Investigating Pitting Corrosion of Stainless Steel and Aluminium Using Scanning Vibrating Electrode Techniques and Electrochemical Noise Measurement

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Abstract: In this study, the susceptibility of aluminium (Al) and steel SS304 to pitting corrosion in FeCl₃ and NaCl solutions with varying pH values were investigated using electrochemical noise (ECN) measurement and Scanning Vibrating Electrode Technique (SVET). Preliminary surface drop test of SS304 in FeCl₃ solution and total immersion test of Al in NaCl solution indicated higher pitting of samples at scratches simulated by scribe. Total immersion test on both SS304 and Al indicated that protecting a sample with epoxy-resin mold does not completely eliminate crevice attack. ECN tests results showed that Al suffered severe time-variant corrosion in FeCl₃ solution unlike SS304. The results of SVET tests conducted after 24h immersion indicated that the early pits captured on Al samples were more intense than the later ones. Higher corrosion rates were obtained for samples in NaCl solution of pH 2.89 compared to pH 7.21, indicating that pitting increased with acidity. SVET maps obtained corroborated Scanning Electron Microscopy (SEM) images of the sampled surfaces. However, though SVET was unable to capture pitting of SS304 in the solutions, SEM image revealed pitting activity on the sample in FeCl₃ solution.

Keywords: Scanning Vibrating Electrode Technique (SVET), Electrochemical Noise (ECN), Coefficient of Variation (CoV)

1. Introduction

Due to massive socio-economic loss and the attendant safety problems, corrosion has been extensively studied in an attempt to minimize its hazardous effects and thus possibly prevent its occurrence [1]. The corrosion engineer is thus embattled with the task of eliminating this economic wastage. It is evident that the earth crust is made up of 70% of marine environment, besides; the atmosphere sustains dense mass of water vapour and dissolved chemicals which are products of global industrialization. Most engineering materials are exposed to this harsh environment which tends to be corrosive [2]. Sea water contains a higher concentration of dissolved salts (predominantly NaCl) than any other natural liquid, except body fluid, salt lake and saline springs. Thus, it can be considered to be responsible for a vast percentage of the world corrosion wastage. Hence, many expensive equipment like the drill ships, ocean liners, offshore rigs, automobiles, underground/underwater gas and oil pipelines suffer massive attacks from this corrosion phenomenon and as such put the lives of crew onboard at risk [3].

To underscore the corrosion phenomenon, in 1975 a comprehensive study of metallic corrosion was carried out by the National Bureau of Standards (NBS) and Battelle memorial institute in response to a congressional directive and a published figure of seventy billion dollars ($70billion) was reported as the loss for metallic corrosion [4], an enormous amount of capital capable of salvaging the dwindling economy of some third world nations.

Metals are the most widely used engineering materials, of which steel and aluminum are among the top ranking in utility. Corrosion which is generally defined as the
degradation of a metal in reaction with a corrosive environment [1], [2], [3], does not spare these metals. There are basically different kinds of corrosion attacks, of which stainless steel and aluminum are greatly attacked by localised forms of corrosion (crevice and pitting corrosion) [1], [5]. These forms give no noticeable indication and many function as crack initiation sites which may propagate into material failure when subjected to mechanical working conditions. To investigate the susceptibility of these materials to pitting corrosion in an environment, weight loss index has been used, and this involves the resulting difference in weight of materials after exposure of the metal to the environment under service conditions for specific time periods. This procedure is slow, uneconomical and a poor representation for evaluating rates for metals in most slow corroding environment. Besides, very small localised mass losses are scarcely and inaccurately captured and quantified. Hence the need for the development of techniques which could capture and quantify these localised activity, and subsequently used to predict future events (accelerated testing).

Pitting can be a very destructive form of corrosion in engineering structures particularly if it causes equipment perforations, although minor pitting void of causing any noticeable perforation is most times tolerated in engineering practice for economic reasons. But were economic index cannot be matched to safety, need for exposing pits even at nanoscales then arise. In confirmation, corrosion pits were identified as crack origins on a test on aluminum in 3.5% wt NaCl solution (model sea water) [6]. It is not surprising that there has been an enormous research effort towards developing techniques to handle this problem of quickly assessing and predicting the susceptibility of a metal to pitting.

