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# A Novel Non-Destructive Sensing Technology for On-Site Corrosion Failure Evaluation of Coatings

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**ABSTRACT** The quality of coating and the resulting rate of corrosion of the underlying metal substrate can be measured by a variety of corrosion measurements (Tafel, electrochemical impedance spectroscopy) by using standard laboratory electrochemical cells. However, there is always a need of low cost, portable, and non-destructive electrochemical cells, which can be used on-site field for condition monitoring of large structures for example bridges and large infrastructures, complex operating systems as aircrafts, precision machines, petrochemical processes, automotive, and locomotives. This research has developed state of the art cells fabricated by using a special magnetic aluminum compound ( $Al_nX_n$ ), which is highly electrically conductive and corrosion resistive. The research has commissioned for deploying this novel sensing technology for micro-defects detection, corrosion rate measurement, and condition assessment of the defected coatings. Tafel measurement facilitated by these non-destructive cells is used to detect micro-defects and corrosion rate measurement while electrochemical impedance spectroscopy measurement is facilitated to measure the coating condition. This technology has been successfully tested and commissioned on automotive, hazardous compartments with polymeric coatings and bridges to assess their coating condition in terms of their structural integrity. Post design testing involved the installation of these cells, running diagnostics, data acquisition, and macrographs to predict structural defects and the resulting corrosion rate. This technology enables the design process to incorporate operational conditions and fully realize more durable and reliable solutions to be applied to high-value large structures and complex interacting systems. Current developments in corrosion condition monitoring especially cost effective and non-destructive techniques to assess structural integrity beneath nonconductive and polymeric coating were long awaited. This reported development will revolutionize durability and reliability assessment techniques to enhance safety and mitigate catastrophic failures for cost savings and avoiding fatal accidents.

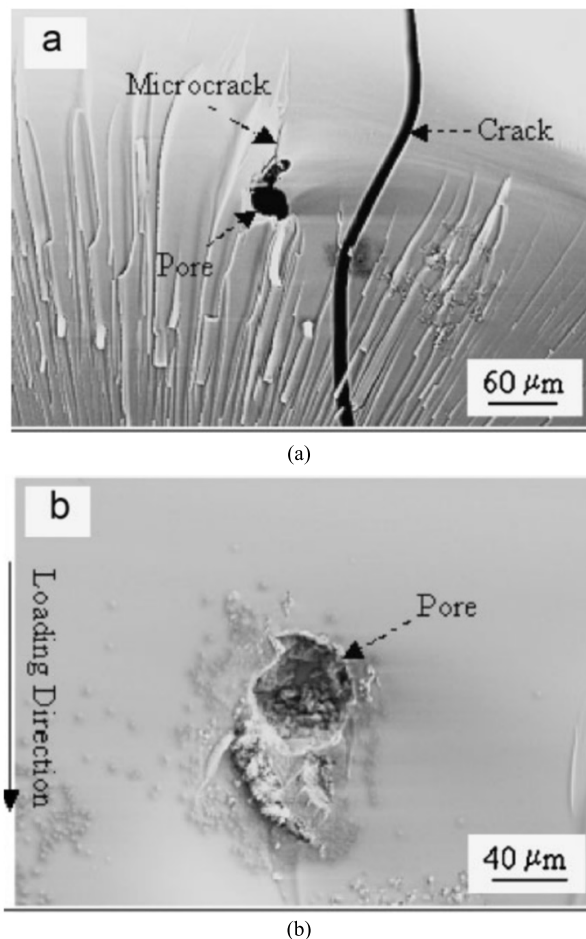
**INDEX TERMS** Coating, corrosion, non-destructive testing, electrochemical impedance spectroscopy (EIS) measurement, electrochemical cell, micro-cracks.

## I. INTRODUCTION

### A. ELECTROCHEMICAL TESTS

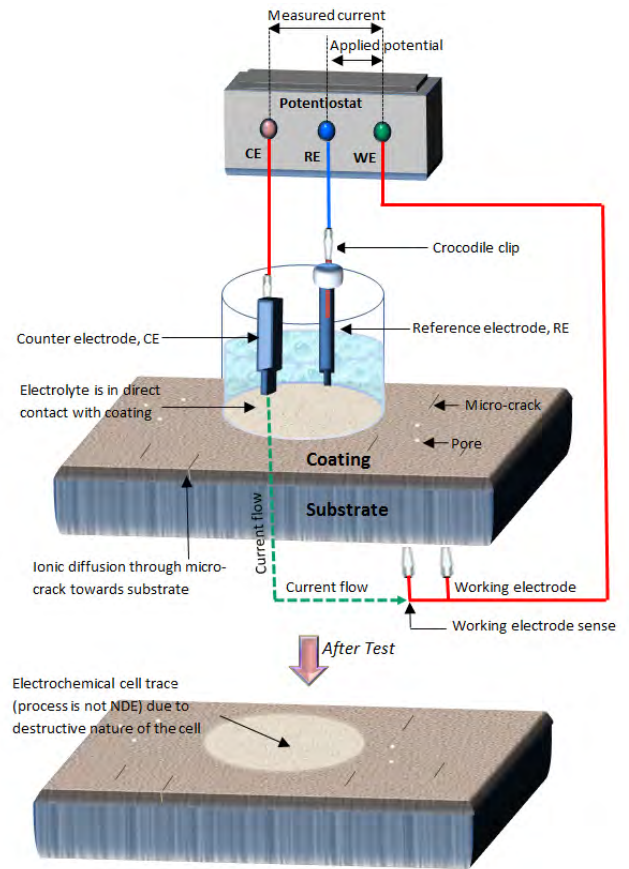
To protect objects made of metal, they are coated with corrosion resistant materials such as polymeric and metal coatings. There are a number of reasons why a particular coating may fail. It is fairly common to ascribe these failures to one of four causes, namely, mis-application of the coating, defective coating, the wrong choice of coating, or exposure to an unanticipated environmental excursion [1]. During its service life, the coating undergoes changes in mechanical properties leading to formation of micro-defects including both pores and/or micro-cracks which upon mechanical and/or thermal

loading subsequently propagate and expose substrate to atmospheric moisture and oxygen as shown in Fig. 1. This action results in accelerated dis-bonding of the paint and flake formation from the coating-substrate interface [2]. Fig. 1 shows the pore-like defect in a polymeric coating. In Fig. 1 (a), a micro-crack can be seen that initiated from a pore and propagated a short distance [3]. These cracks typically propagate roughly parallel to the loading direction. Thus, macroscopic splitting failure under compression likely result from pores (Fig. 1 (b)). For practical applications, it is desirable to deposit continuous, defect-free coatings to achieve optimal performance.



**FIGURE 1.** SEM fractographs demonstrating (a) a pore-like flaw, micro-crack, and crack on the fracture surface, and (b) a pore-like flaw, which resulted in macroscopic splitting failure under compression [3].

There are numerous porosity tests in the literature. The most latest, powerful, rapid and reliable porosity testing includes the electrochemical techniques [4]–[6]. Electrochemical techniques can be used to expose the underlying substrate in a carefully chosen electrolyte. An appropriate electrolyte will allow contact with the substrate through pores and/or micro-cracks to monitor the corrosion resistance of coating [7]. Electrochemical tests include both DC (direct current) and AC (alternating current) tests. DC measurements contain, such as Tafel extrapolation [8], cyclic voltammetry [9], linear polarization resistance (LPR) [10] and corrosion potential monitoring [11]. The AC electrochemical measurement includes electrochemical impedance spectroscopy (EIS) [12]–[14]. EIS can provide a powerful tool for electrochemical measurements on coating integrity. Both DC and AC measurements can be performed rapidly with standard electrochemical instrumentation i.e. electrochemical cell and potentiostat. These measurement methods (DC and AC) are much reliable and have been used to determine the percentage through-porosity in coatings on steel and aluminium substrates. However, such techniques require a proper laboratory setup including potentiostat,



**FIGURE 2.** Schematic diagram for standard electrochemical cell setup for performing the electrochemical analysis of coating.

electrochemical cells with proper electrode configuration. Also such analysis requires appropriate environmental conditions to reduce noise by using Faraday cage [15]. In electrochemistry standard laboratory electrochemical cells are used to characterize and analyze micro-defects in protective coatings by using above standard electrochemical techniques [16]–[19]. Therefore, the application of these techniques is rather limited to in-lab samples testing and cannot be performed where direct coating condition monitoring of large structures is required for example on-site field testing of oil & gas pipelines, bridges and automobiles. Therefore, there is a need for a simple and easily accessible electrochemical cell which can facilitate the above electrochemical techniques to detect micro-defects and analyze the condition of coatings of large structures during operation.

