Mechanism of Corrosion in Production Wells using Stainless Steel: Review

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Abstract: It is essential for the designers, engineers and field operators, who design and fabricate to be duly mindful of corrosion so as to enjoy undisturbed production, since most field materials are subjected to corrosion. This will prolong the commercial existence of production facilities. This paper provides an all-round review of challenges of corrosion while producing and proffers antidotes. The mechanism of corrosion was studied with the different kinds of corrosion that is experience in the Petroleum industry.

Keywords: Corrosion, Mechanism, Production, Wells, Steel, Rate

1. INTRODUCTION

Corrosion is a process that is natural which changes the state of a metal into the form that has chemical stability like as sulphide, hydroxide, or oxide. Its reaction looks like the one which takes place in a battery. Considering steel material in an acidic environment, at the anode, the iron dissolves and at the cathode, hydrogen is formed. An electrical path (e.g., water) is required.

\[
\text{Fe(s) + } 2\text{H}_2\text{O(l)} \rightarrow \text{Fe(OH)}_2(\text{aq}) + \text{H}_2(\text{g}).
\]

Figure 1: Mechanism of Corrosion and the environment it occurs (scib.azhar.live, gibsonstainless.com) and capex.com.ng.

Corrosion has damaging attack on a material by reacting with in the environment (Roberge, 2000) and an inherent automatic hazard connected with production and transportation facilities in the petroleum industry (Kermani and Smith, 1997). Corrosion possesses broad consequences on the integrity of materials used in the oil and gas sector. Such operations include exploration, development, production, transportation and refinery. Corrosion of steel takes place at relative humidity range of about 80% and temperature range of about 32°F (TERESA, 2013). Corrosion worsens in aggressive surroundings (i.e., Saline, Sulphide and/or acidic medium) with high temperatures and high pressures (HPHT) are situations to be confronted by materials in several production wells. As such, the facilities and control for corrosion used under such tough situations and places must be very dependable due to the high cost of maintenance or repair in inaccessible locations.

Types of Corrosion

Corrosion is grouped into eleven classes based on the nature of attack and the kind of surrounding of exposure.

Figure 1:
- Water: Freshwater, Saltwater, Acidic, Basic
- Iron: Iron Oxide
- Corrosion: Anode, Cathode

References:
- Roberge, 2000
- Kermani and Smith, 1997
- TERESA, 2013

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a) General or Uniform corrosion:
   It is the commonest kind of corrosion. Examples include rusting of: underground pipelines, submerged materials and steel bridges. Its control is mostly by painting the surface or using layers of sacrificial metal like steel and zinc.

![Figure 2: Instance of general or uniform corrosion (nace.org)](image)

b) Pitting Corrosion:
   It is a type of corrosion that occurs in carbon and low-alloy steels with scaling tendency at slow speed, CO₂ partial pressure and increasing temperatures of dew in oil or gas production well.

![Figure 3: Instances of Pitting Corrosion (webwormcpt)](image)

c) Erosion corrosion:
   It is a corrosion that takes place by removing the passive layer of products continuously from the walls of the pipe and its accessories (Popoola et al. 2013).

![Figure 4: Instance of Erosion corrosion (arab-oil-naturalgas.com)](image)

d) Stress corrosion Cracking:
   This is a type of corrosion that is localized causing cracks and breakages in metallic materials simultaneously from the action of tensile stress and corrosion itself.
e) Oxygen Corrosion:

It is a kind of corrosion that occur in productions well caused by fluids from casing, open hatches, leaking pump seals and process vents. Oxygen speeds up the damage of metal, anodically as a depolarizer and as an electron acceptor in cathodic reactions, (Weeter, 1965).

f) Galvanic Corrosion:

It is the kind of corrosion that occurs from the exposure of two different metallic materials that has differing electrochemical potential in an electrolytic medium. According to Nalli, 2010, the metallic material with the less or the most negative electrochemical potential becomes the anode and starts corroding.

g) Crevice Corrosion:

This is the type of corrosion which is localized and occurs in places that have narrow clearances, gap or crevices in the metal making room for fluid stagnancy. Roberge, 2000 posited that it is caused by the differences in concentration of corrodents over a metal surface.

h) Mesa Attack:

This is an unstable protective carbonate film formed from corrosion that takes place in fluid flow with low-to-medium. TERESA, 2013 confirmed that an exposed area operates as an anode of a galvanic cell while the film-covered areas operate as the cathode.
i) Hydrogen Embrittlement:
This is the type of corrosion that occur metallic alloys like low-alloy steels and carbon steel with atomic hydrogen which harshly depreciates the fracture resistance of its strength in solid solution especially during manufacturing operations and environmental exposure.

\[
\begin{align*}
H^+ (aq) + e^- & \rightarrow H \\
H + H & \rightarrow H_2 (g)
\end{align*}
\]

Figure 10: Instance of Hydrogen Embrittlement (boulonsplus.net and engineer-educators.com)

j) Sulphide Stress Cracking (Sour Corrosion):
This is a kind of corrosion that happens biologically in biogenic formations from the actions of microbes or sulphate reducing bacteria (SRB) which reduces existing sulphate ions to \(H_2S\) gas. TERESA, 2013 contributed that one of the source of bacterial contamination of aquifers are introduced through water injection that is not adequately treated.

