Electrochemical characterisation of the porosity and corrosion resistance of electrochemically deposited metal coatings

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1. Introduction

Through-porosity is an important parameter that should be assessed to estimate the life expectancy of coated components and their quality. Porosity in the coating is strongly related to the corrosion resistance of the substrate [1]. For example, a galvanic cell can be formed in an electroless nickel (EN) coating on aluminium or steel immersed in an acid. This galvanic cell promotes substrate dissolution and hydrogen evolution (or oxygen reduction) on the nickel coating. The noble nickel coating can also be dissolved in the acidic environment increasing the pore size and number, accelerating localised corrosion of the (base) substrate.

In the case of electroless coatings, the corrosion resistance depends on the phosphorous content [2]; for example, high corrosion resistance in acidic environments have been achieved at >10% phosphorous content while low (<4%) phosphorous content coatings tend to perform well in alkaline electrolytes [3] and at elevated temperatures [4].

For comparable pre-treatment schedules, the porosity of EN coatings has been shown to be much lower than that in nickel electrodeposits from a Watts bath [5]. The porosity of EN coatings is linked to a number of factors that decrease the corrosion resistance of the coated substrate, such as: aged baths [6], inadequate substrate pre-treatment [7,8], surface roughness [9–11] and coating microstructure [12]. Porosity can be affected by heat treatment due to the shrinking of the coating when its structure changes from amorphous to crystalline during the curing process, exposing part of the substrate [13].

1.1. Porosity tests

There are numerous chemical and electrochemical porosity tests in the literature. Chemical porosity tests include neutral and acetic acid salt spray, sulfur dioxide exposure, the use of a ferroxyl indicator and corrodkote corrosion tests [10,14]. Such tests are designed to attack the substrate revealing the corrosion occurring through the pores. However, corrosion products could block the pores and give false readings while very fine pores are hard to penetrate by the electrolyte and may be missed. These tests often take a long time and visual results can be difficult to quantify and reproduce [15]. In contrast, electrochemical technologies can be rapid and more reliable. A classical example is the electrographie test where a current is passed between the test specimen (anode) and an inert cathode. An absorbent paper soaked in an electrolyte containing a colorimetric indicator is sandwiched between the two electrodes. The current reveals coloured areas on the paper due to the reaction between the indicator in the electrolyte and the dissolved metal ions from the test specimen showing the location of the pores. A disadvantage is that very small pores can be missed with this technique; it is also important to chose appropriate electrolyte (and indicator) composition and the duration of the test. Other electrochemical tests, such as Tafel extrapolation, cyclic voltammetry, linear polarization resistance...
and corrosion potential monitoring, are more quantitative and can be performed rapidly with standard electrochemical instrumentation [16–18]. Such techniques have been used to determine the percentage through-porosity in electroless nickel coatings on steel and aluminium substrates [19–22].

Electrochemical methods can be used to expose the underlying substrate in a carefully chosen electrolyte. An appropriate electrolyte will allow contact with the substrate through pores in the deposit and with the coating material. The two metals form a bimetallic corrosion cell. If the coating is noble respect to the substrate, the deposit should be pore-free to avoid a high cathode to anode area ratio, which can lead to intense, localised corrosion of the substrate. This is especially true for thin immersion deposits. Continuous electroless deposition of metal may slowly fill any pores left in previous coating layers [4].

In many cases, the general relationship between the porosity ($P$) and a uniformly deposited coating of thickness ($x$) takes the form [23]:

$$ P = A x^n $$

(1)

where $A$ and $n$ are empirical constants that depend on pre-treatment, plating time and process conditions [5]. Fig. 1 shows an example of the relationship between percentage porosity and deposit thickness. The porosity was determined by cyclic voltammetry carried out on nickel-coated steel samples. The charge under the anodic peaks when the potential was swept towards negative values was measured for different thickness of nickel coated samples. The potential range and the electrolyte selected are such that anodic dissolution of the substrate occurs but nickel dissolution is negligible or occurs outside the potential range, at more positive potentials. The charge under the anodic peak without nickel coating was considered to represent 100% porosity while the charge exhibited by the samples with nickel coating was linearly correlated to the percentage of porosity [22]. The log–log plot of percentage porosity vs. nickel deposit thickness corresponds to the logarithmic form of Eq. (1):

$$ \log P = \log A + n \log x $$

(2)

Such a plot shows reasonable linearity over a thickness range between 0.2 and 6 μm with a correlation coefficient, $r^2$ of 0.997. If $x$ has units of μm, the values of the constants $A$ and $n$ are 12 and 1.35, respectively. The size, geometry and chemical homogeneity of the pores are not taken into account in this equation but it provides a reasonable approximation of the porosity over a wide range of coating thickness, for a uniform thickness specific coating substrate-test protocol.

