Factors Affecting Nucleation and Growth of Chromium Electrodeposited from Cr\textsuperscript{3+} Electrolytes Based on Deep Eutectic Solvents

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ABSTRACT

Chromium was electrodeposited from deep eutectic solvents-based Cr\textsuperscript{3+} electrolytes on HB-pencil graphite electrode. Factors influencing the electrochemical behavior and the processes of Cr nucleation and growth were explored using cyclic voltammetry and chronoamperometry techniques, respectively. Cr\textsuperscript{3+} reduction was found to occur through an irreversible diffusion-controlled step followed by another irreversible one of impure diffusional behaviour. The reduction behavior was found to be greatly affected by Cr\textsuperscript{3+} concentration, temperature, and type of hydrogen bond donor used in deep eutectic solvents (DESs) preparation. A more comprehensive model was suggested and successfully applied to extract a consistent data relevant to Cr nucleation kinetics from the experimental current density transients. The potential, the temperature, and the hydrogen bond donor type were estimated to be critical factors controlling Cr nucleation. The nucleation and growth processes of Cr from either choline chloride/ethylene glycol (EG-DES) or choline chloride/urea (U-DES) deep eutectic solvents were evaluated at 70°C to be three-dimensional (3D) instantaneous and diffusion-controlled, respectively. However, the kinetics of Cr nucleation from EG-DES was found to be faster than that from U-DES. Cr nucleation was tending to be instantaneous at higher temperature, potential, and Cr\textsuperscript{3+} concentration. Cr nuclei electrodeposited from EG-DES were characterized at different conditions using scanning electron microscope (SEM). SEM images show that high number density of fine spherical nuclei of almost same sizes was nearly obtained at higher temperature and more negative potential. Energy dispersive spectroscopy (EDS) analysis confirms that Cr deposits were obtained.

GRAPHICAL ABSTRACT

Keywords: Chromium, Electrodeposition, Nucleation, Deep eutectic solvent, Chronoamperometry

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1. Introduction

Chromium and its alloys are one of the very important and widely used coatings in industry with the purpose of enhancing corrosion resistance “especially at high temperature”, hardness, abrasion and wear resistance, lubricity, and decorative characteristics such as surface texture and brightness [1,2]. It is usually applied to the various metallic substrates via electrochemical deposition. Cr coatings are of two types: one of them is decorative with (< 1.5 μm) thickness which is usually applied over another under-layer coatings such as copper and nickel. The other one is hard with (2.5-500 μm) or greater thickness and with (850-1200 HV) hardness which is usually applied on the base metal directly [1].

Indeed, chromium has many oxidation states; however, hexavalent chromium (Cr$^{6+}$) salts are the most widely used for Cr electrodeposition from aqueous solutions. These Cr$^{6+}$ aqueous baths have several drawbacks that necessitate a significant improvement in Cr electrodeposition technology. Some of the drawbacks are of technical type such as the concomitant hydrogen evolution reaction resulting in a low cathodic reaction efficiency (7-15%) [3], deposit embrittlement, and wasteful energy usage. Others relate to environmental and health issues, where Cr$^{6+}$ is categorized worldwide as highly toxic and carcinogenic agent and it has come under legal restriction by organizations around the world [4].

The need for safe and capable alternative bath technology for Cr electrodeposition becomes more rising. One of such alternatives which still under research and improvement is trivalent chromium (Cr$^{3+}$) based electrolytes [5-7]. It is less toxic and eco-friendly. Although the aqueous type of Cr$^{3+}$ electrolytes has already been used in industry for many years, it results in lower deposit properties such as crystallinity [8], hardness, and thickness (< 100 μm) [9], especially for the hard Cr type, than the traditional Cr$^{6+}$ aqueous electrolytes. Furthermore, Cr$^{3+}$ is barely reduced in aqueous electrolytes due to the formation of stable inner-complex [Cr(H$_2$O)$_6$]$^{3+}$ which is only reduced at potential more negative than that of water reduction leading to intense hydrogen evolution and higher pH value near the cathode [10]. Cr$^{3+}$ ions are easily oxidized to Cr$^{6+}$ at inert anodes during the electrodeposition process resulting in corruption of the electrolyte chemistry. It was reported that the addition of organic complexants to Cr$^{3+}$ aqueous electrolytes with low (Cr$^{3+}$/Ligand) ratio facilitates Cr$^{3+}$ reduction resulting in thicker deposits and this was explained by the formation of other less stable Cr$^{3+}$ complexes [11-13]. Certainly, the final Cr layer deposited from these electrolytes has high carbon content coming from organic additives reduction resulting in a weak deposit performance [14].

Recently, Non-aqueous electrolytes have arisen as an alternative technology for metals electrodeposition [15]. Non-aqueous electrolytes from which Cr has been electrodeposited include broad family composed of underlying categories such as molten salts-based electrolytes [16], traditional organic solvents-based electrolytes [17], and ionic liquids-based electrolytes [18].

In the last two decades, Ionic liquids attract researchers as solvents for Cr electrodeposition due to their wider potential window, good electrical conductivity, low volatility, and low hydrogen evolution compared to aqueous electrolytes [19,20]. Low chemical stability, high cost, and moderate toxicity of ionic liquids push researchers to try using a new class of green solvents known as deep eutectic solvents (DESs) [21]. DESs have the advantages of wider potential window, low toxicity, low environmental bad impact, low cost, and high chemical stability. DESs were reported to be classified into four types [22]. Abbott et al is considered as one of the pioneers in using DESs in purpose of Cr electrodeposition [23,24]. DESs based on choline chloride (ChCl)/hydrogen bond donor (HBD) combinations “known as DES of type III” have been used in the present work as promising solvents for Cr electrodeposition. Ethylene glycol (EG) and urea (U) have been used as HBDS with ChCl to obtain their relevant DES used to electrodeposit Cr from. Seldom reports have explored the mechanism and kinetics of Cr electrodeposition from these solvents [25,26], which are very important to get the knowledge about the optimum conditions for the desired performance.

