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¹ CO₂-Corrosion of Carbon steel: 53 54 ² The Synergy of Chloride Ion 55

Concentration and Temperature ⁵⁶₅₇ on Metal Penetration.

Frederick Pessu, Richard Barker and Anne Neville. 5

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22 ABSTRACT

23 This paper investigates the synergy of chloride ion concentration an $\mathscr{B}3$ 24 temperature on the general and pitting corrosion of carbon steel in $\mathrm{CO}_2 84$ 25 saturated environments. Experiments were conducted over 168 h in $\5 26 different concentrations of NaCl brines (1, 3.5 and 10 wt. %) an 86 27 temperatures (30, 50 and 80°C) with the aim of elucidating the combined effect of changes in chloride ion concentration an $\frac{87}{2}$ 28 29 general and pitting corrosion. This also includes a correlation with th $\overset{89}{=}$ 30 formation and properties of ${\sf FeCO}_3$ corrosion products. Linea 9031 polarisation resistance (LPR) was implemented to monitor the 9132 electrochemical responses. Corrosion product characteristics and 33 morphologies were studied through a combination of scanning electron 9334 microscopy (SEM) and X-ray diffraction (XRD). Pitting corrosion94 35 evaluation was conducted through the application of 3D surface95 36 37 profilometry to study pit geometries such as the depth and diameter96 38 The results show that general and pitting corrosion is strongly correlated 97to the synergistic effects of changing chloride ion concentration and 9839 temperature in carbon steel as a result of their combined influence o_1^{99} 40 ferrite (Fe) dissolution and FeCO $_3$ formation. This represents a paradig 10041 shift from the already established mechanisms on chloride ion and 1000 and 10000 and 1000 and 10000 and 1000 and 1000 an 42 43 temperature effects on passive alloys such as stainless steel. Increasing chloride ion concentration and temperature up to 10 wt. % NaCl and 50 44 80°C, respectively is observed to increase the rate of Fe dissolution and $\frac{103}{104}$ formation of semi-protective FeCO₃ corrosion products, leading to the 45 46 increase manifestation and severity of pitting corrosion. The results also 47 show a "threshold chloride concentration" exists at 30°C, above which 48 there is no significant increase in corrosion rate. However, such 49 "thresholds effect" were not observed at higher temperatures 50 evaluated in the range of chloride concentration considered in this 51 110 52 study. 111

INTRODUCTION

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Corrosion of pipeline steel in oilfields is usually characterized by a complex material degradation mechanism. In most oilfield scenarios, the corrosion damage mechanisms often experienced are driven by synergistic interactions of various process parameters, especially in its applications in CO₂ and/or H₂S-saturated environments. Corrosion damage in such environments is usually dominated by, but not limited to general and/or pitting corrosion^[1]. Chloride ions have been considered as an important environmental parameter that can affect the process of pitting corrosion of carbon steels in oilfields by destabilising the characteristic protectiveness of a FeCO3 corrosion product^[2, 3]. Incidences of a combination of pitting and general corrosion failures in CO₂-saturated environment are presently of significant concern^[4]. This is because of the lack of fundamental understanding of localized/pitting corrosion components in the degradation process of carbon steels in such environments, and the difficulty associated with predicting the dominant nature and risks of localized and/or pitting corrosion of carbon steels.

For active materials such as carbon steel, the prospect of significant general corrosion makes the mechanism of pitting corrosion complex, as both corrosion process are capable of occurring at the same time on a particular region of pipeline steel. Some of these complexities are possibly linked to the contributing driving force for general and pitting corrosion of carbon steels in these environments [5-7]. In such environments, the ionic strength contribution of dissolved chloride ions in the brine phase is also important^[8]. It has been reported that the pitting corrosion process of carbon steel in CO₂ environments is driven by localized breakdown of protective and/or formation of a nonprotective FeCO₃^{[9, 10])}. It has also been shown in previous publications^{[6,} ^{11]} and by the work of other authors^[7, 12], that factors such as temperature, pH, and exposure time can influence the CO₂ corrosion process, associated film formation characteristics and morphology. These factors could also influence the tendency for pitting corrosion of carbon steel to occur in these environments.

Water chemistry is considered one of the most influential parameters affecting CO₂ corrosion. The mechanism of CO₂ corrosion is known to be dependent on many interrelated physicochemical factors whose interactions with steel in CO₂-saturated environments is considered very important^[13]. Oilfield produced water is usually characterized by a high amount of dissolved salts, with chloride ions being the most prominent ^[14-16]. The role of dissolved chloride in accelerating corrosion reactions^[13], and hence pitting corrosion of passive alloys such as stainless steel are already widely recognized^[15]. For stainless steels, chloride ion concentration in the solution is considered as a decisive parameter in defining the pitting resistance of stainless steel^[15]. The autocatalytic process of pit propagation in stainless steels is known to be driven by chloride ions^[17] acting to reduce the local pH and prevent the re-passivation of active pits.

In carbon steel, chloride ions have been reported to be more aggressive than other halides because of their smaller atomic diameter, and as such can fundamentally be considered as an aggressive agent of corrosion capable of destabilising corrosion product layers ^[2, 13, 16]. However, the mechanism by which this effect is manifested in carbon steel has also remanined elusive. There is a significant lack of understanding of the role of chloride ion concentration on pitting corrosion of carbon steel where there is significant contribution of general corrosion. Most existing models of the effect of chloride ion across various temperature ranges in carbon steel corrosion is still built on established understaning of the effect of chloride ion on passive alloys. The most recent studies

1 on this subject have consistently offered conflicting viewpoints while 42 mostly focusing exclusively on general corrosion. There has been little 53 or no attention to the effect of chloride ion on the combined evolution 5657

4 of general and pitting corrosion of carbon steel.

58 5 Experiments by Luo and Singh^[8] on carbon steel in simulated fuel-grade59 6 ethanol (SFGE) show an increase in corrosion rate with increasing 60 7 chloride ion from 0 to 160 mg/L of NaCl at room temperature. This trent 61 8 is also consistent with similar observations reported by Papavinasam et 9 $al^{[9]}$, who varied chloride ion concentration from 10,000 ppm to 120,00 **6**2 10 ppm (other operational variables associated with these experiment 6311 were not disclosed). Contrasting viewpoints were present from 6412 experiments by Jiang et al^[18] at 80°C, Fang et al^[19] at 25°C and da⁶⁵ 13 Chagas Almeida^[20] at 24°C. All three authors concluded that corrosion 6614 rate decreases with increasing chloride ion concentration, while Jiang e6715 al^[18] and Fang et al^[19] suggested that the risk of pitting corrosior $\mathbf{68}$ 16 decreases with increase in chloride ion concentration. However, Fang et 17 $al^{[19]}$ also suggested that the presence of chlorides gave an indication o69 18 an increase in solution conductivity and ionic strength, but stops shor7019 of confirming a possible effect of lower solution resistance (linked $t \overline{q} 1$ 20 solution conductivity/ionic strength) on the general corrosion rate. 72

73 21 A recent study by Gao et al^[2] also reported on the destructive74 22 capabilities of increasing chloride ion on pre-formed FeCO3 at 80°C75 23 Changes in chloride ion were achieved in this study by the introduction 7624 of NaClO₄ into a corrosive environment after the formation of protective 7 25 FeCO₃. The result from this study shows that Cl⁻ ion and/or ionic strength 78 26 can destabilize an already formed protective corrosion product laye79 27 leading to pit initiation. However, it does not explain the evolution \circ 80 28 the pitting process with consideration for the general corrosion proces 81 29 that supports pitting corrosion and drive metal penetration for carbor82 30 steel materials^[6]. 83 84

The work presented in this paper is focused on understanding how85 31 changes in chloride ion concentration in combination with changes in 8632 33 temperature can influence the corrosion mechanisms ang7 34 characteristics of carbon steels in an unbuffered CO2-saturate data 35 environment. This study aims to elucidate the synergy of two criticago 36 environmental variables; temperature and chloride ion concentration on the general and pitting corrosion process of carbon steels in CO_{91} 37 38 environments. 92 39