2. Electrochemical techniques

In the case of a noble metal coating, the simplified electrode reactions during corrosion of an iron substrate in acid are typically:

Anodic reaction : $\text{Fe}^{2+} \rightarrow \text{Fe}^{3+}$ $E^0 = -0.447$ vs. SHE

(3)

Cathodic reaction : $2\text{H}^+ + 2\text{e}^- \rightarrow H_2$ $E^0 = 0.000$ vs. SHE

(4)

or : $1/2\text{O}_2 + 2\text{H}^+ + 2\text{e}^- \rightarrow \text{H}_2\text{O}$ $E^0 = 1.229$ vs. SHE

(5)

Corrosion cell reaction : $\text{Fe} + 2\text{H}^+ \rightarrow \text{Fe}^{2+} + \text{H}_2$

(6)

These reactions can be used to estimate the percentage of porosity. The most common electrochemical techniques used for this purpose are summarized in Table 1 and are explained in the Section 2.1.

In this paper, the emphasis is on direct current (d.c.) techniques. The reader is referred elsewhere for studies of porosity using a.c. techniques such as electrochemical impedance spectroscopy [24–26]. EIS can provide a powerful tool for electrochemical measurements on metal coating integrity but presents the problem of interpretation of complex data. For example, Zhang et al. suggested an equivalent circuit to measure the morphological properties of plasma [24].

![Fig. 1. Log–log plot of percentage porosity vs. coating thickness for electroless nickel deposits on steel in 0.125 mol dm$^{-3}$ H$_2$SO$_4$ at 295 K. Data obtained by cyclic voltammetry. •) Nahlé et al. [22]; and ○) recent data obtained by Walsh et al. [unpublished work].](image-url)
sprayed thermal barriers coatings. The porosity of the coating was proportional to the ceramic capacitance while the pore shape was estimated by the pore resistance. Ahm et al. [25] evaluated the porosity of WC–Cr–Al, N coatings on high speed steel using EIS. Creus et al. [26] determined the porosity of 3–10 μm Al, Ti, TiN and CrN protective coatings on steel after 1 h immersion in 3% NaCl from the faradic capacity of the coated steel. However, the porosity measurements critically depended on the immersion time which determines the charge transfer resistance and can only be accurate if rapid dissolution of the substrate and accumulation of the corrosion products in the pores can be eliminated [25]. Creus et al., [26] found that the porosity obtained from d.c. techniques such as polarization resistance, mixed potential theory and polarization curves and EIS were similar. They also suggested that sophisticated methods such as EIS could be avoided, in some cases, simple (direct current) polarisation resistance methods can give precise porosity values.

2.3. Cyclic voltammetry

The charge under the curve of the anodic peaks can be related to the percentage of substrate exposed to the electrolyte. An example of this method was briefly considered in the introduction for electroless nickel coating on steel. The samples were prepared by immersing the steel strips in an electroless nickel bath at different times. Samples immersed for 2.14 and 30 min developed Ni coatings of approximately 0.4, 2.8 and 6 μm thickness, respectively. Voltammograms of each coated sample were obtained by sweeping the potential towards negative values at a linear sweep rate of 50 mV s⁻¹ between −0.5 and +0.5 V vs. SCE in 0.1 mol dm⁻³ H₂SO₄ and in 0.1 mol dm⁻³ KFe(CN)₆ electrolyte at 298 K. The curves showed an anodic peak whose charge depended on the nickel coating thickness and the charge when no nickel was deposited was considered as 100% porous. The sample immersed in the nickel electroless bath for 30 min covered the whole steel surface and was considered 0% porous as the anodic peak showed none or little charge. Under these considerations the porosity of the 0.4 and 2.8 μm Ni thickness coated samples was 56% and 97%, respectively [22]. This technique can be easily carried out with simple electrochemical equipment in approximately 10 min.

2.4. Current-time transients

Controlling the anodic potential in the points where the slope of the anodic polarization curve, η vs. j, changes, can yield information of the reactions taking place on the surface. The levels of dissolved metal coating (Ni) and the underlying metal (Fe) will determine the predominant anodic dissolution process and can be directly related to the porosity of the coating deposit [33, 34].

