# Effect of Inhibitors on Weld Corrosion under Sweet Conditions Using Flow Channel

Khaled Alawadhi, Abdulkareem Aloraier, Suraj Joshi, Jalal Alsarraf

Abstract-The aim of this paper is to compare the effectiveness and electrochemical behavior of typical oilfield corrosion inhibitors with previous oilfield corrosion inhibitors under the same electrochemical techniques to control preferential weld corrosion of X65 pipeline steel in artificial seawater saturated with carbon dioxide at a pressure of one bar. A secondary aim is to investigate the conditions under which current reversal takes place. A flow channel apparatus was used in the laboratory to simulate the actual condition that occurs in marine pipelines. Different samples from the parent metal, the weld metal and the heat affected zone in the pipeline steel were galvanically coupled. The galvanic currents flowing between the weld regions were recorded using zero-resistance ammeters and tested under static and flowing conditions in both inhibited and uninhibited media. The results show that a current reversal took place when 30ppm of both green oilfield inhibitors were present, resulting in accelerated weld corrosion.

*Keywords*—Carbon dioxide, carbon steel, current reversal, inhibitor, weld corrosion.

## I. INTRODUCTION

CARBON dioxide (CO<sub>2</sub>) corrosion of carbon steel pipelines and equipment in the oil and gas industry has received much attention in recent years because of an increased tendency to inject CO<sub>2</sub> into oil wells in order to reduce the viscosity of oil and increase production [1], [2]. It is very well known that CO<sub>2</sub> corrosion causes failure of pipelines and structural components in petroleum production processes, resulting in significant economic and manpower losses by catastrophic accidents and the pollution of water resources and the environment [3], [4].

Despite the fact that carbon steel has low resistance to  $CO_2$  environments, it is widely used in the petroleum industry mainly due to economic reasons [5]. Galvanic corrosion attack on welds in steel components and subsequent failure of the structures has been quite common in seawater pipelines. It is well known that galvanic corrosion can occur between the different regions of a weld due to differences in composition and microstructure [6].

Attempts to control preferential weld corrosion have

K. Alawadhi and J. Alsarraf are with the Department of Automotive and Marine Technology, The Public Authority for Applied Education and Training, P.O. Box 42325, Shuwaikh, Kuwait City 70654, Kuwait (e-mail: paaetq8@hotmail.com).

A. Aloraier is with the Department of Manufacturing and Welding Technology, The Public Authority for Applied Education and Training, P.O. Box 42325, Shuwaikh, Kuwait City 70654, Kuwait.

S. Joshi is with the Department of Mechanical and Industrial Engineering, Concordia University, 1515 St. Catherine Street West, Montréal, Québec, H3G 2W1.

It is acknowledged that the research described in this paper was supported by Clariant Oil Services.

previously involved making minor additions of noble metals (Ni, Cr, Mo and Cu) in order to make the weld metal cathodic with respect to the adjacent parent metal and the heat affected zone. Nevertheless, these additions must be made with caution since over-alloying can result in enhanced HAZ corrosion [7].

Using inhibitors is one of the most important applications that have been used widely in corrosion protection of carbon steel in  $CO_2$  environments. However, in some cases, the use of inhibitors to control internal pipeline corrosion has the effect of reversing the galvanic currents between the weld regions, such that the weld metal becomes anodic and accelerated rates of galvanic corrosion take place. An example of severe preferential weld corrosion (PWC) in a pipeline containing hydrocarbons and brine saturated with  $CO_2$  is shown in Fig. 1 [8].



Fig. 1 Severe preferential weld corrosion in carbon steel pipeline carrying inhibited oil and gas [8]

Weld corrosion has been investigated by a range of experimental techniques, such as a flow channel [9], a recirculating flow loop [10] and rotating cylinder electrodes [11]. There are only limited, documented studies concerning the corrosion of welded steel exposed to marine environment in general and in the oil and gas environment in particular.