40 Materials and Experimental Procedure

The purpose of this work is to investigate the general and localized 9641 corrosion processes of carbon steel in three different CO_2 -saturate 9742 environments, containing 1, 3.5 and 10 wt. % NaCl, each a 9843 44 temperatures of 30°C, 50°C and 80°C. The focus of this study is to understand how general corrosion (related to ferrite dissolution), FeCO 45 formation and pitting corrosion processes of carbon steel are influence 10046 by changes in chloride ion concentration in synergy with temperature01 47 The partial pressure of CO_2 gas in the gas phase of the corroside 0248 103 49 environment at different temperature are provided in Table 1 104 <u>Materials:</u> X65 carbon steel samples were used as the working $\frac{100}{100}$ 50 electrodes within a three-electrode cell in each experiment. The stee 19651

composed of a ferritic/pearlitic microstructure. The nominal 7 52 108 53 composition of X65 steel is provided in Table 2. 109 The carbon steel was sectioned into 10 mm x 10 mm x 5 mm samples. Wires were soldered to the back of each test specimen and then embedded in a non-conducting resin. Prior to the start of each experiment, test samples were wet-ground up to 1200 silicon carbide grit paper, degreased with acetone, rinsed with distilled water and dried with compressed air before immersion into the test brine. A surface area of 1 cm² was exposed to the electrolyte per sample and 10 samples were immersed in each 2 litre vessel.

Brine preparation and test conditions: Three different NaCl brine solutions were used for all experiments. The test solution was saturated with CO₂ for a minimum of 12 h (usually overnight) prior to starting each experiment to deaerate the test environment. CO₂ was also bubbled into the system throughout the duration of every test and all tests were conducted at atmospheric pressure. The experiment matrix is shown in Table 3.

In-situ electrochemical measurements: Twin test cells were used in each experiment to ensure repeatability of corrosion rate measurements. Corrosion rates were recorded from two samples per test cell. 10 samples, each of 1 cm² surface area, were immersed into a 2 litre test solution. The samples used for monitoring electrochemical responses remained inside the test cells until the end of the experiment to allow full collection of electrochemical data over the 168 h. Corrosion rate measurements were conducted using both DC and AC measurements with an ACM potentiostat. DC Linear polarisation resistance (LPR) measurements were performed by polarising the working electrode ± 15 mV about the open circuit potential (OCP) at a scan rate of 0.25 mV/s to obtain polarisation resistance measurements (R_p). Tafel polarisation measurements were performed on freshly ground samples in separate tests to determine anodic and cathodic Tafel constants. This was combined with Faraday's Law and the measured values of R_p to estimate general corrosion rates. Tafel plots were obtained by performing anodic and cathodic sweeps ±250 mV about the OCP at a scan rate of 0.25 mV/s. Both anodic and cathodic sweeps were performed on separate samples in the same test vessel to ensure reliable measurements. Tafel polarisation measurements were performed after 7 h in separate experiments to those samples from which corrosion measurements were recorded over 168 h. Electrochemical impedance spectroscopy measurements were made to account for solution resistances. To ensure that the measured solution resistance truly reflects the ionic strength of electrolyte, the proximity between reference, counter and working remained consistent in each test and for the duration of the tests. This was achieved by adding the samples through designated holes to ensure that all working electrodes are equidistant from the reference electrode, especially the sample used for electrochemical measurements.

Characterisation of FeCO₃ corrosion products and pitting corrosion: Corrosion tests were conducted for 168 h with the aim of monitoring the changes in different morphologies of corrosion products and assessing their influence on the evolution of general corrosion and the surface pitting at chosen test conditions. Corrosion product characterisation was performed using Scanning Electron Microscopy (SEM) and X-ray Diffraction (XRD). XRD patterns were collected using a Bruker D8¹ equipped with a LynxEye² detector, using a 90 position auto sampler and employing Cu K α radiation with an active area of 1cm^2 programmable di-vergence slits. Scans were performed over a range 20 = 10 to 70° at a step size of 0.033 per second and a total scan time of

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1 ~50 minutes. The results were analysed using X'Pert 3 HighScor54 2 software and compared with individual crystal standards from an in-

3 house database.

4 Surface profilometry was implemented in this study to quantify pittin565 attack. Samples removed at different time intervals from the same tes §7 6 conditions were cleaned to remove all traces of corrosion products using 8 7 Clarke's solution before using profilometry to determine the extent o59 8 pitting corrosion on the steel surface. Clarke's solution consisted of 20 60 9 antimony trioxide + 50 g stannous chloride + 1000 ml 38% hydrochlori61 10 acid, prepared in accordance with ASTM⁴ Standard G1-03. The sample 2 11 surfaces were wiped with a cotton pad soaked in Clarke's solution63 12 washed, air dried, and carefully stored in a de-aerated chamber to avoid 64 13 oxidation prior to surface analysis. Pit depth measurements wer65 14 performed in alignment with ASTM standard G46-94^[21]. NPFLEX⁵ 3166 15 interferometer was used for obtaining the discrete geometry of pits or 67 16 over 81% of the steel surface (the remaining 19% represents the edge6817 of the sample) to represent the entire exposed surface area. Pits wer6918 identified based on carefully chosen thresholds. The thresholds were $\overline{\mathcal{O}}$ 19 chosen with reference to the surface roughness of generally corrode $\sqrt[3]{1}$ 20 areas surrounding the pits. This enabled pits with distinct pit parameter $\overline{y}2$ 21 of interests; pit-mouth diameters and pit depths, to be identified an $\sqrt{3}$ 22 quantified. ASTM G46-94 stipulates that an average depth of the 107423 deepest pits and the depth of deepest pit (based on relative pit depth 75 24 measurement after removal of corrosion products) should be used for 76 25 evaluating and quantifying the extent of pitting corrosion damage. The 726 measured pits are ranked according to how deep they are across the 2827 entire exposed surface. The 10 deepest are used to calculate the 9 28 average pit depth for a specific test condition, according to ASTM80 29 standard G46-94^[21]. This is plotted along with the depth of the deepes81 30 pit identified. On the basis of the recommendation in the ASTM8231 standard G46-94^[21], the pit density was not quantified in this study. 83 32 84

33 Results and Discussion

Stern-Geary constants were calculated using condition specific Tafe8734 constants from polarisation measurements for carbon steel in three different chloride concentrations at each temperature. The Stern-Gear constants are provided in Solubility data for dissolved CO₂ (Table 3) is calculated using experimentally determined Henry's coefficients 35 36 37 constants are provided in Solubility data for discorred 0.21 solutions is calculated using experimentally determined Henry's coefficients for CO₂ in water at different NaCl solutions and at different 93 38 39 40 temperatures by Zheng et al.^[37]. Table 4. Stern-Geary constants (B) were used in Equation 1 to calculate 41 corrosion current and consequently corrosion rate based on a tota9542 96 43 exposed projected surface area of 1 cm² per sample. 97

$$i_{corr} = \frac{1}{2.303R_p} \left[\frac{1}{\beta_a} + \frac{1}{\beta_c} \right]^{-1} = \frac{B}{R_p}$$
 Equation 1

Note: the corrosion rate estimated from linear polarisation (LPR9844 45 measurements captures both the material loss responses occurring 9946 across the entire exposed surface including the localized regions 90 47 Therefore it is not used as a true quantitative measure of contributid 01 48 of general and pitting corrosion in this study. However, the trend in L μQ^2 49 responses (in the form corrosion rate) gives an indication to the exte10350 to which corrosion products is being deposited on the steel surface 10451 impede corrosion process. It is important to note that corrosion produt 0^{5} 52 established as a result will also occur at areas where pitting corrosion 196107 53 likely to manifest. 108

