Does “coating resistance” control corrosion?


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A B S T R A C T

Bacon, Smith and Rugg first identified a link between the DC resistance of a coated panel after exposure to salt solution and the performance of the coating in an exposure test. Mayne explained this as “resistance inhibition”. This paper will consider what coating resistance measured by Electrochemical Impedance Spectroscopy (EIS) tells us about factors controlling the corrosion process. It shows that the activation energy for ion conduction does not correspond with that for the corrosion process, suggesting that the paint resistance we measure is not the controlling factor. Furthermore, tests on a model galvanic cell reveal a significant influence of both electrode polarisation and resistance inhibition.

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1. Introduction

EIS has now become a standard technique for characterisation of organic coatings, and has proved valuable in assessing their performance [1,2]. The impedance behaviour of coated panels can be measured during immersion in a test environment or at intervals during cyclic testing. The coating resistance and other useful information can be extracted by fitting the impedance response across a range of frequencies to a suitable model circuit [1]. A typical model circuit includes the coating resistance and capacitance and components corresponding to the electrochemical behaviour at the metal-coating interface. The capacitance of the coating changes rapidly during immersion and is often used to measure the rate of water uptake [3]. The coating resistance generally falls over time and is an indication of coating breakdown. Mansfeld [1] identifies the coating resistance as “pore resistance” with conduction by ion transport within pores in the coating. The parameters from the coating-metal interface can in principle quantify the corrosion rate of the underlying metal.

The connection between “coating resistance” and protective performance was identified by Bacon, Smith and Rugg as long ago as 1948 [4]. They carried out DC resistance tests on coated panels over a few weeks of immersion and compared the results with the exposure performance of the same systems. Over 300 samples were tested and it was found that those which performed well outside doors all showed a high resistance (>10⁸ Ω cm²) over the whole period of the laboratory test. With poor coatings, resistance quickly fell by several orders of magnitude. They proposed that “coating resistance” could be used to identify good and bad coatings from laboratory tests.

However, we should recognise that these tests were on coated steel panels and the separate impedances of the coating and the coating-metal interface were not determined. Yet the DC current has to pass through both the resistance of the coating and that of the interface which add together because they are connected in series. If the coating was good and the corrosion rate extremely low, we can infer from the Stern–Geary equation [5] that the polarisation resistance of the interface (inversely proportional to the corrosion rate) must be high and likely to be similar to that of the coating.

Mayne [6] supports the argument that the ionic resistance of a coating is the chief factor providing protection, ruling out transport of oxygen or water as rate-controlling processes, and suggesting a mechanism (“resistance inhibition”) by which the coating resistance greatly reduces the corrosion current. This supposes (based on experimental observations of alkali-filled cathodic blisters separate from the orange rust spots after exposure of painted steel panels to salt solution) that corrosion occurs by an electrochemical mechanism with anode and cathode at different sites beneath the coating. Electrons pass from anode to cathode through the metal, but for current to flow around a complete circuit charge needs to pass, as ions, through the coating at both anode and cathode. Fig. 1 shows a schematic representation of these processes.

Mayne also suggested a way to modify the Evans diagram [7] to include the ohmic potential drop through the coating, iRg. This takes account of the fact that with the coating the anode and cath-
ode are no longer at the same potential, because a potential drop is needed to drive the corrosion current through the paint. The film resistance, $R_f$, consists of two parts, $R_a$, the resistance of the current pathway through the coating above the anode (or anodes) and $R_c$, that above the cathodes. The separate ohmic potential drops can be indicated on the diagram, with the potential measured by a reference electrode in the test solution lying below that for the cathode (if a reference electrode were to be placed below the paint) and above that for the anode. The position of this potential will depend upon the relative magnitudes of $R_a$ and $R_c$, so that if $R_a$ falls as corrosion develops the measured potential of the panel falls, even if the potentials of the anode and cathode remain the same.

In Fig. 2 current has been plotted on a logarithmic scale, to accord with Tafel behaviour, rather than the linear scale used by Mayne.

