Department of Chemistry

Top-of-the-Line Corrosion Control by Continuous Chemical Treatment

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Declaration

To the best of my knowledge and belief this thesis contains no material previously published by any other person except where due acknowledgment has been made.

This thesis contains no material which has been accepted for the award of any other degree or diploma in any university.

Signature: ..................................................

Date: ............................................
Summary

Top-of-the-Line (TOL) corrosion is a severe problem associated with the transportation of wet natural gas. It has the potential to cost the oil and gas industry millions of dollars every year through lost production and the necessity to replace effected pipelines.

Research was carried out in order to better understand and estimate the TOL corrosion risk and its control. The first part of the research focused on the development of a domain diagram. The TOL corrosion domain diagram was designed to gain a better understanding of how carbon dioxide partial pressure, acetic acid concentration, and temperature influence TOL corrosion at a constant condensation rate. It summarises the effects of some important parameters on TOL corrosion. TOL corrosion tests were performed with three different carbon dioxide partial pressures of 5 bar (72.5 psi), 10 bar (145 psi), and 20 bar (290 psi), and different acetic acid concentrations of 0 ppm, 500 ppm, and 1000 ppm. The sample temperature was also varied at 30 °C and 80 °C. The gas temperatures were adjusted to 91 °C and 115 °C to maintain a constant condensation rate in both temperature regimes of 0.40 g/m2/s.

The next part of the research was focused on volatile corrosion inhibitors (VCIs). 16 potential VCI compounds (Aminoethylpiperazine, Aminomorpholine, Aniline, Benzylamine, Cyclohexylamine, Dicyclohexylamine, Diethylamine, Dimethylethylamine, Methylidethanolamine, Methylmorpholine, Methylpiperazine, Morpholine, Octylamine, Picoline, Pyridazine, and Pyridine) were chosen and tested in a variety of TOL corrosion test rigs. It was possible to identify the properties and characteristics of a potential VCI compound in the presence of acetic acid.

The same VCIs were also tested in a rotating cylinder electrode test set-up using electrochemistry, mainly linear polarisation resistance (LPR) and electrochemical impedance spectroscopy (EIS), to test their Bottom-of-the-Line (BOL) inhibition ability as well as their mechanism of inhibition. All of the compounds showed BOL inhibition capability. One compound stood out
as a good film former on carbon steel despite an uncommon molecular structure for a film forming corrosion inhibitor.

Design modifications for the different TOL corrosion set-ups used in the experiments were devised using the data and experience gained in this research. A modified TOL corrosion apparatus was constructed, capable of high pressure, high temperature TOL corrosion testing, with the potential to standardize TOL corrosion and VCI testing was suggested.
Publications, Presentations, Posters


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List of Abbreviations

A  Area
AAS  Atomic absorption spectrometry
A-HCT  Altered Horizontal Cooled Tube
(aq)  Aqueous
B  Beta A value (0.12)
bc  Beta C value (0.12)
BOL  Bottom-of-the-Line
CFP  Cooled Finger Probe
CR  Corrosion rate
DEA  Diethylamine
DMEA  Dimethylethylamine
EDX  Energy dispersive X-ray
EW  Equivalent weight
(g)  Gaseous
IC  Iron concentration
icor  Corrosion current density
IE  Inhibition efficiency
IFM  Infinite focus microscope
I-HCT  Improved Horizontal Cooled Tube
HCT  Horizontal Cooled Tube
Icor  Corrosion current density
K₁  Constant (3.27 *10⁻³ mm g/µA cm yr)
LPR  Linear polarisation resistance
m  Mass
MDEA  Methyldiethanolamine
N  Number of Specimens
Np  Number of pitted specimens
OCP  Open circuit potential
P  Pitting probability
pCO₂  Partial pressure of CO₂
PD  Potentiodynamic
PIG  Pipeline inspection gauge
Ra  Arithmetical mean roughness
RCT  Charge transfer resistance
RP  Polarisation resistance
RS  Solution resistance
RCE  Rotating cylinder electrode
Rz  Ten-point mean roughness
SEM  Scanning electron microscope
t  Time
TA  Temperature Autoclave
TS  Temperature Sample
TOL  Top-of-the-Line
VCI  Volatile corrosion inhibitor
WCO  World Corrosion Organisation
WL  Weight loss

Units

A  Ampere
cm³  Cubic centimetre
°C  Degree Celsius
g  Gram
h  Hours
Hz  Herz (Frequency)
m²  Square metre
mL  Millilitre
mm  Millimetre
μm  Micrometre
mV  Millivolts
ppm  Parts per million
rpm  Revolutions per minute
s  Second
V  Volt
y  Year
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1. Introduction

Corrosion is an often overlooked, but omnipresent phenomenon. It was estimated by the World Corrosion Organisation (WCO) in 2010 that 3% of the world’s GDP, $ 2.2 trillion was the annual direct cost of corrosion worldwide (Hays 2010). This huge number makes the annual corrosion costs in the US oil and gas industry in exploration, production, refining, and pipelines of about $ 12 billion look comparatively small. (Koch et al. 2002). Corrosion in the oil and gas industry is still taken very seriously since millions of dollars can be saved using the right treatments or making the right choices. An even bigger factor is that corrosion is also associated with numerous fatalities, injuries and environmental tragedies all over the world (NTSB 1969-2012) (Bills and Agostini 2009).

To keep incidents to the bare minimum, research is undertaken in many directions; one of which is CO\textsubscript{2} corrosion and its mitigation. CO\textsubscript{2} corrosion, also known as “sweet corrosion” is the prevalent form of internal corrosion for carbon steel pipelines. Carbon steel is the most common pipeline material, despite the fact that it is prone to CO\textsubscript{2} corrosion. Nevertheless, with the right corrosion prevention programs, like inhibition or pH stabilisation, along with corrosion monitoring programs, it is a more cost effective solution than higher alloyed steels.

1.1. Top-of-the-Line Corrosion

“Vapour-phase corrosion due to the condensation of water vapour in the presence of acid gases and in the absence of hydrocarbon condensate has been identified as the corrosion mechanism in the Crossfield gas gathering system”. This was concluded in the investigation of a pipe burst in the Crossfield gas gathering system near Calgary, Alberta, Canada in 1985 (Bich and Szklarz 1988). This was most likely one of the first descriptions Top-of-the-Line corrosion induced failure in a CO\textsubscript{2} dominated environment.
TOL corrosion can occur when hot wet natural gas is transported in a poorly insulated pipeline with a high heat transfer coefficient to the surrounding area; for example subsea pipelines. The gas is cooled down rapidly and the water vapour condenses at the pipe walls. The condensed water is very aggressive and often referred to as “hungry water” due to the lack of buffering agents and dissolved iron. The CO$_2$ dissolves in the water and is converted into carbonic acid, lowering the pH, causing the water to become more aggressive. It is reported that a condensation rate of 0.25 g/m$^2$/s is sufficient to induce TOL corrosion in these conditions (Gunaltun and Larrey 2000). The necessary condensation rate is lowered by a magnitude as soon as a sufficient amount of organic acids are present.

Organic acids, predominantly acetic acid, are a very common by-product in the natural gas production. Concentrations of about a few hundred to a few thousand ppm are not uncommon, and they are reported to increase the corrosion rate and affect the protective scale formation even in relative low concentrations.

**1.2. Top-of-the-Line Corrosion Control**

Many efforts to mitigate TOL corrosion have been suggested and tested. Some of these efforts are more successful and practical than others. As long as a field is in the building stage, the first kilometres can be built of corrosion resistant alloys without heat insulation. Water vapour then can be condensed out so it is no longer present further down the line. Another solution is burying the pipeline deeper into the ocean floor, increasing the heat insulation. These are engineering efforts that can be implemented in newly developed fields but not in already developed and working pipelines. In these cases, inhibition plays a major role in TOL corrosion mitigation.

Major hurdles are the distribution of the inhibitor to the TOL. One of the more common techniques is inhibitor batch treatment. A sticky long chain organic inhibitor is put in between two Pipeline Inspection Gauges (PIGs) and it is run through the pipelines. Unfortunately, the production of the pipe
or even large parts of a field needs to be shut down for this procedure, resulting in a big loss for the operating companies.

Another method would be a rather novel approach using a spray PIG. The PIG uses the Venturi Effect and sprays solution from the Bottom-of-the-Line (BOL) with a high inhibitor concentration to the TOL. The field doesn’t need to be shut down completely, but the pipeline still needs a launching and receiving point for PIGs as well as the risk of a PIG being stuck in a pipeline.

Another promising approach is Volatile Corrosion Inhibitors (VCI). VCIs are commonly known to be small chain amines due to the required volatility. These VCIs are continuously injected into the BOL, utilizing the same injectors installed for conventional BOL corrosion inhibitors. It is not well understood how VCIs work, but commonly, VCIs are thought to evaporate and co-condense at the TOL with all the other fluids and inhibit the CO₂ corrosion directly where it occurs. Nevertheless, most of the commercially available VCIs are not yet effective enough at inhibiting the TOL corrosion for them to be used as the sole countermeasure. So far, they are mainly used in combination with PIGs to extend the cycles in which they have to be used, which is already an enormous benefit.

This research investigates several generic potential VCI compounds; some of them are in use in commercial formulations. As mentioned above, it is not yet clearly known how they reach the TOL (foaming, volatility) and by which mechanism they inhibit (neutralizing, film forming) TOL corrosion.

Different laboratory-based equipment was used to gain a better understanding of TOL corrosion under different conditions and investigate the performance and inhibition mechanism of the different VCIs.
1.3. Research Objectives

- Developing a TOL corrosion domain diagram with variable carbon dioxide partial pressure, acetic acid concentration, and temperature under constant condensation rate
- Gain a better understanding of TOL corrosion testing under high carbon dioxide partial pressures
- Find a working VCI compound able to be used in a formulated commercial VCI
- Examine the mechanism of TOL corrosion inhibition through various VCI compounds
- Identify properties of a potential generic VCI compound
- Improve the existing TOL corrosion test methods
- Develop a TOL corrosion test method with the potential to be accepted as standard test method for VCI evaluation
2. Top-of-the-Line Domain Diagram

2.1. Introduction

2.1.1. CO₂ Corrosion

CO₂ corrosion in the oil and gas industry was discovered for the first time in the 1940s and has been a problem in the industry ever since. Several research teams all over the world keep investigating the phenomenon to gain a better understanding of the mechanisms of CO₂ corrosion (Crolet and Bonis 1983; Sun and Nesic 2006; Tan et al. 2010). Nevertheless, the exact mechanism is still widely debated. The following equations display the trail of CO₂ becoming a corrosive environment for carbon steel in combination with water.

CO₂ in the gas phase and in the liquid phase are in equilibrium.

\[ CO_2 (g) \leftrightarrow CO_2 (aq) \]  
Equation 1

CO₂ then dissolves in the water phase forming carbonic acid.

\[ CO_2 (aq) + H_2O \leftrightarrow H_2CO_3 (aq) \]  
Equation 2

H₂CO₃(aq) dissociates in two steps releasing two H⁺ as seen in the next equations.

\[ H_2CO_3 (aq) \leftrightarrow H^+_2 (aq) + HCO_3^- (aq) \]  
Equation 3

\[ HCO_3^- (aq) \leftrightarrow H^+_2 (aq) + CO_3^{2-} (aq) \]  
Equation 4

The amount of carbonic acid and all subsequent carbonates is highly dependent on the CO₂ partial pressure (Equation 1- Equation 4) which in turn has a direct influence on the corrosivity of the liquid phase.
The exact mechanism how CO₂ corrodes carbon steel is still under debate and is very much dependent on other factors like water chemistry, temperature, pH, partial pressure and others. The overall equation for CO₂ corrosion of carbon steel is also often shown as

\[ Fe + CO_2 + H_2O \rightarrow FeCO_3 + H_2 \]  \hspace{1cm} \text{Equation 5}

Dugstad concludes in his review paper that CO₂ is assumed to contribute to the cathodic reaction rate in two different ways. H₂CO₃ can be directly reduced and the dissociation of H₂CO₃ can serve as a source of H⁺ ions (Dugstad 2006). The equation for direct reduction would look like the following.

\[ Fe + 2H_2CO_3 \rightarrow Fe^{2+} + 2H_2CO_3^- + H_2 \]  \hspace{1cm} \text{Equation 6}

The free iron (II) ion then can react with the carbonate ion to form iron carbonate.

\[ Fe^{2+} + CO_3^{2-} \rightarrow FeCO_3 \]  \hspace{1cm} \text{Equation 7}

The iron carbonate then accumulates in the water phase. With ongoing corrosion, the concentration of iron carbonate in solution will increase until it hits the saturation limit and an iron carbonate film starts forming on the carbon steel surface. The precipitation rate is a function of supersaturation and temperature (Sun and Nesic 2006). It was shown that FeCO₃ is more soluble at lower temperatures and therefore higher FeCO₃ concentrations are necessary to reach supersaturation at lower temperatures (Johnson 1991; Dugstad 1998). Once an iron carbonate film has formed it stays protective even at a much lower supersaturation (Dugstad 1998). The iron carbonate then acts as a protective corrosion product layer retarding further corrosion. Once again, the protectiveness of the corrosion layer depends, strongly on environmental parameters like surface condition, temperature, pH and others (Nesic et al. 2001; Nesic et al. 2002).
2.1.2. Effect of Acetic Acid on CO₂ Corrosion

Acetic acid is a weak organic acid that provides a single hydrogen ion upon dissociation. The connection between organic acids and CO₂ corrosion was reported for the first time as early as 1944 by Menaul in the Oil and Gas Journal (Menaul 1944). Later it has been shown that mainly formic, acetic, propionic and, butyric acid are present with acetic acid being by far the most prevalent at up to 90% (Dougherty 2004). Studies have shown that there is very little difference in the corrosion behaviour of the diverse low molecular weight organic acids (Fajardo et al. 2007). Therefore, the focus here will be on the most abundant, acetic acid, and its contribution to CO₂ corrosion.

Over the years many different even contradictory theories were put forward to explain the contribution of acetic acid to CO₂ corrosion. In 1999 it was published that acetate (CH₃COO⁻), which is dissociated acetic acid, increases the corrosion rate (Hedges and McVeigh 1999). At the same conference it was concluded that it is in fact not the acetate but just the concentration of free acetic acid that increases the corrosion rate (Crolet et al. 1999). Many experiments and publications later, it can be concluded that the latter theory is the right one (George and Nesic 2007).

Acetic acid is reported to increase the cathodic corrosion reaction of bare steel in two ways. It is believed that acetic acid supplies H⁺ ions to the corrosion site (by dissociation) very similar to CO₂ corrosion. Additionally, it is directly reduced on the steel surface (Garsany et al. 2002; Crolet and Bonis 2005) in form of Equation 8.

$$CH₃COOH + 2e^- → H₂ + CH₃COO^-$$  \hspace{1cm} \text{Equation 8}$$

In real life applications, iron carbonate scale is often present on the metal. The effect of acetic acid on the growth, protectiveness and thickness of an iron carbonate layer is also the subject of contradicting publications. In 1999, it was published that acetic acid reduced the thickness and protectiveness of the corrosion layer (Hedges and McVeigh 1999). Some more recent publications state that there is no significant effect on the iron carbonate scale formation and its protectiveness (Nafday and Nesic 2005).
Another approach to explain the contribution of acetic acid in relation to corrosion product layers was made by Fajardo et al. by arguing that acetic acid initiates a “scale undermining effect”. Thicker iron carbonate scales in presence of organic acids were found. It was concluded that plenty of iron carbonate was formed but due to the higher corrosion rates initiated by acetic acid, it took longer for the film to take a foothold retarding further corrosion. (Fajardo et al. 2007).

Acetic acid also seems to induce localized corrosion in different environments. It was reported that acetic acid induces a local potential difference promoting the initiation of localized corrosion (Amri et al. 2009). A higher concentration was also connected to an increased pit penetration rate (Gulbrandsen and Bilkova 2006). Acetic acid has also be found to exhibit corrosion inhibitive effects on the anodic part (Gulbrandsen and Bilkova 2006), but it still increases the overall corrosion rate.

Crolet et al. showed also that it is possible, at least in low CO\textsubscript{2} partial pressures, that genuine acetic acid corrosion can occur and replace iron carbonate with iron acetate. Iron acetate has a much greater solubility decreasing the protectiveness of the scale drastically (Crolet et al. 1999).

It has also been reported that acetic acid strongly contributes to, or even is necessary for Top-of-the-Line corrosion (Dougherty 2004).

2.1.3. Top-of-the-Line (TOL) Corrosion

Many TOL corrosion problems have been reported, especially in the Asia-Pacific region (Gunaltun et al. 1999; Gunaltun 2006; Gunaltun et al. 2010). TOL corrosion has been cited as one of the most complex forms of corrosion in the oil and gas industry (Bailey 2010). It is a specific form of CO\textsubscript{2} corrosion limited to the production and transportation of wet gas in carbon steel pipelines with a stratified flow regime (Figure 1).

TOL corrosion was reported for the first time in 1981 by Paillassa et al. in a H\textsubscript{2}S containing gas field in France (Paillassa et al. 1981). It didn’t take too long before the first case was described in a CO\textsubscript{2} corrosion dominated gas
field (Bich and Szklarz 1988). Studies have been conducted and awareness has been raised ever since, but TOL corrosion is still not yet fully understood.

A key factor to consider in trying understanding TOL corrosion is the pipe and gas temperature. The most incidences of TOL corrosion are in the first kilometres of carbon steel pipelines where the gas is still comparatively hot and moist and the pipe is cold. These conditions are found in poorly insulated subsea pipelines, river crossings or upheaval buckling sections. This combination results in a high rate of condensation. Reportedly the condensation rate needs to be above 0.15-0.25 g/m²/s to initiate TOL corrosion in a low organic acid environment (Gunaltn and Belghazi 2001). It also has been reported that the condensation rate can be even as low as 0.025g/m²/s with very high (2500 ppm) concentrations of organic acids (Gunaltn et al. 2006) making TOL corrosion possible even in rather well insulated pipelines.

The condensing water is often referred to as “hungry water” because no buffer and low pH values are present. Co-condensing organic acids and dissolved CO₂ decrease the pH of the condensate even further making it even more aggressive. A race between iron carbonate saturation (and therefore
iron carbonate precipitation) and dilution by further condensation commences when water first condenses. The carbon steel pipeline corrodes, saturating the condensed water with iron carbonate. The constant supply of condensing liquid keeps diluting the solution. Eventually a droplet will form and either rinse down the side wall or drop off the TOL. In the case of a low condensation rate, the pH will increase and a protective corrosion product layer can form, mitigating further corrosion (Andersen et al. 2007). At high condensation rates this is not the case and corrosion keeps going, rather unrestricted.

Temperature also plays a major role in the formation of corrosion products. As for general CO₂ corrosion, the protectiveness of a FeCO₃ layer depends on the temperature of the substrate on which it is grown. As described in section 2.1.1, a protective FeCO₃ layer is more likely to form at elevated temperatures due to the supersaturation levels required.

The concentration of acetic acid at the TOL is driven by the liquid/ vapour equilibrium of free acetic acid at the Bottom-of-the-Line (BOL) and total acetic acid at the TOL. The distribution of free acetic acid at the TOL depends then mainly on the pH of the condensed liquid (Hinkson et al. 2010). The percentage of free acetic acid at the TOL or the BOL can be calculated by the following formula:

$$\% \text{ CH}_3\text{COOH} = \frac{100}{\left(1 + 10^{\text{pH}-\text{pKa}}\right)}$$  \hspace{1cm} \text{Equation 9}

with $\% \text{ CH}_3\text{COOH}$ is the percentage of undissociated acetic acid, pH of the solution, and pKa of the acid, in this case acetic acid. It is possible to calculate the percentage of undissociated acetic acid for the necessary pH range and plot it in a graph (Figure 2). It can be seen that the dissociation of acetic acid is highly dependent on the pH of the solution.

Besides the temperature, condensation rate and acetic acid concentration the gas velocity also plays a role in TOL corrosion. At low velocities, stagnant droplets are formed whereas at higher gas velocities the droplets are sliding along the TOL and eventually slide to the BOL. For stagnant droplets it is
easier to form a protective \( \text{FeCO}_3 \) layer whereas sliding droplets are generally not in contact long enough with the steel and rather form a non-protective layer (Singer et al. 2009). In many of the case studies localized corrosion was reported at the TOL. There seems to be a strong correlation between localized attack and acetic acid and temperature (Singer et al. 2009).

Many different laboratory test setups have been developed to investigate TOL corrosion. Large flow loops are used to replicate field conditions as closely as possible under laboratory conditions (Nyborg et al. 2009; Singer et al. 2010) but they are a large, labour intensive piece of equipment and expensive to maintain and run. Then there are many test setups replicating TOL corrosion in bench top experiments for easier handling and greater flexibility (Pots and Hendriksen 2000; John et al. 2009; Chen et al. 2011). Many TOL corrosion experiments are even conducted with a specimen submerged in a glass cell, similar to BOL corrosion tests (Amri et al. 2011). It is difficult to implement all factors into a single test rig and therefore priorities and research objectives have to be defined.

![Figure 2: Acetic acid dissociation curve](image.png)
2.2. Experimental

2.2.1. Pressure Reactor and Corrosion Sample

The test reactor is a slightly modified version of a commercially available 316 stainless steel 2 L PARR high pressure reactor. It is built to withstand up to 130 bar pressure at 350° C. It is equipped with an internal purge tube and a gas outlet, both sealed with a high pressure valve. It is also fitted with a rupture disc that releases the internal pressure in case it exceeds its maximum limit. Two independent pressure gauges are fitted as well; an analogue gauge and a digital gauge which is connected to a PARR control unit. The control unit is connected to a thermocouple to measure temperature, a pressure gage to read pressure inside the reactor, and a furnace to control the temperature. An upper and lower safety limit for both the temperature and the pressure can be set, where the control unit cuts off the power and the reactor cools down to prevent the pressure or temperature from rising further.