Some of these techniques include atomic force microscopy (AFM) [7], [8], scanning electrochemical microscopy (SECM) [9], scanning tunneling microscope (STM) [10], electrochemical noise (ECN) [11], [12], Electrochemical Impedance spectroscopy (EIS) [13], and Scanning reference electrode technique (SRET) [15]. The Scanning vibrating electrode technique (SVET) [16], adopted in this study belong to this SRET group, all of which function under the principle of their ability to pick up little changes in electrochemical activity within a localised pit and amplifying it [18], [19]. This study examines the susceptibility of two metals, stainless steel and Al, to pitting using SVET and the ECN box measurements in chloride environments to mimic seawater conditions.

2. Materials and Methods

2.1. Materials

2.1.1. Stainless Steel (SS304) and Aluminium Samples

Small blocks of aluminium (Al) (99.95% purity) and stainless steel (SS304) were cut to (15.4 x 15.4) mm cross-section. Electrical connectivity of samples was made by lead soldering a conductive copper wire to the rear side of the sample. For firm positioning of sample during SVET test, samples were embedded in an inert epoxy resin mould (Figure 1(a)), made from a mixture of 5:1 wt (epoxy resin: resin hardener) ratio, which was hardened for 20-24h before removal. This mould is to protect sample edges from possible crevice corrosion attack when in contact with the electrolyte during experiments.

The surface of the sample is treated to a fine finish by grinding on silicon carbide paper in the sequence of 240, 600, 1200, 2400 Grit size and polishing using diamond compound slurry in the sequence of 6µ to 1µ. This keeps the surface homogeneous and eliminates contours which could damage the fragile platinum probe tip, hence giving wrong / false output signals. It reduces occlusions that may pose as sources of pseudo localised corrosion.

![Figure 1. (a) Prepared Sample embedded in epoxy-resin mould; (b) Drop of 10% FeCl₃ on sample of finely scribed SS304.](image-url)

2.1.2. Electrolyte Solution

3% NaCl solution was prepared by dissolving 30g of laboratory sodium chloride salt in 1000ml of laboratory tap water introducing small drops of HCl (molar concentration unknown) into the solution to obtain the desired pH-value. 10% FeCl₃·6H₂O (≡ 6% FeCl₃) was prepared by dissolving 10g of hydrated FeCl₃·6H₂O in 90ml of lab water. Hydrolysis of 6% FeCl₃ produces an acidic solution of 1.2 – 1.3 pH value. The Fe³⁺ acts as the oxidizer by reduction to Fe²⁺ and the Cl⁻ is the pitting agent. A combination of this two qualities results in the aggressive environment for testing the resistance for the metals towards pitting [6].
2.2. Methods

2.2.1. Total Immersion Tests
- A 3-face Al sample, prepared to give three different surface profiles (plain, scribed and indented), was immersed in 3% NaCl and observed for 24h.
- Unscribed samples of SS304 and Al were totally immersed in 10% FeCl₃·6H₂O for 48h.
- Similarly, scribed samples (moppet scribing) of same samples were immersed in 10% FeCl₃·6H₂O (≡ 6% FeCl₃) and 3% NaCl.

2.2.2. Surface Drop Tests
- Two samples of SS304, (one with scribe, and one without scribe), were tested for corrosion by dropping 10% FeCl₃·6H₂O on the surface and left for 1h as in Figure 1(b).
- Also, similar experiments were done using Al sample in FeCl₃ and NaCl solution.

2.2.3. ECN Measurement

The Potentiostat (Gill AC, ACM Instruments, Cumbria – UK) with ancilliary equipment were connected as in Figure 2 and used for ECN current and potential-time measurements.
Scribed samples of Al and SS304, with edges or interface between epoxy and metal covered using an epoxy/beeswax mixed to 3:1 wt ratio, were connected to the working-electrode WE1 terminal of the Noise box (ACM Instruments, Cumbria-UK) from the rear part of the sample. Measurements were carried out against a non-corroding reference electrode, at rates of 0.2s per point for 3000 or 2000 sampling points, giving sampling periods of 600s or 400s, respectively. Solutions used include 10% FeCl₃ and 3% NaCl with pH values 7.21, 6.21, 5.01, 3.12 and 2.35. Readings were taken at 0h, 1h, 3h, 6h, 24h, 27h, 48h and 51h intervals after immersion.

