# MPC Journal

Copy of e-mail Notification

MPC Published by ASTM

Dear Author,

YOUR PAGE PROOF IS AVAILABLE IN PDF FORMAT; please refer to this URL address http://115.111.50.156/jw/AuthorProofLogin.aspx?pwd=5ddad2de8d9b&CA=AT

Login: your e-mail address Password: 5ddad2de8d9b

The site contains 1 file. You will need to have Adobe Acrobat Reader software to read these files. This is free software and is available for user downloading at http://www.adobe.com/products/acrobat/readstep.html.

This file contains: Adobe Acrobat Users - NOTES tool sheet A copy of your page proofs for your article

Please read the page proofs carefully and:

1) indicate changes or corrections using e-annotation;

- 2) answer all queries;
- 3) proofread any tables and equations carefully;
- 4) check that any special characters have translated correctly.

Special Notes:

Your edits must be returned (uploaded or emailed) within 48 hours of receiving page proofs to review. Please notify Denise Stanley (denise.stanley@cenveo.com) immediately if you anticipate a delay in returning your edits.

Your Login and Password are valid for a limited time. Your prompt attention to and return of page proofs will help expedite publication of your work. Thank you for your cooperation.

If you have any questions regarding your article, please contact me. PLEASE ALWAYS INCLUDE YOUR ARTICLE NO. (MPC20120016.R1) WITH ALL CORRESPONDENCE.

Posting Author Papers on Website Policy—Author's Rights as Stipulated in the Accepted Electronic Agreement during Paper Submission:

"The right to post the pre-print version of the Work on your website or your employer's website with reference to the publication by ASTM as the copyright holder. This preprint will be sent to you by the copyeditor. This version

# **MPC** Journal

Copy of e-mail Notification

does not include the final edits. Such preprints may be posted as electronic files on the Author's own website for personal or professional use, or on the Author's internal university or corporate networks/intranet, or secure external website at the Author's institution, but not for commercial sale or for any systematic external distribution by a third party (eg: a listserver or database connected to a public access server). Prior to publication, the Author must include the following notice on the preprint: "This is a preprint of an article accepted for publication in (Journal Title Copyright @ (year) (copyright ASTM International, West Conshohocken, PA)". After publication of the Work by ASTM International, the preprint notice should be amended to read as follows: "This is a preprint of an article published in (include the complete citation information for the final version of the Work as published in the print edition of the Journal)" and should provide an electronic link to the Journal's WWW site, located at the following ASTM URL: http://www.astm.org. The author agrees not to update the preprint or replace it with the published version of the Work."

Sincerely,

Denise Stanley, Journal Production Manager E-mail: Denise.Stanley@cenveo.com



# REPRINTS ORDER FORM

For reprints of articles and/or papers from ASTM Journals, Selected Technical Papers (STPs), book chapters and *Standardization News.* 

Note: Authors receive a 25% discount

			•	-		
Reprinted article a	and cover with typeset t	itle of article and a	author's name			
Reprinted article, r	no cover					
Cover: (check one)						
Page length of article:			Number of copies: (25 copies minimum)			
Paper Title: Publication:	month, and volume number; S	STP number; or stock n	umber)			
Date:	Pnone:	err	iali:	Fax:		
				E		
			State:	Zıp:		
Address:			04-4-4	7:		
Organization:						
<b>B</b> ILL <b>To:</b> Name:						
Country:						
City:			State:	Zip:		
Address:						
Organization:						
<b>S</b> нıр <b>To:</b> Name:						

QUESTIONS ABOUT YOUR ORDER? Call Customer Relations at 610/832-9585, weekdays 8:30am-4:30pm ET



# **REPRINTS PRICE SCALE\***

# COLOR REPRINTS

Copies Pages	100	200	500	1000	2000
2	\$1,102	\$1,323	\$1,543	\$1,764	\$1,984
3-4	\$2,116	\$2,221	\$2,326	\$2,425	\$2,824
5-8	\$2,168	\$2,278	\$2,394	\$2,499	\$3,024
9-12	\$3,307	\$3,470	\$3,638	\$3,801	\$4,126

# **BLACK & WHITE REPRINTS**

Copies Pages	25	50	100	200	300	500	1000	2000
1-4	\$136	\$152	\$168	\$184	\$199	\$226	\$310	\$472
5-8	\$195	\$226	\$257	\$278	\$299	\$352	\$499	\$709
9-12	\$225	\$257	\$289	\$336	\$357	\$425	\$577	\$787
13-16	\$263	\$294	\$325	\$394	\$430	\$520	\$751	\$1,118
17-20	\$278	\$336	\$394	\$472	\$514	\$635	\$887	\$1,307
21-24	\$301	\$373	\$446	\$535	\$583	\$724	\$1,003	\$1,475
25-28	\$331	\$404	\$478	\$583	\$646	\$808	\$1,165	\$1,795
29-32	\$383	\$441	\$499	\$651	\$724	\$913	\$1,323	\$2,058
Cover (optional)	\$100	\$115	\$136	\$157	\$178	\$220	\$373	\$598

\* Prices are subject to change without notice.

ASTM International Customer Relations department will finalize your reprint quote including freight charges. ASTM International will contact you with the quote price.

# AUTHOR QUERY FORM

	Journal: Mater. Perform. Character	Please provide your responses and any correc- tions by annotating this PDF and uploading it to ASTM's eProof website as detailed in the Welcome email.
INTERNATIONAL	Article Number:	
Standards Worldwide	MPC20120016	

Dear Author,

Below are the queries associated with your article; please answer all of these queries before sending the proof back to Cenveo. Author please indicate the correct color processing option from the list below:

- 1. Author, please confirm Figure number(s) that should appear as color in print. Please know that any associated mandatory fees will apply for figures printed in color.
- 2. Author, please confirm Figure number(s) that should appear as color online only, there will be no fees applied.
- 3. Author, your paper currently does not include any color figures for online or print. If color is needed please indicate which figures it should be applied to and whether it is color in print or online.

Location in article	Query / Remark: click on the Q link to navigate to the appropriate spot in the proof. There, insert your comments as a PDF annotation.
AQ1	Please advise whether reference 32 was meant here instead of 22.
AQ2	Please advise – reference 54 citation deleted here as there is no reference 54 in the list.
AQ3	For reference 1, please provide dates and location of meeting.
AQ4	For reference 20, please check standard number, year, and name of organization.
AQ5	Please provide all authors for references 38 and 49.
AQ6	For the website references, please check that the date the sites were last accessed are correct. If
	not, please provide correct dates. Please provide the date for all website references.

Thank you for your assistance.

