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Studies of Aqueous Hydrogen Sulfide Corrosion in Producing SAGD Wells

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Studies of Aqueous Hydrogen Sulfide Corrosion in Producing SAGD Wells

by

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A THESIS
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Abstract

In this research lab and field corrosion coupon testing was completed to determine corrosion rates on relevant Steam Assisted Gravity Drainage (SAGD) metals. Evaluating how rates varied with well depth, environment, and a dominating corrosion mechanism.


Corrosion rates decreased from 0.0178mm/y at the wells bottom to 0.0145mm/y at higher elevations. This corresponded to a decrease in iron sulfide (FeS) scale content from the well bottom upwards formed through a solid-state reaction between aqueous H2S and the metal. High average corrosion rates of 0.263mm/y were measured in the lab, compared to the low 0.0183mm/y in field studies. This difference is due to the inhibiting effects of oil in the field which inhibits corrosion rates and the longer test duration.
Acknowledgements

I would like to express my sincere gratitude to my respected supervisor, Dr. Frank Cheng, for his instruction, encouragement and support throughout my whole research career here at the University of Calgary. His enthusiasm, diligence, perception and professional attitude to science and research have been inspiring and guiding me to move forwards in my future life and career. This work was also supported by Suncor Energy, Cormetrics Ltd, Corr Science, and the University of Calgary. Their support and cooperation has been greatly appreciated through the research process.

Appreciations are also worth extending to the members in my group, Dr. Qiang Li, Yuanchao Feng, Yuan Li, Shan Qian, and Yao Yang for their unforgettable support and generous help in this work.
Dedication

This work is dedicated to my parents, as well as Leon Fedenczuk, Kristina Hoffmann, and Hugo Acosta-Ramirez for pushing me to pursue graduate studies and their support throughout.
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<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>AP</td>
<td>Anodic Protection</td>
</tr>
<tr>
<td>BHP</td>
<td>Bottom Hole Pressure</td>
</tr>
<tr>
<td>(c_{b,H_2S}, c_{s,H_2S})</td>
<td>(H_2S) concentrations in the bulk solution and locally at the steels surface respectively, (\text{mol/m}^3)</td>
</tr>
<tr>
<td>CP</td>
<td>Cathodic Protection</td>
</tr>
<tr>
<td>CPP</td>
<td>Cyclic Potentiodynamic Polarization</td>
</tr>
<tr>
<td>CRA</td>
<td>Corrosion Resistant Alloy</td>
</tr>
<tr>
<td>(CR_{H_2S})</td>
<td>Corrosion rate as (\text{mol/m}^2\text{s})</td>
</tr>
<tr>
<td>CS</td>
<td>Carbon Steel</td>
</tr>
<tr>
<td>(D_{(H_2S)})</td>
<td>Diffusion transfer coefficient of (H_2S) in water</td>
</tr>
<tr>
<td>(E_{\text{prot}})</td>
<td>Re-Passivation Potential</td>
</tr>
<tr>
<td>(E_{\text{Pit}})</td>
<td>Pitting Potential</td>
</tr>
<tr>
<td>ESP</td>
<td>Electric Submersible Pump</td>
</tr>
<tr>
<td>GLV</td>
<td>Galvanized Coating</td>
</tr>
<tr>
<td>HE</td>
<td>Hydrogen Embrittlement</td>
</tr>
<tr>
<td>HIC</td>
<td>Hydrogen Induced Cracking</td>
</tr>
<tr>
<td>HTHP</td>
<td>High Temperature High Pressure</td>
</tr>
<tr>
<td>(k_m(H_2S))</td>
<td>Mass transfer coefficient of (H_2S) in liquid boundary layer in (\text{m/s})</td>
</tr>
<tr>
<td>LPR</td>
<td>Linear Polarization Resistance</td>
</tr>
<tr>
<td>mKB</td>
<td>Meters Kelly Bushing</td>
</tr>
<tr>
<td>mm/y</td>
<td>Millimeter per year</td>
</tr>
<tr>
<td>MPY</td>
<td>Mils per year</td>
</tr>
<tr>
<td>OCP ((E_{\text{OC}}))</td>
<td>Open Circuit Potential</td>
</tr>
<tr>
<td>(P_{\text{CO}_2})</td>
<td>(CO_2) Partial Pressure</td>
</tr>
<tr>
<td>(P_{H_2S})</td>
<td>(H_2S) Partial Pressure</td>
</tr>
<tr>
<td>PP</td>
<td>Potentiostatic Polarization</td>
</tr>
<tr>
<td>ppmv</td>
<td>parts per million by volume</td>
</tr>
<tr>
<td>PTFE</td>
<td>Polytetrafluoroethylene</td>
</tr>
<tr>
<td>SAGD</td>
<td>Steam Assisted Gravity Drainage</td>
</tr>
<tr>
<td>SCC</td>
<td>Stress Corrosion Cracking</td>
</tr>
<tr>
<td>TSR</td>
<td>Thermochemical Sulphate Reduction</td>
</tr>
<tr>
<td>VCI</td>
<td>Volatile Corrosion Inhibitors</td>
</tr>
<tr>
<td>WL</td>
<td>Weight Loss</td>
</tr>
<tr>
<td>ZRA</td>
<td>Zero Resistance Ammeter</td>
</tr>
<tr>
<td>(\varphi)</td>
<td>Tortuosity of outer film</td>
</tr>
<tr>
<td>(\varepsilon')</td>
<td>Porosity of outer film</td>
</tr>
<tr>
<td>(\delta_{OS})</td>
<td>Outer film thickness, (\text{m})</td>
</tr>
</tbody>
</table>
Chapter One: Introduction

1.1 SAGD background

SAGD is a high temperature oil recovery technique used to produce bitumen from relatively deep, usually greater than 200 m [1], heavy oil sands deposits. SAGD is a relatively new and advanced form of oil recovery, where a pair of horizontal wells are drilled into the reservoir with a steam injection well drilled only a few meters above a second production well [2]. Steam is injected into the upper well at temperatures at or above 240°C to heat the bitumen and reduce its viscosity from approximately being close to that of a solid (>1,000,000 cp) down to ~20 cp so that it can drain and be produced economically using one of a variety of artificial lift methods. Figure 1.1 shows a typical SAGD drilling and production description diagram.

Figure 1.1 SAGD well pair diagram
With the first full scale SAGD asset having now produced for only 10-15 years, this is a relatively new method of producing oil compared to conventional oil and gas assets which have been producing for 60+ years. As SAGD assets begin to age, downhole well corrosion, well integrity concerns, and other issues are being found and discussed. Well integrity and corrosion concerns can be an issue at several locations within a well, including internally and externally on tubing and casing strings as well as on wellhead components. Downhole completion and tubular materials used in SAGD are similar to those used in conventional oil and gas assets. Materials used include a variety of carbon steels, metal alloys, and corrosion resistant alloy’s (CRA’s). The advantage and reason many operations use carbon steel is due to their lower costs. The disadvantage of using standard carbon steel tubulars is that they are susceptible to corrosion, specifically corrosion caused by aqueous acidic gases.

In SAGD, a process known as aquathermolysis [3, 4] takes place and leads to the formation of aqueous acidic gases, including H₂S and CO₂, both of which are known to be highly corrosive when dissolved in water. Such conditions can be present in SAGD wells, specifically in the annulus of production wells where these gases are able to concentrate in a low flowing or stale environment.

The combination of exposures to aqueous acidic gases, elevated temperatures (>200°C), and relatively high pressures (>2000 kPa) are known to accelerate localized and general corrosion. Corrosion is likely initiated and accelerated by the presence of galvanic and crevice corrosion cells formed between components. Corrosion control measures commonly used in the oil and gas industry include promoting the formation of passive scales, corrosion inhibitors, upgraded CRA materials, and altering environmental
conditions. Given the relatively new application and use of SAGD, corrosion in this type of producing environment is not a well understood issue, and limited research has been completed to date to understand the main corrosion mechanisms, corrosion rates, and how to prevent it. The results presented in this thesis are based on the completion types and operating conditions present at Suncor Energy’s Firebag SAGD asset.

1.2 SAGD well completion diagram information

Figure 1.2 shows an image of a typical SAGD well completion at Firebag. Red indicates the production tubing where the emulsion is produced to surface. The green area is the annulus between the production tubing and production casing. The blue line is the instrumentation (pressure measurement and electrical power lines)

![Figure 1.2: Typical Firebag production well completion diagram](image)

In this completion, an electric submersible pump (ESP) is used to artificially lift fluids from the reservoir to surface for further processing in the central plant. Alternative
artificial lift methods include a process known as gas lift which utilizes injecting an inert gas to lift fluids to the surface by reducing the downhole fluid density and the pressure required to lift the fluid to surface. Inert gases commonly used are methane or natural gas.

Tubular and completion components and materials commonly used in SAGD are shown in Table 1.1:

Table 1.1: Completion materials and locations of use [5-8]

<table>
<thead>
<tr>
<th>Material</th>
<th>Location of Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>J-55/K-55/TN-55TH/L-80</td>
<td>Casing and Tubing</td>
</tr>
<tr>
<td>Galvanized J-55 and/or Galvanized K-55</td>
<td>Clamps and Cables</td>
</tr>
<tr>
<td>Deloro-40 HF</td>
<td>Gate Valve Gate Face</td>
</tr>
<tr>
<td>Stellite-6HF</td>
<td>Gate Valve Gate Seat</td>
</tr>
<tr>
<td>1018 Carbon Steel</td>
<td>General use</td>
</tr>
</tbody>
</table>

1.3 SAGD produced fluid and gas information

Table 1.2 and Table 1.3 list the typical produced fluid and gas compositions from a SAGD well during oil production [9, 10].

Table 1.2: Average Firebag emulsion composition

<table>
<thead>
<tr>
<th>Component</th>
<th>Relative Concentration (mass %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>50-95</td>
</tr>
<tr>
<td>Oil</td>
<td>5-30</td>
</tr>
<tr>
<td>Gas</td>
<td>1-15</td>
</tr>
<tr>
<td>Solids</td>
<td>&lt;5</td>
</tr>
</tbody>
</table>
Table 1.3: Average Firebag produced gas composition

<table>
<thead>
<tr>
<th>Compound</th>
<th>Concentration (mol fraction %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH$_4$</td>
<td>40-70</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>35-55</td>
</tr>
<tr>
<td>H$_2$S</td>
<td>1-15</td>
</tr>
<tr>
<td>N</td>
<td>&lt;1.5</td>
</tr>
<tr>
<td>H</td>
<td>&lt;1</td>
</tr>
</tbody>
</table>

As can be seen, the emulsion consists mostly of water with lesser amounts of oil, gas, and solids, which are typically sands produced from the reservoir. The produced water is a combination of condensed water from the re-produced steam as well as formation (connate) water already in place. Produced water samples have shown to be non-saline, salt free, with limited amount of chlorides present. Both salt and chlorides are known to accelerate corrosion of metals. Produced gases from the reservoir consist mostly of methane (CH$_4$), relatively high concentrations of CO$_2$, and in the case of Suncor’s Firebag Asset, higher concentrations of H$_2$S. Lesser amount of nitrogen (N$_2$) and hydrogen (H$_2$) are also produced. The ratio and quantities of these gases can vary from site to site depending on reservoir properties, production conditions, and asset age [4].

Aqueous CO$_2$ and H$_2$S are produced from oil and gas fields around the world and are known to cause corrosion related completion, facility, and reliability issues. This type of corrosion has been well studied in conventional oil and gas assets and has led to numerous material, corrosion inhibitor, and production operation advancements to prevent and mitigate the corrosion process.

The corrosive environment in producing SAGD assets is different and unique from most conventional assets. This is due to the combination of high temperatures, relatively
high pressures, the relative quantities of CO₂ and H₂S, and the completion designs used. For this reason, further studies are required to understand the potential for corrosion.

1.4 Research objectives

The objective of this research is to better understand the corrosion rates present in the annulus of producing SAGD wells through lab and field tests, and to determine the likely corrosion mechanisms in order to evaluate the severity of corrosion in SAGD wells. Progress has been made in the following areas:

- The potential corrosion rates in the annulus of producing SAGD wells on common completion materials. Materials include carbon steel, J-55, K-55, TH-55TH, and galvanized materials, and different hard facing overlay finishes
- A better understanding of the corrosion product characteristics and composition
- Changes in corrosion product composition with well depth
- Changes in corrosion rates at different well depths
- Relating findings and results to known corrosion mechanisms to recommend mitigating strategies and measures

1.5 Structure of thesis

The thesis contains seven chapters, with Chapter 1 giving an introduction of the research background and objectives.

Chapter 2 is a comprehensive review of literature relevant to this research. This includes a discussion on aqueous CO₂ and H₂S corrosion mechanisms, corrosion rates,
scale products, and a review of relevant High Temperature High Pressure (HTHP) literature in combined CO₂ and H₂S gaseous systems.

Chapter 3 outlines the lab and field experimental setups, including material specimens used and the preparation of the material samples and solutions. This includes the autoclave testing parameters, field production conditions, measurement techniques and analysis methods used.

Chapter 4 presents the lab and field testing results on materials commonly used in SAGD production wells, including the corrosion rates and scale products in varied corrosion scenarios.

Chapter 5 outlines the possible corrosion mechanism present to better understand how to possibly predict and prevent corrosion.

Chapter 6 presents conclusions and recommendations based on the research completed.
Chapter Two: Literature review

2.1 A brief introduction to corrosion during oil production

In 2016 a NACE Impact study was completed, which estimated the annual global cost of corrosion to be US$2.5 trillion, of which approximately $5.6B can be attributed to the oil and gas industry [11]. Corrosion is present throughout the production, processing, and consumption cycles of hydrocarbon products. This includes corrosion in downhole wells, processing facilities, refineries, pipelines, and shipping/distribution assets throughout their life cycles from initial construction to final asset abandonment and many years afterwards. While preventing and eliminating corrosion can be very difficult and costly, controlling the corrosion rate through improved products, facility design, and chemicals are the common and economical options to delay the process to a manageable level compared to eliminating it. Corrosion engineers are therefore increasingly involved in designing corrosion prevention systems to estimate their costs and the useful life of equipment. [11].

2.1.1 Corrosion as a key mechanism resulting in well failures during oil production

It is integral that producing oil and gas companies take the steps and prevention measures to ensure that well integrity is maintained to prevent well failures throughout the lifecycle of an asset. One of the most common definitions of well integrity is given by NORSOK D-010 as the “application of technical, operational and organizational solutions to reduce risk of uncontrolled release of formation fluids throughout the life cycle of a well” [12]. Uncontrolled releases of a fluid from a well can occur for several
reasons and happen below or above ground level. Well failure mechanisms leading to uncontrolled releases can be initiated and occur due to operating or maintenance procedures not being followed, improper well design and construction, and corrosion. Historical rates of well “failures” in oil and gas fields can vary from a few percent of wells to >40% depending on the wells operating conditions and procedures followed [13, 14].

Corrosion is a leading cause of failure as it can attack every component in a well at all stages of its life cycle. This includes all downhole completion components and tubulars from the time a well is initially drilled through to its final abandonment. Corrosion is a common cause of well failures due to the number of initiating and propagation mechanisms that exist and the extreme conditions present. These include the presence of oxygen, acidic gases, water, solids, microbes, varied materials and stresses, chemicals, and elevated temperatures and pressures. All of which are known to initiate and accelerate corrosion.

2.1.2 Primary corrosion types

There are 8 known general forms of corrosion. In the oil and gas industry all forms can be found throughout the production to consumption process of oil products [15]. These main forms of corrosion are discussed next along with a brief description of the mechanism and their effects in oil wells and assets. The 8 forms are uniform, galvanic, crevice, pitting, intergranular, de-alloying, erosion, and stress corrosion cracking.
Uniform corrosion, or general corrosion, is defined as a type of corrosion that is uniformly distributed over the entire exposed surface of a metal. Uniform corrosion also refers to the corrosion that proceeds at approximately the same rate over the exposed metal surface without appreciable localization. It is recognized by a roughening of the surface and usually by the presence of corrosion products. The mechanism of the attack typically is an electrochemical process that takes place on the surface of the material. Differences in composition or orientation between small areas on the metal surface create local anodes and cathodes that facilitate the corrosion process. The anodic reaction in the corrosion process is always the oxidation reaction (M – “Metal”):

\[ M \leftrightarrow M^+ + e^- \]  \hspace{1cm} (2-1)

In acidic environments, i.e., pH < 7, the cathodic process is mainly the reduction of hydrogen ions:

\[ 2H^+ + 2e \leftrightarrow H_2 \]  \hspace{1cm} (2-2)

In alkaline and neutral, environments with a pH \( \geq 7 \), the reduction of dissolved oxygen is the predominant cathodic process:

\[ O_2 + 2H_2O + 4e \leftrightarrow 4OH^- \]  \hspace{1cm} (2-3)

With uniform distribution of cathodic reactants over the entire exposed metal surface, reactions (2-2) and/or (2-3) take place in a "uniform" manner and there is no preferential site or location for cathodic or anodic reaction. The cathodes and anodes are located randomly and alternating with time. This results in a relatively uniform loss of thickness of the metals [16]. Uniform corrosion is most often caused by an inappropriate material selected for use in a corrosive environment. General corrosion can be tolerated because the effect of metal loss is relatively easy to assess and allowances can be made in
the initial design. Protective coatings and/or cathodic protection (CP) are particularly
effective in controlling uniform corrosion. Such protection methods, individually or in
combination, are commonly used to protect above and below ground oil and gas pipelines
from uniform corrosion.

*Galvanic corrosion* occurs due to the electrical potential difference that exists
when dissimilar metals come into contact in a conductive medium, i.e., electrolyte. One
metal acts as the anode and the other as the cathode. The electrolyte provides a means for
ion migration whereby metallic ions move from the anode to the cathode. The anode
metal dissolves into the electrolyte, and deposit collects on the cathodic metal.

*Crevice corrosion* refers to the localized attack on a metal surface at, or
immediately adjacent to, the gap or crevice between two joining surfaces. The gap or
crevise can be formed between two metals or a metal and non-metallic material. Outside
the gap or without the gap, both metals are resistant to corrosion. The damage caused by
crevise corrosion is normally confined to one metal at a localized area within or close to
the joining surfaces.