2.5. Corrosion potential measurements

The open-circuit corrosion potential of porous deposits on a metal surface varies between the potential of the coating and the substrate. These measurements can therefore be used to assess whether the substrate is fully encased (pore-free deposit) and thereby resistant to corrosion in a given electrolyte or not. For example, the corrosion potentials E⁻ cor of aluminium (100% porous) and a 24 μm thick electroless nickel coating on aluminium (assumed to be pore free) are −0.86 V and −0.42 V vs. SCE, respectively. Similarly, the corrosion potentials of mild steel and a 24 μm (100% pore-free) nickel coating mild steel in 0.125 mol dm⁻³ H₂SO₄ are −0.25 and −0.47 V vs. SCE, respectively. Values of the corrosion potentials between these limits can be correlated to the degree of porosity. For thin <6 μm, coatings, other effects such as oxide formation and the stronger activity of the substrate should be taken into account [20].

3. Experimental details

3.1. Composition and plating conditions for aluminium substrates

Aluminium alloy plates (3105 H24) of 20×50×1 mm size were used for the corrosion studies. The composition of the alloy was 6.08% Si, 0.70% Fe, 0.30% Cu, 0.30–0.80% Mn, 0.20–0.80% Mg, 0.20% Cr, 0.40% Zn and 96.4% wt. Al. Each sample was pretreated by an alkaline degrease (pH 7.2–8) at 323–343 K [35], rinsed with deionised water and etched in acid, desmutting solution, typically 50% HNO₃ with added H₂SO₄ and fluoride ions to remove metallic oxides [36]. The aluminium substrates were then immersed twice in a zinicate solution, rinsed in deionised water and acid etched in between. The full pre-treatment process is described in detail elsewhere [37]. The pre-treated aluminium substrates were plated with a commercial electroless nickel plating process (Fidelity 5010) at various times. A temperature of 363–365 K, the plating rate was approximately 4–6 μm h⁻¹ and controlled deposition times
were chosen to produce nickel coated samples of 0.5, 1, 3, 6, 12 and 24 µm thickness.

3.2. Composition and plating conditions for mild steel substrates

Panels of mild steel (Pyrene steel — 040A10 BS 970:1983) of 75×25×1 mm size were used. Prior to the electroless plating, the panels were degreased in a 10% concentration of a commercial alkaline soak cleaner (Fidelity 3152) for 10 min at 323–333 K, followed by 1 min rinse in deionised water and 2 min immersion in 50% HCl solution. The process was repeated until complete removal of oxide surfaces was achieved. The pre-treatment of the mild steel is fully described elsewhere [20]. The panels were then coated in a high-phosphorus, electroless nickel plating solution to achieve 1, 3, 6, 12, 18 and 24 µm coating thickness.

3.3. Composition and plating conditions for ABS substrates

Panels of ABS plastic, size 75×25×1 mm, were used. The panels were degreased with a commercial alkaline soak cleaner Durarep, (Duratech Industries Ltd.) rinsed with deionised water, acid etched (providing a good key for subsequent metallic coating) and rinsed again. The panels were then immersed in a stannous chloride/HCl solution followed by activation of the polymer surface with PdCl2 and a final double rinse with deionised water before applying a 30 µm electroless copper deposit. This was followed by immersing the panels in a gold containing solution for controlled periods of time to achieve gold deposit of nominally 0, 75, 100 and 190×nm thickness. The ABS samples were rinsed in deionised water, washed with ethanol, dried with a hot air and stored in a dessicator. The ABS pre-treatment is fully described elsewhere [1].

3.4. Choice of electrolyte

The nature of the electrolyte is critical to keep the corrosion of the deposit as low as possible. It is assumed that the current flowing from the electrode is due to the corrosion of the substrate together with dissolved ions passing through the existing pores of the coating. Electrochemical measurements of the three samples described above were carried out at 295 K as follows: (i) aluminium substrate samples in 0.85 mol dm$^{-3}$ (5%) sodium chloride electrolyte, (ii) mild steel coupons in 0.125 mol dm$^{-3}$ sulfuric acid and (iii) gold copper-coated ABS plastic samples in a 0.1 mol dm$^{-3}$ NH4BF4 electrolyte.

The chemicals used in the experiments were analytical grade and the solutions were prepared with deionised water. All the experiments were carried out in triplicate using a saturated calomel reference electrode (SCE) and a 4 cm$^2$ Pt mesh counter electrode.

3.5. Tafel extrapolation, anodic polarisation and current–time transients

A PAR EG & G potentiostat (model 273A) was used to apply a linear potential sweep rate of 1 mV s$^{-1}$ for all the electrochemical tests. The experimental arrangement including a two-compartment electrochemical glass cell (300 cm$^3$) containing 3-electrodes as illustrated in Fig. 2. This cell was used for the majority of the experiments. The linear polarization resistance (LPR) and corrosion potential measurements were carried out in a 250 cm$^3$ beaker. The
plated samples were polarised from $-0.25$ to $+0.25$ V vs. SCE (Tafel measurements) and $-0.25$ to $+0.16$ V vs. SCE (anodic polarisation measurements). Current vs. time transients were obtained at various constant potentials, previously identified on the anodic polarisation curves, for 60 min.