This model will not fit all circumstances, for instance Kendig [8] observed with a thin coating of polybutadiene on steel that cathodic disbonding by cathodic alkali took place around dark rust spots, showing that a direct ionic pathway existed between anode and cathode. A similar situation exists with neutral blisters where anode and cathode co-exist within the blister. Doherty demonstrated by scanning Kelvin probe (SKP) and scanning acoustic microscopy (SAM) that anodic blisters could develop alongside, and connected to, cathodic blisters then as rust precipitated out became cathodes that triggered the development of fresh anodic blisters [9]. In these situations the cell current has no need to pass through the paint.

What we wish to highlight with Mayne’s model (see Fig. 1) is that in the path for the corrosion current $R_a$ and $R_c$ are connected together in series, so if one is much bigger than the other then the larger one controls the corrosion rate, yet when we measure the resistance between the metal and the solution (as we do with EIS or similar methods) then the resistances are connected in parallel and the smaller of the two will control the measured value. The implication is that we can’t directly measure the parameter we really need to know.

The aim of this paper is to develop this argument and see if this difficulty is real. It will bring together recent published research results from Oxford students [10–14] and put them into context. I will not try in this short paper to bring together the many other pieces of supporting (or opposing) evidence that can be culled from the work of other researchers.

2. Methods and materials

The first part of this work [10] was carried out to investigate the behaviour of an organic coating on galvanised steel and better understand how the galvanic cell established when a small area of zinc is removed by damage (or at a cut edge) would operate if anode and cathode were both covered by a polymeric coating. Is the current large enough to provide sacrificial protection? To understand this situation separate electrodes of mild steel (Q-panel) and pure zinc (Advent Research Materials Ltd.) were mounted side-by-side in cold-setting resin (Lecoset), ground flat, and coated with a thin (dft = 45 μm) film of poly(vinyl butyral) using a spreader bar. Full details have been published elsewhere [10]. When the sample was immersed face down in 1% by weight NaCl solution this arrange-

![Fig. 1. A schematic representation of a corrosion cell beneath an organic coating to illustrate the operation of resistance inhibition.](image1)

![Fig. 2. Mayne’s modification of an Evans diagram to allow for coating resistance.](image2)
Fig. 3. Current changes in a zinc-steel bi-electrode when switched between in and out of the test solution (after days immersion) [10].

3. Results

Fig. 3 shows a ZRA current trace after almost 13 days of immersion. In this test the sample has been removed from the solution and dried with tissue in order to determine whether current can pass from anode to cathode along the paint-metal interface or only flows through the coating and solution as in Mayne’s model. Although a small residual current does flow after removal from solution (labelled 4), it is clear that the almost all of the current (labelled 1) flows via the solution while the sample is immersed. The current traces show large spikes when the sample is re-immersed (labelled 5) which will be argued later to be an indication of polarisation of the electrode process (but were previously linked with water loss during the dry period [10]).

Fig. 4 shows separate Nyquist plots for the steel and the zinc and the result for the coupled pair, all taken on the 13 day. In this instance the zinc shows two semi circles, whereas the steel gives only one, but the impedance of the coated steel is very much greater than that of the zinc. Thus if the pair is coupled the impedance response is virtually identical to that of the zinc, because the low impedance of the zinc + coating short circuits that of the steel + coating. These preliminary experiments confirm that the highest resistance in the current pathway for a zinc–steel couple cannot here be determined from EIS on the coupled electrodes. The same would be true for a piece of coated galvanised steel with the underlying steel exposed.

The plots in Fig. 4 show results for immersion in solutions with different salt concentrations – used to ascertain the mode of conduction in the coating. The zinc sample shows that coating resistance (from the high frequency semi-circle) increased as the salt solution was diluted: D type behaviour [16] as expected for this coating, but the steel showed little change. It is of particular interest to note that when sucrose was added to the most dilute salt solution (which will lower water activity), there is a further increase in resistance, showing that for D films resistance is influenced by water activity, the controlling factor with I films, as well as ion concentration.