#### II. EXPERIMENTAL WORK

#### A. Welded Pipeline Steel

All tests were carried out on samples machined from welded X65 steel pipe with a chemical composition of 0.08% C, 1.6% Mn, and 0.3% Si. The steel had been thermomechanically control-rolled with a fine-grained ferritic microstructure and a hardness of 200-210 Hv. The double-vee weld was produced by the submerged arc process, with a relatively high heat input (5-10 J/mm) and a hardness of 223-230 Hv. A sample was cut from the weld, polished and etched to identify the positions of the parent metal (PM), heat affected zones (HAZ) and weld metal (WM), as shown in Fig.

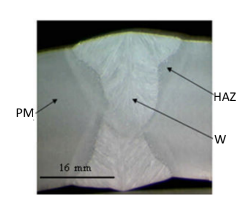


Fig. 2 Sections of the weld (parent metal, heat affected zone and weld metal)

#### B. Flow Channel Apparatus

The working electrodes were machined from commercial steel pipe grade API X65 and had wires attached to carry the electrochemical signals. The surface preparation was carried out by grinding/polishing with 1200 grit of silicon carbide paper and then degreasing with isopropanol to remove any dust and scratches in order that only the examined corrosion remained in the cell. The three regions were then separated by carefully cutting along the boundaries and were reassembled by mounting in epoxy resin. A Perspex plate was attached to the samples, separated by a 1mm thick rubber gasket, to form a flow channel through which brine saturated with CO<sub>2</sub> was pumped using a peristaltic pump, as shown in Fig. 3.

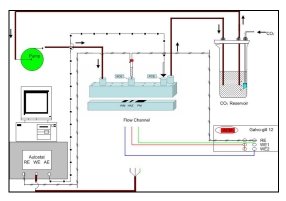


Fig. 3 Flow loop assembly

The areas of the exposed parent metal, heat affected zone and weld metal were in the ratio of 8:1:2 to represent the approximate proportions of each region in the vicinity of a typical pipeline weld.

The wall shear stress for the channel was calculated from (1) [12]:

$$\tau = f \frac{V^2}{8g} \tag{1}$$

where v is the flow velocity and f is the friction factor.

#### C. Electrochemical Measurements

The galvanic currents between each weld region were recorded every 60 s during the test using a multi-channel zero resistance ammeter (ACM Instruments GalvoGill 12) connected to a data logging PC. The currents from the parent metal to the weld metal and from the HAZ to the weld metal were recorded on two channels and, as the three electrodes were in the short-circuit condition, their individual galvanic currents were established from (2)

$$I_{PM} + I_{HAZ} + I_{WM} = 0 \tag{2}$$

## D. Experimental Conditions

The experiments were carried out at ambient temperature (20°C) in a flow channel containing artificial seawater as shown in Table I, saturated with CO2 with absence and presence of 30 ppm by volume of a 'green' oilfield corrosion inhibitor. The flow channel was equipped with a fixed rubber hose for inlet and outlet of flow, while the glass cell (reservoir) was filled at 0.8L capacity of 3.5% artificial seawater with chemical composition, as shown in Table I, and fitted with a suitable cover lid with 3 inlets to allow for the CO<sub>2</sub> injection as well as the discharge and suction of the electrolyte solution. Moreover, a peristaltic pump was used to maintain a uniform flow circulation and direction of 0.6 ms<sup>-1</sup>, which corresponded to a shear stress of 2.56 Pa. In one experimental run, galvanic currents were recorded at 60 s intervals throughout the test. Performances of CORRTREAT 05-193 and CORRTREAT CT04-020 [13] inhibitors as shown in Table II, supplied by Clariant Oil Services, UK were tested and compared in order to determine the more efficient inhibitor and whether they would provide enough corrosion protection to be used in real conditions.

TABLE I CHEMICAL COMPOSITION OF ARTIFICIAL SEAWATER Element Cr Ph Se Ca Si Mn 0.0002 0.0009 Ppm 114 0.008 2.9 0.0004 Р ν Zn S Element Co Fe 0.0066 0.009 0.0019 0.005 0.0005 0.00039 Ppm Note: ppm = parts per million

 TABLE II

 PRODUCTS COMPARED WITH GENERIC NAME AND BASIC CHEMISTRY

Product	Active Ingredient
Corrosion inhibitor A	Ethanediol, thioglycolic acid and
CT04-020	phosphoric acid ester
Corrosion inhibitor B	Neutralizing amines and synergists in a
05-193	water/glycol based solvent package.