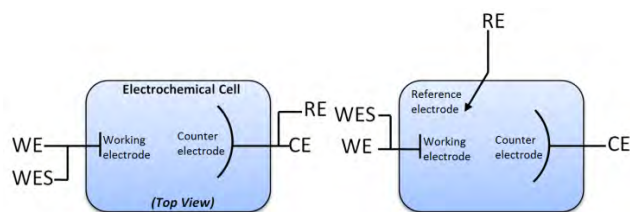
## B. STANDARD ELECTROCHEMICAL CELL

A typical electrochemical setup for coatings consists of an electrochemical cell and a potentiostat. In testing practice, a standard electrolytic cell (the system under investigation) is setup consisting of an electrolyte, a reference electrode (RE), a counter electrode (CE), working electrode sense (WES) and the coated sample of interest as shown in Fig. 2. The coated sample is called the working electrode (WE) and can be seen to possess micro defects providing the pathways

for corrosive ionic species towards the substrate. The WES is required when measuring currents in excess of 200mA. All the electrodes are connected to an electronic instrument called as potentiostat. The potentiostat applies the potential and analyses the response of the system to determine the electrochemical response of the system. The reference and counter electrodes are placed in the electrolyte, generally a solution that most closely resembles the actual application environment of the material being tested.

The complete cell is placed above the coated sample (working electrode) such that the electrolyte comes in contact with the coated side of sample.

In the electrolyte, an electrochemical potential (voltage) is measured by the potentiostat as an energy difference between the reference and the working. The current flows between counter and working electrode (shown by green line) which follows the path as: counter electrode  $\rightarrow$  electrolyte  $\rightarrow$  coating  $\rightarrow$  substrate  $\rightarrow$  working electrode. Although the standard electrochemical cell is significantly used among wider scientific community however the destructive nature of cell due to the direct contact of electrolyte with the coating limits its application when on-site field measurements are required as shown in Fig. 2. The coating area exposed to the electrolyte can lose its adhesion and can later on result in complete failure during application.



**FIGURE 3. Schematic overview of the (a) 2-electrode setup (b) 3-electrode setup.**

The most basic form of the cell has two electrodes (2-electrode configuration) as shown in Fig. 3 (a). The potentiostat cell cables WE+WES leads are connected together to working electrode and cell cable CE+RE leads are connected to counter electrode as there is no reference electrode. It is also used in measurements of electrochemical measurements at high frequencies ( $>100$  kHz), especially when the thickness of coating is very large or coating is in very good condition (defect free - ideal) [20]. A three-electrode configuration for an electrochemical cell is most common for typical electrochemical applications. The cell cables WE+WES leads are connected together to working electrode however due to the addition of reference electrode cell cable CE connects to counter electrode and cable RE connects to reference electrode. A third additional electrode (the reference electrode) is used to determine the potential across the electrochemical interface accurately as shown in Fig. 3 (b). Such setup gives reliable results for damaged coatings which poses large defects [20]. Cell schematic in Fig. 2 is a three electrode configuration [20].

Previous investigations on coating life assessment and corrosion damage measurement of the substrate in real time were reported in [21]–[28], [30]–[34]. To determine the quality of the coating and the resulting rate of corrosion of the underlying metal substrate, a variety of corrosion measurements can be done (Tafel, EIS). To facilitate these measurements, in the present work, we develop an on-site field electrochemical sealed cell which can easily be used to perform the corrosion rate analysis of the underlying substrate and non-destructive coating quality assessment. The cell can detect the micro-defects in coating. The cell can be used in applications like infrastructure, automotive and aerospace.

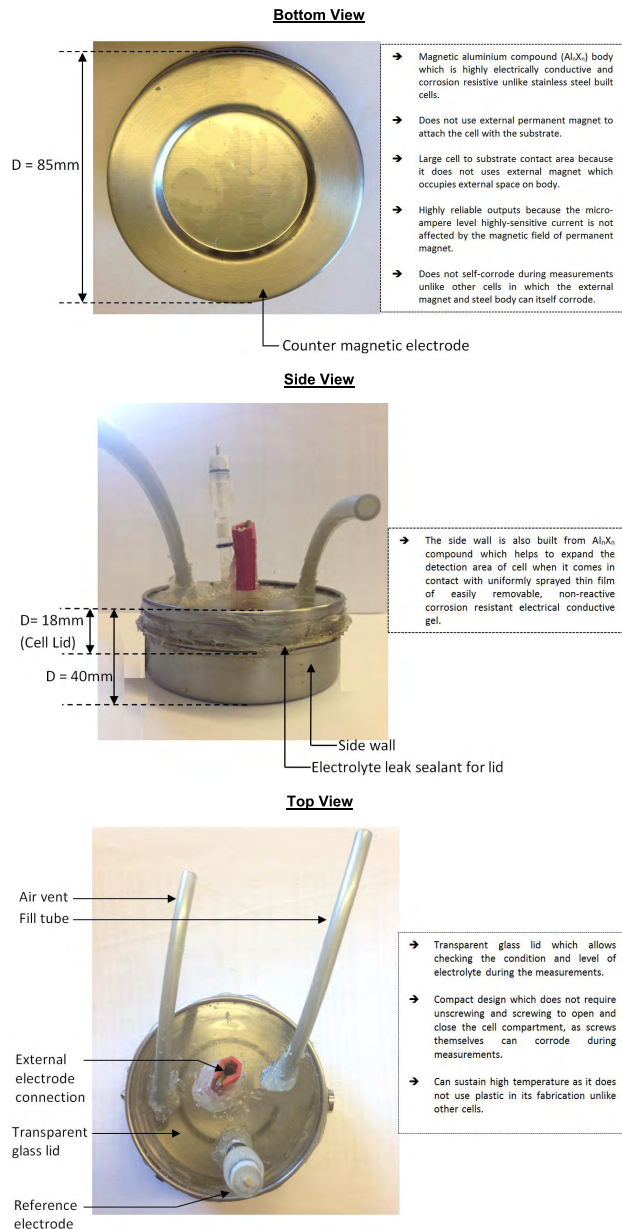
Sealed magnetic cell which is fabricated using stainless steel and can clamp with the substrate is already commercially available but the downside of such cell is that it uses an external permanent magnet to clamp the cell with the substrate. There are three downsides of using external permanent magnet, firstly it reduces the contact area of cell with the substrate by occupying a large area on cell's outer body, secondly the magnetic field of permanent magnet directly affects the micro-ampere level highly-sensitive current which can reduce the reliability of results, thirdly, the external magnet can itself corrode during the corrosion evaluations. Another downside is the use of stainless steel in fabrication as it has poor electrical conductivity and relatively low corrosion resistance. These drawbacks have been addressed by developing a sealed magnetic case of magnetic aluminium compound  $Al_nX_n$  which has high electrical conductivity, high corrosion resistance and strong clamping power. The  $Al_nX_n$  built cell does not need external permanent magnet. Also  $Al_nX_n$  built cell in combination with a specialised gel can be used to detect the defects (pores and micro-cracks) and condition of coating. The  $Al_nX_n$  cell shares some common manufacturing features with commercially available cell for example inlet filling tube and outlet air vent tube and the electrode positioning however  $Al_nX_n$  built case and its distinct working mechanism by using the specialised gel significantly improves the efficiency and reliability of  $Al_nX_n$  cell.