Corrosion rates and ferrite dissolution

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Corrosion rate calculated from LPR measurements and measured in-situ pH of the bulk solution are presented in Figure 1(a) and (b), Figure 2(a) and (b) and Figure 3(a) and (b) for 30, 50 and 80°C, respectively. In Figure 1(a), the first 24 h shows approximately equal corrosion rate of ~1.1 mm/yr for tests in 1 and 10 wt. % NaCl at 30°C, and a slightly higher corrosion rate for test in 3.5 wt. % NaCl at ~1.5 mm/yr. This observation indicates a "threshold effect" which can be linked to the "salting out effect" of chloride concentration on dissolved CO₂ gas^[3] (provided in Table 3). Decreasing dissolved CO₂ with increasing chloride concentration reduces the driving force for the cathodic reaction^[1]. Salting-out effect describes the decrease in solubility due to the presence of a salt in a solute-solvent system^[3]. However, increase in the ionic strength and hence the conductivity of the electrolyte due to higher chloride concentration could also offer a counter effect of increase in corrosion rate. This could be the reason why initial corrosion rate is higher in 3.5 wt. % NaCl. At low temperature (30°C) and atmospheric pressure, it is believed that the "salting out effect" of NaCl on dissolved CO₂ is more dominant than the electrolyte conductivity effect^[3], especially in the initial stages of the corrosion process. Electrolyte conductivity is related to the rate of transfer of electrons. Other researchers ^[3, 13, 22] have also reported on the threshold effect at lower temperature. According to Fang et al.^[22], at 20°C and after 24 h, corrosion rate decreases by 50% with increase in chloride concentration from 3 to 10 wt.%, which is similar to results shown in Figure 1(a) after 24 h. Eliyan et al.^[13] reported an increase in corrosion rate with increase in chloride concentration in terms of NaCl from 0 to 1.5 wt.% at 20°C, after which the corrosion rate decreases steadily to 3.5 wt.% concentration and remained constant up to a concentration of 8 wt.%, while Liu et al.^[23], observed peak corrosion rate at 2.5 wt. % of Cl⁻ concentration for tests carried out between 0-15 wt. % Cl- at 2 MPa, 100°C for 72 h in CO₂ saturated systems. Within the same 24 h period, there is no evidence of "threshold effect" with respect to chloride concentration as shown in Figure 2(a) and Figure 3(a) for tests at 50 and 80°C respectively. The absence of the threshold effect is due to the dominance of the ionic strength and hence the electrolyte conductivity effect of the environment on the redox charge exchanges that drives ferrite dissolution and corrosion process at high temperature. Referring to Table 3, the salting effect on dissolved CO₂ is significant. However, at high temperatures, this is suppressed and the increased aggressiveness of the corrosion environment represents a synergy of chloride concentration and temperature on the corrosivity of water chemistry ^{[8,} 16, 24]

According to Figure 1(a), the "threshold effect" disappears at 30°C after 24 h, resulting in a continuous increase in corrosion rate in 10 wt.% NaCl until an equal corrosion rate is attained for tests in in 3.5 and 10 wt.% NaCl. The change in the corrosion rate profile between 24-168 h now follows an expected trend of increased solution conductivity and aggressiveness (decreasing solution resistance); also reported by Fang et al^[19], with increase in chloride concentration (measured solution resistances were 27.5 ± 0.5 ohms.cm², 9.0 ± 0.2 ohms.cm² and 4.1 ± 0.2 ohms.cm² for 1, 3.5 and 10 wt.% NaCl, respectively at 30°C from this study). With respect to increasing aggressiveness of corrosion environments, it is expected that charge-transfer will increase, leading

⁵ Trade name

³ Trade name

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1 to increasing corrosion rate as shown in the later stages of corrosion rate602 in Error! Reference source not found.(a). While the time-dependen613 disappearance of the "threshold effect" remains unclear, it is believe dc_2 4 that the overall corrosion rate behavior is the result of the competitiv $\mathbf{63}$ 5 effect of "salting out effect" on dissolved CO_2 and aggressiveness of the 646 corrosion environment and potentially other secondary effects tha 657 have not been considered in this study. The work by das Chagas Almeid $\mathbf{a}\mathbf{b}$ 8 et al^[20] at CO₂ partial pressure of 100 kPa and 30 MPa distilled water an $\mathbf{67}$ 9 18.7 wt.% NaCl (3.2M NaCl) at 24°C shows that at higher pressure, the 6810 corrosion rate was higher with high NaCl. This is also an evidence t $\mathbf{69}$ 11 support the findings in the paper that the "threshold effect" is limite $\overline{J}0$ 12 by a range of other environmental parameters within a corrosion 7113 system. It is important to note that the results of Fang et al.^[22] and Li \overline{J}^2 14 et al.^[23] were based on a 24 h and 72 h exposure, respectively, while th $\sqrt{3}$ 15 exposure time for the experiment by Elivan et al^[13] was not stated. 74 75

16 76 17 The corrosion rate measured for the tests conducted at 50 and 80° C wa37consistent throughout the duration of the experiments (see Figure 2(a7818 19 and Figure 3(a) for 50 and 80°C, respectively), showing strong direct79 20 correlation with increase in aggressiveness of environment. The solution \$021 resistance of corrosion environment at 80°C was measured at 14.8 \pm 0. \pounds 1 22 ohms.cm², 4.9±0.6 ohms.cm² and 2.6±0.0 ohms.cm² for 1, 3.5 and 10 wt82 23 % NaCl, respectively. This is significantly higher than at lowe 83 24 temperatures (30°C). This observation indicates the effect of therma&425 activation of chloride ions at corrosion interfaces at high 8526 temperatures^[16], leading to higher rates of ferrite dissolution. Th\$627 difference in the corrosion behaviour at 50 and 80°C is primarily in term\$728 of the magnitude of the corrosion rate and the time it takes to peal 8829 corrosion rate values. Such difference clearly highlights the synergy o8930 chloride ions and temperature in driving the Fe dissolution process90 31 Corrosion rate peaked within the first 11 h (at ~ 3.2 mm/yr, 5.6 mm/y9132 and 7.0 mm/yr) for tests in 1, 3.5 and 10 wt. % NaCl at 80°C and withir 9233 the first 60 h (at \sim 2.2mm/yr, 3.5 mm/yr and 5.3 mm/yr) for tests in 193 34 3.5 and 10 wt. % NaCl at 50°C. The relatively faster rate to peak corrosion 9435 rate and reduction of corrosion rate with time observed at 80°C than aheta 536 50 and 30°C can be attributed to the rate of Fe²⁺ generation from Fe 37 dissolution and saturation of corrosion interface for FeCO₃ formation^[996] 38 (although amorphous in nature and semi-protective due to th 97 39 relatively low but continuously evolving pH (shown in Figure 1(b), Figurheta 840 2(b) and Figure 3(b) for 30, 50 and 80°C, respectively)). This is discusse $\oplus 9$ 41 in detail in the next section. 100 101

42 Corrosion product formation

103 The effect of chloride concentration and temperature on the formati $\frac{100}{100}$ 43 and evolution of corrosion product layers is shown by the SEM $\mathsf{imag}_{195}^{105}$ 44 provided in Figure 4, Figure 6 and Figure 8 for 30, 50 and 8040645 respectively and the XRD patterns in Figure 5, Figure 7 and Figure 9 $\mathrm{f}4\mathrm{p}7$ 46 1 and 10 wt.% NaCl solutions at for 30, 50 and 80°C, respectively. Befo $\frac{108}{100}$ 47 discussing the corrosion product formation process in detail, it $109\,$ 48 important to establish the relationship between Fe dissolution, FeC $\frac{10}{2}$ 49 50 formation and *in-situ* pH changes with time shown in Figure 1(b), Figure 1 51 2(b) and Figure 3(b) for 30, 50 and 80°C, respectively. It is believed that 2the reduction of corrosion rate is due to the formation of corrosi $\frac{1}{2}$ 52 products, specifically FeCO₃. However, the extent of such reduci $\frac{1}{2}$ 53 effect is determined by the competing effect of rate of release of Fe 15 54 and ${\tt FeCO}_3$ formation at the corrosion interface and correlated $\frac{1}{10} \frac{1}{6}$ 55 evolving bulk and surface $pH^{[25]}$. According to Guo et al.^[25], increasi 56 *in-situ* pH is an indication that the rate of Fe dissolution is greater than 1857 the rate of precipitation of FeCO₃. In-situ pH will remain constant $\frac{1}{2}$ 58 decreases when the rate of Fe dissolution equals or is lower than the 12059