For the first time, a disposable pencil graphite electrode (PGE) has been used in this work to study the mechanism and kinetics of Cr electrodeposition. PGE is appropriate to investigate the nucleation and growth processes occurring at high negative potentials with lower contribution from the hydrogen evolution. Moreover, commercial availability, low cost, and low background current give PGE a greater
advantage over the widely used glassy carbon or pyrolytic graphite electrodes [27]. Cu and Pd nucleation and growth mechanisms were studied using PGE by R. Majidi M and Rezaei M, respectively [28,29].

In this work the electrochemical behavior of Cr\(^{3+}\)/DES on PGE surface under different conditions has been studied using cyclic voltammetry (CV) technique. Factors affecting nucleation also have been studied from current density transients obtained under different conditions using chronocoulometry (CA) technique. The deposition on a foreign substrate is well known to occur through three dimensional (3D) nucleation process [30]. Several theoretical models have been reported to predict the behavior of nucleation mechanism. In this work a two suitable theoretical models were used to get qualitative and quantitative information about the Cr nucleation and growth process happened during Cr electrodeposition on PGE surface: one of these models was proposed by Sharifker and Hills (SH) which is based on two limiting cases of 3D-instantaneous (3D-IN) and progressive (3D-PN) nucleation and growth [31]. SH-model has been used just for qualitative estimation. The other model based on the idea of Palomar-Pardavé et al which indicates that the deviation of experimental transients from theoretical ones predicted by SH-model is due to the fact of presence of concurrent electrochemical processes other than nucleation and growth processes during electrodeposition process [32]. Non-linear fitting of a model considering the other individual contributions was used to get quantitative data about the factors affecting nucleation and growth of Cr on PGE. Deconvolution method has been used to get information about the current density transients’ individual contributions.

2. Experimental

2.1 Electrolyte preparation

Choline chloride/ethylene glycol (EG-DES) and choline chloride/urea (U-DES) deep eutectic solvents were prepared by mixing ethylene glycol (El-Nasr pharmaceutical chemicals co., +99%) and Urea (BDH Chemicals Ltd., +99.5%) with choline chloride (LobaChemie Pvt. Ltd., +98%) in a 2:1 molar ratio, respectively at 80\(^\circ\)C and stirring till clear homogeneous solutions were obtained. Various Cr\(^{3+}\) electrolytes were obtained by dissolving different moles of chromium chloride hexahydrate salt (CrCl\(_2\cdot6\)H\(_2\)O) (LobaChemie Pvt. Ltd., +93%) into DESs at 80\(^\circ\)C and stirring till clear homogeneous solutions were obtained. To prevent the moisture effect as possible, all the electrolytes were put directly after preparation in tightly-closed vials under N\(_2\) gaseous atmosphere and put in a desiccator till measurements.

2.2 Electrochemical experiments

A 10 ml-cell containing three electrodes was used to investigate Cr electrodeposition. Platinum and silver wires were used as counter electrode and quasi reference electrode (QRE), respectively. HB-type pencil lead (PGE) (KOH-I-NOOR, Czech Republic) of 2 mm diameter was used as the working electrode. Based on the elemental analysis reported by Skrzypczyńska K. et al. [33], PGE is chemically composed of C, O, Si, and Al as major elements and K, Ca, Mg, Ti, and Fe as minor elements.

For each experiment the electrochemical measurements were conducted using a computer controlled potentiostat / galvanostat model 273 A from Princeton Applied Research, Oak Ridge, TN, USA with software of 270/250-PAR.

PGE was treated as follows: wet-polished with 1500 grade emery paper, rinsed with double distilled water, and then treated by immersion in concentrated HCl for 1 min, rinsed with double distilled water, ultrasonically cleaned with ethanol, rinsed with double distilled water, and finally mirror-polished with a butter paper. The electroactive surface area of PGE has been determined experimentally using Randles-Sevick equation at 25\(^\circ\)C:

\[
I_p = 2.69 \times 10^5 n^{3/2} A D^{1/2} c^{1/2} v^{1/2}
\]

Where \(I_p\) is peak current (A), \(n\) indicates the number of electrons transferred \((n=1\ for\ K_{4}[Fe(CN)]_{6}\)), \(A\) is the surface area of the electrode (cm\(^2\)), \(D\) is the diffusion coefficient \((D = 6.3 \times 10^{-6} \text{ cm}^2 \text{s}^{-1} \text{ for} K_{4}[Fe(CN)]_{6}\)), \(c^o\) is the concentration of electroactive species (mol cm\(^{-3}\)), and \(v\) is the scan rate (V s\(^{-1}\)).

By recording CVs at different scan rates for 0.01 mol dm\(^{-3}\) potassium ferrocyanide solution and estimating the slope of \(I_p\ vs. v^{1/2}\) plot, the electrode surface area was estimated to be 0.0362 cm\(^2\).

Cyclic voltammetry experiments were performed in order to assess the electrochemical behavior.
Nucleation and growth processes were studied by the chronoamperometry technique.

All the electrochemical experiments were implemented under N\textsubscript{2} gaseous atmosphere. The cell temperature was controlled with the aid of a thermostated hot plate and a thermometer immersed in the sample during each experiment. The solvent current density background was subtracted from all recorded current density transients in this work. Furthermore, the potential error resulting from the uncompensated resistance was considered and the corrected potential scale was used for all experiments. All figures were constructed using OriginPro 2018 software.

2.3 Surface characterization

Deposits morphology was studied using scanning electron microscopy (SEM) (QUANTA, FEG 250, Japan). Their elemental compositions were analysed using energy dispersive spectroscopy (EDS) attached with the SEM.