3.6. Linear polarisation resistance and corrosion potential

Identical coupons of the samples coated with electroless nickel were covered with an insulating masking tape leaving 1 cm² area exposed and were attached to a twin electrode LPR meter (SSL Ltd.). The meter was internally calibrated to give an average corrosion reading in terms of the estimated penetration rate in mm yr⁻¹. In parallel with the LPR readings, corrosion potential ($E_{corr}$) measurements ($±0.002$ V) vs. SCE were taken over 96 h, using a high impedance digital voltmeter (Iso-Tech IDM 201).

4. Results and discussion

4.1. Porosity of electroless nickel coatings on an aluminium alloy substrate

4.1.1. Anodic polarisation

Fig. 3 shows the logarithm of the current density vs. potential for the aluminium alloy samples. The curves represent the bare alloy and samples covered with 0.5, 1.0, 3.0 and 24 μm electroless nickel (EN) coating in 0.85 mol dm⁻³ (5%) NaCl solution. All curves have a horizontal region of steady current as the potential increases indicating the formation of a passive film on the nickel deposits. The passive film for the 24 μm thick coating, which was assumed to be pore free, appears to
break down at about +0.25 V vs. SCE before the thinner coatings. Despite
this increase of current probably due to the gradual opening of pores on
the coating, the current is still 10⁻⁴ A cm⁻² at this potential which shows
the substantial protection provided by the EN coating. The bare
aluminium alloy at this potential corrodes at 0.1 A cm⁻². The current of
thinner coatings lies between the 10⁻⁴ and 10⁻³ A cm⁻² at this potential
but they break down at approximately +0.75 V vs. SCE generating a
dissolution current of approximately 0.01 A cm⁻², which is still an order of
magnitude lower than the current generated at the bare aluminium alloy.

4.1.2. Dissolution potential measurements

Fig. 4 shows the potential (E), of the EN-coated aluminium alloy
samples vs. the deposit thickness at selected current densities. The
samples were immersed in 0.85 mol dm⁻³ (5%) NaCl solution. The
open-circuit potential (OCP) of the samples after 15 min and 24 h of
the deposition, increased from ≈−0.9 V for the bare aluminium
substrate (see Fig. 3) to −0.3 and −0.4 V vs. SCE respectively when the
nickel coating was ≥ 5 μm thick. This potential difference is due to the

Table 2
Tafel analyses for mild steel and various electroless nickel deposits in 0.1 mol dm⁻³
H₂SO₄ at a temperature of 295 K

<table>
<thead>
<tr>
<th>Deposit thickness, x/μm (Letter in Fig. 7 a)</th>
<th>Corrosion potential, E_cor vs. SCE/V</th>
<th>Corrosion current density, j_cor/A cm⁻²</th>
<th>Mean corrosion rate/mm yr⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 (uncoated mild steel)</td>
<td>−0.540</td>
<td>61</td>
<td>0.75</td>
</tr>
<tr>
<td>1 (a)</td>
<td>−0.450</td>
<td>30</td>
<td>0.37</td>
</tr>
<tr>
<td>3 (b)</td>
<td>−0.425</td>
<td>30</td>
<td>0.37</td>
</tr>
<tr>
<td>6 (c)</td>
<td>−0.420</td>
<td>30</td>
<td>0.37</td>
</tr>
<tr>
<td>12 (d)</td>
<td>−0.390</td>
<td>9</td>
<td>0.11</td>
</tr>
<tr>
<td>18 (e)</td>
<td>−0.230</td>
<td>6</td>
<td>0.089</td>
</tr>
<tr>
<td>24 (f)</td>
<td>−0.165</td>
<td>7</td>
<td>0.083</td>
</tr>
</tbody>
</table>

Fig. 7. a) Cathodic Tafel extrapolation plots of mild steel samples coated with various
thicknesses of electroless nickel deposits immersed in 0.125 mol dm⁻³ H₂SO₄ at 295 K.
Deposit thickness: a) 1 μm, b) 3 μm, c) 6 μm, d) 12 μm, e) 18 μm and f) 24 μm. b) Corrosion
potential and corrosion current vs. deposition time obtained from a Tafel plot (Figure a):
 a) E_cor, b) j_cor.

