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XU, L., PI, L., DOU, Y., CUI, Y., MAO, X., LIN, A., FERNANDEZ, C. and PENG, C. 2020. Electroplating of thick hard chromium coating from a trivalent chromium bath containing a ternary complexing agent: a methodological and mechanistic study. ACS sustainable chemistry and engineering [online], 8(41), pages 15540-15549. Available from: https://doi.org/10.1021/acssuschemeng.0c04529
Electroplating of Thick Hard Chromium Coating from a Trivalent Chromium Bath Containing a Ternary Complexing Agent: A Methodological and Mechanistic Study

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ABSTRACT:

Trivalent chromium electroplating is considered a greener alternative to hexavalent chromium electroplating. However, the electrochemical reduction of Cr(III) in aqueous solution is extremely unstable and the thickness of the coating can hardly grow beyond 10 \( \mu \)m, which limits its industrial applications. We herein report a trivalent chromium bath containing a ternary complexing agent, which yields bright and uniform chromium coatings with thickness exceeding 30 \( \mu \)m, as well as high hardness and corrosion resistance. The electroplating behaviors and first principal calculations reveal that the Cr(II) intermediate plays a vital role in sustained electroplating in Cr(III) baths. The composition of complexing agents has a profound effect on the geometry and electron accepting ability of the Cr(II) complex ions. The failure to grow thick Cr coating is attributed to the accumulation of inactive hydroxo-bridged complexes. In the bath with ternary complexing agent, the Cr(II) complex ions are dsp\(^3\) or dsp\(^2\) hybridized with high electron affinity, which can be readily reduced to metallic Cr. The bath also shows a strong competitive ligand-binding effect that renders Cr(II) to preferentially bind with an organic ligand rather than a hydroxy ligand. Furthermore, the planar geometry of dsp\(^2\) hybridization does not favor the formation of hydroxo-bridged complexes because of the steric hindrance effect. The above merits of the ternary complexing agent lead to sustained electroplating, and thick Cr coatings are obtained.

KEYWORDS: trivalent chromium electroplating, complexing agent, steric hindrance, electron affinity, divalent intermediates

INTRODUCTION

Hard chromium (HC) coating has been widely used for protective and functional purposes in mechanical parts of aerospace, automotive, and petrochemical industries due to its multiple desired properties, such as high hardness, low friction coefficient, superior wear, and corrosion resistance.\(^1\)\(^-\)\(^3\) The thickness of HC coating is usually in the range from several tens
to several hundreds of micrometers.^^4,5\^\textsuperscript{4,5} Despite its widespread use, most high-quality HC coatings are electrodeposited from aqueous electroplating baths containing highly toxic hexavalent chromium, i.e., Cr(VI). With the increasing environmental concerns, strict regulations, or complete ban has been or will be enacted on hexavalent chromium electroplating worldwide.^^6,7\^\textsuperscript{6,7} As a more environmentally benign alternative, trivalent chromium electroplating (TCE) has been studied extensively in recent decades.^^8,9\^\textsuperscript{8,9} Despite the success in depositing thin decorative Cr coatings (0.1−2 μm), it remains a challenge as how to obtain high-quality thick HC coatings from a Cr(III) bath. The low current efficiency or high overpotential is commonly observed which is attributed to the formation of stable coordination of Cr(III) with water molecules. Furthermore, the deposition rate suffers from a rapid decrease over time and the obtained maximum coating thickness is generally less than 10 μm,^^10−12\^\textsuperscript{10−12} making the product unsuitable for functional applications. It is postulated that the fast deactivation is mainly due to the following reasons. During the electrodeposition process, hydrogen evolution reaction causes a local pH increase near the cathode, resulting in the formation of an electroactive intermediate, i.e., hydroxo-complexed chromium(II) ions.^^13−15\^\textsuperscript{13−15} These complex ions can be easily converted into inert hydroxy-bridged compounds through an olation reaction and hinder the continuous reduction of Cr(III).^^16\^\textsuperscript{16} As a result, both the current efficiency and deposition rate drop rapidly and the coating thickness can hardly increase.

Numerous endeavors have been made to deposit HC by changing the solution chemistry and process conditions. Ionic liquids (ILs) have attracted increasing interest as media for TCE to avoid parasitic hydrogen evolution.^^17−20\^\textsuperscript{17−20} Although the electrochemical window of ILs is wide, its high viscosity, poor conductivity, and high cost limit its practical applications. Other endeavors have been devoted to electrodeposition of Cr alloys (e.g., Cr−C, Cr−Ni, and Cr−Co) from aqueous solutions containing chromium(III) and other reducible components.^^21−24\^\textsuperscript{21−24} However, the properties of these composite coatings are only similar to certain aspects of HC. There is no alloy reported in literature that could adequately replace HC in overall performances.