2.2.4. SVET Measurement

(a) SVET Point-In-Space (PIS) Calibration
The Scanning Vibrating Electrode Technique (SVET) apparatus (SVP100, Uniscan Instruments, Buxton-UK) was connected-up with its ancillary equipment as shown in Figure 2 for the SVET experimental-runs.

(b) SVET Scans
To accommodate the Perspex box introduced, the set-up shown in Figure 3 was used for the experiment. Samples were earthed through the electrometer box in accordance with Original Equipment Manufacturers (OEM) provision for free corrosion test-earthed sample [20].
samples at pH 2.89 and 7.21, at 0h and 24h after immersion. The scan specifications used are the same for step-(a) above.

3. Results and Discussions

3.1. Total Immersion

The results of the immersion tests are shown in the Figure 4.

The SS304 immersed in FeCl₃ precipitated with heavy pitting at the crevices. The Al sample immersed in 10% FeCl₃ precipitated with the evolution of H₂ gas and general uniform corrosion were visually observed, no pits were found on scanning. White formations suspected to be corrosion products of Al₂O₃ were formed both at sample-epoxy interface indicating some form of crevice attack on all three profiles (Figure 4) and also at the near central location for the scribed surface. Bubbles were also observed but more with FeCl₃ solution. SS304 did not show any form of activity even after several weeks of immersion in NaCl solution.

Artificial scratches are used to simulate the mechanical or chemical breakdown of the passive oxide film, exposing the metal to the corrodent. The picture of Figure 4a shows the visible formation of pits on the scribed surface as compared with the other profiles, despite the crevice attack at the sample-epoxy interface. Similar experiment depicted in Figure 5 showed the heavy concentration of pits along the scribe as compared to the other parts of the metal surface. The change in colour from brown to green observed during this test was simply as a result of the reduction of ferric ions (Fe³⁺) to ferrous ions (Fe²⁺), indicating that there was actually corrosion which lead to the pits being formed. The embedded sample of Figure 4b indicates the formation of crevice attack on sample, which tends to cathodically protect the sample surface from possible pitting. This condition is not very favorable for the pitting experiment; as such this informed the need to protect the interface again with an epoxy-beeswax mix.

3.2. Surface Drop Test

On dropping the FeCl₃ solution on the scribed SS304 and Al samples, a few visible gas bubbles evolved from the Al sample after 300s and continued thereafter, no pits were found on microscopic observation and scanning using the optical scanner. The SS304 sample exhibited some localised formations or pits concentrated along the artificial scratch or scribe as shown in Figure 8 after 0.5h. In both cases there was a change of colour from brown FeCl₃ to green, which indicates the reduction of Fe₃⁺ into Fe₂⁺, due to corrosion or metal oxidation [5].

Scanning the drop on metal surface would have been a better way of getting SVET to pick pits along the scribe, but there exists a limitation, in the sense that the second probe tip is not in communication with the bulk solution as it is fitted above the main platinum tip, to sample the pH of the bulk solution [20]. This led to the use of the smaller Perspex box (Figure 2b) of volume 3cm³ compared to 4000 cm³ of SVET basin. Besides, the economy of size, the box can also accommodate very acidic electrolytes which evolve unhealthy and dangerous fumes. Communication of the second probe tip was not a limitation for the
modification as can be seen from Figure 2 and Figure 3.

3.3. Electrochemical Noise (ECN) Tests

3.3.1. Current Density - Time Plot

Since noise is random, an algebraic expression which defines the amplitude-time dependence of a particular source is impossible; and this means that noise is a non-deterministic process [21]. Fluctuations in corrosion potential are random; hence ECN are classified as non-solution combination are displayed as sample graphs in Figures 6, 7.

Figure 6. Current-time record for SS304 in 10% FeCl$_3$.6H$_2$O immediately after immersion.