Materials Performance and Characterization, Vol. 2, No. 1 Paper ID MPC20120016 www.astm.org

Adil Saeed,<sup>1</sup> Zulfiqar A. Khan,<sup>2</sup> and Eliza L. Montgomery<sup>3</sup>

# Corrosion Damage Analysis and Material Characterization of Sherman and Centaur— The Historic Military Tanks

**REFERENCE:** Saeed, Adil, Khan, Zulfiqar A., and Montgomery, Eliza L., "Corrosion Damage Analysis and Material Characterization of Sherman and Centaur—The Historic Military Tanks," *Materials Performance and Characterization*, Vol. 2, No. 1, 2013, pp. 1–16, doi:10.1520/MPC20120016. ISSN 2165-3992.

**ABSTRACT:** A study of corrosion damage and material characterization of two historic military tanks, the Sherman and Centaur is reported. Experiments were conducted to analyse surface corrosion and corrosion propagation from surface to sub-surface. Significant surface corrosion was found, and this phenomenon was further facilitated by delamination failure mechanisms. Corrosion depth for the Sherman was approximately 110  $\mu$ m, where sulphide inclusions were detected in the sub-surface analysis. The Centaur's analysis showed corrosion pits at 100  $\mu$ m depth. These pits possess random geometrical configurations with evidence of sulphur, sodium, and calcium.

**KEYWORDS:** corrosion, material characterization, museum environment, military vehicles, sulphide inclusions

# Introduction

The Tank Museum in Bovington, United Kingdom has one of the largest collections of military 11 tanks from the First and Second World Wars and from post-war conflicts. These historic vehicles 12 are at risk of aging because of corrosion. Historic military vehicles and other large mechanical mu-13 seum artefacts are key entities, which provide a wealth of information and insight into past design 14 processes, design methods, materials, and manufacturing techniques. Historic vehicles in the Tank 15 Museum at Bovington are kept in two distinct environments, indoors (controlled) and outdoors 16 (uncontrolled). Some of the vehicles run occasionally after being kept in the controlled environ-17 ment in the museum. This temporary change in environment of the vehicles combined with other 18 operating factors poses significant risk of failures. 19

The indoors environment refers to the controlled environment inside the Tank Museum. The 20 lighting source is natural daylight combined with fluorescent lighting, which emits a low amount 21 of ultraviolet (UV) light. The temperature is controlled in winter only and humidity is not con-22 trolled during any season. The outdoor environment refers to the uncontrolled environment that 23 exists naturally in Bovington, United Kingdom. Bovington is in the county of Dorset on the south-24 western coast of England and is located approximately 9 km north of the English Channel. 25

<sup>1</sup>Sustainable Design Research Centre, School of Design, Engineering and Computing, Bournemouth Univ. Talbot Campus, Poole, Dorset BH12 5BB, United Kingdom (Corresponding author), e-mail: asaeed@bournemouth.ac.uk

<sup>3</sup>Ph.D., Materials Testing and Corrosion Control Branch, National Aeronautics and Space Administration, Kennedy Space Center, FL 32899, United States of America, e-mail: eliza.l.montgomery@nasa.gov

Copyright © 2013 by ASTM International, 100 Barr Harbor Drive, PO Box C700, West Conshohocken, PA 19428-2959.

1

2

3

4

5

6

7

8

9

Manuscript received June 11, 2012; accepted for publication December 20, 2012; published online xx xx xxxx.

<sup>&</sup>lt;sup>2</sup>Ph.D., Sustainable Design Research Centre, School of Design, Engineering and Computing, Bournemouth Univ. Talbot Campus, Poole, Dorset BH12 5BB, United Kingdom, e-mail: zkhan@bournemouth.ac.uk

58

59

### PROOF COPY [MPC20120016]

2

#### MATERIALS PERFORMANCE AND CHARACTERIZATION

Structural deterioration through corrosion damage in large metal structures such as ships, aircraft, and bridges, is a colossal predicament [1,2], and large metal museum artefacts are no exception to this. The dilemma of corrosion damage in historic military tanks in the Tank Museum can be classified as one of the major perils to their structural integrity. 29

This paper presents results of corrosion on the surface, its propagation from surface to sub-30 surface, and the subsequent damage to the structural integrity of the Sherman M4A1 and/or the 31 Centaur A27L. Both the Sherman and Centaur participated in the Second World War and post-32 war conflicts, encountered opponents' explosive attacks, and operated in a wide variety of terrains 33 ranging from desert to Eastern Europe and therefore accumulated various types of structural dam-34 age during their service lives. Now kept in a museum, these vehicles still go through structural deg-35 radation because of corrosion. Sustainable methods are required to diminish corrosion and 36 corrosion-related problems in these historic vehicles to preserve them for the current and coming 37 generations keeping their cultural biography intact. 38

This is the first research conducted to evaluate corrosion on historic military tanks kept in the <sup>39</sup> Tank Museum at Bovington. There is insufficient data on the tanks concerning their past exposure <sup>40</sup> during the wars, their locations, and their operating environments, and it is beyond the scope of <sup>41</sup> this research work to track their service history. However, war-related damage is significant on <sup>42</sup> many of the tanks and the majority of the tanks display failures in the protective coatings. <sup>43</sup>

It is important to slow down the process of structural degradation caused by corrosion failures 44 or stop it wherever possible. It is, however, of critical importance that such measures are sustain-45 able in terms of not compromising the historic and cultural heritage of these vehicles by excessive 46 repair, replacements, and other maintenance techniques such as coatings. The sustainable conser-47 vation of these artefacts would include full understanding of the critical parameters that are respon-48 sible for initiating and accelerated propagation of corrosion. Linking these critical attributes to 49 controlled environment facilities to optimise their design for achieving these goals economically is 50 the objective of the present research. 51

It is, however, not possible to achieve controlled environments for all 300 plus vehicles; therefore, alternative methodologies can be devised for vehicles kept in uncontrolled facilities to slow down the process of structural degradation to minimise historic loss.

Findings from the undertaken research will enable the design of a sustainable framework of conservation of these vehicles in both controlled and uncontrolled environments. 56

# **Experimental Methodology**

Sample Selection

### Sherman M4A1

Sherman M4A1 was designed by Lima Locomotives in 1941–1942 in the United States and was utilised by the British Army in the Second World War [3,4]. This Sherman was provided by Royal Armoured Corps Gunnery School, Lulworth Camp, United Kingdom in 1955 to the museum. No images were taken with a view to identify specific areas of corrosion at that time or afterwards. This tank operated occasionally and is now inside the museum in a controlled environment. Inside the museum, the temperature is kept under control in winter only, and no control of relative humidity is exercised in any weather. The Sherman experiences medium levels of corrosion.