Alternatively, the use of multicomponent complexing agents is regarded as the most effective and convenient approach for improving the properties of aqueous plating bath as well as the quality of the deposits.^^25−31\^\textsuperscript{25−31} The electrolyte containing more than one ligand may strengthen the ligand exchange reaction and regulate the overpotential for metal deposition, which ultimately improves the stability and efficiency of electroplating. This method has also been studied for Cr(III) plating. For instance, Cr(III) baths containing both formic acid and urea have yielded thick Cr coatings with a rather high current efficiency.^^32−34\^\textsuperscript{32−34} Nevertheless, these deposits commonly show inadequate properties such as hardness and corrosion resistance, making them unsuitable for tribological and anticorrosion applications. While the search for optimum bath composition for HC electroplating is mostly intuitive, the underlying mechanism also remains debatable.^^35−37\^\textsuperscript{35−37} The role of divalent chromium intermediates in the electrode position reaction has not been elucidated. Moreover, the mechanism of depositing thick uniform coating from Cr(III) baths containing a multicomponent complexing agent is yet to be elucidated. In this paper, a trivalent Cr bath containing ternary complexing agent is proposed for hard chromium electrodeposition. The performances of the plating solution and the chromium deposit are investigated and compared with those in the literature. Furthermore, the role of the Cr(II) intermediates is comprehensively studied through first-principles calculations, and the mechanism for depositing thick uniform high-quality chromium coatings is discussed.
**Electrodeposition of Cr.** The basic composition and concentration of the trivalent chromium bath containing ternary complexing agent are summarized in Table 1. All solutions were prepared using deionized water and AnalaR grade chemicals. The pH was adjusted to 1.5 by addition of H$_2$SO$_4$ or NaOH. After mixing of all components, the solution was heated at 60° C for 20 min, then cooled and aged for 24 h. The two-electrode electroplating was carried out in a temperature-controlled glass cell without separation of electrode compartments. Dimensionally stable Ti/IrO$_2$ anode (DSA) and brass (S = 0.1 dm$^2$) were used as the anode and cathode, respectively. The surface of the brass substrate was polished with magnesium oxide and then rinsed with hydrochloric acid solution and distillate water. Prior to galvanostatic chromium plating, the optimum current density range of the plating bath was determined using a 267 mL standard Hull Cell at a cell current of 10 A for a duration of 10 min. A brass plate (75 × 100 mm) served as the substrate, and the length of bright Cr-covered area was measured after electrolysis. The corresponding current density was evaluated by the following empirical equation (German standard DIN50957, eq 1):

$$j(x) = I(5.10 - 5.24 \log x)$$  \hspace{1cm} (1)

where $x$ indicates the distance (cm) along the cathode from the high current edge, $j(x)$ is the local current density (A/dm$^2$) at distance $x$, and $I$ is the total electrolytic current (A). The galvanostatic chromium deposition was carried out at 30 °C for different duration time of 2, 5, 10, 15, 20, 25, 30, and 35 min.

**Characterization.** The thickness of the Cr coatings deposited on brass substrate was measured by a TT260 thickness detector (Shidai Instrument, China) based on the Eddy current principles. The current efficiency of chromium electrodeposition was calculated as the ratio between the experimental and theoretical weight uptakes after electrodeposition. Vickers hardness (HV) of chromium coatings were evaluated by a microhardness tester with a load of 100 g. The surface morphology of deposits was studied by SIGMA scanning electron microscope (Zeiss, German). The crystal structure of the coating was characterized by a SHIMADZU XRD-6000 diffractometer (Japan) employing Cu K radiation ($\lambda = 0.154$ nm) with 20 canning step of 2°/min and angle range of 10° ~ 80°. Chemical composition of the chromium deposit was examined using an Aztec Energy X-Max energy-dispersive spectrometer (Oxford Instruments, UK) with an acceleration voltage of 20 kV.

**Electrochemical Measurements.** Electrochemical measurements were conducted in a three-electrode cell using a CHI660D electrochemical workstation (Chenhua, China). A platinum foil (1 × 1 cm) or the Cr-coated brass sheet (1 × 1 cm) served as the working electrodes for the polarization curve test of the bath and corrosion behavior characterizations of the Cr coatings, respectively. The counter and reference electrode were a large area platinum foil and a saturated calomel electrode (SCE), respectively. Cathodic polarization was performed in the Cr(III) baths at a scan rate of 5 mV/s. The Tafel plot was obtained in 3.5% NaCl aqueous solution at a scan rate of 2 mV/s from −0.3 to 0.3 V versus the open circuit potential (OCP). Electrochemical impedance measurement was also carried out in 3.5% NaCl aqueous solution and recorded in the frequency range from 10−2 ~ 105 Hz with an AC amplitude of 5 mV. All experiments were carried out at room temperature.
Density Functional Theory (DFT) Calculations. Geometry optimizations and single-point energy calculations of the molecular structures were performed with the Gaussian16 A03 program at the BP86/def-SVP level of theory. Taking into account the solvation effect of water, solvent model density (SMD) continuum models were utilized to simulate the coordination geometry of Cr$^{2+}$ with four typical complexing agents of oxalate (HOOCCOO$^-$, OA-H), tartrate (HOOC(CHOH)$_2$COO$^-$, TA-H), formate (HCOO$^-$, FA-H), and urea (CO(NH$_2$)$_2$, U-H). The resulting stable aquo-complexed chromium(II) are denoted as [Cr(OA-H)(H$_2$O)$_5$]$^+$, [Cr(TA-H)-(H$_2$O)$_5$]$^+$, [Cr(FA-H)(H$_2$O)$_5$]$^+$, and [Cr(U-H)(H$_2$O)$_3$]$^{2+}$, respectively. After deprotonation, the above four complexes are transformed to [Cr(OA)(H$_2$O)$_5$], [Cr(TA)(H$_2$O)$_5$], [Cr(FA-H)(H$_2$O)$_4$OH], [Cr(U-H)(H$_2$O)$_4$OH]$^+$, as revealed by the DFT calculations.