The reactor in its original state as supplied by the manufacturer is equipped with an internal cooling coil through which water is pumped. This cooling coil was removed and the TOL corrosion set up was installed instead. A 13 cm long ¼ inch carbon steel tube (ASTM-A179 2005), bent into a rectangular U-shape using a tube bender, was attached as corrosion sample using stainless steel Swagelok fittings, Swagelok elbows, and Teflon ferrule tube fittings.

After the tube was bent into shape, a heat treatment was conducted. The sample was placed inside a furnace for 90 minutes at 650 °C to remove residual stresses in the material (ASM 1998) induced by the bending process.

An inhibited cooling fluid (5% sodium nitrite in water) was pumped through the test sample at a controlled temperature to initiate a temperature difference but prevent internal corrosion (Hayyan et al. 2012). Due to the lower temperature of the carbon steel pipe, condensation occurred on the
outside of the pipe, triggering condensed water corrosion as an analogue of TOL corrosion.

1 cm of each side of the carbon steel tube was inside the fittings; therefore, 11 cm of the tube can be considered as the sample length, producing a total surface area of 22.14 cm². The chemical composition of the sample (as provided by the manufacturer) is listed in Table 1 below.

<table>
<thead>
<tr>
<th>[%]</th>
<th>C</th>
<th>Mn</th>
<th>P</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test samples</td>
<td>0.079</td>
<td>0.429</td>
<td>0.014</td>
<td>0.007</td>
</tr>
<tr>
<td>ASTM A179</td>
<td>0.06-0.18</td>
<td>0.27-0.63</td>
<td>&lt;0.035</td>
<td>&lt;0.035</td>
</tr>
</tbody>
</table>

Figure 3: Schematic and image of the Horizontal Cooled Tube test set-up
2.2.2. Preparation of a TOL Corrosion Test

The test solution (650 mL of milliQ water) was pre-purged with high purity CO₂ (99.99% pure) for at least two hours inside the body of the pressure reactor. In the meantime, the carbon steel tube (already bend and heat treated) was internally and externally sandblasted in a Hafco Sandblasting cabinet to ensure a smooth and clean surface. Throughout the duration of the research, the same sandblasting garnet product was used; old and abraded garnet was constantly filtered out of the sandblaster cabinet and the garnet was periodically topped up with fresh product to ensure a reproducible surface finish. After the sample was sandblasted it was put upside down into a beaker of acetone and treated for 5-10 minutes in an ultrasonic bath to thoroughly clean the specimen inside and out. The cleaned sample was then rinsed with milliQ water and acetone and subsequently dried in an oven for 20 minutes at 70 °C. After the 20 minutes it was left in a vacuum desiccator to cool down to room temperature to be weighed.

After the sample was weighed, it was attached to the head of the reactor using the Teflon and stainless steel tube connections. The appropriate amount of acetic acid was added to the solution. The head of the pressure reactor was then attached to the body and tightened using the split ring and drop band according to the specifications provided by PARR. To remove any residual oxygen in the system, the closed up pressure reactor was allowed to purge for another 10 minutes with CO₂ through the purge tube.

After the 10 minutes of pre-purging, the reactor was pressurised to 20 bar and subsequently depressurised using the same high purity CO₂. This pressurizing – depressurizing cycle was repeated four times to reduce the oxygen to a minimum. During the fifth pressurisation the system was allowed to equilibrate for 20 minutes at 20 bar.

After the 20 minutes, the reactor was sealed and placed into a controlled heating sleeve and heated to the appropriate temperature of the test (91°C or 115 °C). After the temperature was reached, cooling of the tube was started and the test was considered to be initialized.
The procedure for low pCO$_2$ testing was the same as above up to the pressurizing and depressurizing cycles. After the fourth cycle to 20 bar, the pressure was reduced to 5 or 10 bar (depending on the test) and held for 20 minutes to equilibrate. Afterwards, the pressure was topped up to 20 bar with nitrogen and again held for 20 minutes.

The residual oxygen was measured using an Orbisphere for some tests to verify whether or not the method described above is achieves the intent of an oxygen free environment. The oxygen concentration was verified to be below 4 ppb in all tested cases.

### 2.2.3. Corrosion Rate Determination

The corrosion rate was determined in two ways, by weight loss and by analysing the iron concentration in the bulk solution. Clarke’s solution was used to determine the weight loss of the samples. The solution was prepared in batches and used for several specimens.

#### 2.2.3.1. Clarke’s Solution

Every batch of Clarke’s solution was prepares using the chemicals as stated in the standard ASTM-G1. The sequence of preparation was as the following:

- 50 g of stannous oxide and 20 g of antimony trioxide was weight on an analytical balance and dissolved in the HCl
- The volumetric flask was filled up with HCl to exactly 1 L
- A stirrer bar was placed into the volumetric flask which was subsequently covered and stirring was conducted for 24 hours
- After 24 hours, the solution was filtered using a vacuum filter unit and filter paper sheets
- The last step might be repeated in case the solution was not clear
• The filtered Clarke’s solution is ready to use and can be stored in a sealable glass bottle

2.2.3.2. Corrosion Rate by Weight Loss (WL)

The mass \( m_1 \) of the sample was taken before the test using an analytical balance. Every time before the weight of a sample was taken a standard weight of exactly 20.0000 g was weighed to ensure the analytical balance was properly calibrated, not out of zero, and working well.

After the test, the sample was carefully removed from the test setup and cleaned using DI water and acetone. Afterwards the sample was heated in an oven for 20 minutes at 70 °C to thoroughly dry and was then stored in a vacuum desiccator to cool down to room temperature. After the sample cooled down to ambient temperature the mass \( m_2 \) of the sample was taken with an analytical balance.

![Figure 4: Mass loss vs. cleaning cycle diagram (ASTM G1)](image)
The sample still has the full layer of corrosion product that needs to be removed. This is performed using a procedure based on ASTM-G1. The sample was dipped into Clarke’s solution for 30 seconds at room temperature. The sample was cleaned, rinsed with DI water and acetone, heated in the oven for 20 min at 70 °C, left to cool down in the desiccator, and weighed on the analytical balance. The procedure was repeated several times to produce a mass loss vs. cleaning cycles diagram (Figure 4). This diagram will show 2 straight lines AB and BC. The latter will correspond to corrosion of the metal after removal of corrosion products. The mass of corrosion product will correspond approximately to point B (ASTM-G1 2003).

The final mass of the sample will be:

\[ m_3 = m_2 - m_{\text{point B}} \quad \text{Equation 10} \]

Therefore, the total mass loss due to corrosion can be calculated by the following equation:

\[ WL = m_1 - m_3 \quad \text{Equation 11} \]

Where WL is the total weight loss, \( m_1 \) the mass of the sample before- and \( m_3 \) the mass of the sample after the corrosion test.

Equation 12 was used to calculate the corrosion rates (CR) by weight loss in mm/year where \( K \) is a constant (8.76*10^4), WL is the mass loss in g, \( A \) is the sample area in cm², \( t \) is the exposure time in hours and \( \rho \) is 7.85 g/cm³ (density) (ASTM-G1 2003). The CR by weight loss is an average corrosion rate over the entire test period, which in this case is 7 days.

\[ CR = \frac{(K \times WL)}{(A \times t \times \rho)} \quad \text{Equation 12} \]

### 2.2.3.3. Corrosion Rate by Iron Concentration (AAS)

Corrosion takes place on the external surface of the cooled tube. A small amount of the corrosion product forms an iron carbonate corrosion product
layer, but most of the iron ends up in a droplet of condensed liquid forming on the tube, which eventually detaches and falls into the bulk solution.

A daily ritual of sample-taking was conducted. First, approximately 10 mL of bulk solution was removed to rinse the residual solution off the sampling tube and to determine the pH of the solution. Then, another 10 mL was taken and used to measure the iron concentration by means of atomic absorption spectrometry (AAS). The samples were preserved in 1.85 mL of concentrated 32% HCl. The overall HCl concentration in the solution was then 5 %. The samples were stored until there was a small batch (approximately 40 – 50). They were then diluted 1:50 (sometimes 1:100) using a 5 % HCl in milliQ water solution to decrease the iron concentration to less than 10 ppm.

AAS is a comparative technique, standard solutions were necessary to produce a standard curve comparing absorption to a concentration. 5 standards were produced by diluting a 100 ppm iron solution to 2, 4, 6, 8, and 10 ppm.

The standard solutions are measured before the actual samples. A calibration curve is plotted as shown in Figure 5 where a known iron concentration is related to the absorbance (the absorbance can change and depends on different factors like the adjustment of the flame or quality of the lamp measuring the absorbance; therefore it has to be prepared every time the machine is used). It is now possible to correlate the measured absorbance of a sample and correlate it by means of the calibration curve to an iron concentration.
The same AAS machine (Spectra AA), software (Spectra AA), and standard solutions were used throughout the testing for the domain diagram. A new calibration curve was measured before the machine was used.

After the iron concentration in the bulk solution was determined, the mass loss was calculated and then Equation 12 was used to calculate the corrosion rate.

2.2.4. Condensation Rate Determination

The condensation rate was determined experimentally using a beaker to collect the condensed liquid of the test sample. The beaker was covered with a Teflon lid, allowing the condensed liquid to go into the beaker but minimizing evaporation off the beaker during the test. The amount of liquid was measured and the condensation rate was calculated using Equation 13.

\[
\text{Condensation Rate} = \frac{m}{A \times t} \times 1000
\]

Equation 13

where \( m \) is the mass of condensed liquid in kilogram, \( A \) is the area of condensation in square meters and \( t \) the test duration in seconds.
Despite all efforts to minimize evaporation off the beaker, evaporation still occurred. Therefore, the same Teflon covered beaker was filled with a known amount of liquid and put into the autoclave at different temperatures to measure the amount of evaporation. An evaporation rate in grams per hour was calculated and later added on to the calculated condensation rate. Both the condensation and evaporation tests were conducted for about 20 hours.

2.2.5. Test Conditions

Each test has a unique combination of acetic acid concentration, pCO$_2$, and temperature regime whereas other parameters like condensation rate, test duration and overall pressure were kept constant. The acetic acid concentration is the total acetic acid by volume. Table 2 below shows all parameters used in the test. The parameters were chosen to be as close to real conditions as possible; restricted due to technical limitations, practicality and health and safety regulations.

<table>
<thead>
<tr>
<th>Table 2: Top-of-the-Line domain diagram test conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test duration</td>
</tr>
<tr>
<td>Total pressure at $T_{room}$</td>
</tr>
<tr>
<td>Condensation Rate</td>
</tr>
<tr>
<td>Bulk solution</td>
</tr>
<tr>
<td>pCO$_2$</td>
</tr>
<tr>
<td>Acetic acid</td>
</tr>
<tr>
<td>Temperature regime 1</td>
</tr>
<tr>
<td></td>
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<tr>
<td>Temperature regime 2</td>
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<td></td>
</tr>
</tbody>
</table>

The condensation rate was chosen to be well above the commonly accepted and well documented threshold for TOL corrosion in literature of 0.25 g/m$^2$/s (Gunaltun and Larrey 2000). The sample temperature of 30 °C
couldn’t be lowered due to the limitation of the circulating pump which had no cooling ability.

Acetic acid concentrations relate to the total acetic acid in solution. It has to be differentiated between total acetic acid and free acetic acid. The term total acetic acid will refer to the amount of acetic acid that is added into the solution in the beginning of the test and will include dissociated \((H^+ + CH_3COO^-)\) and undissociated acetic acid \((CH_3COOH)\). Free acetic acid will just refer to the undissociated acetic acid \((CH_3COOH)\). The amount of free acetic acid depends on the pH of the solution, the higher the pH the lower the free acetic acid concentration. Due to the high pressure nature of the tests and constantly changing pH it was more practical to work with total acetic acid and calculate free acetic acid after the tests.

More than 50 corrosion tests (including duplicates and triplicates) were conducted to gain the results for this domain diagram.

### 2.2.6. Corrosion Sample Identification

The sample names used in this chapter are arranged by the acetic acid concentration, \(CO_2\) partial pressure and temperature regime. The next two examples will demonstrate how the names are produced and how the information can be read.

Example:

500C10T9130: 500 ppm acetic acid; 10 bar \(pCO_2\); \(T_A\): 91 °C and \(T_S\): 30 °C

Or

0C5T11580: 0 ppm acetic acid; 5 bar \(pCO_2\); \(T_A\): 115 °C and \(T_S\): 80 °C

where \(T_A\) stands for the autoclave temperature and \(T_S\) for the sample temperature.
2.2.7. Surface Investigation

To evaluate the surface two different techniques have been used; a scanning electron microscope (SEM) and an Alicona Infinite Focus Microscope (IFM). The SEM was a Zeiss Evo with an Energy Dispersive X-ray (EDX) detector. The accelerating voltage, spot size and working distance are shown individually on each picture. In general, pictures taken by the SEM are in much higher magnification than pictures taken by the IFM. The focus of all surface investigations was on the lower side of the sample where the condensation rate accumulated.

The IFM is an optical microscope that is able to adjust all three axes (X, Y and Z axis) automatically in very small increments. A particular volume (X, Y and Z direction) is set by the user to be captured and the microscope takes the pictures. Then, software puts the pictures together and by just using the part of the picture that is in focus, the software develops a 3 dimensional model of the specimen. Using this technique it is possible to capture a magnified picture over a large area and depth, especially important for the curvature of the ¼ inch carbon steel tubes.

Using the software of the microscope, it was then possible to subtract the curvature of the sample and transform it digitally into a flat data set. By doing so, it was possible to measure the depths and size of the localized corrosion. Unfortunately, it was only possible to use this technique on limited parts of the sample due to the severe curvature of the sample in all three dimensions.

It was possible to calculate an important value; the probability that pits will be initiated in the given circumstances (ASTM-G46 2005).

The pitting probability (P) is calculated as

\[ P = \frac{N_P}{N} \times 100 \quad \text{Equation 14} \]

with \( N_P \) as the number of pitted specimens and \( N \) as total number of specimen in the given test.
2.3. Results and Discussion

2.3.1. Horizontal Cooled Tube Test development

The Horizontal Cooled Tube (HCT) test is a high pressure TOL corrosion bench top test developed and continuously improved at Curtin University. Originally it started as the so called “U-tube test” (Figure 6) with a vertical ¼ inch (0.635 cm) carbon steel tube (John et al. 2009). The water was condensing on the tube surface just to immediately run down the pipe wall to the tip of the “U”. The vertical shape made it impossible for the droplets to be saturated with iron carbonate and eventually form an iron carbonate layer.

A very similar effect was present at the bottom of the pipe despite that the droplets gathered at that spot. The condensation from the entire surface congregated there and even with an overall rather low condensation rate, with all condensation collecting at one spot, droplets were detaching too rapidly to form any protective layer. In that case, just the spot where the droplet formed can be defined as TOL corrosion.

Figure 6: Early version of the TOL corrosion test set up
The design was modified so that the $\frac{1}{4}$ inch tube was moved from the vertical into a horizontal position in order to spread the condensation over the entire surface of the tube. In consequence, the TOL corrosion was also more distributed over the entire sample surface increasing the actual surface area of TOL corrosion testing.

### 2.3.2. Top-of-the-Line Corrosion Domain Diagram

The Top-of-the-Line (TOL) Domain Diagram was developed to gain a better understanding of TOL corrosion in presence of different CO$_2$ partial pressures (pCO$_2$), acetic acid concentrations and temperature regimes. The condensation rate was kept constant to eliminate its contribution and focus on the other variables. Figure 7 shows the average corrosion rates of weight loss and AAS, which were performed as duplicate or triplicate if necessary. The corrosion rates by weight loss and by AAS were very consistent with each other; hence, the average was calculated and displayed in the same diagram.

![Figure 7: TOL domain diagram results; average CR determined by means of WL and AAS](image)
It is eye catching (and probably expected) that the corrosion rate increases with increased concentrations of acetic acid and decreases with increased sample temperature. In the following sections the participation of the different parameters on the corrosion rate will be discussed in more detail.

2.3.2.1. Effect of Temperature

In order to keep the condensation rate constant, it was not possible to vary the sample temperature and the gas temperature separately. Corrosion and corrosion product layer formation takes place at the actual steel sample, therefore the sample temperature is the important factor in this test and the gas temperature was adjusted accordingly.

In Figure 7, the effect of temperature is very obvious. The low temperature samples show in every case approximately double the corrosion rate of the respective high temperature sample. This can also be seen in the overall average of all low and high temperature tests of 0.61 mm/y and 0.3 mm/y, respectively. Therefore it can be said that the sample temperature played the most crucial role in these TOL corrosion studies. This finding is consistent with recently published data obtained using a different test apparatus but similar conditions (Ojifinni and Li 2011).

A protective iron carbonate corrosion product layer cannot form at low temperatures and with 30 °C, the critical temperature is not reached (Nazari et al. 2010). In Figure 8 the weight of the corrosion product scale of each condition is displayed. With overall averages of 0.011 and 0.039 g the high temperature samples developed 3.5 times more scale than the low temperature ones. After that scale has been developed on the surface, it protects the surface from further uniform corrosion. A surface layer for a low and high temperature sample can be seen in Figure 9; acetic acid and pCO$_2$ are the same in both samples, 1000 ppm and 20 bar, respectively. The images of the surface are representative for low and high temperature samples and it does not change significantly with varying acetic acid or CO$_2$ concentrations. It can be seen that the high temperature sample forms a very dense, thick, and protective iron carbonate layer (cubes in Figure 9.4) which
protects the surface of further uniform corrosion. An EDX picture and result can be seen in Figure 10 taken from another high temperature sample, indicating the presence of iron carbonate (FeCO$_3$).

The low temperature sample does not form such a protective layer. In some parts of the surface the iron is still fully exposed and iron carbonate forms only sporadically and in a thin layer. This can be seen in the EDX measurements shown in Figure 11. A better protection against uniform corrosion at higher temperatures goes along the lines of other studies (Kinsella et al. 1998; Ojifinni and Li 2011).

The very same iron carbonate corrosion layer that protects from uniform corrosion at higher temperatures makes the sample more prone to pitting and mesa corrosion. A representative sample is shown in Figure 12.2. Over 70 % of the high temperature samples show pitting and or mesa attack on the surface whereas there is no pitting corrosion or mesa attack visible on any of the low temperature samples (a representative sample shown in Figure 12.1) regardless of acetic acid concentration or CO$_2$ partial pressure (Localized corrosion will be discussed further in section 2.3.2.7).

![Figure 8: TOL domain diagram results; average weight of corrosion product](image-url)
Figure 9: SEM images from low (1, 2 - 1000°C/130°C) and high (3, 4 - 1000°C/1580°C) temperature samples

Figure 10: EDX measurements on a high temperature sample
Figure 11: EDX measurements on a low temperature sample on two different spots

Figure 12: HCT samples;
1 - low temperature (500°C20T9130) without localized corrosion
2 - high temperature (500°C20T11580) with localized corrosion
2.3.2.2. Effect of CO₂ partial pressure (pCO₂)

The effect of pCO₂ isn’t as obvious in Figure 7 as the effect of temperature. Therefore, the data was rearranged and displayed in Figure 13. It is shown that both high and low temperature samples follow the same trend with increasing pCO₂.

In the tests without acetic acid (the first three blue and orange columns) the corrosion rate increases with increasing pCO₂. In the case of the low temperature samples, the average corrosion rate increases over the 7 day period from 0.47 mm/y to 0.49 mm/y for 5 bar, 10 bar and 20 bar pCO₂, respectively. A similar increase can be observed for the high temperature samples where the corrosion rate stays constant for 5 bar and 10 bar at 0.20 mm/y and then increases for 20 bar pCO₂ to 0.25 mm/y. These results are consistent with many other studies where an increase of pCO₂ increases the corrosion rate due to a higher concentration of carbonic acid and therefore a lower pH in the liquid phase.

Within the tests containing 500 ppm acetic acid (the centre three blue and orange columns) the corrosion rate slightly decreases with rising pCO₂ at low temperature and basically stays constant for the high temperature testing. The effect of a falling corrosion rate is even more pronounced with a 1000 ppm acetic acid concentration.

The corrosion rate of the low temperature samples at 1000 ppm decreases from 0.80 mm/y to 0.66 mm/y and stays more or less constant at 0.67 mm/y with 5 bar, 10 bar, and 20 bar pCO₂, respectively. At high temperature virtually the same decrease of corrosion rate can be observed. The corrosion rate is 0.43 mm/y, 0.34 mm/y, and 0.35 mm/y for 5 bar, 10 bar, and 20 bar pCO₂.
The effect of a declining corrosion rate at the TOL with an increasing pCO$_2$ at high acetic acid concentration has not been reported before in the literature. Some publications would even suggest the opposite by saying that a higher pCO$_2$ decreases the pH in the condensed liquid which in turn locally re-associates acetic acid which then increases the corrosion rate. This effect is not observed in the present study.

It can be concluded that an increase in pCO$_2$ without any acetic acid present results in a decreasing pH of the condensed liquid which results in a more severe environment and therefore in an increased corrosion rate. This is in line with previously published data, experimental and predictive.

A different mechanism must come into play at medium and high acetic acid concentrations. More experiments need to be conducted in similar conditions with focus on the formation and structure of the protective film formed. At this stage is can be anticipated that the protective film that forms at elevated acetic acid concentrations becomes more protective with an increasing pCO$_2$. This might be the case due to the fact that the negative influence of a higher pCO$_2$, the lower pH in the condensed liquid, is not as
pronounced at higher acetic acid concentrations due to an already low pH in these conditions. A protective corrosion product layer was observed in previous studies, even in lower temperatures (Zhang et al. 2007).