## III. RESULTS

#### A. Measurements under Uninhibited Conditions

The galvanic current measurements for both the inhibitors (A and B) are shown in Fig. 4 under uninhibited conditions. The current behavior is observed to be similar, with the parent metal and the HAZ both being anodic (positive currents) with respect to the weld metal at test temperature. Clearly, it was

favorable for the weld metal to be the cathodic (negative currents) component in the couple as this condition ensured that its corrosion rate was reduced by sacrificial protection of the other weld regions.

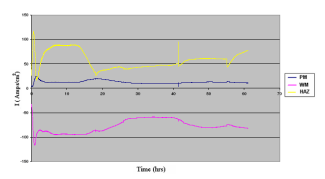
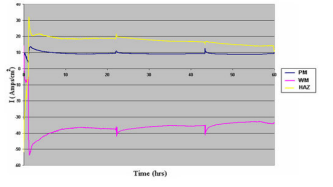
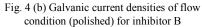


Fig. 4 (a) Galvanic current densities of flow condition (polished) for inhibitor A [13]





## B. Measurements under Inhibited Conditions

The presence of 30ppm of both oilfield corrosion inhibitors caused a marked change in the galvanic corrosion behavior, as shown in Fig 5. At the beginning of the exposure, both inhibitors appear to be unstable due to current reversal. However, after exposure for almost 20 hours, galvanic current for both inhibitors can be considered as stable since current reversal behavior disappeared.

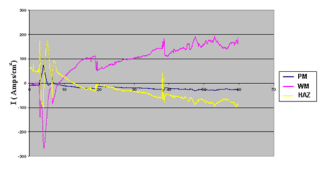


Fig. 5 (a) Galvanic current densities of flow condition using inhibitor (A)

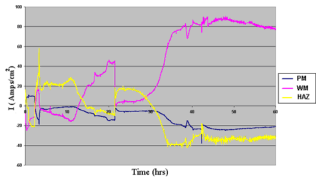


Fig. 5 (b) Galvanic current densities of flow condition using inhibitor (B)

In terms of galvanic currents, inhibitor B seems to be more efficient as it enables reduction of galvanic corrosion given that the values are quite small compared to the values obtained for inhibitor A. Besides, with inhibitor A, the peaks representing reversals reached high values (more than 100  $\mu$ A/cm<sup>2</sup> which is equivalent to more than 1 mm/y). This is not the case observed with inhibitor B.

The galvanic current for inhibitor A was particularly high on the weld metal than for inhibitor B, suggesting that the inhibitor A film was more adversely affected by flow causing preferential removal of the inhibitor film from the WM compared to inhibitor B.

After 20 hours of exposure, both the inhibitors had similar behavior, with the WM becoming anodic and both PM and HAZ becoming cathodic. Galvanic currents gradually rose for inhibitor A toward anodic side for the WM, while for inhibitor B, the WM remained anodic but the currents were fairly constant over the test period.

It would appear that, in this case, film removal was most pronounced on the WM for both inhibitors but was more severe with inhibitor A compared to inhibitor B. Clearly, for such current reversal to take place, further damage to the inhibitor film must have occurred. This situation is undesirable and would be expected to result in localized weld corrosion of the type that has sometimes been reported to occur in service.

## C. Measurements for Open Circuit Potential

The OCP values with a potential shift to positive values in the absence of inhibitors for the galvanic couples tested (parent metal and weld metal) are shown in Fig. 6. This variation of open circuit potential with time reveals the corrosivity of the galvanic couples tested. Initially the OCP for both parent metal and weld metal were recorded as -655mV and -500mV respectively, but after a certain time, both potentials move fast toward negative values around -720mVand reached a reasonably steady value as a result of the breakdown of the film inhibitor. It was noticed that both galvanic coupled potentials fluctuate continuously and many positive and negative peaks in high ranges are observed due to localized corrosion of both parent metal and weld metal.

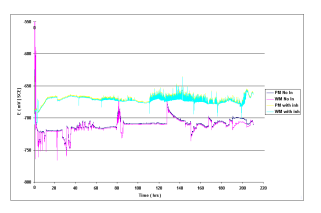


Fig. 6 Variation of the open-circuit potential as a function of time under uninhibited and inhibited conditions (Inhibitor B)

In contrast, with the presence of the inhibitor, the steady state potentials for the galvanic couples tested were shifted towards the noble direction and the potentials remained noiseless for the first 110 hrs, and then an increase was observed with reasonably steady fluctuating peaks in a small range as a result of the breakdown of the film inhibitor.

