Deposition of Nanocrystalline Zinc-Nickel Alloys by D.C. Plating in Additive Free Chloride Bath

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Nanocrystalline zinc-nickel alloy coatings were electro-deposited in chloride bath containing ZnCl2 (50–200 gL−1), NiCl2·6H2O (50–200 gL−1) and H3BO3 (40 gL−1) at 45°C. The initial alloy composition, as investigated by voltammetry, chronopotentiometry or EDX analysis, exhibited an pronounced dependence on the composition of plating bath. However, the alloy composition tended to approach an upper limit of 18 atomic percent when deposits grew thicker, for most of the plating bath compositions. The process was clearly the anomalous codeposition, which was considered to be due to the formation of an adsorbed layer of Zn(OH)2 containing nickel cations, as an intermediate product. The dissolution potentials of the fcc, and (pure) porous nickel (fcc) with increasing order of phase (Ni5Zn21) for about 12–18 atomic percent nickel and a mixture of γ- and ♂-phases (hcp) for nickel atom percent below 8. The grain size decreased from about 50nm to 20nm with decrease in nickel content away from the stoichiometric composition of Ni5Zn21 along with a decrease in hardness from about 450KHN to 150KHN. The decrease in hardness with a decrease in grain size seems attributable to the grain boundary sliding as predominant deformation mechanism. Anodic dissolution of the alloys involved pronounced dezincification, forming α-phase (fcc) that dissolved at the end. [doi:10.2320/matertrans.MER2007022]

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1. Introduction

Electrodeposition is a versatile materials processing technique finding applications ranging from miniature electronic devices to giant industrial structures as functional and/or aesthetic coatings. Both aqueous1) and non aqueous2,3) solutions have been employed for the purpose. However, the aqueous solutions have the special advantage of room temperature operation and are, therefore, preferred for large scale applications.

Zinc has been the most widely used sacrificial coating for the protection of steel from corrosion. The addition of small quantity of iron group metals such as nickel (10–18%) to zinc deposits has been known to increase its corrosion resistance along with an improvement in mechanical properties.4) In salt spray tests, the ability of electro-deposited Zn-Ni alloy coatings to protect steel from corrosion is found to be 3–4 times better than pure zinc coating, although the cost is only 2 times higher. On the other hand, the hardness of electro-deposited zinc alloys containing 10-20% nickel is 3–6 times higher than that of pure zinc deposit.5) Ductility of these alloys is also better than zinc.6) Improved corrosion resistance and hardness (wear resistance) also allow using relatively thinner deposits required for better formability and weldability.7) The alloy coatings also allow spot welding because of higher melting temperature of the intermetallic phases.8)

Mostly, the electrolytic codeposition of Zn-Ni alloys is anomalous in nature which means the deposition of a relatively noble metal, i.e., nickel, is hindered before the deposition of other alloy constituent, i.e., zinc, also starts taking place.9) Different alkaline and acid type plating baths are in use.10) The acid type baths are further subdivided into chloride type and sulfate type baths. The sulfate type baths are used for high-speed alloy plating with simple shapes whereas chloride baths are preferred for higher throwing power to plate intricate shapes.11) The higher micro-throwing power of chloride bath also ensures an improved corrosion resistance.12)

Apart from electroplating, a number of authors have performed electrochemical studies on Zn-Ni system, involving voltammetry or chronopotentiometry. However, any systematic study to explain the anodic peaks (waves) in the voltammograms and anodic plateaus in the chronopotentiograms remained absent after the work of Swathirajan,9) who was mostly followed to describe the dissolution as occurring in three main stages, namely dezincification/dissolution of η- (ZnNi), hcp plus γ- (Ni5Zn21), bcc phases, α-phase (NiZn), fcc, and (pure) porous nickel (fcc) with increasing order of dissolution potentials. The dissolution potential of the presumed porous nickel, as formed by prior dezincification, was significantly lower than that of the electrodeposited nickel. Recently, Petrauskas et al.10,11) identified dissolution of β- (NiZn) at the potentials previously assigned to the dissolution of porous nickel. Therefore, it is required that more investigations are performed to clarify the dissolution behavior of Ni-Zn alloys in different electrolytes.