Figure 7. Current-time record for SS304 in 10% FeCl$_3$.6H$_2$O 24h after immersion.

3.3.2. Statistical Evaluation

The statistical parameters of the current transients such as, Mean, standard deviation, Kurtosis k, and Skewness sK were evaluated based on the analysis tool pack add-in package in excel platform. From the results of the descriptive statistics of the current transients by ECN measurements of SS304 in 10% FeCl$_3$.6H$_2$O from 0-51h immersion time, the following were observed:

- A steady decline in mean $\mu$ and variance $\sigma^2$ within 0 - 24h of immersion;
- Erratic standard deviation $\sigma$, Kurtosis K and Skewness sK within 0 - 24h of immersion;
- CoV, and standard deviation $\sigma$ declined within 0 - 3h immersion before a steady rise.
- From the results of the descriptive statistics of the current transients by ECN measurements of Al in 10% FeCl$_3$.6H$_2$O from 0-51h immersion time, the following were observed:
  - CoV, Standard deviation $\sigma$ and Kurtosis K increased within 0 - 3h before decreasing;
  - Skewness sK increased within 0-6h before decreasing, but;
  - Mean $\mu$ was observed to be erratic.

From the K values obtained the dominant corrosion mechanism for Al in 10% FeCl$_3$.6H$_2$O were the intergranular corrosion (K > 6) for time t < 51h immersion and general / uniform corrosion (K < 6) after time t ≥ 51h immersion. However, for SS304 in 10% FeCl$_3$.6H$_2$O the dominant mechanism is intergranular (K > 6). This corroborates the work of Cappeln et. al [22] who performed ECN measurements on AISI347, 10CrMo910, 15Mo3, and X20CrMoV121 steels in molten NaCl-K$_2$SO$_4$ at 630°C. Different types of current noise were identified for pitting, intergranular and peeling corrosion. The corrosion mechanism was the so-called active corrosion (that is, the corrosion proceeds with no passivation due to the influence of Cl), characterized by the formation of volatile metal chlorides as a primary corrosion product. An empirical separation of general and intergranular corrosion using kurtosis K was done. It was found that average k > 6 indicated intergranular corrosion and average k < 6 indicated general corrosion. The response time for localized corrosion detection in in-plant monitoring was approximately 90min on this basis.

3.3.3. Change in Coefficient of Variation (CoV) with pH-Values for SS304 and Al in 3% NaCl Solution

| SS304 in 3% NaCl solution | pH of bulk solution | Coefficient of variation (CoV) |
|---------------------------|---------------------|-------------------------------|
| Time (h)                  | 7.21                | 6.21                          |
| 0                         | 3.7278              | -1.6809                       | 8.8435           | 2.8988  | 2.358  |
| 3                         | 3.3761              | 0.1566                        | 2.1654           | 1.2303  | 0.4913 |
| 6                         | 2.0800              | 0.1352                        | 0.6432           | 1.0260  | 0.8979 |
| 24                        | 1.7849              | -1.0158                       | 0.0923           | 0.2417  | 0.8459 |
| 27                        | -0.8557             | -0.2045                       | 2.7301           | 6.9464  | 0.1838 |

| Al in 3% NaCl solution    | pH of bulk solution | Coefficient of variation (CoV) |
|---------------------------|---------------------|-------------------------------|
| Time (h)                  | 7.21                | 6.21                          |
| 0                         | -12.3020            | 7.0471                        | 10.8339          | 1.0124  | 3.9322 |
| 3                         | -6.2100             | -2.0082                       | 3.2116           | 3.4187  | 0.4901 |
| 6                         | 4.7811              | 0.1336                        | 0.0751           | 0.4217  | 0.2268 |
| 24                        | 0.2046              | 0.1309                        | 0.3146           | 0.1332  | 4.9236 |

Evaluation of CoV variation with time at different pH-values of the bulk solution shown above indicate a steady decline in CoV with immersion time for pH-values of 7.21 and 5.01 for both SS304 and Al in 3% NaCl.