2.3.2.3. **Effect of Total Acetic Acid**

The influence of acetic acid on TOL corrosion rate was observed to be very strong. Figure 7 clearly displays that within the same CO₂ partial pressure and temperature regime, the corrosion rate increases with increasing acetic concentration, except for one value which is 0C20T9130. This test was repeated 5 times and always gives a higher corrosion rate compared to the tests with acetic acid.

The average of all tests with the same total acetic acid concentration shows the same trend; the more acetic acid, the higher the corrosion rate (Figure 14). The low temperature tests show an average corrosion rate of 0.54 mm/y, 0.59 mm/y and 0.71 mm/y as an average of 0 ppm, 500 ppm and 1000 ppm of acetic acid, respectively. High temperature tests give an average corrosion rate of 0.22 mm/y, 0.30 mm/y and 0.38 mm/y for 0 ppm, 500 ppm and 1000 ppm acetic acid, respectively (Figure 14).

At high temperatures, the corrosion rate correlates linearly with the total acetic acid concentration. This can be seen by looking at the R-squared value. R-squared is a statistical tool that indicated how well a trend line, in this case a linear trend line, fits the values with 0 being no fit at all and 1 being a perfect fit. The high temperature samples display a perfect linear fit with an R-squared value of 1. The low temperature tests don’t seem to follow a linear increase in corrosion rate against acetic acid concentration. It shows a 0.05 mm/y increase from 0 ppm to 500 ppm acetic acid and a 0.12 mm/y increase from 500 ppm to 1000 ppm acetic acid, resulting in an R-squared value of 0.9465 (Figure 14).
By calculating the average without the unusually heavily corroded sample 0C20T9130 for the average corrosion rate without acetic acid at low temperatures a linearity can be observed (Figure 15). The average corrosion rate would be 0.48 mm/y for no acetic acid and would display an increase of 0.11 mm/y from 0 ppm to 500 ppm acetic acid and a 0.12 mm/y increase from 500 ppm to 1000 ppm acetic acid. The R-squared value in this case is 0.9994 which is a much better linear fit than the R-squared value of 0.9465 of the previously calculated CR.

Taking these values into consideration, the low temperature samples can be seen as more susceptible to an increasing acetic acid concentration displayed by the gradient of the corrosion rate.

The scale formation seems to be also affected by the acetic acid concentration as displayed in Figure 8. In 4 out of 6 sets (at 20 bar pCO2 and low temperature and all tests at high temperature) the weight of scale formed on the samples is the lowest at 0 ppm acetic acid, followed by 1000 ppm acetic acid and the highest weight of scale is at 500 ppm acetic acid. Most of them results are within the error bars of the tests, but a trend is clearly visible.
2.3.2.4. **Effect of Free Acetic Acid**

The free acetic acid can be calculated from the total acetic acid and the pH value (For more information see Equation 9 and Figure 2). The amount of free acetic acid was therefore calculated using the measured pH of the samples. Unfortunately, during the beginning of the testing, the pH was not measured for all tests; therefore not all samples can be evaluated here. Nevertheless, in Figure 16 the corrosion rates are plotted against the free acetic acid concentration for those experiments for which data was available. The blue and orange data points represent low and high temperature samples, respectively. Symbols of the same colour and shape are taken from the same test.

As a test progresses, the free acetic acid goes down due to a rising pH in the bulk solution. The corrosion rate also decreases with the decreasing free acetic acid over time in an almost linear way. For any given set of conditions the corrosion rate is higher for a higher free acetic acid concentration, which is again in line with previous published data (Singer et al. 2004; George and Nesic 2007; Singer et al. 2009). This goes along with the observation of a
higher average corrosion rate with an increasing total acetic acid concentration.

The data also reveal that the HCT TOL corrosion test rig in its current state is not able to maintain a constant specific free acetic acid concentration. Corrosion products in a droplet at the TOL constantly fall into the bulk solution, changing the pH and therefore the free acetic acid concentration. This could be addressed by slight modifications to the set-up, which will be discussed further in section 4.2 on page 132 onwards.

![Graph](image)

**Figure 16: Free acetic acid concentration against corrosion rate – high vs. low temperature**

### 2.3.2.5. The effect of bulk pH

The pH in the bulk solution was neither a constant nor a controlled variable of the test; it was the result of all test parameters and it was constantly changing throughout every test. The pH at the beginning of each test was essentially dependent on the acetic acid concentration and the pCO$_2$.

For the iron concentration measurements, a bulk solution sample was taken every 24 hours. The pH and iron concentration were measured and the corrosion rate calculated for this sample. Using this data, it was possible to
correlate an average corrosion rate over 24 hours for a representative pH range. This range of pH was determined from the measurement at the beginning and end of each 24 hour period. In Figure 17 (low temperature) and Figure 18 (high temperature) the corrosion rates are plotted against the pH range, regardless of what day of the 7 day test period the measurement was recorded. In the graphs the total acetic acid concentration is differentiated using different colours.

Regardless of the original total acetic acid concentration and the day of sampling, both, in the low and high temperature test the corrosion rate against pH graphs follow almost an exponential decay curve as indicated in red. Also at both temperatures, comparing tests where acetic acid is present and absent in the same diagram, it seems that there are numerous outliers in the tests without acetic acid. Most of the outliers of the low temperature testing are from first two days of testing 0C20T9130 sample with an unusually high corrosion rate especially in the beginning of the test. Nevertheless, the overall picture follows the same trend.

![Corrosion rate against pH for all low temperature samples; differentiated only in 0 ppm, 500 ppm and 100 ppm of acetic acid](image)

*Figure 17: Corrosion rate against pH for all low temperature samples; differentiated only in 0 ppm, 500 ppm and 100 ppm of acetic acid*
The pH has a very strong influence on the amount of free acetic acid; with the data shown in this section, it can be assumed that the corrosion rate is highly affected by both the pH itself and the resulting decrease of free acetic acid.

2.3.2.6. **Surface Roughness**

Using the Infinite Focus Microscope (IFM) it was possible to capture a large portion of the curved surface and transform it back into a flat area suitable for further data processing. Surface roughness was calculated but no pattern was observed in these results. For example, in Figure 19 the results of two surface roughness measurements are shown. Both measurements are performed on the same sample on the same spot with a totally different outcome. The results of Figure 19 are representative for most measurements. Because all samples are slightly different and the flattening of the surface worked slightly different for each sample, it was not possible to gather a representative and comparable roughness values.
2.3.2.7. **Localized Corrosion**

The evaluation for localized corrosion was performed also by means of the IFM but much more successfully than the roughness measurements. The localized attack was usually wide and flat bottomed (Figure 20) which allows easy examination with an optical microscope. Due to issues indicated in the previous section on “surface roughness” the values should be regarded as an approximation rather than an exact value of the pit depth.

Temperature has the most dominant effect on localized attacks within the tested parameters. Of all the samples tested at low temperature, just two show localized attacks. One sample was exposed to 0 ppm and the other one to 500 ppm of acetic acid; however, both samples were exposed to a high CO$_2$ partial pressure (20 bar). In both instances, the localized corrosion is very shallow (< 30 μm) and rather wide. The pitting probability for all low temperature samples is 7 % and therefore very low compared to the high temperature samples with 70 % pitting probability.

As indicated by the very high pitting probability, much more pitting was evident at high temperatures. All of the different parameter combinations showed one or more samples with localized corrosion except for 500 ppm acetic acid at pCO$_2$ = 10 bar.

In general, the localized attack was also more severe at high temperatures. The shape of most of the pits was wide with steep walls and a flat bottom, it
can be considered as mesa corrosion (Nyborg and Dugstad 2003). For mesa attack to form, a partially protective corrosion product film needs to form, which is the case at high temperatures in this test (Nyborg 1998). During the evaluation of the mesa corrosion it was observed that the number of localized attacks per sample increases with increasing acetic acid concentration regardless of the pCO₂.

The connection of acetic acid and pCO₂ to the width and depth of the localized attacks is not straightforward. Figure 21 plots the average depth of the localized corrosion attack for each condition. It can be seen that at low pCO₂ (5 bar) the pitting depth is significantly higher with 1000 ppm of acetic acid compared to 0 ppm and 500 ppm acetic acid. The total opposite effect was observed at high CO₂ partial pressure (20 bar) where the average depth of the localized corrosion is reduced with increasing acetic acid concentration. At 10 bar pCO₂, the depth of the localized corrosion increased slightly from 0 ppm to 1000 ppm acetic acid concentration but dropped to 0 μm at 500 ppm acetic acid (The occurrence of localized attacks is given as a pitting probability and therefore it might be coincidental that there is no pitting).

Figure 22 displays the results from Figure 21 in a rearranged form to emphasize the effect of pCO₂. A very similar trend is visible as seen with the acetic acid concentration. At 0 ppm acetic acid, the average localized corrosion depth increases with increasing pCO₂, whereas the average depth decreases at 1000 ppm acetic acid with increasing pCO₂. It is also slightly higher at 500 ppm acetic acid across the range from 5 to 20 bar pCO₂.

Regardless of the depth and size, an increase of localized attacks could be observed with increasing acetic acid concentration within the same CO₂ partial pressure range.

There are many contradictory publications available concerning the influence of acetic acid concentrations and pCO₂ at low pressures for BOL and TOL corrosion (Crolet et al. 1999; Mendez et al. 2005; Nafday and Nesic 2005; Amri et al. 2010; Fajardo 2011). The results indicate that there might be a more complicated relation between acetic acid, higher CO₂ partial
pressures, and pitting depth. It is very important to note that there is a pitting probability, it means that there is a high chance that pitting will occur, but it is not a pitting certainty. This may lead to the contradicting results in comparable conditions, especially in laboratory tests which are usually short in duration.

Figure 20: Example of localized corrosion
1) SEM image
2) IFM image (1.3 mm x 1 mm)

Figure 21: Average depth of the localized corrosion in different conditions; emphasising the effect of acetic acid
Figure 22: Average depth of localized corrosion in different conditions; emphasising the effect of pCO₂
2.4. Conclusions

The corrosion at the TOL is affected by many parameters.

- The main influence on TOL corrosion within the investigated parameters of this study had the pipe temperature. A dense and protective corrosion layer formed at high temperatures.

- Localized attacks were observed regularly at the higher temperatures.

- A high \( \text{CO}_2 \) partial pressure (20 bar) has a positive influence at high acetic acid concentrations (1000 ppm).

- A very important parameter is the pH of the bulk solution, which is directly related to the concentration of free acetic acid. The lower the pH, the more acetic acid is present in its undissociated state and therefore increases the corrosion at the TOL.

- No localized corrosion was observed for medium \( \text{pCO}_2 \) (10 bar) and medium acetic acid concentration (500 ppm).

- The lowest average depth of localized corrosion of the high temperature samples was observed for low and medium \( \text{pCO}_2 \) (5 and 10 bar) with no acetic acid as well as for high \( \text{pCO}_2 \) (20 bar) with a high acetic acid concentration (1000 ppm).

- The correlation between higher \( \text{pCO}_2 \) and acetic acid concentrations on TOL corrosion needs further investigation

- Influence on TOL corrosion of

  \[ \text{Temperature} > \text{acetic acid concentration} > \text{pCO}_2 \]
3. Generic Volatile Corrosion Inhibitor Compound Investigation

3.1. Introduction

3.1.1. CO₂ Corrosion Inhibition

Carbon steel is widely used as pipeline steel in the oil and gas industry for a vast variety of applications. Many of the environments are corrosive for carbon steel and therefore carbon steel requires protection. There are two main approaches protecting against CO₂ corrosion of carbon steel; corrosion inhibition or pH stabilisation. Inhibition by pH stabilisation increases the pH of the formation water to 6.5–7.5 which facilitates the formation of a protective iron carbonate layer on the steel surface, reducing Bottom-of-the-Line (BOL) corrosion considerably (Dugstad et al. 2003). The pH stabilisation method can be used in combination with glycol to prevent the formation of hydrate (Dugstad and Dronen 1999). In this way, the pH stabilising agents can be reclaimed with the glycol and it becomes a more cost efficient way of inhibition (Halvorsen and Andersen 2003). The limiting factor for this technique is formation water; it cannot be used with large quantities of formation water due to the formation of calcium carbonate scale close to the pipeline inlet at elevated pH levels (Nyborg 2009).

Corrosion inhibition is one of the most common corrosion control methods for oil and gas production and transportation (Kapusta 1999). The inhibitors are usually long chained organic compounds. These compounds attach their polar head to the carbon steel surface and form a physical barrier between the steel and the corrosive environment. The corrosion inhibitors were thought to form a mono layer, but more recent studies have shown that above a critical concentration they can adsorb to the surface in several “shapes” like spheres, rods, or double layers. The exact mode of adsorption to the steel highly depends on the type and concentration of the inhibitor (John
et al. 2006; Bosenberg et al. 2007). Tests have shown that inhibitors are effective in quantities as low as 10 ppm (Thomas 2000) and even lower at 0.5 ppm (Jovancicevic et al. 2000). The film formed by many inhibitors is proven to be very stable and cannot easily be removed at shear stresses commonly found in pipelines (Xiong et al. 2011).

The corrosion inhibitors are usually applied continuously by injection into the liquid phase of the pipeline. Consequently, the inhibitors only prevent corrosion in the liquid phase of the pipeline; therefore they are commonly referred to as Bottom-of-the-Line inhibitors. The gas phase above the liquid phase is beyond the influence of normal BOL inhibitors which leads to the potential risk of Top-of-the-Line (TOL) corrosion in wet gas pipelines.

3.1.2. Top-of-the-Line Corrosion Inhibition

The inhibition of TOL corrosion is a common problem in the transportation of wet gas. There are several different approaches to the problem.

The best known and longest used inhibition technique is the inhibitor batch treatment. A batch of inhibitor with a high film persistency is carried through the pipeline between two PIGs reaching and applying inhibitor to the TOL. The treatment has to be repeated periodically, otherwise the corrosion protection will decrease significantly over time (Punpruk et al. 2010). There are several problems with the batch treatment; not all pipelines are prepared to launch PIGs or the PIG launcher might be too small for the large volume of inhibitor required (Gunaltun and Belghazi 2001). Most importantly, the production has to be shut down to perform the operation which results in a huge production loss for the operating company.

In 2003, a novel approach using a PIG was presented, the TLCC-PIG (Gunaltun and Payne 2003). The TLCC-PIG is launched into the pipeline during production. The gas pressure propels the PIG through the pipeline. Utilizing the Venturi Effect, the TLCC-PIG sprays the conventionally inhibited formation water from the BOL to the TOL. The production does not have to be stopped entirely, which is a major advantage over the conventional PIGs. Nevertheless, some lines require cleaning before a TLCC-
PIG can be run to prevent the nozzles being jammed (Freeman 2009). Also, the possibility of a TLCC-PIG getting stuck in a pipeline is still present which would result in a huge loss of production and incur recovery costs.

The best solution to inhibit the TOL would be a continuous chemical treatment that is able to reach the TOL without the assistance of a PIG. Modes of transport to the TOL for continuous chemical treatment include foaming, spreading, and volatility.

Foaming inhibitors are generally not in use because the foam will cause problems in the processing of the formation water and natural gas or even require the addition of anti-foaming agents, increasing the total cost. Nonetheless, solely from the point of corrosion inhibition, published results indicate that foaming does work in the laboratory as a mode of inhibitor transport (Jevremovic et al. 2012).

The spreading inhibitor model was mainly researched and published by Günter Schmitt at al. where a spreading agent was reported to be able to carry the inhibitor to the TOL against gravity (Scheepers 2001; Schmitt et al. 2001). No literature could be found of a field trial performed using spreading inhibitors.

Volatile Corrosion Inhibitors (VCIs) are thought to be a promising way forward of continuous chemical TOL corrosion mitigation. VCIs are injected into the liquid phase of the pipeline, similar to conventional BOL corrosion inhibitors. It is believed that the VCIs evaporate and co-condense at the TOL and inhibit corrosion. Due to their volatility, VCIs are able to reach the entire surface of the carbon steel pipeline, including complex shaped parts and crevices (Andreev and Kuznetsov 1998). Unfortunately, VCIs are not as frequently researched as conventional BOL corrosion inhibitors. Therefore, the actual mechanism of inhibition of most individual VCI compounds is not known. Some commercially available VCIs are not even volatile and they simply stabilize the pH at the BOL. A higher pH at the BOL leaves less organic acid available for the TOL and therefore less TOL corrosion is occurring (Gunaltun et al. 2010). BOL pH stabilisation is indeed a way to mitigate TOL corrosion, but the inhibitor cannot accurately be called a VCI.
Reliable field data is also rarely published, and when it is, the identity of the compounds is often kept confidential (Gunaltun 2006; Pou 2009). Nevertheless, a paper has been published with an inhibition efficiency of about 70 % in a field trial (Punpruk et al. 2010). The scarce amount of information does leave enormous room for improvement and research for VCIs.

In the following sections, the TOL corrosion inhibition potential of a variety of amines is investigated by different methods. Amines with a variety of molecular structures, chemical, and physical properties were chosen to shed light onto TOL corrosion inhibition by VCIs.
3.2. Experimental

3.2.1. Generic Volatile Corrosion Inhibitor Candidates

Different potential generic VCI compounds were investigated for their ability to inhibit TOL corrosion (Table 3). Some of the VCIs were chosen because they are used in VCI applications outside the oil and gas industry. Some others were already used in commercially available VCIs in the oil and gas industry but were never thoroughly investigated. Again others were chosen due to their similarity in molecular formulation (small amino molecules) or physical properties (e.g. boiling point) (Bastidas et al. 1990; Petersen et al. 2004; Bhargava et al. 2009).

All 16 generic VCIs are amines or feature amine functional groups and therefore they are all bases. Amines of different classes were chosen including primary-, secondary-, tertiary- and cyclic amines. Some of the generic VCIs also have the characteristic of a combination of the classes mentioned. An extreme example combining all classes is 1-(2-aminoethyl)piperazine. It has three nitrogen atoms; a primary, a secondary and a tertiary one in a form of a ring (cyclic) with a tail. A graphic illustration of the different classes and their typical chemical structures can be found in Table 4. Dark grey atoms represent Carbon (C), small light grey atoms are Hydrogen (H), blue atoms are Nitrogen (N) and red atoms are Oxygen (O) in all graphics used hereafter.

The results of all compounds in every test are duplicated. If the results are not satisfactory, a triplicate was conducted. The numeric value presented for each individual test is the average of successful tests. It was deemed not practical to show individual error bars in the diagrams since it would either clutter the diagram in its current state or it would have been necessary to show the results individually, which in turn would have multiplied the amount of diagrams approximately by 8. Additional to the vast amount of diagrams, it would have made a visual comparison between the individual VCIs much harder, if not impossible.
A list of the investigated generic VCIs including their molecular formula can be found in Table 3. A table with the basic physical and chemical properties and a basic description for each compound can be found in the results and discussion (in section 3.3.2, Table 11 and in section 3.3.3, Table 16) for the respective compound. All compounds were sources from Sigma Aldrich where also all the MSDS sheets can be accessed.

The GVCIs were tested using a variety of test setups to investigate their TOL and BOL inhibition properties. Most of the generic VCIs underwent all tests, but some limitations apply due to a shortage of generic VCI or equipment usability. The different testing procedures will be described in the following sections.

**Table 3: List of GVCIs investigated**

<table>
<thead>
<tr>
<th>Name</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-(2-Aminoethyl)piperazine</td>
<td>C6H15N3</td>
</tr>
<tr>
<td>4-Aminomorpholine</td>
<td>C4H10N2O</td>
</tr>
<tr>
<td>Aniline</td>
<td>C6H7N</td>
</tr>
<tr>
<td>Benzylamine</td>
<td>C7H9N</td>
</tr>
<tr>
<td>Cyclohexylamine</td>
<td>C6H13N</td>
</tr>
<tr>
<td>Diethylamine (DEA)</td>
<td>C4H11N</td>
</tr>
<tr>
<td>Dicyclohexylamine</td>
<td>C12H23N</td>
</tr>
<tr>
<td>N,N-Dimethylethylamine (DMEA)</td>
<td>C4H11N</td>
</tr>
<tr>
<td>N-Methyl diethanolamine (MDEA)</td>
<td>C5H13NO2</td>
</tr>
<tr>
<td>4-Methylmorpholine</td>
<td>C5H11NO</td>
</tr>
<tr>
<td>Methylpiperazine</td>
<td>C5H12N2</td>
</tr>
<tr>
<td>Morpholine</td>
<td>C4H9NO</td>
</tr>
<tr>
<td>Octylamine</td>
<td>C8H19N</td>
</tr>
<tr>
<td>4-Picoline</td>
<td>C6H7N</td>
</tr>
<tr>
<td>Pyridazine</td>
<td>C4H4N2</td>
</tr>
<tr>
<td>Pyridine</td>
<td>C5H5N</td>
</tr>
</tbody>
</table>
3.2.2. Horizontal Cooled Tube (HCT) Test

Most of the experimental procedures and materials used to evaluate VCIs using the HCT test setup are identical to the procedures and materials used to develop the TOL domain diagram. Therefore, detailed information about the pressure reactor, corrosion samples, preparation of a test, and corrosion rate determination can be referred back to the experimental section of the previous chapter (Section 2.2 Experimental).

3.2.2.1. VCI Charging Process

A 150 mL high pressure liquid charging pipette manufactured by Swagelok was used to insert the VCI into the pressurized reactor. A tedious procedure
had to be followed to maintain the lowest possible oxygen contamination in the test chamber.