Among chloride baths for deposition of Zn-Ni alloys, addition of ammonium chloride or acetate has been common. Other additives are also added for improving the leveling. On the other hand, Erb et al.12,13) has employed boric acid without addition of ammonia or acetate. They were able to prepare good deposits by pulse plating. They were also able to control the grain size to nano-scale range. It is worth-mentioning that nanocrystalline alloys exhibit extremely high corrosion resistance comparable to extremely homogeneous amorphous counterparts,14–19) and extra-ordinary high strength with reasonable ductility depending on the grain...
size and deformation mechanism.20,21) Electrochemical study and DC plating in the bath based on zinc and nickel chlorides and boric acid without any addition has not yet been focused upon to the best of our knowledge.

The purpose of the present study is to explore the electrodeposition of nanocrystalline Zn-Ni alloys by DC plating in the additive free chloride baths. Particular emphasis has been laid on effect of concentrations of electroactive species (salts) on the composition, texture and quality of the deposit in addition to anodic dissolution behavior.

2. Experimental

Electrochemical measurements and electrodeposition were performed using Potentiostat/Galvanostat AMEL Model-2051 and Function Generator AMEL Model-568. Different compositions of plating bath were employed as shown in Table 1. For referring to the bath compositions, concentration of ZnCl₂ and NiCl₂·6H₂O will be written in the parenthesis as numerator and denominator, respectively, wherever found necessary. The analytical grade chemicals were used along with doubly distilled water to prepare the solutions. The pH was measured by a pH meter, (Model PH 330/SET – 0, 82362 WTW, Germany). For all measurements, the initial pH of the bath was 3.6 ± 0.1 and it remained almost same during whole experiment. All the experiments were performed in a 100 ml beaker with an electrolyte volume of about 80 ml. No agitation was used and the temperature was maintained at 45 ± 0.5 °C by placing the cell in a hot water bath.

For electrochemical studies (cyclic voltammetry & reverse chronopotentiometry), a platinum strip (with an exposed area of 1 cm²) was used as a working electrode. It was cleaned with conc. H₂SO₄, and degreased with acetone using ultrasonic vibrator followed by rinsing in distilled water. Another platinum strip and a saturated calomel electrode (SCE) were used as counter and reference electrodes, respectively. The scan rate for cyclic voltammetry was 0.05 Vs⁻¹. All the potentials in this paper are presented with respect to the SCE.

For electrodeposition of alloy coatings, commercial purity cold rolled copper sheet with an exposed area of 2 cm² was used as a substrate. The copper substrate was mechanically polished using emery papers up to fine grit of 1200, followed by ultrasonic cleaning in acetone. Before deposition, the copper substrate was chemically etched in 10% sulfuric acid for about 5–6 seconds followed by rinsing with distilled water. Perfect adhesion of the deposits with the substrate was thus ensured. Chemical composition of the electrodeposits was determined using Energy Dispersive X-ray Spectroscopy (EDS) system attached with Scanning Electron Microscope (SEM), LEO 440. The analysis was performed at 2 or 3 locations on the surface of the sample. The variation of composition on a given sample was found to be less than 1%. Average values, thus obtained, have been reported in the results. The compositions of the alloys presented in the following sections are given in atomic percent. X-ray diffractometer (XRD, Rigaku Model DMAX III) with Cu Kα radiation was used to obtain XRD patterns using standard θ-2θ geometry. X-Ray peak broadening was used to estimate the grain size with the help of Scherrer’s formula.22)

3. Results

3.1 Electrochemical study

3.1.1 Voltammetry

Figure 1 shows typical cyclic voltammograms obtained on platinum electrode in baths 1 and 2 at 45 °C. The scan rate was 0.05 Vs⁻¹ and the reversal potential was −1.25 V (In the parenthesis, concentrations (gL⁻¹) of ZnCl₂ and NiCl₂·6H₂O have been presented in the form of numerator and denominator, respectively).

![Image](https://via.placeholder.com/150)

Table 1 Composition of plating baths employed for the study.