Fig. 8. a) Log current density vs. potential for a 1 μm thickness electroless nickel-12%
phosphorus deposit on mild steel. b) Current vs. time at the potential indicated in the
points of figure a: (1) −0.17 V, (2) +0.2 V and (3) +0.6 V vs. SCE. The electrolyte
(0.125 mol dm⁻³ H₂SO₄ at 295 K), was analysed for Ni²⁺ and Fe²⁺ ions for all experiments
at the points A, B, C and D.

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fact that the open-circuit potential immediately after the deposition was taken from the anodic polarisation curve and the equilibrium, which takes approximately 15 min, had not been reached. The potential values, extrapolated from the anodic polarisation plots of Fig. 4 at 0.05 (0.1 mA) and 0.5 mA cm$^{-2}$ (1 mA) current densities are approximately +0.580 and +0.350 V vs. SCE, respectively when the coating thickness was >5 μm.

Longer term corrosion tests over 96 h are presented in Fig. 5. The corrosion potentials for the 3, 6 and 24 μm coating thickness were higher than for the bare aluminium substrate and the 5 μm coating and remained fairly constant for approximately 48 h. After this time, the samples coated with 3 and 6 μm EN developed pores resulting in a potential decreasing. The 12 μm (not shown) and 24 μm thickness EN nickel coatings show no evidence of corrosion over the time period considered.

4.1.3. Linear polarisation resistance and corrosion current monitoring

The linear polarisation measurements of the EN coated Al samples immersed in 0.85 mol dm$^{-3}$ (5%) NaCl solution can be seen in Fig. 6. The curve shows that the corrosion rate decreases drastically to +0.52 mm yr$^{-1}$ for the bare aluminium, to +0.01 mm yr$^{-1}$ for nickel coatings of more than 3 μm thickness. The results are comparable with a Tafel analysis which showed that the corrosion rate decrease from 0.04 to 0.01 mm yr$^{-1}$. The slightly higher initial results are attributed to the 24 h stabilisation required for the 2-probe LPR method in comparison to a typical time of 15 min required for a Tafel analysis. The 3 μm EN thickness showed an averaged penetration rate of less than 0.1 mm yr$^{-1}$ by both methods and the corrosion rates indicate substantial decrease in the porosity of the nickel coating. These corrosion rates compare values reported by Zeller of less than 1 mm yr$^{-1}$.[38].

4.2. Porosity measurements on electroless nickel deposits on a mild steel substrate

4.2.1. Tafel extrapolation (log current density vs. potential) diagrams

Pyrene steel and electroless nickel coated pyrene steel samples were examined. The values of the corrosion current density obtained from Tafel plots were to calculate the corrosion resistance of different nickel coating thickness [39]. Some typical results are shown in Fig. 7 a) and in Table 2.

Fig. 7 a) shows that the 24 μm thickness EN deposit on mild steel gave rise to cathodic Tafel slopes between −0.19 to −0.20 V. In comparison, pyrene steel and the other EN deposits show lower Tafel slopes of approximately 0.10 to 0.14 V. This variation is due to the incomplete reduction of residual oxide films during cathodic polarisation. These films are not efficient cathodes as the oxide free surfaces (i.e., a lower $j_0$) and indicate that hydrogen reduction is not the only cathodic reaction; reduction of surface oxides might also occur. For thicker deposits, the value of $E_{cor}$ is just above the hydrogen evolution line ($\text{H}^+$/H$_2$) as a $E$-pH diagram. This will result in insufficient hydrogen being generated when the electrode was polarised to approximately 0.20 V below the value of $E_{cor}$ for the removal of the oxide film. The levels of current for hydrogen reduction at these potentials will be in the order of 0.01 μA cm$^{-2}$. Table 2 shows that the value of $j_{cor}$ for mild steel was 61 A cm$^{-2}$, while for a 24 μm electroless nickel deposit was 7 A cm$^{-2}$. Thin EN coatings tend to be porous and give intermediate values of $j_{cor}$ and $E_{cor}$. The corrosion resistance of these coatings depends on the exposed area of steel at the base of the pores.

Fig. 7 b) shows the effect of the EN deposition time (hence deposit thickness) on both, corrosion potential $E_{cor}$, and corrosion current density $j_{cor}$. The corrosion potential changed from −0.45 to −0.20 V vs. SCE as the deposition time increased from 0 to 120 min while the corrosion current density decreased from approximately 0.30 to 0.005 mA cm$^{-2}$ over the same period of time.

4.2.2. Anodic polarisation and current vs. time transients

4.2.2.1. 1 μm electroless nickel coating. Fig. 8 a) shows the log current density vs. potential curve for 1 μm thickness electroless nickel coating on mild steel substrate. The numbers on the Figure indicate the changes in the slope of the anodic polarisation plot and the points at which a constant electrolysis was carried out. The currents at these points were monitored vs. time and the results are plotted in Fig. 8 b), where the curves are identified with the numbers given to the points on Fig. 8 a). The electrolyte was analysed for nickel and iron content at the end of the electrolysis to determine the predominant anodic dissolution process, i.e., nickel or iron. The results are summarised in Table 3.