rate of FeCO₃ precipitation^[25]. Increase in Fe dissolution removed more H⁺ from the environment, creating a charge imbalance. H⁺ removed by the cathodic reaction is replenished by H₂CO₃ acting as a buffer. The balance of rate of removal of H+ and replenishment by H₂CO₃ could either increase or decrease the pH depending on the direction of change of charge neutrality. Of all the pH data shown in Figure 1(b), Figure 2(b) and Figure 3(b) for 30, 50 and 80°C respectively, a period of constant pH is established to confirm the scenario of equal rate of Fe dissolution and FeCO₃ formation except for test in 10 wt. % NaCl at 80°C. Prior to the end of the test in 10 wt. % NaCl at 80°C, there was an initial increase in pH from ~3.9 at the start to ~5.6 after 50 h, corresponding to periods of increasing Fe dissolution and followed by a drop in pH after ~50 h. This is attributed to the scenario where the rate of precipitation of FeCO₃ becomes greater than the rate of generation of Fe^{2+[25]}. It also indicates attainment of saturation of the corrosion interface far above the saturation limit for FeCO₃. This scenario was only observed in the case of 10 wt. % NaCl at 80°C and not at lower concentration of NaCl and/or lower temperatures. This is a direct effect of the aggressiveness of the environment to drive Fe dissolution process, FeCO₃ formation, thus, confirming the synergy of chloride ions and temperature. It is also important to note that the results presented in Figure 1(b), Figure 2(b) and Figure 3(b) for 30, 50 and 80°C respectively in this study and the observation in Guo et al^[25] are based on experiments in closed systems. In such systems the exchange of interfacial ionic species for FeCO₃ with the bulk environment is limited and hence rate of saturation of the interface can be directly correlated to the rate of Fe dissolution with the consequence of inducing real-time buffering effect on the bulk pH shown in Figure 1(b), Figure 2(b) and Figure 3(b). It is believed that in actual oilfield pipe interfaces, mass transfer effect will likely be substantial leading to a different effect of Fe dissolution on changes in in-situ pH. Particularly as mass transfer effect will remove Fe²⁺ from the interface while also replenishing the aggressive ions that drives Fe dissolution. Thus it is likely that longer exposure time will be required to achieve similar bulk pH changes observed in this study. In some cases this may only occur during pipe shut-in or in the event of restriction to pipe flow.

From the SEM images in Figure 4, Figure 6 and Figure 8 for 30 (after 168 h), 50 (after 7 and 168 h) and 80°C (after 7 and 168 h), respectively, it is evident (especially from Figure 6(a)-(f) and Figure 8(a)-(f) for tests at 50 and 80°C after 7 and 168 h) that the formation of FeCO₃ is usually preceded by Fe dissolution, leaving behind an iron carbide (Fe₃C) network^[1, 5, 6, 26]. There is also evidence of a polycrystalline form of FeCO₃ after 168 h (FeCO₃ was identified using XRD as shown in Figure 5, Figure 7 and Figure 9 for test at 30, 50 and 80°C respectively in 1 and 10 wt. % NaCl solutions). The nature of FeCO3 formed across the three temperatures of interest after 168 h shows a transition from amorphous and/or polycrystalline to crystalline FeCO3 in the direction of increase in both temperature and chloride concentration. At each temperature, the transition from amorphous and/or polycrystalline to crystalline FeCO₃ also occurs in the direction of increasing chloride concentration. This is most evident for test in 10 wt. % NaCl; Figure 4(c), Figure 6 (f) and Figure 8 (f) for 10 wt. % at 30, 50 and 80°C, respectively. These observations correlate with the trend of increase in high initial corrosion rate (Fe dissolution) due to the aggressiveness (presence of chloride ion at high temperatures) of the environment. This same aggressiveness is driving the evolution of FeCO₃ from an amorphous and/or polycrystalline to easily identifiable FeCO₃ crystals. XRD patterns shown in Figure 5, Figure 7 and Figure 9 for test at 30, 50 and 80°C, respectively in 1 and 10 wt. % NaCl solutions consistently show an increase in the intensity of peaks for FeCO₃ with increase in chloride ion concentration. This is a direct correlation between progressive increase of Fe dissolution (related to

1 increased corrosion rate) with the morphology of FeCO₃ corrosion 602 product and the transition from amorphous and/or polycrystalline t $\mathbf{61}$ 3 crystalline FeCO₃ over 168 h. It is therefore evident that in an un624 buffered and closed corrosion system and in the absence of any pre635 formed FeCO₃ (typical for new oilfield flow lines), the effect of chlorid 646 concentration is strongly associated with its influence on the electrolyti $\delta 5$ 7 strength and aggressiveness of the corrosion environments. This als668 imply that $FeCO_3$ film formation can also be enhanced with increase in 679 chloride ion concentration by the same mechanism, leading to increase 10 in the interfacial saturation level for formation of $FeCO_3^{[5]}$. This effect is 11 shown in this study to be further enhanced by increase in temperatur6912 via thermal activation of chloride ion ^[16, 27]. Increasing temperatura 70 13 accelerates most of the chemical, electrochemical and transport 71processes at the corrosion interface; H^+ reduction and H_2CO_3/HCO_372 14 reduction (or its buffering effect)^[28, 29]. Thermal activation of chloride⁷³ 15 16 ions at higher temperatures therefore impacts the rate of 17 electrochemical activation of carbon steel to lose more Fe2+ and as shown by the results in this study also contributes towards the 18 precipitation of $FeCO_3^{[16, 27]}$. These trends represents the synergy that 19 exist between temperature and chloride ion concentration in terms $o\frac{7}{78}$ 20 metal penetration from general corrosion rate and rate of saturation o_{fg} 21 22 corrosion interface. This synergy is summarised in Figure 10 in terms of 23 the estimated thickness loss due to general corrosion from LPRO 24 measurements after 72 and 186 h. Estimated thickness loss presenteg in Figure 10 is based on LPR measurements on the assumption tha 8225 corrosion rate captured from the electrochemical measurement ig3 26 27 general across the entire corrosion interface. Although this could als **Q**₄ 28 takes into account the localized electrochemical activity related to 85 29 pitting corrosion, it provides a useful baseline for qualifying the 86 aggressiveness of the bulk environment and its transition with time. The 30 31 pitting corrosion contribution to LPR measurement is assumed to be relatively small in comparison to the general corrosion process taking 32 place across the entire surface. This is evident from the data on \widetilde{gg} 33 estimated thickness and the pitting corrosion analysis presented later in $\tilde{b_0}$ 34 35 this paper. 91 92

36 The results reported thus far in this paper is in contrast to the genera93 37 viewpoints on the influence of chloride ion on the process of corrosion 38 product formation by several authors $^{[2,\ 16,\ 30]}.$ Chloride ions have ofter 9439 been linked to the destabilisation of FeCO $_3$ corrosion products; after i9540 is formed, and increasing the solubility product and saturation limi9641 required for the formation of FeCO₃ corrosion products. It is importan $\frac{97}{2}$ to note that in the findings of Gao et al^[2] that also support this 42 viewpoint, FeCO_3 corrosion products were initially formed at a high^{99} 43 initial pH of 6.6 and Fe²⁺ concentration of 100 ppm in a 0.1-1 wt. % NaCl 44 45 solution before subjecting the corrosion media to higher chloride ig concentration. While the work of Gao et al^[2] may have shown that the source of t 46 higher chloride ion concentration could destabilise the corrosid 47 products, the results presented in this work provides usering 103 48 understanding of the effect of chloride ion concentration on $\frac{160}{1000}$ dissolution and FeCO₃ corrosion product evolution in CO₂ corrosion $\frac{160}{1000}$ 49 50 51 systems. The result presented here is particularly relevant for CO ĥ07 saturated corrosion environments where the corrosion productor formation process is slowly preceded by electrochemical dissolution $\frac{107}{109}$ Fe²⁺ as shown by the results Figure 6 and Figure 8 at 50 (after 7 and 168) 52 53 54 55 h) and 80°C (after 7 and 168 h), respectively.