RESULTS AND DISCUSSION

Electrodeposition of Thick Hard Chromium Coating. As reported, both the current density and deposition time are important process parameters for trivalent chromium electroplating. The optimum current density range of the plating bath was determined by Hull cell tests. Figure 1a shows a Hull cell panel plated with chromium coating from the bath containing a ternary complexing agent of oxalic acid, tartaric acid, and urea (denoted as OA-TA-U). The shiny Cr-covered area on the left shows that chromium was successfully electroplated on the brass sheet and the optimum current density range was identified between 30 and 55 A/dm$^2$ according to eq 1. Thus, a reasonable current density of 35 A/dm$^2$ was chosen for the subsequent experiments. Figure 1b shows that the thickness of Cr coating increases linearly with deposition time within 15 min. Beyond 15 min, the coating thickness also increased as the plating proceeded, though at a slower rate. Hard chromium with a thickness exceeding 30 μm was obtained with a moderate deposition time of 35 min. The rate of chromium deposition was reported to diminish rapidly over deposition time and usually after several minutes of electrolysis no mass gain was observable. Nevertheless, the deposition rate in this work did not decrease significantly in the first 30 min, and thick deposits could be obtained upon prolonged deposition time. An obvious decline in deposition rate did occur at 30 min, and the coating barely grew after 35 min. The morphology of the Cr coating (Figure 1c) indicates that the Cr(III) electroplating features a typical nucleation–growth mechanism. Within the first 2 min of plating, ultrathin Cr layer grew uniformly on the substrate. From 5 to 20 min, the existing crystal nuclei grew over time and new ones were formed. Concomitantly, microscopic fissures were observed on the Cr deposits. As the plating proceeded, the coating morphology at 25 and 30 min became similar to that observed at 5 min, suggesting that large nodular-like structures (20 min) were covered by newly grown fine grains and the coating defects were significantly leveled off. This was due to the relatively low nucleation and growth rate, which was consistent with the observed rate decline in Figure 1b. At 35 min, the fissures were mostly healed and the coating became dense and uniform. Meanwhile an obvious decrease in current efficiency was also observed (Figure 1b). Figure 1d shows the crosssection microstructure of the coating and the corresponding elemental mapping images of Cu and Cr, confirming that a homogeneous and dense chromium coating with a favorable thickness of ca. 30 μm was successfully obtained after 30 min of electroplating. Compared with the rapid growth of Cr coating in hexavalent baths (typically 0.2–0.6 μm/min), the time duration for electroplating in the Cr(III) bath was notably prolonged.
Characterization of the Plating Solutions and the Cr Coatings. To further explore the practicality of the ternary complexing agent bath for HC plating, both the solution and the deposits were characterized and compared with the following three baths: (1) binary complexing agent bath containing oxalic acid and tartaric acid (denoted as OA-TA), (2) hard chrome bath containing formic acid and urea (FAU), and (3) decorative chrome bath containing formic acid (FA). Detailed composition of the four baths is listed in Table S1. The key parameters of the plating process and the Cr coatings are compared in Table 2. The optimum current density ranges for the baths of OA-TA, FA-U, and FA were determined by Hull cell tests (Figure S1), and thus the Cr plating was carried out at 45, 20, and 5 A/dm², respectively. The Cr(III) plating bath with the OA-TA-U ternary complexing agent exhibits the highest deposition rate of 1.01 μm/min. Its current efficiency was 19.18%, higher than that for a typical hexavalent chromium electrodeposition (12% ~ 18%). FA-U solution shows even higher current efficiency of 28.12% but both the deposition rate and the coating thickness are lower. Bath solution with FA alone exhibits minimal deposition rate and the coating could barely grow after 10 min of plating. The OA-TA-U solution not only yields the highest coating thickness but also the highest coating hardness of 866 HV (Figure S2). SEM and optical images (Figure 2a–d) show the coating obtained with the OA-TA-U solution exhibits smooth and shiny metallic appearance as well as uniform and dense micromorphology (Figure 2a). However, the coatings deposited from OA-TA or FA-U solution exhibited a rough surface and nodular structure with microscopic cracks (Figure 2b,c). Moreover, both coatings show a dull and dark surface, caused by a high surface roughness or the presence of impurities such as carbon and sulfur. The thin coating deposited from FA (Figure 2d) shows typical features of decorative chromium coating with shiny metallic luster, good