3. Results and Discussion

3.1 Cyclic voltammetry study

The influences of potential scan rate, temperature, HBD type used in DES preparation and Cr\textsuperscript{3+} concentration on cyclic voltammogram of Cr\textsuperscript{3+} in DES at the PGE surface were assessed.

3.1.1 Effect of potential scan rate

As shown in Fig. 1 only two well-defined cathodic peaks, peak (I) at range (-0.8 to -1.0 V) which can be attributed to reduction of Cr\textsuperscript{3+} to Cr\textsuperscript{2+}, and peak (II) at range (-1.5 to -1.8 V) can be attributed to reduction of Cr\textsuperscript{2+} to Cr\textsuperscript{0}, implying a well-known two-step mechanism of reduction [11]:

\begin{align*}
\text{Cr}^{3+} + e^- & \rightarrow \text{Cr}^{2+} \quad \text{Peak I} \\
\text{Cr}^{2+} + 2e^- & \rightarrow \text{Cr}^{0} \quad \text{Peak II}
\end{align*}

No anodic peaks appear. It can be assumed that the chromium electrodeposition shows totally irreversible reduction process.

Fig. 1 presents the effect of increasing potential scanning rate (v) from 0.02 to 0.4 Vs\textsuperscript{-1} on behavior of the cathodic peaks (I, II) using 0.1 mol dm\textsuperscript{-3} Cr\textsuperscript{3+}/EG-DES on PGE at 30°C. For Peaks (I, II) the peak potential (E\textsubscript{p}) shifts negatively and the peak current density (j\textsubscript{p}) increases, with the increase of v. As shown in Fig. 2a, j\textsubscript{p} values of peak (I) varies linearly versus v\textsuperscript{1/2} with the best fit \([R^2 = 0.98874]\) implying that reduction of Cr\textsuperscript{3+} to Cr\textsuperscript{2+} (Peak I) is a typical diffusion-controlled process. Although j\textsubscript{p} values of peak (II) varies linearly versus v\textsuperscript{1/2} \([R^2 = 0.97544]\), the resulting line is not the best fit with data points as that of peak (I). Further inspection was done by plotting j\textsubscript{p} values of peak (II) versus v as displaced in Fig. 2b and the resulting fitting line suggests that j\textsubscript{p} values of peak (II) varies linearly versus v with the best fit \([R^2 = 0.99684]\). Thus one can conclude that reduction of Cr\textsuperscript{2+} to Cr\textsuperscript{0} (peak II) may be not pure diffusion-controlled process. The diffusionless behavior is often attributed to presence of an accompanying adsorption process and equation 2 can be used to justify Fig. 2b:

\[
j_p = \frac{n^2F^2}{4RT} \Gamma^* v \\
\]  

(2)

Where \(\Gamma^*\) (mol cm\textsuperscript{-2}) is the surface coverage of the adsorbed species, R (8.314 J mol\textsuperscript{-1} K\textsuperscript{-1}) is the ideal gas constant, \(n\) is the electron transfer number, \(T\) (K) is the absolute temperature, F (96487 C mol\textsuperscript{-1}) is the Faraday constant, and \(j_p\) (A cm\textsuperscript{-2}) is the peak current density. The value of \(\Gamma^*\) calculated from equation 2 is \(6.302 \times 10^{-9}\) mol cm\textsuperscript{-2}.

So that, one can assume that the whole process of chromium electrodeposition is totally irreversible diffusion-controlled process with adsorption contribution. Moreover, using the Laviron’s equation 3 [34] the transfer coefficient (\(\alpha\)) and the standard rate con-
stant \((k_s)\) of the second Cr reduction step corresponding to Peak II were estimated:

\[
E_p = E^0 - \frac{RT}{nF} \ln(\alpha n F / RT k_s) + lnv \tag{3}
\]

Where \(E_p\) (V) is the cathodic peak potential, \(E^0\) (V) is the formal potential, \(\alpha\) is the transfer coefficient, \(k_s\) (cm \(s^{-1}\)) is the standard rate constant of the surface reaction.

Using Fig. 1 the value of \(E^0 = -1.53\) V was estimated from the intercept of \(E_p\) vs. \(v\) plot on the ordinate by extrapolating the line to \(v = 0\) and from the plot of \(E_p\) vs. \(ln v\), the values of \(\alpha = 0.12\) and \(k_s = 0.3\) cm \(s^{-1}\) were obtained from the slope and intercept, respectively.

Since, (peak I) represents a pure diffusion-controlled irreversible step of \(\text{Cr}^{3+}\) to \(\text{Cr}^{2+}\) reduction process equations 4 and 5 can be used to calculate the transfer and diffusion coefficients of \(\text{Cr}^{3+}\) based on data obtained from Fig. 1 and 2a [35]:

\[
E_p = E^0 - \frac{RT}{anF} \left[ \frac{0.00534 \times F}{RT} + \ln \left( \frac{\pi D \alpha n F}{(RT)^{1/2}} \right) + \frac{lnv}{2} \right] \tag{4}
\]

\[
j_p = 0.4958 n F c^0 \frac{\alpha n F D v}{RT} \tag{5}
\]

Where \(j_p\) (A cm\(^{-2}\)) is the cathode peak current density, \(c^0\) (mol cm\(^{-3}\)) is the concentration of metal ions, \(D\) (cm\(^2\) s\(^{-1}\)) is the diffusion coefficient.

The values of \(\alpha = 0.19\) and \(D = 8.5 \times 10^{-7}\) cm\(^2\) s\(^{-1}\) were obtained from the slopes of \(E_p\) vs. \(ln v\) and \(j_p\) vs. \(v^{1/2}\) plots, respectively. The value of \(D\) is slightly lower than the reported value \(3.5 \times 10^{-6}\) cm\(^2\) s\(^{-1}\) obtained in 0.05 mol dm\(^{-3}\) \(\text{Cr}^{3+}/\text{EG-DES}\) at 70°C [26] and this result is logical due to lower temperature (30°C) and high viscosity of electrolyte used here in our experiment.

