later on in the reverse cycle.

On the other hand, the increase of Phen concentration resulted in a shrinkage of the BS domain. At the lowest employed concentration of Phen, that is, 0.5 mmol L$^{-1}$, only

\begin{figure}
\centering
\includegraphics[width=\textwidth]{Figure2}
\caption{Cyclic voltammograms at $dE/dt = 0.01$ V s$^{-1}$ under different external resistances at Phen = 2.0 mmol L$^{-1}$ and $R_{\text{ext}} = (a) 7, (b) 14, (c) 21, and (d) 28$ $\Omega$ cm$^2$ and under the effect of the concentration of Phen at $R_{\text{ext}} = 28$ $\Omega$ cm$^2$ and Phen = (e) 0.5, (f) 1.0, (g) 1.5, and (h) 2.0 mmol L$^{-1}$.}
\end{figure}
the lower limit is observed and an expressive hysteretic profile does not arise (Figure 2e). When the concentration of Phen is increased to 1.0 mmol L\(^{-1}\) a different scenario appears and the BS is clearly established under these conditions (Figure 2f). Note that the approximation of the upper and lower limits to an invariable potential became even more evident in larger concentrations of Phen, that is, 1.5 or 2.0 mmol L\(^{-1}\) (Figure 2g–h). No significant changes were observed in these two concentrations, which indicate a surface saturation of the inhibiting species above of 1.5 mmol L\(^{-1}\).

A summary of the effect of \(R_{\text{ext}}\) and concentration of Phen on the bistable behavior in the electrodeposition of Cu is shown in terms of a bifurcation diagram in Figure 3. At first glance, its shape has a considerable resemblance with the bifurcation diagrams of systems belonging to an N-NDR category.\(^{26}\) In fact, Nakanishi et al.\(^{45}\) have observed electrochemical oscillations in both current and potential, which is a good indicative that the cathodic deposition of Cu perturbed with Phen remains in the hidden N-NDR (HN-NDR) subsystem. H means that the NDR can be partially hidden from an intermediate that blocks the surface and reduces the global rate in lower potentials.\(^{23,24}\)

Additionally, Figure 3 highlights the saddle-node bifurcation\(^{1} as the lines surrounding the domains of the BS and how it is strongly modified by the increase of the concentration of Phen. A straight green line represents the slight suppression of the current when [Phen] = 0.5 mmol L\(^{-1}\) is added into the electrolyte. The contraction of the BS domains reaches a maximum around 1.5 mmol L\(^{-1}\), and negligible changes are observed when compared to the domain formed at 2.0 mmol L\(^{-1}\) (Figures 2g,h and S2 in the Supporting Information). The calculated areas of these diagrams are 7.9 and 6.3 V \(\Omega\) cm\(^{2}\) for 1.0 and 2.0 mmol L\(^{-1}\) of Phen, respectively. This compression in the bifurcation diagram by inhibiting species is in agreement with the numerical simulations performed by Nascimento et al.\(^{56}\) A considerable shrinkage of the Hopf domain was observed as the irreversible surface poisoning evolved. The dynamics changed accordingly and the chaotic oscillations were dramatically suppressed.

The saddle-node bifurcation can be confirmed by the EIS under potentiostatic control when the imaginary part of the impedance is zero as the frequency of perturbation tends to zero as well,\(^{41}\) that is, \(Z'(\omega) = 0\) for \(\omega = 0\). Figure 4 depicts Nyquist plots for the electroreduction of Cu without (black lines) and with (red lines) Phen molecules in the concentration of 2.0 mmol L\(^{-1}\) at the applied voltages: (a) \(E = 0.24\) V, (b) \(E = 0.12\) V, and (c) \(E = -0.05\) V. The insets correspond to a “zoom in” on the plates (a–c). Sinusoidal amplitude of perturbation is ±10 mV from 50 kHz to 0.5 mHz.