## II. PORTABLE ELECTROCHEMICAL CELL (PEC)

### A. FABRICATION

The PEC system unlike standard electrochemical cell is a portable cell with a magnetic body which can be attached to any horizontal, vertical and upside down metal object. In this way the object constitutes the bottom of the PEC and the object itself can be used as an electrode. The cell can be filled with the electrolyte (3.5% NaCl solution) by using a tube that leads directly into the cell. A similar tube allows air to escape to ensure a completely filled cell in order to prevent air pockets inside the cell compartment. The inlet filling tube is positioned in a way such that it is always lower in level compared to outlet/ air vent tube. Another purpose of inlet and outlet tubes is for cleaning the cell by flushing the used electrolyte out and filling it with fresh electrolyte. PEC design does not allow the electrolyte to come in direct contact with the coating due to magnetic aluminium compound  $Al_nX_n$





**FIGURE 4.** Bottom view, side view and top views of portable electrochemical cell (PEC).

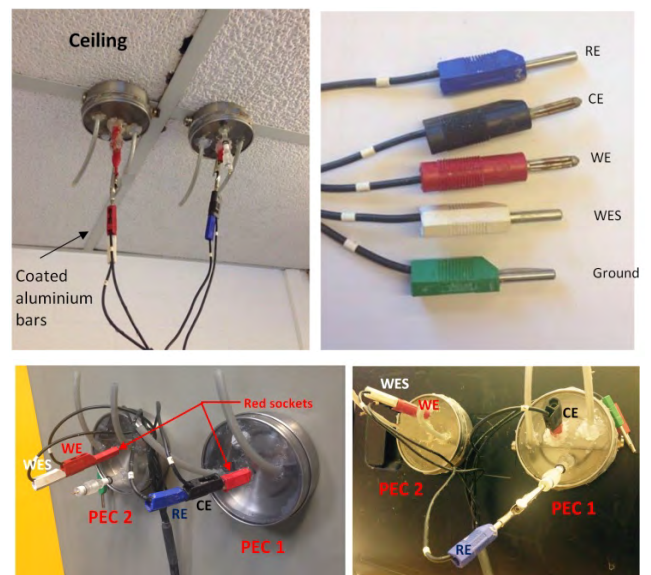
base of cell which ensures non-destructive monitoring of coating condition.  $Al_nX_n$  has been specifically chosen for fabricating PEC to obtain optimum corrosion monitoring data due to its exceptional electrical-conductive and corrosion resistance properties.  $Al_nX_n$  is prepared by doping aluminum with ferrite nanoparticles [35].

There are two main parts of PEC: (i) the bottom and (ii) the cell lid. The circular bottom of PEC consists of  $Al_nX_n$  area which constitutes the counter electrode as shown in Fig. 4 (bottom view).  $Al_nX_n$  electrode is magnetic which allows the cell to be easily placed on the metal surface. From bottom to top this consists 2 layers: the  $Al_nX_n$  container and the cell lid (side view). The  $Al_nX_n$  container holds the electrolyte such that the container is tightly sealed to the lid using

silicone gel to avoid any electrolyte leakage. The reason for selecting  $Al_nX_n$  as a counter electrode is its light weight, corrosion resistance and high electrical conductance especially for sensing  $\mu$ Amp and PicoAmp corrosion currents for high impedance coatings (defect free). The top lid holds the electrode connector, 2 filling holes and a reference electrode. The reference electrode (saturated calomel) is placed near the counter electrode for accurate measurement.

## B. DESCRIPTION OF USE

PEC has an advantage over standard electrochemical cell that because of its completely sealed design it can be attached to metal objects with any orientation (vertical, horizontal, upside down) which allows its restriction free use in the laboratory as well as on the field as shown in Fig. 5 (top left). The object it has been attached to can be used as counter (or working) electrode. This object can be both coated and an uncoated structure. For non-destructive testing of coated object (without damaging the coating), an additional PEC is used as a second electrode which works in combination with the first PEC. This combination allows the condition monitoring of large coated structures with no possibility to reach the metal substrate. The two PECs in combination can work both in 2-electrode and 3-electrode configuration by using the leads shown in Fig. 5 (top right).



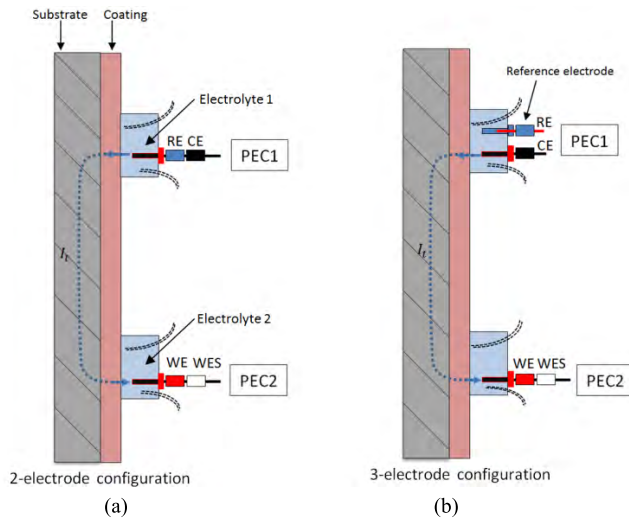
**FIGURE 5.** The top left image shows the corrosion monitoring of steel coated ceiling bar. The top right image shows the type of leads. Two PEC are being used together for analysing the condition of coating in a two electrode configuration in top right and three electrode configuration in bottom left. This non-destructive system is able to detect the micro-defects over the large surface area of coating.

Fig. 5 (bottom left) shows the two electrode configuration in which the base metal cannot be accessed, the potentiostat cell cable CE+RE leads are connected together to the stainless steel electrode of PEC1 (red socket). The potentiostat cell cable WE+WES leads are connected together to the stainless steel electrode of PEC2 (red socket). In the shown 2-electrode

connection no reference electrode is used. However, if reference electrode is used to form a 3-electrode configuration, a reference electrode in PEC 1 is connected by using cell cable RE lead as shown in Fig. 5 (bottom right). The cell cable CE lead is connected to counter electrode in PEC 1 while cell cable WE+WES leads are connected together to the working electrode of PEC 2.

### C. WORKING MECHANISM

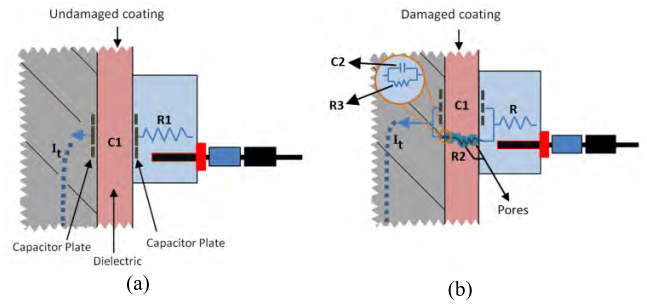
When both PEC's are attached to the polymeric coated steel an electrochemical cell is formed between the  $Al_nX_n$  electrodes in the back of both PEC's. Connecting the potentiostat cell cable WE+WES leads to the  $Al_nX_n$  electrode of PEC2, and the potentiostat cell cable leads CE+RE to the  $Al_nX_n$  electrode of PEC1, a 2- electrode configuration is formed. If reference electrode is connected to cell cable lead RE, a 3-electrode cell is formed. Schematic representations of both the 2- and 3- electrode configurations are shown in Fig. 6 (a-b) below. Like standard cell, in PECs, the current also flows between working and counter electrode. However, unlike standard cell it has to pass through the coating twice.



**FIGURE 6.** Schematic representations of (a) 2- electrode configuration PECs and (b) 3- electrode configuration PECs.

Using two PEC's results in a the path of current  $I_t$  that flows from electrode PEC 1 → electrolyte1 → coating → substrate → coating → electrolyte 2 → electrode PEC 2.