Crevice corrosion is initiated by a difference in concentration of chemical
constituents, often oxygen, which set up an electrochemical concentration cell. In the
case of oxygen this is a differential aeration cell. Outside of the crevice, at the cathode,
the oxygen content and the pH are higher, but chlorides are lower. Chlorides are also able
to concentrate inside the crevice, at the anode, worsening the situation. Ferrous ions form
ferric chloride and attack stainless steels rapidly. The pH and the oxygen content are
lower in the crevice than in the bulk water solution, just as they are inside a pit. The pH
inside the crevice may be as low as 2 when the bulk solution is neutral. Once a crevice
has formed, the propagation mechanism for crevice corrosion is the same as for pitting
corrosion. The major factors influencing crevice corrosion are:

- Crevice type: metal-to-metal, metal-to-non-metal
- Crevice geometry: gap size, depth, surface roughness
- Material: alloy composition (e.g. Cr, Mo), structure
- Environment: pH, temperature, halide ions, oxygen

_Pitting corrosion_ is a form of extremely localized corrosion attack that results in
holes in the metal which are commonly found in oil and gas operations. A pitting
corrosion example is in Figure 2.1 from a field failure inspection on a section of
galvanized power cable.

*Figure 2.1: Magnified cable pitting corrosion example (10X magnification) [17]*

These holes may be small or large in diameter but in most cases, they are relatively
small. Pits are sometimes isolated or so close together that they look like a rough surface.
Generally, a pit may be described as a cavity or hole with the surface diameter about the same as or less than the depth. Pitting is one of the most destructive and insidious forms of corrosion. It is often difficult to detect pits because of their small size and because the pits are often covered with corrosion products. Pitting is particularly aggressive because it is a localized and intensive form of corrosion, and failures often occur with extreme suddenness. Pitting corrosion can be initiated several ways including:

- Localized chemical or mechanical damage to a protective passive film
- Water chemistry factors which can cause breakdown of a passive film are acidity, low dissolved oxygen concentrations (which reduce the protectiveness of oxide films) and high concentrations of chloride [18]
- Localized damage to, or poor application of, a protective coating
- The presence of non-uniformities in the metal structure of the component, e.g. non-metallic inclusions

*Intergranular corrosion* is localized attack along grain boundaries, or immediately adjacent to grain boundaries, while the bulk of the grains remain largely unaffected. The microstructure of metals and alloys is made up of grains, separated by grain boundaries.

Grain boundary effects are of little or no consequence in most applications or uses of metals. If a metal corrodes, uniform attack occurs since grain boundaries are usually only slightly more reactive than the matrix. However, under certain conditions, grain interfaces are very reactive and intergranular corrosion is caused. Intergranular corrosion attack is usually related to the segregation of specific elements or the formation of a
compound in the boundary. Corrosion then occurs by preferential attack on the grain-boundary phase, or in a zone adjacent to it that has lost an element necessary for adequate corrosion resistance - thus making the grain boundary zone anodic relative to the remainder of the surface. The attack usually progresses along the narrow grain boundary and, in a severe case of grain-boundary corrosion, entire grains may be dislodged due to complete deterioration of their boundaries.

*Selective leaching* is the removal of one element from a solid alloy by corrosion processes. The most common example is the selective removal of zinc in brass alloys (dezincification). Similar processes occur in other alloy systems in which aluminum, iron, cobalt, chromium, and other elements are removed. Selective leaching is the general term to describe these processes [19, 20].

*Erosion-corrosion* is the acceleration or increase in rate of deterioration or attack on a metal because of relative movement between a corrosive fluid and the metal surface. Generally, this movement is quite rapid, and mechanical wear effects or abrasion are involved. Metal is removed from the surface as dissolved ions, or it forms solid corrosion products which are mechanically swept from the metal surface. The increased turbulence caused by pitting on the internal surfaces of a tube can result in rapidly increasing erosion rates and eventually a leak.

Materials selection plays an important role in minimizing erosion corrosion damage. Caution is in order when predicting erosion corrosion behavior based on hardness. High hardness in a material does not necessarily guarantee a high degree of resistance to erosion corrosion.
Erosion corrosion is characterized in appearance by grooves, gullies, waves, rounded holes, and valleys, and usually exhibits a directional pattern. In many cases, failures caused by erosion corrosion occur in a relatively short time, and they are unexpected largely because corrosion evaluation testing is completed under static conditions or because the erosion effects were not considered. Prevention methods include increasing material thickness in areas prone to erosion, improved materials, removing particles and solids, and improving system design to reduce fluid velocity as well as preventing turbulent flow.

*Stress-corrosion cracking* (SCC) refers to material cracking caused by the simultaneous presence of a tensile stress, a specific corrosive medium, and a susceptible material. The stresses that cause SCC can be produced as a result from externally applied stresses or from residual stresses introduced during manufacturing. The stresses required to cause SCC are typically small and well below the materials yield strength in an aggressive aqueous environment. Cracks often initiate at pre-existing surface flaws/discontinuities, corrosion pits, or dislocation slip lines and steps. Crack propagation is typically intergranular or trans-granular in nature but it can also be a combination of the two. Examples of each are shown in Figure 2.2 [19].
During SCC, the metal or alloy is virtually unattacked over most of its surface while fine cracks progress through it. This cracking phenomenon has serious consequences since it can occur at stresses within the range of typical design stress.

2.1.3 Main corrosion control measures

Significant measures are taken to prevent, limit, and delay corrosion. These include: corrosion inhibition, coating applications, improving or altering materials, cathodic and anodic protection system implementation, and by altering and improving the environment and conditions present.

A corrosion inhibitor is a substance which, when added in a small concentration to an environment, reduces the corrosion rate of a metal exposed to that environment. Inhibitors often play an important role in the oil and gas industries throughout extraction.
and processing. Here, and in many other industries, they have been considered to be the first line of defence against corrosion due to the extreme environments and processes required as well as material limitations. Inhibitors often work by adsorbing themselves on the metallic surface and protecting the metallic surface by forming a film. Inhibitors are normally distributed and dispersed from a solution but they can also be included during a protective coatings formulation. Inhibitors slow corrosion processes in one of three ways:

- Increasing the anodic or cathodic polarization behavior
- Reducing the movement or diffusion of ions to the metallic surface
- Increasing the electrical resistance of the metallic surface

**Anti-corrosion coatings** are primarily applied for the protection of solid metal surfaces, and occasionally some non-metals. The main functions of surface coatings are to act as an insulating barrier that eliminates the likelihood of contact between a substrate and electrolytes in the working environment, thus preventing the likely electrochemical reaction that can cause corrosion. When selecting a suitable corrosion protection system one must consider the nature of the corrosive environment faced by the assets to be protected, such as pipelines, structures, vehicles, ships, bridges, plant equipment, and buildings as well as the environment in which it is located. Desirable coating characteristics include: effective electrical insulator and moisture barrier, its ability to resist holiday development, good adhesion to pipe and resistance to dis-bonding, ease of repair, good hardness/abrasion, soil stressing, and penetration resistance.

In selecting an appropriate material for corrosion control, the material’s corrosion resistance is an important property to consider. The materials most resistant to
corrosion are those for which corrosion is thermodynamically unfavorable. An optimum material is the least expensive material that can provide the performance which is safe to install and operate, suitable to the application, and economic. To select a material, it is best to review past materials, specific test and material data, and industry experiences. It is also important to consider passive metals, non-metallic metals, and materials close together in the galvanic series when designing a system. Commonly used materials in specific environments are shown in Table 2.1.

Table 2.1: Common corrosive environments and corrosion resistant material

<table>
<thead>
<tr>
<th>Number</th>
<th>Environment</th>
<th>Material</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Nitric Acid</td>
<td>Stainless Steels</td>
</tr>
<tr>
<td>2</td>
<td>Caustic (high pH)</td>
<td>Nickel and Nickel Alloys</td>
</tr>
<tr>
<td>3</td>
<td>Hydrofluoric Acid</td>
<td>Monel (Ni-Cu)</td>
</tr>
<tr>
<td>4</td>
<td>Hot Hydrochloric Acid</td>
<td>Hastelloys (Ni-Cr-Mo)</td>
</tr>
<tr>
<td>5</td>
<td>Dilute Sulfuric Acid</td>
<td>Lead</td>
</tr>
<tr>
<td>6</td>
<td>Non-staining Atm Exposure</td>
<td>Aluminum</td>
</tr>
<tr>
<td>7</td>
<td>Distilled Water</td>
<td>Tin</td>
</tr>
<tr>
<td>8</td>
<td>Hot Strong Oxidizing Solution</td>
<td>Titanium</td>
</tr>
<tr>
<td>9</td>
<td>Ultimate Resistance</td>
<td>Titanium</td>
</tr>
<tr>
<td>10</td>
<td>Concentrated Sulfuric Acid</td>
<td>Steel</td>
</tr>
</tbody>
</table>

*Cathodic protection* (CP) has become a widely used and cost effective method for controlling the corrosion of metallic structures in electrically conducting environments. Such environments contain enough ions to conduct electricity such as soils, seawater and basically all natural waters. Cathodic protection reduces the corrosion rate of a metallic structure by supplying electrical current (or free electrons) from an alternate source. This reduces the structures corrosion potential, bringing the metal closer to an immune state. This converts all the anodic (active) sites on the metal surface to cathodic (passive) sites.
From thermodynamics point of view, the application of a CP current basically reduces the corrosion rate of a metallic structure by reducing its corrosion potential towards its immune state.

*Anodic protection* (AP) is a technique to reduce the corrosion of a metal surface by passing an external current to it to cause its electrical potential to become passive and therefore preventing corrosive attack. AP is used to protect metals that exhibit passivation in environments whereby the current density in the freely corroding state is significantly higher than the current density in the passive state over a wide range of potentials. AP is often used to protect mild steel storage vessels containing concentrated acidic or alkaline fluids such as concentrated sulfuric acid [21-23]. AP is only applicable to metals which show active-passive behaviour and can handle more aggressive environments as compared to CP.

Preventing corrosion in the *design phase* is the best method and required an understanding of its potential and to plan for it by involving a corrosion engineer for a complete project review and analysis.

### 2.2 CO₂ corrosion

Metals, including steels, do not corrode in dry CO₂ gas. However, corrosion can occur in the presence of water where aqueous CO₂ forms into carbonic acid which can be highly corrosive, which is often present in oil and gas production systems.
2.2.1 CO₂ corrosion in the oil and gas industry

CO₂ corrosion is by far the most prevalent form of corrosion attack encountered in upstream oil and gas production [24]. It has been studied extensively in a variety of conditions and environments, both in surface facilities and downhole, due to its prevalence and the severe consequences of a failure [25]. CO₂ is produced from, and at times re-injected into [26], oil and gas reservoirs in corrosion promoting conditions. This has led to corrosion studies being completed in many different environmental conditions.

CO₂ on its own is a gas, but in the presence of free water can form carbonic acid. Water and water condensates are present in oil and gas production and injection operations leading to the formation of carbonic acid.

A variety of methods are used to prevent and reduce CO₂ corrosion. Facilities and operations often rely on the formation of a passive scale, siderite or magnetite, to form. These are known to be stable, adhesive, and non-conductive and prevent corrosion under the right conditions. As long as the film remains intact it is highly protective. In the presence of high velocity turbulent fluids, solids, or other scale impurities the film breaks off and leads to highly accelerate corrosion rates. Other methods include using inhibitors and scavengers, increasing pH to buffer the carbonic acid, elevated temperatures and flow path design to reduce turbulence. Pitting and mesa corrosion are commonly used terms to describe the morphology and appearance of CO₂ corrosion [27].

2.2.2 The CO₂ corrosion mechanisms

Due to the prevalence in CO₂ corrosion occurrence, numerous CO₂ corrosion models have been developed and proposed to predict it. An original and one of the most
widely accepted models for CO$_2$ corrosion was presented in 1975 by DeWaard and Milliams [28]. It outlined a direct reduction of carbonic acid and was based on the dependence of temperature and partial pressure of CO$_2$. Since its original development in 1975, multiple revisions have been made. Alternative models available include Norsok M-506 Model, Hydrocor, Corplus, Cassandra, KSC Model, Multicorp, and several others [29]. These utilize different approaches to account for oil welling factors, protective film formation and adhesion, and the effects of impurities. The fundamental mechanism and reactions for CO$_2$ corrosion are as follows [30].

\[
CO_2(g) \leftrightarrow CO_2(aq) \quad (2-4)
\]

\[
CO_2(aq) + H_2O \leftrightarrow H_2CO_3 \quad (2-5)
\]

\[
H_2CO_3 \leftrightarrow H^+ + HCO_3^- \quad (2-6)
\]

\[
HCO_3^- \leftrightarrow H^+ + CO_3^{2-} \quad (2-7)
\]

Iron carbonate scales can form chemically as above, as well as electrochemically during the corrosion of the steels. The anodic reaction is primarily the oxidation of iron:

\[
Fe \leftrightarrow Fe^{2+} + 2e \quad (2-8)
\]

Cathodic reactions depend on the solutions pH. These are:

\[
H^+ + e \rightarrow 0.5H_2 \quad (pH < 4) \quad (2-9)
\]

\[
H_2CO_3 + e \rightarrow HCO_3^- + 0.5H_2 \quad (4 < pH < 6) \quad (2-10)
\]
For pH > 6 the direct reduction of bicarbonate ions and water can be important at low CO₂ partial pressures, P\text{CO₂}.

\[
2\text{HCO}_3^- + 2e \rightarrow 2\text{CO}_3^{2-} + H_2 \quad \text{(2-11)}
\]

\[
2\text{H}_2\text{O} + 2e \rightarrow 2\text{OH}^- + H_2 \quad \text{(2-12)}
\]

The formation of an iron carbonate scale (FeCO₃) occurs either chemically or electrochemically when the bicarbonate and carbonate ions [HCO₃⁻] and [CO₃²⁻] exceed the solubility limit. This progresses as follow:

Chemical: \[
\text{Fe}^{2+} + \text{CO}_3^{2-} \leftrightarrow \text{FeCO}_3 \quad \text{(2-13)}
\]

Electrochemical: \[
\text{Fe} + \text{HCO}_3^- \leftrightarrow \text{FeCO}_3 + \text{H}^+ + e \quad \text{(2-14)}
\]

### 2.2.3 Factors which affect CO₂ corrosion

The main factor which affects corrosion caused by CO₂ is its ability to form an iron carbonate scale. Depending on the type and quality of the scale, it can become very protective and reduce the rate of underlying steel corrosion by blocking the surface and preventing corrosive species from reaching it. The protectiveness of the iron carbonate scale depends on a number of factors. Factors include fluid flow rates, temperature, pressure, pH, as well as the presence of condensates, chlorides, salts and other impurities present in a system.

One of the main factors is the local system temperature. Generally, at ambient temperatures below 60°C a non-passive scale can form. This scale is relatively light, thin, non-adhesive, and therefore not protective. When the temperatures are increased to above
60°C the formed iron carbonate scale is adhesive and dense, providing an effective passive film to protect the underlying steel. At more extreme temperatures, above 150°C, the scales formed remain protective.

The formation of iron carbonate scale and its structure is also dependent on solution pH. In acidic solutions (pH < 5) a scale is usually not formed. A compact and protective scale is generally formed in near-neutral or alkaline solutions with a pH > 7 [31].

The scales formation and protectiveness also depend on the partial pressure of $CO_2$, $P_{CO_2}$. A general rule of thumb used for predicting $CO_2$ corrosion is based on the partial pressure of $CO_2$ ($P_{CO_2} = \text{fractional mole } % \times \text{system pressure}$) when [31]:

- $P_{CO_2}$ is greater than 30 kPa, corrosion is very likely
- $P_{CO_2}$ is between 7-30 kPa, corrosion is possible
- $P_{CO_2}$ is less than 7 kPa, corrosion is not likely

It is interesting to note that in SAGD steam injection systems with elevated temperatures, between 80°C to 250°C, that $Fe_5O_4$ could also be formed due to reaction of $FeCO_3$ with water, rather than with $O_2$, because the oxygen content is negligibly low in the steam [32].

### 2.3 $H_2S$ corrosion

Extensive studies have shown that the kinetics of $H_2S$ corrosion are controlled by the presence of an iron sulfide, $FeS$, corrosion product film. $H_2S$ corrosion is more
localized than $CO_2$ corrosion and increasing $H_2S$ would not necessarily cause an increase in general corrosion rates, but rather acceleration in localized corrosion rates leaving a material susceptible to catastrophic failure. Forms of $H_2S$ corrosion include aqueous $H_2S$ corrosion, Hydrogen Induced Cracking (HIC), Hydrogen Embrittlement (HE), and Sulfide Stress Corrosion Cracking (SSCC). $H_2S$ corrosion appears to be of major concern in SAGD operations, specifically at Suncors Firebag operation.

HIC can occur in the presence of $H_2S$ corrosion due to the formation of the $FeS$ scale and release of hydrogen atoms into the materials matrix. HIC is caused by the blistering of a metal due to a high concentration of hydrogen atoms diffused into the steels matrix. When a critical amount of hydrogen is accumulated, HIC cracks initiate and propagate [33].

HE occurs when hydrogen gas is absorbed, and diffuses into the metal at local defects. This creates local increases in internal stress causing the material to fail at stresses much less than the materials yield stress [34].

SSCC is generally encountered in high strength steels with Rockwell hardness above 22 and occurs in $H_2S$ containing environments. It is similar to HE with the exception of having $H_2S$ being present as the source of Hydrogen. HE and SSCC can be differentiated through applying current. If the specimen becomes anodic, the attack is SSCC. If cracking accelerates in the opposite direction and hydrogen gas forms, the attack is HE [34].

Depending on the environment, different corrosion films can form. Each film having distinct properties and morphology [35]. Environmental conditions which can affect these include $H_2S$ concentration, temperature, pressure, $H_2S$ activity, pH, fluid flow
rate and composition. Forms of scales include mackinawite, pyrrhotite, pyrite, troilite, and cubic \( FeS \). The morphology of \( FeS \) will vary depending on the environmental conditions present.

### 2.3.1 \( H_2S \) corrosion in the oil and gas industry

It is well known that sour oil and gas reservoirs can produce hydrogen sulfide and lead to the corrosion of downhole tubulars and surface facility infrastructure. Given the prevalence of \( H_2S \) corrosion and its severe failure consequences extensive reviews and laboratory research [36] has been completed to understand and prevent corrosion in sour oil and gas operations [37]. Sour oil and gas fields have now been operating for more than 50 years and are classified as ones in which gas phase concentrations of \( H_2S \) are greater than 3 parts per million by volume (ppmv) [38].