### 3.1.2 Effect of Temperature

The recorded CVs for neat EG-DES vs. the EG-DES containing \(\text{Cr}^{3+}\) show a clear difference. On one hand, the CVs of the EG-DES in Fig. 3a show no peaks in both polarization directions. The current density is constant from starting point of recording (-0.1 V) to about (-1.2 V), then the current density start to increase rapidly due to choline cation decomposition followed by hydrogen evolution [36,37]. Raising the temperature causes noticeable increase in current density in cathodic direction after (-1.2 V). This is due to the enhancement of choline cation decomposition and hydrogen evolution by thermal energy.

On the other hand, the CVs of 0.1 mol dm\(^{-3}\) \(\text{Cr}^{3+}/\text{EG-DES}\) electrolyte in Fig. 3b show two well-defined cathodic peaks. One can notice that raising the temperature results in a gradual increase in \(j_p\) values for cathodic peaks (I, II). This behavior is due to the increase of mass transfer rate [38]. Also, a shift to more positive potentials for both cathodic peaks is observed. This may be attributed to the decrease in energy barrier required for chromium reduction as a result of change in structure of Cr species formed in DES which accompanying the temperature changes [23,39]. This conclusion is supported by the change in color of \(\text{Cr}^{3+}/\text{EG DES}\) electrolytes as shown in Fig. 3.

By calculating \(D_{\text{Cr}^{3+}}\) at different temperatures (30, 50, and 70°C) based on data from Fig. 3b and equa-
tion 5, the activation energy of Cr\textsuperscript{3+} species diffusion ($E_D$) of value 4.3 kJ mol\textsuperscript{-1} and the pre-exponential factor ($A_D$) of value 1.7 cm\textsuperscript{2} s\textsuperscript{-1} were deduced using equation 6 [38] from slope and intercept of ln ($D_{\text{Cr}^{3+}}$) vs. T\textsuperscript{-1} plot, respectively.

$$D_{\text{Cr}^{3+}} = A_D \exp(-E_D/RT) \quad (6)$$

3.1.3 Effect of hydrogen bond donor type used in DES preparation

The type of HBD used in DES preparation strongly affects physical properties of the resulting DES such as viscosity and conductivity which further affects mass transport and electrochemical behavior. Cyclic voltammograms were recorded at 70°C on PGE at scan rate 0.05 V s\textsuperscript{-1} for 0.7 mol dm\textsuperscript{-3} Cr\textsuperscript{3+} dissolved in EG-DES (see Fig. 4a) and in U-DES (see Fig. 4b). It was found, as shown in Fig. 4, that the characteristic cathodic peaks (I, II) of Cr\textsuperscript{3+} reduction in EG-DES that appear at -0.75 V and -1.384 V, respectively, are more sharp and higher in $j_p$ values than those obtained in U-DES which appear at -0.9 and -1.53 V, respectively. This can be attributed to the lower viscosity and higher mass transport of EG-DES than that of U-DES at the same temperature (70°C). Thus, one can conclude that EG-DES can be a promising solvent for Cr electrodeposition.

3.1.4 Effect of Cr\textsuperscript{3+} concentration

To the best of our knowledge, the effect of using higher concentrations on electrochemical behavior of Cr\textsuperscript{3+}/EG-DES electrolyte is investigated for first time in our work. Fig. 5 shows the effect of increasing concentration of Cr\textsuperscript{3+} in EG-DES at different temperatures.
It is obvious that at the same temperature, the peak potential ($E_{p}$) and the peak current density ($j_{p}$) of all cathodic peaks shift to negative and to higher values, respectively, with raising Cr$^{3+}$ concentration. Occasionally, a new cathodic peak (III) at range (-1.30 to -1.39 V) was appeared at concentrations higher than 0.1 mol dm$^{-3}$ at the same temperature (30°C), and this can be explained by the change in coordination sphere surrounding Cr$^{3+}$ centers [23] resulting in new Cr$^{2+}$ species having lower reduction potential. This concentration effect was indicated by the color change of Cr$^{3+}$/EG-DES with the change in Cr$^{3+}$ concentration under the same temperature as in Fig. 5d. At higher temperatures (50°C), $j_{p}$ of Peak (II) begins to diminish, and Peak (III) becomes more sharp and greater in the peak current density ($j_{p}$) value as presented in Fig. 5b. This may be justified by the conversion of the hardly reduced Cr$^{2+}$ species corresponding to Peak (II) to the easily reduced Cr$^{2+}$ species corresponding to Peak (III) with increasing temperature. At 70°C, Peak (II) disappears completely, and Peak (III) becomes more sharp and higher in $j_{p}$ value as introduced in Fig. 5c. Thus, it can be concluded that there are many species of Cr$^{3+}$ formed in EG-DES solvents and the coordination sphere surrounding Cr$^{3+}$ centers can be modified according to the concentration of Cr$^{3+}$ and the temperature of electrolyte.

3.2 Chronoamperometry study
The influences of step potentials, HBD type used in DES preparation, temperature, and Cr$^{3+}$ concentration on Cr nucleation and growth were investigated using the recorded current density transient curves. Chronoamperomery experiments were executed by stepping the potential of the working electrode from the value (-0.2 V) at which no reduction of Cr$^{3+}$ occurs to potentials sufficiently negative to initiate reduction which corresponds to peak (III) as shown in Fig. 5c.