There was poor pitting characteristics of SS304 in pH 7.21 and pH 5.01. The pitting characteristics was fairly good at pH 6.21 during the early stages of immersion, although the physical observation of the current time records indicate that these transients are small compared to those of FeCl$_3$ which indicating a more corrosive environment for pitting. After
about 24h immersion, records show that better pitting of SS304 occurred at pH 3.09 followed by 5.01 which was indicated by the sharp rise in the CoV values as a first and simple evaluation. Those at pH 6.21 and 2.35 are fairly equal.

Similar observation were made for Al in 3% NaCl solution, at pH 7.21 where sharp rise in CoV in the first 6h was observed, followed by sudden decline while that of pH 3.12 and 2.35 showed some fine consistency after 24h immersion.

Based on this deductions from the ECN test, it became necessary to carry out the SVET test on 3% NaCl at pH 3.12, 2.89 and 7.21 (neutral solution).

3.4. SVET Test

3.4.1. Point-in-Space (PIS) Calibration Test

The line scan maps obtained from the PIS calibration tests are shown in Figure 8, a scan of 1 in 10 line scans per solution.

![Figure 8](image)

Figure 8. (a) Map showing 10 line scans for PIS calibration in 10%FeCl₃ solution; (b) Probe voltage signal versus displacement for a line scan from (a).

The values of the potential drop or probe peak signal as extracted from the line maps which are used for computation of the calibration factor (CF) for the different electrolytes. An initial probe reading before sudden drop = 174.6*10⁻⁹V, applied current = 0.025mA=25μA and applied current density = 79.46598856mA/cm² were utilized to obtain the CF in accordance with OEM recommendation (0-150mA/cm²) within which a linear relationship exist between the measured probe output and the local current density [9]. The CF for different solutions includes:

- Solution A (10% FeCl₃, H₂O at pH 1.30), 10 line scan Average Epis = -1.049mV, Eeff = -1.049mV; CF = 75727.719953mA/cm²/V;
- Solution B (3% NaCl at pH 7.21), 10 line scan Average Epis = -1.784mV, Eeff = -1.784mV; CF = 44531.868684mA/cm²/V;
- Solution C (3% NaCl at pH 2.91), 10 line scan Average Epis = -1.286mV, Eeff = -1.286mV; CF = 61765.550600mA/cm²/V;
- Solution D (3% NaCl at pH 2.41), 10 line scan Average Epis = -0.593mV, Eeff = -0.593mV; CF = 134003.426824mA/cm²/V;

The SVET test commenced with the performance of a PIS calibration using line scan rather than the conventional area scan. Probe peak output signal voltage values were averaged to reduce the error associated with inconsistent readings of probe signal. The PIS showed that for a particular solution, the signal increases with increasing pH values. This is principally due to the variation in the resistance posed by the positively and negatively charged ions as they move freely in solution.

3.4.2. SS304 in 10% FeCl₃·6H₂O - SVET Test Versus SEM Images of Sample Surfaces

i. SVET Scan

No visible pit formation was picked from the SVET equipment for 40 line-scans of SS304 from 0h to 96h immersion time in 10% FeCl₃·6H₂O as the line data showed mere straight lines.

ii. SEM Micrographs

At the end of the line scans carried out on the SS304 sample using SVET, no pit was picked up by the equipment, but the sample scanned using SEM revealed the formation of some pits along and outside the scribe as shown in Figure 9 and Figure 10.
Similarly, no pit was captured after area scans of the SS304 sample in 10% FeCl$_3$.6H$_2$O using SVET, but the sample scanned using SEM revealed the formation of some pits as shown in Figure 11.

**Figure 10.** SEM Micrograph showing pits on SS304 sample after SVET line scan in 10% FeCl$_3$.6H$_2$O (SVET scan line 2).

**Figure 11.** SEM Micrographs showing pits on SS304 surface after a SVET area scan in 10% FeCl$_3$.6H$_2$O.

### 3.4.3. Al in 3% NaCl

#### (i) SVET Scan

The SVET was able to pick-up and map the pits as they form, grow and repassivate on Al sample in 3% NaCl for pH 7.21 and pH 2.89 (Figure 12). Only the first formations are shown.

