ELECTROCHEMICAL IMPEDANCE SPECTROSCOPY ON THERMAL AGEING EVALUATION OF EPOXY COATING CONTAINING ZINC RICH PRIMER

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Abstract
This research concentrates on the thermal ageing of a full 3-coat system with sacrificial pigment (zinc rich) primer on mild steel where the temperature dependence test is conducted to explore the correlation between the coating resistances with the corrosion rate underneath the coating. Electrochemical impedance spectroscopy (EIS) is introduced over a range of temperature to extract activation energies for the rate of controlling processes in the corrosion reactions. It is expected that the zinc rich primer does not form a barrier coating for the metal substrate rather it will be the most active component of the substrate in the electrochemistry. Full 3-coat system with zinc rich primer show the extracted activation energy from coating resistance is significantly lower than the activation energy extracted from the charge transfer resistance. This suggested that the coating resistance from EIS cannot be controlling the corrosion reaction. The activation energies generated for the corrosion process here (78–97 kJmol⁻¹) are very much higher than those of ion transport through the coating (19–37 kJmol⁻¹) during early immersion. Further interesting findings come from the activation energy trends over time particularly for the corrosion process which shown that the value is decreasing where at the end of exposure, the activation energy values for coating and charge transfer resistance become quite similar. It is suggested that at this stage ion transport in the coating might be controlling the corrosion process unlike at the beginning; the activation energy is getting smaller due to coating degradation.

Index Terms: epoxy coating, electrochemical impedance spectroscopy, thermal effect, zinc rich primer

1. INTRODUCTION
Temperature changes impose a direct impact on the corrosive protective performance of coatings; especially in an environment where the value of temperature keeps changing. For example, a natural gas pipeline network often has continuous changes in temperature, with possible range as high as 135°C in the vicinity of a natural gas well to an ambient temperature at a pumping station.

Previous work [1, 2] has established that it could be possible to separate resistances for the coating and the coating-metal interface from EIS spectra. The plotted Arrhenius of logarithm resistance against reciprocal of temperature yielded larger activation energies for the corrosion process than for ion conduction in the coating, showing that the coating resistance could not be responsible for controlling the corrosion rate. Here further coatings, including a full 3-coat system with sacrificial pigment primer are studied. In this work, the electrochemical activity of the zinc rich primer (coupled to the steel substrate) plus barrier properties of those two topcoats were in concern. Thus, steel with zinc rich primer (ZRP) alone was tested first to identify characteristics of its electrochemical (sacrificial action) behavior.

2. EXPERIMENTAL PROCEDURE
A commercial zinc rich primer with 50µm thickness was tested. Coatings were applied to grit-blasted mild steel panels by air spraying. Further tests were conducted on a full 3-coat system consisting of a zinc rich pigmented 2-component epoxy primer (50µm), a low volatile organic content (VOC), high-solid, high-build epoxy barrier coat pigmented with micaceous iron oxide (180µm) and a 2-component acrylic polyurethane topcoat (60µm), prepare and applied similarly. The edges of all panels were protected with a thick high-performance epoxy coating. The electrochemical measurements were made using a three-electrode cell with vertical working electrode of area 40 cm². The reference electrode was a saturated calomel electrode (SCE) and the counter electrode a platiniised titanium electrode of area of 9 cm². The samples were exposed in hot 3% NaCl solution (800 cm³), made from analytical grade chemicals and deionised water, held at constant temperature in a water bath (or slowly
cooled in an insulated box). Tests were conducted in duplicate. Impedance measurements were taken at the free corrosion potential using a Gill AC computer-controlled potentiostat with a high-impedance paint buffer (ACM Instruments), using a 20 mV sine amplitude and a frequency range from 0.1 Hz to 30 kHz. Samples were cooled from 50°C down to ambient over (typically) 4 h. After the samples had been tested, the cell was replaced in the hot water bath and kept at 50°C until the next measurement.