Fig. 5. Cyclic voltammograms recorded on PGE using 0.1, 0.3, and 0.5 mol dm$^{-3}$ Cr$^{3+}$/EG-DES at (a) 30°C (b) 50°C and (c) 70°C, with scan rate 0.05 Vs$^{-1}$. (d) represents photographic images showing color of different concentrations of Cr$^{3+}$/EG-DES at 50°C.
3.2.1 Qualitative estimation using the dimensionless SH-model

3.2.1.1 Effect of temperature

The effect of temperature on the mechanism of nucleation and growth of Cr on HB-PGE was investigated by performing CA experiments at two different temperatures (50 and 70°C) using 0.1 mol dm⁻³ Cr³⁺/EG-DES as shown in Fig. 6. The resulting current transients are typical of nucleation and growth process, in which at very short times the first current density decay can be attributed to charge-discharge equilibrium of double layer, adsorption-desorption equilibrium, and/or Cotterellian processes specially in multi step reduction process as in our case. Then, the current density increases to the maximum (jₘ) which corresponds to (tₘ), then the current decreases exponentially tending to a plateau.

As introduced in Fig. 6, as the applied potential becomes more negative, the values of jₘ increases and those of tₘ decreases. Also, it was found that increasing electrolyte temperature from 50 to 70°C leads to an increase in current of nucleation and growth of Cr on HB-PGE.

The effect of temperature on type of nucleation and growth of Cr on PGE was studied qualitatively by building dimensionless plots using jₘ and tₘ obtained from current density transients shown in Fig. 6 and comparing it with the dimensionless SH-model. Its two limiting cases of instantaneous (IN) and progressive (PN) can be expressed by two equations 7 and 8:

\[ \left( \frac{t}{t_m} \right)_{3D-\text{IN}} = \frac{1.9542}{v/v_m} \left( 1 - \exp \left[ -1.2564\left( \frac{t}{t_m} \right) \right] \right) \]

\[ \left( \frac{t}{t_m} \right)_{3D-\text{PN}} = \frac{1.9542}{v/v_m} \left( 1 - \exp \left[ -1.2564\left( \frac{t}{t_m} \right) \right] \right) \]

Fig. 6. Two families of current density transients obtained from chronoamperometry experiments for different step potentials using a 0.1 mol dm⁻³ Cr³⁺/EG-DES at (a) 50°C and (b) 70°C. (c) and (d) represent the corresponding dimensionless plots of the experimental current density transients and the theoretical ones of SH-model for instantaneous (IN) and progressive (PN) of 3D nucleation.
The fitted model can be deconvoluted into its individual contributions which can give information about processes occurring in parallel with the nucleation and growth process. taking into account the present case of two-step reduction mechanism, the cotterellian process $j_{\text{catt}}(t)$ of Cr$^{3+}$ reduction to Cr$^{2+}$, Langmuir type of adsorption-desorption process $j_{\text{ads}}$ deposition on PGE at 70°C was investigated as shown in Fig. 8. In the current density transients presented in Figs. 8a, 8b, and 8c, there is an increase in current density as Cr$^{3+}$ concentration increases. To get more information about nucleation and growth type a comparison between the experimental current density transients and theoretical 3D nucleation model. One can conclude, as in Figs. 8d, 8e, and 8f, that at the same temperature (70°C) and potential step, as the concentration increases, the nucleation and growth mechanisms shift from 3D-PN to 3D-IN type for ($t/t_m > 1$), the experimental curves is well coincided with the theoretical 3D-IN one for ($t/t_m < 1$) at all Cr$^{3+}$ concentrations.

### 3.2.2 Quantitative estimation using a non-linear fitting and deconvolution method

As shown in the previous section, Comparisons between dimensionless experimental current density transients and theoretical ones of SH-model, suggest that this model cannot adequately describe the nucleation and growth mechanism due to the complexity of real mechanism which incorporate many contributions other than the pure limiting IN and PN mechanisms. The early stages (the first current decay) of experimental current density transients may have many contributions from charge-discharge equilibrium of electrical double layer, adsorption-desorption equilibrium on electrode surface, and/or Cotterellian processes especially in multi-step reduction process as previously stated. The stage after $t_m$ may also have contributions from proton reduction and/or 2D-IN processes. All these contributions can combine to give the experimental current density transients. Thus, based on the idea of Palomar-paradávé et al.[32], a non-linear fitting technique is used to get the typical equation which can properly describe the experimental current density transients.

The resulting equation parameters can be used to calculate important kinetic parameters such as nucleation rate constant ($A$), the number density of active sites ($N_0$), the product ($AN_0$), and the diffusion coefficient ($D$) of the electroactive species.

The fitted model can be deconvoluted into its individual contributions which can give information about processes occurring in parallel with the nucleation and growth process. taking into account the present case of two-step reduction mechanism, the cotterellian process $j_{\text{catt}}(t)$ of Cr$^{3+}$ reduction to Cr$^{2+}$, Langmuir type of adsorption-desorption process $j_{\text{ads}}$
and proton reduction $j_{PR}(t)$ were suggested to be considered beside the 3D nucleation and growth $j_{3D}(t)$. So that the overall current density can be described by:

$$j_{total}(t) = j_{Ads}(t) + j_{Cott}(t) + j_{PR}(t) + j_{3D}(t)$$  \hspace{1cm} (9)

This model can be expressed in a parameterized format as follows:

$$j_{total} = K_1 \exp(-K_2 t) + P_0 t^{-1/2} + (P_4 + P_4 t^{-1/2})$$

$$ \times \left[ 1 - \exp \left( -P_2 \left( t - \frac{1 - \exp(-P_3 t)}{P_3} \right) \right) \right]$$

$$j_{Ads}(t) = K_1 \exp(-K_2 t)$$  \hspace{1cm} (10-a)

$$K_1 = E/R_s$$  \hspace{1cm} (10-b)

$$K_2 = 1/R_s C$$  \hspace{1cm} (10-c)

$$j_{Cott}(t) = P_0 t^{-1/2}$$  \hspace{1cm} (10-d)

$$P_0 = zFD_C^{1/2} \left( C_0^{1/2} / \pi \right)$$  \hspace{1cm} (10-e)

$$j_{PR}(t) = P_4 \theta(t)$$  \hspace{1cm} (10-f)

$$P_4 = (2C_0 M/\pi p) Z_{PR} F K_{PR}$$  \hspace{1cm} (10-g)

$$\theta(t) = \left( 1 - \exp \left( -P_2 \left( t - \frac{1 - \exp(-P_3 t)}{P_3} \right) \right) \right)$$  \hspace{1cm} (10-h)

$$j_{3D}(t) = P_4 t^{-1/2} \theta$$  \hspace{1cm} (10-i)

$$P_1 = zFD_C^{1/2} C_0^{1/2}$$  \hspace{1cm} (10-j)