High purity CO₂ was connected to the top of the pipette (1). All valves were opened and CO₂ gas flowed through the pipette to remove oxygen. After a few minutes, the tube fitting (4) was attached to the valve of the pressure reactor while gas was still flowing through. After the tube fitting was connected to the valve of the pressure reactor, the lower valve of the high pressure pipette (3) was closed, leaving the tube (between 3 and 4) slightly under pressure but free of oxygen. Then, the pressure from the rest of the pipette was released and the CO₂ was disconnected and the top valve (2) was removed from the pipette. 100 mL of a pre-sparged 0.1 % NaCl solution and 1000 ppm of the VCI (0.75 mL) was inserted into the pipette and sparged for 10 minutes using high purity CO₂. After the 10 minutes of sparging, the sparge tube was removed and the top valve (2) was reattached to the pipette. The reattachment of the valve was the only step where a very small amount of oxygen ingress might have occurred. Then the CO₂ line was reconnected (1) and the pipette was pressurized to a higher pressure as the TOL test reactor. The lower valve (3) was opened, than the valve at the TOL test reactor was opened and the whole volume of the pressure pipette was transferred into the TOL test reactor.

After the process, a sample of the bulk solution was taken to have a baseline reading of iron from the bulk solution at the moment the VCI was inserted.

Figure 23: Liquid charging pipette
3.2.2.2. **Test Conditions**

The test conditions used for generic VCI testing can be seen in Table 5. All test parameters were kept constant throughout all generic VCI tests to allow maximum comparability.

<table>
<thead>
<tr>
<th>Table 5: Test conditions for generic VCI testing using the HCT test</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Test duration</strong></td>
</tr>
<tr>
<td><strong>Pre-corrosion</strong></td>
</tr>
<tr>
<td><strong>Bulk solution</strong></td>
</tr>
<tr>
<td><strong>Condensation rate</strong></td>
</tr>
<tr>
<td><strong>Total pressure/pCO₂ @ T_{room}</strong></td>
</tr>
<tr>
<td><strong>Acetic acid conc.</strong></td>
</tr>
<tr>
<td><strong>VCI conc.</strong></td>
</tr>
<tr>
<td><strong>Temperature regime</strong></td>
</tr>
</tbody>
</table>

3.2.3. **Cooled Finger Probe (CFP) Test**

The CFP approaches the testing of VCI from a totally different angle. Tests are performed under ambient pressure using a combination of corrosion under TOL conditions and electrochemistry in the condensed liquid. The test method was developed and performed at Nalco Energy Solutions in Singapore (Gough et al. 2009).

3.2.3.1. **CFP Corrosion Sample and Conditions**

The CFP test chamber is a slightly modified 2 L glass cell and a glass head with jointed fittings (Figure 24). Water saturated CO₂ was constantly sparged into the cell though the bulk solution which consisted of a 0.1 % sodium chloride solution with 1000 ppm acetic acid. The exiting CO₂ flowed through a bubbler to maintain ambient pressure and prohibiting oxygen from entering the cell. The bulk solution was heated using a hot plate connected to a thermocouple in the solution to control the solution temperature.
An externally cooled LPR probe was used for testing. Three 1018 grade carbon steel pins (ASTM-A29 2011) were attached to the LPR probe and subsequently cleaned for an hour in a cleaning alcohol. The carbon steel pins were commercial test samples commonly used for this type of probe and for each test a new set was used.

The LPR probe was suspended over a beaker inside the class cell. Liquid condensed at the three carbon steel pins corroding them under TOL conditions for 22 hours a day. The condensed liquid was then collected in the beaker. For 2 hours a day, the LPR probe was immersed into the collected condensed liquid in the beaker and LPR scans were performed using a Gamry Reference 600 potentiostat. The carbon steel pins are 4.75 mm in diameter and 30.5 mm long resulting in a total surface area for all three pins of 14.03 cm².

After the LPR scans were performed, the condensed liquid was sampled from the beaker. It was analysed for acetic acid (by means of gas chromatography), iron concentration (by means of a Hach spectrophotometer), and pH. Every 48 hours, a 25 mL sample from the bulk solution was taken and also analysed for acetic acid, iron and pH. Any liquid that was removed from the cell in any way was replenished into the bulk liquid with a pre-sparged 0.1 % sodium chloride, 1000 ppm acetic acid solution.

The CR calculated by means of LPR and IC does not necessarily give the same values of CR because different corrosion mechanisms are responsible for the corrosion. LPR essentially just measures the corrosivity of the condensed liquid after the 22 hours of TOL corrosion. The CR by means of IC on the other hand, is directly affected by severity of the TOL corrosion which is, in turn, highly affected by the condensation rate. Nevertheless, both follow the same trend indicating whether or not a VCI compound has inhibiting properties, which can be seen in the “Results and Discussion” section.

In addition to the LPR tests and iron concentration (IC) measurements, the corrosion rate was also determined by means of weight loss of the carbon steel pins.
The acetate concentration was determined by means of ion chromatography. With the acetate concentration and the pH it was then possible to calculate the concentration of the dissociated and undissociated acetic acid; in other words, how much of the measured acetate is present in the solution as acetic acid and therefore contributes to TOL corrosion (more in section 2.1.3).

\[
\% \text{CH}_3\text{COOH} = \frac{100}{1 + 10^{\text{pH} - \text{pK}_a}} \quad \text{Equation 9}
\]

The test duration was 7 days, where the first two days were dedicated to a pre corrosion of the sample under TOL conditions but without VCI in the system. After the two days of pre-corrosion, the generic VCI compound was inserted into the bulk solution.

As the condensed liquid removed from the cell, the condensation rate was calculated daily and was found to be between 0.25 g/m\(^2\)/s and 0.40 g/m\(^2\)/s using the following equation:

\[
\text{Condensation Rate} = \frac{m}{A \times t} \times 1000 \quad \text{Equation 13}
\]

where \(m\) is the mass of condensed liquid in kilograms, \(A\) the sample area in square meters and \(t\) is the time in seconds over which the condensation occurs (here 22 hours = 79200 s).

<table>
<thead>
<tr>
<th>Table 6: Test conditions for the cooled finger probe tests</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Test duration</strong></td>
</tr>
<tr>
<td><strong>Pre-corrosion</strong></td>
</tr>
<tr>
<td><strong>Bulk solution</strong></td>
</tr>
<tr>
<td><strong>Condensation rate</strong></td>
</tr>
<tr>
<td><strong>Total pressure</strong></td>
</tr>
<tr>
<td><strong>pCO(_2)</strong></td>
</tr>
<tr>
<td><strong>Acetic acid conc.</strong></td>
</tr>
<tr>
<td><strong>VCI conc.</strong></td>
</tr>
<tr>
<td><strong>Temperature regime</strong></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>
Figure 24: Image and schematic of the Cooled Finger Probe test set up
3.2.4.  Altered Horizontal Cooled Tube (A-HCT) Test

An alternative version of the HCT test was invented to minimize the effect of the VCI on the pH of the bulk solution. In this way, the volatility and inhibition effect directly at the TOL can be investigated.

To achieve this objective, two stainless steel cups were attached to the inside of the reactor. A schematic and a photograph are shown in Figure 25. The top cup is designed to collect any condensation and corrosion product to prevent it from mixing with the bulk solution and altering the pH. The lower cup holds a VCI solution and separates the VCI from the bulk solution, again to prevent any changes in pH. The lower cup is partly submerged in the bulk solution to maintain the same temperature. The VCI concentration of 1000 ppm is calculated by means of the total solution in the pressure reactor. It takes the 750 mL of bulk solution as well as the 100 mL of solution in the lower beaker into account, totalling 850 mL. Therefore, 0.85 mL of VCI was inserted in the lower beaker.

The contents of the top cup were removed daily and a 10 mL bulk solution sample was taken at least every second day. Both samples were analysed for their iron concentration by means of AAS and the pH was measured.

The bulk solution and condensed liquid samples are drawn by opening the valve of the autoclave and the high pressure inside the autoclave forces the fluid out. Because of large sample sizes (approx. 85 mL per day) and therefore large pressure drop, the internal pressure was topped up to the original pressure of the autoclave after two days. The test conditions for the A-HCT testing are displayed in Table 7.
Table 7: Test conditions for generic VCI testing using the A-HCT test

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test duration</td>
<td>5 days (approx. 90 h)</td>
</tr>
<tr>
<td>Pre-corrosion</td>
<td>No pre-corrosion</td>
</tr>
<tr>
<td>Total liquid in vessel</td>
<td>850 mL 0.1 % NaCl</td>
</tr>
<tr>
<td>Condensation rate</td>
<td>0.4 g/m²/s</td>
</tr>
<tr>
<td>Total pressure/pCO₂ @ T_{room}</td>
<td>20 bar (290 psi)</td>
</tr>
<tr>
<td>Acetic acid conc.</td>
<td>1000 ppm</td>
</tr>
<tr>
<td>Total VCI conc.</td>
<td>1000 ppm</td>
</tr>
<tr>
<td>Temperature regime</td>
<td>91 °C T_{Autoclave}</td>
</tr>
<tr>
<td></td>
<td>30 °C T_{Sample}</td>
</tr>
</tbody>
</table>

Figure 25: Schematic and photograph of the A-HCT test setup
3.2.5. Rotating Cylinder Electrode (RCE) Test

RCE tests were performed to investigate the inhibition effect of the VCI on the BOL corrosion. The aim also was to investigate the mechanism of inhibition for the given VCIs.

500 mL of a 0.1 % NaCl solution with 1000 ppm of acetic acid was used to perform all electrochemical tests. The solution was pre-sparged for about 60 minutes with CO$_2$ and continuously sparged throughout the test at a rate of approximately 50 ml/min. A silver silver-chloride reference electrode and a platinum mesh counter electrode were used in each test. Low rotation speeds of 200 rpm were used throughout the testing (Figure 26).

A 1030 grade carbon steel sample with a density of 7.87 g/cm$^3$ was used as the RCE sample. Samples were machined from a rod and subsequently stored in a desiccator until they were used. It was wet ground using water with 120, 320 and 600 grit silicon carbide grinding paper and rinsed with water and acetone. Afterwards, it was put into a beaker containing acetone and sonicated in the ultrasonic bath for 10-15 minutes. Then, it was rinsed again with fresh acetone, dried, and screwed onto the RCE sample holder between two Teflon washers to prevent crevice corrosion. The diameter and height were measured to calculate the sample area (approximately 3 cm$^2$) and the test was started.

A Gamry Reference 600 potentiostat with Gamry software was utilized to run a test sequence consisting of an open circuit potential (OCP) for 30 minutes, followed by alternating linear polarisation resistance (LPR) scans and electrochemical impedance spectroscopy (EIS), and finally a potentiodynamic (PD) scan. The entire test duration was 18 hours.

1000 ppm of VCI was inserted into the system after 4 or 5 LPR and EIS tests (approximately 4 hours) to record the blank corrosion rate for every sample. The pH was measured right before and after the VCI was inserted. Two different approaches have been tested for every VCI sample:

- After the pre-corrosion the VCI was injected into the solution. The VCI raises the pH of the solution and inhibits the corrosion of the steel
sample. The overall inhibition performance of the VCI was verified in this way.

- In other tests, the VCI was also injected into the bulk solution after the pre-corrosion. The difference was that the pH was adjusted back to a value before the VCI was inserted into the system (pH 3.20) using HCl and NaOH. In this way, the inhibition by neutralisation was eliminated and the ability of the VCI to inhibit by a different mechanism was investigated. It can be assumed that the residual inhibition mechanism was due to film forming.

In the following “Results and Discussion” section both results are distinguished by the addition “+ HCl”.

It was possible to determine the ratio of inhibition by film forming (Inhibition efficiency of the pH adjusted tests “+HCl”) and neutralisation (difference in inhibition efficiency of pH adjusted and non-adjusted tests). Inhibition efficiency will be introduced in section 3.2.7.

Figure 26: Rotating cylinder test set up
3.2.5.1. **Linear Polarisation Resistance (LPR)**

The OCP was allowed to stabilize for 30 minutes before LPR measurements were initiated. The LPR was measured ±10 mV around the OCP with a scan rate of 0.125 mV/s. Beta A and Beta C values of 0.12 were assumed (Ogundele and White 1986) to calculate the Stern-Geary constant (B) of 26.05 using the following equation

\[ B = \frac{ba \times bc}{2.303 (ba + bc)} \]  

Equation 15

The calculated Stern-Geary constant was then used to calculate the corrosion current density \( i_{corr} \) in the following equation

\[ i_{corr} = \frac{B}{R_p} \]  

Equation 16

where \( R_p \) is the polarisation resistance measured in the LPR test.

An example of a LPR scan is displayed in Figure 27. The polarisation resistance \( R_p \) has to be gained out of each of the LPR plots to calculate the \( i_{corr} \) using Equation 16. \( i_{corr} \) is then used to calculate the CR in Equation 17.

\[ CR = K_1 \times \left( \frac{i_{corr}}{\rho} \right) \times EW \]  

Equation 17

where \( K_1 \) is a constant of 3.27 \( \times 10^{-3} \) mm g/\( \mu A \) cm y, the constant is needed that the resulting CR is given in mm/y, \( i_{cor} \) is the corrosion current density in \( \mu A/cm^2 \), \( \rho \) the density of the material in g/cm\(^3\), in this case 7.87 g/cm\(^3\), and 27.92 was used as the equivalent weight (EW) (ASTM-G102-89 2010). The calculated CRs are then plotted as a function of time to evaluate the VCIs. It is possible to run subsequent tests because LPR is essentially a non-destructive technique (Fontana and Staehle 1980; Enos 1997; Mansfeld 2005).
Figure 27: Linear polarisation resistance (LPR) plot of the blank sample – 500 mL of 0.1 % NaCl solution with 1000 ppm acetic acid at 200 rpm

LPR is not able to distinguish between charge transfer resistance and solution resistance and is better suited for solutions with a high conductivity (and therefore low solution resistance). To verify the LPR results, EIS measurements were performed using the RCE test set up.

3.2.5.2. **Electrochemical Impedance Spectroscopy**

After an initial LPR test a subsequent EIS test was performed (both, LPR and EIS were continuously alternated). EIS measurements were performed between 10000 Hz and 0.01 Hz frequency with an AC voltage of 10 mV rms and 10 points/decade were recorded. The focus of the investigation was on the Nyquist plot (Figure 28). The advantage of EIS is the possibility to clearly distinguish between solution resistance \( R_S \) and charge transfer resistance \( R_{CT} \) where \( R_S \) should not be incorporated in the calculation of the CR.

Figure 28 is an actual blank EIS measurement of the investigation. It can be seen that the solution resistance is quite high, but more on this topic in the Results and Discussion of this chapter.

ZView software (Scribner Associates Inc.) was used to extract \( R_S \) and \( R_{CT} \) from the recorded data. The CR was then calculated using Equation 15,
Equation 16, and Equation 17. All results are displaying the evolution of the CR over time and no single EIS plots are shown anymore.

![Figure 28: EIS (Nyquist) plot of the blank sample – 500 mL of 0.1 % NaCl solution with 1000 ppm acetic acid at 200 rpm](image)

### 3.2.5.3. Potentiodynamic Scan

At the end of each test sequence a potentiodynamic scan was performed from -250 mV to +500 mV around the OCP with a scan rate of 0.167 mV/s.

### 3.2.6. Corrosion Rate

All TOL corrosion test methods feature two or more ways to determine the CR of the samples. In all cases, the CR was calculated by means of weight loss (WL) and by means of the iron concentration (IC) in the solution. In the case of the CFP, LPR measurements were used as an additional method to calculate the corrosion rate. The formulas used to calculate the CR by WL and IC can be found in section 2.2.3 on page 15 “Corrosion Rate Determination”. To calculate the CR by means of LPR and EIS the Equation 15, Equation 16, and Equation 17 were used.
3.2.7. Inhibition Efficiency

The inhibition efficiency (IE) was calculated for every compound in every TOL corrosion test. The higher the IE, the more efficient the inhibitor works. It is calculated by the following equation:

\[
IE = \left( \frac{CR_{Blank} - CR_{Inhibited}}{CR_{Blank}} \right) \times 100
\]

Equation 18

with IE given in percent (Durnie et al. 2001). The unit of CR is irrelevant as long it is used consistently; here mm/y was used.

To calculate the IE the pre-corrosion values were not taken into account for electrochemical (LPR and EIS) and iron count data. Technically, the longer a pre-corrosion would last, the lower the IE of an inhibitor would be. By not taking the pre-corrosion into account, this effect can be eliminated and the IE data is more representative for the individual generic VCI compound.
3.3. Results and Discussion

In this section, the performance of the individual GVCI compounds will be discussed separately for each test setup. Due to the vast amount of compounds and test methods corresponding results to discuss, they were split into two groups based on their overall performance.

The bottom 8 performing TOL corrosion inhibitors will be discussed in a group and the focus will be on some key attributes of the different compounds.

The top 8 performing TOL corrosion inhibitors will be discussed individually for each test setup. The aim is to propose a mechanism of inhibition for each compound.

All VCIs are compared to the respective blank test for each set up which will be presented in the following section.

3.3.1. Blank Test – No Inhibitor

To define an uninhibited CR, blank tests for each set up were performed. The blank tests were identical to all following VCI tests except for the insertion of the actual VCI into the system. All upcoming inhibitor efficiency (IE) measurements will refer to the blank test of the respective set up.

3.3.1.1. Horizontal Cooled Tube - Blank

The results for the blank test in the HCT test can be seen in Table 8 and Figure 29. The CR during the first two days is around 1.5 mm/y and falls on day three to approximately 0.6 mm/y where it plateaus until the end of the test. This is a huge drop for an uninhibited system. The main reason for the declining CR is the neutralization of the bulk solution via the iron corrosion product in the droplets which reduces the volatile amount of acetic acid and therefore the amount of acetic acid available for TOL corrosion. Nevertheless, the average corrosion rate by weight loss is 0.84 mm/y and via
iron concentration it is 0.71 mm/y. The weight loss via iron concentration does not include the first day, which explains the slightly lower average CR.

The relationship between the pH of the bulk solution and the CR at the TOL becomes clear in Figure 29. The CR is the highest during the first 48 hours of the test (above 1.4 mm/y), during this time the pH increases rapidly from pH 3.2 to pH 4.5. As the pH is above 4.5, the CR drops down to approximately 0.5 mm/y and stays rather constant while the pH still rises, but not as fast as in the beginning.

The blank CR and pH graphs will be in displayed in the diagrams of the VCIs for the purpose of comparison in a grey colour. The blue line is a “trend line” to highlight development of the data points over time. The average CRs will be used to calculate the IE of the VCIs.

<table>
<thead>
<tr>
<th>Table 8: Corrosion rates and inhibition efficiencies from the blank HCT test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iron conc. (IC)</td>
</tr>
<tr>
<td>CR [mm/y]</td>
</tr>
<tr>
<td>IE [%]</td>
</tr>
</tbody>
</table>

Figure 29: Corrosion rate by iron concentration, pH, and corrosion rate trend line from the HCT blank test (without VCI)
3.3.1.2. Cooled Finger Probe - Blank

The blank results for the corrosion test using the CFP are displayed in Figure 30. The CR by means of LPR is higher throughout the test compared to the CR by means of the IC (as described in section 3.2.3.1). Both, the CR by means of LPR and IC follow the same trend though. They slightly decline during day 1-4 and stay levelled at about 0.8 mm/y and 0.4 mm/y, respectively.

The yellow curve in Figure 30 is the calculated condensation rate for the respective time period. It stays fairly constant throughout the entire test. The condensation rate slightly increases from day 2 to day 3 which can immediately be seen in the CR by means of IC, which decreases at a slower rate compared to the day before or day after. It is well documented in the literature that the condensation rate has a strong effect on the CR at the TOL. At high condensation rates the corrosion product containing droplet is diluted quickly by condensing liquid, which in turn, makes it hard form a protective corrosion product layer since it the iron super saturation level cannot be reached.

In Figure 31 the pH of the bulk solution and the condensed liquid are displayed. The pH of the bulk solution stays very constant around pH 3. The pH of the bulk solution stays throughout the experiment because, in contrast to the HCT test (Section 3.3.1.1) because the condensed liquid with the corrosion product are collected in an isolated beaker totally separating bulk solution and condensed liquid. The pH of the condensed liquid also stays relatively constant but slightly declines from pH 5.3 on the first day to pH 4.9 on the last day.

The CR by weight loss and average CRs by means of LPR and IC are shown in Table 9. The pre-corrosion was not taken into account for CR by LPR and IC to be able to calculate the true IE in the later tests.

The curves in Figure 30 and Figure 31 will be displayed in grey in all upcoming graphs of VCI compounds and the values of Table 9 will be used to calculate the IE for the VCI compounds.
Table 9: Corrosion rates and inhibition efficiencies from the CFP blank test

<table>
<thead>
<tr>
<th>LPR</th>
<th>Iron conc. (IC)</th>
<th>Weightloss (WL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CR [mm/y]</td>
<td>0.78</td>
<td>0.40</td>
</tr>
<tr>
<td>IE [%]</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Figure 30: Corrosion rate and condensation rate of the blank sample from the A-HCT test

Figure 31: pH of the bulk solution and condensed liquid of the blank sample from the A-HCT test
3.3.1.3. Altered Horizontal Cooled Tube – Blank

The blank CR of the A-HCT test slightly decreases from day 1 to day 2 from 1.5 to 1.4 mm/y but then increases again up to 1.9 mm/y by the end of the test as seen in Figure 32. The development of the CR correlates very well with the measured condensation rate which essentially stays constant on day 1 and 2 around 0.4 g/m²/s but then increases up to 0.6 g/m²/s. This increase of condensation rate might be explained with the decrease of pressure due to the sampling process. The observed increase in CR generated by an increase in condensation rate goes along with a similar observation in the blank test of the CFP in section 3.2.3.1 Therefore, the close relationship of condensation rate and CR is witnessed in two blank tests in different test set-ups.