Sample was collected from the hull (turret area) of the Sherman. The exposed surface of the 67 sample had a coating. However, the inner side was corroding and any previously applied coating 68 had completely failed. 69

J\_ID: MPC DOI: 10.1520/MPC20120016 Date: 5-January-13

PROOF COPY [MPC20120016]

#### SAEED ET AL. ON CORROSION DAMAGE ANALYSIS

Total Pages: 16

#### Centaur A27L

This tank was designed by the Leyland Motors in the United Kingdom in 1942-1945 and served in 71 the Second World War [3,4]. This tank was given to the museum by the Ministry of Defence 72 (MOD) in 1994. Again, there was no corrosion study conducted at the time of accession. Since 73 1994, this vehicle is stationary outside the museum in an uncontrolled environment. The Centaur 74 is stationed outside the museum and is exposed to environmental conditions such as direct UV 75 light from the sun, fluctuations in temperature and humidity, and varying times of wetness. The 76 Centaur is undergoing extensive structural degradation. Figure 1 show the Centaur's armoured 77 skirt, extensively affected by corrosion. 78

A sample was collected from the armoured skirt of the Centaur. The sample was coated on the 79 exposed surface, whereas the inner side (towards the tracks) was corroding with little evidence of 80 previously applied coating.

#### Samples' Materials Characterisation

Both samples were sectioned and x-ray fluorescence (XRF) was conducted at the cross section 83 (newly exposed surface) of the samples to obtain the materials composition. XRF results illustrated 84 that Sherman composition was approximately matching AISI 1000, and the Centaur was matching 85 AISI 4000 series steel [5,6]. These materials were processed before/during the Second World War. 86



FIG. 1—Corrosion in Centaur's armoured skirt to a significant level.

ID: veeraragavanb Time: 19:00 I Path: //xinchnasjn/ASTM/3B2/MPC#/Vol00201/120037/APPFile/AT-MPC#120037

3

70

82

J\_ID: MPC DOI: 10.1520/MPC20120016 Date: 5-January-13

#### Stage: Page: 4 Total Pages: 16

# PROOF COPY [MPC20120016]

4

#### MATERIALS PERFORMANCE AND CHARACTERIZATION

# Sample Preparation

For sub-surface analysis, samples were sectioned, and then mounted in bakelite. Polishing was performed using silicon carbide abrasive papers. Samples were conditioned with a diamond suspension spray of 3  $\mu$ m, rinsed with water, and then dried. No chemical treatment on samples was performed.

# Sample Analysis

92

105

106

87

Scanning electron microscopy (SEM) and energy dispersive x-ray spectroscopy (EDS) were performed under high vacuum using field emission gun scanning electron microscopy (FEGSEM) at 20 kV. Samples were conductive and therefore required no carbon coating for the analysis. EDS calibration was conducted with Cu standard, a normal practice for EDS calibration. SEM examination obtained micro-structural characteristics of the samples in high resolution. Through EDS, a qualitative and quantitative determination of the elements was obtained and evaluated from the energy spectrum versus relative counts of the detected x rays [7,8].

First, SEM and EDS analyses were performed on the corroded surfaces of each tank, which had 100 no effective protection (paints/coatings). Second, analysis was performed at the cross section of the 101 samples, where corrosion propagation from surface to sub-surface, corrosion pitting (a form of 102 localised corrosion confined to a small area, which results in cavities), and cracking were all 103 prevalent.

# **Results and Discussion**

#### Sherman M4A1

On the corroding surface of the Sherman in total six elements, Iron (Fe), Calcium (Ca), Silicon 107 (Si), Aluminium (Al), Sodium (Na), and Oxygen (O), were identified at three spectra. EDS results 108 of these three spectra are provide in Table 1. In spectrum 1, high proportion of O (46.01 wt. %) was 109 recorded. Fe (44.22 wt. %) was recorded lowest at spectrum 2.

X-ray fluorescence (XRF) results at the cross section of the sample illustrated Fe (98.30 wt. %), 111 Mn (0.63 wt. %), and Si (0.65 wt %) as the main elements [5]. Si originates from alloying elements. 112 The detection of Al on the corroding surface could be attributed to the Al-based paint protection 113 applied at some stage. Ca and Na in spectra 1 and 2 are surface contaminants and originate from 114 previous/current atmospheres. 115

Further investigations were conducted to obtain corrosion propagation from surface to subsurface, shown in Fig. 2. Significant corrosion was recorded that has resulted in delamination of 117 corrosion residues from the surfaces. The surface was affected to a depth of almost 110  $\mu$ m into the 118 bulk metal, where sub-surface cracks were also identified. Corrosion was investigated at the cross 119 section of the corroding edge, labelled as spectrum 1, shown in Fig. 2(*a*), and at the bulk metal 120

				ē ;					
Sample: Sherman-corroding Surface (Results in Weight %) Processing Option: All Elements Analysed/Normalised									
1	46.01	0.00	3.02	4.62	1.19	45.13			
2	41.55	1.05	6.55	5.67	0.93	44.22			
3	39.98	0.00	1.46	1.16	0.00	57.38			

TABLE 1—Sherman EDS point analysis of corroding surface.

ID: veeraragavanb Time: 19:00 I Path: //xinchnasjn/ASTM/3B2/MPC#/Vol00201/120037/APPFile/AT-MPC#120037

#### PROOF COPY [MPC20120016]

#### SAEED ET AL. ON CORROSION DAMAGE ANALYSIS



FIG. 2—Sherman cross section—rectangular area shows EDS analysis area (approximately 820  $\mu$ m<sup>2</sup>): (a) CP to 110  $\mu$ m, and materials' delamination, and (b) sub-surface crack propagation.

where no corrosion was occurring, labelled as spectrum 2, shown in Fig. 2(*b*). In spectrum 1, Fe 121 (60.84 wt. %), Mn (0.45 wt. %), Ca (0.40 wt. %), and O (38.30 wt. %) were detected. The presence of 122 a high amount of oxygen and traces of Ca show that drastic corrosion has occurred. Oxygen is the 123 signature for iron oxide (FeO) corrosion product formation, and Ca is known to be present in 124 atmospheric salt deposits as well as sea salt [9,10]. At spectrum 2, where no corrosion has propa-125 gated yet, only Fe (99.41 wt. %) and Mn (0.58 wt. %) were detected.

Figure 2 demonstrates corrosion on the Sherman resulting in cracks and delamination of the 127 material. Such surfaces, when exposed to stresses, will become one possible cause of structural failure [11], these are referred to as both mechanical (stresses because of dynamic and/or static load-129 ing) and corrosion-induced (stress corrosion cracking) stresses.