Nyquist plots of the electrodeposition of Cu in three different applied voltages of 0.24, 0.12, and −0.05 V which are located at the SSI, the BS domain, and the SSII, respectively. Black lines indicate the cathodic reduction of Cu without the presence of Phen, whereas in red lines, Phen is present in the reaction media. We modeled the frequency responses by equivalent circuits for the impedances shown in Figure 4a,c.
and extracted the respective electrical parameters (Figure S3 and Table S1 in the Supporting Information).

Figure 4a shows that the process has distinct time scales with the presence of two semicircles, one at higher and another at lower frequencies. They are related to two limiting steps which compete simultaneously and are affected by the applied voltage—high frequencies describe activation control, whereas low frequencies describe mass transport control. In the latter, the Warburg impedance shows a decreasing imaginary component, resulting in the observed trend (Figure 4a, black line). It has been also ascribed that low-frequency loops are strongly dependent on the growth mode of copper deposits. There are different interpretations for the inductive behavior in the lowest frequencies, but it can represent the adsorption—desorption of inhibiting species, for example, anions from the electrolyte or a nucleation process (Figure 4a, black line). The Ohmic drop ($R_{ct}$) shows very low values of around 2.2 Ω cm$^2$ in all experiments.

Interestingly, the first semicircle is not affected by the presence of the Phen but causes a significant increase in the real and imaginary components of the low-frequency loop (Figure 4a, red line). As the dominant loop belongs to the higher frequency spectrum, the global faradaic reaction is not highly affected by the Phen molecules under the voltage of 0.24 V. Figure 1b reflects this description where no significant suppression is observed and similar charge transfer resistances ($R_{ct}$) of 10 Ω cm$^2$, with and without the presence of Phen, corroborates with this scenario. Nevertheless, there is a slight increase in the constant phase element from 21.4 to 29.9 μF cm$^{-2}$. For the second process located in lower frequencies, the polarization resistance increases from 2.1 to 8.4 Ω cm$^2$, reflecting the surface inhibiting effect of the Phen (Figure 4a, red line).

The dynamic instability induced by Phen becomes even clearer in the Figure 4b. The first observation is the anticlockwise movement in lower frequencies on the Nyquist plot and the subsequent intersection in the real axis at 1.5 mHz which is related to the Hopf bifurcation, that is, the imaginary components showed an evident capacitive effect of 390–687 mF cm$^{-2}$ when Phen is added into the reaction media.

### Gravimetric Monitoring

In this section, we will provide further insights on the nature of the adsorbed species and it will allow us to propose a mechanistic interpretation of how BS appears in this system. Figure 5 shows the $j$ vs $E$ curves (dashed line) during the electroreduction of Cu at $dE/dt = 0.02$ V s$^{-1}$ followed by the corresponding mass changes profiles (full lines) obtained with the aid of the EQCN. The system was perturbed with small additions of Phen as final concentrations of 0 (black), 0.5 mmol L$^{-1}$ (green), 1.0 mmol L$^{-1}$ (blue), and 2.0 mmol L$^{-1}$ (red). [H$_2$SO$_4$] = 0.5 mol L$^{-1}$ and [CuSO$_4$] = 0.5 mol L$^{-1}$.

![Figure 5](image-url)
observation suggests that there is a strong adsorption of species dissolved in the electrolyte during the negative sweep. This abrupt current suppression is detailed in Figure 6. When it is turned to positive direction, the adsorbed species are released from the surface and the Cu deposition occurs quicker.

Focusing on the potential interval of 0–0.20 V, it was possible to calculate the mass/charge (m/z) ratios from j vs E and Δm vs q curves in the negative sweep as represented by the arrows. The colors assignments are the same of the Figure 5. The experimental m/z ratio for the electrodeposition of Cu without the presence of Phen (Figure 6a,e) is 332 μg C⁻¹, which agrees with the theoretical value of 329 μg C⁻¹ and retracts the electroreduction of Cu²⁺ to Cu⁰ on the surface. In this calculation, we used the molar mass of 63.5 g mol⁻¹ for Cu and 2 electrons for the faradaic reaction, that is, 63.5 g mol⁻¹/2 × 96 485 C mol⁻¹. As long as the Phen concentration

Figure 6. (a–d) j vs E and (e–h) Δm vs q in the potential interval of 0–0.20 V. The arrows indicate the direction of the sweep. The curves follow the color code used in the text for Phen concentrations: 0 (black), 0.5 mmol L⁻¹ (green), 1.0 mmol L⁻¹ (blue), and 2.0 mmol L⁻¹ (red). The gray bands represent the potential window, where mass/charge ratios were calculated.