adhesion, and low roughness. The homogeneous microstructure of this coating may also derive from its low thickness. Despite the appealing appearance, the extremely thin Cr coating is not suitable for anticorrosion and tribological applications. Therefore, the Cr coating deposited from the OA-TA-U solution is the only one to meet the two important requirements of hard chromium, i.e., sufficient thickness and desired appearance and microstructure. The crystal structure of the coatings was further probed by X-ray diffraction (XRD). As shown in Figure 2e, three peaks at 42.2°, 49.2°, and 72.2° are assigned to the (111), (200), and (220) facets of the brass substrate. The presence of these sharp substrate peaks in the samples from FA, FA-U, and OA-TA baths can be attributed to (1) the thickness of the coating being less than the penetration depth of the X-ray or (2) the presence of microscopic cracks within the Cr coatings. In contrast, these substrate peaks can be hardly detected in the sample from the OA-TA-U bath, confirming that high-quality thick Cr coatings were obtained. The magnified spectrum of OA-TA-U in Figure 2f shows a broad peak located at around 45°. This peak is attributed to amorphous/microcrystalline Cr (110), which is a typical feature of Cr coating electrodeposited from trivalent chromium baths. For the FA-U sample, the diffraction peaks of Cr are sharper, indicating a higher degree of crystallinity compared to the OA-TA-U sample. The higher crystallinity is probably a result of increased crystal size which leads to reduced compactness of the coating. The linear sweep voltammetry (LSV) curves (Figure 3a) reveal that the electroreduction process in Cr(III) baths consist of three regions, in good agreement with the literature. In the first potential region, from 0 to −0.6 V for the OA-TA-U and OA-TA baths and 0 to −0.75 V for the FA-U and FA baths, no cathodic current is detected because there is no Faradaic reaction and the electrode is ideally polarized. In the second region, from −0.6 to −1.1 V for the OA-TA-U and OA-TA baths and −0.75 to −1.3 V for the FAU and FA baths, the current increases with potential until reaching a limiting diffusion current plateau, indicating that Cr(III) reduction has occurred. At potentials more negative than −1.1 V and −1.3 V, respectively, a rapid increase of the curve slope is observed for all electrolytes and the main cathodic reaction is hydrogen evolution.
Compared with the baths of FA-U and FA, an obvious increase of the limiting diffusion current density was observed in the LSV curves of OA-TA-U and OA-TA, suggesting that higher current density can be employed in galvanostatic chromium plating, which is consistent with the results in Table 2. However, because of the lower current efficiency, more severe hydrogen evolution would occur in the OA-TA bath. The corrosion resistance of the coatings was evaluated with potentiodynamic polarization test coupled with electrochemical impedance measurement. The corrosion current density ($I_{\text{corr}}$) and corrosion potential ($E_{\text{corr}}$) are calculated from the intercept of the Tafel slopes (Figure 3b) and listed in Table S2. Among the four samples, chromium deposited from a OA-TA-U bath exhibits the lowest value of $I_{\text{corr}}$ and thus the lowest corrosion rate. The sample deposited from the FA bath exhibits the most positive value of $E_{\text{corr}}$, which can be interpreted as a mixed potential of chromium and brass corrosion resulting from the incomplete coverage of the substrate by the ultrathin chromium coating. All the four samples exhibit two capacitive loops in the Nyquist plots (Figure 3c and inset). The bode plots (Figure S3) reveal two time constants for the thin Cr coating from the bath containing FA alone. In contrast, the rest of the three samples show three time constants because the defects (e.g., micropores and microcracks) result in more complex reaction interfaces in thicker chromium coatings. Therefore, two different equivalent circuits (Figure 3d,e) are used to fit the experimental data of the thick and thin chromium coatings, respectively. In both models, $R_S$ is the electrolyte resistance. $CPE_{dl}$ and $R_{\text{ct}}$ are the double layer capacitance and the charge transfer resistance of the corrosion reaction. $R_1$ and $CPE_1$ represent the resistance and capacitance of an oxide layer. $R_2$ and $CPE_2$ are the resistance and capacitance associated with porous structure and defects of the coating. The Warburg impedance $W$ is a diffusion impedance element. The fitted curves from the respective equivalent circuit show high conformity with the experimental data (Figure S4). The simulated equivalent circuit parameters for the four samples are listed in Table S3. The coating deposited from the OA-NAU bath exhibits significantly higher $R_{\text{ct}}$ value than the other three, verifying its better corrosion resistance.