Sulphide inclusions were identified in the cross section from the Sherman, shown in Fig. 3. 131 Spectrum 1 results showed Fe only 5.31 wt. % with a high amount of sulphur (S) (35.25 wt. %), 132 other elements found at spectrum 1 were Mn (57.71 wt. %) and Fluorine (F) (1.70 wt. %). Spectrum 133 2, which was conducted at the metal, demonstrated Fe (99.24 wt. %) and Mn (0.75 wt. %). Spec-134 trum 3 was conducted at the second inclusion and showed Fe (24.14 wt. %) with S (26.37 wt. %), 135 Mn (46.95 wt. %), and F (2.52 wt. %). F can be classified as a surface contaminant that has possibly come into contact with the tank's surfaces during the war. F belongs to a halogen group of elements and is highly poisonous yellow gaseous element. Elemental F and F ion are highly toxic and can even react with inert noble gases such as krypton and xenon [12].

Sulphur in the Sherman originates from the steel-making process and Mn is an alloying 140 agent [13], being added to steel to avoid FeS formation and to form MnS for the purpose of 141

#### 6 MATERIALS PERFORMANCE AND CHARACTERIZATION



FIG. 3—Sherman cross section—sulphide inclusions and EDS (point analysis) at 3 spectra.

segregating S. MnS has a higher melting point and is chemically stable, and also during hot rolling 142 prevents the formation of FeS along the grain boundaries [14,15]. MnS inclusions tend to provide 143 better machining, increased wear resistance of the components, and reduced costs during product 144 manufacturing [16]. However, MnS inclusions cause the initiation of corrosive pits in bulk metal, 145 such as carbon steel and low carbon steel [17]. In addition, combined with corrosive environments, 146 these inclusions result in accelerated pit formation and deterioration of the overall corrosion resist-147 ance of the metals [17–19].

The temperature in the museum is kept between  $18-25^{\circ}$ C in winter. The buildings where the 149 tanks are housed are not humidity controlled apart from being enclosed and protected from direct 150 atmospheric moisture. Recording of temperatures and relative humidity (RH) started in October 151 2011 in five locations and another five locations were included in early 2012 in the museum. The 152 recent monitoring, over a period of 4 months, indicate temperature fluctuations from  $17^{\circ}$ C to  $20^{\circ}$ C 153 and daily range of RH was reported to be from a low of 40 % to a high of 80 %. All of the 10 154 locations detected conditions where RH reached above 70 % at various intervals of time during a 155 24 h period.

There is plenty of evidence of condensation inside some areas in the museum where water regularly runs off the tank's surfaces. Figure 4 illustrates the temperature and RH recorded in the WW2 158



FIG. 4—Temperature (a), and RH (b) graphs in WW2 Hall where Sherman is kept.

192

hall where the Sherman is kept. In first reading during 25/2010/2011-16/2011/2011, highest and 159 lowest temperatures recorded were  $\sim 23^{\circ}$ C and  $\sim 16.5^{\circ}$ C with highest RH being  $\sim 80$  % and the 160 lowest being  $\sim 53$  %. In the second reading (Dec. 1, 2011– Jan. 15, 2012), the highest and lowest 161 temperatures were  $\sim 18.50^{\circ}$ C and  $\sim 14^{\circ}$ C, respectively, as shown in Fig. 4. RH recorded during the 162 same duration was just above 65 % RH as the highest and  $\sim 30$  % RH the lowest. 163

Corrosion is hugely influenced by the environment; in turn, this is attributed to the atmospheric 164 pollutants [13]. Condensation is known to occur when temperatures exceed 0°C with 80 % RH 165 [20], thus leading to a longer time of wetness (TOW) and atmospheric corrosion [21,22]. There-166 fore, during these temperature and RH combinations some degree of corrosion is expected in tanks 167 in the museum. In addition, the critical RH will also decrease significantly in the presence of air-168 borne as well as surface pollutants [23].

Under wet/damp conditions ( $T \ge 0$  and RH > 80 %) and when the air is saturated with water 170 vapour, dissolution of MnS inclusions are possible leading to the formation of corrosion pits. Sulphide inclusions identified in the bulk metal of the Sherman are likely points of weakness along the 172 grain boundaries and will promote pitting corrosion; dissolution is expected at the edge of the 173 inclusions through the formation of rust of sulphur over the inclusions and at the surrounding 174 bulk steel. Changes in the shapes of the inclusions and the formation of microscopic cracks are also likely. The metal matrix is exposed when MnS dissolution occurs; the consequent dissolution products, i.e., elemental sulphur, thio-sulphate ion ( $S_2O_3^{2-}$ ), hydrogen sulphide (H<sub>2</sub>S), and hydrogen 177 sulphide ion (HS<sup>-</sup>) create a corrosive environment, encouraging pit propagation, and faster corrosion will also result [14,19,22,24–26]. Steel, stainless steel, and copper are all known to suffer from 179 stress corrosion cracking and sulphide stress cracking because of free sulphides [27].

The identification of Na, Ca, and F on the surfaces is not favourable, whether accumulated from 181 the air or the places where the tanks operated in the past. These will form corrosive electrolytes on 182 the bare surfaces of the Sherman, consequently accelerating corrosion. Rusting will form on the 183 surfaces from the precipitation of ferric oxide (Fe<sub>2</sub>O<sub>3</sub>). This rust is composed of the first oxyhydroxide lepidocrocite ( $\gamma$ -FeOOH) and goethite ( $\alpha$ -FeOOH), which consists of pores, cracks, and 185 have poor adherence. Such rust layers formed on steel do not have the ability to form a protection 186 against the corrosive products reaching the steel substrate and therefore will not effectively restrict 187 the diffusion process [13,28–31].

Under current environmental conditions, when the RH reaches above 80 % and in the presence 189 of surface contaminants, it is likely that corrosion in the Sherman will prevail resulting in structural 190 failures.

#### Centaur A27L

Ultrasonic scanning to record the material loss because of corrosion has already been reported for 193 the Centaur [5]. Table 2 illustrates results from ultrasonic scanning for the maximum and minimum remaining thicknesses at five points each. The difference between maximum and minimum 195 points provides relative measure of corrosion as less thickness indicates more corrosion and vice 196

TABLE 2—Centaur ultrasonic scanning results of the five lowest and highest remaining thicknesses.

Centaur Sample's Dimension Length: 240 mm, Width: 96 mm										
Remaining thickness in mm	2.30	3.60	3.80	3.90	4.0	6.80	6.90	7.0	9.20	9.80
No. of occurrences of the remaining thicknesses	1	2	5	3	2	374	410	201	2	1

#### 8 MATERIALS PERFORMANCE AND CHARACTERIZATION



FIG. 5—Centaur armoured skirt sample—corroding surface and EDS spectra locations.

versa. At the time of manufacture, the maximum and minimum armour thicknesses for the Centaur were 76 mm and 20 mm, respectively [3,4]. In the collected sample from the armoured skirt the highest thickness was recorded as 9.80 mm. No higher value was detected after repeated scanning. Values in Table 2 provide an understating of the armoured decay as the thickness has been reduced significantly. Material loss of 7.50 mm has been recorded between the maximum and minimum remaining thicknesses of 9.80 mm and 2.30 mm, respectively.