Several mechanisms exist which can lead to the release and production of \( H_2S \) from oil and gas reservoirs. As noted in section 1.3 one common mechanism which can release \( H_2S \) when high temperature processes are used, such as SAGD and Cyclic Steam Stimulation (CSS), is through a process known as aquathermolysis. Elevated temperatures decompose the organic sulfur compounds found in oil sands bitumen to release a variety of organic and inorganic compounds including sulfur and the autocatalytic reduction of \( H_2S \). Hydrogen sulfide can also form and be produced through the introduction of foreign microbes to a reservoir, specifically Sulfur Reducing Bacteria (SRB). SRB’s consume sulfur in the reservoir to produce \( H_2S \). This is also known as reservoir souring. Foreign bacteria can be introduced to a reservoir through the injection of water back into the reservoir. Water is often injected during well workover operations,
for reservoir pressure maintenance, and for the purposes of disposing of produced waters. Other non-microbiological souring mechanisms include Thermochemical Sulphate Reduction (TSR), dissolution of pyritic materials, and redox reactions involving bisulphite oxygen scavengers [36, 38, 39].

Materials used in sour oil and gas operations continue to be dominated by carbon steels, although more recently through the advancement of CRA’s and improved CRA material economics, their use is becoming more common for major pieces of equipment [36]. Commonly used CRA’s include:

- Martensitic Stainless Steels
- Austenitic Stainless Steels
- Duplex and Super Duplex Stainless Steels
- Nickel Alloys
- Super Austenitic Stainless Steels
- Non-metals

It is important to note that CRA’s should be carefully selected for use as some may cause extra, unforeseen material failure mechanisms. In some situations, it may be appropriate to use a combination of CS and CRA’s where high corrosion rates can be expected. The can include downhole liners and tubular components, as well as on aging assets after reservoir souring or higher water cuts are noticed.
2.3.2 The H₂S corrosion mechanisms

Many studies have been done to understand the mechanism for H₂S Corrosion [40-42]. It is generally accepted that the corrosion of steels in aqueous H₂S environments proceeds initially by a very fast direct heterogeneous chemical reaction at the steel surface to form a solid adherent corrosion layer. The type of layer can depend on many factors, temperature being a main one. At lower temperatures mackinawite often forms, but at elevated temperature pyrrhotite, pyrite, or cubic FeS can form [41]. The overall reaction is a solid-state reaction and can be written as:

\[
Fe + 2H₂S \leftrightarrow FeS₃(s) + H₂
\]  

(2-15)

Under standard conditions, H₂S absorbs onto steel surface and reacts very fast to form a thin protective mackinawite film. The mackinawite’s scale formation is known to be a “Solid State Reaction” as both the initial and final states of Fe are solid. This initial mackinawite formed is a very thin (<< 1 µm) [43], but dense scale acts as a solid state diffusion barrier for the sulfide species involved in the corrosion reaction.

The thin mackinawite film continuously goes through a cyclic process of growth, cracking and delamination. This generates the outer sulfide layer which thickens over time (typically >1 µm), presenting a diffusion barrier. This outer sulfide layer is more porous and rather loosely attached. Over time it may crack, peel and spall when aggravated by flow. The form of iron sulfide scale may change, depending on a variety of production parameters including temperature and H₂S activity. Variations of FeS scale that can form include mackinawite, pyrite, pyrrhotite, triolite, and cubic FeS.
Due to the presence of the thin passive inner mackinawite film and the outer sulfide layer, it is assumed that the corrosion rate of steels in H$_2$S solutions is always under mass, flux, transfer control [41]. The total flux of H$_2$S through the scale is a combination of:

- Convective diffusion through the mass transfer boundary layer
- Molecular diffusion through the liquid in the porous outer layer
- Solid state diffusion through the inner mackinawite film

The morphology of corrosion scales on carbon steels in sour environments can take many shapes and forms. The most common of which is that of a pit, which could be a single large wide pit, or a cluster of smaller pits. Failures often initiate in these areas. Corrosion pit morphology variations in carbon steels can be due to several factors which include the microstructure of the steel, variations in the materials surface finish, and local variations in water chemistry. Mechanisms for pit development on ferritic-pearlitic steel surfaces are due to big differences in ferrite grain size and the distribution of pearlite between different carbon steels with similar elemental composition. During exposure in the field, differences in service history can be an important factor, e.g., production shutdowns and stagnant conditions, as they have been found to enhance and accelerate the pitting initiation process.

Lab morphology testing usually finds that edge and crevice corrosion near test coupon holders are sometimes observed in sour corrosion tests. These attacks can be so severe that the corrosion on the rest of the surface is overshadowed. While it may be impossible to completely avoid artificial corrosion attacks, the possibility of such attacks
should be considered when designing the experimental setup, evaluating of the results, and in making field comparisons [35].

2.3.3 Factors which affect H$_2$S corrosion

A variety of factors can affect the occurrence and rate of H$_2$S corrosion. In most environments and solutions, a very thin and protective iron sulfide film, often mackinawite, is able to form. The rate of corrosion is highly dependent on the formation of this passive film. Alternate forms of FeS scales commonly found include pyrrhotite, pyrite, and cubic iron sulfide. These scales are more porous and have a higher porosity, permeability, and are less adhesive than mackinawite. A variety of factors contribute to the ability for an adhesive scale to form and protect the steel. Contributing factors include material and electrolyte properties, fluid flow rates, as well as the solution pH, and partial pressure of H$_2$S ($P_{H_2S}$).

The material being used has a large effect on the occurrence and rate of corrosion. Proper material selection plays a large role in preventing H$_2$S corrosion from initiating, giving rise to the development of the NACE standard: NACE MR0175/ISO 15156 [44]. This standard was developed to determine the severity of an environment to corrosion, specifically SCC, depending on solution pH and $P_{H_2S}$. Four regions are outlined, which describe the severity of the corrosion environment and recommends material and design specifications, as shown in Figure 2.3 [44].
In region 0 no precautions are required; and regions 1, 2, and 3 all require material and design precautions to be taken.

The electrolyte’s conductivity plays a role in the rate of corrosion. As conductivity increases there will be an increase in the ion flow through the electrolyte and therefore increased corrosion current [45].

Fluid flow rates affect corrosion rates by preventing or promoting the formation of a passive scale. Higher flow rates can prevent scales from properly adhering to a materials surface. Under low flow or stagnant conditions solution impurities can be present and prevent the formation of passive films, progressing and even accelerating corrosion.
Temperature also plays a large role in determining the rate of corrosion when H$_2$S is present. The study of corrosion related to H$_2$S has mainly been focused on HIC and SSCC at ambient temperatures. Limited testing has been done to understand aqueous H$_2$S corrosion rates at moderate or elevated temperatures. Figure 2.4 and Figure 2.5 [46] show how corrosion rate and the scale formed depend highly on temperature as well as H$_2$S activity. H$_2$S activity is a measure of the "effective concentration" of a species in a mixture.

![Figure 2.4: Corrosion rate vs temperature][46]
As temperature increases from ambient the corrosion rates continue to increase and peaks at 40°C. Afterwards corrosion rates decrease rapidly when the temperature is up to 60°C, followed by a more gradual decrease between 60°C-90°C as temperature increases. It was proposed that corrosion rates initially increased due to the diffusion enhancement of corrosive species and the increase in both the $H^+$ concentration and the rate constant of corrosion reaction. High transport resistance of the corrosion film and the decreasing $H^+$ concentration were responsible for the decrease in corrosion rate with further increase of temperature above 40°C. In addition, the corrosion film formed likely has a more dominant role in the decrease in corrosion rate than the dilution of $H^+$ concentration. Depending on the solution temperature, pH, and the length of exposure time, different corrosion films and products have been observed. At temperatures below 40°C, lab testing commonly produces the iron sulfide film mackinawite. Above 40°C a
combination of mackinawite and cubic iron sulfide (cubic FeS) has been observed. Cubic FeS is known to be a more adhesive and dense scale than mackinawite.

The presence of chlorides and other corrosive impurities can also play a large effect in determining the ability for a passive corrosion film to form and on the rate of corrosion of a materials surface.

2.4 Synergistic effects of H$_2$S and CO$_2$ on steel corrosion

Studies of combined aqueous CO$_2$ and H$_2$S corrosion have been investigated for over 60 years, starting with investigations on carbon steel pipelines [47]. More recently an increase in the amount of research in this area has been completed in different operating conditions and system types [48-50]. Small amounts of H$_2$S in a system have been found to change the corrosion mechanism and rate [51]. A shift in the preferred corrosion mechanism and product will occur when the corrosion kinetics of FeS formation becomes favorable to that of FeCO$_3$.

To simplify the analysis and understanding of the corrosion mechanism present in systems which include both CO$_2$ and H$_2$S, a partial pressure relationship between CO$_2$ and H$_2$S was developed. This was done to determine how much H$_2$S is required relative to CO$_2$ to turn a sweet system into a sour one with the H$_2$S concentration > 0.5ppm. A ratio of the partial pressures of CO$_2$/H$_2$S = 500 was found to represent the transition point from being CO$_2$ to H$_2$S dominant. When the partial pressure ratio is < 500 an iron sulfide corrosion product can be expected to initiate [52] and dominate at a ratio < 20. Above 500 an iron carbonate scale should prevail. Between a ratio range of 20 to 500 a transition from being fully CO$_2$ dominate to H$_2$S dominate exists. This ratio has become a
commonly used “rule of thumb” for a conversion from sweet to sour corrosion since its proposal at NACE CORROSION 1983 [53]. It is important to note that this ratio was developed under unique conditions to show that for the appropriate solubility products of ionization constants, and equilibrium constants, the co-existence of siderite and iron sulfide at 25°C is indicated when the partial pressures of CO₂/ H₂S = 500. Temperature and other solution imperfections were not considered in the conclusion that iron carbonate should prevail when the partial pressure of CO₂/ H₂S ≥ 500, and iron sulfide can be expected with a CO₂/ H₂S ratio < 500.

Because of its popularity in use, this value has been recently re-evaluated and many questions and concerns have come up [54]. The transition point has shown to be highly sensitive to the thermodynamic value inputs. Slight changes in the solubility product and equilibrium constant values used will lead to extremely large changes in the transition value. Temperature has shown to have little effect on the ratio in the range of 20°C-100°C, assuming a mackinawite corrosion product. If the corrosion product changes from mackinawite to pyrrhotite the ratio could increase from 500 to roughly 2500. Furthermore, a 1% change in the free energy value of iron carbonate can increase the value to over 7600 [54].

2.5 Summary of literature review

Corrosion is known to occur and have caused well failures in the oil and gas industry including in SAGD wells. Well failures have a significant economic, regulatory, and environmental impact on an operating company’s bottom line, as well as on the SAGD and oil and gas industry due to reputation and social rights. Based on the literature review
completed and industry experience, no research has been completed to better understand corrosion in the high temperature high pressure conditions found in SAGD, specifically to those at Suncors Firebag asset. Component and well failures have occurred due to corrosion which is an economic, environmental, and regularity hazard to operations. The goal of this research is thus to:

- Better understand corrosion on wellhead gate valve gate face materials and evaluate an alternate material to use for prevention
- Better understand the severity of corrosion on downhole completion components under the given conditions which can be expected in SAGD operations
- Understand where the most severe corrosion is likely to occur
- Determine the dominating corrosion mechanisms present
- Understand how corrosion can change on different materials and at different locations in a well

Having a more thorough understanding of these factors will lead to improved well design, operating practices, and interventions to prevent corrosion occurring in SAGD wells.
Chapter Three: Experimental testing

3.1 Introduction

To better understand corrosion rates and mechanisms in the annulus of producing SAGD wells, experiments were conducted in both lab and field environments. Lab testing was completed using HTHP autoclaves for real time and average corrosion rate measurements on selected materials in simulated downhole conditions. Materials tested were those of commonly used wellhead and other completion materials setup to represent conditions found in SAGD wells.

Field testing was completed to better understand corrosion in real world applications. Having a better understanding of the corrosion rates, scale composition, and morphology from field samples can help to better understand how and why corrosion is occurring, where it can be expected to be of the highest concern, and how to prevent it. Field testing consisted of installing corrosion coupons on specially designed clamps used to secure the ESP’s cable to the external of the production tubing from surface to where the ESP is landed. Similar materials and corrosive setups were constructed in the clamps as was used in the autoclaves to allow for accurate and direct comparisons between the field and lab tests. This will allow for more accurate lab tests to be set up in the future for simulation testing. The following sections give a detailed explanation of the testing procedures and setups used, measurements taken, and analysis completed.
3.2 Lab testing

HTHP autoclave testing was conducted to understand corrosion in SAGD wells on commonly used wellhead and completion materials in controlled setups with real time measurements. Materials tested include those of wellhead gate valve materials, production tubular materials, and selected other completion component materials. Materials tested, the testing completed, and location of use in application are summarized in Table 3.1.

Table 3.1: Autoclave testing summary

<table>
<thead>
<tr>
<th>Materials</th>
<th>Measurements</th>
<th>Test Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wellhead Materials</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1018 Carbon Steel</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td></td>
</tr>
<tr>
<td>Deloro-40 HF</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Cyclic Potentiodynamic Potential</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Potentiostatic Potential</td>
<td></td>
</tr>
<tr>
<td>Stellite-6 HF</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Cyclic Potentiodynamic Potential</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Potentiostatic Potential</td>
<td></td>
</tr>
<tr>
<td>Downhole Materials</td>
<td></td>
<td></td>
</tr>
<tr>
<td>J-55</td>
<td>- Weight Loss</td>
<td>- Crevice CR</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Galvanic CR</td>
</tr>
<tr>
<td>K-55</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td>- Crevice CR</td>
</tr>
<tr>
<td></td>
<td>- Zero Resistance Ammeter</td>
<td>- Galvanic CR</td>
</tr>
<tr>
<td>TN-55TH</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td>- Crevice CR</td>
</tr>
<tr>
<td></td>
<td>- Zero Resistance Ammeter</td>
<td>- Galvanic CR</td>
</tr>
<tr>
<td>GLV-K55</td>
<td>- Linear Polarization Resistance</td>
<td>- Crevice CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td>- Galvanic CR</td>
</tr>
<tr>
<td>GLV-J55</td>
<td>- Weight Loss</td>
<td>- Crevice CR</td>
</tr>
</tbody>
</table>
The appendix shows the materials mechanical properties, and references [5, 6, 8, 55] include material chemistries and general information as provided by the coupon manufacturers and suppliers. It is important to note that J-55, K-55, and TN-55TH are all similar materials manufactured to API specification 5CT with the exception that the minimum tensile strength of the J-55 is lower than the K-55. Also TN-55TH is a proprietary thermal use carbon steel product supplied by Tenaris [5]. L-80 is a steel alloy also manufactured to API specification 5CT with controlled yield and hardness properties [56]. Test setups of general corrosion testing, as well as galvanic and crevice corrosion testing cells were constructed to represent downhole corrosion conditions.

Carbon steel was selected for testing as a baseline material due to its extensive use in oil and gas industry. Carbon steel overlaid with Deloro-40 HF or Stellite-6 HF was tested to compare their ability to prevent corrosion. These materials are used in wellhead gate valves on the gate face component which is used to isolate surface lines from the downhole fluids. This is depicted in Figure 3.1.
The gate face is directly exposed to production fluids in stale or flowing conditions. Having the ability to properly operate these valves and achieve isolation is integral to maintaining well integrity. A Deloro-40 HF is currently being used in the valves on the gate face but found to be corroding and causing the valves to leak internally when exposed to fluid flow. The Stellite-6 HF was tested to determine its effectiveness in preventing corrosion, and the valves from leaking, as a potential future material upgrade change.

The K-55 and J-55 materials are commonly used in the oil and gas industry as a downhole tubular material. They are carbon steels manufactured to a specific API standard for use in oil and gas. This standard outlines select material properties which must be met and maintained for use in the oil and gas industry. More recently, tubing suppliers have started manufacturing their own proprietary thermal grades of J-55/K-55, tubulars, one of which is a TN-55TH. It is now commonly used in SAGD and selected for
its improved thermal properties. TN-55TH has tighter tolerances on its chemical composition, as well as physical dimension and mechanical properties compared to standard API grade materials. L-80 is a third standard API carbon steel used in SAGD and was also tested. Compared to J-55/K-55, L-80 is a quenched and tempered material, not rolled or normalized as J-55/K-55 are, and has a more restrictive chemistry and higher yield and tensile strength properties. Finally, galvanized versions of J-55/K-55 materials were tested as it is an alloy used on select downhole components including ESP cables, and cable clamps.

The HTHP autoclaves used were constructed of Hastelloy 276-C alloy and have a capacity of approximately 300 mL. All testing was carried out with 250 mL of field simulated brine in each cell. The temperature of the fluid in the autoclave is sensed by a thermistor probe, accurate to +/- 2°C [57, 58], held at the center of the cell by a Hastelloy sleeve. Charging of the autoclave was done using an offset Hastelloy tube fitted with a pressure gauge and sour-service valve. Each cell is also equipped with a pressure relief valve used for purging test liquids directly in the cell. Figure 3.2 shows the assembled autoclaves used for testing.
3.2.1 Wellhead material autoclave testing

The following section outlines the autoclave specifications used in testing, condition and brines used for tested, materials tested, and the measurements taken.

3.2.1.1 Autoclave Setup and Measurements

A three-electrode autoclave assembly, supplied by Cormetrics Ltd, was used with the electrodes suspended from the lid of the autoclave. The bottom was kept clear for a teflon coated magnetic stir-bar used to mix the fluids [59]. The hanging electrodes are configured to that of an equilateral triangle. Disc electrodes were 20 mm in diameter × 3 mm in height for the working and counter electrodes. This set-up was used to simulate the conditions of the valve gate where it is retracted into the valve body during normal operations. In all tests the reference electrode was made of a Hastelloy 276-C material.
The temperature of the fluid in the autoclave is sensed by a thermistor probe [57, 58], held at the center of the cell by a Hastelloy thermowell. Charging of the autoclave was by means of an offset Hastelloy tube, fitted with a pressure gauge and sour-service valve. Each cell is also equipped with a pressure relief valve which is used when purging the test liquids directly in the cells.

Weight loss (WL) and Linear Polarization Resistance (LPR) measurements were taken to measure corrosion rates on all materials. Further cyclic potentiodynamic polarization (CPP) and potentiostatic polarization (PP) testing was also completed to more accurately differentiate the performance between the Deloro-40 HF and Stellite-6 HF materials. LPR and Zero Resistance Ammeter (ZRA) measurements were obtained at 30 minute intervals by connecting each cell to a Gamry PC4-300 potentiostat and controller [60], via a Gamry ECM8 multiplexer [61]. Data acquisition was performed by means of Gamry’s DC105 software package. The LPR test setup included using a grooved teflon washer mounted on the top surface of the disc to create a crevice corrosion condition that is conducive to localized corrosion activity. This assembly is shown in Figure 3.3.