**Mechanism for Deposition of Thick Chromium Coatings from Cr(III) Baths.** The above characterization indicates that the bath containing an OA-TA-U ternary complexing agent has clear advantages in depositing thick Cr coatings. The underlying mechanisms need to be further elucidated. In aqueous solution, $\text{Cr}^{3+}$ is coordinated with six water molecules to form $[\text{Cr}(\text{H}_2\text{O})_6]^{3+}$ which exhibits regular-octahedron structure with $\text{Cr}^{3+}$ located at the center. The electrons on the cathode can hardly reach the inner d-orbits of $\text{Cr}^{3+}$ during the electrodeposition because of the surrounding water molecules. Ionic complexing agents ($L^-$), such as HCOO$^-$, HOOCCOO$^-$, and $\text{C}_2\text{H}_4\text{NO}_2^-$, can compete with water molecules to coordinate with $\text{Cr}^{3+}$, forming cocoodinated $[\text{CrL}(\text{H}_2\text{O})_5]^{2+}$. The large-sized ionic ligand generates a steric hindrance effect so $[\text{CrL}(\text{H}_2\text{O})_5]^{2+}$ presents an irregular-octahedron structure, in which the distances between $\text{Cr}^{3+}$ and the surrounding water molecules are expanded (eq 2). Therefore, upon addition of the ligand, $\text{Cr}^{3+}$ ions gain immediate contact with the cathode, i.e., easier to receive electrons from the cathode. Previous studies have suggested that electrodeposition of chromium involves two consecutive steps: one-electron reduction of trivalent chromium complex $[\text{Cr(III)L(H}_2\text{O})_5]^{2+}$ (eq 3), followed by a two-electron reduction of the as-formed divalent chromium complex $[\text{Cr(II)L(H}_2\text{O})_5]^+$ to metallic Cr (eq 4). Concomitantly, hydrogen evolution reaction in eq 5 occurs at the same applied potential, leading to an increase of local $\text{pH}$. 
\[ [\text{Cr(III)}(\text{H}_2\text{O})_6]^{3+} + \text{L}^- \rightarrow [\text{Cr(III)}\text{L}(\text{H}_2\text{O})_3]^{2+} + \text{H}_2\text{O} \]  
\[ (2) \]

\[ [\text{Cr(III)}\text{L}(\text{H}_2\text{O})_3]^{2+} + \text{e}^- \rightarrow [\text{Cr(II)}\text{L}(\text{H}_2\text{O})_3]^+ \]  
\[ (3) \]

\[ [\text{Cr(II)}\text{L}(\text{H}_2\text{O})_3]^+ + 2\text{e}^- \rightarrow \text{Cr}^0 + 5\text{H}_2\text{O} + \text{L}^- \]  
\[ (4) \]

\[ 2\text{H}^+ + 2\text{e}^- \rightarrow \text{H}_2(\text{g}) \]  
\[ (5) \]

The intermediate divalent chromium complex is of great significance during the deposition process.\textsuperscript{14,15} Due to the high pH values in the vicinity of the cathode, the coordinated water molecules of the divalent chromium complex could be partially converted to OH groups (eq 6). In such a situation, the metal electrodeposition proceeds with the participation of the electroactive hydroxo-complex of divalent chromium (eq 7).\textsuperscript{44} However, long-term electrolysis gives rise to excessive formation of an unstable hydroxo-complex of divalent chromium (i.e., [Cr(II)L(H\textsubscript{2}O\textsubscript{4}OH)], which can be easily converted into a hydroxo-bridged dimer through olation reaction (eq 8). Once the oligomers are formed, the reaction can go

\[ 2[\text{Cr(II)}\text{L}(\text{H}_2\text{O})_3\text{OH}] \rightarrow [\text{L}(\text{H}_2\text{O})_3\text{Cr(II)}] + [\text{Cr(II)}(\text{H}_2\text{O})_3\text{L}] + 2\text{H}_2\text{O} \]  
\[ (8) \]

further to produce inert polymers of higher molecular weight which act as a barrier that hinders the deposition process.\textsuperscript{16,48} Therefore, it is reasonable to speculate that the suppression of nonelectroactive intermediate of divalent chromium may be the critical factor in sustained deposition of chromium. To shed light on the effects of different complexing agents on the redox abilities of intermediate chromium(II) complexes, DFT calculations were performed.

\[ [\text{Cr(II)}\text{L}(\text{H}_2\text{O})_3]^+ \rightarrow [\text{Cr(II)}\text{L}(\text{H}_2\text{O})_4\text{OH}] + \text{H}^- \]  
\[ (6) \]

\[ [\text{Cr(II)}\text{L}(\text{H}_2\text{O})_4\text{OH} + 2\text{e}^- \rightarrow \text{Cr}^0 + 4\text{H}_2\text{O} + \text{L}^- + \text{OH}^- \]  
\[ (7) \]