The corroding surface was measured at three different spectra 1, 2, and 3 as shown in Fig. 5. 203 The experimental results of the state of corrosion measured at these spectra are presented in 204 Table 3. The surface was actively corroding and the protective coatings were severely degraded. 205 Experimental results from the above three spectra have shown that four elements Fe, Si, Al, and O 206 were present. Spectrum 1 indicated a high amount of corrosion product because of a high concentration of O, which was responsible for the oxidation of Fe. Fe was recorded 56.25 wt. % at this 208 spectrum. The experimental results have found other elements such as Al and Si within the same 209 corrosion spectrum. Results from spectrum 2 has shown highest corrosion damage comparing to 210 spectra 1 and 3. The highest Fe concentration has been recorded in spectrum 3 with Si, Al, and O. 211

Previous studies [6] have shown elements such as Fe, Cr, Mn, Ni, and Si through XRF analysis 212 of the cross section of the Centaur sample. Again Si originates from alloying elements and Al as a 213 residue from passivation treatment (Al-based coating) applied during/after the Centaur service life. 214

Corrosive pits of random sizes were identified approximately 100  $\mu$ m deep in the cross section 215 of the sample illustrated in Fig. 6. Nickel (Ni), Fe, Mn, Chromium (Cr), Ca, S, Si, Al, and O were 216

Sample: Centaur-corroding Surface (Results in Weight %)									
Processing Option: All Elements Analysed/Normalised									
Spectrum No.	0	Al	Si	Fe					
1	36.58	3.97	3.18	56.25					
2	36.88	22.34	5.04	35.72					
3	28.06	2.15	4.09	65.67					

TABLE 3—Centaur EDS points analysis results of corroding surface.

#### PROOF COPY [MPC20120016]

AQ1

#### SAEED ET AL. ON CORROSION DAMAGE ANALYSIS



FIG. 6—Centaur cross section—location of EDS points, corrosion propagation, corrosion pits, and cracks.

identified at four spectra and their results are given in Table 4. Spectra 1, 2, and 3 are of the corrosion product layer, which clearly shows the high amount of oxygen versus elemental iron. Fe was recorded only 18.75 wt. % in the crack in spectrum 2 as shown in Fig. 6. O, Ca, and S are present in considerable amounts and will form localised zones of corrosion to propagate any cracks preferentially across the bulk metal grain boundaries leading to delamination of the surface material. Spectrum 4 was selected at the bulk metal and has been located approximately  $50 \,\mu$ m below the interface of the corrosion product layer and the bulk metal. Results from this spectrum have shown alloying elements only. Corrosion propagation of approximately  $50 \,\mu$ m deep from surface to subsurface was observed in this sample.

A sub-surface crack was identified in the cross section of the Centaur, shown in Fig. 7. Results 226 from the EDS investigations are provided in Table 5. Spectra were measured at the crack and at the 227 bulk area surrounding this crack. At spectra 1 and 2, only the alloying elements were identified and 228 no O or Al was detected. However, in the crack, O was present. It is likely that this crack will prop-229 agate when subjected to stresses and/or fatigue. 230

Figures 6 and 8 illustrate a localised form of corrosion pits associated with surface corrosion. <sup>231</sup> Pits are widespread and their formation into cracks/cavities can be observed in Fig. 8 [32]. The formation of pits in two phases, (a) meta-stable and (b) stable pits, has been extensively reported <sup>233</sup>

				4	· ·	5			
	Sample: Centaur—Cross Sectional Surface (Results in Weight %)								
Processing Option: All Elements Analysed/Normalised									
Spectrum No.	О	Al	Si	S	Ca	Cr	Mn	Fe	Ni
1	40.39	0.29	3.39	0.40	1.71	1.30	2.00	49.88	0.60
2	39.86	1.05	14.12	0.29	0.47	1.35	24.06	18.75	0.00
3	41.14	0.00	0.57	0.76	1.85	5.52	0.68	47.94	1.50
4	0.00	0.00	0.41	0.00	0.00	1.87	0.46	96.50	0.74

TABLE 4—Centaur EDS (point analysis) results of cross section.

#### 10 MATERIALS PERFORMANCE AND CHARACTERIZATION



FIG. 7-Centaur cross section sub-surface crack and EDS points.

[14,25,33–36]. Initially because of non-linear coupling between the dissolution of the metal and the 234 electrolyte composition, the formation of the pit takes place: (a) after which pit may propagate for 235 a short period and then die-down (meta-stable pit), or (b) it may continue propagating indefinitely, 236 known as stable pit, depending on the local fluctuations in the hydrodynamic boundary layer 237 [14,33,34].

The Centaur is exposed to rain, sea winds, and temperature fluctuations. Consequently, with <sup>239</sup> time, the sizes of the pits in Figs. 6 and 8 are expected to increase sidewise in the corresponding <sup>240</sup> plane as well as in depth, depending on (a) pitting potential, (b) re-passivation potential, and (c) in-<sup>241</sup> hibition potential and their properties [37,38]. A corrosive environment, under fatigue loads and <sup>242</sup> the localised pitting in the Centaur could lead to the formation of large cavities and could become <sup>243</sup> one possible form of structural failure even though considerable material remains on the surfaces <sup>244</sup> [22,39,40]. Furthermore, in its current environment, corrosion pits in the Centaur (Figs. 6 and 8) <sup>245</sup> will continue their electrochemical reactions and cause inter-sub-granular corrosion, resulting in <sup>246</sup> stress corrosion cracking (SCC) when exposed to stresses [40,41]. Climatic effects such as oxygen, <sup>247</sup> water, and salts are able to promote surface corrosion [21,39] and can penetrate into the subsurface layers through the corrosion cracks (Fig. 5) delivering rapid corrosion. <sup>249</sup>

Significant precipitation in Bovington for 18 days of each month [42] on average has been 250 reported in terms of fog, rain, and/or snow. The relative humidity averages around 80 % for most 251

				5 5					
Sample: Centaur—Cross Sectional Surface (Results in Weight %)									
Processing Option: All Elements Analysed/Normalised									
Spectrum No.	О	Al	Si	Cr	Mn	Fe	Ni	Мо	
1	0.00	0.00	0.00	1.94	0.74	96.63	0.67	0.00	
2	0.00	0.00	0.42	1.90	0.47	96.42	0.76	0.00	
Crack	4.36	0.86	1.58	1.87	1.23	88.21	0.65	1.20	

TABLE 5—Centaur EDS results of sub-surface crack and bulk metal.