Figure 33 shows that the pH of the bulk solution and condensed liquid stay constant over time. The bulk solution increases from pH 3.3 to 3.6 whereas the pH of the condensed liquid decreases from pH 6.1 to pH 5.8.

The average CRs are listed in Table 10. As with the other test methods, the average CRs are used to calculate the IE for the VCI compounds and the necessary blank curves will be displayed in grey for a good comparison.

<table>
<thead>
<tr>
<th>Iron conc. (IC)</th>
<th>Weightloss (WL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CR [mm/y]</td>
<td>1.61</td>
</tr>
<tr>
<td>IE [%]</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 10: Corrosion rates, inhibition efficiencies and scale from the blank A-HCT test.
**Figure 32:** Corrosion rate and condensation rate from the blank A-HCT (without VCI)

**Figure 33:** pH of the bulk solution and condensed liquid from the A-HCT blank sample
3.3.1.4. **Rotating Cylinder Electrode**

The blank CRs by means of LPR can be seen in Figure 34 as the scatter diagram. Two different tests were conducted; the first test was left alone from beginning to end (black graph labelled 'Blank - LPR') and in the second, the pH was slightly adjusted to pH 3.20 after 5 hours using HCl. The adjustment was later performed with all VCIs as well to eliminate the neutralising effect of the VCI to the solution.

In both blank tests, the CR stays constant over the entire test period at around 1.0 mm/y. The pH at the start of both experiments was 3.25 and at the end of the blank test, it was 3.48. The slight increase in pH was due to the corrosion products in the solution. These curves are used as reference for later LPR measurements.

LPR measurements were alternated with EIS measurements throughout the tests. The EIS results for the blank tests can also be seen in Figure 34 as line curves. Both blank tests show a CR of around 2.5 mm/y. It can be seen that the CRs calculated by means of EIS are about 2.5 times higher than the CRs calculated by LPR. As explained in the

LPR is not able to distinguish between charge transfer resistance and solution resistance and is better suited for solutions with a high conductivity (and therefore low solution resistance). To verify the LPR results, EIS measurements were performed using the RCE test set up.

Electrochemical Impedance Spectroscopy section 3.2.5 before, EIS can differentiate between $R_P$ and $R_S$ and therefore calculates a more exact CR where LPR includes $R_S$ to $R_P$ resulting in a higher resistance and therefore lower CR. Nevertheless, LPR is a standard method to determine the CR in the field and laboratory which will therefore be continued throughout the VCI investigation as comparison.
A potentiodynamic scan from -0.25 to +0.5 V around the OCP was performed at the end of every test procedure. The scans can be seen below in Figure 35 with an \( E_{\text{corr}} \) of -0.67 V for the blank and 0.63 V the blank with HCl sample. The potentiodynamic scans of the VCIs will be compared to the respective blank tests.
3.3.1.5. **Conclusion on the Blank Tests**

The Blank tests are going to be used as a baseline and reference for all tests including VCIs. For the blank TOL corrosion samples, it can be concluded that the CR is very responsive to the condensation rate. An impact of the condensation rate on the CR was observed in two tests, the CFP and the A-HCT test, which are also the only tests where the condensation rate is being monitored.

CR rate values of LPR and EIS tests do not match well in the blank test. Nevertheless, both techniques indicate a stable CR over time in the blank tests. Since LPR is a standard test in many laboratories, both techniques will be continued to investigate the VCIs and the electrochemical techniques.
3.3.2. Low performing VCIs - Results

In the following section, 8 of the generic VCI compounds will be discussed. These 8 compounds were the compounds with the lowest performance in regards of TOL corrosion inhibition. The molecular structures (displayed using the free molecule editor software Avogadro 1.0.3) and some of the basic properties of each molecule can be found in Table 11. The properties of the molecules were taken from the respective MSDS sheets accessed via the homepage of the supplier, Sigma Aldrich (MSDS 2009; MSDS 2010; MSDS 2011; MSDS 2011; MSDS 2012; MSDS 2012; MSDS 2012; MSDS 2012).

The molecular structure is displayed as grey, blue, red, and small light grey spheres representing carbon, nitrogen, oxygen, and hydrogen, respectively.

All of the compounds are amines or involve an amino group. Amino groups can be found either as primary, secondary, tertiary, cyclic amino group, or in any combination. Within the compounds discussed in this section, aminomorpholine is the only molecule that contains an oxygen atom.

The grey graph in every diagram represents the blank test and every VCI compound has its designated colour/marker combination throughout the entire “Low performing VCI” discussion.
Table 11: Structures and properties of all low performing VCI compounds

<table>
<thead>
<tr>
<th></th>
<th>Aminoethyl-piperazine</th>
<th>Aminomorpholine</th>
<th>Aniline</th>
<th>Dicyclohexylamine</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Formula</strong></td>
<td>C6H15N3</td>
<td>C4H10N2O</td>
<td>C6H7N</td>
<td>C12H23N</td>
</tr>
<tr>
<td><strong>Molecular weight [g/mol]</strong></td>
<td>129.2</td>
<td>102.14</td>
<td>93.13</td>
<td>181.32</td>
</tr>
<tr>
<td><strong>Boiling point [°C]</strong></td>
<td>218-222</td>
<td>168</td>
<td>184</td>
<td>256</td>
</tr>
<tr>
<td><strong>Vapour pressure [mm Hg]</strong></td>
<td>0.076 (@ 20 °C)</td>
<td>1.65 (@ 25 °C)</td>
<td>1.00 (@ 35 °C)</td>
<td>0.034 (@ 25 °C)</td>
</tr>
<tr>
<td><strong>Purity [%]</strong></td>
<td>99</td>
<td>97</td>
<td>99</td>
<td>99</td>
</tr>
<tr>
<td><strong>Molecular Structure</strong></td>
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<td><img src="image2" alt="Structure" /></td>
<td><img src="image3" alt="Structure" /></td>
<td><img src="image4" alt="Structure" /></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Octylamine</th>
<th>Picoline</th>
<th>Pyridazine</th>
<th>Pyridine</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Formula</strong></td>
<td>C8H19N</td>
<td>C6H7N</td>
<td>C4H4N2</td>
<td>C5H5N</td>
</tr>
<tr>
<td><strong>Molecular Weight [g/mol]</strong></td>
<td>129.24</td>
<td>93.13</td>
<td>80.09</td>
<td>79.1</td>
</tr>
<tr>
<td><strong>Boiling point [°C]</strong></td>
<td>175-177</td>
<td>145</td>
<td>208</td>
<td>115</td>
</tr>
<tr>
<td><strong>Vapour pressure [mm Hg]</strong></td>
<td>1.00 (@ 20 °C)</td>
<td>4.00 (@ 20 °C)</td>
<td>2.33 (@ 25 °C)</td>
<td>18.00 (@ 25 °C)</td>
</tr>
<tr>
<td><strong>Purity [%]</strong></td>
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<td>99</td>
<td>98</td>
<td>99</td>
</tr>
<tr>
<td><strong>Molecular Structure</strong></td>
<td><img src="image5" alt="Structure" /></td>
<td><img src="image6" alt="Structure" /></td>
<td><img src="image7" alt="Structure" /></td>
<td><img src="image8" alt="Structure" /></td>
</tr>
</tbody>
</table>
3.3.2.1. **Horizontal Cooled Tube**

The HCT was used as a first evaluation of the VCIs. Due to a very limited availability, aminomorpholine was not tested in the HCT test.

The evolution of the CR over time is displayed in Figure 36. It is visible that all VCIs follow more or less the same pattern of CR over time, except for three candidates. The majority of the VCI candidates significantly decrease the CR as soon the VCI was inserted into the autoclave. Nevertheless, tests that involve picoline, pyridazine or pyridine do not follow the pattern mentioned before. The CR of the test including pyridine declines slightly after it was inserted, but levels out around 0.80 mm/y and even increases on the last day. A very similar trend is shown by picoline and pyridazine except that CRs for both decrease only during the last day. All three compounds show a negative IE by means of WL and a very weak or even negative IE by means of IC (Table 12). The neutralisation curves of these three VCIs couldn’t be more different as seen in Figure 37. Pyridazine has the weakest neutralisation effect on the bulk solution of all tested VCIs (from pH 3.3 to pH 4.1), whereas picoline and pyridine belong to the VCIs with the greatest neutralising effect with an increase to pH 5.2 and pH 4.9, respectively (blank pH 3.5).

Within the group of the low performing VCIs aminoethylpiperazine, aniline, dicyclohexylamine, and octylamine were among the better performers in the HCT test set up. All tests follow the same trend in which the CR drops directly after the VCI was inserted into the system. Aminoethylpiperazine and dicyclohexylamine manage to drop the CR by IC to 0.74 mm/y and 0.60 mm/y on the day after insertion, respectively. Both VCIs maintain a CR around 0.50 mm/y from day 3 until the end of the test. Aniline and octylamine follow a very similar trend but show greater fluctuation from day 3 to the end. The average CR by means of IC decreases through the VCIs aminoethylpiperazine, aniline, dicyclohexylamine and octylamine to 0.59 mm/y, 0.52 mm/y, 0.50 mm/y to 0.47 mm/y, respectively (Table 12) compared to a blank CR of 0.71 mm/y. The same trend can be seen with the CR by means of WL.
Aminoethylpiperazine achieves the highest pH neutralisation of the bulk solution but performs as the worst VCI in this group of four. Straight after injection the pH increases to pH 5.5 and at the end of the experiment is was at pH 5.9. The three other VCIs, aniline, dicyclohexylamine, and octylamine all affect the pH of the bulk solution the in a similar way. Straight after insertion, the pH of all three is around 4.6, and at the end of the test, the pH ends up around 5.1.

Table 12: Corrosion rates and Inhibition efficiency by weight loss and iron count from the HCT test

<table>
<thead>
<tr>
<th></th>
<th>CR [mm/y]</th>
<th>IE [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IC</td>
<td>WL</td>
</tr>
<tr>
<td><strong>Aminoethylpiperazine</strong></td>
<td>0.59</td>
<td>0.75</td>
</tr>
<tr>
<td><strong>Aniline</strong></td>
<td>0.52</td>
<td>0.71</td>
</tr>
<tr>
<td><strong>Dicyclohexylamine</strong></td>
<td>0.5</td>
<td>0.64</td>
</tr>
<tr>
<td><strong>Octylamine</strong></td>
<td>0.47</td>
<td>0.54</td>
</tr>
<tr>
<td><strong>Picoline</strong></td>
<td>0.66</td>
<td>1</td>
</tr>
<tr>
<td><strong>Pyridazine</strong></td>
<td>0.61</td>
<td>1.6</td>
</tr>
<tr>
<td><strong>Pyridine</strong></td>
<td>0.96</td>
<td>1</td>
</tr>
<tr>
<td><strong>Blank</strong></td>
<td>0.71</td>
<td>0.84</td>
</tr>
</tbody>
</table>

Figure 36: Corrosion rates from the HCT test – VCI was injected right after the first measurement
3.3.2.1.1. **Conclusions from the Horizontal Cooled Tube Test**

Most of the tested VCIs display their very good ability of neutralisation in this test. All VCIs increase the pH straight away. None of the tested compounds displays good inhibition ability in this test. It can be concluded that bulk neutralisation alone does not determine the ability of a good VCI. This can be seen especially in the case of aminoethylpiperazine and picoline, which raise the pH the highest throughout the test but exhibit a comparatively poor IE. The best performing VCIs of this group in the HCT test are octylamine and dicyclohexylamine with only less than 35 % IE.
3.3.2.2. Cooled Finger Probe

The same VCIs were tested in the CFP set up and will be discussed in this section. Aniline was not available for this test; therefore aminomorpholine was included in this series of experiments. Using the CFP, it was possible to track the evolution of the CR over time by means of two different methods; LPR (Figure 38) and iron count (Figure 39). Additionally the CR was also measured by means of WL and the condensation rate of every 24 hour period recorded.

It is worth reiterating that the VCI was injected into the system after 48 hours, which means that the first CR affected by the VCI is after 72 hours. Also important to note is the fact that Figure 38 and Figure 39 have a different scale on the Y-axis. In general, all CR calculated by means of IC are lower compared to the CR calculated by means of LPR. Also, the pH of the bulk solution of the CFP test is lower compared to the HCT test since the entire corrosion product is collected in the condensed liquid beaker, and therefore does not elevate the bulk pH.

Similar to the HCT test, most of the tests follow a similar pattern where the CR by means of LPR decreases after the VCI was inserted and then stays relatively level. Tests containing aminomorpholine and picoline as VCI do not follow this pattern. The CRs by means of LPR for both tests stay above the blank CR throughout the test. Picoline also has a higher CR throughout by means of IC and the CR by IC of aminomorpholine equals the blank at the end of the test. Both VCIs display a very poor IE overall, with both at negative IE values (Table 13).

The correlation of condensation rate and corrosion rate by means of IC for picoline seems very counterintuitive. When the condensation rate increases, the CR decreases; and when the condensation rate decreases, the CR increases throughout the test. Picoline is the only investigated compound in the CFP test that has that kind of behaviour. It seems that picoline affects the volatility of acetic acid. The test with picoline has an average acetic acid concentration in the bulk solution (Figure 43) but an unusually high concentration of acetic acid in the condensed liquid (Figure 44). There might
be a connection between the unusual condensation rate – CR correlation and the high acetic acid concentration in the condensed liquid, but the connection cannot be fully explained.

Picoline and aminomorpholine neutralise the bulk solution from pH 3 to a pH of 3.8 and pH 4.4, respectively (Figure 41). There is no impact on the pH of the condensed liquid at the TOL where both remain around pH 5 throughout the test (Figure 42).

Octylamine, pyridazine and pyridine are the next three VCIs discussed more detail. The pre-corrosion by means of LPR for octylamine and pyridazine is, as expected, very similar to the blank test. Pyridine starts with a lower CR compared to all the other VCIs due to a slightly lower acetic acid concentration in the bulk solution and lower condensation rate at the beginning of the test (Figure 40 and Figure 43). Nevertheless, as soon the VCI was inserted, the CR by means of LPR for octylamine decreases in a step and levels out at 0.60 mm/y. The CR for both pyridazine and pyridine continues to decrease slowly until it reaches a CR by means of LPR of 0.46 mm/y.

The CR by means of IC of all three tests seems to be unaffected by the injection of the VCI. The CRs fluctuate around the blank CR throughout the entire test period and seem to be more affected by the condensation rate than the VCIs. This can also be seen in the IE values where octylamine and pyridine display a very poor performance by means of IC (-2% and 2% IE).

The pH adjustment of the bulk solution is very different for all three VCIs. Octylamine increases the pH from pH 2.8 to pH 4.2, pyridine increases the pH just to 3.5, and pyridazine has virtually no effect on the pH in the bulk solution. Interestingly, the progression of the pH in the condensed liquid is very similar to the pH of the condensed liquid of the blank test for all three tests.

The final set of VCIs discussed are aminomorpholine and dicyclohexylamine which are also the better VCIs within this low performing VCI group.
Dicyclohexylamine starts equivalent to the blank test during the pre-corrosion as seen in Figure 38. As soon as the VCI is inserted after 48 hours, the CR decreases to 0.61 mm/y and continues the downward trend until the end where a CR of 0.43 mm/y is reached. The CR of aminoethylpiperazine also ends up at a very similar value (0.39 mm/y), but in this case, the CR drops in one step (as soon the VCI was inserted), going from 0.70 mm/y to 0.42 mm/y, and staying essentially constant until the end of the test.

The CR chart by means of IC is again very responsive to a change in the condensation rate. This can be seen especially well in the curve of aminoethylpiperazine after 48 hours, but also throughout the entire graph. Nevertheless, as soon the VCIs were inserted into the system, both graphs stay well below the blank curve.

The very good performance of aminoethylpiperazine here might be explained by its extraordinary ability of neutralisation. The pH of the bulk solution was raised from 2.9 to 5.1, which in turn raised the pH of the condensed liquid to an uncommonly high level in the range from pH 5.2 to pH 5.8. The acetic acid concentration in the bulk solution is average compared to most other tests, but the acetic acid in the condensed liquid is much lower. This is probably due to the high pH in the bulk solution dissociating more of the acetic acid into acetate which is not volatile and therefore can’t alter the pH in the condensed liquid of the TOL. In the case of aminoethylpiperazine, the neutralisation of the bulk solution might have played a major role in the inhibition of TOL.

Dicyclohexylamine does not have the same neutralisation ability as aminoethylpiperazine, and only raises the pH in the bulk solution from pH 2.8 to pH 4.0. Therefore, the effect on the pH of the condensed liquid is not as strong, but it still maintains a higher pH than the blank test.

The acetic acid concentration in the bulk solution for all VCIs in this chapter is very consistent, except for aminomorpholine which is slightly higher than all others. Also, the acetic acid concentration in the condensed liquid is very consistent for most of the tests. Aminoethylpiperazine is an exception with a very low acetic acid concentration in the condensed liquid. Picoline and
aminomorpholine both show a high concentration of acetic acid in the condensed liquid. The acetic acid concentration of all other tests is between 650 ppm and 800 ppm in the condensed liquid at the end of the test.

Table 13: Corrosion rates and Inhibition efficiency by LPR, weight loss, and iron count from the CFP test

<table>
<thead>
<tr>
<th></th>
<th>CR [mm/y]</th>
<th>IE [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LPR  IC  WL</td>
<td>LPR  IC  WL</td>
</tr>
<tr>
<td>Aminoethylpiperazine</td>
<td>0.40 0.27 1.30</td>
<td>49 32 34</td>
</tr>
<tr>
<td>Aminomorpholine</td>
<td>0.95 0.35 1.89</td>
<td>-22 13 4</td>
</tr>
<tr>
<td>Dicyclohexylamine</td>
<td>0.49 0.34 1.18</td>
<td>37 14 40</td>
</tr>
<tr>
<td>Octylamine</td>
<td>0.59 0.41 1.69</td>
<td>24 -2 14</td>
</tr>
<tr>
<td>Picoline</td>
<td>1.00 0.56 1.95</td>
<td>-29 -39 1</td>
</tr>
<tr>
<td>Picoline</td>
<td>0.50 0.31 1.42</td>
<td>36 22 28</td>
</tr>
<tr>
<td>Pyridazine</td>
<td>0.46 0.39 1.40</td>
<td>41 2 29</td>
</tr>
<tr>
<td>Pyridine</td>
<td>0.78 0.40 1.98</td>
<td>- - -</td>
</tr>
<tr>
<td>Blank</td>
<td>0.78 0.40 1.98</td>
<td>- - -</td>
</tr>
</tbody>
</table>

Figure 38: Corrosion rates by LPR from the CFP test
Figure 39: Corrosion rates by iron count from the CFP test

Figure 40: Condensation rate from the CFP test
Figure 41: Bulk pH from the CFP test

Figure 42: Condensed liquid pH from the CFP test
**Figure 43:** Acetic acid concentration in the bulk solution

**Figure 44:** Acetic acid concentration in the condensed liquid
3.3.2.2.1. **Conclusion Cooled Finger Probe**

In this test, it was shown again that most of the compounds have a strong neutralising effect on the bulk solution. This alone is, in most cases, not sufficient to inhibit TOL corrosion, but in the case of aminoethylpiperazine it plays a major role due to the extent of neutralisation.

The CR by means of IC is very closely related to the corrosivity of the condensed liquid measured by means of the LPR. There is a very strong response of CR by means of IC to the condensation rate visible in most of the samples.

The best performing VCI of this group in the CFP test is aminoethylpiperazine. The comparatively good performance of aminoethylpiperazine is can be explained with the high pH in the bulk solution and condensed liquid with a resulting low concentration of free acetic acid. Nevertheless, the overall performance of all compounds is not satisfactory with none of the IE above 50%.
3.3.2.3. **Altered Horizontal Cooled Tube**

Most of the VCIs of this section were also tested in the A-HCT test. The difference to the normal HCT test is that the condensed liquid is collected in a beaker in order to keep it off the bulk solution. In theory, the VCI should be kept away from the bulk solution to keep the acetic acid fully dissociated, only testing the effect of the VCI directly on the TOL. Unfortunately, due to the handling, mainly the purging, and the pressurising with CO2 into the autoclave, the separation was not achieved in all tests. It is assumed that if the separation was properly achieved, the pH of the bulk solution should be reasonably low (which is the case in some tests). Some tests though, have a comparatively high pH in the bulk solution after 24 hours. Therefore it is assumed that during the pressurising (due to the bubbling) some of the bulk solution flooded the beaker with VCI, partly mixing it into the bulk solution.

To start the discussion, the VCIs resulting in the highest and lowest CRs will be considered first; pyridazine and aminoethylpiperazine. Both VCIs seem to be the odd ones out therefore they are discussed together. The CRs of pyridazine and all other tested VCIs can be seen in Figure 45. The CR and condensation rate is very close to the respective blank rate throughout the entire test (Figure 46). Also, the bulk pH and condensed pH are very similar to the respective blank pH (Figure 47, Figure 48). Pyridazine has essentially no effect on any parameter of this test. This can also be seen in the IE by means of IC and WL; exactly 2% for both.

Aminoethylpiperazine, on the other hand, is seemingly the best performing VCI in this test despite a comparatively high condensation rate. Unfortunately, due to the very high neutralisation ability of aminoethylpiperazine, a small volume is enough to elevate the bulk pH and neutralizes the acetic acid into acetate. The bulk pH was already at 5.6 after 24 hours, and it is strongly assumed that it happened due to the previously explained problem with the pressurising. Therefore, the results of aminoethylpiperazine from this test have to be excluded from the “rating” and cannot be compared to the blank results.
The four other VCIs display very comparable corrosion inhibition behaviours, at least by means of IC. Picoline and pyridine neutralize the bulk solution in the same way. Both start after 24 hours at pH 4.6 and end up at pH 4.3. These are the only bulk solutions which become more acidic over time. The CR of picoline stays constant around 0.8 mm/y whereas the CR of pyridine stays just below 1.00 mm/y. The average CRs and IEs can be seen in Table 14 where it becomes obvious that both VCIs display reasonably good IE by means of IC of 40-50 %. The IE by means of WL, on the other hand, does not match the IC measurements of both VCIs. The pH of the condensed liquid is unaffected by both VCIs.