When the potential was fixed at −0.17 V, (point 1 on Fig. 8 a) the current density was +3 mA cm$^{-2}$ as seen on curve (1) Fig. 8 b). At this potential there should be zero nickel dissolution whereas the iron substrate can be anodically dissolved through the pores of the coating. The data in Table 3 show a concentration of 16.5 ppm iron dissolved while no nickel was detected at the end of the electrolysis. Point 2 on Fig. 8 a) corresponds to a metastable region for iron passivation. The potential at this point corresponds approximately to +0.20 V vs. SCE and the curve (2) on Fig. 8 b) shows the current vs. time plot at this potential.

The repeated fluctuation of the current readings over the period of the test clearly demonstrates the unstable nature of the film formed on the exposed iron surface at the base of the pores. This current is due to the anodic dissolution of iron as no nickel was detected in the electrolyte. At these potentials, the electroless nickel surface was still covered with a

| Table 3 Concentration of dissolved nickel and iron at significant changes of the slope in the anodic polarisation curve (numbered 1–9 on Figs. 8, 9 and 10) |
|-----------------|-----------------|-----------------|-----------------|
| Deposition time (thickness of EN deposit) | Position on figures and potential (point) E vs. SCE | Concentration of metal ion in 0.125 mol dm$^{-3}$ H$_2$SO$_4$ electrolyte/ppm | Measurements for the area under the current-time curves |
|                 |                 | Calculated Ni | Experimental Fe | Ni | Charge, q/A s | Current density, j/mA cm$^{-2}$ | Stripping time/min |
| 5 min (1 μm) Fig. 8a) and 8b) | (1) −0.17 V | – | 16.5 | 0.0 | 10.6 | 2.96 |
| 15 min (3 μm) Fig. 9a) and 9b) | (2) +0.20 V | – | 44.9 | 0.0 | 24.0 | 6.60 |
| 120 min (24 μm) Fig. 10a) and 10b) | (3) +0.60 V | 4.5 | 5.4 | 3.9 | 8.3 | 2.46 |
|                             | (4) −0.17 V | – | 8.3 | 0.0 | 5.1 | 1.41 |
|                             | (5) +0.20 V | – | 8.7 | 0.0 | 5.2 | 1.46 |
|                             | (6) +0.60 V | 9.9 | 0.5 | 9.7 | 11.6 | 3.21 |
|                             | (7) 0.00 V | – | 0.0 | 0.0 | 0.14 | 0.04 |
|                             | (8) +0.40 V | 5.4 | 0.0 | 4.5 | 5.3 | 1.48 |
|                             | (9) +0.85 V | 36.6 | 0.65 | 30 | 36.2 | 10.0 |

Electrical charge, current density and theoretical times for stripping are taken from the current vs. time plots of various electroless nickel coating thickness on a mild steel substrate in 0.1 mol dm$^{-3}$ H$_2$SO$_4$ at 295 K.
phosphorus type film. Table 3 shows that the iron content in the acid electrolyte was 44.9 ppm, which is appreciably higher than the concentration observed at $-0.17$ V vs. SCE, 16.5 ppm. The current increase and decrease around the point 3 on Fig. 8a) indicates that the protective film on the surface of the EN broke down at $\approx +0.6$ V vs. SCE. Curve (3) in Fig. 8b) shows the current vs. time at this potential. The current increase in the region (A)–(B) of curve (3) can be due to two factors: a) breakdown of the phosphorus type film in the electroless nickel surface i.e., the current rise is the result of nickel dissolution through the phosphorus film or b) dissolution of iron at the base of the pores until the value of $i_{\text{crit}}$ has been exceeded and the iron began to passivate. Following these processes the electroless nickel deposit dissolved in the regions (B) to (C) with the current falling as the area of electroless nickel diminished.