<table>
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<th>No</th>
<th>Conc. of ZnCl₂ (gL⁻¹)</th>
<th>Conc. of NiCl₂·6H₂O (gL⁻¹)</th>
<th>Boric acid (gL⁻¹)</th>
<th>Referred in the text as</th>
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<td>40</td>
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<td>0</td>
<td>200</td>
<td>40</td>
<td>Bath 2 (0/200)</td>
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<td>200</td>
<td>200</td>
<td>40</td>
<td>Bath 3 (200/200)</td>
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<td>200</td>
<td>50</td>
<td>40</td>
<td>Bath 4 (200/50)</td>
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<td>40</td>
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<td>50</td>
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<td>Bath 6 (50/50)</td>
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<td>100</td>
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<td>Bath 8 (150/150)</td>
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<td>100</td>
<td>40</td>
<td>Bath 9 (200/100)</td>
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<tr>
<td>10</td>
<td>100</td>
<td>200</td>
<td>40</td>
<td>Bath 10 (100/200)</td>
</tr>
</tbody>
</table>

Fig. 1 Typical cyclic voltammograms obtained on platinum electrode in baths 1 and 2 at 45 °C. The scan rate was 0.05 Vs⁻¹ and the reversal potential was −1.25 V (In the parenthesis, concentrations (gL⁻¹) of ZnCl₂ and NiCl₂·6H₂O have been presented in the form of numerator and denominator, respectively).
The dissolution potential of the alloy constituents depend on the alloy composition. This is a typical hindrance in the deposition of nickel. This is in agreement with a similar shift of reversal potential. Therefore, the nature (and the chemical composition) of the deposit seems to remain same in spite of extending the deposition potential in the negative direction, suggesting no appreciable effect of the deposition potential on the alloy composition. This behavior may also be related with anomalous codeposition.

3.1.2 Chronopotentiometry

Figure 4(a) shows typical reverse chronopotentiograms obtained on platinum in different solutions. Three anodic plateaus related with alloy dissolution are observed, namely I, II, and III, which seem to correspond to the main anodic peaks of the voltammograms, i.e., A, B, and C, respectively. The plateau I in case of bath 4 appears at the same potential where pure zinc is dissolved in bath 1. Hence, it corresponds to the dissolution of η-phase (which is η-Zn (hcp) containing dissolved nickel). The same plateau appears at slightly higher potentials in the other baths containing higher concentration of nickel chloride, indicating a gradual rise in the nickel content of the η-phase with an increase in nickel concentration in the bath. This is in agreement with a similar shift of peak A in the voltammograms (Fig. 2). Furthermore, the relative amount of η-phase decreases with increase in nickel concentration of the plating bath as indicated by the decrease in charge associated with plateau I in chronopotentiograms. These observations are in agreement with voltammetry.

Figure 4(b) shows a typical reverse chronopotentiogram obtained in bath 9. The deposition time was 1.8 ks. The plateau I is almost absent, which suggests that the η phase does not form after electrodeposition for 1.8 ks in this bath, in contrast to the deposition for 180 s. In order to determine the phases that dissolve at the plateau II and III, various samples were prepared by deposition for 1.8 ks in the same solution followed by dissolution for various time intervals, as corresponding to 1-5 positions indicated on the chronopotentiogram (Fig. 4(b)). The XRD patterns exhibited by these
samples along with their compositions are shown in Fig. 4(c). The \( \eta \)-phase is not observed after electrodeposition for 1.8 ks, in agreement with the chronopotentiogram (Fig. 4(b)). It may be noticed that \( \gamma \)-phase (an intermetallic phase also known as \( \text{Ni}_5 \text{Zn}_{21} \)) is present at plateau II, although the nickel content of the deposit reaches above 40% as examined by EDX analysis. Thedezincification of \( \gamma \)-phase is also confirmed from shift of X-ray reflections towards higher angles as shown in Fig 4(c). The alloy composition at plateau III lies above 60% Ni. The corresponding XRD pattern indicates the formation of \( \alpha \)-phase (i.e., \( \alpha \)-Ni (fcc) with large concentration of zinc in the solid solution). Hence the dezincification of \( \gamma \)-phase at plateau II leads to the formation of \( \alpha \) phase. In turn, the \( \alpha \) phase undergoes further dezincification at plateau III. Considering this dissolution behavior, the main anodic peaks B and C in the voltammograms (Fig. 2) can also be assigned to the dezincification and dissolution of \( \gamma \) and \( \alpha \)-phases, respectively.