#### SAEED ET AL. ON CORROSION DAMAGE ANALYSIS 11



FIG. 8—Centaur cross section—corrosion pits in random shapes and geometries.

of the year; however, peaks of 100 % and lows over a range of 40 %–60 % occur [43]. Average high 252 and low temperatures for around 30 years (1981–2010) were 14.03°C and 8.11°C, respectively, 253 whereas daily average fluctuations between 7°C to 22°C in summer and 2°C to 12°C in winter 254 were observed [42,44]. 255

There are 30 Acid and Aerosols Monitoring Network (AGANet) sites in the United Kingdom 256 [45]. There is no such monitoring centre at Bovington. Therefore, the AGANet site at Goonhilly 257 Downs was used to understand the air quality with respect to average deposition of salts and pollu-258 tants. Goonhilly Downs is situated in the Lizard Peninsula in Cornwall, England and is approxi-259 mately 6 km north of the English Channel. Similar environmental conditions are expected in 260 Bovington, which is around 9 km from the same marine environment.

The average salts and pollutants recorded between 2006 and 2009 were: chlorides at  $4.0 \,\mu g/m^3$ , 262 sulphur dioxide at  $0.080 \,\mu g/m^3$ , sulphites at  $1.00 \,\mu g/m^3$ , nitrates at  $3.00 \,\mu g/m^3$ , sodium at 263  $2.00 \,\mu g/m^3$ , and no calcium and magnesium were found [45]. Sulphur dioxide and sulphite levels 264 are shown to be relatively low, though they are only indirectly related to the effects of sulphur 265 dioxide on corrosion but the actual amount of hydrated sulphur dioxide deposited on metal 266 surfaces is important. The actual amount of sulphur deposited on the outdoor tanks will be less 267 than the amount found in the aerosols. The atmospheric chloride concentration is low, as expected 268 for a normal rural area.

The presence of S-containing compounds in the air and presence of S at the cross section of the 270 Centaur will stimulate corrosion. The rust formed on the Centaur will absorb  $SO_2$  and water from 271 the air and the result would be the formation of sulphate-containing electrolytes. When corrosion 272 occurs, the formation of iron sulphate (FeSO<sub>4</sub>) takes place, which then reacts with oxygen resulting 273 in iron oxy-hydroxide. Consequently, local acidity is increased by the regeneration of sulphate [13]. 274

$$\text{FeSO}_4 + 1.5\text{H}_2\text{O} + 0.5\text{O}_2 \rightarrow \text{FeO}(\text{OH}) + 2\text{H}^+ + \text{SO}_4^{2-}$$

The electrochemical mechanism developed by the continuous wet–dry cycles in this natural environment, via condensation, precipitation, heat, and wind would play a significant role in the corrosion of the tanks [46,47]. Once the paint barrier is damaged and the underlying carbon steel 277

#### 12 MATERIALS PERFORMANCE AND CHARACTERIZATION

substrate is exposed to the environment, the activity of the corrosion reactions would be almost 278 continuous. The oxidation of the metal is known to occur in an accelerated manner during wet 279 conditions. Furthermore, the highest rate of corrosion takes place just before drying starts, i.e., 280 when the liquid layer is thin (before evaporation) and the oxygen transport through the electrolyte 281 film is maximum [13,48].

The addition of chlorides and sulphurs, in the form of sulphates, sulphides, sulphites, and sulphurous salts, in the atmospheric environment, increases the corrosion rate [49–52]. Salts play 284 three main roles in accelerating corrosion: (a) they are imperfections on the surfaces and thereby 285 points of corrosion nucleation, (b) they form a hygroscopic salt film on the substrate surface that 286 keeps the surface wet for longer periods of time because the salt layer causes the surface to remain 287 wet at a lower relative humidity compared to clean surfaces, and (c) salts act to increase the conductivity of the thin electrolyte film that drives the electrochemical reaction, i.e., corrosion [52,53]. 289

Bovington is around 9 km from the sea (English Channel), therefore, the electrolytic layer 290 formed on the surfaces of the tanks will be conductive because of the possibility of soluble chlorides 291 and sulphates; this may result in the breakdown of protection [13,22]. Once the breakdown of the 292 protection has occurred, corrosion products, airborne salinity, and particles of hygroscopic salts 293 will influence the rate of corrosion and as a consequence deterioration of the Centaur will prevail. 294

#### Conclusions

This research work identified corrosion pits, sulphide inclusions, sub-surface corrosion cracking, 296 and progressive general corrosion on the surfaces for both the Sherman and Centaur tanks. Study 297 of the environments both inside and outside the museum has been conducted to relate corrosion 298 activity within the context. 299

Corrosion residues were relatively thick on the surfaces of the Sherman, whereas the phenomof sub-surface corrosion pits was widespread in the Centaur. Corrosion damage in the Centaur is critical and can be classified as a major risk to its structural integrity. It is of great interest that tanks that display surface contaminants should be conditioned. For the museum artefacts, preventative methods that could compromise the historic distinction of the vehicles are not viable options. However, for their longevity, it is important to eliminate corrosion contaminants from the surfaces, and the control of temperature and humidity is needed.

Close proximity of Bovington to the English Channel results in extensive rainfall, winds, temperature fluctuations, and high humidity together with high salt content throughout the year. For the outdoor tanks, it is important that any surfaces without adequate protection be protected against such conditions, so their structural deterioration through corrosion can be minimised. 310

The novelty of the current research lies in the fact of fully characterising structural materials <sup>311</sup> where information and specifications were not complete or scarce. The current research provides <sup>312</sup> understanding of the failure mechanisms because of corrosion and linkages to the materials' characteristics. These results will inform an optimised design solution of controlled environment and <sup>314</sup> preventative measures for the vehicles kept in uncontrolled environments. <sup>315</sup>

#### Acknowledgments

This research is jointed funded by the Tank Museum at Bovington, United Kingdom and Bourne- 317 mouth University. The writers thank Mike Hayton and Erik Blakeley, and the Tank Museum, 318 Bovington for their contributions. 319