**Figure 12.** (a) Map showing 20 line scans for Al in 3% NaCl at pH 2.89 after 24h immersion; (b) line data for line 16.

#### (ii) SEM Micrographs

At the end of the line and area scan carried out on the Al sample using SVET, the sample scanned using SEM revealed the formation of some pits as shown in Figure 13.

**Figure 13.** SEM image of Al in 3% NaCl solution at pH 2.89 showing distribution of corrosion products along scribe and possible occlusion of pits.
3.5. Computation of Corrosion Rate Due to Pitting

Corrosion rates were calculated from equation (1) and plotted against time over the entire scanning and presented as Figure 14 for Al but that for SS304 is not presented because of the null values obtained.

\[
E_{\text{corr}} = |E_{\text{corr}} - E_{\text{pitting}}| \quad (1)
\]

![Figure 14. Corrosion rate Versus time for Al and SS304, in 3% NaCl (pH 2.89 and 7.21) using SVET after 24h immersion.](image)

The ability of a metal to pit in an environment has been regarded as a probabilistic issue [23]. It is then necessary to define the best probabilistic conditions upon which pitting may occur before any scanning. Scanning technique adopted is important because of the repassivation of pits which is also probabilistic in itself, picking up pits formation and growth before death. There is every tendency that probe gets to pit probabilistic in itself, picking up pits formation and growth before formation or while pit is repassivated [24]. In an attempt to increase the chance of capturing a pit as they form, in the total immersion test it was found that the best surface profile for a sample being used in this kind of experiment is the one with a scribe on it, which is in line with the postulate of Pessall and Liu [25].

The ECN transients consist of uni and bi-directional transients, that is, anodic (positive) and cathodic (negative). Transients depict the nucleation, temporary growth and repassivation of small pits. Negative transients indicate the breakdown of passive films which are turned into active metal dissolution [26]. On observation of Figure 12, the cathodic current transients of SS304 reached values as small as \(-3\times10^{-8}\) mA/cm\(^2\) immediately after immersion, with also more spikes as compared with that after 24h immersion, where a single transient of comparatively larger value of \(-2.5\times10^{-6}\) mA/cm\(^2\) was obtained. This suggests that the SS304 pits with time in this solution as bigger transients are formed, that is larger metal dissolution. The current time record of Al (plot not included) showed numerous spikes indicating that the electrolyte is more corrosive to Al than SS304; although transients indicate pit formations but the randomness suggests that general kind of corrosion is more prominent than pitting. After 24h immersion the record describes the same condition of corrosion activity although more general in this case than in the one encountered immediately after immersion as can be shown by the cluster of smaller transients.

A study using the CoV, indicate that SS304 has a better pitting tendency than Al in FeCl\(_3\) solution, the steep slope observed in the value of the CoV for Al 6h immersion could be as a result of general form of corrosion rather than pitting. This repeated after 48h; in essence Al suffers from severe general corrosion and pitting in FeCl\(_3\) solution unlike the SS304 with profile indicating gradual pitting process across the immersion period.

The test on SS304 in pH 2.89 within the short period yielded no result with SVET. Also, microscope observation revealed no pits. This explains the resistance to natural pitting by SS304 in sea water condition and the inability for the CI\(^+\) present to break the passive film formed by the Cr element in the metal. However several tests performed on SS304, were able to get it to pit by the passage and withdrawal of an anodic current through the sample [24], [27]. On test with ferric chloride solution, SVET could not also pick and map any pits, but the SEM image in this instance revealed the formation of pits as in Figures 12, 13. The inability of SVET to pick and map these pits could be due to earlier arrival of probe at pitting site before formation or late arrival after pit repassivation.