Figure 7. (a) j vs E, (b) Δm vs E, and (c) Δm vs q in the potential interval of 0–0.20 V in the absence of copper sulfate. The curves follow the color code used as previously: 0.5 mmol L⁻¹ (green), 1.0 mmol L⁻¹ (blue), and 2.0 mmol L⁻¹ (red). [H₂SO₄] = 0.5 mol L⁻¹.
increases, the $m/z$ values increases as well. In this respect, we can safely consider that the process attributed to the adsorption, responsible to generate the instability in the negative sweep, takes place more intensely following the ascending order: $[\text{Phen}] = 0.5 \text{ mmol L}^{-1}$ and $352 \mu\text{g C}^{-1}$ (Figure 6b), $[\text{Phen}] = 1.0 \text{ mmol L}^{-1}$ and $392 \mu\text{g C}^{-1}$ (Figure 6c), and $[\text{Phen}] = 2.0 \text{ mmol L}^{-1}$ and $416 \mu\text{g C}^{-1}$ (Figure 6d). The effect of adsorption of protonated Phen was studied separately, vide Figure 7.

Nakanishi et al. have postulated that the reduced form of the adsorbed metallic complexes $[\text{Cu(I)}(\text{Phen})]_2^{2+}$ could be the responsible for the inhibition effect, but it was observed mainly in neutral media. In acidic environment, the ligands could be totally protonated, which would weaken the coordination bonds between $\text{Cu}^{2+}$ and Phen and, therefore, generate uncoordinated ligands molecules. According to this description, Phen might also participate in the adsorption process causing the inhibition effect simultaneously with the adsorption of the metallic complexes. Figure 7a depicts cyclic voltamograms of the system $\text{CuH}_2\text{SO}_4,\text{Phen}$ under different concentrations of Phen. In this potential region, there is no $\text{Cu}$ dissolution (Figure S4 in the Supporting Information) and the anodic and cathodic currents can be associated with desorption and adsorption processes, respectively. Large currents are observed in higher concentrations of Phen.

Figure 7b shows that the respective mass profiles corroborate with the processes discussed previously. Note, however, that the mass scale is now in nanograms. In the positive sweep, the mass variations decrease for $E > 0.17 \text{ V}$ and increase in the negative sweep starting at $E < 0.19 \text{ V}$. In this respect, one can conclude that in NDR region observed during copper electrodeposition, the Phen molecules (or their protonated form) interact with the surface competing for adsorption sites. We have tried to explain the nature of Phen adsorption by $m/z$ ratios in the potential interval of 0.07–0.15 V, cf. Figure 7c. Straight lines with a good fit were obtained and correspond to $166 \mu\text{g C}^{-1}$ (green line), $170 \mu\text{g C}^{-1}$ (blue line), and $181 \mu\text{g C}^{-1}$ (red line).

As observed, these values do not show big discrepancies between each other and are approximately 10 times lower than those expected to the Phen adsorption in a one-electron process, thus suggesting unknown charge transfer processes occurring simultaneously at this potential window (Figure S4 in the Supporting Information). In certain extension, the adsorption of protonated Phen can be considered a fast equilibrium with the dissolved molecules in the interface, which modifies the total coverage of the adsorbed species and ends up with underestimated values of $m/z$. In the next sections, we will provide a more realistic information, or physical sense, by converting $m/z$ ratios in apparent molar mass and, finally, propose a kinetic mechanism to the emergence of the BS.