Dicyclohexylamine and octylamine are the last VCIs to be discussed in this group. The CR of the test containing octylamine stays around 0.80 mm/y throughout the first three days and increases slightly to 0.88 mm/y on the last day of the test. The pH in the bulk solution started just below pH 4 but did rise up to pH 4.7 after 90 hours. The pH of the condensed liquid was not affected by octylamine and stays at the same level as the blank test over the 90 hours test period. As with the picoline and pyridine, the IE by means of IC and WL do not match up. Dicyclohexylamine was the overall best performing VCI in this test group. Despite an increasing condensation rate after 48 hours, the CR stayed constant and even dropped slightly at the end of the test from a constant 0.75 mm/y to 0.64 mm/y. The pH in the bulk solution started as the second lowest of the tested VCIs with pH 3.7 and rose up to pH 4.5 over time. The pH in the condensed liquid started on day 1 around pH 6 (as the blank test did), but then dropped over time down to pH 4.8. This might be the result of an increasing condensation rate and a constant or slightly declining CR; more condensed water with less corrosion product will result in a lower pH. The IE by means of IC and WL is 55 % and 56 %, respectively, which are the highest IEs within the tested VCIs in this test. All results of dicyclohexylamine combined (higher condensation rate – constant CR – declining pH in the condensed liquid) indicate that dicyclohexylamine inhibited the sample by forming a film directly on the surface.
Table 14: Average corrosion rates and inhibition efficiencies by iron concentration and weight loss from the A-HCT test

<table>
<thead>
<tr>
<th></th>
<th>CR [mm/y]</th>
<th>IE [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IC</td>
<td>WL</td>
</tr>
<tr>
<td><strong>Aminoethylpiperazine</strong></td>
<td>0.47</td>
<td>0.54</td>
</tr>
<tr>
<td><strong>Dicyclohexylamine</strong></td>
<td>0.72</td>
<td>0.57</td>
</tr>
<tr>
<td><strong>Octylamine</strong></td>
<td>0.80</td>
<td>1.46</td>
</tr>
<tr>
<td><strong>Picoline</strong></td>
<td>0.81</td>
<td>1.29</td>
</tr>
<tr>
<td><strong>Pyridazine</strong></td>
<td>1.58</td>
<td>1.28</td>
</tr>
<tr>
<td><strong>Pyridine</strong></td>
<td>0.95</td>
<td>1.03</td>
</tr>
<tr>
<td><strong>Blank</strong></td>
<td>1.61</td>
<td>1.31</td>
</tr>
</tbody>
</table>

Figure 45: Corrosion rate by IC from the A-HCT test
Figure 46: Condensation rates from the A-HCT test

Figure 47: Bulk pH from the A-HCT test
3.3.2.3.1. Conclusion Altered Horizontal Cooled Tube

With this test set-up, the element of neutralisation should have been eliminated. Due to the difficulties in the preparation and handling of the test, it was not entirely achieved for all VCIs.

Aminoethylpiperazine was taken out of the equation in this test. Most of the other VCIs show significant discrepancies between the average CR by means of IC and WL. Nevertheless, dicyclohexylamine as a VCI works very well in this set-up. The starting pH was reasonably low and the CR stays constant despite the rising condensation rate as more and more dicyclohexylamine reached the TOL sample through the vapour phase.

It also proves the concept of the test to be a viable set-up to investigate the volatility and effectiveness of VCIs.

Figure 48: Condensed liquid pH from the A-HCT test
3.3.2.4. **Rotating Cylinder Electrode Testing**

The rotating cylinder electrode test set up was used with alternating LPR and EIS tests, with a potentiodynamic scan at the end of the testing period. For all tests, pre corrosion was allowed for approximately 5 hours. All tests displayed in Figure 52 were treated additionally with HCl for the purpose of eliminating the neutralizing effect of the VCI. All tests were performed with 1000 ppm acetic acid and 200 rpm rotation speed. In this section, all VCIs are discussed separately in respect to their performance in the different RCE tests.

In general, the CRs calculated by LPR (Figure 49; Figure 52) follow the same trend as the CRs calculated by EIS (Figure 50; Figure 53) with and without HCl. The big difference in the results of LPR and EIS in these tests is the actual values of the pre-corrosion. After the VCIs were injected into the system, the calculated CRs of both are quite similar to each, compared to the values before the VCI was added. This happens mainly because without VCI in the system the ratio of R\textsubscript{S} to R\textsubscript{CT} is too large. The value for R\textsubscript{S} in the given test solution is relatively high and the R\textsubscript{CT} for the bare carbon steel is relatively low. In that case, LPR detects a high resistance (R\textsubscript{F} + R\textsubscript{CT}) resulting in a lower calculated CR associated with a low level of reliability of the LPR measurements.

After a working VCI was injected, R\textsubscript{CT} increases significantly. The better the inhibition, the more R\textsubscript{CT} increases. R\textsubscript{S} decreases in most cases due to a better conductivity of the solution. Therefore, the ratio of R\textsubscript{S} to R\textsubscript{CT} decreases significantly, and so does the discrepancy in the CR calculated by LPR, bringing EIS and LPR results closer together. It can be seen in Table 15 that once the IE was calculated, the values by LPR and EIS are very comparable.

Aminoethylpiperazine was the amine with the highest neutralisation ability at the TOL. In these tests as well, aminoethylpiperazine raised the pH higher than the other VCIs. At the beginning of all tests, the pH is around 3.15; with aminoethylpiperazine inserted the pH climbed around 5.2. Nevertheless, an inhibition effect could not be achieved with this compound. Straight after injection, the CR increased constantly over the next 20 hours to over
2.00 mm/y as seen in Figure 49 and Figure 50. When HCl was added to decrease the pH back to a pre VCI value, the CR calculated by LPR dropped briefly from just above 1.00 mm/y to 0.65 mm/y but then came back up to the blank CR. It seems that the negative effect seen without HCl was gone, but no inhibition effect was observed. The CR calculated by means of EIS did show a much more positive evolution with an IE of 64 %.

All other tested VCIs exhibited a very different behaviour compared to aminoethylpiperazine in all tests. Dicyclohexylamine, octylamine, and pyridine have an IE by LPR of 86 %, 56 %, and 89 %, respectively if neutralization of the solution is permitted (Table 15; Figure 49). The IE calculated by LPR was confirmed by EIS where 93 %, 78 %, and 93 % was calculated for dicyclohexylamine, octylamine, and pyridine, respectively.

In the potentiodynamic scans in Figure 51 it can be seen that all of VCIs (except Octylamine) shift the corrosion potential to a more noble value. Octylamine did not affect the corrosion potential at all. It can also be observed that all other VCIs (except Pyridazine) lower the cathodic part of the scan and leave the anodic side virtually unaffected. Pyridazine displays in the potentiodynamic scans the biggest shift to a more positive corrosion potential and Pyridazine decreased the corrosion current to the lowest value of all VCI compounds by lowering the anodic and cathodic parts of the potentiodynamic curve.

In Figure 52 and Figure 53 it can be seen that the compounds still display a comparatively good inhibition in the presence of HCl. In the order, pyridine, octylamine to dicyclohexylamine, the IE calculated by LPR increases from 25 % to 41 % to 55 %. The IE’s by EIS are even higher with 56 %, 63 %, and 77 % for the same order of VCIs. The IEs of the tests with HCl are all lower compared to the tests without HCl, but a good part of inhibition remains. The potentiodynamic scans, including HCl, also show that the anodic reaction is unaffected and the cathodic corrosion current lowered.

The last amine to be discussed in this section is also the best performing amine in presence of HCl in the RCE test set up. Pyridazine decreased the CR by LPR to 0.10 mm/y (0.24 mm/y by EIS) and therefore achieved an IE of
93 % by LPR and 90 % by EIS without the addition of HCl. The pH increased to 3.75 after pyridazine was inserted, a very minor change compared to the other amines. Pyridazine is the only compound that significantly changes the shape of the potentiodynamic curve; the corrosion potential is shifted from -0.67 V to -0.52 V, the cathodic corrosion current is lower than the blank and the anodic side is also slightly lower than the blank.

The addition of HCl affects the performance of pyridazine and lowers the IE calculated by LPR and EIS to 78 % and 84 %, respectively. It is the best IE after the addition of HCl and the appearance of the potentiodynamic scan is virtually unaffected by HCl. The corrosion potential is still located at -0.52 V (Figure 54). Pyridazine displays superior BOL inhibition properties (in respect of the molecule size) as tested with the RCE test set up.

Table 15: Inhibition efficiencies by LPR and EIS from the RCE test without and with (+HCl) pH adjustment

<table>
<thead>
<tr>
<th></th>
<th>IE by LPR (Figure 49)</th>
<th>IE by EIS (Figure 50)</th>
<th>IE by LPR + HCl (Figure 52)</th>
<th>IE by EIS + HCl (Figure 53)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aminoethyl-piperazine</td>
<td>--</td>
<td>-</td>
<td>10</td>
<td>64</td>
</tr>
<tr>
<td>Dicyclohexylamine</td>
<td>86</td>
<td>93</td>
<td>55</td>
<td>77</td>
</tr>
<tr>
<td>Octylamine</td>
<td>56</td>
<td>78</td>
<td>41</td>
<td>63</td>
</tr>
<tr>
<td>Picoline</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Pyridazine</td>
<td>86</td>
<td>90</td>
<td>78</td>
<td>84</td>
</tr>
<tr>
<td>Pyridine</td>
<td>89</td>
<td>93</td>
<td>25</td>
<td>56</td>
</tr>
</tbody>
</table>
Figure 49: Corrosion rates by means of LPR from the RCE test (without pH adjustment)

Figure 50: Corrosion rates by means of EIS from the RCE test (without pH adjustment)
Figure 51: Potentiodynamic scans from the RCE test (without pH adjustment)

Figure 52: Corrosion rates by means of LPR from the RCE test (pH adjusted +HCl)
Figure 53: Corrosion rates by means of EIS from the RCE test (pH adjusted +HCl)

Figure 54: Potentiodynamic scans from the RCE test (pH adjusted +HCl)
3.3.2.4.1. **Conclusions from the Rotating Cylinder Test**

Much information was gathered by means of the RCE tests. Excluding aminoethylpiperazine, all the tested VCIs inhibited the corrosion in this test. All of the other VCIs displayed some kind of alternative inhibition in addition to the neutralisation. It can be assumed that a protective film was formed on the surface of the corrosion sample.

Pyridazine, as one of the worst performers in the actual TOL tests, displayed a superior performance as film forming BOL corrosion inhibitor.

3.3.2.5. **Discussion and Conclusion “Low performing VCIs”**

None of the tested VCIs was a satisfactory TOL corrosion inhibitor by itself. All of the tested VCIs except of Pyridazine display a high ability to increase the pH of an acidic solution. Due to the overall lack of satisfactory TOL inhibition, it can also be concluded that pH neutralisation is not the predominant inhibition mechanism in TOL corrosion inhibition by VCIs.

RCE testing showed that all of the tested VCI compounds have a good BOL corrosion inhibition ability. Even after the solution was neutralized using HCl, much of the inhibition ability was still remaining indicating that a large portion of inhibition is by film forming rather than neutralisation.

In general, it can be concluded that the poor TOL inhibition in the tested conditions is a result of the high boiling point and rather low vapour pressure of all low performing VCIs.

3.3.2.5.1. **Aminoethylpiperazine**

Aminoethylpiperazine is an excellent pH neutralizer. It significantly increased the bulk pH in all TOL tests as well as in the RCE tests. It also displayed, in comparison to the other VCIs tested in this section, good TOL inhibition properties. Nevertheless, the CR by means of RCE doubled to 2 mm/y after the VCI was inserted into the system. Therefore, the compound cannot be used in the tested environments and concentrations.
3.3.2.5.2. *Aminomorpholine*

Aminomorpholine was the most expensive compound and only a very limited amount was available for testing. Therefore, testing was stopped after it failed to show TOL corrosion inhibition properties in the CFP.

3.3.2.5.3. *Aniline*

Aniline was only tested in the HCT test and therefore the conclusions are very limited. Within the tested VCIs in the HCT, it was an average performer.

3.3.2.5.4. *Dicyclohexylamine*

Dicyclohexylamine isn’t volatile enough in the given conditions to have a good effect on TOL corrosion. Despite a good inhibition in the A-HCT test, it didn’t inhibit as well in the other TOL corrosion tests. An IE of up to 93 % was achieved in the RCE test. Dicyclohexylamine seems to be a film forming and neutralising amine that is not volatile enough to be part of a successful VCI formulation that inhibits TOL corrosion.

3.3.2.5.5. *Octylamine*

The vapour pressure and volatility of octylamine is also not sufficient to be effective at the TOL. The BOL inhibition, in comparison to the other VCIs tested in this section is also underwhelming.

3.3.2.5.6. *Picoline*

Picoline has a negative effect on TOL corrosion. In the HCT and CFP test the CR was increased with the injection of picoline. Therefore, it was not further investigated on its BOL corrosion inhibition properties.
3.3.2.5.7. *Pyridazine*

Pyridazine is the only tested compound that does not have a significant neutralisation effect on the solution. It does not inhibit in either the HCT or A-HCT set up, but shows average TOL inhibition properties in the CFP. It was the best performing compound (including all amines, meaning also the high performing amines discussed later) in the RCE test in the presence of HCl. The IE was still between 78– 84 % with HCl and up to 90 % without HCl. It seems to be mainly a film forming inhibitor, limited to the BOL. It is assumed that the two N atoms stick to the metal surface separating it from the corrosion media. Further investigation might reveal possible fields of application for a small molecular, film forming amine. Unfortunately, this is outside the scope of this research.

3.3.2.5.8. *Pyridine*

The TOL inhibition of pyridine was not satisfying, but Pyridine showed decent BOL inhibition properties without HCl in the system. An IE of 93 % was achieved. The large part of the BOL inhibition properties was lost with HCl being included in the testing indicating the inhibition is mainly achieved by neutralisation.
3.3.3. High Performing VCIs - Results

In the following section, 8 of the generic VCI compounds will be discussed. These 8 compounds were the compounds with the best performance in regards of TOL corrosion inhibition. The molecular structures (displayed using the free molecule editor software Avogadro 1.0.3) and some of the basic properties of each molecule can be found in Table 16. The properties of the molecules were taken from the respective MSDS sheets accessed via the homepage of the supplier, Sigma Aldrich (MSDS 2011; MSDS 2011; MSDS 2011; MSDS 2011; MSDS 2011; MSDS 2011; MSDS 2011; MSDS 2012).

The molecular structure is displayed as grey, blue, red, and small light grey spheres representing carbon, nitrogen, oxygen, and hydrogen, respectively.

All of the compounds are amines or involve an amino group. Amino groups can be found either as primary, secondary, tertiary, cyclic amino group, or in any combination. Three of the compounds contain oxygen; MDEA with two oxygen atoms, methylmorpholine and morpholine with one oxygen atom each. The molecular weight of all compounds lies between 73.14 and 119.16 g/mol and the boiling points are between 36 and 248 °C.

The grey graph in every diagram represents the blank test, and every VCI compound has its dedicated colour/marker combination throughout the entire “High Performing VCI” discussion. Where necessary, some graphs have been split up into two for clarity.
Table 16: Structures and properties of all high performing VCI compounds

<table>
<thead>
<tr>
<th></th>
<th>Benzyamine</th>
<th>Cyclohexyl-amine</th>
<th>Diethylamine (DEA)</th>
<th>Dimethyl-ethylamine (DMEA)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Formula</strong></td>
<td>C₇H₉N</td>
<td>C₆H₁₃N</td>
<td>C₄H₁₁N</td>
<td>C₄H₁₁N</td>
</tr>
<tr>
<td><strong>Molecular weight [g/mol]</strong></td>
<td>107.15</td>
<td>99.17</td>
<td>73.14</td>
<td>73.14</td>
</tr>
<tr>
<td><strong>Boiling point [°C]</strong></td>
<td>184-185</td>
<td>134</td>
<td>55</td>
<td>36-38</td>
</tr>
<tr>
<td><strong>Vapour pressure [mm Hg]</strong></td>
<td>0.662 ( @ 25 °C)</td>
<td>10 ( @ 22 °C)</td>
<td>418 ( @ 20 °C)</td>
<td>237 ( @ 25 °C)</td>
</tr>
<tr>
<td><strong>Purity [%]</strong></td>
<td>99</td>
<td>99</td>
<td>99</td>
<td>99</td>
</tr>
<tr>
<td><strong>Molecular Structure</strong></td>
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<td><img src="image2" alt="Structure" /></td>
<td><img src="image3" alt="Structure" /></td>
<td><img src="image4" alt="Structure" /></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Methyl-diethanol-amine (MDEA)</th>
<th>Methyl-morpholine</th>
<th>Methyl-piperazine</th>
<th>Morpholine</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Formula</strong></td>
<td>C₅H₁₃NO₂</td>
<td>C₅H₁₁NO</td>
<td>C₅H₁₂N₂</td>
<td>C₄H₉NO</td>
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<tr>
<td><strong>Molecular Weight [g/mol]</strong></td>
<td>119.16</td>
<td>101.15</td>
<td>100.16</td>
<td>87.12</td>
</tr>
<tr>
<td><strong>Boiling point [°C]</strong></td>
<td>246-248</td>
<td>115-116</td>
<td>138</td>
<td>126-130</td>
</tr>
<tr>
<td><strong>Vapour pressure [mm Hg]</strong></td>
<td>&lt;0.01 ( @ 20 °C)</td>
<td>18 ( @ 20 °C)</td>
<td>7.5 ( @ 20 °C)</td>
<td>10.4 ( @ 25 °C)</td>
</tr>
<tr>
<td><strong>Purity [%]</strong></td>
<td>99</td>
<td>99</td>
<td>99</td>
<td>99</td>
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<tr>
<td><strong>Molecular Structure</strong></td>
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<td><img src="image6" alt="Structure" /></td>
<td><img src="image7" alt="Structure" /></td>
<td><img src="image8" alt="Structure" /></td>
</tr>
</tbody>
</table>
3.3.3.1. **Horizontal Cooled Tube**

In this section, the performance of the high performing VCIs is discussed. In Figure 55 and Figure 56 the CRs by means of IC are displayed and Table 17 lists the average CRs by means of IC and WL as well as the IEs of all VCIs.

During the first 24 hours without VCI in the system, all tests averaged a CR of around 1.50 mm/y. As soon as the VCIs were injected into the test, the CRs drop to approximately 0.60 mm/y. The CRs kept declining until day three (72 hours) in all tests.

Benzylamine stays constantly just below 0.40 mm/y after 72 hours until the end of the test. A very similar behaviour can be observed for the CR of cyclohexylamine, DEA, DMEA and methylpiperazine. They all levelled around 0.40 mm/y until the end of the testing period.

MDEA, methylmorpholine and morpholine display a slightly different behaviour. The CR of all three tests continues to decline slightly until the end of the test.

Figure 57 displays the pH evolution of each test. All tested compounds alter the pH of the bulk solution in a very similar fashion. The pH at the very beginning of each test is 3.2. The pH of the test straight after the VCIs were injected is in a range of 4.9 to 5.3 which is approximately 1.5 pH units higher than the blank pH. At the end of the test, the pH ranges between 5.4 and 5.7 (blank pH 5.0). The pH in the blank test increased 1.5 pH units from day two to day seven, whereas the VCI tests increased about 0.5 pH units during the same time period.
Table 17: Corrosion rates and inhibition efficiencies by iron concentration and weight loss from the HCT test

<table>
<thead>
<tr>
<th></th>
<th>CR [mm/y]</th>
<th>IE [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IC</td>
<td>WL</td>
</tr>
<tr>
<td>Benzylamine</td>
<td>0.45</td>
<td>0.62</td>
</tr>
<tr>
<td>Cyclohexylamine</td>
<td>0.44</td>
<td>0.65</td>
</tr>
<tr>
<td>DEA</td>
<td>0.42</td>
<td>0.58</td>
</tr>
<tr>
<td>DMEA</td>
<td>0.40</td>
<td>0.59</td>
</tr>
<tr>
<td>MDEA</td>
<td>0.47</td>
<td>0.64</td>
</tr>
<tr>
<td>Methylmorpholine</td>
<td>0.39</td>
<td>0.66</td>
</tr>
<tr>
<td>Methylpiperazine</td>
<td>0.40</td>
<td>0.65</td>
</tr>
<tr>
<td>Morpholine</td>
<td>0.45</td>
<td>0.63</td>
</tr>
<tr>
<td>Blank</td>
<td>0.71</td>
<td>0.84</td>
</tr>
</tbody>
</table>

Figure 55: Corrosion rates from the HCT test (High performing VCIs Part 1)
3.3.3.1.1. **Conclusion Horizontal Cooled Tube**

All VCIs discussed here have potent pH neutralisation abilities, and despite the variety of compounds, it was not possible to really distinguish them by
IE. All of the compounds inhibit the simulated TOL corrosion by means of IC in the HCT test by 35–46%. Tests using other apparatuses are described in the following sections to help identify mechanisms of inhibition of the compounds.