A metal covered with an undamaged coating generally has very high impedance resulting in negligible current  $I_t$  flow mostly in PicoAmp as shown in the equivalent circuit in Fig. 7 (a). The model includes a very high resistor R1 (due to electrolyte resistance) and a very low coating capacitance C1 in series. In the case of a coated metal in contact with the electrolyte through PEC's counter electrode (at bottom), the substrate metal is one plate, the coating is the dielectric, and the electrolyte in PEC is the second plate as shown in Fig. 7 (a). The coating capacitance relationship is given by



**FIGURE 7.** Equivalent circuit in 2-electrode configuration (a) undamaged coating (b) damaged coating.

the following equation [36]

$$C1 = \frac{\epsilon_0 \epsilon_r A}{d} \quad (1)$$

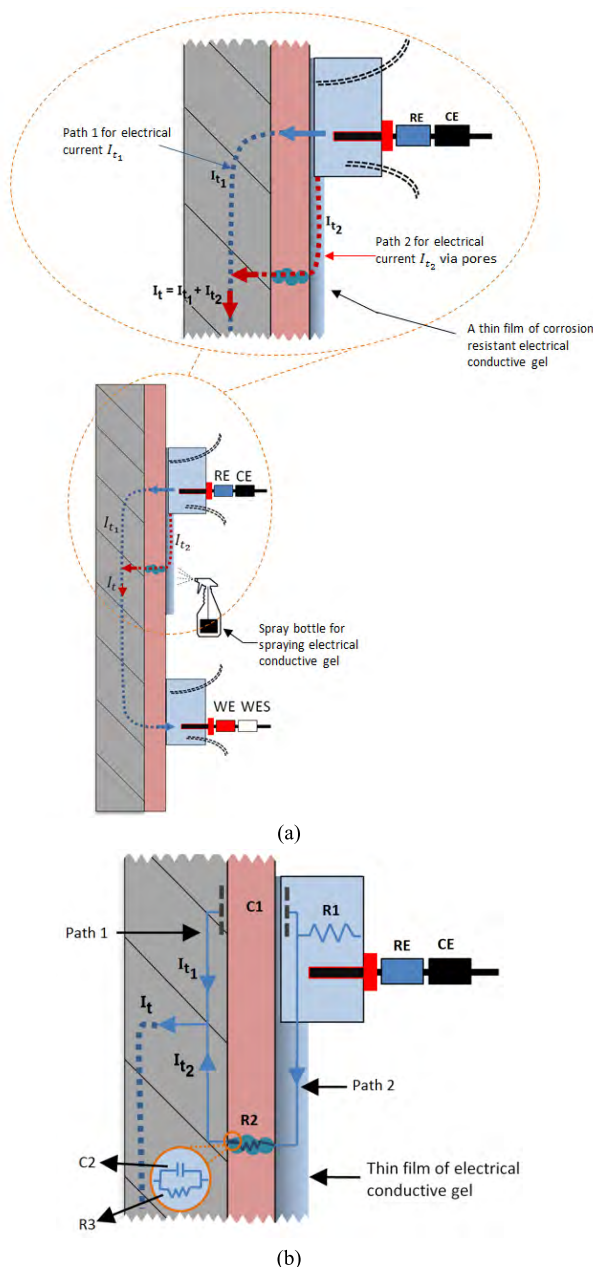
Where  $\epsilon_0$  is the electrical permittivity,  $\epsilon_r$  is the relative electrical permittivity (dielectric), A is the surface of one plate and d is the distances between two plates. Capacitors in electrochemical analysis often do not behave ideally. Instead, they act like a constant phase element (CPE) as defined below [37]. The impedance of a capacitor  $Z_{CPE}$  can be expressed as:

$$Z_{CPE} = \frac{1}{(j\omega)^\alpha C1} \quad (2)$$

Where,  $\omega$  is called the 'Frequency Response' and  $\alpha$  is the exponent equaling 1 for a capacitor [36]. Thick, high quality coatings characteristically have almost infinite impedance and very low capacitance as clear from eq. 2. It is obvious that their high impedance results in very small response AC currents  $I_t$ , especially at low frequencies where impedance element in the model dominates. On a more subtle level, their low capacitance results in small AC currents as can be seen from eq.3.

$$I_t = \frac{E}{Z_{CPE}} \quad (3)$$

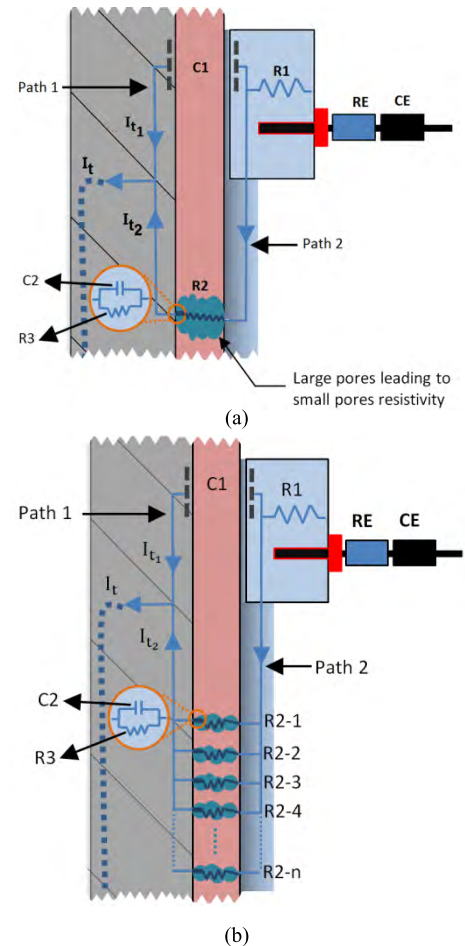
Where  $E_t$  is the potential at time t. Most coatings degrade with time, resulting in more complex behavior. The degradation results in the opening of pore(s) and/or micro crack(s) in the coating. As pores in the coating begin to expand over time, the resistance R2 associated with these pores (also called pores resistivity) increases. Also degradation results in the increase in the coating's capacitance C1. An increase in coating capacitance represents deterioration of the coating's ability to shield the metal substrate from the environment. The equivalent circuit for this situation is in Fig. 7 (b). On the metal side of the pore, it is assumed that an area of the coating has delaminated and a pocket filled with corrosive solution from an outer environment has formed as shown by orange circle in Fig. 7 (b). This solution has reached to the bare metal through pores. The interface between this pocket of solution and the bare metal is modelled as a double layer capacitance C2 in parallel with a kinetically controlled charge transfer resistance R3. Once the model has been fitted to the data,



**FIGURE 8.** (a). Schematic showing that the (a) large sized pores have smaller  $R_2$  which accounts for large current  $I_{t2}$ . (b) the magnitude of current  $I_{t2}$  through path 2 increases with the number of pores 'n' in this path.

changes in the model's parameters offer insight into the health of the coating.

The above setup shown in Fig. 6 and 7 can only monitor the condition and detect pores in the coating area which is directly under the PEC. This is because that only the coating area which is directly under the PEC acts as a conductive region for the current  $I_t$  between the counter electrode and metal substrate. The pores which are not located under the PEC will not be detected. In order to detect such pores an alternative conductive path (path 2) is needed between the



**FIGURE 9.** (a) Schematic showing that the (a) large sized pores have smaller  $R_2$  which accounts for large current  $I_{t2}$ . (b) the magnitude of current  $I_{t2}$  through path 2 increases with the number of pores 'n' in this path.

electrode of PEC and the substrate such that the current can flow from electrode to pore and then from pore to substrate via path 2 as shown in Fig. 8 (a). Path 2 is created by uniformly spraying a thin film of easily removable, non-reactive corrosion resistant electrical conductive gel on coating, which comes in contact with the pores forming a closed path between counter electrode and substrate. After the gel has been sprayed, there will be two parallel paths: path 1 and path 2. Path 1 is via undamaged coating region therefore it has very high impedance resulting in negligible current flow  $I_{t1}$  while path 2 contains pores causing low impedance which results in large current flow  $I_{t2}$  as shown in Fig. 8 (b). The net current  $I_t$  which flows between PEC 1 and PEC 2 is given as:  $I_t = I_{t1} + I_{t2}$ .

The magnitude of current  $I_{t2}$  which flows through path 2 depends on the pore resistivity  $R_2$ . Smaller  $R_2$  (meaning large sized pores) accounts for large current  $I_{t2}$  which increases the net current  $I_t$  and vice versa as shown in Fig. 9 (a). Similarly the magnitude of current  $I_{t2}$  through path 2 depends on the number of pores 'n' in this path, each pore having resistance  $R_{2-1}$ ,  $R_{2-2}$ ,  $R_{2-3}$  up to  $R_{2-n}$  respectively as shown



in Fig. 9(b). The larger the number of pores, the larger is the current  $I_{t2}$  which consequently results in larger net current  $I_t$  as show in Fig. 9(b).