Fig. 7. Two families of current density transients obtained from chronoamperometry experiments for different step potentials at 70°C using a 0.5 mol dm$^{-3}$ Cr$^{3+}$ dissolved in (a) EG-DES and in (b) U-DES. (c) and (d) represent the corresponding dimensionless plots of the experimental current density transients with theoretical ones of SH-model for instantaneous (IN) and progressive (PN) of 3D nucleation.
For all previous equations, $E$ (V) is the applied potential, $R_s$ (V A$^{-1}$ cm$^2$) is the electrolyte’s resistance, $C$ (V$^{-1}$ A cm$^{-2}$ s) is the double layer capacitance, $zF$ (C) is the molar charge transferred during the process, $D_{Cr^{3+}}$ and $D_{Cr^{2+}}$ are the diffusion coefficients for Cr$^{3+}$ and Cr$^{2+}$ species respectively, $c_o$ is the bulk concentration of active species, $M$ (g mol$^{-1}$) and $\rho$ (g cm$^{-3}$) are the atomic mass and density of Cr$^0$, $N_0$ (cm$^{-2}$) is the number density of active sites, $A$ (s$^{-1}$) is the nucleation rate constant.

**3.2.2.1 Effect of step potential**

Fig. 9 shows a comparison of experimental current density transients with theoretical ones of SH-model for instantaneous (IN) and progressive (PN) of 3D nucleation.
density transients recorded on PGE using 0.1 mol dm\(^{-3}\) Cr\(^{3+}\)/EG-DES at the same temperature (70\(^\circ\)C) but at different step potentials and with those got by the non-linear fitting of the parametric equation 10 to the experimental data. It is noticed that at any potential used, the resulted curve is well fitted to the experimental current density transient overall the time \((t)\) range. It is worth noting that there is a difference in the individual contributions obtained by the deconvolution of the fitted curve, with the increase of step potential used. At the lowest potential used (-1.25 V) proton reduction contribution \(j_{PR}(t)\) is negligible, and then it begins to appear and increase with the rise in potential values. Contributions due to Cotterellian \(j_{Cott}(t)\) adsorption \(j_{Ads}(t)\) and 3D-nucleation and growth \(j_{3D}(t)\) concurrent processes are observed for all different potentials. The \(t_m^{3D}\) and \(j_m^{3D}\) corresponding to the maxima of current density transient contributions of 3D nucleation and growth are shifted to lower and higher values, respectively, with increasing the step potential. The estimated constants of equation 10 by fitting the experimental transients through applying the proposed model 9 are presented in Table 1a. The kinetic parameters \(D_{Cr^{3+}}, D_{Cr^{2+}}, A, N_0\) and \(AN_0\) are also calculated based on these constants. As presented in Table 1a, the average diffusion coefficient of Cr\(^{3+}\) is \(5.7 \times 10^{-7}\) cm\(^2\)s\(^{-1}\) and that of Cr\(^{2+}\) is \(2.515 \times 10^{-7}\) cm\(^2\)s\(^{-1}\) which is slightly lower than the results reported by Mejía-Caballero I et al. [26].

It has been found that the values of nucleation kinetic parameters \(A, N_0\) and \(AN_0\) significantly increase with increasing the applied step potential, indicating that the

![Fig. 9.](image)
kinetics of nucleation during Cr electrodeposition is greatly affected by the applied step potential.

3.2.2.2 Effect of hydrogen bond donor used in DES preparation

Fig. 10 presents experimental current density transients obtained at the same Cr\(^{3+}\) concentration (0.5 mol dm\(^{-3}\)) and the same temperature (70°C) but at different HBDs namely EG (see Fig. 10a) and U (see Fig. 10b) using step potentials (-1.38 and -1.53 V), respectively. One can observe larger \(t_m\) and lower \(j_m\) when U is used instead of EG as HBD. This behavior can be attributed to lower nucleation kinetics which are indicated by the kinetic parameters \(A\)

![Table 1](image)

Table 1. The best fitting parameters obtained through the non-linear fitting of equation 10 to the experimental current density transients recorded on PGE at (a) different step potential using 0.1 mol dm\(^{-3}\) Cr\(^{3+}\)/EG-DES at 70°C (b), different HBD using 0.5 mol dm\(^{-3}\) Cr\(^{3+}\) dissolved in EG-DES (step potential -1.38 V) and in U-DES (step potential -1.53 V), (c) different concentrations of Cr\(^{3+}\)/EG-DES at step potential -1.25 V at 70°C, and (d) different temperatures at step potential -1.30 V using 0.1 mol dm\(^{-3}\) Cr\(^{3+}\)/EG-DES

![Fig. 10](image)