![Figure 3.3: Crevice washer and electrode assembly](image)

For this testing, the working electrode consisted of 1018 carbon steel which had a total exposed surface area of 5.94 cm². This was used for all corrosion rate calculations.
The electrodes were solvent rinsed and weighed prior to the test. Weight loss corrosion rates are reported as an average of the working electrode based on a total exposure period of 163 hours (~7 day). In the CPP and potentiostatic polarization tests the working disc electrode was mounted on the electrode poles and sealed off on the bottom using a PEEK cap. PEEK is an inert and biocompatible polyetheretherketone tubing material [62].

Electrodes were solvent rinsed and weighed prior to the test.

During the CPP scans, electrochemical measurements were obtained at 1 second intervals, and 15 minute intervals for the PP testing. This was done by connecting each cell to a Gamry PC4-300 potentiostat and controller, via a Gamry ECM8 multiplexer. Data acquisition was by means of Gamry’s DC105 software package.

3.2.1.2 Coupon materials and autoclave setup

Table 3.2 lists the materials tested, where they are used in application, and test measurements taken.

<table>
<thead>
<tr>
<th>Material</th>
<th>Location of Use</th>
<th>Analysis Completed</th>
</tr>
</thead>
<tbody>
<tr>
<td>1018 Carbon Steel</td>
<td>Gate valve internal components</td>
<td>LPR, WL</td>
</tr>
<tr>
<td>Deloro-40 HF</td>
<td>Gate valve gate face overlay</td>
<td>LPR, WL, CPP, PP</td>
</tr>
<tr>
<td>Stellite-6 HF</td>
<td>Gate valve gate face overlay (for potential use)</td>
<td>LPR, WL, CPP, PP</td>
</tr>
</tbody>
</table>

For LPR testing the disc style coupons were milled from the materials and mounted in the crevice corrosion device. CPP testing used disc style coupons milled from each material and secured on the electrode poles with the PEEK cap. The Deloro-40 HF and Stellite-6 HF materials were made using these materials matching the required coupon configuration but tapping the threaded connection within the discs was not able to
be completed. To allow for the CPP measurements, a silver adhesion paste was utilized to allow for electrical continuity between the disc and mounting rod. This alternative to the threaded connection is commonly used in electrochemical testing and has proven to be a comparable alternative. Based on the data generated and stable open circuit potential it is believed that the test set-up worked properly.

3.2.1.3 Solution and testing conditions

The synthetic brine used in the autoclave was prepared based on a water sample analysis from a SAGD well. The brines composition is displayed in Table 3.3.

**Table 3.3: Synthetic brine composition**

<table>
<thead>
<tr>
<th>Sodium (mg/L)</th>
<th>Potassium (mg/L)</th>
<th>Magnesium (mg/L)</th>
<th>Calcium (mg/L)</th>
<th>Chloride (mg/L)</th>
<th>Bicarbonate (mg/L)</th>
<th>Sulfate (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>266.0</td>
<td>15.5</td>
<td>3.5</td>
<td>9.98</td>
<td>141.0</td>
<td>450.0</td>
<td>43.0</td>
</tr>
</tbody>
</table>

This synthetic brine was pre-purged with pure CO₂ for 2 hours prior to use in testing and had a pH of 5.50 after pre-purging. Acidic gas quantities of 6.9% mole of CO₂ and 6.48% mole of H₂S with a system pressure of 3000 kPa was used based on a well gas analysis. Autoclave charge pressures used are listed in Table 3.4. Due to the relatively short test duration, the pressures are believed to have reduced an insignificant amount during the test. This can be considered negligible.

**Table 3.4: Autoclave charge pressures**

<table>
<thead>
<tr>
<th>Total Pressure (kPa)</th>
<th>CO₂ (kPa)</th>
<th>H₂S (kPa)</th>
<th>N₂ (kPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3000</td>
<td>207</td>
<td>194</td>
<td>2599</td>
</tr>
</tbody>
</table>
A testing temperature of 170°C was used. This is the maximum autoclave operating temperature. The sealed autoclaves were placed inside individual mantles and heated to the set test temperature.

### 3.2.2 Downhole material autoclave testing

Autoclave corrosion testing was completed on select commonly used completion materials including K-55, GLV-J55, and TN-55TH. K-55 and TN-55TH are downhole tubular materials, and the GLV-J55 is an ESP cable and ESP cable clamp material.

To understand general corrosion rates on individual materials the autoclaves were setup for LPR corrosion rate measurements and pre/post-test weight loss measurements and analysis. Crevice corrosion rate tests were also performed by setting up the coupons to be in close contact in the autoclave. This testing was designed to evaluate the severity of localized corrosion attack in crevices created between materials in contact downhole. Localized crevices could be an area of potentially severe corrosion attack. Material combinations used in this testing include K-55+TN-55TH, K-55+GLV-J55 and K-55+K-55. Analysis completed includes visual observations and general weight loss corrosion rate analysis.

Finally, galvanic corrosion analysis was completed to evaluate the effects of dissimilar metals coming into contact on corrosion rate. A potentiostat connected to the autoclave in ZRA mode was used to evaluate the corrosion current density, which is directly related to corrosion rate, in the material. In ZRA mode the working and counter electrodes are shorted together so that there is a zero-net voltage drop across the cell. Thereby the current flowing between the two electrodes due to the potential difference can be measured. The direction of current flow is indicated by a positive or negative
current density. If the potential difference between the two materials is great enough the more electronegative metal of the couple will preferentially corrode.

3.2.2.1 Autoclave setup and measurements

For the LPR and ZRA testing, a three-electrode assembly was suspended from the lid of the autoclave as shown in Figure 3.4 and Figure 3.5, while keeping the bottom clear for a teflon-coated magnetic stir-bar.

Figure 3.4: Autoclave lid for general LPR testing
Figure 3.5: Galvanized J-55 & K-55 coupon pair tied at the base

The configuration of the electrodes is that of a closely spaced equilateral triangle, with each cylindrical electrode having dimensions of 63 mm x 38 mm. The reference electrode was made from Hastelloy 276-C, while the working and counter electrodes were the materials being evaluated. The inlet tube used to purge and charge the cell with fluids is directly beneath the reference electrode. The electrodes are electrically isolated from the autoclave cell via the PEEK shafts visible in the photograph. Electrodes were solvent rinsed and weighed prior to the commencement of the test period. The surface area of each rod electrode utilized in the corrosion rate calculations was 7.92 cm². LPR and ZRA measurements were obtained at 30-minute intervals by connecting each cell to a Gamry PC4-300 potentiostat and controller, via a Gamry ECM8 multiplexer. Data acquisition was by means of Gamry’s DC105 software package.
To measure the current due to galvanic coupling, a potentiostat in ZRA mode was used. The corrosion rate, measured via weight loss, and visual observations of each electrode from the galvanic pair are reported.

For crevice testing the strip coupons were attached to the short PEEK insulators suspended from the lid of the autoclave. The top of each coupon pair was mounted on the side of the PEEK insulator and the bottom is held together to create a contact point and artificial crevice. The coupon dimensions are 1.27 cm x 7.62 cm x 0.16 cm (WxLxT) with 0.64 cm diameter mounting holes located 1.27 cm from either end. The coupons were solvent rinsed and weighed prior to testing. The total exposed surface area of 20.9 cm² was used for weight loss corrosion rate calculations.

The crevice testing setup was provided in Figure 3.5. Here, strip style coupons were tied tightly together at the bottom and then mounted from short PEEK shafts attach to the lid of the autoclave. A high temperature strip of 100% PTFE was used to attached the coupons to each other and mount them from the lid of the autoclave. After preparing the 3 different crevice setups, a 250 mL volume of the purged synthetic brine, composition shown in Table 3.6 in the following section, was added to each autoclave before assembling with the lids and sealed together. Finally, the charged autoclave cells were placed inside individual heating mantles with magnetic stirrers and brought up to a final test temperature of 170°C. A magnetic stir-bar was also present in the autoclave so that fluids can be gently stirred throughout test duration of the testing.

3.2.2.2 Coupon materials and testing setup

Rod style electrodes were milled from K-55 and TN-55TH samples with two of the milled K-55 electrodes being electro-galvanized. The rod style electrodes were used
in the general and galvanic corrosion tests. Additionally, strip style coupons were milled from tubular samples for use in the crevice testing. These coupons were milled to the same physical dimensions as the GLV-J55 and K-55 coupons. Table 3.5 summarizes the materials tested, their location and application for use, and the measurements taken for testing.

**Table 3.5: Materials and location of use**

<table>
<thead>
<tr>
<th>Material</th>
<th>Location and Application for Use</th>
<th>Analysis Completed</th>
</tr>
</thead>
<tbody>
<tr>
<td>K-55</td>
<td>Tubular Material</td>
<td>LPR, Crevice &amp; Galvanic</td>
</tr>
<tr>
<td>TN-55TH</td>
<td>Tubular Material</td>
<td>LPR, Crevice &amp; Galvanic</td>
</tr>
<tr>
<td>GLV-J55</td>
<td>Clamp and ESP Cable</td>
<td>Crevice</td>
</tr>
<tr>
<td>GLV-K55</td>
<td>Clamp and ESP Cable</td>
<td>Galvanic</td>
</tr>
</tbody>
</table>

3.2.2.3 Solution and testing conditions

A condensed water sample from the field was provided for compositional analysis to determine its composition and the quantities of calcium, chloride, bicarbonate, and sulfate present. Using this compositional analysis, a synthetic brine was prepared for use in the autoclave tests, and its composition is shown in Table 3.6.

**Table 3.6: Synthetic brine composition**

<table>
<thead>
<tr>
<th>Calcium (mg/L)</th>
<th>Chloride (mg/L)</th>
<th>Bicarbonate (mg/L)</th>
<th>Sulfate (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>9.98</td>
<td>141</td>
<td>450</td>
<td>43</td>
</tr>
</tbody>
</table>

The brine was pre-purged with CO₂ for 2 hours to deoxygenate the brine before testing. The pH of the brine was 4.36 after the pre-purge for the LPR and Crevice testing, and 4.34 in the galvanic testing. The gas composition used in testing was 30% molar CO₂.
and 8% molar H$_2$S based on a system pressure of 1000 kPa. Autoclave charge pressures used in the experiment are listed in Table 3.7.

**Table 3.7: Autoclave charge partial pressures**

<table>
<thead>
<tr>
<th>Total Pressure (kPa)</th>
<th>CO$_2$ (kPa)</th>
<th>H$_2$S (kPa)</th>
<th>N$_2$ (kPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1000</td>
<td>300</td>
<td>80</td>
<td>620</td>
</tr>
</tbody>
</table>

ZRA experiments were run over a 4-day exposure while LPR and crevice tests were completed with 7-day exposure times. A maximum autoclave testing temperature of 170°C was used. This is below maximum SAGD operating temperatures which can be more than 220°C and is due to autoclave material limitations.

### 3.3 Field testing

Field corrosion coupon testing has been completed to determine the effects of corrosion under actual operating conditions for longer duration periods. In this analysis corrosion coupons were manufactured and installed in the annulus of producing SAGD wells. The purpose of this testing was to determine corrosion rates on commonly used SAGD completion materials at different elevations in the production annulus as well as for corrosion rate and mechanism determination and comparisons to the controlled lab testing.

Here corrosion coupons were installed in specially designed production tubing instrumentation clamps used to fasten downhole instrumentation lines to the production tubing. These clamps are shown in Figure 3.6 and were manufactured to protect the coupons during installation and to prevent unwanted coupon contact with other materials.
Figure 3.6: Clamp and coupon installation setup

Five sets of clamps with coupon holders were installed at select well elevations. This was done to determine variations in corrosion rate and mechanism at different elevations due to the presence of varying fluid compositions and flow conditions. This is shown in Figure 3.7 and Table 3.8. The measure mKB “meters kelly bushing” is a depth measurement taken from a reference location, the kelly bushing, on a drilling rig. It is the measure of a well’s depth along the wellbore.

Figure 3.7: Coupon installation depths
Table 3.8: Corrosion coupon installation depths

<table>
<thead>
<tr>
<th>Component</th>
<th>Depth (mKB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ESP Depth (mKB)</td>
<td>537.58</td>
</tr>
<tr>
<td>ESP Depth (mTVD)</td>
<td>334.00</td>
</tr>
<tr>
<td>Joint 2</td>
<td>505.15</td>
</tr>
<tr>
<td>Joint 12</td>
<td>372.47</td>
</tr>
<tr>
<td>Joint 23</td>
<td>223.25</td>
</tr>
<tr>
<td>Joint 24</td>
<td>209.76</td>
</tr>
<tr>
<td>Joint 25</td>
<td>196.17</td>
</tr>
</tbody>
</table>

Each clamp had three separate corrosion cell setups on it. Setups included standalone material testing on the J-55 and L-80 materials, as well as a galvanic/crevice setup between J-55 and GLV-J55 coupons. Three of the clamps were installed on tubing joints at, above, and below where the annular liquid level is believed to be. Due to varying operating conditions in a well, the range of fluid level elevation is approximately 30 m. A fourth clamp was installed half way between the ESP and liquid level. And the fifth clamp was installed on the tubing joint above the ESP. The fluid level top within the well can be estimated using surface and ESP pressure measurements. Field testing work is summarized in Table 3.9.
Table 3.9: Field coupon testing summary

<table>
<thead>
<tr>
<th>Materials</th>
<th>Measurements</th>
<th>Analysis</th>
<th>Test Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>J-55</td>
<td>WL</td>
<td>XRD, SEM, EDX</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Morphology</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Scale Composition</td>
</tr>
<tr>
<td>L-80</td>
<td>WL</td>
<td>XRD, SEM, EDX</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Morphology</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Scale Composition</td>
</tr>
<tr>
<td>GLV-J55</td>
<td>WL</td>
<td>XRD, SEM, EDX</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Morphology</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Scale Composition</td>
</tr>
</tbody>
</table>

3.3.1 Coupon materials, clamp design, and installation design

Table 3.10 lists the materials analysed in the downhole corrosion tests.

Table 3.10: Downhole corrosion coupons

<table>
<thead>
<tr>
<th>Material</th>
<th>Location of Use</th>
<th>Analysis Completed</th>
</tr>
</thead>
<tbody>
<tr>
<td>J-55</td>
<td>Tubular Material</td>
<td>Corrosion Rate and Morphology</td>
</tr>
<tr>
<td>L-80</td>
<td>Tubular Material</td>
<td>Corrosion Rate and Morphology</td>
</tr>
<tr>
<td>GLV-J55 + J-55</td>
<td>Clamp and ESP Cable</td>
<td>Corrosion Rate and Morphology</td>
</tr>
</tbody>
</table>

J-55 and L-80 were tested due to their common use in oil and gas tubular applications and the combination of the GLV-J55+ J-55 was to better understand galvanic and crevice corrosion between completion and tubular components. K-55 was not tested due to sample availability and its compositional similarity to J-55. Enough so that comparisons to the K-55 coupon corrosion rates from the lab testing can be made. J-55 and L-80 materials have both mechanical and chemical differences. L-80 was tested for it being a higher carbon content steel with increased strength and hardness properties to
understand the effects of the SAGD conditions present on this type of material. See appendix for general material property differences.

Five specially designed instrumentation clamps were installed in a well for a total of 20 coupons. Coupons were bolted on the clamp and separated from the clamp using a teflon spacer. In the galvanic and crevice testing setup between the GLV-J55 and J-55, no washer or separator was used to isolate each coupon. The same teflon spacer was still used to isolate it from the clamp.

Coupon samples were manufactured to the NACE Standard RP0775-2005 “Preparation, Installation, Analysis, and Interpretation of Corrosion Coupons in Oilfield Operations” [63]. The corrosion coupon dimensions were as follows:

1. J-55 and GLV-J55 coupons for crevice/galvanic testing: 1/16" X 1/2" X 3" with two 1/4" diameter holes. Each hole located 1/2” in from the end and centered. The coupon corners are squared.
2. L80 and J-55 coupon: 1/8" X 1/2" X 3" with two 1/4" diameter holes. Each hole located 1/2” in from the end and centered. The coupon corners are squared.

### 3.3.2 Downhole operating conditions

Figure 3.8 to Figure 3.10 show the wells operating conditions throughout the testing period. Including the estimated average liquid level in the well, surface and bottom hole pressure (BHP), as well as the downhole ESP and casing surface temperatures respectively. Figure 3.8 also shows the estimated liquid level in the wells annulus with the coupon elevations. The joint 2 coupon is located just above the ESP, continually submerged in the emulsified fluid. The joint 12 coupon was submerged in oil
for the majority of the coupons installation. The coupons on joints 23, 24, and 25 were likely both submerged and above the liquid level throughout the testing.

The liquid level in a well varies depending on the wells operating conditions and setpoints. Changes in ESP speed, production tubing and casing surface pressure setpoints, all of which cause BHP deviations, will lead to changes in the fluid level due to differential pressure changes between the BHP and casing surface pressure. Changes in annular pressures or temperatures, because of well operating condition changes, can also lead to an increase or decrease in the annular gas flow and condensation rates based on changing saturation conditions. During the testing these parameters were out of my control as they are adjusted to optimize the wells operation and oil production based on plant and site requirements. As can be seen in Figure 3.9 the casing surface pressure setpoint is steady and that changes in downhole operating parameters is what more commonly dictates the liquid level in the well. Understanding the liquid level elevation is important as corrosion issues have been found here in other wells based on past failure analysis.

Figure 3.10 shows the wells downhole and casing surface temperate trends. This shows downhole temperatures being at 200°C for much of the testing period and up to 215°C for periods of time. The operating BHP is shown to be between 2000 kPa-2500 kPa for the majority of the trial, with saturation temperatures of approximately 212.4°C to 223.9°C respectively. This indicates that at the ESP fluids were for the most part in a liquid form.

Casing temperatures can give an indication as to when the wells annulus may be flowing gas or not. Casing annular gas flow meter data is not available in this well. The
steam saturation temperature at 1000 kPa is approximately 180.0°C. When operating conditions and temperatures reach this level, that annular fluid, containing water, will begin to vaporize and the casing will likely start flowing fluids to surface and be produced. Some of the fluids may condense as temperature and pressure drop from the liquid level to surface. This will cause condensation and possibly further corrosion at higher well elevations.