The optimized geometrical structures of divalent chromium complexes and the corresponding visual HOMOs are shown in Figure 4a. Unlike the octahedral geometry of the sixcoordinated Cr(III) complex, the Cr(II) atoms are five coordinated and present the geometry of a distorted trigonal bipyramid. In the Cr(II) complexes of [Cr(OA-H)(\text{H}_2\text{O})_5]^+, [Cr(TA-H)(\text{H}_2\text{O})_5]^+, and [Cr(U-H)(\text{H}_2\text{O})_5]^{2+}, the organic ligands (oxalate, tartrate, and urea) bind to the metal center in a bidentate fashion, with three water molecules taking part in the coordination. In the case of formic acid, Cr(II) ion coordinates with one monodentate formate ligand and four water molecules to form the [Cr(FA-H)(\text{H}_2\text{O})_5]^{2+} complex. The calculated distances between the uncoordinated water molecules and the Cr(II) ion are all larger than 3 Å, implying negligible complexation abilities toward Cr(II). The HOMOs of five-coordinated Cr(II) complexes are mainly contributed by the dz\textsuperscript{2} orbitals of chromium and the 2p orbitals of the coordinated oxygen, leading to dsp\textsuperscript{3} hybridization and a distorted trigonal bipyramid.
To evaluate the electron-accepting ability of the divalent chromium complexes, vertical electron affinity (VEA) and adiabatic electron affinity (AEA) analysis has also been carried out based on the optimized geometries.\textsuperscript{49,50} VEA is calculated as the energy difference between the cationic and neutral chromium species with both at the optimized geometry of the cations (E1 – E2), while AEA is calculated as the energy difference between the cationic and neutral chromium species at their respective optimized geometries (E1 – E3) (Figure 4c). The calculated VEA and AEA are all positive (Table S4), indicating that the neutral species is lower in energy than the corresponding cationic species, hence the divalent chromium complexes are unstable and prone to accepting electrons. Moreover, [Cr(OA-H)(H$_2$O)$_5$]$^+$ and [Cr(TA-H)(H$_2$O)$_5$]$^+$ have higher values of VEA and AEA, thus a stronger ability to accept electrons and can be more easily reduced on the cathode. The fast electroreduction of Cr(II) in the bath containing oxalic acid or tartaric acid may effectively reduce the generation of chromium(II) hydroxo-complexes which have an adverse effect on sustained electrodeposition by formation of dimers or polymers (eq 8).

As discussed above, the long-term electrolysis often leads to deactivation of chromium(II) complexes. The effects of hydrogen evolution reaction and associated pH change on redox abilities of chromium(II) complexes need to be taken into account. At high pH, the divalent chromium complexes of [Cr(OA-H)(H$_2$O)$_5$]$^+$, [Cr(TA-H)(H$_2$O)$_5$]$^+$, [Cr(FA-H-)(H$_2$O)$_5$]$^+$, and [Cr(U−H)(H$_2$O)$_5$]$^{2+}$ tend to lose a proton to form new intermediate species (denoted as [Cr(OA)- (H$_2$O)$_5$], [Cr(TA)(H$_2$O)$_5$], [Cr(FA-H) H$_2$O)$_4$OH], and [Cr(U−H)(H$_2$O)$_4$OH]$^+$, respectively). Their optimized geometrical structures and corresponding visual HOMOs are shown in Figure 4b. Protons are preferentially lost from the organic ligand (OA-H and TA-H) in the [Cr(OA-H)(H$_2$O)$_5$]$^+$ and [Cr(TA-H)(H$_2$O)$_5$]$^+$ complexes. In the case of [Cr(FAH)( H$_2$O)$_5$]$^+$ and [Cr(U−H) (H$_2$O)$_4$OH]$^{2+}$ complexes, coordinated water molecules lose a proton to generate a OH ligand. Thus, the hydroxo-complexes (i.e., [Cr(FA-H)(H$_2$O)$_4$OH] and [Cr(U−H)(H$_2$O)$_4$OH]$^+$) can be largely generated in the bath containing formic acid or urea with the increase of pH value. The deprotonation products of [Cr(OA)(H$_2$O)$_5$], [Cr(TA)(H$_2$O)$_5$], and [Cr(FA-H) (H$_2$O)$_4$OH] maintain the five-coordinated distorted trigonal bipyramid configuration. However, as shown in Figure S5, different from the five coordinated [Cr(U−H)(H$_2$O)$_5$]$^{2+}$ complex, the Cr(II) ion in the [Cr(U−H)(H$_2$O)$_4$OH]$^+$ complex is four-coordinated with one monodentate urea ligand, one hydroxyl ligand, and two water molecules, which presents the geometry of square planar with dsp$^2$ hybridization. Its HOMOs are mainly contributed by the d($x^2$−$y^2$) orbitals of chromium and the 2p orbitals of the coordinated oxygen.
The calculated VEA values of divalent chromium complexes before and after losing a proton are summarized in Figure 4d and Table S4. In comparison to $[\text{Cr}(L-H)(\text{H}_2\text{O})_5]^n$ (L represents OA, TA, FA, or U, $n = 1$ or 2), the corresponding $[\text{Cr}(L)(\text{H}_2\text{O})_5]^{(n-1)}$ or $[\text{Cr}(L-H)(\text{H}_2\text{O})_4\text{OH}]^{(n-1)}$ all show a reduced value of VEA, suggesting a lower electron-accepting ability. Thus, the deprotonation products are less electroactive, and their cathodic reduction is more difficult. Furthermore, the formate hydroxo complex (e.g., $[\text{Cr}(\text{FA}-\text{H})(\text{H}_2\text{O})_4\text{OH}]$) could be largely accumulated and hence the polymerization reaction occurs, which hinders the reduction process of chromium(II). Different from the formate ligand, although the urea hydroxo complex of $[\text{Cr}(\text{U}-\text{H})(\text{H}_2\text{O})_4\text{OH}]^+$ exhibits the lowest VEA value, the thickness of Cr coating from the OA-TA bath is lower than the OA-TA-U bath. This is probably due to the different bonding type of $[\text{Cr}(\text{U}-\text{H})(\text{H}_2\text{O})_4\text{OH}]^+$ (dsp² hybridization instead of dsp³). The new structure has a more pronounced steric hindrance effect which impedes the formation of hydroxo bridged complex through olation reaction.