3.3.3.2. Cooled Finger Probe

Using the CFP test, it is possible to study the VCIs in more detail because a wider variety of results are obtained. The discussion of the VCIs will be divided into two sections for clarity reasons in the graphs: first benzylamine, cyclohexylamine, DEA and DMEA and then further down MDEA, methylmorpholine, methylpiperazine and morpholine. An overview of the results can be found in Table 18.

<table>
<thead>
<tr>
<th></th>
<th>CR [mm/y]</th>
<th>IE [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LPR</td>
<td>IC</td>
</tr>
<tr>
<td>Benzylamine</td>
<td>0.40</td>
<td>0.27</td>
</tr>
<tr>
<td>Cyclohexylamine</td>
<td>0.46</td>
<td>0.28</td>
</tr>
<tr>
<td>DEA</td>
<td>0.41</td>
<td>0.21</td>
</tr>
<tr>
<td>DMEA</td>
<td>0.50</td>
<td>0.31</td>
</tr>
<tr>
<td>MDEA</td>
<td>0.50</td>
<td>0.25</td>
</tr>
<tr>
<td>Methylmorpholine</td>
<td>0.49</td>
<td>0.31</td>
</tr>
<tr>
<td>Methylpiperazine</td>
<td>0.45</td>
<td>0.37</td>
</tr>
<tr>
<td>Morpholine</td>
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<td>0.28</td>
</tr>
<tr>
<td>Blank</td>
<td>0.78</td>
<td>0.40</td>
</tr>
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</table>

Benzylamine was the best performing VCI by LPR in the CFP test. The average CR of the pre corrosion period was 1 mm/y which dropped down to 0.5 mm/y the day after the VCI was injected and declined further to 0.38 mm/y the next two days and ended up at 0.33 mm/y (Figure 58). A similarly good performance was seen with the CR by IC where it decreases from a pre-corrosion average of 0.45 mm/y to a CR of 0.27 mm/y (Figure 59). The bulk pH did rise from pH 2.8 to pH 4.4 with benzylamine injected (Figure 61) and the condensate pH was 0.4 units higher than the blank.
during the first 24 hours of injection, but it stayed close to the blank pH during the rest of the test with a maximum difference of 0.2 units (Figure 62). The acetate concentration in the bulk solution slightly increases over time from 1160 ppm on day two to 1330 ppm on day four and stayed constant afterwards (Figure 63). By using Equation 9 it is possible to calculate the acetic acid concentration of this test. There is 933 ppm acetic acid in the bulk solution after 96 hours (blank test 872 ppm) and still 940 ppm of acetic acid at the very end of the test (blank 1100 ppm). So, despite the elevated pH, the acetic acid concentration with VCI is very comparable to the acetic acid concentration of blank test. This means the bulk neutralisation effect is eliminated here. The acetate concentration increases slightly because a 1000 ppm acetic acid concentration solution is used to top up sampled liquid and accumulates acetates. Due to the VCI, the acetate concentration in the condensed liquid decreased from 800 ppm during the first two days down to 500 ppm over the next two days (Figure 64). Acetic acid was constantly topped up and therefore the concentration in the condensed liquid increased back up to 650 ppm. Despite the increase, the pH and CR was virtually unaffected which gives a positive indication for benzylamine as a VCI.

Cyclohexylamine and DEA exhibit almost the same characteristics as described for benzylamine. The CR by LPR of the test with cyclohexylamine drops from 0.80 mm/y down to 0.52 mm/y and keeps decreasing to 0.41 mm/y at the end of the test (Figure 58). The CR by means of IC also decreases over time from 0.44 mm/y during pre-corrosion down to 0.31 mm/y to a final 0.27 mm/y (Figure 59) with a very constant condensation rate (Figure 60). The CR by LPR of DEA falls from 0.78 mm/y the second day of pre-corrosion to 0.42 mm/y the first day of VCI in the system and keeps decreasing to 0.35 mm/y at the end of the test. DEA changes the CR by means of IC to the lowest CR of all tested VCIs in this system. The CR drops from 0.38 mm/y to 0.22 mm/y down to a final 0.19 mm/y. The pH, acetate and acetic acid concentration evolution is almost exactly the same as described before for benzylamine and will not be described again.
The last VCI in this group of four is DMEA. Within this group and test, DMEA is the least effective performing VCI, despite the comparatively low acetic acid concentration (Figure 63). After the VCI was injected, the CR by LPR decreased to 0.57 mm/y and kept decreasing to 0.44 mm/y at the end of the test. The CR by means of IC levelled around 0.31 mm/y after the VCI was injected with an average condensation rate. As mentioned, the acetic acid concentration of the bulk solution and therefore also in the condensed liquid was lower than the blank test. Despite the slightly less aggressive conditions cyclohexylamine and DEA did not perform as well as benzylamine.

Figure 58: Corrosion rates by LPR from the CFP test – Part 1
Figure 59: Corrosion rates by iron concentration from the CFP test – Part 1

Figure 60: Condensation rates from the CFP test – Part 1
Figure 61: pH of the bulk solution from the CFP test – Part 1

Figure 62: pH of the condensed liquid from the CFP test – Part 1
Figure 63: Acetic acid concentration of the bulk solution from the CFP test – Part 1

Figure 64: Acetic acid concentration in the condensed liquid from the CFP test – Part 1
The second set of four VCIs includes MDEA, methylmorpholine, methylpiperazine and morpholine. The evolution of the CR by means of LPR for all four VCIs is nearly identical. During the pre-corrosion the CR is around 1.00 mm/y and it falls to 0.6 mm/y, 0.55 mm/y, 0.47 mm/y and 0.43 mm/y for MDEA, methylmorpholine, methylpiperazine and morpholine, respectively. The CR for all four VCIs falls further over the next three days between 0.43 mm/y and 0.50 mm/y (Figure 65).

The CRs by means of IC are more separated on first appearance (Figure 66). Methylpiperazine stands out with a higher CR than the other VCIs. By comparing the condensation rates of the VCIs, it becomes clear that the condensation rate of methylpiperazine is higher than all others (Figure 67). Also seen in all other tests before, the CR is affected very much by the condensation rate. Despite the higher condensation rate, methylpiperazine still keeps the CR below the blank CR. A similar effect can be seen by methylmorpholine where the condensation rate briefly increases after 96 and 120 hours. With the increase in condensation rate, the CR increases, too. The CR by IC for MDEA, methylmorpholine and morpholine decreases from around 0.40 mm/y pre corrosion to 0.27 mm/y corrosion with VCI. At the end of the test, all three show a CR around 0.27 mm/y (Figure 66). The acetate concentrations are higher than the blank and therefore, the acetic acid concentrations are again (as with the first set of four VCIs) very comparable with the blank test. The pH of the bulk solution is between 4.3 and 4.6 the day of VCI injection and falls to between pH 4.1 and 4.4 until the end of the test (Figure 68). The pH of all four VCIs in the condensed liquid the day after injection is also very similar and lies between pH 5.3 and 5.4. It falls to pH 5.0 at the end of the test with MDEA, methylmorpholine and morpholine (Figure 69). The pH of methylpiperazine falls to pH 4.77 a day before the end and stays constant on the last day. This might be in correlation with the increase of condensation rate during the last two days.

The average CRs and IEs in Table 18 on page 103 also indicates how similar the tested VCIs in this section are especially in regards of the CR and IE by means of LPR.
Figure 65: Corrosion rates by LPR from the CFP test – Part 2

Figure 66: Corrosion rates by iron concentration from the CFP test – Part 2
Figure 67: Condensation rates from the CFP test – Part 2

Figure 68: pH of the bulk solution from the CFP test – Part 2
Figure 69: pH of the condensed liquid from the CFP test – Part 2

Figure 70: Acetic acid concentration of the bulk solution from the CFP test – Part 2
3.3.3.2.1. **Conclusion Cooled Finger Probe**

All of the “high performing” compounds tested in the CFP display very similar properties to each other. For example, the pH of the bulk solution of each of the tested compounds the day of injection is between pH 4.3 and pH 4.6 and at the end of the test, the pH is between pH 4.1 and pH 4.5. The pH of the condensed liquid (except for DMEA) also lies in a very tight range, between pH 4.8 and pH 5.1 at the end of the test. More importantly, the evolution of the CRs and the resulting IEs are very similar, too. It can be concluded that all of the tested VCIs in this section work well in inhibiting TOL corrosion in the given conditions provided by the CFP test.

3.3.3.3. **Altered Horizontal Cooled Tube**

In this section, the performance of the “high performing VCIs” in the A-HCT will be discussed. The average CRs and IEs are listed in Table 19 and it is obvious that the performance of the VCIs show more variation than with the CFP test in the section before. Therefore, the results of the “high performing VCIs” are displayed in one graph for each parameter: CRs, condensation
rates, bulk pH, and condensed pH (Figure 72, Figure 73, Figure 74, and Figure 75).

The bulk pH is an important indication in these results. It should start low which would ensure that the VCI and the bulk solution have good separation. The tests discussed in this section have a low starting pH except for methylmorpholine and DEA. Both of those VCI tests start up with a bulk pH of over pH 4 and therefore the results need to be evaluated with care and it might be necessary to exclude the results from the following discussions (Figure 74).

The average CR of benzylamine by means of IC and WL is 1.14 mm/y and 1.12 mm/y, respectively resulting in an IE of 29 % and 14 %. The CR starts at 1.30 mm/y and declines on day two to 0.80 mm/y with the condensation rate staying constant around 0.44 g/m²/s. On day three, the CR increases back up to 1.25 mm/y and then just increases slightly to 1.32 mm/y on day four as the condensation rate also increases up to 0.60 g/m²/s on day three and four. The condensed pH stays around pH 6.0 throughout the test. The high condensed pH is very similar to the blank version and is mainly due to the reasonably high corrosion rate. Benzylamine is not a suitable VCI in this test set up.

Cyclohexylamine displays just slightly better inhibition than benzylamine. The CR declines from 1.35 mm/y on day one to 1.01 mm/y on day three and increases again to 1.25 mm/y during the last 24 hours (Figure 72). A slightly increase in condensation rate is also observed in this test from 0.33 g/m²/s to 0.49 g/m²/s over the four days of testing. The effect on the pH of the condensed liquid is also limited, and the pH stays constantly around 6.0.

As mentioned in the beginning of this section, the bulk pH of DEA starts off high. Additionally, the condensation rate is comparatively low (0.32 g/m²/s-0.43 g/m²/s) which results inevitably in a low CR. The same can be said about methylmorpholine. It has a high pH in the bulk solution at the beginning and the condensation rate is even lower. Both results will not be in the assessment of the performance of these VCIs.
DMEA shows a very good IE by means of IC and WL of about 64 % and 58 %, respectively (Table 19). The CR starts at 0.72 mm/y and decreases to a final 0.49 mm/y on day four. The condensation rate starts low around 0.32 g/m²/s but doubles over the next days to 0.65 g/m²/s but the CR stays low. The pH in the bulk solution starts at a very low pH of 3.4 and increases to pH 5.6 over the test duration. The huge increase indicates that DMEA has comparatively high volatility. The pH of the condensed liquid starts slightly lower than the pH of the other VCIs at pH 5.7 and decreases to pH 4.8 at the end of the test. All results of this test together indicate very good inhibition behaviour of DMEA.

MDEA also displays very good IEs by means of IC and WL of 56 % and 60 %, respectively. With a rising condensation rate and a consistently low bulk pH, the CR decreases from 1.1 mm/y in the beginning of the test to 0.39 mm/y after three days. As the condensation rate keeps increasing, the CR also increases to a final of 0.70 mm/y.

Methylpiperazine has the highest IE by means of IC in the A-HCT test of all tested VCIs with in IE of 65 % (excluding DEA and methylmorpholine for the high pH and low condensation rates). The CR by IC declines during the test from 0.78 mm/y to 0.34 mm/y by the end. The condensation rate increases from 0.33 g/m²/s to 0.50 g/m²/s during the test and is comparatively low after four days. The pH of the bulk solution increases from pH 3.9 to pH 5.3 and the pH in the condensed liquid decrease from pH 6.2 to pH 5.3.

The pH of the bulk solution and condensed liquid of morpholine is essentially the same as the just described methylpiperazine. The condensation rate in the case of morpholine was higher than in any of the other VCIs tests. An IE of 49 % and 42 % for IC and WL, respectively, is a very good result given the slightly harsher circumstances. The CR after day two is constant around 0.70 mm/y.
Table 19: Average corrosion rates and inhibition efficiencies by iron concentration and weight loss from the A-HCT test

<table>
<thead>
<tr>
<th>Compound</th>
<th>CR [mm/y]</th>
<th>IE [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IC</td>
<td>WL</td>
</tr>
<tr>
<td>Benzylamine</td>
<td>1.14</td>
<td>1.12</td>
</tr>
<tr>
<td>Cyclohexylamine</td>
<td>1.16</td>
<td>1.00</td>
</tr>
<tr>
<td>DEA</td>
<td>0.47</td>
<td>0.54</td>
</tr>
<tr>
<td>DMEA</td>
<td>0.58</td>
<td>0.56</td>
</tr>
<tr>
<td>MDEA</td>
<td>0.71</td>
<td>0.53</td>
</tr>
<tr>
<td>Methylmorpholine</td>
<td>0.43</td>
<td>0.59</td>
</tr>
<tr>
<td>Methylpiperazine</td>
<td>0.56</td>
<td>0.72</td>
</tr>
<tr>
<td>Morpholine</td>
<td>0.82</td>
<td>0.76</td>
</tr>
<tr>
<td>Blank</td>
<td>1.61</td>
<td>1.31</td>
</tr>
</tbody>
</table>

Figure 72: Corrosion rate by iron concentration from the A-HCT test
Figure 73: Condensation rate from the A-HCT test

Figure 74: Bulk pH from the A-HCT test
3.3.3.3.1. **Conclusions Altered Horizontal Cooled Tube**

The A-HCT test must be performed with utmost care to produce reliable results. Nevertheless, it seems to be a great test for research purposes under high pressure due to unique set of data it produces.

All of the tested high performing VCIs have a positive impact on the CR in the A-HCT test.

3.3.3.4. **Rotating Cylinder Electrode**

The last experiments to be discussed are the RCE tests of the high performing VCIs. In Figure 76 and Figure 77 the CR is displayed calculated by means of LPR and EIS, respectively. The CRs at the beginning of the test (approximately 5 hours) are the blank CRs of the respective tests; around 1.00 mm/y for LPR and 2.5 mm/y for EIS measurements. While tests were running, the VCI was injected into the system to alter the CR.

The characteristic of the CR evolution was the same for every compound; the CR dropped drastically to a lower value and remained there until the end of
the test. The CR over time curves calculated from the LPR tests are shown in Figure 76 and for EIS they are displayed in Figure 77. Most of the resulting CRs at the end of the tests are within a narrow window. The CR calculated by LPR for the tests with Cyclohexylamine, DEA, DMEA, MDEA, methylmorpholine, and morpholine all drop abruptly to a lower CR as soon the VCI was injected and stabilizes between 0.40 mm/y and 0.75 mm/y. A very similar picture can be seen in the evolution in the CR by calculated by EIS where the resulting CRs dropped down between 0.59 mm/y and 0.86 mm/y. The IEs of the mentioned generic VCIs are in the range of 34 % - 58 % and 66 % - 75 % by LPR and EIS, respectively (Table 20).

The potentiodynamic scans are displayed in Figure 78. None of the VCIs significantly affects the potential or the anodic side of the curve in any way. The maximum difference in corrosion potential from the blank is less than 25 mV. All of the VCIs slightly lower the cathodic current.

To investigate the mechanism of inhibition of the VCIs, a second test was run where the effect of the VCI on the pH was eliminated by adding an appropriate amount of HCl to adjust the pH back to pH 3.15. The CR graphs are shown in Figure 79 and Figure 80 and the IEs are listed in the second column of Table 20. The CRs calculated by LPR indicate a large drop in inhibition as the HCl is added. In Figure 79 it can be seen that the curves are spread out more than without HCl and the IEs are much lower between 0% - 32 %. The same tests investigated by EIS indicate a better inhibition ability of the compounds. The resulting CRs are altogether higher than without HCl added into the system, but with IEs between 34 % - 66% much more efficient than calculated by LPR. In this case, the EIS results are more trustworthy. As described under Rotating Cylinder Electrode (RCE) Test in section 3.2.5, LPR cannot distinguish between solution resistance (Rs) and charge transfer resistance (R_CT). In the experiments involving addition of VCI and HCl the charge transfer resistance increases due to inhibition but the effect is diminished in the LPR measurements by a smaller solution resistance due to the addition of VCI and more importantly HCl and NaOH to adjust the pH. The amines have a higher film forming ability as seen with the LPR tests.
The two remaining VCI compounds are benzylamine and methylpiperazine. Benzylamine displays very good inhibition as calculated by both LPR and EIS with a final CR of 0.12 mm/y and 0.13 mm/y and an IE of 88 % and 97 %, respectively. Methylpiperazine actually increases the CR significantly in all tests without HCl in the cell and therefore it was not possible to calculate an IE for this compound. Almost surprisingly, methylpiperazine does have a positive effect on the CR in the tests with HCl included. It shows a calculated IE of 44 % and 72 % by LPR and EIS, respectively. The compound decreases the CR significantly straight after it was injected into the system, but the CR increases over time as calculated by both, LPR and EIS.

Benzylamine on the other hand still displays a very good IE with HCl in the system. The IE calculated by EIS is 79 % and therefore this is the most effective of the high performing VCIs in the presence of HCl.

In Figure 81, the potentiodynamic scans are displayed of the tests with HCl. None of the curves are affected in a major fashion by any of the VCIs including HCl. The corrosion potential is shifted not more than 40 mV, this to a more cathodic value, though. Nevertheless, neither the cathodic nor the anodic corrosion current is significantly affected and stays very close to the blank curve.

<table>
<thead>
<tr>
<th>[%]</th>
<th>IE by LPR (Figure 76)</th>
<th>IE by EIS (Figure 77)</th>
<th>IE by LPR + HCl (Figure 79)</th>
<th>IE by EIS + HCl (Figure 80)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzylamine</td>
<td>88</td>
<td>97</td>
<td>59</td>
<td>79</td>
</tr>
<tr>
<td>Cyclohexylamine</td>
<td>42</td>
<td>70</td>
<td>0</td>
<td>34</td>
</tr>
<tr>
<td>DEA</td>
<td>40</td>
<td>66</td>
<td>10</td>
<td>39</td>
</tr>
<tr>
<td>DMEA</td>
<td>34</td>
<td>67</td>
<td>19</td>
<td>55</td>
</tr>
<tr>
<td>MDEA</td>
<td>51</td>
<td>75</td>
<td>33</td>
<td>66</td>
</tr>
<tr>
<td>Methylmorpholine</td>
<td>47</td>
<td>72</td>
<td>23</td>
<td>61</td>
</tr>
<tr>
<td>Methylpiperazine</td>
<td>-</td>
<td>-</td>
<td>44</td>
<td>72</td>
</tr>
<tr>
<td>Morpholine</td>
<td>58</td>
<td>-</td>
<td>32</td>
<td>65</td>
</tr>
</tbody>
</table>
Figure 76: Corrosion rate by LPR from the RCE test (without pH adjustment)

Figure 77: Corrosion rate by EIS from the RCE test (without pH adjustment)
Figure 78: Potentiodynamic scans from the RCE test (without pH adjustment)

Figure 79: Corrosion rate by LPR from the RCE test (pH adjusted +HCl)
Figure 80: Corrosion rate by EIS from the RCE test (pH adjusted +HCl)

Figure 81: Potentiodynamic scans from the RCE test (pH adjusted +HCl)
3.3.3.4.1. Conclusions Rotating Cylinder Electrode

The high performing VCIs show a very positive behaviour in the RCE test. All of the tested VCIs show an IE of over 66 % calculated by EIS. The addition of HCl distinguishes the different modes of inhibition of the different VCIs. IEs with HCl range from 34 % - 79 % calculated by EIS. In general, the low performing VCIs are better BOL inhibitors than the high performing VCIs.

EIS investigation is much better suited to the conditions tested in this investigation. 0.1 % NaCl solution is not conductive enough to gain trustworthy results by means of LPR. Especially in TOL corrosion investigations where condensed waters are replicated or directly measured, LPR tests seem to be misleading and inhibition properties of VCIs can be underestimated.

On the positive side, LPR overestimates the CR which is the more conservative outcome and therefore preferred by pipeline operators. Unfortunately, a VCI compound which actually works to protect against TOL corrosion, or even a fully formulated VCI, may be rejected as an option to inhibit the TOL because of a misleading measurement in the laboratory or in the field since many of the monitoring techniques utilise LPR to monitor the CR in the field.

3.3.3.5. Discussion and Conclusions “High Performing VCIs”

In this section, all the results for the individual VCIs will be discussed including the results of the individual test methods. The EIS results are used to draw conclusions for the RCE test.

3.3.3.5.1. Benzylamine

Benzylamine was an overall well performing compound despite its high boiling point and low vapour pressure. TOL corrosion was successfully inhibited in the HCT and CFP with a better performance in the LPR test than the WL or IC measurements. The performance in the A-HCT test was not very good.
The RCE tests with and without HCl were the best of the high performing VCIs with an IE of 97 % and 79 % with and without HCl as calculated by EIS, respectively. The very high IE with HCl in the system indicates that benzylamine is forming a stable film on the steel surface that separates the steel form the corrosive media around.

Due to the low vapour pressure, high boiling point and the results gained in the TOL corrosion test set-ups, it is concluded that the inhibition by benzylamine is mainly by neutralizing the bulk solution and not by forming a film on the TOL. Nevertheless, RCE tests at the BOL indicate that benzylamine is also form a film on the surface.