**Imaging Analysis.** Corroborating with the previous measurements, this section will support the overall discussion by extracting surface images with the aid of the SEM. Nakanishi et al. have inspected the electrodeposit surface and observed two types of surface morphologies during the electrochemical oscillations: leaflet-like particles of a few micrometers in size and dense round-shaped leaflets, which are associated with low and high current states, respectively. Cu leaflets would occur in an anisotropic crystal growth by a selective adsorption on a specific crystalline face inhibiting the global reaction. Interestingly, in a bistable state, the addition of Phen resulted in a completely different morphology, reflecting the emergence of structured surfaces driven by distinct dynamic instabilities.

**Figure 8** depicts SEM images of the surface at three applied voltages: $0.24 \text{ V (SSI)}$, $0.12 \text{ V [bistability, upper (BSu) and lower (BSd) branches, and } -0.05 \text{ V (SSII).]}$. For each experiment, we cycled 20 times between $-0.1$ and $0.3 \text{ V at } \text{d}E/\text{dt} = 0.01 \text{ V s}^{-1}$, pulled out the WE in the desired voltage and sequentially disconnected it from the potentiostat. This procedure assured a good reproducibility. The images show that the surface morphology of the electroplated deposit depends on the applied voltage and the presence of Phen in the reaction media. Regarding the SSI, in the potential region, where Phen alters slightly the current density (Figure 1b), a fine globular structure is clearly seen. As the potential slowly decreases, this fine structure moves toward a roundish nodule morphology, vide BSd. Then, after the abrupt suppression of the current by the adsorption of reaction intermediates, the system reaches a state where massive deposition of Cu takes place.

The image taken in the SSII represents this situation and evidences the production of thick Cu films, which are coarse-grained and uniform along the electrode. Similar observations have been reported in the electrodeposition of copper. Finally, in the reverse scan, the surface structure gradually changes to a finer globular formation again as seen in BSu. However, this return is not totally completed to the SSI state and visual differences are observed in the images found at BSu and BSd. Overall, the BSd surface structure has a lower number of fine nodulations and embranchments compared to BSu. The system is very sensitive to the initial conditions and, for this reason, exhibits a memory effect. In fact, this characteristic is a strong evidence of the presence of BS. According to the surface inhibition provoked by the addition of Phen, these molecules seem to prevent the formation of coarse grained structures.
because it demonstrates a substantial inhibiting effect in the reaction of Cu deposition.

**Mechanistic Interpretation.** The initial voltammetric characterization presented in Figure 1 attests the inhibition effect of the Phen in the electrodeposition of Cu. In fact, it has been proposed that metallic complexes have a strong adsorption on the surface, which reduce the total active area.\(^{45}\) Phen molecules have also a favorable interaction with adsorption on the surface, which reduce the total active area. It has been proposed that metallic complexes have a strong adsorption on the surface, which reduce the total active area.\(^{45}\)

First, it is important to consider that the reaction media, where the study was conducted, is highly acidic with \([\text{H}_2\text{SO}_4]\) = 0.5 mol L\(^{-1}\) and \([\text{CuSO}_4]\) = 0.5 mol L\(^{-1}\) or pH \(\rightarrow 0\). As the pK\(_a\) of the \([\text{H}(\text{Phen})]\) and \([\text{H}2(\text{Phen})]\) species are 5.12 and 2.11, respectively,\(^{42,45}\) the Phen molecules are protonated in the nitrogen atoms. Hence, for pK\(_a\) > pH, it is expected to observe the following reactions in the bulk of the solution

\[
\text{Phen}_{(aq)} + \text{H}^+_{(aq)} \rightleftharpoons [\text{H}(\text{Phen})]^{+}_{(aq)} \quad (r1)
\]

\[
[\text{H}(\text{Phen})]^{+}_{(aq)} + \text{Phen}_{(aq)} \rightleftharpoons [\text{H}(\text{Phen})]^{+}_{(aq)} \quad (r2)
\]

For the sake of simplicity, we neglected the contribution of the \([\text{H}(\text{Phen})]^{+}_{(aq)}\) species in the analysis, although it has been also detected experimentally in a self-stacking mechanism.\(^{75,76}\) Because copper sulfate is also added into the electrolyte, and possibly, those small changes in the reaction pathway.