The change in the net current  $I_t$  is monitored by using Tafel measurement. This change is monitored corresponding to changing linear potential scan from E start to E end on the counter electrode. The results are displayed as I – E graph on computer software interfaced with potentiostat. The scanning is performed while increasing the effective area of thin film of corrosion resistant electrical conductive gel by spraying. As soon as the conductive gel comes in contact with the pores, a closed path (path 2) is formed between the counter electrode and the substrate which results in the rapid initiation of current flow through path  $I_{t2}$ . The initiation of current  $I_{t2}$  adds up with the negligible current flow  $I_{t1}$  which is flowing through non-degraded high impedance path (path 1) resulting in the increase in net current as:  $I_t = I_{t1} + I_{t2}$ . This sudden rise in  $I_t$  can be observed in the form of large trough in I - E graph. This large trough clearly indicates that the gel has encountered a defected area which can be treated. The Tafel slope is calculated from I - E graph to calculate the corrosion rate of coated substrate.

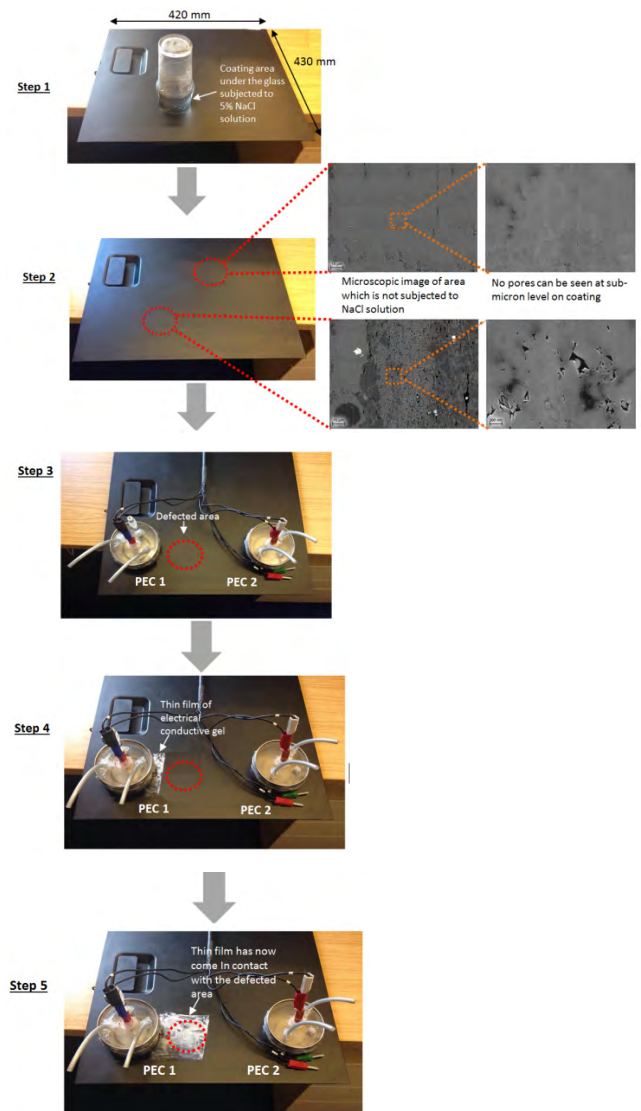
Once the defected area has been detected by using Tafel measurement, the same equivalent circuit (path 1+ path 2) is also used for EIS measurement to estimate the values of equivalent circuit parameters: electrolyte resistance R1, coating capacitance C1, pore resistance R2, double layer capacitance C2 and charge transfer resistance R3 by using Nyquist and Bode plots. These parameters help to identify the condition of coating.

The graphical analysis for Tafel and EIS measurement is discussed in the results section in detail. Thus, the complete coating condition monitoring analysis can be divided in to two stages: (1) Detecting the micro defects by using Tafel measurement and (2) Identifying the condition of coating through parametric evaluation of equivalent circuit by using EIS measurement.

### III. EXPERIMENT

A five step experimental study was performed by using a 420mm × 430mm aluminium sample coated with polymeric epoxy resin in black as shown in Fig. 10. The equivalent weight and density of aluminium are 26.9g and 2.7 g/cm<sup>3</sup> respectively. For the most part, epoxy coatings are used as functional coatings for substrate protection where corrosion resistance, impact resistance, and adhesion are essential. This coating is normally used in industrial equipment, automotive components, metal furniture and appliances [38]. The objective of this experiment was to detect invisible micron and sub-micron level defects (pores) and to determine the condition of defected coating by using a custom developed PEC.

In the first step, a selected area on the coated sample was subjected to 5% NaCl solution for 72 hours as shown in Fig. 10 (step 1). This process was performed to degrade the specific area of coating under the glass which resulted in the opening of pores in the coating. In step 2, the microscopic



**FIGURE 10.** The five step experimental study performed to analyse the working of PECs.

analysis of the subjected area was performed which was compared with the microscopic images taken from the non-subjected area of coated sample. In step 3, the two PECs were installed on the sample in 2-electrode configuration. In next step 4, a thin film of corrosion resistant electrical conductive gel was sprayed on coating by using a precise gel coat spray gun. The gel is non-corrosive, non-reactive and can easily be removed by using a dry wipe. The gel was sprayed in a stepwise manner such that the coverage area of gel was slowly increased until it incorporated the degraded area beneath it. At the same time the Tafel scan was also performed, in which the potential on counter electrode was increased from −3 V (E start) to +3 V (E end) which resulted in the corresponding change in net current  $I_t$ . The scan rate (from −3 to +3V) was set to a low value of 20 mV/sec which resulted in large scan duration giving enough time to increase the coverage area of gel gradually. In step 5, as the conductive gel came in contact

with the degraded area it resulted in the formation of a closed path (path 2) between the counter electrode of PEC 1 and the substrate. It is at this specific instance that the sudden rise in  $I_t$  is observed in the form of large trough in  $I - E$  graph indicating the location of defected area on coating.

EIS measurement was then performed on this defected sample measuring the parametric values of capacitances ( $C_1$ ,  $C_2$ ) and resistors ( $R_1$ ,  $R_2$  and  $R_3$ ) of the equivalent circuit to determine the condition of coating. AC measurement on coated sample by using two PECs in a 2-electrode configuration can be used for EIS measurements where an AC signal is applied around OCP (Open Circuit Potential). Such a measurement will result in  $Z^*$  coating impedance ( $Z_{CPE}$ ), because the path of the current passes the coating twice. In current experimental setup, the parametric values are calculated by applying AC signal with frequency varying from 0.06 Hz to 1MHz.

Before performing the above 5 step experimental procedure, the Tafel and EIS measurements of the undamaged coating sample (prior to NaCl exposure) was also performed by using PECs in order to conduct the comparative analysis between the results of damaged and undamaged coating sample.

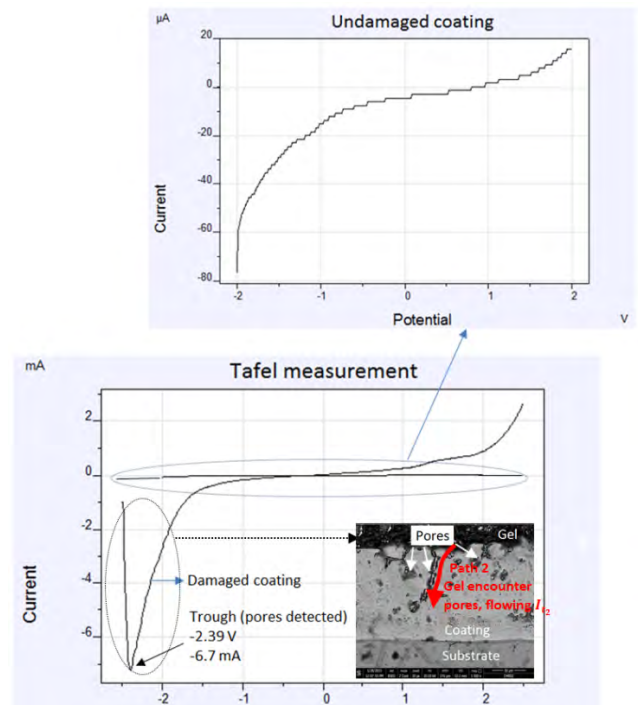
#### IV. RESULTS

The results obtained from the above experiment were divided in to two stages: (1) defect detection stage by using Tafel measurement and (2) parametric values evaluation stage of equivalent circuit by using EIS measurement.