56 Pitting corrosion

- 57 The results discussed in this section is focused on the relationship $\frac{111}{12}$
- 58 between pitting corrosion of carbon steel and the synergetic influence 13
- 59 of chloride ion concentration and temperature. This is also discussed 114

the context of overall metal penetration. The depth of the deepest pit, average depth of pit and the aspect ratio based on the deepest pit on ~81% of the entire exposed 1 cm² sample over 168 h for different chloride concentration are provide in Estimated thickness loss (μ m) based on LPR measurements showing the synergistic effect of chloride ion concentration and temperature on carbon steel penetration after (a) 72 h and (b) 168 h. *Note: severity of material loss in magnitude increases in this order; Green* \rightarrow *Amber* \rightarrow *Red.*

Figure 11, Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 30°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. *Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits*

Figure 12 and Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 50°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. *Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.*

Figure 13 for 30, 50 and 80°C, respectively. Since the depths of pits are characterized after removal of corrosion products, it is important to note that the contribution of Fe dissolution to metal penetration is already removed in the data shown in Estimated thickness loss (μ m) based on LPR measurements showing the synergistic effect of chloride ion concentration and temperature on carbon steel penetration after (a) 72 h and (b) 168 h. *Note: severity of material loss in magnitude increases in this order; Green* \rightarrow *Amber* \rightarrow *Red.*

Figure 11, Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 30°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. *Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits*

Figure 12 and Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 50°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. *Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.*

Figure 13. The diameter of the deepest pit for 1 and 3.5% wt. NaCl at 30, 50 and 80°C were used in combination with the depth of deepest pit to estimate the aspect ratio. The aspect ratio is defined as the ratio of the diameter (Dia.max) and depth (d_{max}) of the deepest pit on an entire exposed surface area. It is mathematically presented in Equation 2. It highlights the role and scope of influence of general corrosion in the pitting corrosion process. The aspect ratios of 1 and 3.5 wt. % NaCl are used to show how the pit diameters evolves during pitting corrosion process. 1 and 3.5% NaCl were chosen because pit diameter for 10 wt. NaCl was not recorded during measurements.

$$Aspect\ ratio\ = \frac{Dia_{max}}{d_{max}}$$
 Equation 2

When aspect ratio increases, it indicates that pit diameter increase at a higher rate than pit depth. When aspect ratio remains constant, either the pit diameter and pit depth are growing at proportional rate or both processes are being retarded at an equal rate. However, when aspect ratio decreases, it indicates that pit depth is increasing at a higher rate

than the pit diameter attack around the surrounding surface of an activ $\mathbf{58}$ 1 2 59 pit. 60 It has already been shown that Fe₃C exposure results from Fe dissolution $\pounds 1$ 3 4 especially in the early stages of the corrosion process^[1, 5, 26]. Therefore 62 5 it is believed that the initial stages of pitting corrosion (pit initiation), i $\mathfrak{G3}$ 6 related to the role of revealed Fe₃C in establishing distinct local anodi647 and cathodic sites (galvanic corrosion cells) across the corroding surface 65 8 It has previosly been shown that the overall progression of pitting b9 corrosion is linked to the nature, morphology and protective capabilitie of FeCO₃ formed over time^[6]. FeCO₃ formed in this case is either an 6810 11 amorphous and/or polycrystalline (semi-protective) FeCO₃ corrosion 69 12 product layer capable of supporting the progress of pitting corrosion^{[9,} ^{31]}. The initial Fe dissolution and establishment of Fe₃C is believed to b φ_0 13 critical in the establishment of local anodic and cathodic sites. The 1 14 15 protectiveness and/or semi-protectiveness of $FeCO_3$ is therefore hinged 2 16 on the distribution of saturated species of Fe²⁺ and CO_3^{2-} at the corrosion **7**3 17 interface for the formation of FeCO₃. The distribution of FeCO₃ acros \overline{a} 4 18 the corroding interface is likely to anchor preferentially on Fe₃C due $t\sigma^{5}$ 19 its lower overpotential for the cathodic reactions (HCO_3^{-}/CO_3^{2-}) than 20 $Fe^{[1, 32]}$. A combination of such Fe_3C -FeCO₃ distribution can lead to th $\cancel{2}6$ 21 establishment of galvanic effects between adjoining Fe-rich regions and 722 Fe₃C-FeCO₃ rich regions, thereby accelerating the dissolution of Fe b $\sqrt{28}$ 79 23 accelerating the cathodic reaction adjacent to it. 80 By correlating the corrosion rate data in Figure 1, Figure 2, and Figure $\1 24 25 for 30, 50 and 80°C, respectively with pitting corrosion data provided in Estimated thickness loss (μ m) based on LPR measurements 8226 showing the synergistic effect of chloride ion concentration an \mathscr{B}^3 27 temperature on carbon steel penetration after (a) 72 h and (b) $16\4 28 h. Note: severity of material loss in magnitude increases in this85 29 86 30 order: Green \rightarrow Amber \rightarrow Red. 87 Figure 11(a) and (b), Pitting corrosion in X65 carbon steel in different 31 NaCl solutions over 168 h, at 30°C (a) Deepest pit, (b) averages 32 depth of pits and (c) Aspect ratio based on the diameter of deepes 33 depth of pits and (c) Aspect ratio based on the diameter of accepting pit. Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the 92 34 35 36 standard deviation of 10 deepest pits 93 Figure 12(a) and (b), and Pitting corrosion in X65 carbon steel in $\frac{94}{2}$ 37 different NaCl solutions over 168 h, at 50°C (a) Deepest pit, (b25 38 average depth of pits and (c) Aspect ratio based on the diamete 96 39 of deepest pit. Note: Pit depth quantified after removal of corrosion 97 40 product layer and Error bars on average depth of pits represent $\98 41 42 the standard deviation of 10 deepest pits. 99 100 43 Figure 13(a) and (b) for 30, 50 and 80°C, respectively, it is evident the 10^{10}

the rate of Fe dissolution (and revealing of Fe₃C) and supersaturation 10244 45 the corrosion interface is strongly linked with the evolution of pitting 3 46 corrosion over 168 h. The depths of pits (average and deepest) aft10447 removal of corrosion products consistently show lowest values after 10548 h for tests in 10 wt.% NaCl; ~13 μm at 30°C, ~22 μm at 50°C and ~24 $\mu h 06$ 49 at 80°C. These values were observed to be either lower than or equal 10750 the pit depth for 1 wt. % NaCl; ~13 μ m at 30°C, ~30 μ m at 50°C and ~2D851 μ m at 80°C. These observation shows that at high chlorid 40952 concentration and higher temperature the depth of pits are lower d14053 to higher rate of Fe dissolution which produces Fe²⁺ for the formation 111 54 FeCO₃. The pit depth is lower at lower chloride concentration and low $\frac{1}{2}$ 55 temperature due to the lower rate of Fe dissolution and hence low 1356 rate of FeCO₃ formation. This is consistent with the previous discussion 11457 on how increasing chloride concentration and temperature increases

the rate of saturation of corrosion interface for promoting the formation of FeCO₃. The implication of this effect is that with increasing exposure time, the rate of FeCO₃ formation increases as interfacial saturation increases, especially for high chloride and temperature systems. This also increases the protection from FeCO₃ covered areas, which are electrochemically distinct but connnected in a galvanic cell with locally corroding anodes, to support pit growth as shown by the increase in the depth of pits after 168 h in Estimated thickness loss (μ m) based on LPR measurements showing the synergistic effect of chloride ion concentration and temperature on carbon steel penetration after (a) 72 h and (b) 168 h. *Note: severity of material loss in magnitude increases in this order; Green* \rightarrow *Amber* \rightarrow *Red*.