Therefore, as shown in Figure 5, there are generally two routes for the electroreduction of Cr(II) complexes to metallic Cr⁺: (i) a direct route through highly electroactive $[\text{Cr}(\text{LH})(\text{H}_2\text{O})_5]^n$ complexes and (ii) an indirect route through the less electroactive $[\text{Cr}(\text{L})(\text{H}_2\text{O})_5]^{(n-1)}$ or $[\text{Cr}(\text{L-H}-)(\text{H}_2\text{O})_4\text{OH}]^{(n-1)}$ complexes. In the traditional bath containing formic acid (Figure 5a), because of the low VEA value of $[\text{Cr}(\text{FA}-\text{H})(\text{H}_2\text{O})_5]^+$ and $[\text{Cr}(\text{FA-H})(\text{H}_2\text{O})_4\text{OH}]^+$, chromium-(II) hydroxo complex can be largely generated and accumulated after a few minutes of electrodeposition (route 2), leading to the formation of inert polymers through an olation reaction. Inversely, the adsorbed polymers act as a barrier that dramatically decrease the diffusion of ions from bulk solution to the electrode surface, further hindering the electroreduction of reactive chromium(II) complexes (route 1). As a result, the current efficiency drops and the coating thickness could barely increase. In contrast, in the bath containing the ternary complexing agent (Figure 5b), the adverse olation reactions could be greatly alleviated and thus the diffusion of chromium ions is not conspicuously impeded, which can also be evidenced by the enhanced limiting diffusion current density shown in Figure 3a. At the initial deposition stage (the first 15 min of electroplating in Figure 1b), the electroactive chromium(II) complexes ([Cr(OA-H)(H_2O)_5]^+, [Cr(TA-H)(H_2O)_5]^+, and [Cr(U-H)(H_2O)_4OH]^+) are rapidly reduced through route 1 and the deposition rate of chromium does not decrease. After a period of electroplating (15−30 min of electroplating in Figure 1b), small amounts of deprotonated intermediate products ([Cr(OA)(H_2O)_5], [Cr(TA)(H_2O)_5], and [Cr(U-H)(H_2O)_4OH]^+) are also formed through route 2. These less reactive Cr(II) complexes cannot accumulate in large quantities because of its moderate electroreduction to metallic chromium. The deposition rate shows a trend of slight decrease in this stage due to the low VEA value of the deprotonated complexes. However, after the long-term electrolysis (30 min or longer), large amounts of deprotonated complexes would present in the bath due to the increased pH value. At this point, chromium(II) hydroxo complex could be generated through further deprotonation and harmful polymerization reactions may occur. The electroplating rate becomes extremely slow and the chromium coating virtually stops growing.

It is also observed that thick chromium coating with uniform appearance can be hardly obtained in the trivalent chromium bath just containing a single ligand of oxalic acid, tartaric acid, or urea (Figures S6 and S7). The formation of a less electroactive hydroxo-complex decreases the throwing power and covering power of the bath, resulting in uneven distribution of the cathodic current density. The multicomponent bath such as the OA-TA-U may strengthen the ligand exchange reaction and cause competitive complexation of Cr(II) with organic ligands and hydroxy ligands, which further suppresses the hydroxo-complex formation upon sustained electroplating.
CONCLUSIONS
In summary, a novel trivalent bath comprising oxalic acid, tartaric acid, and urea (OA-TA-U) is proposed for electroplating of hard chromium. Shiny metallic coatings with thickness exceeding 30 μm can be obtained from the bath, and the current efficiency of Cr deposition does not significantly decrease during relatively long-term electrolysis. Compared with OA-TA, FA-U, and FA bath, coatings deposited from OA-TA-U solution exhibits advantages in deposition rate, microstructure, hardness, and corrosion resistance. First-principles calculations suggest that rational selection of complexing agents promotes formation of intermediate Cr(II) complexes with higher value of VEA and AEA. Furthermore, the suppression of formation of nonelectroactive hydroxo bridged complexes is found to be the critical factor in sustained Cr(III) deposition. In this regard, the ternary complexing agent offers two benefits: (1) reducing the formation of inactive chromium(II) hydroxo-complex because of the competitive complexation between organic and hydroxyl ligands and (2) further inhibiting the olation reaction due to the synergistic steric hindrance effect of chromium(II) hydroxo-complex. This work provides a mechanistic understanding of the Cr(III) electroreduction process, as well as a practical guidance on electrolyte design for hard chromium plating.