Additional current vs. time transients (not shown) were recorded at points (C) and (D) on Fig. 8b). The analysis of the electrolyte at point (C) showed no dissolved iron, which discounts the hypothesis b) above. Fluctuation of the current in the region (A) to (C) is due solely to the dissolution of nickel from the deposit. The current density value at point (D) on Fig. 8b) corresponds to the passivation current $i_{\text{pass}}$ for iron in the acid electrolyte at this potential. Analysis of the solution at the end of the electrolysis on point (D) indicates that 3.9 ppm for nickel and 5.4 ppm for iron are dissolved in the electrolyte (see Table 3). This result shows that, for a 1 μm thick coating, the Ni breaks down after approximately 5 min following an applied potential of +0.6 V vs. SCE. Dissolution and passivation of the iron substrate surface follow.
4.2.2.2. 3 μm electroless nickel coating. A similar analysis of the 3 μm thick EN deposit when the potential was held at +0.17 V vs. SCE (point 4 on Fig. 9a) shows that the corrosion current and the iron content of the electrolyte at the end of the electrolysis were 1.41 mA cm$^{-2}$ and 8.3 ppm, respectively (curve (4) on Fig. 9b). These values are lower than those found for the 1 μm nickel deposit at the same potential (see Table 3). When the potential was held at +0.2 V vs. SCE (point 5 on Fig. 9a) similar current vs. time transient with rapid fluctuations of current (similar to the case of the 1 μm thick deposit) were observed (curve (5) on Fig. 9b). The concentration of dissolved iron was 8.7 ppm, which was significantly lower than for the 1 μm deposit, 44.9 ppm (see Table 3). This may be due to the tendency of the iron surface to be covered with a passive film at longer periods of time, which is reflected in the low values of current at both −0.17 and +0.20 V vs. SCE (Table 3). The concentration of dissolved iron at these potential was 8.3 ppm and 8.7 ppm respectively, reinforcing the suggestion of a passive film. When the potential was set at +0.60 V vs. SCE (point 6 on Fig. 9a) the 3 μm thick nickel coating showed a fluctuation of the current, indicating breakdown of the coating (curve (6) in Fig. 9b). The concentration of Ni in the electrolyte at the end of the electrolysis was 9.7 ppm in comparison to the value of 3.9 ppm found for the 1 μm thick deposit with only 0.5 ppm of Fe$^{2+}$ at the same potential. These results show that the 3 μm thick nickel coating protected the iron substrate more effectively than the 1 μm one.

4.2.2.3. 24 μm electroless nickel coating on mild steel. For a 24 μm thick EN deposit, the potential was held at 0 V vs. SCE, corresponding to point 7 in Fig. 10a). At this potential the phosphorus type film suppresses nickel dissolution and no nickel ions were detected in the electrolyte as it is shown in Table 3. As expected, the current density at this potential is very low; <0.04 mA cm$^{-2}$ (curve (7) Fig. 10b). Further increase in the potential to +0.4 V vs. SCE (point 8 Fig. 10a) showed no Fe$^{2+}$ ions at the end of the electrolysis and 4.5 ppm of Ni ions in the electrolyte (curve (8) Fig. 10b). At this potential, the phosphorus film covering the surface of the 24 m deposit begins to break down. The coating is expected to be essentially pore-free and this is reflected in the results of the current vs. time plots at the potentials indicated as points 7 and 8 in Fig. 10a). At these potentials the concentration of iron was below the detection limit of the analytical method (0.005 to 0.01 ppm). Increasing the potential to

Table 4

<table>
<thead>
<tr>
<th>Material</th>
<th>Open-circuit potential, $E_{oc}$ or corrosion potential, $E_{cor}$ vs. SCE/V</th>
<th>Integrated area under anodic polarisation curve (charge per unit area)/mC cm$^{-2}$ (Fig. 11)</th>
<th>Integrated area of $j$ vs. $t$ transients/mC cm$^{-2}$ (Fig. 12)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gold foil</td>
<td>0.120</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>190 nm Au deposit</td>
<td>0.120</td>
<td>6.4</td>
<td>0.02</td>
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<tr>
<td>100 nm Au deposit</td>
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<td>6.9</td>
<td>0.51</td>
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<tr>
<td>75 nm Au deposit</td>
<td>0.148</td>
<td>10.1</td>
<td>0.93</td>
</tr>
<tr>
<td>30 μm electroless copper deposit</td>
<td>0.250</td>
<td>52.2</td>
<td>1.20</td>
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</table>

Fig. 11. Anodic polarisation curves for bare electroless copper and various submicron thick immersion gold deposits on copper-coated ABS polymer in 0.1 mol dm$^{-3}$ NH$_4$BF$_4$ at 295 K. a) Bare electroless copper deposit on ABS, b) 75 nm, c) 100 nm and d) 190 nm are the nominal thickness of Au coatings.

Fig. 12. Current density vs. time curves for an electroless copper-covered ABS substrate and various immersion gold deposits on the copper in 0.1 mol dm$^{-3}$ NH$_4$BF$_4$ electrolyte, at 295 K. ● Bare electroless Cu on ABS of 30 μm thickness. Thickness of Au coatings: ○ 75 nm, ■ 100 nm, and □ 190 nm.