Figure 11(a) and(b), Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 30°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits

Figure 12(a) and (b), and Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 50°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. *Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.*

Figure 13(a) and (b) for 30, 50 and 80°C, respectively (10 wt.% NaCl; ~26 μ m at 30°C, ~52 μ m at 50°C and ~31 μ m at 80°C compared to 1 wt.% NaCl; ~15 μ m at 30°C, ~52 μ m at 50°C and ~34 μ m at 80°C). Between 72 h and 168 h and for test at 80°C the size of pit increased by ~30% for test in both 1 wt.% (from ~27 μ m to ~34 μ m) and 10 wt.% NaCl (from~24 μ m to ~31 μ m).

Although the final depth of pit is slightly higher in 1 wt.% NaCl, the results also shows that while increasing chloride concentration could increase the rate of FeCO₃ (although semi-protective) formation to support pit growth, its effect on Fe dissolution also supresses pitting corrosion contribution to total metal penetration. The reverse effect is observed with decreasing chloride concentration. For tests in 10 wt.% NaCl and between 72 and 168 h, pit depth only increased from ~13 μm to ~26 μ m at 30°C, ~22 μ m to ~52 μ m at 50°C and ~24 μ m to ~31 μ m at 80°C, representing 50%, 136% and 30% increase for 30, 50 and 80°C respectively. At 30°C, low temperature supresses the aggressive effect of chloride ions needed to increase interfacial supersaturation and form semi-protective FeCO₃ that supports pit growth, especially as this temperature records the lowest general corrosion rate. At 80°C, the aggressiveness of chloride ions are enhanced to drive Fe dissolution (manifesting mainly as high general corrosion). This will lead to the formation of semi-protecive FeCO₃ capable of supporting significant pit growth. However, the size of pit is reduced by the receeding corroding interface due to general corrosion. At 50°C, the highest pit growth percentage increase was recorded because of the balancing effect of both Fe dissolution and FeCO₃ formation to support pit growth at this intermediate temperature. This is consistent with the general characterisation of FeCO₃ formed at temperatures ≤60°C as nonprotective^[1]; a precursor of pitting corrosion attack^[9]. The results discussed in this section is considered a disntinctive corrosion characteristics of active metals such as carbon steel^[7] and clearly shows that chloride concentration and temperature can act in synergy to determine the corrosion mechanism that dominates metal penetration

in CO_2 containing corrosion environments, especially when the 571 2 58 corrosion environment is un-bufferred. 59 The aspect ratios for 1 and 3.5 wt.% NaCl are presented in Estimate 603 4 thickness loss (μ m) based on LPR measurements showing the 1 5 synergistic effect of chloride ion concentration and temperature or 62 6 carbon steel penetration after (a) 72 h and (b) 168 h. Note: severit 63 of material loss in magnitude increases in this order; Green -64 7 65 8 Amber \rightarrow Red. 66 67 9 Figure 11(c), Pitting corrosion in X65 carbon steel in different NaCl 10 solutions over 168 h, at 30°C (a) Deepest pit, (b) average depth of 8 pits and (c) Aspect ratio based on the diameter of deepest pitgo 11 Note: Pit depth quantified after removal of corrosion product layer 12 and Error bars on average depth of pits represents the standard 13 14 deviation of 10 deepest pits 72 Figure 12(c) and Pitting corrosion in X65 carbon steel in different $\frac{73}{74}$ 15 NaCl solutions over 168 h, at 50°C (a) Deepest pit, (b) average 75 16 17 depth of pits and (c) Aspect ratio based on the diameter of deepest pit. Note: Pit depth quantified after removal of corrosion product, 18 layer and Error bars on average depth of pits represents the 78 19 20 standard deviation of 10 deepest pits. 79 Figure 13 (c) for 30, 50 and 80°C, respectively. Generally, an increase in $\frac{80}{4}$ 21 aspect ratio implies that the diameter of pits is getting broader while the 82 22 23 pit depth is becoming shallower. This is usually the case when there is 33 24 significant contribution of general corrosion to metal penetration such 34

25 that the surrounding surfaces around the pit is receding. When the 85 26 aspect ratio is constant, it implies that either the rate at which the pit is 86 27 growing depth-wise is equal to the rate at which the pit diameter is broadening by general corrosion or both processes are being retarded at an equal rate. When the aspect ratio is decreasing, it implies that the 28 29 rate of pit growth is higher than the rate of diameter broadening due to $\frac{88}{2}$ 30 general corrosion. On the premise of this, it is evident from Estimated thickness loss (μ m) based on LPR measurements showing the 31 32 synergistic effect of chloride ion concentration and temperature of 2 carbon steel penetration after (a) 72 h and (b) 168 h. *Note: severity* 23 33 34 35 of material loss in magnitude increases in this order; Green -94 36 Amber \rightarrow Red. 95

37 Figure 11(c) that within the first 36 h, general corrosion dominated the 38 material loss process in 3.5 wt. % NaCl than in 1 wt. % NaCl. This is shown 97 39 by the higher aspect ratio. However, substantial pit growth is observed 8 40 for 3.5 wt. % NaCl between 36-72 h after which the pit and diametegg 41 broadening rate remained almost the same. This coincides with the $\eta = 0$ 42 period of stable corrosion rate due to the establishment of semi-43 protective FeCO₃ on the corroding surface. For 1 wt. % NaCl, aspect ratio increased significantly with time until 120 h, showing a period where general corrosion dominates metal penetration. At 50°C, aspect ration 44 45 after 7 h show a substantial decrease (increase in pit depth compared 2 46 to increase in pit diameter) in 1 wt. % NaCl while it remained constant 4 3.5 wt. % NaCl. This corroborates the increase in general corrosing 4 47 48 05 49 contribution with increase in chloride concentration (3.5 wt. % NaCl) high temperature to cause pit depth to be increasing at the same rate 50 51 as the diameter is increasing. This can be described as the "broadening of pit". At 80°C, the aspect ratio decreases after 7 h (3.5 wt.%) and 340752 (1 wt.%) to indicate a period of higher pit growth rate than the rate 108 increase of diameter until after 72 h when the general corrosion rate 53 54 becomes dominant. At this stage the diameter of pit increases fast ± 10 55 56 than pit growth and the surface of pit will recede faster.

The results from this study, particularly the aspect ratio shows the effect of highly dynamic and complex interfacial phenomena that supports pitting corrosion process. General corrosion is able to impose limitations on pitting corrosion behavior of carbon steel to such an extent that the pit growth rate become seemingly retarded depending on the conditions. Pit retardation can also be due to localized self-healing of pits due to local saturation with precipitable anions ^[6, 12]. Chloride ion concentration and temperature have been shown in this study to have strong effect in how all of these interfacial phenomena act in synergy to determine the relative dominance of a suit of corrosion mechanisms driving metal penetration.

The results discussed above also explains the complex means by which chloride ions influences the pitting corrosion characteristics of carbon steel materials in un-buffered CO₂ corrosion environments. They are consistent with the evolution of pitting corrosion from a starting point of Fe dissolution (and Fe₃C revelation) and FeCO₃ formation. Therefore, in conditions where FeCO₃ has not been pre-formed as is the case in this study, the effect of chloride concentration on the general and pitting corrosion characteristics is therefore correlated on the corrosivity of the environment and how it translates into the different processes that leads to pitting corrosion; Fe dissolution and formation of nonprotective FeCO₃. This is consistent with the understanding that if FeCO₃ does not form, metal corrosion will be completely dominated by general corrosion^[9] and if a completely protective and non-porous FeCO₃ layer forms without any local galvanic cell and/or local anode, pitting corrosion is almost absent. Thus, any corrosion scenario that falls in between these absolutes will lead to a certain degree of pitting corrosion. However, once a protective FeCO₃ is formed, the mechanism of influence of chloride ion may change to become more of a destructive agent to the integrity of protective $FeCO_3^{[2]}$.