![Figure 3.8: Annular fluid level](image)

Figure 3.8: Annular fluid level
Figure 3.9: Bottom hole and casing surface pressures

Figure 3.10: Operating temperatures
3.3.3 Field testing analysis

Similar to the lab testing, selecting the proper corrosion analysis techniques to analyze the corrosion coupons is important to accurately determine corrosion rates and mechanisms. Corrosion coupon analysis techniques used include:

- Visual inspections
- Weight loss analysis for averaged corrosion rates
- X-ray diffraction (XRD) for corrosion scale composition: Determines the atomic and molecular structure of a crystal. Unidentified compounds can be noted when homogeneous and single phase materials are present.
- Scanning Electron Microscopy (SEM): Produces sample images containing information about the sample's surface topography
- Energy-Dispersive X-ray Spectroscopy (EDS): An analytical technique used for the elemental analysis of a sample
  - **NOTE:** EDS is typically only completed to support XRD analysis should clarifications be required or XRD discrepancy readings found. No XRD issues were noted and therefore this data not displayed.
Chapter Four: Testing results and analysis

This section discusses and analyzes the testing results for the lab and field tests completed. Analysis completed includes corrosion rate, current, and potential testing. Also completed were morphology imaging, visual inspections, and comparative scale compositional analysis.

Corrosion rate calculations and analysis were done using NACE standard SP0775-2013 [63]. This standard outlines how corrosion rates are calculated, and categorizes corrosion severity based on corrosion coupons installed in oil and gas operations. The categories are presented in Table 4.1.

<table>
<thead>
<tr>
<th>Category</th>
<th>Corrosion Rate Range (mm/y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>&lt;0.025</td>
</tr>
<tr>
<td>Moderate</td>
<td>0.025-0.15</td>
</tr>
<tr>
<td>High</td>
<td>0.16-0.25</td>
</tr>
<tr>
<td>Severe</td>
<td>&gt;0.25</td>
</tr>
</tbody>
</table>

4.1 Lab testing results and analysis

This section presents the results from the wellhead [64] and downhole material testing [65, 66] completed in a controlled laboratory environment. Wellhead material weight loss, LPR, potentiostatic polarization, and CPP testing results on 1018 carbon steel, Deloro-40 HF, and Stellite-6 HF materials are presented. The downhole material testing results from the K-55, TN-55TH, and GLV-J55 materials are also shown along with a discussion on the findings.
4.1.1 Wellhead material testing results

The LPR and weight loss corrosion rate tests are summarized in Table 4.2 and shown in Figure 4.1 and Figure 4.2 to display the corrosion rate resting completed using LPR. Post-testing photographs are shown in Figure 4.3.

Table 4.2: Corrosion rate data summary

<table>
<thead>
<tr>
<th>Material</th>
<th>Initial LPR (MPY/mm/y)</th>
<th>Final LPR (MPY/mm/y)</th>
<th>Weight Loss (mg)</th>
<th>Weight Loss CR (MPY/mm/y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1018 Carbon Steel</td>
<td>70.43/1.79</td>
<td>15.05/0.38</td>
<td>28.4</td>
<td>9.66/0.25</td>
</tr>
<tr>
<td>Deloro-40 HF</td>
<td>0.00</td>
<td>1.38/0.035</td>
<td>0.10</td>
<td>0.03/0.00076</td>
</tr>
<tr>
<td>Stellite-6 HF</td>
<td>0.61/0.015</td>
<td>0.95/0.024</td>
<td>-10.60</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Figure 4.1: Wellhead material LPR autoclave crevice corrosion testing
The 1018 carbon steel weight loss corrosion rate was calculated to be 0.25 mm/y (9.66 MPY). The initial LPR corrosion rate was 1.789 mm/y (70.43 MPY), which then trended down sharply to ~0.635 mm/y (25 MPY) as shown in Figure 4.1. This was followed by a gradual decrease for the remainder of the 7-day test period. This spike followed by a decrease indicates initially high corrosion rates followed by the formation
of a passivating corrosion scale. It is believed that FeS scales become stable and uniform after ~24h [50]. This is observed on the 1018 CS samples, but not on the HF materials. This is likely due to their corrosion resistant properties and a longer time required for a passive film to form. A moderate etching attack was noted in the creviced areas of the coupon, as well as the exposed bottom surface indicating no preferential corrosion had occurred in the creviced areas compared to the flat surfaces. Although preferential corrosion can be expected, it is likely to have not occurred here due to the steels susceptibility to rapid rates of initial corrosion and uniform scale formation. In this relatively undisturbed environment the passive scale remains intact and limits any preferential corrosion from occurring.

The Deloro-40 HF material had a very low initial weight loss corrosion rate of 0.001 mm/y (0.03 MPY). The recorded weight loss of 0.1 mg is equal to the accuracy of the analytical balance used to weigh the electrodes and may be considered negligible. The Deloro-40 HF LPR corrosion rate displayed a peak level of ~0.152 mm/y (6 MPY) before decreasing to ~0.0254 mm/y (1 MPY) by the end of the test period. No visible corrosion activity was noted on the electrode surface when observed under 10 times magnification. Deloro-40 HF peak corrosion rates were reached 3-4 days into the testing, as compared to the carbon steel which peaked in <1 day. Past research has shown it difficult to determine the exact mechanism of corrosion and corrosion resistance for HF alloys [67, 68]. A hypothesis in the longer time taken to reach peak rates before decreasing to stabilize is that it could be due to the lower corrosion susceptibility of the Deloro-40 HF material more time required for a passive scale to form. But once formed, becomes a more passive system. The Stellite-6 HF material also had a very low LPR corrosion rate and trended
below 0.0254 mm/y (1 MPY) for the duration of the test. The Stellite-6 HF measured a negative weight loss reading of -10.6 mg. This was not expected as it indicates that the sample had gained weight and more scale had formed than Stellite-6 HF material corroded. Similar to the Deloro-40 HF trend, it took several days for peak corrosion rates to be reached before decreasing because of the time required for a passive scale to form.

Both the Deloro-40 HF and Stellite-6 HF materials were strong performers in resisting corrosion. To better differentiate the Deloro-40 HF and Stellite-6HF materials for use in application, further CPP and potentiostatic polarization testing was completed to better understand the materials tendency to pit and ability to re-passivate should it occur. A summary of the results are listed in Table 4.3 and Table 4.4 with the Deloro-40 HF and Stellite-6 HF CPP data trends shown in Figure 4.4 and Figure 4.5.

**Table 4.3: CPP testing summary**

<table>
<thead>
<tr>
<th>Material</th>
<th>Pit Potential (mV vs Eoc)</th>
<th>Apex Potential (mV vs Eoc)</th>
<th>Re-passivation Potential (mV vs Eoc)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deloro-40 HF</td>
<td>82</td>
<td>581</td>
<td>-3.9</td>
</tr>
<tr>
<td>Stellite-6 HF</td>
<td>194</td>
<td>635</td>
<td>60</td>
</tr>
</tbody>
</table>

**Table 4.4: PP testing summary**

<table>
<thead>
<tr>
<th>Material</th>
<th>Initial Current Density (uA/cm²)</th>
<th>Final Current Density (uA/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deloro-40 HF</td>
<td>500.5</td>
<td>11.7</td>
</tr>
<tr>
<td>Stellite-6 HF</td>
<td>236.2</td>
<td>-29</td>
</tr>
</tbody>
</table>
The Deloro-40 HF trend shows an initially gradual linear response to the applied anodic potential. An inferred pitting potential ($E_{pit}$, point of pitting initiation) is seen at 82 mV versus the Open Circuit Potential (OCP, $E_{OC}$). Increasing anodic potentials were
applied to the electrode until a peak current density was achieved. The point at which this occurred for the Deloro-40 HF was of 5 mA/cm² and 581 mV, this is the apex potential. The potential was then swept in a cathodic direction to 100 mV below (cathodic to) the EOC. The reverse sweep for the Deloro-40 HF material did not intersect with the anodic sweep. When the reverse sweep does not intersect with the anodic sweep, this indicates a positive hysteresis loop and that if any pitting were to occur that it would progress in an aggressive manner as the material is less likely to quickly form a passive film [69].

The Stellite-6 HF material had a more prominent pitting potential of 194 mV as shown in Figure 4.5. Although the rapid increase in current density over a relatively small potential change is indicative of a tendency to pit, the potential at which this occurred is more than double that of the Deloro-40 HF indicating it to be a more resistant alloy to corrosion and pitting. Additionally, a higher apex potential of 635 mV was observed, as in Figure 4.5, for the Stellite-6 HF in comparison to the Deloro-40 HF. This indicates the material is able to withstand more anodic (aggressive) potentials before the peak current density is reached and is indicative of a more noble metal. During the cathodic sweep the plotted line intersected with that of the anodic sweep indicating a negative hysteresis loop. This type of loop is indicative of a more corrosion resistant material, as any pitting that may occur is likely to passivate rather than progress further [70, 71]. The point at which current density reaches its lowest value on the reverse scan is labelled as the re-passivation potential (E_{Pr ot}). The Stellite-6 HF material displayed a higher E_{Pr ot}, which again is indicative of a material that will quickly re-passivate and prevent further pit growth.
The potentiostatic polarization testing results in Figure 4.6 show the Deloro-40 HF material had a higher initial current density in response to the constant applied potential of 60 mV than did the Stellite-6 HF. Its current density also trended higher for most of the test. The downward trend towards lower current density values indicates the formation and presence of a passivating film. In both the LPR and PP testing corrosion rates started increasing after 10 hours of testing before reaching peak corrosion rates/current densities and slowly trending downwards.

The Stellite-6 HF had an initial current density of 236.2 μA/cm², roughly half that of the Deloro-40 HF. The current response trended down throughout and eventually read negative (cathodic) current values. These cathodic current values are unexpected in response to an applied anodic potential but is likely due to the presence of a highly passivating and resilient corrosion scale on the electrode surface.

**Figure 4.6: Deloro-40 HF and Stellite-6 HF potentiostatic test data**
4.1.2 Downhole material testing results

Table 4.5 and Figure 4.7 summarize the LPR corrosion rate data for both the K-55 and TN-55TH materials.

Table 4.5: Downhole material LPR corrosion rate summary

<table>
<thead>
<tr>
<th>Electrode Material</th>
<th>LPR Corrosion Rate</th>
<th>Weight Loss</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Initial (MPY/mm/y)</td>
<td>Final (MPY/mm/y)</td>
</tr>
<tr>
<td>K-55</td>
<td>7.42/0.188</td>
<td>9.57/0.243</td>
</tr>
<tr>
<td>TN-55TH</td>
<td>7.74/0.196</td>
<td>9.6/0.243</td>
</tr>
</tbody>
</table>

Figure 4.7: LPR K-55 and TN-55TH corrosion rate vs time

Both of the LPR corrosion rate trends initially increase rapidly, followed by a decrease, and then slowly trend upwards to a stable and equilibrium corrosion rate. The corrosion mechanism present is believed to be that of a solid-state reaction where initially a very adhesive, thin, and protective scale forms. This is followed by a sequence of internal stresses cracking the scale causing a re-growth which leads to a secondary less passive outer scale to form [72] and why the corrosion rates slowly increased before coming to a slightly higher and stable corrosion rate.
The LPR corrosion rate for the K-55 material trended from an initial 0.188 mm/y (7.42 MPY) up to a final stable corrosion rate of 0.243 mm/y (9.57 MPY). The low and stable LPR trend indicates the presence of a passivating corrosion scale. The electrodes had an overall moderate surface etch with a corresponding weight loss corrosion rate of 0.298 mm/y (11.73 MPY). There was no indication of localized corrosion activity on the electrode surface as seen in Figure 4.8.

The TN-55TH sample had a very similar LPR trend to that of K-55. The LPR corrosion rate trended higher for the majority of the test but they were nearly identical by the end of the exposure period. The electrode weight loss was slightly higher at 0.319 mm/y (12.56 MPY) which corresponds well with the slightly higher LPR trend. There was no indication of localized corrosion activity as the electrodes had a similar overall moderate surface etch as seen in Figure 4.8.

These trends are opposite to that of the 1018 CS trends found in the wellhead testing. A difference between these tests is the pH of the brine. The wellhead materials brine was pH 5.50 and the downhole testing at pH 4.36. The lower pH can delay the formation of a passive scale and allow corrosion rates to increase until the corrosion product solubility limit is reached for a protective film to form.
Table 4.6 summarizes the crevice corrosion rate test data for the K-55+TH-55TH, and K-55+GLV-J55 crevice couples and Figure 4.9 shows post-test images.

Table 4.6: Downhole material crevice corrosion rate testing summary

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Corrosion Rate (MPY/mm/y)</td>
<td>9.5/0.241</td>
<td>9.69/0.246</td>
<td>10.87/0.276</td>
</tr>
</tbody>
</table>
The K-55+TN-55TH coupon pair had very similar weight loss corrosion rates with the K-55 at 0.241 mm/y (9.50 MPY) being slightly lower than the 0.246 mm/y (9.69 MPY) for the TN-55TH. Although the TN-55TH had a slightly higher weight loss corrosion rate, the difference in corrosion rate is not significant enough to determine its corrosion performance versus that of a K-55 alloy. The two coupons had a similar overall
moderate surface etch. No localized or preferential attack was noted at the point of contact or crevice formed by the coupon pair as Figure 4.9 indicates. This is unexpected as creviced conditions are known to accelerate corrosion rates. The similar corrosion rates and no visual localized attack is likely due to the relatively short exposure period, similar materials, and static testing conditions. These factors will not allow for enough of a difference in the bulk fluid to localized area fluid properties and therefore variations in corrosion rates between the open and creviced surfaces.

Crevice corrosion testing indicated the K-55+ GLV-J55 coupon pair had very different weight loss corrosion rates. The K-55 weight loss corrosion rate of 10.87 MPY (0.276 mm/y) was double that of the GLV-J55 at 5.95 MPY (0.151 mm/y). The K-55 had an overall moderate surface etch with a shinier appearance due to the formation of a Zn based film. The low weight loss corrosion rate observed in the GLV-J55 coupon corresponds well with the observed physical condition of the post-test coupon in Figure 4.9. There was no visually identifiable localized attack on either coupon. The low corrosion rate noted on the galvanized coupon is likely due to zinc passivation. Galvanized materials protect the base metals in two ways, being a barrier and the formation of a zinc patina [73]. The zinc patina is formed by the conversion of the galvanized coating into corrosion products. Typically zinc oxide (ZnO) and zinc hydroxide (Zn(OH)₂) are initially formed. Later in the corrosion cycle these products interact with carbon dioxide in the environment to form zinc carbonate (ZnCO₃). ZnCO₃ is a passive, stable film that adheres to the galvanized surface to protect the base galvanized material. It is not water soluble and will not dissolve into the bulk fluid. In the conditions tested the galvanized zinc coating does not react with H₂S to form a zinc
sulfide (ZnS) and hydrogen gas, or ZnO reacting with $H_2S$ to form a zinc sulfide and water. The temperatures of formation for these products are $>450^\circ$C, which is typically not found in oil and gas wells [74]. Also under the conditions tested ZnS is insoluble which does not promote the corrosion process to occur. Weight loss corrosion rates on the galvanized components should be taken with caution due to the removal of the galvanized coating when cleaning off the scales.

Similar to the K-55+TN-55TH pairing, the K-55+K-55 coupon pair had nearly identical weight loss corrosion rates of 8.48 MPY & 8.49 MPY (0.215 mm/y). Neither coupon in this pair displayed any localized or preferential corrosion attack due to the artificial crevice. Both coupons had an overall moderate surface etch as Figure 4.9 depicts. This is again expected due to the materials similarity.

Table 4.7 summarizes the galvanic corrosion rate test results and Figure 4.10, Figure 4.11, and Figure 4.12 illustrate the test period trends and post-test surface images.

| Table 4.7: Downhole material galvanic testing corrosion rate summary |
|-------------------------|---------------------|---------------------|
| **Coupon Pair** | **K-55+TH-55TH** | **K-55+ GLV-J55** |
| **Individual Coupon** | K-55 | TN-55TH | K-55 | GLV-J55 |
| **Corrosion Rate (MPY/mm/y)** | 18.18/0.462 | 17.87/0.454 | 10.11/0.257 | 9.3/0.236 |
Figure 4.10: Galvanic current density vs time

Figure 4.11: Galvanic coupled open circuit potential vs time
The K-55+TN-55TH galvanic pair resulted in a fluctuating galvanic current graph. The measured current density fluctuated from an initial 18.54 μA/cm² to -6.55 μA/cm² over the first 10 hours as Figure 4.10 indicates. The current then trended up
towards a net zero current flow between the coupled electrodes. Positive current readings indicate a net current flow from the K-55 to the TN-55TH. Negative current readings indicate a net current flow from the TN-55TH to the K-55 coupon. The current density readings are low and do not indicate a strong galvanic interaction where one metal is clearly corroding preferentially over the other. Galvanic corrosion typically results in significantly higher forward current flow. The coupled OCP versus the reference electrode increased from an initial -387 mV to stabilize around -140 mV (Figure 4.11). A less anodic potential generally indicates decreasing corrosivity. In harsh environments a potential difference < -150 mV is desirable, which is observed here [75].

The weight loss corrosion rates of 18.18 MPY (0.462 mm/y) for the K-55 and 17.87 MPY (0.454 mm/y) for the TN-55TH are higher than those recorded in the LPR and crevice tests. This increase is likely partially due to the reduced exposure time in the galvanic testing of 90 hours versus 165.5 hours in the LPR and crevice testing.

The current flow between the K-55 and GLV-K55 was very low. It trended from -0.76 μA/cm² to +0.58 μA/cm². Positive current readings indicate a net current flow from the K-55 electrode while negative readings correspond to a net current flow from the GLV-K55 electrode. The low current reading trends recorded indicate galvanic corrosion between the two materials was minimal. The coupled K-55 and GLV-K55 OCP stabilized at ~ -40mV versus the reference electrode. This potential is significantly less anodic than the K-55+TN-55TH couple. This is expected based on the galvanized zinc coating and formation of ZnO as noted above. The weight loss of the K-55 electrode when coupled with the GLV-K55 was significantly less than the K-55 electrode when coupled with TN-55TH as was displayed in Table 4.6. Again, the galvanized coating
further limited any galvanic current, and corrosion, from occurring between the dissimilar galvanized finish to the K-55 sample.