Fig. 10. Comparison between the experimental current density transients recorded on PGE using 0.5 mol dm\(^{-3}\) Cr\(^{3+}\) dissolved in (a) EG-DES (step potential -1.38 V) and in (b) U-DES (step potential -1.53 V) and their respective theoretical ones obtained through the nonlinear fitting of equation 10 to the experimental data. The individual contributions are shown in different line styles (see figure legends).
and $AN_0$ (see Table 1b) obtained from the well-fitted theoretical transients. The greater value of $N_0$ parameter in case of U-HBD than that in case of EG-HBD as presented in Table 1b, can be attributed to the higher potential (-1.53 V) used for recording current transients in case of U-DES than the potential (-1.38 V) used in recording current transients in case of EG-DES. This behavior is agreed with the fact of strong dependence of $N_0$ on applied potential, where the former increases with increasing the later. As shown in Fig. 10, when EG is used as HBD the well-fitted theoretical current density transient are deconvoluted into four curves representing the contributions from Cotterellian process of $\text{Cr}^{3+}$ reduction to $\text{Cr}^{2+}$, adsorption process, proton reduction process and 3D nucleation and growth. While, when U is used as HBD the well-fitted theoretical current density transient is deconvoluted into only three curves representing the contributions from adsorption process, proton reduction process and 3D nucleation and growth. The absence of cotterellian contribution of $\text{Cr}^{3+}$ reduction to $\text{Cr}^{2+}$ can be attributed to passivation of electrode surface by the competing adsorption.

Furthermore, the high viscosity of U-DES compared to EG-DES leads to small concentration gradient and large diffusion layer thickness. This behavior

![Graph image](image-url)

**Fig. 11.** Comparison between the experimental current density transients recorded on PGE at step potential -1.25 V using (a) 0.1 mol dm$^{-3}$, (b) 0.3 mol dm$^{-3}$, and (c) 0.5 mol dm$^{-3}$ $\text{Cr}^{3+}$/EG-DES at 70°C, and their respective theoretical ones obtained through the nonlinear fitting of equation 10 to the experimental data. The individual contributions are shown in different line styles (see figure legends).
is supported by the broad reduction peak I (see Fig. 4b). The proton reduction contribution in U-DES is noticed to be greater than that of EG-DES. This can be explained by the cathodic shift in the reduction potential ($E_P$) of $\text{Cr}^{2+}$ to $\text{Cr}^{0}$ when U-DES instead of EG-DES is used as shown in Fig. 4, at which a concurrent proton reduction and hydrogen evolution process occurs.

3.2.2.3 Effect of $\text{Cr}^{3+}$ concentration

By comparing the experimental current density transients (see Fig. 11) obtained at different concentrations using PGE at the same step potential (-1.25 V) and temperature (70°C), it was observed that however $j_m$ increases with increasing concentration, $t_m$ has almost the same value. Deconvolution of the well fitted theoretical transients resulted in individual contributions of adsorption process and cotterellian process beside the main 3D nucleation and growth process for $\text{Cr}^{3+}$ concentration of 0.1 and 0.3 mol dm$^{-3}$, but at 0.5 mol dm$^{-3}$ the adsorption contribution disappears and instead a proton reduction contribution appears. As introduced in Table 1c $A$ increases while $N_0$ decreases with increasing concentration from 0.1 to 0.3 mol dm$^{-3}$ due to the increase of electrode surface covered by adsorbed species then the effect is reversed for both $A$ and $N_0$ at 0.5 mol dm$^{-3}$ which may attributed to the change in contribution percentage of other concurrent processes. But, $AN_0$ markedly decreases with increasing the $\text{Cr}^{3+}$ i.e. the rate of nucleation is decreased with increasing $\text{Cr}^{3+}$ concentration and then tends to stability due to the competition with other processes which have high effect on the exposed active centers.

3.2.2.4 Effect of temperature

It has been found that the electrolyte temperature has a marked effect on nucleation and growth mechanism as introduced in Fig. 12. $t_m$ and $j_m$ are shifted towards lower and higher values, respectively. With the temperature rise from 50 to 70°C. At lower temperature (50°C) the theoretical 3D nucleation and growth contribution curve has no characteristic sharp nucleation peak i.e. lower nucleation rate, which is supported by data in Table 1d. It is also noticed that at lower temperatures (50°C) the adsorption process is greatly favored but at high temperature (70°C) the proton reduction and cotterellian process of $\text{Cr}^{3+}$ reduction into $\text{Cr}^{2+}$ are favored over adsorption. As shown in Table 1d $A$, $N_0$ and $AN_0$ significantly increase with increasing the temperature.

3.3 Morphology and elemental composition

The surface of PGE was investigated by SEM and EDS techniques before and after electrochemical nucleation and growth of Cr from a quiescent $\text{Cr}^{3+}$/EG-DES. The neat PGE surface in Fig. 13a has the well-known layer structure of graphite. EDS analysis

![Fig. 12. Comparison between the experimental current density transients recorded on PGE at step potential -1.30 V using 0.1 mol dm$^{-3}$ $\text{Cr}^{3+}$/EG-DES at (a) 50°C and (b) 70°C and their respective theoretical ones obtained through the nonlinear fitting of equation 10 to the experimental data. The individual contributions are shown in different line styles (see figure legends).](image_url)
proved the aforementioned composition of HB-type of PGE. It is composed of Si, Al and O beside C as major elements (see Fig. 13c). When chronoamperometry experiment executed by stepping the potential of PGE to more negative value (-1.38 V) was done for 60 sec, Cr nuclei were born instantenously and grown in a 3D mode as shown in Fig. 13b. The appearance of characteristic Cr peaks in EDS analysis in Fig. 13d proves that the obtained deposits are mainly Cr. Peaks of Si, Al, O and C elements belonging to the substrate (neat PGE) appear because the layer of Cr nuclei are very thin and X-ray used in EDS is highly penetrating.

Morphology of Cr nuclei formed in the initial stages of electrodeposition process is crucial to determine nucleation and growth type under different conditions. Effect of temperature on Cr nucleation and growth as shown in Figs. 14a and 14b was obtained at 50°C and 70°C from Cr3+/EG-DES under same other conditions. At low temperature (Fig. 14a), Cr nuclei are of irregular shapes and inhomogeneous size distribution compared to that of greater spherical ones of homogeneous size distribution obtained at higher temperature (Fig. 14b). At lower temperature the deposits are patchy and compact. Further, at higher temperature number density of Cr nuclei is greater. These results agree with the obtained electrochemical and theoretical results where $A, N_0$ and $AN_0$ significantly increase with increasing the temperature.