Cu\(^{2+}\) (aq) + 2[H(Phen)]\(^+\) (aq) \rightleftharpoons [Cu(II)(Phen)]\(^{2+}\) (aq) + 2H\(^+\) (aq) \quad (r3)

which shows an oxidized form of the metallic complex as the dominant species in the solution. It is proposed that it can adsorb on the surface and compete in parallel with the deposition of Cu.

The electrochemical reduction of Cu in the solid/liquid interface takes place as

\[
\text{Cu}^{2+}_{(aq)} + 2e^- \rightarrow \text{Cu}^0 \quad (r4)
\]

This reaction proceeds through two steps, and the rate-determining step is considered to be the first electron transfer by the bivalent ion.\(^{77}\) We write this reaction in a short form \(r4\). On the other hand, the electrons transfer in the metallic complex happens in an adsorption pathway\(^{45}\)

\[
[\text{Cu}(\text{II})(\text{Phen})^2]^{+}_{(ad)} + e^- + * \rightleftharpoons [\text{Cu}(I)(\text{Phen})^2]^{+}_{(ad)} \quad (r5)
\]

in our electrochemical mechanism, asterisk means free active sites on the surface. The nature of the adsorbed complex is still not clear but experimental evidences in different substrates, concentrations, and solution pH have shown to be \([\text{Cu}(\text{I})(\text{Phen})]\)\(^{78,79}\). Nevertheless, in very strong acidic media as ours, the protonation of the nitrogen in the Phen molecules is energetically favorable. Because of the positively charged reactants, the formation of the coordinated bonds is not easily performed and a complex dissociation would occur as well.

Naturally, the formation of the NDR is severely dependent on the pH, being its appearance facilitated when pH increases.\(^{45}\)

This reaction intermediate has a short-lived state, and it is rapidly reduced to metallic Cu in lower potentials.\(^{76,79}\)

\[
[\text{Cu}(I)(\text{Phen})^2]^{+}_{(ad)} + e^- \rightarrow \text{Cu}^0 + 2\text{Phen}_{(aq)} + * \quad (r6)
\]

Then, larger concentrations of Phen result in steeper slopes \(j\) vs \(E\) during the inhibition process and a shift to less negative potentials. As the dynamic equilibrium of the reaction \(r5\) has an impact on the total coverage of the adsorbed metallic complexes, the mass variations would be associate mainly with the electrochemical deposition of Cu (the reduction evolving a metallic complex should shift the onset potential, but it is not verified in our experiments). Although the concentration of Phen is much lower than Cu\(^{2+}\)\(^{74}\), a fast equilibrium might be observed by those remaining protonated molecules which are not coordinated with Cu.

\[
[\text{H}(\text{Phen})]^{+}_{(aq)} + e^- + * \rightleftharpoons \text{Phen}_{(ad)} + \text{H}^+_{(aq)} \quad (r7)
\]

We can now use the m/z ratios and convert them to apparent molar mass (M\(_{app}\)), considering the faradaic process. Figure 6 provides the following information in each concentration of Phen: 352 \(\mu\)g C\(^{-1}\) (0.5 mmol L\(^{-1}\)), 392 \(\mu\)g C\(^{-1}\) (1.0 mmol L\(^{-1}\)), 416 \(\mu\)g C\(^{-1}\) (2.0 mmol L\(^{-1}\)). We already know the faradaic reaction of Cu\(^{2+}\) has a theoretical value of 329 \(\mu\)g C\(^{-1}\), and hence, the experimentally measured m/z ratios represent mostly the reaction \(r4\). This observation is in agreement with the large difference between the concentrations of copper sulfate and Phen in the reaction media and the massive electrochemical deposition of Cu during the gravimetric measurements. In this respect, the faradaic reactions \(r5-r7\) participate only with 23 \(\mu\)g C\(^{-1}\) (0.5 mmol L\(^{-1}\)), 63 \(\mu\)g C\(^{-1}\) (1.0 mmol L\(^{-1}\)), and 87 \(\mu\)g C\(^{-1}\) (2.0 mmol L\(^{-1}\)) because of the discount of 329 \(\mu\)g C\(^{-1}\) from the total value.