Fig. 13. Immersion gold deposits on an electroless copper of 30 μm thickness deposited on an ABS plastic substrate. Assessment of the extent of deposit porosity at 295 K.
+0.65 V vs. SCE causes a dramatic increase in current density to 10 mA cm$^{-2}$ (curve (9) Fig. 10 b) which indicates a rapid dissolution of the deposit. Point 9 on Fig. 10 a confirms that the dissolution of the coating is rapid at this potential as the $i_{\text{crit}}$ value for EN dissolution is approached. This is reflected in the high concentration of nickel ions in the electrolyte, 30 ppm (see Table 3). The substrate at this potential is still protected by the Ni coating as is reflected by the low concentration of Fe$^{2+}$ ions.

4.2.2.4. Predicted and experimental concentrations of Ni$^{2+}$ in the electrolyte. To verify that the high currents observed at +0.6 V and +0.65 V vs. SCE are due to dissolution of the electroleless nickel coating, the predicted amount of nickel for each deposit thickness that would dissolve into the electrolyte as Ni$^{2+}$ was calculated from Faraday's law (assuming 100% efficiency). It was also possible to determine the time taken to completely dissolve the deposit for the current values measured (see Table 3). The area under the current vs. time curve is equivalent to the electrical charge passed per mole, i.e., A s mol$^{-1}$. For example, a 1 µm deposit at a fixed potential of +0.6 V, gave a value of 8.3 coulombs. This is equivalent to a current density of 2.5 mA cm$^{-2}$ (Table 3), which would result in the complete dissolution of the deposit from a 1 cm$^2$ area in 30.3 min. This should have resulted in a predicted concentration of 4.5 ppm of nickel in the 300 cm$^3$ electrolyte (Table 3). Experimentally, a value of 3.9 ppm was observed, which is in reasonable agreement with the predicted value and supports the argument that EN deposit corrodes at this potential. It was also possible to determine the time taken to completely dissolve the deposit for the current values measured (see Table 3). The area under the current vs. time curve is equivalent to the electrical charge passed per mole, i.e., A s mol$^{-1}$. For example, a 1 µm deposit at a fixed potential of +0.6 V, gave a value of 8.3 coulombs.

The electrochemical porosity tests highlighted in this paper included d.c. techniques which are easier to interpret and provide similar results as the EIS technique without the complications. Such techniques included: linear polarisation resistance, Tafel extrapolation, anodic polarisation and current vs. time transients. The test were carried out in specifically chosen electrolytes, as this choice is critical for porosity determination, i.e., the electrolyte must maximise the corrosion of the substrate with minimal destruction of the coating material.

The determination of porosity by electrochemical techniques can be an effective alternative to more conventional chemical tests. d.c. tests identified the presence of pores on the coatings of aluminium, mild steel and Cu-plated ABS polymer substrates, illustrating that these electrochemical techniques have a wide variety of applications, which are not specific to one coating/substrate combination.

### List of symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A$</td>
<td>Constant in Eq. (2)</td>
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<tr>
<td>$K$</td>
<td>Constant in Eq. (8)</td>
<td>V</td>
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<tr>
<td>$E^0$</td>
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<td>$E$</td>
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<td>Corrosion potential</td>
<td>V</td>
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<tr>
<td>$E_{meas}$</td>
<td>Measured potential</td>
<td>V</td>
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<tr>
<td>$E_{MP}$</td>
<td>Mixed potential</td>
<td>V</td>
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<tr>
<td>$j$</td>
<td>Current density</td>
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<tr>
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<td>Corrosion current density</td>
<td>A cm$^{-2}$</td>
</tr>
<tr>
<td>$j_{crit}$</td>
<td>Critical current density</td>
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<td>$j_0$</td>
<td>Exchange current density</td>
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<tr>
<td>$N$</td>
<td>Constant in Eq. (1) and (2)</td>
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<tr>
<td>$P$</td>
<td>Porosity of the coating</td>
<td>Dimensionless</td>
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<td>$q$</td>
<td>Electrical charge</td>
<td>C</td>
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<tr>
<td>$x$</td>
<td>Deposit thickness</td>
<td>nm and µm</td>
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<tr>
<td>$\beta_a$</td>
<td>Anodic Tafel slope</td>
<td>V decade$^{-1}$</td>
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<tr>
<td>$\beta_c$</td>
<td>Cathodic Tafel slope</td>
<td>V decade$^{-1}$</td>
</tr>
</tbody>
</table>
$\eta_c$ Cathodic polarization $V$

$\eta_a$ Anodic polarization $V$

References