3.1 Tests with Zinc Rich Primer (ZRP) Alone

3.1.1 Visual Inspection

Optical micrographs as seen in Figure 2 shows the results of galvanic activity on the surface of the ZRP coating surface before and after exposure up to 22 days in 3% NaCl solution. Figure 2b shows the presence of small ‘white’ particles, probably zinc oxide which starts to build up on the coating surface after 2 days of exposure. By day 19, the presence of blisters was noted and with time, these blisters grow, and unlike normal blisters, they are rough and not smooth (Figure 2c).

3.1.2 Preliminary Analysis for Coated Panel Exposed at 21°C

After an hour of exposure, Nyquist plot shows two semicircles (Figure 3), similar finding to the study conducted by Meroufel and Touzain [3]. This indicates that the coating is highly porous and permeable. The potentials recorded were -0.96 V_SCE and -0.98 V_SCE which suggests that zinc particles are already starting to corrode.

3.1.3 Open Circuit Potential Measurements

The duration of protection potential for steel (E_pp) is given as the time during which the potential remains lower than free corrosion potential of steel (-0.65V_SCE). However it is often taken in practical as being the time which the potential remains lower than -0.85V_SCE [9]. The ZRP coated panel potentials are monitored up to 22 days and presented in Figure 4.
Fig. 4: Evolution of corrosion potential measured at 21°C with time for ZRP coating (2 replicates) exposed in 3% NaCl solution for 22 days at 50°C

Due to the high permeability of ZRP coatings, immediately upon immersion in the electrolyte the potential is about -0.98V SCE, a value which is close to the potential of zinc in sea water. This indicates clearly that the zinc particles were actively consumed to provide initial galvanic protection to the steel. Notice that the potential passed the $E_{PP}$ value by 6 days. By 22 days, contact between zinc and steel may have been lost as the potential recorded is close to the corrosion potential of steel.

3.1.4 Temperature Dependence of Impedance

The effect of changing temperature on the EIS response of ZRP coating in 3% NaCl solution after 7 and 14 days exposure at 50°C are shown in Figure 5 and 6 respectively. It seems that the effect of changing temperature is barely noticeable on the impedance spectra after 1 and 4 days of immersion.

However by day 7 the effect of changing temperature becomes much clearer. Even though the effect of temperature on the size of the ‘semicircle’ is apparent in these figures, Nyquist plots are difficult to fit. Only the first semicircle could be fitted using “circle fit” from the ACM software. Logarithm of coating resistance is then plotted against reciprocal of temperature (Figure 7) and the activation energies measured are in the range of 32–34 kJmol⁻¹.

Fig. 5: Nyquist plots for ZRP coating at various temperatures after 7 days of exposure at 50°C

Fig. 6: Nyquist plots for ZRP coating at various temperatures after 14 days of exposure at 50°C

Fig. 7: Arrhenius plot for ZRP coating after (a) 7 days and (b) 14 days of exposure at 50°C
3.2 Tests on Full System

3.2.1 Open Circuit Potential Measurement

Figure 8 displays the open circuit potential (OCP) measured over time for coated panels (duplicate sets) exposed at 21 and 50°C. It is interesting to note that the two panels tested at 21°C reacted differently during early immersion.

During the first 21 days of immersion Panel 1 had an unstable potential which varied between -0.3 and -0.8 V_{SCE} before reaching a stationary value close to the corrosion potential of steel. In contrast Panel 2 shows potential values close to the zinc potential during early immersion but rapidly increased reaching the corrosion potential of steel by 21 days. Panels 3 and 4 tested at 50°C gave comparable potential values. Their values were fluctuating in the range of -0.45 to -0.65 V_{SCE} which is similar to the potential for a coated steel substrate. In this study, the number of days during which the coated panel exhibits more negative potential than E_{PP} (-0.85 V_{SCE}) in 3% NaCl solution is used for judgment of how long zinc rich primer coatings provide full cathodic protection to the steel substrate.