These m/z ratios are quite small, and if the conversion is made for a 1 electron transfer reaction, they can be associated with M\(_{app}\): 2.2, 6.1, and 8.4 g mol\(^{-1}\), respectively. The results point out that none of the metallic complex intermediates, containing Cu (63.5 g mol\(^{-1}\)) or Phen (180.2 g mol\(^{-1}\)), could be adsorbed on the surface because of their high molar mass. Note that our model does not take into account the participation of water molecules and (bi)sulfate anions dissolved in the electrolyte, and possibly, those small changes could be also related to it. Performing the calculation using the derivative \(d\Delta m/dC\) vs E along 0–0.2 V (Figure SS in the Supporting Information), it is possible to obtain point-by-point the m/z ratios. Clearly, the reaction \(r4\) is dominant and takes place over the total interval of 0–0.2 V, and the m/z ratios slightly diverge from 329 \(\mu\)g C\(^{-1}\) in the abrupt current suppressions shown in Figure 6.

On the basis of these experimental results, we can conclude that the adsorbed intermediates do not alter mass variations significantly during the formation of the NDR but can inhibit the reduction of copper ions and modify drastically the global faradaic reaction as observed in Figures 1 and 5. It was found that Phen molecules are adsorbed vertically on Cu(111) with their nitrogen atoms facing the substrate.\(^{71}\) The molecules were stacked to form polymer-like chains because of the \(\pi\)-electron attraction in the aromatic ring among them. The adlayer of Phen formed on Cu surface changes depending on the applied voltage and it is considered more organized in positive potentials. If the electrolyte is kept in acidic conditions, the molecular plane of Phen is inclined toward...
the metal surface, which could further enhance the inhibition factor.

Our experimental results converge to the idea that the inhibiting effect of the global faradaic reaction could be related with the adsorption of Phen—the protonated Phen molecules coordinate with Cu²⁺ in a negligible extension because of the highly acidic media (two positively charged reactants). Then, reactions r₅ and r₆ should not be present and the equilibrium in the reaction r₃ is definitely shifted to the left. Additionally, the adsorption of protonated Phen on the Cu surface might occur in a fast equilibrium with the molecules on the interface, impacting on the total coverage of the adsorbed species and resulting in underestimated values of m/z, vide Figures 6 and 7. It might be also considered a weak adsorption driven possibly by physical interactions, which does not contribute to meaningful mass variations, but still affects the Faradaic reaction of the electrochemical reduction of Cu.

**CONCLUSIONS**

We have studied the effect of Phen in the electrochemical reduction of Cu under strong acidic media by means of different surface techniques. Overall, Phen molecules are found to be protonated and the coordination with copper ions dissolved in the solution seems to be unfavorable. The Phen molecules adsorb on the surface and inhibit the reduction of Cu²⁺ from the solution. This blocking process alters significantly the area of the electrode and induces an autocatalytic evolution of the double layer potential. As experimentally observed, this dynamic behavior promotes the formation of a bistable domain, delineated by a saddle-node bifurcation, in a wide range of controlling parameters, such as the external resistance, the applied voltage, and the concentration of Phen.

The increase of the concentration of Phen favors a shrinkage of the bifurcation diagram and shifts it to less negative potentials. Besides voltammetric characterization, the EIS measurements also converge to a conclusion of an inhibiting process. The instability generated by adsorption was confirmed by the hidden negative impedance, that is, Δm versus E and dΔm/dC vs E plots.

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