Figure 14(a) and (b), presents evidence of the relationship between the synergy of chloride ion concentration and temperature on total metal penetration after 72 h and 168 h. This is also presented in terms of the relationship between pitting factor and the synergistic effect of chloride ion and temperature in Matrix to qualify the effect of synergy of chloride ion concentration and temperature on total metal penetration of carbon steel after (a) 72 h and (b) 168 h. Note: severity of material loss in magnitude increases in this order; Green \rightarrow Amber \rightarrow Red.

Figure 15(a) and (b). The concept of pitting factor (P_f) has been introduced in previous publications $^{[6,\,11]}$ and used to reflect the relative contribution of corrosion damage mechanisms; general and pitting corrosion to metal penetration. It is defined in ASTM standard G46-94 $^{[21]}$ as:

Pitting Factor;

Where P_d is the deepest metal penetration (µm) for the whole exposed surface area (sum of depth of deepest pit (d_{max}), (after removal of corrosion products) plus the average estimated metal penetration (µm) from general corrosion rate measurement (termed P_u), i.e; $P_d = P_u + d_{max}$. A pitting factor of 1 represents general corrosion; i.e., $P_d = P_u$, and $d_{max} = 0$. The greater the (d_{max}), the greater the pitting factor.

 $P_f = \left(\frac{P_d}{P_u}\right)$

Referring to Matrix to qualify the effect of synergy of chloride ion concentration and temperature on total metal penetration of carbon steel after (a) 72 h and (b) 168 h. *Note: severity of material loss in magnitude increases in this order; Green* \rightarrow *Amber* \rightarrow *Red.*

Figure 15 and according to ASTM standard G46-94^[21], a higher pittin561 2 factor indicates a lower contribution to metal penetration from genera 57 3 corrosion. The "threshold effect" can be observed in the pitting facto $\overline{58}$ 4 data at 30°C and slowly disappears with increasing temperature a595 higher temperature within the first 36 h (Matrix to qualify the effect o60synergy of chloride ion concentration and temperature on tota61 6 7 metal penetration of carbon steel after (a) 72 h and (b) 168 h. Note62 8 severity of material loss in magnitude increases in this order 63 64 9 Green \rightarrow Amber \rightarrow Red. 65 Figure 15(a)), while the effect of general corrosion is evident in the 6610 lower pitting factor after 72 and 168 h for all conditions (Matrix t_{0}^{57} 11 qualify the effect of synergy of chloride ion concentration and 12 temperature on total metal penetration of carbon steel after (a) 72^{69} 13 h and (b) 168 h. Note: severity of material loss in magnitude 7014 71 15 increases in this order; Green \rightarrow Amber \rightarrow Red. 72 73 16 Figure 15(b) and (c)). At the higher temperature of 80°C, the results on 74 pitting factor shown in Matrix to qualify the effect of synergy of 5 17 chloride ion concentration and temperature on total metabo 18 penetration of carbon steel after (a) 72 h and (b) 168 h. Note 19 severity of material loss in magnitude increases in this order 78 20 21 Green \rightarrow Amber \rightarrow Red. 79

22 Figure 15 suggests that the pitting characteristics of carbon steel is

limited by the extent of general corrosion and FeCO₃ formation. While1 References 23 24 the actual ionic/electrochemical interaction of Cl⁻ within an active pit on

carbon steel still remain unclear and could not be established in this 82 1. 25

26 study, it is clear that such interaction will be different from the auto $\overline{g_3}$

27 catalytic changes in local chemistry and pit propagation in stainless steeba

28 in the presence of Cl⁻. Therefore, it is believed that the increase in the

total metal penetration with increase in chloride ion concentration and 29

temperature as shown in Figure 14 is related to the process of therma $\overline{66}$ 30

31 activation.

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32 Conclusions

- 89 33 The effect and synergy of chloride concentration and temperature or 90
- 34 carbon steel corrosion behavior; general and pitting corrosion in CO_291
- 35 containing corrosion environment have been investigated over 168 h.
- 36 The key conclusions from this work are presented herein:
- 37 ٠ Chloride ion concentration and temperature car94
- 38 independently and synergistically influence the overall 39 corrosion characteristic; general and pitting corrosion og 40 carbon steel to determine the extent of metal penetration 96 41 This is correlated on their influence on the rate of ferrite7 42
 - dissolution, and FeCO₃ formation kinetics and properties.
- 44 Generally, the rate of general corrosion increases with $\bar{q}q$ * 45 temperature and chloride concentration. However, at low temperatures, the effect of chloride ion concentration 10146 47 strongly linked to the "salting out effect" on CO₂ gas solubility and aggressiveness of the environment. The competition p_{02} 48 7. these phenomena activates the emergence of the "threshold 5349 effect" on the influence of chloride ion concentration $\dot{q} \vec{b} \vec{4}$ 50 51 corrosion rate at lower temperatures, particularly in the early 52 stages of the corrosion process. 105 8. 53
 - 106
- 54 At higher temperature, the "Arrhenius effect" of temperature ٠ causes the thermal activation of Cl⁻ species to increase the 55

aggressiveness of the corrosion environment and driving the Fe dissolution process for FeCO₃ formation. The synergy of thermal activation of chloride species increases rate of metal corrosion; including general and pitting corrosion. This synergy, especially with increasing chloride concentration and temperature, promotes the transition from amorphous and/or polycrystalline FeCO₃ to crystalline FeCO₃ due to increased saturation of corrosion interface with ferrous ions.

- The extent of metal penetration from pitting corrosion is also observed to be related to the ability of chloride ions and temperature effect to act synergistically to promote the formation of semi-protective FeCO₃. The rate of formation of this type of FeCO₃ increases with increasing chloride concentration, especially with increasing temperature from 30 to 50°C and supporting higher pit penetration depth.
- Based on pitting factor analysis, the "threshold effect" of chloride ions is also evident in the pitting initiation process; remaining still dominant at 30°C. At higher temperatures (80°C) and higher chloride concentration (10 wt. % NaCl), the threshold effect is replaced by the masking effect of pit penetration depth by the higher contribution from general corrosion to highlight the impact of the synergy of chloride and temperature on total metal penetration.

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13 **FIGURE CAPTIONS**

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- Figure 1: Plots of (a) corrosion rate and (b) in-situ bulk pH fo70 14 corrosion system of X65 carbon steel in 1, 3.5 and 10 wt. % NaC/71 15 72 16 soln. saturated with CO₂ at 30°C.
- 73 Figure 2: Plots of (a) corrosion rate and (b) in-situ bulk pH for 17 corrosion system of X65 carbon steel in 1, 3.5 and 10 wt. % NaC(⁴ 75 18
- 19 soln. saturated with CO₂ at 50°C.
- Figure 3: Plots of (a) corrosion rate and (b) in-situ bulk pH for corrosion system of X65 carbon steel in 1, 3.5 and 10 wt. % NaC78 20 21 22 soln. saturated with CO2 at 80°C.
- 23 Figure 4: SEM of corrosion products formed on X65 carbon steel
- 24 at 30°C after 168 h in (a) 1 wt. % NaCl soln., (b) 3.5 wt. % NaCl 25 soln., and (c) 10 wt. % NaCl soln.
- 26 Figure 5: XRD patterns for (a) 1 wt. % and (b) 10 wt. % NaCl soln.
- 27 for 168 h at 30°C (Note that the intensity scale is arbitrary). XRD79
- 28 patterns are referenced to the literature [33-36]. 80

29 Figure 6: SEM of corrosion products formed on X65 carbon steel 30 at 50°C in (a) 1 wt. % NaCl soln., 7 h, (b) 1 wt. % NaCl soln., 168 31 h (c) 3.5 wt. % NaCl soln., 7 h (d) 3.5 wt. % NaCl soln., 168 h, (e) 81

- 32 10wt. % NaCl soln., 7 h, and (f) 10 wt. % NaCl soln., 168 h.
- 82 33 Figure 7: XRD patterns for (a) 1 wt. % and (b) 10 wt. % NaCl soln for 168 h at 50°C (Note that the intensity scale is arbitrary). XRD⁸³ 34 35 patterns are referenced to the literature [33-36].