According to these results, it is evident that OCP of these panels never lies in cathodic protection region (below -0.85 V_{SCE}). The potential measured for these coatings will be however a compromise between the zinc-steel galvanic couple (anode and cathode) as suggested by Mayne [10]. Anode potential may give reading at -0.95 V_{SCE} and cathode potential may be a very much higher.

3.2.2 Effect of Signal Amplitude – A Test of the Circuit Model

ZRP (primer only) gave two semicircles (Figure 3) and here again two semicircles were observed (Figure 9) for the full system. The first semicircle was expected to be the top coating over the zinc rich epoxy primer and the second semicircle corresponded to the electrochemical activity of the zinc rich primer coupled to the steel substrate. To be certain a test is conducted to investigate the effect of changing the signal amplitude on the semicircles as a way of identifying the different features displayed in Nyquist plots. EIS spectra were taken at amplitudes ranging from 20 mV to 120 mV. It is clear from Figure 9 that changing the applied potential only changes the shape of the second semicircle.

3.2.3 Temperature Dependence of Coating Parameters

Figures 10 to 11 show the effect of changing temperature on the EIS response of full system coating after 21 and 35 days exposure at 50°C. Similar to epoxy-phenolic paint and full system with an inhibitive primer as studied previously [1, 2] the size of the semicircles decreases as temperature rises.

These semicircles were fitted to the equivalent circuit in Figure 1 and this procedure give the most accurate determination of coating resistance (R_{p}) at high frequency and charge transfer resistance (R_{ct}) at low frequency. Then the logarithm of R_{p} and logarithm of R_{ct} were plotted against reciprocal of temperature to determine activation energies for conduction in the coating and the corrosion reaction. A straight line was fitted to the data points in Figure 12 to calculate the activation energies by multiplying the slope with the gas constant, R.
The effect of temperature on ion transport through the film and the corrosion process are summarized in Figure 12 which shows the trend of activation energy determined for ion conduction in the film ($R_p$) and corrosion process ($R_{ct}$). The activation energy for the corrosion process is very much higher than ion conduction during earlier immersion.

If we compare these result with zinc phosphate full system coating as reported previously [1, 2, 11] we see that $R_p$ values for 2 coats of zinc rich full system are similar to those for the first semi-circles on the thicker zinc phosphate full system coating; suggesting that the interpretation offered here is correct. However the high activation energy values seen in the zinc phosphate full system coating were not seen here.

Fig- 10: Nyquist plots at various temperatures for zinc rich full system coating after 21 days of exposure at 50°C

Fig- 11: Nyquist plots at various temperatures for zinc rich full system coating after 35 days of exposure at 50°C

Fig- 12: Arrhenius plots of $R_p$ and $R_{ct}$ for zinc rich full system coating after (a) 21 days and (b) 35 days of exposure at 50°C

It is evident from Figure 13 that the activation energy for the corrosion process decreasing overtime, and at the end of the exposure, they are quite similar. It is suggested that at this stage ion transport in the coating might be controlling the corrosion process. In the beginning the activation energy values get smaller over time because of coating degradation.

Fig- 13: Evolution with time of activation energy for ion conduction and corrosion process for zinc rich full system coating (duplicate sets)
CONCLUSIONS

In contrast with the zinc phosphate full system coating [1], zinc rich full system gave the same conclusion as the thinner coatings [2] where the coating resistance from EIS cannot be controlling the corrosion reaction. The activation energies generated for the corrosion process here (78–97 kJmol-1) are very much higher than those of ion transport through the coating (19–37 kJmol-1) during early immersion (Figure 13).

Further interesting findings come from the activation energy trends over time particularly for the corrosion process where the value is decreasing, so that at the end of exposure the values for Rp and Rct become quite similar. It is suggested that at this stage ion transport in the coating might be controlling the corrosion process unlike at the beginning; the activation energy is getting smaller due to coating degradation.

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