3.2 Electroplating

The alloy composition after electroplating at 700 Am\(^{-2}\) in different solutions is shown in Fig. 5. The nickel content of the alloys tends to increase with the time of deposition. It is also influenced by bath composition particularly for short-time deposition, in agreement with voltammetry and chronopotentiometry. Nevertheless, the maximum nickel content in the alloy is about 18% as observed in a number of baths for deposition times of 1.08 and 1.8 ks. Consequently, about 18% nickel seems to be an upper limit in the additive free chloride baths employed in this work.

Figure 6 shows typical XRD patterns of the alloy deposits from different baths at a current density of 700 Am\(^{-2}\), as a function of deposition time.
shown in Fig. 6 (a-d) typically exhibit the formation of only γ
d phase in a variety of baths. This may suggest that η-phase is
formed in these baths only at initial stages and then it dies out
due to preferential growth of γ-phase, due to increase in
nickel content of alloy with deposition time. In bath 4, which
contains ZnCl₂ (200 gL⁻¹) and NiCl₂·6H₂O (50 gL⁻¹), both
η and γ-phases are obtained in thick deposits as shown in
Fig. 6(e). This may be related with the fact that nickel
content of the electrodeposits decreases with deposition time
(Fig. 5), in contrast to general behavior of other baths. The
nickel rich α-phase is not found in the electrodeposits. Hence,
it is confirmed that α-phase forms only by anodic dezinci-
fication of γ-phase during cyclic voltammetry and reverse
chronopotentiometry. It may be noticed that the relative
intensities of (330) and (600) reflections of γ-phase vary,
suggesting a change in the preferred texture from one

electrodeposit to the other. In general, (600) reflection increases with deposition time (Fig. 6(a-d)). When different solutions are compared, it is strongly revealed that the nickel-rich baths favor (600) texture while the zinc-rich baths favor the (330) texture. This is true even for the electrodeposits with similar composition but obtained in different baths.

Typical SEM images of the deposits are shown in Fig. 7. Relatively smoother deposits accompanied by finer morphological features are formed in the nickel rich baths. The surface patterns primarily correspond to the protuberances resulting from the growth pattern of the deposits. It was not possible to resolve grain structure by SEM. Figure 7 clearly shows that the deposit quality strongly depends on the composition of plating bath, although it has less pronounced effects on the alloy composition.

The grain sizes were estimated from X-ray peak broadening with the help of Scherrer’s formula as given below:

\[
d = \frac{0.94 \lambda}{B \cdot \cos \theta}
\]

where
\[
d = \text{grain size (nm)}
\]
\[
\lambda = \text{wavelength of X-ray used (nm)}
\]
\[
\theta = \text{diffraction angle (degrees)}
\]
\[
B = \text{full width half maximum (FWHM) of X-ray diffraction peak (rad.)}
\]

As shown in Fig. 8, the grain size varies with the alloy composition although it remains in the nano-scale range. The grain size decreases with a decrease in nickel content of the deposit. In this way, the smaller grain size of about 24 nm has been obtained in \(\gamma\)-phase when it is highly deficient in nickel with respect to its stoichiometric requirement (19.3% Ni). Hence, large concentration of point defects (substitutional and/or vacancies) seems attributable to a decrease in the grain size.

Hardness is also found to be a function of alloy composition as shown in Fig. 9. It is observed that with increasing amount of nickel in the deposit, the hardness increases sharply from 150KHN at 12% Ni to 450KHN near stoichiometric composition of \(\text{Ni}_5\text{Zn}_{21}\). Comparison of Figs. 8 and 9 indicates that the hardness increases with an increase in grain size. Hence, the behavior is opposite to Hall-Petch relationship, suggesting the grain boundary sliding as a primary mode of deformation for these nanocrystalline alloys.