##### A. TAFEL MEASUREMENT

From the above experimental setup, the results are shown in Fig. 11 below. It can also be seen from the trend of damaged coating that the Tafel scanning was performed from  $-3$  V to  $+3$  V at the same time when the coverage area of gel was increased by spraying (as depicted in steps 4 and 5 of Fig. 10). As the coverage area was increased during scan, the gel encountered the pores at  $E = -2.39$  V which resulted in the sudden drop in net current  $I_t$  to a level  $-6.7$  mA and then the subsequent exponential rise in  $I_t$ . This sudden drop and rise in  $I_t$  was observed in the form of a trough, indicating the detection point of pores. The exponential rise in  $I_t$  is associated with the formation of low impedance conductive path 2 through pores resulting in large current flow  $I_{t2}$ . The cross-sectional microscopic image of coating-substrate interface on the right of graph shows the instance when the gel encountered the pores and resulted in the formation of a trough due to development of path 2 causing large current flow  $I_{t2}$ .

Comparison between the Tafel measurements from damaged (after NaCl exposure) and the undamaged (prior to NaCl exposure) coating can also be seen in the graph. Undamaged coating possesses only one high impedance path (path 1) for current flow  $I_{t1}$ , therefore the magnitude of net current  $I_t$  is extremely small, mainly in  $\mu A$ 's as can be seen in Fig. 11. However, damaged coating possesses two paths (path 1 +



**FIGURE 11.** Tafel measurements for both the damaged (bottom graph) and undamaged (top graph) coatings. The graph for the damaged coating shows the trough which indicates the detection point of pores on damaged coating during Tafel scanning.

path 2), therefore, the magnitude of net current  $I_t$  is high because of the formation additional low impedance path 2 (due to presence of pores) eventually adding more current:  $I_t = I_{t1} + I_{t2}$ , mainly in mA's. It can also be seen that for damaged coating the current reached to maximum peak current of 2.4 mA which is certainly very high current indicating the presence of low resistive large sized pores within the damaged area.

"Corrosion rate" analysis of the damaged coating was performed to determine the corrosion rate of epoxy coated aluminium substrate. The rate of corrosion can be analyzed by using any one of three approaches:

- Slope analysis
- Tafel analysis
- Model analysis

In this study all three approaches were used for in-depth analysis of corrosion rate. The comparison of the results obtained from all three approaches helped to validate the accuracy of results.

At first, the 'slope analysis' was performed as shown in Fig. 12. Slope analysis identifies the corrosion potential and the polarisation resistance  $R_p$  at the zero current intersection point. The calculated quantitative results ( $E_{corr}$ ,  $I_{corr}$ ,  $R_p$  and  $C$  Rate) from slope analysis can be seen under the 'Pol Res' column of table in Fig. 13.

The 'Tafel analysis' was performed on the damaged coating to calculate the corrosion parameters from the 2 Tafel lines as shown in Fig. 14. The calculated quantitative results



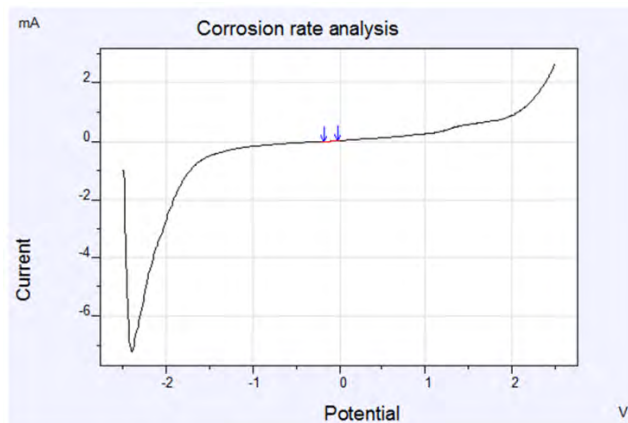


FIGURE 12. Measurement of corrosion rate by using 'slope analysis' approach.

	Pol. Res.	Tafel data	Model data
E. corr V	-0.1036	-0.4461	-0.0990
i cor. A	3.33E-6	3.807E-5	1.884E-5
I cor. A/cm <sup>2</sup>	3.33E-6	3.807E-5	1.884E-5
R <sub>p</sub> Ohm	0.06521	0.06719	0.06746
b <sub>a</sub> V/dec		1.703	0.617
b <sub>c</sub> V/dec		0.900	0.557
C. Rate mm/y	0.3867	0.4422	0.2188

☒ Automatic analysis  
☐ Slope Analysis  
☐ Tafel Analysis  
☐ Model Analysis  
☐ minimize logarithmic errors

FIGURE 13. The calculated quantitative results from slope analysis is shown under the 'Pol Re' column of table.

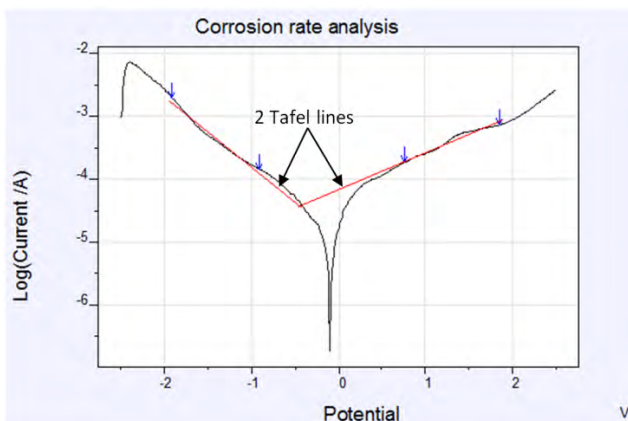


FIGURE 14. Measurement of corrosion rate by using 'tafel analysis' approach.

from Tafel analysis are shown in the second column of table 'Tafel data'. Further the corrosion parameters were calculated with a numerical analysis of the available data by using

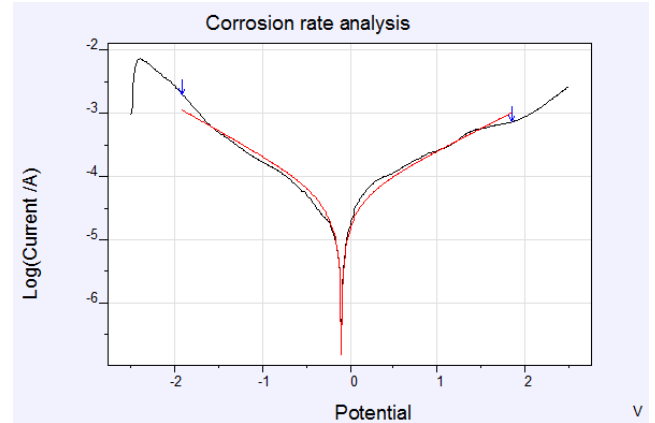


FIGURE 15. Calculation of corrosion rate by using 'Model analysis' approach.

'Model analysis' as shown in Fig. 15. Model analysis utilises the material constants i.e. surface area, equivalent weight and density (already identified in previous section) to calculate the corrosion data. The Model analysis is expected to yield the best results in most cases, because it applies complete Butler-Volmer to full data range therefore model data results are most reliable compared to slope analysis and Tafel analysis.

Comparing the corrosion parametric values obtained from all three approaches (Table in Fig. 13) shows a minute difference in the values of certain parameters. The analysis of  $R_p$  shows values close to one another in between 0.0652 and 0.0674 Ohm for all three approaches. Likewise corrosion rate shows values in between 0.2188 and 0.3867 mm/y. The table also identifies and compares the values of Tafel constants ( $b_a$ ,  $b_c$ ),  $I_{corr}$  and  $V_{corr}$ .