316

AQ2

J\_ID: MPC DOI: 10.1520/MPC20120016 Date: 5-January-13

SAEED ET AL. ON CORROSION DAMAGE ANALYSIS 13

320

# References

AQ3

[1]	Mined A. C. "Comparing in the Militage" Comparing Fundaments and Industria. ACM Ltd.	
[1]	Vinod, A. S., Corrosion in the Military, Corrosion: Environments and Industries: ASM Int.,	321
[2]	Vol. 15C, 2006, pp. 126–135.	322 0
[2]	Moreneid, S., Drozdz, S., Hock, V., and Addott, W., Measuring Rates and Impact of Corro-	323
[2]	sion on DOD Equipment, Corros. Mil. II, vol. 38, 2008, pp. 163–181.	324
[3]	Chamberlain, P. and Ellis, C., British and American Tanks of World War II: The Complete	325
	Illustratea History of British, American, and Commonwealth Tanks, 1939–1945, Cassell & Co.,	326
[4]	London/New York, 2000.	327
[4]	Jackson, K., Tanks and Armoured Fighting venicles, Parragon, Bain, 2007.	328
[5]	Saeed, A., Knan, Z., Garland, N., and Smith, K., Material Characterisation to Understand	329
	Various Modes of Corrosion Fanures in Large Mintary Venicles of Historical Importance,	330
	Fifth International Conference on Computational Methods and Experiments in Materials	331
[6]	Characterisation, Kos, Greece, June 2011, pp. 13–15.	332
[6]	saeed, A., Knan, Z., Clark, M., Nel, N., and Smith, K., Non-Destructive Material Character-	333
	Destruct Test Courd Mariter Val 52 2011 pp 282 296	334
[7]	Destruct. Test. Cond. Monitor., Vol. 55, 2011, pp. 582–586.	335
[/]	Engel, L. and Kingele, H., An Alus of Metal Damage Surface Examination by Scanning Elec-	336
[0]	Scholl N. Marting D. V. Bodymann E. Duhnau H. H. Kishn D. and Schrover A. "The	337
[0]	High Energy Materials Science Recombine at DETPA III." Stress Fuel Mater Using Neutrons	220
	Sinc Padiat Vol 571 572 2008 pp 261 266	240
[0]	Blücher D. B. Svensson J. F. and Johansson J. G. "The Influence of CO. AlCl. 6H.O.	241
[2]	MgCl. 6H-O Na-SO, and NaCl on the Atmospheric Corrosion of Aluminum " Corros Sci	341
	Vol 48 2006 pp 1848-1866	3/12
[10]	Finlayson-Pitts B I and Pitts I N Atmospheric Chemistry: Fundamentals and Experimental	343
[10]	Techniques Wiley New York 1986	344
[11]	Chao I. Candevila-Montes C. and González-Carrasco I. I. "On the Delamination of FeCrAl	346
[11]	ODS Allovs" Mater Sci Eng A Vol 515 2009 pp 190–198	347
[12]	Hylander I. D. "Global Mercury Pollution and Its Expected Decrease after a Mercury Trade	348
[12]	Ban " <i>Water Air Soil Pollut</i> Vol 125 2001 pp 331–344	340
[13]	I von S. B. "Corrosion of Carbon and Low Alloy Steels" <i>Shreir's Corrosion</i> T. I. A. Richard-	350
[10]	son Ed Elsevier New York 2010 pp 1693–1736	351
[14]	Williams D F Kilburn M R Cliff I and Waterhouse G I N "Composition Changes	352
[14]	around Sulphide Inclusions in Stainless Steels and Implications for the Initiation of Pitting	352
	Corrosion "Corros Sci Vol 52 2010 pp 3702–3716	354
[15]	Kiessling R and Lange N. Non-Metallic Inclusions in Steel 2nd ed. Metals Society London	355
[15]	1078	355
[16]	Pereira A A Boehs I and Guesser W I "The Influence of Sulfur on the Machinability of	257
[10]	Grav Cast Iron FC25" I Mater Proc Technol Vol 179 2006 pp 165–171	358
[17]	Wranglen G "Pitting and Sulphide Inclusions in Steel" Corros Sci Vol 14 1074 pp.	350
[1/]	331-349	360
[18]	Bento I M V Pena A Lino C M and Pereira I A "Determination of Ochratovin A	361
[10]	Dento, J. H. T., Leno, G. H., and Ferena, J. A., Determination of Oematoxin A	501

[10] Bento, J. M. V., Fena, A., Eno, C. M., and Ferena, J. A., "Determination of Octratoxin A 361 Content in Wheat Bread Samples Collected from the Algarve and Bragança Regions, Portugal: 362 Winter 2007," *Microchem. J.*, Vol. 91, 2009, pp. 165–169.

ID: veeraragavanb Time: 19:01 I Path: //xinchnasjn/ASTM/3B2/MPC#/Vol00201/120037/APPFile/AT-MPC#120037

- 14 MATERIALS PERFORMANCE AND CHARACTERIZATION
  - [19] Jeon, S.-H., Kim, S.-T., Lee, I.-S., and Park, Y.-S., "Effects of Sulfur Addition on Pitting Corro- 364 sion and Machinability Behavior of Super Duplex Stainless Steel Containing Rare Earth Met- 365 als: Part 2," *Corros. Sci.*, Vol. 52, 2010, pp. 3537–3547.
  - [20] ISO 9223, "Corrosion of Metals and Alloys—Corrosivity of Atmospheres—Classification, 367
     Determination and Estimation," International Organization for Standardization, Geneva, 368
     Switzerland, 2012, p. 15.
  - [21] Cai, J. P. and Lyon, S. B., "A Mechanistic Study of Initial Atmospheric Corrosion Kinetics 370 Using Electrical Resistance Sensors," *Corros. Sci.*, Vol. 47, 2005, pp. 2956–2973.
     371
  - [22] Schweitzer, P. A., Fundamentals of Corrosion: Mechanisms, Causes, and Preventative Methods, 372
     CRC, Boca Raton, FL, 2010.
     373
  - [23] Trethewey, K. R. and Chamberlain, J., Corrosion: For Students of Science and Engineering, 374
     Longman Scientific & Technical, Harlow, U.K., 1988.
     375
  - [24] Kart, H. H., Uludogan, M., and Cagin, T., "DFT Studies of Sulfur Induced Stress Corrosion 376 Cracking in Nickel," *Comput. Mater. Sci.*, Vol. 44, 2008, pp. 1236–1242.
  - [25] Williams, D. E., Mohiuddin, T. F., and Zhu, Y. Y., "Elucidation of a Trigger Mechanism for 378
     Pitting Corrosion of Stainless Steels Using Submicron Resolution Scanning Electrochemical 379
     and Photoelectrochemical Microscopy," J. Electrochem. Soc., Vol. 145, 1998, pp. 2664–2672. 380
  - [26] Eklund, G. S., "Initiation of Pitting at Sulphide Inclusions in Stainless Steel," J. Electrochem. 381
     Soc., Vol. 121, 1974, pp. 467–473.
     382
  - [27] Greenwood, N. N. and Earnshaw, A., *Chemistry of the Elements*, 2nd ed., Butterworth-Heinemann, Oxford, 1997.
     384
  - [28] Castaño, J. G., Botero, C. A., Restrepo, A. H., Agudelo, E. A., Correa, E., and Echeverría, F., 385
     "Atmospheric Corrosion of Carbon Steel in Colombia," *Corros. Sci.*, Vol. 52, 2010, pp. 216–223. 386
  - [29] Kamimura, T., Hara, S., Miyuki, H., Yamashita, M., and Uchida, H., "Composition and Pro-387 tective Ability of Rust Layer Formed on Weathering Steel Exposed to Various Environments," 388 *Corros. Sci.*, Vol. 48, 2006, pp. 2799–2812.
  - [30] Dillmann, P., Mazaudier, F., and Hœrlé, S., "Advances in Understanding Atmospheric Corro- 390 sion of Iron: I. Rust Characterisation of Ancient Ferrous Artefacts Exposed to Indoor Atmos- 391 pheric Corrosion," *Corros. Sci.*, Vol. 46, 2004, pp. 1401–1429.
     392
  - [31] Ma, Y., Li, Y., and Wang, F., "The Effect of β-FeOOH on the Corrosion Behavior of Low Carbon Steel Exposed in Tropic Marine Environment," *Mater. Chem. Phys.*, Vol. 112, 2008, pp. 394 844–852.
  - [32] Williams, D. E., Westcott, C., and Fleischmann, M., Stochastic Models of Pitting Corrosion of 396
     Stainless Steels: Part 1; Modelling of the Initiation and Growth of Pits at Constant Potential, 397
     UKAEA Atomic Energy Establishment Materials Development Division, Harwell, 1984.
  - [33] Williams, D. E., Westcott, C., and Fleischmann, M., Stochastic Models of Pitting Corrosion of 399 Stainless Steels: Part 2; Measurement and Interpretation of Data at Constant Potential, 400 UKAEA Atomic Energy Establishment Materials Development Division, Harwell, 1984.
  - [34] Williams, D. E., Westcott, C., and Fleischmann, M., "Studies of the Initiation of Pitting Corro-402 sion on Stainless Steels," *J. Electroanal. Chem. Interfacial Electrochem.*, Vol. 180, 1984, pp. 403 549–564.
  - [35] Williams, J. A., "Wear and Wear Particles—Some Fundamentals," *Tribol. Int.*, Vol. 38, 2005, 405 pp. 863–870.
  - [36] Alvarez, M. G. and Galvele, J. R., "Pitting Corrosion," *Shreir's Corrosion*, T. J. A. Richardson, 407
     Ed., Elsevier, New York, 2010, pp. 772–800.
     408