4. Discussion

4.1 Nanocrystallinity of the deposit

Alfantazi and Erb et al.\(^ {12} \) prepared nanocrystalline Zn-Ni coatings in the same electrolyte by pulse plating. However,
we have been successful in obtaining nanocrystalline alloys with a grain size as low as 20 nm, with simple DC plating at 700 Am\textsuperscript{-2}. The grain size has been found to be a strong function of the alloy composition. It reduces with a decrease in the nickel content and deviation from the stoichiometric composition of Ni\textsubscript{3}Zn\textsubscript{21}. It appears that the point defects, formed due to deviation from stoichiometric composition, are mostly accommodated at the grain boundaries resulting in an increased grain boundary area accompanied by a smaller grain size.

There are two modes of deformation of the nanocrystalline alloys that can be identified by the variation of hardness with grain size.\textsuperscript{26} For coarse grained alloys, hardness increases with a decrease in grain size following a famous Hall-Petch relationship\textsuperscript{27,28} as the deformation is based on dislocation glide. Below a critical grain size, however, grain boundary sliding is favored, instead of dislocation glide, resulting in an increase in hardness with an increase in grain size.\textsuperscript{26} We have found that both the hardness and the grain size increase with the nickel content of our alloys. We believe that increase in hardness is not primarily a function of composition as the point defects decrease with increase in nickel content towards stoichiometry. Accordingly, the increase in hardness is attributable to the increase in grain size owing to the grain boundary sliding as a primary mode of deformation. As the intermetallic compounds resist the dislocation movement because of large burger’s vector,\textsuperscript{29} grain boundary sliding may be favored, allowing a relatively large critical grain size at which the primary deformation modes are switched. This seems to be a reason that grain boundary sliding has appeared as a primary mode of deformation at grain sizes of 20–50 nm.

4.2 Phases formed

Several kinds of baths are used for electrodeposition of Zn-Ni alloys. The chloride baths are preferred for improved micro-throwing power.\textsuperscript{4} Our baths contained only chlorides of nickel and zinc along with boric acid. The commonly used additive, i.e., NH\textsubscript{4}Cl, was not used. The deposition temperature was 45 °C. The XRD results reveal that only γ-phase forms in a wide range of composition of plating bath and the alloy composition varies from about 12 to 18% nickel. Only in case of zinc-rich bath 4 where the alloy contains ~5.8% nickel, a mixture of γ and η-phases is formed. Interestingly, δ-phase (Ni\textsubscript{3}Zn\textsubscript{22}) whose composition lies between η and γ-phases has not been observed in our electro-deposited alloys. A large number of authors have observed δ-phase coexisting with either η or γ-phase when the zinc content of the alloy exceeds the solubility limit of zinc in the γ-phase.\textsuperscript{30,31} However, the baths used by them contained either sulfates, or chlorides with NH\textsubscript{4}Cl causing higher pH as compared to our chloride baths. Therefore, lower pH of chloride baths containing boric acid without NH\textsubscript{4}Cl seems useful for preparing single-phase alloys comprising γ-phase. It is worth-mentioning that single-phase alloys are believed to provide better corrosion resistance in comparison with two-phase alloys.