k) CO₂ Corrosion (Sweet Corrosion):
It is a kind of corrosion which involves dissolution of CO₂ in an aqueous phase promoting reaction electrochemically between the steel and this contacting aqueous phase (Dugstad, 1992).
Controlling Corrosion

1. Application of protective coatings to metal surfaces to act as a barrier or perhaps provide sacrificial protection.
2. The addition of chemical species of inhibitors to the environment to inhibit corrosion, alteration of an alloy chemistry to make it more resistant to corrosion (Baojun, 2021).
3. The treatment of the surface of a metal to increase its resistance to corrosion (Barbara and Robert, 2006).
5. Cathodic protection technique (Zaki, 2006).

Selection of appropriate materials (Popoola et al., 2013).

Corrosion Rate

Corrosion rate is corrosion loss (the loss of the amount) of metal per unit time (year). It speeds up by impurities in the atmosphere, i.e. dissolution of metallic material in water (vaporized or condensed) and by dirt or dust which settles on metallic surfaces.

\[
\text{Corrosion Rate of Coupon (Mils per year penetration)} = \frac{(\text{Weight Loss in grams}) \times (22300)}{A \times dt}
\]

Where A is the area of coupon (Sq. in) and dt is change in time (year) (Moural et al., 2008).

Table 1: Recommended Materials in the Oil and Gas industry

<table>
<thead>
<tr>
<th>Material specification</th>
<th>Oil and gas application</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Nickel-iron (incolys) - Alloys of Ni-Fe-Cr</td>
<td>Used in flow lines, manifolds, wellheads, liners, hangers and production tubings that possess high temperature and sour application</td>
</tr>
<tr>
<td>2. Low- and medium-alloy steels</td>
<td></td>
</tr>
<tr>
<td>3. Nickel-chrome (inconels) - alloys of Ni-Cr-Fe</td>
<td></td>
</tr>
<tr>
<td>Nickel steels between (2.5 - 9%) Ni.</td>
<td>Used in pipings, pumps and storage tanks of liquefied natural gas</td>
</tr>
<tr>
<td>Straight chromium steels between (12 - 18%) Cr.</td>
<td>Used in casing pipes, wellheads Christmas trees, valves and downhole rods.</td>
</tr>
<tr>
<td>Duplex stainless steels for:</td>
<td>Used where extreme the level of chlorides is active like pipings, internals of tanks and vessel.</td>
</tr>
<tr>
<td>1. 22% chromium duplex,</td>
<td></td>
</tr>
<tr>
<td>2. 25% chromium super - duplex (Falkland et al., 2011)</td>
<td></td>
</tr>
</tbody>
</table>
**II. LITERATURE REVIEW**

The overall yearly cost of corrosion in the Petroleum production industry is estimated to be US$1.372 billion, broken down into $589 million in surface pipeline and facility costs, $463 million yearly in downhole tubing expenses, and another $320 million in capital expenditures associated to corrosion (Simmons, 2008). The most prevalent form of corrosion in the petroleum industry happens when steel comes in contact with an aqueous medium environment and rusts (Corbin and Willson, 2007). Corrosion that is flow-induced are connected to velocities of high-flow and turbulence. Also, local turbulence can be produced by pits and mesa attack zones with particular geometries. This turbulence, in return, may damage the protective scales that exist and also prevent fresh formation on the exposed metallic surface (TERESA, 2013). Other catastrophic incidences resulting from corrosion failure had been historically recorded. (Popoola et al., 2013). The control for corrosion CO₂ corrosion has been performed by common techniques such as inhibitor injection, use of corrosion resistance alloy and plastic resin coating (Masakatsu, Hideki and Perry, 2000). Acid-corrosion results from air-oxidation of iron (II) in the anode paths, in the absence of OH⁻ ions which results in strongly acidic solutions with pH of 1.41(Tamura, 2008). Currently, there is a very high emphasis on the development of advanced and workable coatings for corrosion protection in various applications of technological (Montemor, 2014).

**III. MATERIAL AND METHODS**

This study targets the mechanism of the different types of corrosion that occur in well of oil and gas production, since this has been a deleterious happens continuously. The review method was applied in this paper.

**IV. DATA AND METHODOLOGY**

Data Review and Analysis

<table>
<thead>
<tr>
<th>Corrosion (mm/year)</th>
<th>Rate (mm/year)</th>
<th>Time (hour)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.9</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>1.8</td>
<td>0.1</td>
<td></td>
</tr>
<tr>
<td>0.9</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>1.7</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>0.4</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>0.3</td>
<td>5</td>
<td></td>
</tr>
</tbody>
</table>

Table 2: Experimental data taken from Nesic and Lee, 2003 illustrating the experimentally measured corrosion rates showing the effect of formation protective scale using the test conditions of temperature at 80OC, pH of 6.6, Partial pressure of CO₂ of 0.54bar, Concentration of Fe²⁺ of 250ppm and Speed of 1m/s.
Table 4: Data taken from Wang, George and Nesic, 2004 illustrating the experimentally measured corrosion rates showing the effect of the partial pressure of CO2 using the test conditions of temperature at 60°C, pH of 5 and Speed of 1m/s.