3.3.3.5.2. Cyclohexylamine

Cyclohexylamine was consistently an average performing VCI. The performance was average in the HCT and CFP but not good in the A-HCT. Due to these results it can be concluded that the boiling point and vapour pressure are still not favourable enough to make cyclohexylamine a very good TOL VCI compound. Most of its inhibition results from the neutralisation of the bulk solution and in the given conditions the volatility is too low so not enough of the VCI compound reaches the TOL.

Cyclohexylamine displayed an IE of 70 % in the RCE test. As soon HCl was added IE dropped to 34 %. This indicates a poor film forming capability of cyclohexylamine.

3.3.3.5.3. Diethylamine (DEA)

DEA was one of the best performing VCIs across all TOL corrosion test methods. It has a strong ability to neutralize the bulk solution. It displayed the best IE by WL in the HCT test. It also decreased the CR by means of IC in the CFP to the lowest value resulting in the highest IE by IC of 46 % (the next best IE by IC is 32 %). The performance in the A-HCT is also very good. Due to the high starting pH and therefore bulk neutralisation in the A-HCT test it can’t be significantly relied on, but the performance in the A-HCT goes along
with the overall performance of DEA. DEA has a low boiling point of 55 °C and a very high vapour pressure of 418 mmHg (at 20 °C). This seems to confirm the necessity of a low boiling point a high vapour pressure as characteristic for a successful VCI for TOL corrosion inhibition. The boiling point of DEA was in between the bulk solution temperature of the TOL tests (80 and 91 °C) and the test sample temperatures (below 55 °C). The boiling point was just right for the compound to evaporate from the bulk solution and condense at the TOL and inhibit corrosion. This could explain the rather good TOL inhibition properties despite the rather poor BOL inhibition properties tested in the RCE. DEA was just right for the conditions in the TOL corrosion tests.

The performance in the RCE test, with an IE of 66 % without and 39 % with the addition of HCl was lower than average. A large part of the inhibition seems to result from film forming.

The results indicate that DEA neutralizes the bulk solution and it seems to be volatile enough to reach the TOL. It then can form a film and neutralize the condensing liquid right at the TOL.

### 3.3.3.5.4. Dimethylethylamine (DMEA)

DMEA has excellent neutralization properties and therefore shows a good inhibition in the HCT test and also the A-HCT test. The performance in the CFP was lower than average.

The IE in the RCE test without HCl was with 67 % also lower than average. The IE with HCl was with 55 % also on the lower end of the VCIs but it also shows that a huge part of the inhibition seems to be a result of film forming.

Additionally to the neutralisation of the bulk solution, DMEA has a very low boiling point and high vapour pressure and is therefore volatile. Similarly to DEA, the BOL inhibition is low compared to the other tested VCIs but the TOL inhibition is rather good. It can again be concluded that the boiling point of DMEA is located just right for this test where due to the high temperature the VCI evaporates from the bulk solution and co-condenses at
the TOL due to the lower temperature. DMEA seems to neutralise the condensed liquid and forms a film on the steel surface.

3.3.3.5.5. **Methyldiethanolamine (MDEA)**

Within the “high performing VCIs”, MDEA has the weakest pH neutralisation ability which can be seen in the bulk pH of the HCT and CFP. The IE by means of IC was the lowest in the HCT test and also the lowest by means of LPR in the CFP. The IE by means of IC in the A-HCT is comparatively low but shows a certain inhibition effect. Nevertheless, the compound shows a good inhibition in the RCE test set up and maintains most of the IE in the tests including HCl with 75 % and 66 %, respectively.

Due to its high boiling point it is not very volatile, but MDEA seems to be protective directly at the TOL as seen in the A-HCT test. Much of the performance comes from its film forming abilities.

3.3.3.5.6. **Methylmorpholine**

Methylmorpholine seems to work well in the HCT and A-HCT with 46 % and 73 % IE by IC, respectively. Both values are the highest in the respective test set up. In the CFP test, the performance was average except for the WL data with an IE of 30 % which is just 1 % short of morpholine as the best performing VCI. The LPR performance was relatively good with an IE of 72 % without HCl and 61 % with HCl.

Methylmorpholine shows potential in the pressurised test set ups. It shows volatility but most of the performance comes from neutralisation of the bulk solution. In the RCE test it shows a pronounced film forming ability.

3.3.3.5.7. **Methylpiperazine**

Methylpiperazine is one of the better performing VCIs in all tests. In the CFP though, it reaches an IE of 43 % by means of LPR and only 8 % and 7 % of IE by means of IC and WL, respectively, both the lowest measured values. A
comparatively high condensation rate is responsible for the high CR rates by means of IC and WL. Taking all test results into account, methylpiperazine performed well as a VCI.

Methylpiperazine seems to increase the CR in the given conditions and concentration in the RCE test without HCL. When HCl was added into the test as a neutraliser, the IE was comparatively good at 72 %.

Due to the good inhibition of methylpiperazine, the increasing CR at the BOL inhibition should be investigated further. Experiments at a lower concentration should be valuable.

Methylpiperazine is a volatile compound with good film forming, neutralizing, and inhibiting properties.

3.3.3.5.8.  **Morpholine**

Morpholine is the smallest cyclic compound in the high performing section. It is an average performing compound with solid IEs throughout the tests. In the A-HCT test it was challenged with the highest condensation rates which probably resulted in a lower IE.

In the RCE test with HCl morpholine showed a residual IE of 65 % indicating a high possibility of film forming.

The volatility of morpholine is sufficient to reach the TOL and inhibit there directly by neutralisation and film formation. As with all other high performing VCIs, morpholine also neutralises the bulk solution.
3.3.4. Conclusion Generic VCI Compounds

Several generic VCI compounds were tested in a variety of test set ups. Unfortunately, the “silver bullet” against TOL corrosion was not within the tested compounds.

Nevertheless, some important data about the chemical compounds was gathered during the experiments. The conclusions of the separate compounds have been presented after the tests under “Low performing VCIs” and “High performing VCIs”.

In general, it seems that the boiling point is one very important property of a successful VCI compound. Most of the low performing VCI compounds have high boiling points. The lowest boiling point was 115 °C for pyridine, with all other low performing compounds having a boiling point of 145 °C and higher, up to 256 °C. In the high performing VCIs, the compounds with the highest boiling points were MDEA, benzylamine and then cyclohexylamine where two of the three performed poorly in the A-HCT test where volatility is a key property. DEA and DMEA, the two compounds with the lowest boiling point (and highest vapour pressure) were performing well in the TOL tests despite both were exhibiting a “lower than average” IE in the RCE tests. In other words, this means that the actual inhibition ability of the compounds is low compared to the other contestants but due to the “right” boiling point the compounds are able to evaporate from the bulk solution and re-condense at the TOL at a higher rate and therefore perform the inhibition to a more satisfactory grade. These results highlight the importance of choosing the right compound for the right circumstance.

The chemical structure is another important property. Four out of six low performing cyclic amines contain a benzene ring where just one of the high performing VCIs has a benzene ring, benzylamine. The benzene ring is not responsible for the high boiling point, but seems to be disadvantageous for TOL corrosion inhibition.
Overall, to be able to inhibit TOL corrosion directly, the boiling point must be low, the volatility high, and the molecular weight low. These necessary properties are usually not found in a film forming compound since these are traditionally long chained polar molecules. Therefore, all of the tested and more successful TOL inhibitors have stronger neutralisation ability than film forming ability. Nevertheless, most compounds also tend to form a film on the surface which does inhibit TOL corrosion to some extent.

A very positive finding was made on BOL corrosion inhibition with pyridazine. Despite the very small size of the molecule a very strong inhibition was revealed in the RCE test set up. More importantly, the inhibition effect is solely due to film forming effects since pyridazine had virtually no effect on the pH and in the RCE test that did include HCl most of the inhibition was still observed. Pyridazine reached an IE of 90 % in the RCE tests with 84 % left with HCl in the system. Compared to many other BOL inhibition compounds 90 % IE at a concentration of 1000 ppm is relatively poor performance. Nevertheless, further research might reveal that pyridazine might be a good candidate for niche applications like under deposit corrosion or it might be useful as an additional compound in a formulated BOL corrosion inhibitor.
4. Top-of-the-Line Corrosion Test Set-ups

4.1. Introduction

There are numerous TOL corrosion test methods in use. Many research groups and virtually every company which produce and sells VCIs have their own method of TOL corrosion testing and VCI testing (Pots and Hendriksen 2000; Gough et al. 2009; Singer et al. 2009; Gunaltun et al. 2010; Kashkovskiy and Kuzmnetsov 2012; Oehler et al. 2012). All of the methods produce different results and are difficult to compare.

Experience with the different set-ups was gained during the research. Advantages and disadvantages of the test methods will be assessed and suggestions for improvements will be made.

An improved set-up will be proposed to be used for future TOL corrosion and VCI research at Curtin University. If accepted by others, it could also cater as a standardized method for VCI research to be able to independently test and compare the efficiency of VCIs in different laboratories.
4.2. Discussion

The experiments discussed in the previous chapters are used and examined from the perspective of the test method rather than the angle of corrosion.

The HCT (introduced in section 2.2), A-HCT (introduced in section 0), CFP (introduced in section 3.2.3) as TOL corrosion test methods and the RCE test (introduced in section 3.2.5) as a BOL or condensed liquid test method are discussed in this section.

All of the TOL corrosion tests used in this research are static systems, which do not take any gas flow into consideration. Any dynamic system is more complicated and consists of more moving parts increasing the initial and running costs of the system. Nevertheless, the results show that the HCT set-up is generally suitable for performing TOL corrosion research. Indeed, all of the test methods are applicable to perform VCI research, each with its own advantages and disadvantages.

4.2.1. Horizontal Cooled Tube (HCT) Test

The HCT test was already largely improved from its original U-tube design, in which the sample was hanging in a U-shape and all the condensate collected at the lowest point as explained in section 2.3.1 and displayed in Figure 6 (John et al. 2009).

One problem is the test sample preparation by sandblasting. An effort was made to keep a constant ratio of fresh to used garnet in the sandblaster to maintain the same sample surface for each test. As the garnet is used over time the grains become smaller and rough edges are smoothened and therefore the surface roughness decreases. A smaller surface area is the result of the reduced surface roughness which decreases the condensation rate which again reduces corrosion. Hence, the consistency of the garnet can’t be guaranteed as it can be by using a new set of grinding paper unless fresh garnet is used at all times. Maintaining fresh garnet at all times would be possible, but was not achievable for the amount of testing during the
research (over 110 tests with a cooled tube set-up) and the very old and basic sandblaster used.

The major disadvantage for the test set-up would be the fact that the condensed liquid drops into the bulk solution, which elevates the pH and therefore decreases the availability of free acetic acid. As can be seen in the CR calculated by means of AAS for the blank sample, the CR drops significantly after two days, most likely due to the rising pH (Figure 82). The composition of condensed liquid and bulk cannot be studied separately for the same reasons.

The biggest advantage of the system clearly is its ability to perform high pressure and high temperature testing. Investigations can be performed at exactly the same conditions as found or expected in the field. By adjusting the temperature of the cooling fluid (therefore the sample temperature) and the autoclave temperature, it is possible to achieve a vast variety of condensation rates. In fact it even allows the adjustment of the condensation rate throughout the experiment. Another positive feature is that the pressure reactor and control unit are basically an “off the shelf” item with just minor modifications. The parts for the HCT could be ordered, built, and used all over the world with minimal effort or modifications. The test sample is also commercially available as a ¼ inch carbon steel tube, which could be exchanged for any available ¼ inch tube material.

In the case of VCI investigation, the high pressure pipette allows for pre-corrosion of the test sample before an inhibiting chemical is injected into the system. It would also be possible to replenish the bulk solution completely, keeping the system under pressure.
4.2.2. Altered Horizontal Cooled Tube (A-HCT) Test

The difference between A-HCT and HCT are the two added stainless steel cups; one to collect the condensed liquid and the second to hold the VCI and separate it from the bulk solution. The test was designed to be used with VCIs, especially to investigate the effect of VCIs directly to the TOL, eliminating neutralisation effects in the bulk solution.

The “weak point” of the set-up is actually the lower beaker that holds the VCI solution. The vessel is pressurized and de-pressured to remove oxygen out of the system. The pressurizing is performed with the sparge tube through the bulk solution of the autoclave. This procedure often splashes up the bulk solution which sometimes floods the content (VCI solution) of the lower beaker in to the bulk solution. The VCI then increases the pH of the bulk solution, which in turn makes the second cup redundant and even worse, this happens in an uncontrolled fashion.

Slight modifications on the equipment, like a conical cup and a gas disperser to avoid the extensive splashing and flooding could substantially increase the value of the test set-up.

Figure 82: CR calculated by IC and pH in the HCT test
Nevertheless, in the blank test, it can be seen that the condensed liquid doesn’t alter the pH of the bulk solution due to the collecting beaker. Therefore the CR stays high (Figure 83). Separating the condensed liquid and bulk solution has been proven to be important for a robust TOL corrosion test method.

![Figure 83: Corrosion rate by IC, condensation rate, condensed pH and bulk pH of the blank test from the A-HCT set-up](image)

### 4.2.3. Cooled Finger Probe (CFP) Test

The CFP test operates under ambient pressure and needs CO₂ sparging continuously to maintain an oxygen-free environment, increasing the running costs of a test. Nevertheless, the CFP test is very well suited for VCI testing and evaluation. The collection cup and LPR probe of the test make it possible to run in-situ electrochemical tests of the condensed liquid, adding another source of information beside WL and IC. The beaker also makes it possible to determine the exact condensation rate of the preceding 24 hour period, identifying irregularities in the test.
The CFP is limited in its application for TOL corrosion research due to the inability to operate at higher pressures, and therefore it is limited to bulk solution temperatures of less than 100 °C.

An improvement to the set up would be the exchange of LPR tests to EIS tests. As seen in the RCE test in the previous chapter, LPR is not well suited in condensed water conditions with low conductivity. Therefore, EIS would improve the significance of the results gained by electrochemical testing of the condensed liquid.

4.2.4. Rotating Cylinder Electrode (RCE) Test

RCE tests are a very useful research tool for most kinds of corrosion, including TOL corrosion. It is possible to replicate the condensed liquid and investigate its corrosivity in a simple set-up. It is also possible to investigate the performance of VCIs as demonstrated in the previous chapter.

In tests with a relatively high solution resistance or in instances where the conductivity of the solution might be changed by the addition of inhibitor or any other substance, EIS should be chosen to investigate the corrosion processes. In the VCI investigation discussed in the previous chapter, the solution resistance ($R_S$) decreased with the addition of VCI and HCl (Figure 84 from $R_{S1}$ to $R_{S2}$). Due to the inhibition effect of the VCI the polarisation resistance ($R_{CT}$) increased (Figure 84 from $R_{CT1}$ to $R_{CT2}$). The inhibition was not very strong, but existed and was recorded by EIS. LPR on the other hand did not record any changes in CR and the inhibitor appeared not to have any effect on the system whatsoever.

Therefore, for future TOL corrosion research, EIS should be used for both, RCE and CFP testing. It reinforces the confidence in the results and increases the amount of valuable information gathered.
Figure 8.4: Nyquist plot of EIS measurement with and without VCI
4.3. Conclusion  Improved TOL Corrosion Research Test Method

Combining all the experience gained by handling different TOL corrosion test methods, an improved set-up for TOL corrosion research and VCI evaluation can be proposed based on the HCT test.

The major problem with the HCT test was that the condensed liquid drips into the bulk solution, increasing the pH due to the dissolved corrosion products, and thus limiting further corrosion. Including a single stainless steel cup into the system that collects the condensed liquid over a 24 hour period and therefore separates the condensed liquid from the bulk solution solves the problem and additionally allows the condensation rate to be monitored throughout the test (Figure 85). This makes it possible to eliminate the neutralizing effect of the condensed liquid on the acetic acid in the bulk solution.

With the improved set-up, TOL corrosion research and VCI research can be carried out in a more complete way. The VCIs can be injected into the bulk solution after a pre-corrosion period using the high pressure pipette, reproducing the conditions of a real pipeline where corrosion may have been occurring for a substantial period of time.

Once the condensed liquid is extracted out of the vessel it can be used for further tests such as electrochemical measurements to investigate the corrosivity of the liquid or spectrophotometry measurements of corrosion products etc.

The improved HCT test combines all advantages such as the high pressure capability, high temperature capability, condensation rate monitoring (including the ability to vary the condensation rate), versatility of the corrosion sample, and maintaining constant bulk pH into one system.

To research the volatility of the VCIs without the neutralising effect of the bulk solution, a second, rimmed beaker and gas dispenser can be added to
the system to separate VCI from bulk solution. This would be an improved version of the A-HCT test, making the test even more versatile.

Figure 85: Schematic for the Improved Horizontal Cooled Tube (I-HCT) TOL corrosion and VCI test method
5. Conclusions and Future Work

Different aspects of TOL corrosion, TOL corrosion inhibition, and TOL corrosion testing have been investigated. The major findings and conclusions will be summarized in this chapter, including a proposal for further future work.

5.1. Conclusions

- High CO2 partial pressure (20 bar) has a positive influence at high acetic acid concentrations (1000 ppm).
- Research for the domain diagram revealed the severity of the influential factors on TOL corrosion from greatest to least:
  
  Temperature > Acetic acid Concentration > CO2 partial pressure

The VCI research stressed the importance of the condensation rate. It was shown to be the most influential factor on TOL corrosion.

- A low boiling point and a high vapour pressure are two key characteristics for a possible VCI candidate. The results received by DEA and DMEA indicate that it would be very advantageous to choose a boiling point between the high bulk solution temperature and the cooler sample/wall temperature. This allows the VCI to evaporate from the bulk solution and co-condense at the sample/wall in sufficient quantities to inhibit TOL corrosion.

- This research showed that a benzene ring is disadvantageous for a VCI candidate since it results in a lower vapour pressure and a higher boiling point.

- Pyridazine is a very good film forming BOL inhibitor. The molecular structure is very small compared to conventional, long chained BOL inhibitors. Pyridazine should be tested on its performance in specific applications like under deposit corrosion.
• EIS is recommended as the preferred technique to investigate simulated and experimentally condensed water applications because it is possible to distinguish between solution resistance ($R_S$) and charge transfer resistance ($R_{CT}$). LPR on the other hand, only measures polarisation resistance, which is $R_s + R_{CT}$ with $R_S$ being comparatively high in the condensed water applications due to low concentration of dissolved salts.

• EIS should be the preferential technique to be used when the solution resistance ($R_S$) is altered during the test by the addition of high concentrations of (volatile) corrosion inhibitors or when pH adjustment is performed. EIS is also recommended when solutions with different $R_S$ values are compared with each other.

• An improved TOL corrosion research method was proposed combining the advantages of the different methods.
  o High pressure, high temperature capability
  o Condensation rate is adjustable due to high variability of cooling and gas temperatures
  o Beaker separates the bulk solution from the condensed liquid
  o Condensed liquid will be sampled every 24 hours and can be used for further tests such as electrochemical tests or spectrophotometry measurements etc.
  o High pressure pipette allows for pre-corrosion before (volatile) corrosion inhibitor is injected into the system
  o Commercially available corrosion sample; any ¼ inch tube material can be used
  o “Off the shelf” set-up with minor modifications necessary
5.2. Future Work

TOL corrosion has a high potential to be “engineered away” in newly developed fields. In new fields with TOL corrosion potential, a corrosion resistant alloy will most likely be chosen for the first kilometres of the pipeline, which would be installed without heat insulation in order to cool down the gas and condense out any water vapour, eliminating any potential for TOL corrosion to occur. Nevertheless, already existing natural gas fields will stay in production for many years to come and therefore TOL corrosion, its mitigation, and prediction need to be well understood.

- TOL corrosion research should continue under different conditions using the proposed I-HCT TOL corrosion test system utilizing the condensed liquid collecting beaker, especially in high pCO\textsubscript{2} environments. With a focus on surface investigation by means of SEM or similar techniques, it could be possible to learn more about the mechanism of corrosion product layer formation and the protectivity under various conditions.

- TOL corrosion in the presence of hydrocarbons and/or monoethylene glycol (MEG) can be performed in the I-HCT system using separation beakers. The co-condensation of any substance at the TOL is an important subject and can be achieved by the I-HCT test.

- TOL corrosion and its mitigation in the presence of H\textsubscript{2}S should be investigated using the Hastelloy version of the I-HCT set-up available at Curtin University. TOL corrosion in the presence of H\textsubscript{2}S is driven by entirely different mechanisms but can be investigated using the same equipment.

- VCI investigation should continue with a focus on low boiling point compounds. They have been proven to be the most efficient VCIs in this research.

- Atomic force microscopy (AFM) is a powerful and commonly used tool for investigating film forming effects of conventional inhibitors. The technique was used in a trial to investigate the film forming ability of a
VCI compound at a concentration of 5000 ppm on a perfectly flat Mica surface \([\text{K}_2\text{Al}_4(\text{Al}_2\text{Si}_6)\text{O}_{20}(\text{OH})_4]\). It was possible to observe a film formation in-situ (Figure 86). The dark brown surface in the images is the background with a VCI layer growing a film on the surface over 3 hours (brighter parts). This research reinforces the findings of the RCE testing, that even very small molecules are able to form a film on a surface; in this case, on Mica.

A variety of possible VCI compounds can be investigated on their film forming abilities on different surfaces in different environments using the AFM.

Time restriction prevented further research during this project. Future research could involve polished steel samples and adhesion force measurements.
Figure 86: AFM images of a generic VCI compound forming a film on a Mica surface over time
6. Bibliography

Every reasonable effort has been made to acknowledge the owners of copyright material. I would be pleased to hear from any copyright owner who has been omitted or incorrectly acknowledged.


Petersen, P. R., S. A. Lordo and G. R. McAteer (2004). "Choosing a neutralising amine corrosion inhibitor." PTQ.