4.3 Anomalous Codeposition

A clear anomalous codeposition has been observed in the simple chloride baths used in this work for alloy deposition. Even a minute cathodic current associated with the deposition of pure nickel has not been observed during cathodic scan before the codeposition of both the alloy constituents commenced (Fig. 2–4). The anomalous codeposition is attributed to the adsorbed layer of Zn(OH)\textsubscript{2} at the surface that is reduced to metallic form to allow the deposition of zinc.\textsuperscript{32} In this way, the Zn(OH)\textsubscript{2} acts as an intermediate product.\textsuperscript{33} The alloy composition has been found to be affected by plating bath composition as far as the thickness is small and zinc content of the alloy is high. However, the nickel content of the alloy keeps increasing with thickness until it reaches about 18% and further increase ceases to occur suggesting a limiting composition that can be formed by DC plating in the chloride bath. The adsorbed Zn(OH)\textsubscript{2} layer is expected to contain nickel cations that are also reduced along with zinc cations to form an alloy layer.\textsuperscript{33} Higher concentration of nickel cations in the adsorbed hydroxide layer due to an increased concentration of nickel chloride (or a decrease in the concentration of zinc chloride) in the plating bath may lead to higher nickel content in the alloy layer. This seems to be a reason for obtaining higher nickel content in the alloy when deposition is carried out in a nickel-rich bath. This is particularly true for short time deposition. We have found an upper limit of nickel content in the alloy to be about 18%. We consider that the adsorbed Zn(OH)\textsubscript{2} layer does not allow its nickel content to exceed a certain limit. Hence its nickel content, apparently, saturates with deposition time (due to enrichment of nickel in the underlying alloy layer) or increased concentration of nickel in the solution. As a result, the nickel content of the alloy formed from nickel-saturated Zn(OH)\textsubscript{2} remains at an upper limit of about 18%.

4.4 Effect of bath composition

In spite of the fact that alloy composition does not change significantly with the composition of plating bath, particularly, when the nickel content reaches its maximum value of about 18%, the morphology of the deposits and crystallographic orientation (texture) strongly depend on the composition of plating bath. The texture is also found to depend on deposition time. It appears that the adsorbed Zn(OH)\textsubscript{2}, which acts as a precursor for electrodeposits, changes its physical form (possibly the crystallographic arrangement of adsorbed ions) with deposition time and composition of plating bath, thus, altering the texture of the electro-deposited alloy.
4.5 Dissolution behavior

The dissolution of the alloys has been studied by cyclic voltammetry and chronopotentiometry. The main anodic peaks in voltammograms and the anodic plateaus in chronopotentiograms correspond to the dissolution of \( \gamma \)-, \( \gamma' \)- and \( \alpha \)- phases in the order of increasing potential. The dissolution also involves dezincification. Furthermore, the formation of \( \alpha \)-phase in our solution at 45 °C has only been possible by dezincification of \( \gamma \)-phase.

It may be noticed that the anodic peak C (Fig. 2) has been assigned to the dissolution of porous nickel by other authors.\(^9\) The lower dissolution potential than electrodeposited (pure) nickel was believed to be due to porous nature. It has been experimentally confirmed in this work that this peak C (and plateau III) involves the dissolution and dezincification of \( \alpha \) phase with high zinc content. Its low (more active) dissolution potential with respect to electrodeposited (pure) \( \alpha \)-phase is primarily attributable to its high zinc content.

5. Summary

1. Zn-Ni alloys with a composition of 5–18% Ni have been formed by DC plating in additive free chloride bath. The estimated grain size has been found to fall between 20–50 nm. The constitutional point defects concentrating at grain boundaries seem responsible for increase in grain boundary area and, hence, a decrease in grain size.

2. The micro hardness ranged between 150 to 450 KHN. It increased with grain size showing a break down of Hall Petch relationship at grain sizes below 50 nm. This behavior has been attributed to grain boundary sliding as a predominant mode of deformation.

3. The composition of thin deposits obtained during voltammetry and short time deposition depended on the composition of plating bath. The composition of thicker deposits showed less pronounced dependence on bath composition, particularly when nickel content reached a limit of about 18%. Increased nickel content in the plating bath resulted in the formation of smoother deposits with relatively enhanced (600) reflections in XRD patterns, in comparison with zinc-rich baths.

4. The results can be explained on the basis of formation of an adsorbed Zn(OH)\(_2\) layer responsible for codeposition. The composition of the electro-deposited alloy follows the changes in the composition of Zn(OH)\(_2\) layer, i.e. its nickel content, which in turn, is controlled by the composition of plating bath and the underlying alloy layer whose composition changes with deposition time. When the nickel content of Zn(OH)\(_2\) saturates at a certain level, the nickel content of the electro-deposited alloy also saturates to about 18%, irrespective of the plating bath composition.

5. Dissolution of the alloys involved pronounced dezincification. It has been revealed that \( \alpha \)-phase forms as a result of dezincification of \( \gamma \)-phase and that the last dissolution peak in voltammograms does not corre-