<table>
<thead>
<tr>
<th>CORROSION RATE (MM/YEAR)</th>
<th>CO₂ PARTIAL PRESSURE (BAR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>3</td>
</tr>
<tr>
<td>20</td>
<td>10</td>
</tr>
<tr>
<td>30</td>
<td>20</td>
</tr>
<tr>
<td>Regression</td>
<td>0.9897</td>
</tr>
</tbody>
</table>

Figure 15: Corrosion rate versus pH

Table 5: Data taken from Nesic, Solvi and Enerhaug, 1995 illustrating the experimentally measured corrosion rates showing the effect of speed in the absence of the scale of iron carbonate using test conditions of temperature at 20°C, Partial Pressure of CO₂ = 1bar, Concentration of Fe²⁺ is less than 2ppm.

<table>
<thead>
<tr>
<th>CORROSION RATE (MM/YEAR)</th>
<th>SPEED (M/S)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>1.3</td>
<td>2</td>
</tr>
<tr>
<td>2.2</td>
<td>4</td>
</tr>
<tr>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>3.2</td>
<td>10</td>
</tr>
<tr>
<td>Regression</td>
<td>0.9517</td>
</tr>
</tbody>
</table>

Figure 16: Corrosion rate versus CO₂ Partial Pressure

Table 6: Data taken from George, Nesic and deWaard, 2004 illustrating the experimentally measured corrosion rates showing the effect of acetic acid using test conditions of temperature at 60°C, pH of 4, and Speed of 0.5m/s.

<table>
<thead>
<tr>
<th>Corrosion Rate (mm/year)</th>
<th>Rate</th>
<th>Acetic Acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>100</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>Regression</td>
<td>0.3324</td>
<td></td>
</tr>
</tbody>
</table>

Figure 17: Corrosion rate versus Speed

Figure 18: Corrosion rate versus Concentration of acetic acid
Table 8: Summary of Regression Results for all parameter plotted against Corrosion Rate

<table>
<thead>
<tr>
<th>S/No.</th>
<th>Corrosion Rate (Mm/Year)</th>
<th>Regression (R²)</th>
<th>Equations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Versus Time (Hour)</td>
<td>0.9475</td>
<td>y = 1.9992e^{0.035x}</td>
</tr>
<tr>
<td>2.</td>
<td>Versus pH</td>
<td>0.8902</td>
<td>y = 21.75e^{0.035x}</td>
</tr>
<tr>
<td>3.</td>
<td>Versus CO₂ Partial Pressure (Bar)</td>
<td>0.9897</td>
<td>y = 11.415ln(x) - 5.0076</td>
</tr>
<tr>
<td>4.</td>
<td>Versus Speed (m/s)</td>
<td>0.9517</td>
<td>y = 1.0581ln(x) + 0.8335</td>
</tr>
<tr>
<td>5.</td>
<td>Versus Acetic Acid Concentration</td>
<td>0.3324</td>
<td>y = 0.0302x + 7.2253</td>
</tr>
</tbody>
</table>

V. RESULTS AND DISCUSSION

In figure 14, the corrosion rate was considered against time. The corrosion rate down-surge with time from onset of 1.9mm/yr to 0.9mm/yr for 1.5 hours. This value increased again and ran up to 1.7mm/yr for another 0.5 years. This subsequently fell straight to 0.4mm/yr for 1 hour. There was a slight increase to 0.3mm/yr another 1 hour. The trend later continued to nose-dive until corrosion rate disappeared totally.

In figure 15, the corrosion rate was considered against pH. The corrosion rate nose-dive proportionally from 2.5mm/yr to 1.0mm/yr in the acidic medium of 4 to 5. This curved up closely to a neutral point of 6 medium values.

In figure 16, the corrosion rate was considered against CO₂ partial pressure. The corrosion rate sky-rocketed proportionally from 9mm/yr to 20mm/yr with 2.5 bar to 10 bar. This further trend till 30mm/year for another 10 bar.

In figure 17, the corrosion rate was considered against speed. The corrosion rate rose very fast proportionately from 1mm/yr to 3mm/yr for speed of 1m/s to 6m/s. This further increased till 3.2mm/yr for another 4m/s.

In figure 18, the corrosion rate was considered against the concentration of acetic acid. The corrosion rate lapped on 0mm/yr for a while before nudging out to 10mm/yr and further decreasing to 9mm/yr in acetic acid concentration.

Also considering table 8, the correlation regression portends threatened conditions from corrosion rate with time. The conditions worsen considering pH conditions and the imposed partial pressure of CO₂. It is also observed that the corrosion rate was also very fast.

VI. CONCLUSION AND RECOMMENDATION

Since corrosion is aggressive in all the selected conditions, there is need for rapid arrest before it goes from been destructive to becoming unmanageable and a disaster. The corrosion control suggested in this paper can be adopted.

REFERENCES

12. Implementation of a New Alloyed Steel for Oil and Gas Production Wells, NACE-00154.