AQ4

SAEED ET AL. ON CORROSION DAMAGE ANALYSIS 15

444

- [37] Baboian, R., Automotive Corrosion Tests and Standards, Society of Manufacturing Engineers, 409 Dearborn, MI, 1996. 410 [38] Askey, A., Lyon, S. B., Thompson, G. E., Johnson, J. B., Wood, G. C., Sage, P. W., et al, "The 411 Effect of Fly-Ash Particulates on the Atmospheric Corrosion of Zinc and Mild Steel," Corros. 412 Sci., Vol. 34, 1993, pp. 1055-1081. 413 [39] Turnbull, A., McCartney, L. N., and Zhou, S., "A Model to Predict the Evolution of Pitting 414 Corrosion and the Pit-to-Crack Transition Incorporating Statistically Distributed Input 415 Parameters," Corros. Sci., Vol. 48, 2006, pp. 2084-2105. 416 [40] Horner, D. A., Connolly, B. J., Zhou, S., Crocker, L., and Turnbull, A., "Novel Images of the 417 Evolution of Stress Corrosion Cracks from Corrosion Pits," Corros. Sci., Vol. 53, 2011, pp. 418 3466-3485. 419 [41] Weather Underground, 2011, http://www.wunderground.com/weatherstation/WXDailyHis-420 tory.asp?ID=IDORSETB5 (Last accessed 21 Nov 2011). 421 [42] Metoffice, 2011, "Station Data," http://www.metoffice.gov.uk/climate/uk/stationdata/hurnda-422 ta.txt (Last accessed 21 Nov 2011). 423 [43] dorsetforyou.com, 2012, "Climate Data for Weymouth, England (1981-2010)," http:// 424 webapps-wpbc.dorsetforyou.com/apps/weather/annualreport.asp. 425 [44] Defra, 2011, "Acid Gas and Aerosol Network (AGANet)," http://uk-air.defra.gov.uk/net-426 works/network-info?view=aganet (Last accessed 24 Nov 2011). 427 [45] Stratmann, M., Bohnenkamp, K., and Engell, H. J., "An Electrochemical Study of Phase-428 Transitions in Rust Layers," Corros. Sci., Vol. 23, 1983, pp. 969-985. 429 [46] Stratmann, M., Bohnenkamp, K., and Ramchandran, T., "The Influence of Copper upon the 430 Atmospheric Corrosion of Iron," Corros. Sci., Vol. 27, 1987, pp. 905-926. 431 [47] Alwash, S. H., Ashworth, V., Shirkhanzadeh, M., and Thompson, G. E., "An Investigation of 432 the Reduction of Oxygen at a Rotating Disc Electrode With Heat Transfer Facilities," Corros. 433 Sci., Vol. 27, 1987, pp. 1301–1311. 434 [48] El-Mahdy, G. A., Nishikata, A., and Tsuru, T., "Electrochemical Corrosion Monitoring of 435 Galvanized Steel under Cyclic Wet-Dry Conditions," Corros. Sci., Vol. 42, 2000, pp. 183-194. 436 [49] Nishikata, A., Yamashita, Y., Katayama, H., Tsuru, T., Usami, a., Tanabe, K., et al, "An Elec- 437 trochemical Impedance Study on Atmospheric Corrosion of Steels in a Cyclic Wet-Dry Con-438 dition," Corros. Sci., Vol. 37, 1995, pp. 2059-2069. 439 [50] Yadav, A. P., Nishikata, A., and Tsuru, T., "Electrochemical Impedance Study on Galvanized 440 Steel Corrosion under Cyclic Wet-Dry Conditions-Influence of Time of Wetness," Corros. 441 Sci., Vol. 46, 2004, pp. 169–181. 442 [51] Schwitter, H. and Bohni, H., "Influence of Accelerated Weathering on the Corrosion of Low-443
  - Alloy Steels," J. Electrochem. Soc., Vol. 127, 1980, pp. 15-20. [52] Corvo, F., Mendoza, A. R., Autie, M., and Betancourt, N., "Role of Water Adsorption and Salt 445 Content in Atmospheric Corrosion Products of Steel," Corros. Sci., Vol. 39, 1997, pp. 446 815-820. 447
  - [53] Neufeld, A. K., Cole, I. S., Bond, A. M., and Furman, S. A., "The Initiation Mechanism of Cor-448 rosion of Zinc by Sodium Chloride Particle Deposition," Corros. Sci., Vol. 44, 2002, pp. 449 555-572. 450

AQ6

AQ5