The effect of Cr3+ concentration on Cr deposits morphology obtained from Cr3+/EG-DES also was studied. In general, one can notice that at higher Cr3+ concentrations (Figs. 14c and 14d) the born Cr nuclei are of greater sizes and lower number density compared to those obtained at lower Cr3+ concentration (Fig. 14b). However, in Fig. 14c, inhomogeneity in nuclei size distribution and presence of some mature nuclei of greater sizes suggesting PN mechanism at lower negative potentials (-1.25 V), almost all nuclei in Fig. 14d are of nearly same size at higher negative potentials (-1.38 V) in accordance with the IN mechanism. Further, the results show formation of more fine nuclei with great number density at high negative potentials. These results are in agreement with the previous electrochemical results where $A, N_0$.

![Fig. 13. SEM images of (a) neat PGE surface, (b) Cr nuclei on PGE electrodeposited at -1.38 V from 0.5 mol dm$^{-3}$ Cr3+/EG-DES at 70°C. (c) and (d) represents the corresponding EDS spectra.](image-url)
and $AN_0$ significantly increase with more negative step potential values. Fig 14 shows that growth of Cr nuclei is following Volmer-Weber mechanism (i.e. 3D-islands are growing in all directions) [30].

4. Conclusions

Cyclic voltammetry study of Cr$^{3+}$ reduction on PGE showed that it occurs through totally irreversible two-step mechanism. The first step includes reduction of Cr$^{3+}$ to Cr$^{2+}$ under diffusion control, while the second step includes reduction of Cr$^{2+}$ to Cr$^0$ under impure diffusion process accompanied by adsorption where the estimated coverage value was $6.302 \times 10^{-9}$ mol cm$^{-2}$ for 0.1 mol dm$^{-3}$ Cr$^{3+}$/EG-DES at 30°C. CVs showed that Cr$^{3+}$ reduction mechanism was significantly affected by the electrolyte temperature, Cr$^{3+}$ concentration and type of HBD used to prepare DES. The appearance of new reduction peak corresponding to reduction of Cr$^{3+}$ to Cr$^0$ at lower cathodic value range (-1.30 to -1.39 V) when higher concentrations of Cr$^{3+}$ are used, suggests the formation of new Cr complexes and exchange of coordination sphere. Further, increasing the temperature resulted in enhancing this new peak and diminishing the old peak appeared at range (-1.7 to -1.4 V) till disappearance. This is implying that Cr$^{3+}$ species concentration and temperature are important to control the type of Cr complex species present in DESs and consequently the electrochemical behaviour and deposition processes of Cr.

Chronoamperometry was used as valuable tool to get important information about the kinetics of Cr nucleation and growth. Although the dimensionless SH-model was not well fitted with all the resulting experimental current transients due to presence of other parallel processes accompanying Cr nucleation and growth processes, it was used as a qualitative tool to elucidate the Cr nucleation type under different conditions. Nucleation of Cr on PGE using either EG or U as HBD was found to be of 3D-IN type. However, kinetics was better incase of EG-DES than incase of U-DES. Cr nucleation was tending to be instantaneous at higher electrolyte temperature, potential, Cr$^{3+}$ concentration. It was concluded that experimental current transients of Cr on PGE does

Fig. 14. SEM images of Cr nuclei on PGE electrodeposited from 0.1 mol dm$^{-3}$ Cr$^{3+}$/EG-DES at (a) 50°C (step potential -1.30 V) and at (b) 70°C (step potential -1.25 V) and from 0.5 mol dm$^{-3}$ Cr$^{3+}$/EG-DES at 70°C using step potential (c) -1.25 V and (d) -1.38 V.
not represent nucleation and growth processes only, but they include many contributions from other concomitant processes including the cotterelian one corresponding to first reduction step of Cr$^{3+}$ to Cr$^{2+}$, adsorption/desorption equilibrium process, proton reduction process. In combination with the deconvolution method a more general and adequate model was suggested based on Palomar-Pardavé methodology. In this suggested model all possible processes which could accompany the Cr nucleation and growth processes were considered. The contributions from these parallel processes and the kinetics of Cr nucleation were estimated quantitatively under different conditions.

Nucleation kinetics of Cr on PGE is mainly controlled by applied potential, temperature, HBD type. The Cr$^{3+}$ concentration does not mainly affect nucleation process, but may greatly affect growth step. Proton reduction process contribution was enhanced in case of using U-DES, higher potential values, higher Cr$^{3+}$ concentrations and higher temperatures. For 0.1 mol dm$^{-3}$ Cr$^{3+}$/EG-DES at 30°C, the diffusion coefficient for Cr$^{3+}$ species was estimated via CV and CA techniques and they have been found to be 8.5 ×10$^{-7}$ and 5.7 ×10$^{-7}$ cm$^2$s$^{-1}$, respectively, while, the calculated diffusion coefficient of Cr$^{2+}$ using CA technique has been found to be 2.515×10$^{-7}$ cm$^2$s$^{-1}$. SEM images show that at higher temperature and potential values, Cr nuclei electrodeposited from EG-DES has high number density of fine spherical nuclei of almost same sizes. Also, nuclei growth is following Volmer-Weber mechanism (i.e.3D-islands are growing in all directions). EDS analysis confirms that high-purity metallic Cr deposits were obtained.

Chronoamperometric technique in combination with the suggested comprehensive theoretical model and the deconvolution method can be a powerful tool for better understanding of the real nucleation and growth processes at the microscopic level which directly control the physical properties of the electrodeposited metal layer. So that, future studies concerning the effect of controlling Cr nucleation and growth on the physical properties of the electrodeposited layer from EG-DES can be suggested.

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