## IV. CONCLUSIONS

- In uninhibited conditions, the weld metal was cathodic to 1. the parent metal and HAZ. This is the ideal solution as corrosion is distributed over the large area of the parent metal.
- 2. The addition of two oilfield corrosion inhibitors caused a current reversal under flowing conditions. The inhibitor film was removed preferentially from the weld metal, so that it became strongly anodic and lead to a condition that would result in severe localized weld corrosion.
- 3. Inhibitor CORRTREAT 10-569 seems more efficient based on the galvanic current results since it displays stable behavior through the exposure time.
- 4 It appears that preferential weld corrosion is caused by unstable conditions in which the inhibitor film is selectively disrupted on the weld metal but remains effective on other weld regions.

## ACKNOWLEDGMENT

The authors would like to acknowledge Dr Mike Robinson for his extensive guidance and Cranfield University for use of their lab facilities.

#### REFERENCES

- [1] A. Ikeda, M. Ueda, and S. Mukai, "CO2 behavior of carbon and Cr steels", in Proc. NACE Corrosion/84, Houston, Texas, 1984.
- [2] X. Jiang, Y. G. Zheng, D. R. Qu, and W. Ke, "Effect of calcium ions on pitting corrosion and inhibition performance in CO2 corrosion of N80 steel", Corrosion Science, vol. 48, pp. 3091-3090, October 2006.
- [3] D. A. Lopez, S. N. Simison, and S. R. de Sanchez, "The influence of steel microstructure on CO2 corrosion. EIS studies on the inhibition efficiency of benzimidazole", Electrochimica Acta, vol. 48, pp. 845-854, February 2003.
- [4] M. B. Kermani, and A. Morshed, "Carbon dioxide corrosion in oil and gas production - a compendium", Corrosion, vol. 59, pp. 659-683, 2003.

- F. Farelas, and A. Ramirez, "Carbon Dioxide Corrosion Inhibition of [5] Carbon Steels Through Bis-imidazoline and Imidazoline Compounds Studied by EIS", Int. J. Electrochem. Sci., vol. 5, pp. 797-814, 2010. C. M. Lee, S. Bond, and P. Woollin, "Preferential weld corrosion:
- [6] effects of weldment microstructure and composition", in Proc. Corrosion/NACE International 2005, Houston, TX, USA, April 2005.
- [7] J. W. Palmer, W. Hedges, J. L. Dawson, "The Use of Corrosion Inhibitors in Oil & Gas Production", NACE International, TX, USA, 2004
- I. G. Winning, D. Mcnaughtan, A.J. McMahon and N. Bretherton, [8] "Evaluation of Weld Corrosion Behavior and the Application of Corrosion Inhibitors and Combined Scale/Corrosion Inhibitors", NACE International, New Orleans, Louisiana, 2004
- [9] E. Gulbrandsen and A. Dugstad, "Guidelines for Filler-Material Selection To Minimize Preferential Weld Corrosion in Pipeline Steels", in Proc. Corrosion 2005/NACE International, Houston, TX, USA, , 2005.
- [10] K. Fushimi, A. Naganuma, K. Azumi, Y. Kawahara, "Current distribution during galvanic corrosion of carbon steel welded with type-309 stainless steel in NaCl solution", Corrosion Science, vol. 50, pp. 903-911, March 2008.
- A. Chalmers, I. G. Winning, D. McNaughtan and S. McNeil, "Erosion-[11] Corrosion of Carbon Steel Pipework on an Offshore Oil", in Proc. Corrosion 2006, San Diego, CA, USA, 2006. K. Alawadhi, "Inhibition of Weld Corrosion in Flowing Brines
- [12] Containing Carbon Dioxide", PhD Thesis, Cranfield University, 2010.
- S. O. Otu, "Evaluation and performance of corrosion inhibitors for [13] down-hole protection of welded steels", MSc Thesis, Cranfield University, 2005.