36 Figure 8: SEM of corrosion products formed on X65 carbon steel 37 at 80°C in (a) 1 wt. % NaCl soln., 7 h, (b) 1 wt. % NaCl soln., 168

- 38 h (c) 3.5 wt. % NaCl soln., 7 h (d) 3.5 wt. % NaCl soln., 168 h, (e)
- 39 10wt. % NaCl soln., 7 h, and (f) 10 wt. % NaCl soln., 168 h.
- 40 Figure 9: XRD patterns for (a) 1 wt. % and (b) 10 wt. % NaCl soln. 41 for 168 h at 80°C (Note that the intensity scale is arbitrary). XRD patterns are referenced to the literature [33-36]. 42
- 84 43 Figure 10: Estimated thickness loss (µm) based on LPF85 44 measurements showing the synergistic effect of chloride ior86 45 concentration and temperature on carbon steel penetration afte87
- 46 (a) 72 h and (b) 168 h. Note: severity of material loss in magnitude 88

47 increases in this order; Green \rightarrow Amber \rightarrow Red.

89 48 Figure 11: Pitting corrosion in X65 carbon steel in different NaCl 49 solutions over 168 h, at 30°C (a) Deepest pit, (b) average depth of 50 pits and (c) Aspect ratio based on the diameter of deepest pit. 51 Note: Pit depth quantified after removal of corrosion product layer 52 and Error bars on average depth of pits represents the standard 53 deviation of 10 deepest pits

Figure 12 : Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 50°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.

Figure 13 Pitting corrosion in X65 carbon steel in different NaCl Soln. over 168 h, at 80°C (a) size of deepest pit and (b) average size of pit. Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.

Figure 14: Matrix to gualify the effect of synergy of chloride ion concentration and temperature on total metal penetration of carbon steel after (a) 72 h and (b) 168 h. Note: severity of material

loss in magnitude increases in this order; Green \rightarrow Amber \rightarrow Red.

Figure 15: Matrix to qualify the effect of synergy of chloride ion concentration and temperature on the contribution to metal penetration from both pitting and generalgeneral corrosion in terms of pitting factor on carbon steel after (a) 36 h. (b) 72 h and (c) 168 h. Note: severity of material loss in magnitude increases in

this order; Green \rightarrow Amber \rightarrow Red.

TABLES

Table 1: Temperature and total partial pressures of CO₂ gas at atmospheric pressure above test solution

Temperature (°C)	CO ₂ partial pressure (kPa)
30	97
50	89
80	55

С	Si	Р	S	Мо	Mn	Ni	Nb	٧	Fe
0.15	0.22	0.023	0.002	0.17	1.42	0.09	0.05	0.06	97.81

Table 3: Dissolved CO₂ at experimental for assessing the effect of temperature and chloride conc. on corrosion of carbon steel.

NaCl Conc.	Temperature (°C)					
(Wt. %)	Dissolved CO ₂ (ppm)					
	30°C	50°C	80°C			
1	1242	763	458			
3.5	977	595	382			
10	594	297	158			

Solubility data for dissolved CO₂ (Table 3) is calculated using experimentally determined Henry's coefficients for CO2 in water at different NaCl solutions and at different temperatures by Zheng et al.^[37].

Table 4: Tafel constants for each experimental conditions investigated for the effect of temperature and chloride conc.

T(°C)			Tafe	el Para	meter	s (mV/	/dec)		
	1 wt.% NaCl			3.5 wt.% NaCl			10 wt.% NaCl		
	βa	βc	В	βa	βc	В	βa	βc	В
80	47	85	13	58	135	18	75	140	21

50	40	120	13	40	168	14	55	160	18
30	35	145	12	33	200	12	43	180	15

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Figure 16: Plots of (a) corrosion rate and (b) in-situ bulk pH for corrosion system of X65 carbon steel in 1, 3.5 and 10 wt. % NaCl solutions saturated with CO₂ at 30°C.



Figure 17: Plots of (a) corrosion rate and (b) in-situ bulk pH for corrosion system of X65 carbon steel in 1, 3.5 and 10 wt. % NaCl solutions saturated with CO2 at 50°C.



Figure 18: Plots of (a) corrosion rate and (b) in-situ bulk pH for corrosion system of X65 carbon steel in 1, 3.5 and 10 wt. % NaCl solutions saturated with CO₂ at 80°C.





(c)

Figure 19: SEM of corrosion products formed on X65 carbon steel at 30°C after 168 h in (a) 1 wt. % NaCl, (b) 3.5 wt. % NaCl, and (c) 10 wt. % NaCl solutions.











Mag = 1.00 KX 20.00 kV

(b)



Mag = 1.00 KX 20.00 kV SE1 10 μm

(c)







Figure 21: SEM of corrosion products formed on X65 carbon steel at 50°C in (a) 1 wt. % NaCl, 7 h, (b) 1 wt. % NaCl, 168 h (c) 3.5 wt. % NaCl , 7 h (d) 3.5 wt. % NaCl, 168 h, (e) 10wt. % NaCl, 7 h, and (f) 10 wt. % NaCl solutions, 168 h.



(b)

Figure 22: XRD patterns for (a) 1 wt. % and (b) 10 wt. % NaCl solutions for 168 h at 50°C (Note that the intensity scale is arbitrary). XRD patterns are referenced to the literature ^[33-36].







(d)





Figure 23: SEM of corrosion products formed on X65 carbon steel at 80°C in (a) 1 wt. % NaCl, 7 h, (b) 1 wt. % NaCl, 168 h (c) 3.5 wt. % NaCl, 7 h (d) 3.5 wt. % NaCl, 168 h, (e) 10wt. % NaCl, 7 h, and (f) 10 wt. % NaCl, 168 h.





Chloride Content	30°C	50°C	80°C			
1% NaCl	21.37	38.60	45.78			
3.5% NaCl	32.77	57.59	74.17			
10% NaCl	30.02	82.62	100.65			
(b)						

Figure 25: Estimated thickness loss (μ m) based on LPR measurements showing the synergistic effect of chloride ion concentration and temperature on carbon steel penetration after (a) 72 h and (b) 168 h. Note: severity of material loss in magnitude increases in this order; Green \rightarrow Amber \rightarrow Red.

Figure 26: Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 30°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.

Figure 27 : Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 50°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.

Figure 28: Pitting corrosion in X65 carbon steel in different NaCl solutions over 168 h, at 80°C (a) Deepest pit, (b) average depth of pits and (c) Aspect ratio based on the diameter of deepest pit. Note: Pit depth quantified after removal of corrosion product layer and Error bars on average depth of pits represents the standard deviation of 10 deepest pits.

Chloride Content	30°C	50°C	80°C
1% NaCl	22.26	48.25	50.62
3.5% NaCl	31.37	53.16	70.49
10% NaCl	24.07	60.47	76.96
	(a	a)	

Figure 29: Matrix to qualify the effect of synergy of chloride ion concentration and temperature on total metal penetration of carbon steel after (a) 72 h and (b) 168 h. Note: severity of material loss in magnitude increases in this order; Green \rightarrow Amber \rightarrow Red.

Chloride Content	30°C	50°C	80°C			
1%NaCl	3.49	3.30	2.94			
3.5%NaCl	2.42	2.90	1.97			
10%NaCl	3.52	1.87	1.63			
(a)						

Temperature (°C)

Temperature (°C)

Chloride Content	30°C	50°C	80°C
1%NaCl	1.73	2.38	1.74
3.5%NaCl	1.67	1.86	1.49
10%NaCl	1.85	1.64	1.31
		-1	

(c)

Figure 30: Matrix to qualify the effect of synergy of chloride ion concentration and temperature on the contribution to metal penetration from both pitting and uniform corrosion in terms of pitting factor on carbon steel after (a) 36 h, (b) 72 h and (c) 168 h. Note: severity of material loss in magnitude increases in this order; Green \rightarrow Amber \rightarrow Red.