Unlike the SVET scan on SS304, scan on pure Al in 3% NaCl at pH 7.21 and 2.89, pits were correspondingly captured and mapped. We noticed that the early captured formations were more intense and gradually die away as the scan proceeds. The reason could be the possible repassivation of these pits or the subsequent closure of pits with or without the dynamic assistance of the probe tip as it scans past the pit, although some of the line scan data indicated pit growth in Figure 14. Another important factor to be considered is the probe waiting time before another line scan run over pit. The SEM image for the Al could only show the corrosion products as they are distributed over the surface, with heavier concentrations suspected to be around pit (s) mouth. The result of the computation done for SS304 and pure Al, from the SVET maps suggest that pure Al is more prone to pitting attack than the SS304. It is also more severely attacked in 3%NaCl solutions with lower pH values or more acidic as can be seen from the computed values of corrosion rate for line 16 which is assumed to have occurred at approximately the same time.

The single line or point analysis could possibly give a wrong indication of the susceptibility, hence a need for the overall study of the entire map over the time of run, will give a more representative result of the entire process and can be used as a better predictive model. The graph of Figure 14 developed for this study, apart from the instantaneous values of the pitting corrosion rate, gives us an overall profile of the formations. It suggest that the corrosion rate varied with over time. It is higher for pure Al in sea water of pH 2.89 than that of neutral sea water. The gradual decline as seen from the graph could be as a result of the pits re-passivation, closure of pits by occluded corrosion products by the probe while scanning or mass transport of the electrolyte during activity. It should be noted that the plot of SS304 showed no variation as no sudden voltage drops indicative of pitting being recorded by the SVET.

Importantly, the use of effective value of sudden voltage
drops, is necessitated by the fact that we are only considering the transients formed as a result of pitting activity (cathodic or anodic), the SVET setting could give an initial non zero reading (E\text{init}) which signifies a false corrosion activity where there is none. The value of voltage drop caused by the sudden anodic dissolution is the steep voltage produced which is termed the effective voltage (E\text{eff}) in this work. This has the property of singling or extracting out the corrosion due to pitting alone, void of other perturbations.

4. Conclusion

- In-situ evaluation of pitting corrosion by ECN and SVET increases the chance of capturing pits formation, growth and repassivation in the corrosion experiments.
- Line scans along a scribe has a greater chance of capturing pits formation more than area scan. Also, the use of reduced electrolyte basin enhanced measurements better flexibility; in terms of the electrolytes, number of requisite experimental runs and probe proximity to the activity sites. Crevice attacks at interfaces could not be eliminated completely; necessitating the covering of interfaces with colophony resin beeswax to reduce crevice attack on sample before immersion into electrolyte.
- The coefficient of variation (CoV) can be used as a screen (sorting) test for pitting susceptibility preparatory to SVET test. The consistence of pitting with respect to CoV was noticed for pH value around 3.09/2.35 after 24h immersion.
- SVET has proven a veritable technique for capturing pitting and pitting-susceptibility for Al though it could not capture the pit formation in SS304. This is attributable to (a) inappropriate and inadequate experimental controls of the system electrochemistry, (b) short time of pit formations and repassivation compared to the experiment time.
- Gas formation and heavy deposit of corrosion product reveals a high overall continuous corrosion reaction in Al. This intense dynamic environment could be a facilitator for the proper functionality of the SVET equipment.
- The signal output of the SVET over a period of time could be used as a measure of susceptibility of a metal to pitting corrosion in a particular environment. It has been shown that Al is prone to heavier pitting in a more acidic environment, and the pH of a corrosive environment drives the pitting susceptibilities of the metals exposed in that environment.
- SVET maps for Al with their corresponding sudden voltage drops, corrosion rate values computed can be used to study the mechanism of H_2 evolution in the pitting process - reduction in pitting corrosion rate denotes reduction in anodic dissolution hence H_2 evolution.
- The effective voltage drop was able to quantify the dissolution associated with only the pitting action rather than a combination of all the parameters that contribute to potential drop.
- Absolute values of transients rather than the number of transients explain better pitting susceptibility.
- ECN and SVET have the capability of capturing and mapping pits in time domains but the former does not give geometric position of pit formation. ECN has a greater chance of capturing pits than SVET but lacks proper quantification of the pits.
- SS304 pits more than Al in FeCl_3, with Al exhibiting general/ uniform corrosion attack.
- Pit formation in SS304 is more difficult to capture than that of Al. Al pits more than SS304 in salt water, where its pitting susceptibility is more in an acidic medium compared to that of a neutral medium.

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