4.1.3 Lab testing discussion and summary

Both the Deloro-40 HF and Stellite-6 HF were strong performers in the LPR and weight loss evaluation methods compared to the 1018 carbon steel. Weight loss testing indicated close to negligible corrosion rates over the 7-day testing period and final LPR corrosion rates of 0.035 mm/y (1.38 MPY) and 0.024 mm/y (0.95 MPY) for the Deloro-40 HF and Stellite-6 HF materials respectively. This compared to the 1018 carbon steel having a weight loss corrosion rate of 0.25 mm/y (9.66 MPY) and 0.38 mm/y (15.05 MPY) for a final LPR corrosion rate reading. It was difficult to separate the performance of the Deloro-40 HF and Stellite-6 HF materials in the initial weight loss and LPR tests. The Stellite-6 HF showed only a slightly improved overall LPR trend. To differentiate further their performance CPP and potentiostatic polarization testing was completed. In the CPP testing the Stellite-6 HF material had superior performance with higher apex of 635 mV and pitting potential of 194.0 mV compared to the Deloro-40 HF with apex and pitting potentials of 581.0 mV and 82.0 mV, respectively. Further re-passivation potentiostatic polarization testing also displayed the Stellite-6 HF materials superior performance. The Stellite-6 HF samples initial PP current density was measured to be 236.2 µA/cm² and trended down for the 4-day test to eventually read cathodic (negative) values to end at -29.0 µA/cm². Whereas the Deloro-40 HF material had an initial current density close to double the Stellite-6 HF of 500.0 µA/cm² with a final reading of 11.7 µA/cm² and remaining anodic throughout the 4-day test period.
In the second set of autoclave testing, downhole completion materials were analyzed in 3 testing setups under SAGD conditions. General corrosion rate testing was done on K-55 and TN-55TH. Crevice corrosion testing was completed on 3 material combinations including K-55+TH-55TH, K-55+GLV-J55, and K-55+K-55, and finally galvanic corrosion testing on the materials and combinations of K-55+TH-55TH and K-55+GLV-K55.

The K-55 and TN-55TH general weight loss corrosion rate measurements indicated mild to moderate corrosion rates of 0.297 mm/y (11.73 MPY) and 0.319 mm/y (12.56 MPY), respectively. The LPR corrosion rates for both alloys were relatively stable and trended closely. Both showed final corrosion rates of 0.243 mm/y (9.60 MPY). Testing results and trends indicate that a passive corrosion scale formed to protect the electrodes from the aggressive conditions and that both materials share similar amounts of corrosion susceptibility.

There was also no evidence of increased corrosion due to the presence of an artificial crevice in any of the three crevice corrosion tests. The K-55 coupons corrosion rates showed similar results in each of the three crevice setups. The average corrosion rate for the K-55 coupons was 0.237 mm/y (9.34 MPY). The TN-55TH had a corrosion rate of 0.276 mm/y, similar to that of the K-55, with no localized corrosion due to crevice attack visually apparent. The GLV-J55 coupon had the lowest corrosion rate, close to half of the K-55 it was creviced with, at 0.137 mm/y (5.41 MPY). This likely indicates the galvanizing coating protected the surface of the coupon from corrosion. No visual localized corrosion damage was noted at the crevice.
Galvanic testing results showed there was no galvanic effect between the K-55 and TN-55TH materials, the corrosion rates were 0.462 mm/y (18.18 MPY) and 0.454 mm/y (17.87 MPY) respectively. This is expected as they have a similar chemical composition, mechanical, and physical properties. A slightly lower corrosion rate was found on the GLV-K55 coupon at 0.236 mm/y (9.3 MPY) compared to 0.257 mm/y (10.11 MPY) on the K-55 coupon. The current flow between the K-55 and GLV-K55 was very low indicating minimal galvanic corrosion between the two materials.

These results show low to moderate corrosion rates could be expected under the conditions tested. Lab testing corrosion rates of 0.254 mm/y (10 MPY) for an uninhibited specimen exposed to test brine could be considered a severe corrosion rate. The K-55+TN-55TH galvanic tests showed corrosion rates up to 0.462 mm/y (18.18 MPY). Although these rates are considered severe, they are not overly concerning based on the field testing, and past field failure analysis. If corrosion rates of 0.254 mm/y (10 MPY) are found in the field, it would be rated as high and of concern. In a lab, the coupons are exposed over a short duration relative to field monitoring which artificially increases the resulting corrosion rate. Other conditions in the field can also help to prevent corrosion, including the presence of oil which acts as a corrosion inhibitor, or other scales. The galvanized materials appeared to prevent corrosion due to the formation of a zinc patina which prevents direct material exposure and was shown to decrease corrosion currents and potentials to levels much less than would be required for corrosion to occur.

In the future, based on economics and funding, completing XRD analysis on lab samples would be beneficial in confirming the corrosion scale compounds present in each test. Specifically confirming the type of Zn filming compound formed.
4.2 Field testing results and analysis

In this section results and analysis of the downhole corrosion coupons are presented and discussed. Analysis completed included weight loss corrosion rates, compositional XRD, EDS, and SEM morphology analysis.

The corrosion coupons were removed from the well on Feb 19th, 2017 (after 330 days of exposure) as per the installation and removal procedure created before the coupons were installed [76].

4.2.1 Downhole material pre and post-test acid washing images

Figure 4.13 - Figure 4.19 show the coupons as received before and after acid cleaning.

![Figure 4.13: Joint 2 Corrosion coupons as received](image)
Figure 4.14: Joint 2 Corrosion coupons post cleaning

Figure 4.15: Joint 23 Corrosion coupons as received
Figure 4.16: Joint 23 Corrosion coupons post cleaning

Figure 4.17: Joint 25 Corrosion coupons as received
Figure 4.18: Joint 25 Corrosion coupons post cleaning

Figure 4.19: Joint 12 and 24 coupons as received (UofC)
These images show similar surface physical characteristics on each coupon materials at the various depths. The lower coupons from the joint 2 and 12 clamps appear to be darker in color compared to the coupons on the joint 23, 24, and 25 clamps. This is likely due to the fluid separation occurring downhole. The lower coupons are likely exposed to an oil + water + gas emulsion or oil on its own, depending on the levels at which fluid separation occurs. The coupons at higher well elevations are exposed to the separated produced water and other less dense compounds in the fluid. At production temperatures and pressures water (~864 kg/m$^3$ [77]) is less dense than oil (~878.1 kg/m$^3$), causing water to be the top fluid in the wells annulus once emulsion separation occurs.

4.2.2 Corrosion rate results and analysis

The following section outlines the corrosion rate calculation and analysis completed on the installed coupons. For calculating corrosion rates a density of 7.86 g/cm$^3$ [78] was used for the J-55 and L-80 coupons. Calculated corrosion rates are all average, uniform, weight loss corrosion rates based on the coupons pre and post-test weigh changes and that they all visually appeared to have corroded uniformly over the entire surface. Select L-80 coupons at higher elevations in the well appear to have experience pitting corrosion. For these corrosion rate calculations, pitting corrosion rate calculation methods were used as per NACE Standards [63].

The reported corrosion rates for the coupled J-55+GLV-J55 coupons should also be considered with care. This is due to difficulty in calculating the exposed surface area for these coupons considering the way in which the J-55 was coupled with the GLV-J55 coupons. Also, only average weight loss corrosion rates were calculated on the
galvanized samples, pitting analysis and pitting corrosion rates were not considered or reported.

Figure 4.20 - Figure 4.23 displays the weight loss and corrosion rate testing results on all coupons. Figure 4.20 and Figure 4.21 show the coupons weight loss and corrosion rate at the various depths.

Figure 4.20: Weight loss by depth and varying material
Figure 4.21: Corrosion rate by depth and varying material

These results show that the single J-55 material experienced the highest corrosion rate at each depth, followed by the L-80 coupons, and finally the second J-55 coupon coupled with the GLV-J55. The much higher corrosion rate and weight loss results for the GLV-J55 coupons is likely due to the sacrificial nature of the zinc coating to the coupled J-55 coupon as well as higher dissolution rate of zinc in hydrochloric acid during the acid cleaning process. For that reason it is difficult to compare its corrosion rate with the others. The galvanized samples appear to experience a similar “corrosion” rate at each installation location.

Figure 4.22 and Figure 4.23 show each materials weight loss and corrosion rate at different well depths.
Figure 4.22: Weight loss by material and varying depth

Figure 4.23: Corrosion rate by material and varying depths

These results show the highest uniform corrosion rates will occur at lower well elevations, closer to the ESP, on all materials and decrease moving up the well from the
ESP to approximately the liquid level. Corrosion rates actually increase on the joint 25 (196.17mKB) clamp coupons for each material. The location/area of the increase is often where past field failures have been found to occur and could be due to higher concentrations of aqueous acidic gases and more a more conductive electrolite with less oil present to coat the metal surfaces. If the coupons were above the liquid level, condensates formed at high well elevations can contact the coupons and cause higher rates of localized corrosion.

The average corrosion rate on the bottom joint 2 coupons was 0.0228 mm/y, and 0.0119 mm/y for the top joint 25 coupons. At each depth the non-coupled J-55 experienced the highest corrosion rates, followed by the L-80, and finally the second J-55 coupon coupled with the GLV-J55 coupon. With the exception of the J-55 coupon from the joint 2 clamp at 505.15mKB (lowest depth), all corrosion rates can be classified as “low” based on standard NACE standards.

The L-80 coupons at higher elevations in the well appear to have experienced pitting corrosion as shown in Figure 4.15. Pitting corrosion rates on the L-80 coupons were 0.113 mm/y (4.45MPY) on the joint 25 clamp at 196.17mKB, and 0.225mm/y (8.85MPY) on the joint 23 clamp at 223.25mKB. The coupons lower in the well did not appear to have suffered from pitting corrosion. It is likely that at higher well elevations the more aqueous environment, increased fluid movement due to vapor flashing, and possible creviced and galvanic conditions could all promote an environment more susceptible to pitting. Pitting corrosion is often of greater concern than uniform corrosion as its rates are higher and often leads to catastrophic material failures [79, 80].
4.2.3 Corrosion scale results and analysis

Visual inspections on the coupons installed in the joint 2, 23, and 25 clamps at depths 505.15 mKB, 223.47 mKB, and 196.17 mKB respectively are documented and presented in Table 4.8 to Table 4.10 before and after acid cleaning. XRD analysis was completed on the J-55 coupons at each depth. XRD analysis was only completed on the one non-coupled J-55 coupon at each location due to its cost.

Visual inspection comments on the joint 12 and 24 clamp coupons at 372.47 mKB and 209.76 mKB are show in Figure 4.24 to Figure 4.31.

Table 4.8: Joint 2 (505.15mKB) coupon observations

<table>
<thead>
<tr>
<th>Alloy Type</th>
<th>Description of Coupon</th>
<th>After Solvent Wash</th>
<th>After Acid Cleaning</th>
<th>Corrosion Scales</th>
</tr>
</thead>
</table>
| J-55       | - Front: Covered with thin grey scale  
             - Back: Covered with thin grey and black scale | General Corrosion | - Pyrrhotite 
             - Magnetite 
             - Quartz |
| L-80       | - Front: Covered with thin grey scale 
             - Back: Mainly covered with thin grey scale | General Corrosion | N/A |
| J-55       | - Front: Covered with thin grey scale 
             - Back: Mainly covered with thin grey scale and heavier deposits on the edges | General Corrosion | N/A |
| GLV-J55    | - Front/Back: Mainly covered with thin grey scale | General Corrosion | N/A |
### Table 4.9: Joint 23 (223.47mKB) coupon observations

<table>
<thead>
<tr>
<th>Alloy Type</th>
<th>Description of Coupon</th>
<th>Corrosion Scales</th>
</tr>
</thead>
</table>
| J-55       | - **Front:** Mainly covered with thin, black scale with some thick, white deposits  
             - **Back:** Mainly covered with thin, black scale with some thick, white deposits | Etched surface with patchy corrosion  
             - Cristobalite  
             - Quartz  
             - Pyrrhotite |
| L-80       | - **Front:** Mainly covered with thick white deposits  
             - **Back:** Mainly covered with thin, black scale and some areas of thick, white deposits | Patchy corrosion with some pitting  
             N/A |
| J-55       | - **Front:** Mainly covered with thin, grey and black scale with some thick, white deposits  
             - **Back:** Mainly covered with thin, grey scale with some thick, white deposits | General corrosion with patchy attack adjacent to the edges  
             N/A |
| GLV-J55    | - **Front:** Mainly covered with thin, black scale with some thick, white deposits  
             - **Back:** Mainly covered with white deposits | General Corrosion  
             N/A |
### Table 4.10: Joint 25 (196.17mKB) coupon observations

<table>
<thead>
<tr>
<th>Alloy Type</th>
<th>Description of Coupon</th>
</tr>
</thead>
</table>
| J-55       | After Solvent Wash:  
              - **Front:** Mainly covered with thick, white and brown deposit  
              - **Back:** Heavily covered with brown deposits  
              After Acid Cleaning: Spots and Patches of Corrosion  
              Corrosion Scales: - Cristobalite  
              - Quartz |
| L-80       | After Solvent Wash:  
              - **Front:** Mainly covered with thick, white deposits  
              - **Back:** Partially covered with thick, white deposits  
              After Acid Cleaning: General corrosion with some patches of attack on the front  
              Corrosion Scales: N/A |
| J-55       | After Solvent Wash:  
              - **Front:** Partially covered with thick, white deposits  
              - **Back:** Covered with white and brown deposits  
              After Acid Cleaning: General corrosion with some patches of attack on the front  
              Corrosion Scales: N/A |
| GLV-J55    | After Solvent Wash:  
              - **Front:** Mainly covered with thin, black scale  
              - **Back:** Mainly covered with thin, black scale and some white deposits  
              After Acid Cleaning: General Corrosion  
              Corrosion Scales: N/A |

All of the coupons show that before acid cleaning a darker grey, brown and or black scale, are present with some samples showing white deposits. It is known that iron sulfide scales, mackinawite, pyrite, triolite, pyrrhotite, have a darker appearance. Mackinawite is grey or black in color, while pyrrhotite has a more bronze/brown appearance [81, 82]. It is also known that magnetite scales have a gray with brownish appearance [83, 84]. Magnetite was only found on the joint 2 J-55 coupon. Although it is not an expected corrosion product it could be formed due to the presence of CO₂ and it being released from the emulsion closer to the ESP at higher concentrations. It is also possible that it converted to magnetite from a sulfide based scale as a result of being exposed to oxygen after being removed from installation [85]. Magnetite has been noted
in past corrosion inspection failure analysis reports [86]. The white scales are likely those of silicate compounds, quartz and/or cristobalite which have a clear or white color [87-90].

After cleaning, except for the L-80 coupons on the higher elevation clamps, the coupons appeared to corrode uniformly. The L-80 coupons located on the higher clamps experienced visible localized pits on the front and side edges. This is similar to past failure analysis and inspection reports results [17] which also indicate pitting corrosion to occur more prolifically at higher elevations in wells. Relative to the other materials, L-80 is a stronger and harder material with no corrosion resisting alloying components and can be more susceptible to pitting corrosion as the pit tip is less likely to re-passivate [79, 80].

SEM images for all coupons on the joint 12 and 24 clamps are shown in Figure 4.24 to Figure 4.31. The J-55 coupon SEM images from these joints are shown in Figure 4.24 and Figure 4.25.
Figure 4.24: Joint 12, J-55 at 372.47 mKB SEM; (Coupon 20)

Figure 4.25: Joint 24, J-55 at 209.76 mKB SEM; (Coupon 16)
These images show that the lower joint 12 J-55 coupon has less scale, corrosion product present and appears to have corroded more than the joint 24 sample. The joint 24 sample was also lighter in appearance and had more complete scale coverage over the samples surface.

Imaging on the L-80 coupons, Figure 4.26 and Figure 4.27, indicate a similar pattern as the J-55 images with the lower joint 12 coupon having a darker, less complete scale coverage and the occurrence of a more uniform corrosion.

Figure 4.26: Joint 12, L-80 at 372.47 mKB SEM; (Coupon 9)
The second J-55 coupons coupled with the GLV-J55 SEM results in Figure 4.28 and Figure 4.29. These indicate a flakier, less compact, scale buildup over a section of the lower joint 12 coupon sample compared to the joint 24 coupon. The joint 24 coupon shows a dense and compact looking structure.
Figure 4.28: Joint 12, J-55 at 372.47 mKB SEM; (Coupon 49)

Figure 4.29: Joint 24, J-55 at 209.76 mKB SEM; (Coupon 33)
Finally the GLV-J55 joint 12 and 24 coupon images in Figure 4.30 and Figure 4.31, show a rougher and more corroded surface on the lower joint 12 coupon as compared to the lighter color and less corroded joint 24 sample.

Figure 4.30: Joint 12, GLV-J55 at 372.47 mKB SEM; (Coupon 47)
These results appear to be in line with the corrosion rate testing analysis. The lower coupons all appeared to have less scale product present which would lead to lesser corrosion protection and a higher corrosion rate. A similar trend was noted on corrosion rates as the joint 12 samples appeared to corrode more compared to the higher ones at joint 24 [91]. XRD results are presented from Table 4.11 to Table 4.21. The intensity plots are also presented from Figure 4.32 to Figure 4.39.