## B. EIS MEASUREMENT

Once the defected area and the corrosion rate of the damaged coating sample has been determined by using Tafel measurement (after performing all five steps), an EIS measurement was done on the same damaged sample in a 2-electrode configuration. The EIS scan was performed between 0.06 Hz start and 1MHz end frequency with 20mV amplitude and  $E = 0V$  vs. OCP (Open Circuit Potential). EIS measurement was performed to determine the parametric values ( $R_1$ ,  $C_1$ ,  $R_2$ ,  $C_2$  and  $R_3$ ) in the equivalent circuit shown below (Fig. 16) by using various plots for example Bode, Nyquist plots. Fig. 16 shows the circuit for the damaged coating accepted by most coating researchers.

The bode plot for both damaged and undamaged coating samples are plotted in the same graph in Fig. 17. It can be seen that initially at low frequency i.e. 0.06Hz ( $10\log(f)/Hz = -1$ ), the corresponding impedance ( $10\log|Z|/Hz$ ) for both damaged and undamaged coating stayed high till  $10\log(f)/Hz = 0.2$ . After this point the impedance for damaged coating drastically decreased while for the undamaged coating it still sustained to the same level as before till  $10\log(f)/Hz = 3.8$  was achieved which

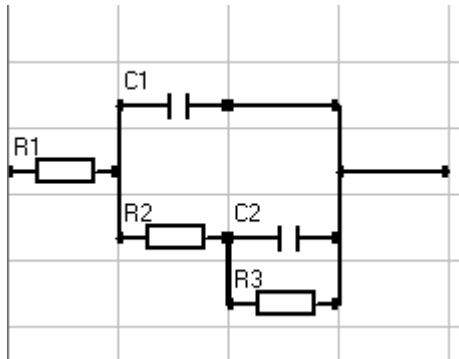


FIGURE 16. Equivalent circuit for the damaged coating.

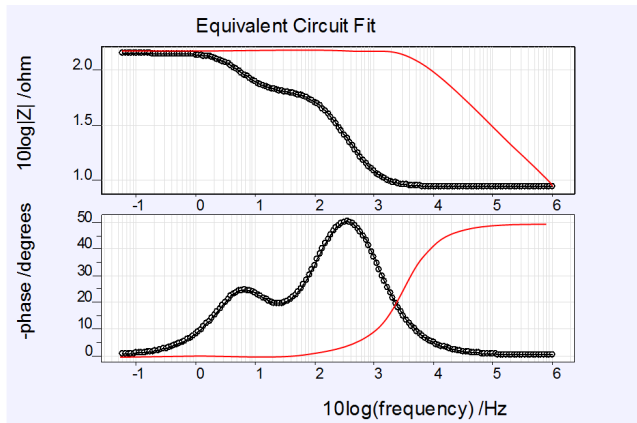


FIGURE 17. Bode plot for the damaged and undamaged coating.

is very high frequency. Similarly, for damaged coating the corresponding phase angle reached to a high level just after start while the corresponding phase angle for undamaged coating remained low till it reached a high frequency of  $10\log(f)/\text{Hz} \approx 2$ . Therefore it can be concluded that the undamaged coating has greater ability to sustain higher frequencies compared to the damaged coating. Bode plots is Fig. 17 works in accordance with the relation shown in eq. 3.

Bode plots are used to evaluate only the impedance behavior of coating w.r.t. varying frequency. Fig. 17 shows a typical Nyquist diagram derived from the experiment. The Nyquist diagram is in the form of a semicircle. The diameter of the semicircle extrapolated in the Nyquist diagram represents the charge transfer resistance  $R_3$  which is equivalent to the polarization resistance  $R_p$  [39]. Therefore the larger the diameter of the semicircle the higher the resistance,  $R_p$ , and hence, the lower the corrosion rate. It is evident from the plots, that the diameter of undamaged coating is far larger (in the power of  $E+03$ ) than the diameter of damaged coating.

The parametric values of equivalent circuit for both damaged and undamaged coating are shown in Fig. 19. Comparative analysis between both type of coatings show that the coating capacitance  $C_1$  for the undamaged coating is low ( $1.722 \text{ E-}08$ ) compared to the damaged coating ( $2.152 \text{ E-}05$ ). Likewise the pore resistance  $R_2$  for the undam-

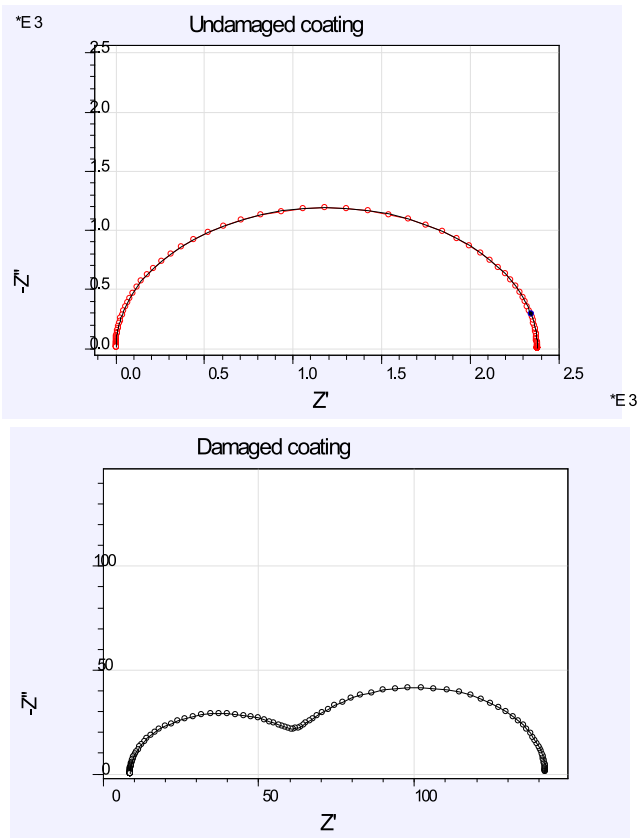


FIGURE 18. Nyquist plot for the damaged (bottom) and undamaged (top) coating.

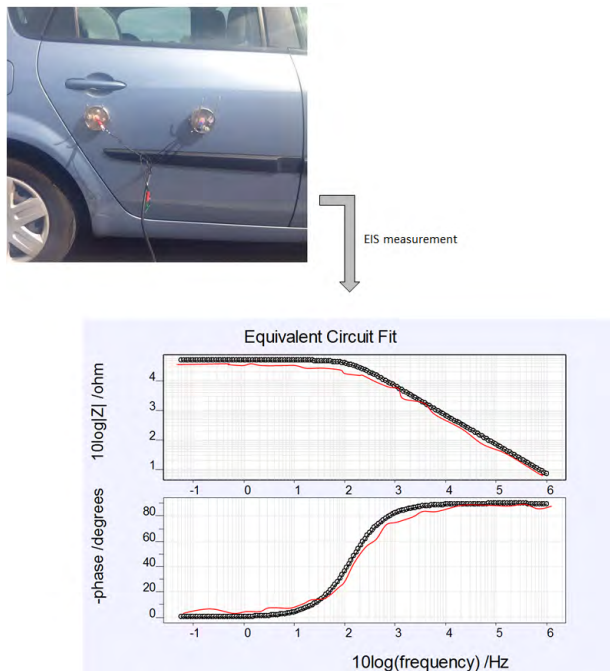
	Par	Fixed	Value	Error%	Min	Max	Unit
Par[1]	R1	<input type="checkbox"/>	8.748E+00	98.74	1.0E-4	1.0E+13	Ohm
Par[2]	R3	<input type="checkbox"/>	1.980E-02	27.75	1.0E-4	1.0E+13	Ohm
Par[3]	R2	<input type="checkbox"/>	5.805E+01	32.98	1.0E-4	1.0E+13	Ohm
Par[4]	C1	<input type="checkbox"/>	2.152E-05	72.09	1.0E-13	1.0E+1	F
Par[5]	C2	<input type="checkbox"/>	5.126E-04	68.98	1.0E-13	1.0E+1	F