Fig. 5 shows similar results for a different zinc-iron couple coated with 40 μm of the vinyl coating after 13 days exposure to salt solution [11]. Here the impedance of the steel (a) is still high but much smaller than the zinc (b). Both show a distinct high-frequency semi-circle arising from the coating resistance but that on the steel
is clearly much smaller. When we examine the plot for the couple (c) we can see that it closely resembles that for steel and is not influenced by simultaneous connection to the zinc (remembering that the potential has been held the same for both).

For the clean steel-rusty steel couple with an epoxy coating, the rusty steel became the anode, but for the first few hours exhibited an impedance $>10^5 \Omega$; however after 24 h the impedance for the rusty steel was clearly much smaller than the clean steel, and remained so as the test progressed (Fig. 6(a) and (b)). When the two electrodes are measured coupled together (c) the plot is the same as the rusty electrode, and the behaviour of the clean steel which is expected to control the corrosion rate, because of its much higher resistance, cannot be detected.

When potential pulse tests were carried out after 7 days exposure and plotted with the potential values corrected for ohmic potential drop in the coating so as to show the true polarisation behaviour, reasonable Tafel lines were fitted (Fig. 7), though care was needed to choose the correct film resistance values [13]. The arrow has been drawn at a couple current of 19 nA and the ohmic potential drop is found to be 145 mV, which agrees quite well with the calculated value [$= \frac{1}{R_\text{film}}$].

At this point both reactions are polarised significantly, but the ohmic potential drop also produces a sizeable reduction in current. Similar results were obtained with Zn-steel and Cu-steel couples.

The tests at elevated temperature on steel panels coated with polyamide cured epoxy and epoxy-phenolic coatings showed a rapid fall in impedance over a few days, yet the same coatings sustained a much higher resistance when tested at room temperature over periods of weeks typical of “good” coatings.

When samples tested at 50 °C were allowed to cool and EIS measurements were made across a range of lower temperatures, then impedance significantly increased. Fig. 8 shows a result for a polyamide cured epoxy coating 75 μm thick after 3 days exposure to 3% NaCl at 50 °C. Curves such as this, where two distinct semi-circles can be seen, were selected to obtain independent measurements of coating resistance and charge-transfer resistance.

By fitting a model circuit (Fig. 9) the resistance of the coating and the charge-transfer resistance of the corrosion reactions at the interface were determined.
Fig. 6. EIS results for a clean steel-rusty steel couple coated with a 86 μm epoxy coating. (a) rusty steel, (b) clean steel, (c) clean and rusty steel couple together, after exposure to NaCl solution for the times indicated.

From these resistance values Arrhenius plots (log R vs 1/T) were made for this coating and the epoxy phenolic to determine activation energies for the rate-controlling processes. In Figs. 10 and 11 a good fit to the Arrhenius equation can be seen for both the coating resistance and the charge transfer-resistance that depends upon corrosion rate. It is evident that for both coatings the process that determines $R_{ct}$ and controls corrosion has a higher activation energy.
energy that for ionic conduction in the coating which determines $K_i$. The activation energies for ionic conduction are similar to those obtained by Mayne for D type samples [17]. Panels of this size would be expected to include D regions in the coating that would control the overall resistance, because of their lower resistance.

4. Conclusions

The EIS tests on bi-electrodes have made it quite clear that in many cases the measured impedance of a coated panel will correspond to whichever electrode (anode or cathode) provides the easiest current path and that if the resistance inhibition model for protection holds for those systems the high-resistance pathway which controls the corrosion current is not measurable. The good agreement between film resistance and performance often found in practice may come about because in a good coating both are high. The tests on coated panels at elevated temperature show that the temperature dependence of the measured impedance does not match that of the corrosion process. This shows that the coating resistance we measure cannot be controlling the corrosion process. However it remains possible that if the higher part of the total resistance between anode and cathode, at either cathode or anode, (which cannot be measured) is in control, then the activation energy for ion transport for that part of the coating could be higher and determine the activation energy for the corrosion process. Higher activation energies have been found for intact free films of paint [15,17].

The potential pulse tests on bi-electrodes have produced examples of “Mayne diagrams” in which the electrode behaviour is credible and the ohmic potential drop that produces resistance inhibition can be measured experimentally. In some cases polarisation of both electrodes is clearly present, so that the corrosion process is under mixed control.

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