Table 4.11: Joint 2, coupon #8, J-55, position #1, deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>65-75%</td>
</tr>
<tr>
<td>Magnetite</td>
<td>Fe₃O₄</td>
<td>10-20%</td>
</tr>
<tr>
<td>Quartz</td>
<td>SiO₂</td>
<td>5-10%</td>
</tr>
<tr>
<td>Unidentified</td>
<td>N/A</td>
<td>1-5%</td>
</tr>
</tbody>
</table>
Table 4.12: Joint 12, coupon #9, L-80, position #2 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>75-85%</td>
</tr>
<tr>
<td>Cristobalite</td>
<td>SiO₂</td>
<td>15-25%</td>
</tr>
</tbody>
</table>

Figure 4.32: Joint 12, coupon #9, L-80, position #2 deposit analysis

Table 4.13: Joint 12, coupon #20, J-55, position #1 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>70-80%</td>
</tr>
<tr>
<td>Cristobalite</td>
<td>SiO₂</td>
<td>4-10%</td>
</tr>
<tr>
<td>Unidentified</td>
<td>N/A</td>
<td>10-20%</td>
</tr>
</tbody>
</table>
Table 4.14: Joint 12, coupon #47, GLV J-55, position #3 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinc Oxide</td>
<td>ZnO</td>
<td>70-80%</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>2-8%</td>
</tr>
<tr>
<td>Cristobalite</td>
<td>SiO$_2$</td>
<td>2-8%</td>
</tr>
<tr>
<td>Unidentified</td>
<td>N/A</td>
<td>10-20%</td>
</tr>
</tbody>
</table>

Figure 4.33: Joint 12, coupon #20, J-55, position #1 deposit analysis

Figure 4.34: Joint 12, coupon #47, GLV J-55, position #3 deposit analysis
Table 4.15: Joint 12, coupon #49, J-55, position #3 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinc Oxide</td>
<td>ZnO</td>
<td>80-90%</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>2-8%</td>
</tr>
<tr>
<td>Unidentified</td>
<td>N/A</td>
<td>2-8%</td>
</tr>
</tbody>
</table>

Figure 4.35: Joint 12, coupon #49, J-55, position #3 deposit analysis

Table 4.16: Joint 23, coupon #9, J-55, position #1 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cristobalite</td>
<td>SiO₂</td>
<td>75-85%</td>
</tr>
<tr>
<td>Quartz</td>
<td>SiO₂</td>
<td>10-20%</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>5-10%</td>
</tr>
</tbody>
</table>

Table 4.17: Joint 24, coupon #18, L-80, position #2 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cristobalite</td>
<td>SiO₂</td>
<td>90-99%</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>1-10%</td>
</tr>
</tbody>
</table>
Figure 4.36: Joint 24, coupon #18, L-80, position #2 deposit analysis

Table 4.18: Joint 24, coupon #16, J-55, position #1 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cristobalite</td>
<td>SiO$_2$</td>
<td>90-99%</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>1-10%</td>
</tr>
</tbody>
</table>

Figure 4.37: Joint 24, coupon #16, J-55, position #1 deposit analysis
Table 4.19: Joint 24, coupon #35, GLV J-55, position #3 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinc Oxide</td>
<td>ZnO</td>
<td>80-90%</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>1-5%</td>
</tr>
<tr>
<td>Unidentified</td>
<td>N/A</td>
<td>5-15%</td>
</tr>
</tbody>
</table>

Figure 4.38: Joint 24, coupon #35, GLV J-55, position #3 deposit analysis

Table 4.20: Joint 24, coupon #33, J-55, position #3 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cristobalite</td>
<td>SiO₂</td>
<td>65-75%</td>
</tr>
<tr>
<td>Zinc Oxide</td>
<td>ZnO</td>
<td>15-25%</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>FeS</td>
<td>1-5%</td>
</tr>
<tr>
<td>Unidentified</td>
<td>N/A</td>
<td>4-10%</td>
</tr>
</tbody>
</table>
Figure 4.39: Joint 24, coupon #33, J-55, position #3 deposit analysis

Table 4.21: Joint 25, coupon #22, J-55, position #1 deposit analysis

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical Formula</th>
<th>Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cristobalite</td>
<td>SiO₂</td>
<td>65-75%</td>
</tr>
<tr>
<td>Quartz</td>
<td>SiO₂</td>
<td>15-25%</td>
</tr>
<tr>
<td>Unidentified</td>
<td>N/A</td>
<td>5-15%</td>
</tr>
</tbody>
</table>

These XRD results indicate that on the non-creviced coupons, not the J-55+GLV-J55, that silicate based scales and the iron sulfide compound pyrrhotite make up most the compounds at each depth. The silicate scales present are quartz and cristobalite. The one exception to this is on the joint 25 J-55 coupon in Table 4.21. Here the only compounds found are silicates and 5-15% of an unidentified compound. Magnetite was also identified on the lowest joint 2 J-55 coupon, not coupled with the GLV-J55 sample. Silicate scales, including quartz and cristobalite, are commonly found compounds in oil and gas wells, lines, and facilities as they are produced from oil and gas formations.
Cristobalite is a high-temperature polymorph of silica and has the same chemical formula as quartz, SiO$_2$, but a different crystal structure [90]. Field scale compositional analysis has shown quartz to be present downhole at multiple locations. Including further down the horizontal section of the well in the liner [92], at the ESP as the joint 2 coupon XRD results also show, or above the ESP as is seen on the coupon samples from joints 23 to 25.

ZnO represents the majority scale compound present on the GLV-J55 coupon as well as the J-55 coupon coupled with it to form a crevice. Galvanized materials are ones that are coated in zinc to reduce corrosion rates by forming a passive ZnO or ZnCO$_3$ scale. The galvanized coating is zinc based which is anodic to steel and would preferentially corrode when directly exposed to the environment. The rapid initial corrosion of the zinc would lead to local saturation of limits being reached and the formation of the passive scale, which if remaining intact, aids in preventing corrosion. The formation of this scale for protection and reducing corrosion processes appears to be the case here with the zinc reacting with the annular fluids to corrode in preference to the base J-55 and the J-55 coupon it was coupled with. This is also shown in the corrosion rate testing from Figure 4.20 to Figure 4.23. The J-55 coupon coupled with the GLV-J55 material corroded less than all others at each location.

The remaining coupons, with the exception of the joint 25 J-55 coupon, showed pyrrhotite to also be present in varying quantities. Pyrrhotites, or other iron sulfide compounds, are expected to occur as it is the likely corrosion reaction product of steel in the presence of aqueous H$_2$S at elevated temperatures [46, 93]. Field failure analysis has also shown iron sulfide scale products present. It is interesting to note that at lower well
elevations FeS appears to be the majority scale compound. Whereas at higher elevations it begins to decrease in quantity and at the top joint 25 coupon it is no longer observed as seen in Table 4.21. This compares well to the corrosion rate testing as at higher well elevations corrosion rates, and therefor corrosion product quantities, decreased. An exception is on the joint 25 coupons which showed an increased in corrosion rates but no FeS scale formation, only silicates and unidentified products. This could be due to corrosion occurring where there was no fluid to saturate FeS for scale formation, or that the unidentified portions are pyrrhotite. As discussed above in the literature review section, pyrrhotite is a corrosion product of carbon steel corrosion under sour conditions and is often found at moderate to high H\textsubscript{2}S concentrations where the corrosion kinetics for FeS scale formation are preferred to \( \text{FeCO}_3 \). Pyrrhotite is an iron sulfide mineral with the formula \( \text{Fe}_{(1-x)}S \) \((x = 0 \text{ to } 0.2)\). It is a nonstoichiometric variant of FeS, troilite [94]. Magnetite is a carbon steel corrosion product often found in systems where aqueous CO\textsubscript{2} is present as is the case in SAGD. The unidentified scale portions refer to one reflection on the diffraction pattern that did not appear to correspond to any of the known compounds [95, 96].

4.3 Testing summary and analysis

This section summarizes the lab autoclave and field testing using corrosion coupons representative of commonly used wellhead and various downhole completion materials.
4.3.1 Autoclave testing

Table 4.22 again outlines the materials tested in the autoclaves, measurements taken, and the type of corrosion being analyzed.

**Table 4.22: Autoclave testing summary**

<table>
<thead>
<tr>
<th>Wellhead Materials</th>
<th>Measurements</th>
<th>Test Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>1018 Carbon Steel</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td></td>
</tr>
<tr>
<td>Deloro-40 HF</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Cyclic Potentiodynamic Potential</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Potentiostatic Potential</td>
<td></td>
</tr>
<tr>
<td>Stellite-6 HF</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Cyclic Potentiodynamic Potential</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Potentiostatic Potential</td>
<td></td>
</tr>
<tr>
<td>J-55</td>
<td>- Weight Loss</td>
<td>- Crevice CR</td>
</tr>
<tr>
<td>K-55</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td>- Crevice CR</td>
</tr>
<tr>
<td></td>
<td>- Zero Resistance Ammeter</td>
<td>- Galvanic CR</td>
</tr>
<tr>
<td>TN-55TH</td>
<td>- Linear Polarization Resistance</td>
<td>- General CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td>- Crevice CR</td>
</tr>
<tr>
<td></td>
<td>- Zero Resistance Ammeter</td>
<td>- Galvanic CR</td>
</tr>
<tr>
<td>GLV-K55</td>
<td>- Linear Polarization Resistance</td>
<td>- Crevise CR</td>
</tr>
<tr>
<td></td>
<td>- Weight Loss</td>
<td>- Galvanic CR</td>
</tr>
<tr>
<td></td>
<td>- Zero Resistance Ammeter</td>
<td></td>
</tr>
<tr>
<td>GLV-J55</td>
<td>- Weight Loss</td>
<td>- Crevise CR</td>
</tr>
</tbody>
</table>

4.3.1.1 Autoclave wellhead material testing

Wellhead material autoclave testing verified 1018 carbon steels susceptibility to corrosion in the conditions tested. LPR measurements indicated corrosion rates between 0.38-1.79 mm/y with average weight loss corrosion rate of 0.25 mm/y. These rates are considered high relative to NACE standards. Deloro-40 HF, which is currently being
used as a gate valve gate face overlay material, experienced LPR corrosion rates between 0-0.024 mm/y with an average weight loss corrosion rate of 0.00076 mm/y which can be considered negligible due to instrumentation measurement error and accuracy limits [60]. Similarly, the Stellite-6 HF overlay experienced low LPR corrosion rates between 0.015-0.024 mm/y, and a negligible weight loss measurement for the same reason as the Deloro-40 HF [97]. To better compare the Deloro-40 HF and Stellite-6 HF materials 4-day CPP and potentiostatic polarization testing was completed to determine their susceptibility to pitting and ability to re-passivate should corrosion occur. These tests indicated that the Stellite-6 HF finish would provide superior protection. It had higher apex and pitting potentials found through CPP evaluations. Further, the re-passivation potential indicates if pitting were to occur, the Stellite-6 HF material should more readily re-passivate to reduce the depth and severity of attack. The Stellite-6 HF also showed a negative hysteresis loop indicating a more corrosion resistant material, compared to the positive loop in the Deloro-40 HF hysteresis loop.

4.3.1.2 Autoclave downhole completion material testing

Autoclave testing was then done on downhole completion materials in general, crevice, and galvanic corrosion setups. General 7-day corrosion rate testing was completed on K-55 and TN-55TH materials through LPR and weight loss analysis. The initial and final LPR K-55 coupon measurements showed rates between 0.188-0.243 mm/y, and a weight loss rate of 0.297 mm/y. The TN-55TH LPR measurements increased from 0.196 mm/y up to 0.243 mm/y with an average weight loss rate of 0.319 mm/y. Both the weight loss and LPR tests showed the TN-55TH material to have slightly
higher corrosion rates. The slightly higher rates can be considered insignificant as they are within measurement device reading accuracy tolerance limits.

In the autoclave crevice corrosion tests, weight loss measurements were utilized to determine the effects of crevices on corrosion rates in a SAGD environment after 7-day exposure. Three crevices setups were made between K-55+TN-55TH, K-55+GLV-J-55, and K-55+K-55 coupon couples. The K-55 and TN-55TH materials weight loss corrosion rates were determined to be 0.241 mm/y and 0.246 mm/y respectively, with moderate surface etching and no localized corrosion noticed at the creviced location. The similar rates are expected as these materials are very similar, they are both carbon steels but the TN-55TH is a patented carbon steel blend with slightly tighter chemistry tolerances compared to standard API K-55 and J-55 materials. The creviced couple of K-55+K-55 materials had identical rates of 0.215 mm/y each as expected given they are the same material. Finally, the K-55+GLV-J55 pair showed very different corrosion rates of 0.276 mm/y and 0.137 mm/y respectively with no localised or preferential attack on either coupon. It is likely that the GLV-J55 sample had reduced corrosion rates due to the formation of a passive ZnO or $\text{ZnCO}_3$ scale. The galvanized coating is zinc based which is anodic to steel and would preferentially corrode when directly exposed to the environment. The rapid initial corrosion of the zinc would lead to local saturation of limits being reached and the formation of the passive scale, which if remaining intact, aids in preventing corrosion.

Finally autoclave tests were completed to understand the effects of galvanic couples on corrosion rates. This testing was done on the same materials as in the crevice testing by measuring the potential difference between the two materials using a
potentiostat in ZRA mode over the 4-day test period. Here two electrode pairs were
galvanically coupled and corrosion rates monitored over a 4-day period. Positive current
readings represent a net current flow from the K-55 and negative current readings
represent a net current flow from the TN-55TH. Weight loss measurements were also
taken and corrosion rates were calculated. The electrode pairs evaluated were K-55+TN-
55TH and K-55+GLV-K55. The galvanic current density between the K-55+TN-55TH
pair varied considerably throughout the test from an initial 18.54 μA/cm$^2$ to -6.55 μA/cm$^2$
over the first 10 hours. The current flow then reversed indicating the TN-55TH electrode
was corroding preferentially. However, the coupled current remained relatively low
throughout and would not be considered high enough to indicate significant galvanic
corrosion was occurring due to dissimilar metal compositions. This is verified by the
corrosion rates found on both the K-55 and TN-55TH materials, with weight loss rates of
0.462 mm/y and 0.454 mm/y respectively. These are again similar materials and similar
rates are expected with no preferential corrosion. It is likely that the weight loss corrosion
rates are higher because of the shorter test duration compared to the previous crevice
corrosion testing. The K-55+GLV-K55 pair had a very low coupled galvanic current;
trending between -0.76 μA/cm$^2$ to 0.58 μA/cm$^2$. The low current readings indicate that
galvanic corrosion between the two was minimal. The coupled open circuit potential
stabilised at ~ -40 mV versus the Hastelloy reference. This potential is significantly less
anodic, corrosive, than the K-55/TN-55TH couple. The weight loss corrosion rates were
also lower for the K-55 and GLV-K55 samples at 0.257 mm/y and 0.236 mm/y
respectively. It is likely that the presence of the galvanized coating again quickly formed
a passive scale very quickly which protected the metals and kept corrosion rates and current densities low throughout the test period.

4.3.2 Field corrosion coupon testing

Corrosion rate and scale analysis was completed on the corrosion coupons installed in a producing SAGD well. XRD results showed the presence of pyrrhotite, quartz, cristobalite, zinc oxide, magnetite, and some unidentified compounds. Pyrrhotite, quartz, and cristobalite are present on most samples in varying concentrations. Zinc oxides are identified on the galvanized coupon sample sets as expected. The scales composition varied in a noticeable manner from the well bottom at the ESP to the liquid level. Coupons from the clamps at the bottom of the well on joints 2 and 12 showed the highest concentrations of pyrrhotite scales of between 65-85%, with 5-25% silicate scales (quartz and cristobalite), and up to 20% of an unidentified compound. Scales on the higher joint 23, 24, and 25 clamp coupons tended to show reduced pyrrhotite concentrations of between 1-10% pyrrhotite, and increased silicate scale concentrations 10-99%. The J-55 coupon in the highest joint 25 clamp showed only the presence of silicate scales, no iron sulfide compounds, with 5-15% of an unidentified compound. Magnetite was also noted on the bottom joint 2 clamp J-55 coupon. Magnetite is not an uncommon compound based on the presence of CO₂ and its ability to corrode carbon steels under the right conditions. It is also possible that it formed after being removed from the well if the pyrrhotite scale was damaged and CO₂ remained trapped under the scale.
The scale composition and concentration results correspond well to the corrosion rate data. On almost all coupon samples and setups corrosion rates decreased from the bottom of the well to the top of the set of coupons. Comparing this to the XRD scale analysis, lesser amounts of iron sulfide corrosion product were noted moving from the bottom to the top of the well. An exception to this was on the top joint 25 coupons. At this location no FeS corrosion products were noted and corrosion rates increased on all materials. It is difficult to determine the corrosion mechanism here given no corrosion products (FeS or \( FeCO_3 \)) were noted, only silicates. But because of the increase in corrosion rates relative to the joint 24 clamp coupons that they were partially in/out of the fluids and that the solubility limits in the surrounding environment were not reached, therefore no corrosion product scale formed.

Corrosion rates also decreased consistently with the standard J-55 material showing the highest rates, followed by the L-80, the second J-55 coupon coupled with the GLV-J55. In each case the galvanized material experienced much higher rates of corrosion due to its anodic properties, as well as to how it was cleaned for scale analysis.

Visual inspections indicated the presence of pitting on only the L-80 coupons which can likely be attributed to it being a harder and stronger material with higher carbon concentrations compared to the J-55, K-55, and TN-55TH materials. The scales were consistently darker in appearance with shades of black, grey, and brown, along with white compounds. Iron sulfide scale have a darker grey/black and brown appearance, magnetite is also a brownish scale, with silicates having a clear and white appearance.
4.3.3 Combined testing analysis and summary

At a first glance comparing the lab test corrosion rate results to the field coupon corrosion rates they do not appear to match. Lab results show much more aggressive corrosion rates compared to the field weight loss testing, up to 10 times higher. This is likely due to the much shorter test exposure period and is common when comparing field to lab data. The downhole coupons were tested for 330 days, compared to the 4 and 7-day lab testing. Also, the autoclave testing brines did not contain oil, a known corrosion inhibitor, and could have led to increased corrosion rates.

It is interesting to note that when comparing the lab results to past field failure analysis completed, some analysis compares well with this testing while other findings are not as expected. Wellhead material testing showed the Stellite-6 HF material to performing exceptionally well compared to carbon steel and better than the Deloro-40 HF. Select wellhead components used contain a Stellite-6 HF overlay, gate valve gate face seat components, have not shown signs of corrosion, like the testing results here. Whereas the gate faces finished with a Deloro-40 HF overlay have experienced pitting corrosion. For this reason it is recommended that a field trial be completed by replacing the Deloro-40 HF overlay with the Stellite-6 HF overlay on the gate valve gate face to determine its performance in real production conditions. It is also recommended that further autoclave and field testing be completed on a third tungsten carbide HF overlay material. This material has been proposed for use by different wellhead and valve suppliers but has not yet been testes in SAGD like conditions.

Lab and field testing and analysis on downhole completion materials have shown that downhole materials are susceptible to corrosion, specifically aqueous $H_2S$ corrosion
when subject to SAGD operating conditions. In general “low” rates of uniform corrosion were found and corrosion rates decreased at higher well elevations up to and just below the annular liquid level. At and just above the liquid level an increase in corrosion rates was found and can be expected in the form of both general uniform and pitting corrosion. These results compare to past downhole component failure analysis which has led to the belief that the most concerning area for corrosion is closer to the top of the annular liquid level due to the presence of water, not oil, and the highest concentrations of aqueous acidic gases. Testing has confirmed this with the “low” uniform corrosion rates calculated at all well depths, but an increase in uniform corrosion rates and formation of corrosion pits on select materials at the top coupons. Pitting corrosion is of much greater concern due to its more severe and often sudden consequences.