	Par	Fixed	Value	Error%	Min	Max	Unit
Par[1]	R1	<input type="checkbox"/>	9.216E-02	1000.00	1.0E-4	1.0E+13	Ohm
Par[2]	R3	<input type="checkbox"/>	7.532E+03	1000.00	1.0E-4	1.0E+13	Ohm
Par[3]	R2	<input type="checkbox"/>	2.382E+03	1000.00	1.0E-4	1.0E+13	Ohm
Par[4]	C1	<input type="checkbox"/>	1.722E-08	1000.00	1.0E-13	1.0E+1	F
Par[5]	C2	<input type="checkbox"/>	6.682E-07	1000.00	1.0E-13	1.0E+1	F

FIGURE 19. EIS results for the damaged (top) and undamaged (bottom) coating.

aged coating is high ( $2.382 \text{ E}+03$ ) compared to the damaged coating ( $5.805 \text{ E}+01$ ). The charge transfer resistance  $R_3$  (equivalent to the polarization resistance  $R_p$ ) for the undamaged coating is high ( $7.532 \text{ E}+03$ ) compared to the damaged coating ( $1.980 \text{ E-}02$ ). As higher  $R_p$  accounts for the low corrosion rate, which can also be seen from  $R_3$  for the undamaged coating that the  $R_3$  is higher compared to the damaged coating. The double layer capacitance  $C_2$  for



**FIGURE 20.** EIS measurement of automotive coating taken by using PEC and standard electrochemical cell (red trend).

	Par	Fixed	Value	Error%	Min	Max	Unit
Par[1]	R1	<input type="checkbox"/>	1.243E-02	-1.00	1.0E-4	1.0E+13	Ohm
Par[2]	R3	<input type="checkbox"/>	1.932E+01	-1.00	1.0E-4	1.0E+13	Ohm
Par[3]	R2	<input type="checkbox"/>	4.968E+04	-1.00	1.0E-4	1.0E+13	Ohm
Par[4]	C1	<input type="checkbox"/>	2.342E-08	-1.00	1.0E-13	1.0E+1	F
Par[5]	C2	<input type="checkbox"/>	6.817E-09	-1.00	1.0E-13	1.0E+1	F

**FIGURE 21.** EIS results of automotive coating calculated by using the measurements of PEC.

the undamaged coating is low ( $6.682 \times 10^{-7}$ ) compared to the undamaged coating ( $5.126 \times 10^{-4}$ ) showing that undamaged coating has higher impedance and therefore more corrosion resistant.

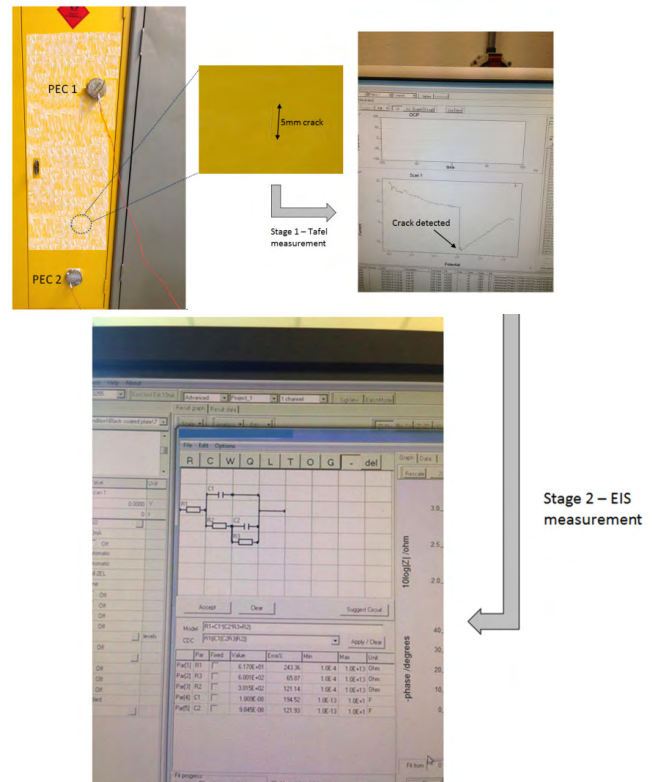
The Error% in Table 19 calculates the error of the fitted parameter for Min and Max parametric values.

## V. ONSITE FIELD TESTING

### A. EXAMPLE 1 - AUTOMOBILE COATING

Measurements in the field have been carried out to test and validate the PEC performance. In first example, the PECs were installed on back door of car coated with (D47 storm blue paint) to assess the condition of coating by using EIS measurement. The same measurements were also done with the standard electrochemical cells and the results were compared as shown in Fig. 20 (Red trend lines). EIS measurement from PECs was then used to calculate the parametric values of the equivalent circuit to measure the condition of coating as shown in Fig. 21.

It can be seen in Fig. 21 that the coating's capacitance C1 is high ( $2.342 \times 10^{-8}$ ) which indicates that the condition



**FIGURE 22.** Tafel scanning performed by using PEC to detect the invisible crack on hazardous compartment coating (Stage 1). EIS measurement is used for calculating the parametric values of the equivalent circuit (Stage 2).

of coating is good and possesses a good ability to shield the metal substrate from the environment.

### B. EXAMPLE 2 - HAZARDOUS COMPARTMENT COATING

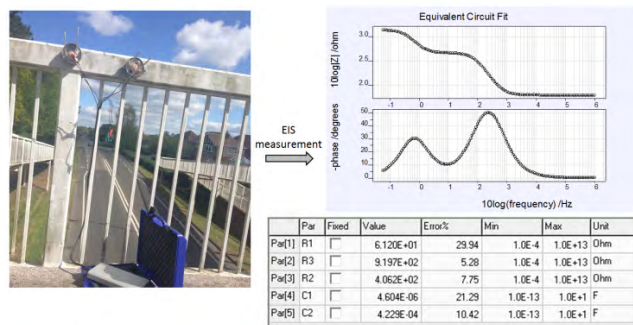
In the second example, PECs were installed on hazardous compartment. Two stage analyses was performed in which the defect in coating was detected by using Tafel measurement in stage 1 and then the coating condition was identified in stage 2 by using EIS measurement as shown in Fig. 22.

The stage 2 measurement shows a low pore resistance R2 ( $3.015 \times 10^2$ ) which indicates that the coating is defected but at the same time it also shows a high coating capacitance C1 ( $9.845 \times 10^{-8}$ ) which points out that the defect is not significant and the overall condition of coating is good. If both C1 and R2 would have been simultaneously low, then this would have been an indication of a degraded coating.

### C. EXAMPLE 3 - OVERHEAD PEDESTRIAN BRIDGE COATING

PECs were installed on overhead bridge to measure the condition of coating as shown in Fig. 23. Visual inspection showed that the coating of bridge was significantly damaged. Non-destructive EIS measurement was performed to cal-





**FIGURE 23.** EIS measurement and result of overhead pedestrian bridge coating taken by using PEC.

culate the parametric values of equivalent circuit which showed considerable low pore resistance R2 (4.062 E+02) and high coating capacitance C1 (4.604 E-06).

The results clearly show that the coating of overhead bridge is significantly degraded as both pore resistance R2 (4.062 E+02) and coating capacitance C1 (4.229 E-04) are low.

## D. CONCLUSIONS

This paper presents the development of low cost, portable and non-destructive electrochemical cell which can be used to assess the quality of coatings. This cell unlike conventional electrochemical cell is portable with magnetic body and because of its completely sealed design it can be attached to any horizontal, vertical and upside down coated metal objects, structures and systems such as steel beam, wall and ceiling, bridges, aircrafts, automobiles and locomotives. In this way the object constitutes the bottom of the cell. The cell can be filled with the electrolyte by using an inlet tube. A combination of two cells can be used for the non-destructive testing of coating with one cell forming the counter electrode while the other forming the working electrode. Such combination develops a close path for the current flow through coating and metal substrate between the two cells. This cell combination facilitates a variety of corrosion measurements (such as Tafel and EIS) to determine the quality of coating where direct access to the metal substrate is not possible. The Tafel measurement is facilitated to detect the micro-defects in coating and to measure the corrosion rate of underlying metal substrate. The EIS measurement is facilitated to measure the corrosion resistant properties of coating. Electrochemical measurements by using custom developed cells are easy, reliable and flexible in terms of on-site field location. This technology can find its application in automotive workshops, aerospace workshops, industry and research departments where onsite field corrosion monitoring is a major challenge.

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