ACKNOWLEDGEMENTS
This work received financial support from the Science and Technology Bureau of Shenzhen (Grant No. JCYJ20170306171540744) and the Science and Technology Bureau of Ningbo (Grant No. 201501CX-C01006).

CONTENT
Supporting Information
Chemical composition of plating baths; corrosion current density and corrosion potential; simulated equivalent circuit parameters; VEA and AEA value of bivalent chromium complexes; Bode and Nyquist plots; SEM and the corresponding optical images of the Cr coatings.

Figures and Tables
Table 1. Basic composition of trivalent Cr plating bath

| Bath composition                   | Function                | Concentration (mol·L⁻¹) |
|------------------------------------|-------------------------|-------------------------|
| Chromium sulfate (Cr₂(SO₄)₃·6H₂O) | Source of Cr            | 0.4                     |
| Sodium sulfate (Na₂SO₄)            | Conducting salt         | 0.3                     |
| Aluminum sulfate (Al₂(SO₄)₃)      | Conducting salt         | 0.15                    |
| Boric acid (H₃BO₃)                | Buffer agent            | 0.65                    |
| Oxalic acid (H₂C₂O₄)              | Complexing agent        | 0.25                    |
| Tartaric acid (C₄H₆O₆)            | Complexing agent        | 0.15                    |
| Urea (CH₄N₂O)                     | Complexing agent        | 0.3                     |

Figure 1. Electrodeposition of hard chromium coating from trivalent chromium bath containing the ternary complexing agent. (a) Determination of the optimum current density range of the plating bath by Hull cell test. (b) Correlation of Cr coating thickness with deposition time. (c) SEM images of chromium coatings obtained at 35 A/dm² with different deposition time. (d) Cross-section microstructure and the corresponding Cr and Cu elemental mapping images of the chromium coating obtained at deposition time of 30 min.
Table 2. Comparison of the main characteristics of different trivalent chromium baths and the resulting Cr coatings.

| Solution | Current density (A/dm²) | Deposition time (min) | Coating Thickness (µm) | Coating Hardness (HV) | Deposition rate (µm/min) | Current efficiency (%) |
|----------|-------------------------|-----------------------|------------------------|-----------------------|--------------------------|------------------------|
| OA-TA-U  | 35                      | 30                    | 30.2                   | 866                   | 1.01                     | 19.18                  |
| OA-TA    | 45                      | 30                    | 20.9                   | 792                   | 0.70                     | 10.33                  |
| FA-U     | 20                      | 30                    | 25.3                   | 630                   | 0.84                     | 28.12                  |
| FA       | 5                       | 10                    | <1                     | -                     | < 0.1                    | -                      |

Figure 2. Morphology and composition characterization of chromium coatings. SEM and the corresponding optical images (inset) of the Cr coatings obtained from (a) OA-TA-U bath, (b) OA-TA bath, (c) FA-U bath, and (d) FA bath. (e) X-ray diffraction patterns of Cr coatings deposited from different baths. (f) The magnified XRD spectrum of OA-TA-U in (e).
Figure 3. Electrochemical characterization of the plating solutions and the chromium coatings. (a) LSV curves of chromium electrodeposition from different baths. (b) Tafel plots and (c) Nyquist plots of the chromium coatings deposited from different baths. Equivalent circuits of impedance behavior of the chromium coatings electrodeposited from the (d) FA bath and (e) OA-TA-U, OA-TA, and FA-U baths. Both the Tafel curves and impedance measurements were conducted in 3.5% NaCl solution at room temperature.
Figure 4. Optimized geometrical structures and visual HOMOs of divalent-Cr complexes of (a) [Cr(L-H)(H_2O)_5]^n and (b) [Cr(L)(H_2O)_5]^{n-1} or [Cr(L-H)(H_2O)_4OH]^{n-1}, L represents OA, TA, FA, or U, n = 1 or 2. (c) Schematic diagram for the calculation of vertical electron affinity (E1 – E2) and adiabatic electron affinity (E1 – E3). (d) Summary of vertical electron affinity energy of divalent-Cr complexes before and after losing a proton.
Figure 5. Schematic illustration of electroreduction process of chromium(II) complexes in trivalent chromium bath containing (a) a formic acid complexing agent and (b) a ternary complexing agent. (i) [Cr(FA-H)(H₂O)₅]⁺, (ii) [Cr(FA-H)(H₂O)₃OH], (iii) [Cr(OA-H)(H₂O)₅]⁺, (iv) [Cr(OA)(H₂O)₅], (v) [Cr(TA-H)(H₂O)₃], (vi) [Cr(TA)(H₂O)₃], (vii) [Cr(U−H)(H₂O)₃]²⁺, and (viii) [Cr(U−H)(H₂O)₄OH]⁺.

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