The “low” uniform corrosion rates at all locations are likely due to the formation of passive scales, the presence of oil at lower well elevations, as well as a reduction in aqueous acidic gas concentrations at lower well elevations due to the partitioning of the gases into the oil phase [98].

Galvanic and crevice corrosion do not appear to be of concern due to similar materials being used and proper material couples made between components to ensure that passive films are able to form. It is important that the films on galvanized materials remain intact, not removed chemically or mechanically, due to the anodic properties of Zn relative to most other metals. If the protective scales are continually removed severe localized corrosion can occur. This has been noted in past failure analysis.

Pitting corrosion at higher well elevations, at and above the liquid level, can be expected on susceptible materials, L-80 in testing, due to higher concentration of aqueous
H₂S, changes in fluid exposure and scale solubility, crevices, and a slightly more dynamic fluid environment with the occurrence of vapor flashing and condensation.

Based on this testing the development and injection of SAGD specific corrosion inhibitor is recommended with a focus should be placed on inhibiting the lower and upper portions of the wells annulus. Different inhibitors and application procedures should be used based on the location of injection and conditions present. It is recommended that to inhibit the lower portion of the well that an inhibitor be injected down the bubbletube, and the upper portion through inhibitor or scavenger injection at the wellhead. It is also recommended that a review of materials being used at higher well elevations be completed to avoid materials susceptible to pitting corrosion.
Chapter Five: Mechanistic aspects of metal corrosion in H$_2$S and CO$_2$ SAGD environments

The final objective of this research was to determine the corrosion mechanisms present in SAGD given the conditions tested. Literature review has shown that limited to no corrosion mechanistic analysis or research has been completed under SAGD type conditions comparable to Firebag. SAGD temperatures can range between 150°C to 240°C, and relatively high concentrations of CO$_2$ and H$_2$S are present. This research shows that the dominating corrosion mechanism present under these environments is that of aqueous H$_2$S corrosion based on the scale compositions, corrosion morphology, past failure analysis, as well as literature review [17, 35, 50, 99]. Because of the elevated concentrations of CO$_2$, one might expect CO$_2$ corrosion to be a dominating mechanism. This testing showed that due to the low ratio of CO$_2$/H$_2$S and the fact that iron sulfide is less soluble than iron carbonate under these conditions, aqueous H$_2$S corrosion is the dominating mechanism [31, 36, 41, 53, 54, 98]. It could also be expected that fluid compositional changes alter the dominating mechanism due to electrolyte property and condition changes; this did not appear to be the case.

As found and noted in the literature review section 2.3.2, the mechanism for hydrogen sulfide corrosion is that of a solid-state reaction mechanism [41, 43, 52, 72, 81] leading to the formation of an iron sulfide corrosion product. Mackinawite is often the first scale formed on the surface, but can quickly transform into alternate forms of iron sulfide products including amorphous ferrous sulfide, cubic ferrous sulfide, smythite, greigite, pyrrhotite, troilite and pyrite. Mackinawite often forms at lower temperatures (T<135°C), but at the elevated SAGD temperatures can rapidly and iso-chemically transition into alternate amorphous iron sulfide products including pyrrhotite as was
found in the testing. It is important to differentiate the type of scale formed as it can influence the corrosion kinetics and rate of corrosion. The following section further discusses this mechanism, factors affecting it, and its implications in SAGD.

5.1 Mechanism of hydrogen sulfide corrosion

The probable mechanism of H₂S corrosion on steel is that of a direct solid state reaction and can be stated as follows [100, 101]:

H₂S dissolution: \( H_2S(g) \leftrightarrow H_2S(aq) \) (5-1)

H₂S dissociation: \( H_2S(aq) \leftrightarrow H^+(aq) + HS^-(aq) \) (5-2)

HS⁻ dissociation: \( HS^-(aq) \leftrightarrow H^+(aq) + S^{2-}(aq) \) (5-3)

H₂S reduction: \( 2H_2S(aq) + 2e^- \rightarrow H_2(g) + 2HS^-(aq) \) (5-4)

FeS formation by precipitation: \( Fe^{2+}(aq) + S^{2-}(aq) \leftrightarrow FeS(s) \) (5-5)

FeS formation by solid state reaction: \( Fe(s) + H_2S(aq) \rightarrow FeS(s) + H_2(g) \) (5-6)

Here the H₂S is initially adsorbed onto the steel to quickly form a thin (<<1 µm), dense, and adhesive iron sulfide film. At lower temperatures (~135°C) the film is likely that of mackinawite [41, 100]. At elevated temperatures between 170°C to 200°C mackinawite will transition into alternate iron sulfide products including pyrrhotite [41]. Past failure analysis and this research have consistently identified pyrrhotite as being a main corrosion products present.

This initial scale formed acts as a diffusion barrier which quickly reduces corrosion rates, but does not completely stop it. Corrosion and scale formation continues
and as it precedes internal stresses and variations in the molar volume ratio of the film to the iron initiates and progresses film micro cracking. As this film breaks off, it continues to re-form. The outer scale is less dense, porous and adhesive than the base layer and continues the formation, growth, and cracking process.

Because of the rapid formation of a protective, passive, film the rate of corrosion is limited to how quickly corrosive species can diffuse through it. Meaning it is influenced mainly by a mass flux transfer control process. The mass flux is controlled by the inner and outer pyrrhotite film layers as well as the liquid boundary layer as follows [43, 72]:

\[
Flux_{H_2S} = k_{m(H_2S)}(c_{(H_2S)} - c_{o(H_2S)}) \quad (5-7)
\]

\[
Flux_{H_2S} = \frac{D_{(H_2S)}\epsilon\phi}{\delta_{OS}}(c_{o(H_2S)} - c_{i(H_2S)}) \quad (5-8)
\]

\[
Flux_{H_2S} = A_{H_2S} \ln\left(\frac{c_{i(H_2S)}}{c_{s(H_2S)}}\right) \quad (5-9)
\]

The process of continued film cracking and re-growth is a steady state one. For that reason the corrosion rate can be estimated by equating these equations to give the final expression [43, 72]:

\[
CR_{H_2S} = A_{H_2S} \ln\left(\frac{\frac{c_{b,H_2S} - Flux_{H_2S}(\frac{\delta_{OS}}{D_{(H_2S)}\epsilon\phi} + \frac{1}{k_{m(H_2S)}})\delta_{OS}}{c_{s,H_2S}}}{\frac{1}{k_{m(H_2S)}}}\right) \quad (5-10)
\]

Where:

\[
CR_{H_2S} : \text{Corrosion rate as mol/m}^2\text{s}
\]

\[
k_{m(H_2S)} : \text{Mass transfer coefficient of H}_2\text{S in liquid boundary layer in m/s}
\]
\( D_{(H,S)} \): Diffusion transfer coefficient of \( H_2S \) in water

\( \varepsilon \varphi \): Porosity and tortuosity of outer film

\( \delta_{OS} \): Outer film thickness, m

\( c_{b,H_2S}, c_{x,H_2S} \): \( H_2S \) concentrations in the bulk solution and locally at the steels surface respectively, mol/m\(^3\)

A difference between the corrosion mechanisms in an \( H_2S \) containing system which does not have \( CO_2 \) and the ones found in SAGD which do have \( CO_2 \) is that additional bulk fluid chemical reactions occur, these include [102]:

The dissociation of dissolved \( H_2S \):

\[
H_2S \rightarrow H^+ + HS^- \quad (5-11)
\]

With solubility product:

\[
K_{H_2S} = \frac{[H^+][HS^-]}{[H_2S]} \quad (5-12)
\]

The dissociation of \( HS^- \) ions:

\[
HS^- \rightarrow H^+ + S^{2-} \quad (5-13)
\]

With solubility product:

\[
K_{HS^-} = \frac{[H^+][S^{2-}]}{[HS^-]} \quad (5-14)
\]

Calculating the solubility products shows that \( H_2S \) gas is approximately 3 times more soluble than \( CO_2 \) gas. The acid created by the dissociation of \( H_2S \) is also about 3
times weaker than carbonic acid. Therefore the effects of each gas on the solutions pH are similar.

5.2 Factors which affect the H₂S corrosion mechanism

The research and literature review completed indicate that several factors can affect the corrosion rate of steels when dominated by aqueous H₂S corrosion. These include fluid composition and flow rate, temperature, H₂S activity, pH, and H₂S concentration.

5.2.1 The effect of flow rate and fluid composition

Literature and field studies have shown that flow rate, and therefore fluid composition, can play a role in corrosion rates. The annular space in a production wells is a low flow environment where fluid phase separation takes place whereas inside the production tubing where emulsified fluids continually flow through it. The continual flow fluid and partitioning of aqueous H₂S into the oil phase and scale formation all appear to limit and prevent corrosion from occurring [98]. To date no severe corrosion has been found on the internal surfaces of the production tubing. The same tubing lines have experienced corrosion on the external surfaces. Because of the fluid composition variation, different corrosion rates have been found at different well locations. Regardless of material, uniform corrosion rates were highest at the lower well elevations (closer to the ESP) and increased as the fluid transitions from an emulsion, to being water at the top of the well before separating into produced gases released form the liquids. The mechanism of aqueous H₂S corrosion does not appear to change as iron sulfide products have been found at all well locations. Research, past failure analysis, and XRD analysis
have shown the presence of pyrrhotite, troilite, and pyrite, at all well depths. Although pyrrhotite was not present on the top coupons in this research, past failure analysis and XRD results indicated that pyrrhotite and other corrosion products can be expected at all well elevations in the presence of steel.

5.2.2 *The effect of H$_2$S concentration*

H$_2$S concentration plays a significant role in corrosion rates due to the inability for a dense iron sulfide scales to form at high concentrations. Higher H$_2$S content leads to the formation of a loose and easily removed film [103]. Also, the concentration and relative partial pressure of H$_2$S compared to CO$_2$ will dictate the active corrosion mechanism. This research and sampling has shown that concentrations of H$_2$S are high enough to have aqueous H$_2$S corrosion be the dominating mechanism and that often a stable and protective iron sulfide scale is able to form. Past failure analysis where siderite and magnetite have been identified is likely due to the partial pressure ratio CO$_2$/H$_2$S to be in the transitional mechanistic zone, not a fully dominating H$_2$S zone. Further sampling should be done to confirm this.

5.2.3 *The effect of temperature*

Research has shown that at lower temperatures (less than ~60°C) that temperature can play an important role in increasing and decreasing H$_2$S corrosion rates due to improved hydrogen diffusion rates. But that above ~40-60°C the effects of temperature are less pronounced. Temperature also appears to impact the type of FeS scale formed, not whether or not a passive scale will form [41]. Some iron sulfide scales
have shown to display improved protective properties, but to no great extent. All are dense and adhere to steel [50, 81, 99].

Although the effects of aqueous H₂S corrosion does not appear to be affected greatly by temperature under SAGD conditions, should the ratio of CO₂/H₂S begin to favor the corrosion kinetics for CO₂ corrosion to dominate it is known that CO₂ corrosion is highly dependent on temperature. At a temperature greater than ~ 60°C a highly passive iron carbonate scale forms to reduce corrosion rates.

In this research temperatures remained well above 150°C for most of the testing period in the well at the liquid level top. Temperatures at the ESP remained above 200°C and at times up to 220°C. This research confirms that at temperatures between 150-220°C similar scales will form, likely that of pyrrhotite. As mentioned, past failure analysis has shown iron sulfide scales to be present at higher well elevations and not just silicate scales.

5.2.4 The effect of pH

Literature review has shown corrosion rates to be highly dependent on the solutions pH [102]. At lower pH values FeS is not as likely to precipitate to form a scale due the high solubility of iron causing continual corrosion to occur at high rates. At slightly higher pH values, between pH ~ 3-5, the iron sulfide films can form and become stable. pH and H₂S partial pressure are often used when selecting materials for sour service applications [44].

Produced fluid analysis from SAGD wells has indicated pH values in the 2-5.5 range. Autoclave testing on the 1018 CS, J-55, and TN-55TH materials show the likely
dependence of corrosion rates on pH based on the LPR trends observed. Testing at pH 4.36 showed an increasing corrosion rate trend before corrosion rates stabilized, compared to the decreasing corrosion rate trend at pH 5.5 levels for the 1018 carbon steel. This indicates a difference in how the iron sulfide scales form in varied conditions before eventually stabilizing at similar rates.

Field test results also confirm this effect based on the “low” rates of corrosion found on all coupons installed and pitting corrosion occurring at higher coupon locations as well as the increase in corrosion rates at the top clamp location. At lower well elevations where an emulsion and oil are present the partitioning of the gases into an oil phase will lead to increased pH values and aid in the formation of a passive scale, as XRD results confirm. Moving up the well and with pH decreasing, as gases are released from the oil phase, less corrosion product was noted. An increase in corrosion rates may be expected due to the reduction in pH, but it is likely that the more dynamic conditions at the lower joint 2 clamp coupons, closer to the ESP, delay the scales rate of formation. At the higher joint 25 clamp the increased acidic gas concentration decreases the pH to a point where the reduced pH and high rate of iron solubility prevent a passive FeS scale from forming.

5.3 Implications of mechanistic research

The implication of this mechanistic research and finding aqueous H₂S corrosion to be the dominating mechanism are as follows:

- Promoting the initial solid state reaction and formation of the initial iron sulfide scale can help to protect the base metal during longer term operation. Ensuring
that this scale is dense and non-porous will limit the inwards flux of fluids or other impurities which can promote further corrosion, although still reduced.

- When both CO₂ and H₂S are present, ensuring that the partial pressure ratio of CO₂/H₂S is higher enough for only iron carbonate scales to form, or low enough that only iron sulfide scales form is important. Avoiding the transitional area where both iron carbonate and iron sulfide scales are present is important as impurities can prevent either scale from proper adhesion and protection.

- Material selection considerations should also be made as some materials are more susceptible to aqueous H₂S corrosion than others. Although all materials tested were affected by corrosion, L-80 proved to be susceptible to pitting corrosion while others did not. This is concerning due to the difficulty in detecting pitting corrosion and how it often leads to sudden failures. Reviewing and selecting materials being used at the liquid level in the well should be considered. Evaluate and select materials which are less susceptible to pitting corrosion caused by aqueous H₂S.

- Also, given the dependence of aqueous H₂S corrosion on pH, inhibition or scavenging should be considered to ensure the systems pH remains in a range of 3-5. Evaluating other methods to control the systems pH should also be considered.

- Literature review and failure analysis have shown that aqueous acidic gas corrosion can be prevented/reduced in the conditions present inside the production tubing lines where a continually flowing emulsion is present. This being due to the formation of a protective scale, oil wetting surfaces, as well as the partitioning
of the gases from the aqueous phase into the oil phase. Although difficult in operation, considerations should be made to either promote the formation and flow of an emulsion from the annular space, or to re-designing well as to not have stagnant fluids present in separated phases.
Chapter Six: Conclusions and recommendations

6.1 Conclusions

The laboratory evidence and field testing results have led to the following conclusions:

- Aqueous H$_2$S corrosion is the dominating corrosion mechanism
- “Low” corrosion rates can be expected due to the formation of pyrrhotite, possibly other FeS scales, which prevent increasing corrosion rates
- Stellite-6 HF material should replace the Deloro-40 HF overlay
- The standard tubular materials J-55, K-55, and TN-55TH coupons consistently showed the highest corrosion rates in both the lab and field testing, up to 0.46 mm/y and 0.0282 mm/y respectively
- Galvanic testing in both the field and lab showed only a slight increase in corrosion rates
- Uniform corrosion rates generally decreased from the bottom coupons in the well to the top on all materials tested
  - At and above the fluid level general and pitting corrosion rates can be expected to increase and is likely due to the combination of increased acidic gas concentrations, water and vapor condensates, and the presence of galvanic/creviced conditions
- Corrosion product compounds and their relative concentrations changed clearly with well depth
6.2 Recommendations for industry and future work

Based on the findings of this research the following recommendations are made.

- Design further laboratory tests to include oil for 3-phase fluid testing, oil + water + gas, with silicate scales present to coat the sample surfaces.
- Designing test loops and autoclave setups for testing at higher temperatures (~200°C), and testing for longer durations of 14 to 21 days.
- Complete further autoclave testing on a third material, tungsten carbide HF overlay. This material has been proposed for use by different wellhead and valve suppliers but has not yet been tried in SAGD like conditions.
- Complete CPP and PP tests on L-80 materials to better understand their pitting characteristics and tendencies in the SAGD environments.
- Further research on how aqueous gas concentrations and pH change when partitioning into the oil phase occurs and evaluate changes at different well elevations.
- Comparing solubility of FeS and silicate scales to better understand which scale will preferentially form in SAGD conditions and what can control it.
- Further review and testing of the corrosion inhibiting effects of silicate scales compared to iron sulfide scales.
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Appendix

Sample material mechanical properties

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<th>Material</th>
<th>Yield Strength (MPa)</th>
<th>Tensile Strength (MPa-min)</th>
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<td>517</td>
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<tr>
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<td>379 – 552</td>
<td>655</td>
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TN-55TH chemistry

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<th>S</th>
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API L-80 Type 1 chemistry

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<tr>
<td>Min</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Max</td>
<td>0.43</td>
<td>1.90</td>
<td>0.030</td>
<td>0.030</td>
<td>0.25</td>
<td>0.35</td>
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</tbody>
</table>

Stellite-6 HF chemical composition

<table>
<thead>
<tr>
<th>Co</th>
<th>Cr</th>
<th>W</th>
<th>C</th>
<th>Other</th>
</tr>
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<tbody>
<tr>
<td>Min</td>
<td>Base</td>
<td>27</td>
<td>4</td>
<td>0.9</td>
</tr>
<tr>
<td>Max</td>
<td></td>
<td>32</td>
<td>6</td>
<td>1.4</td>
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</tbody>
</table>

Stellite-6 HF mechanical properties

<table>
<thead>
<tr>
<th></th>
<th>Hardness</th>
<th>Density</th>
<th>Melting Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imperial</td>
<td>36-45 HRC</td>
<td>0.305 lb/in³</td>
<td>2340-2570°F</td>
</tr>
<tr>
<td>Metric</td>
<td>380-490 HV</td>
<td>8.44 g/cm³</td>
<td>1285-1410°C</td>
</tr>
</tbody>
</table>